OPERATING SYSTEMS
THREE EASY PIECES

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To Vedat S. Arpaci, a lifelong inspiration
Preface

To Everyone

Welcome to this book! We hope you’ll enjoy reading it as much as we enjoyed writing it. The book is called Operating Systems: Three Easy Pieces, and the title is obviously an homage to one of the greatest sets of lecture notes ever created, by one Richard Feynman on the topic of Physics [F96]. While this book will undoubtedly fall short of the high standard set by that famous physicist, perhaps it will be good enough for you in your quest to understand what operating systems (and more generally, systems) are all about.

The three easy pieces refer to the three major thematic elements the book is organized around: virtualization, concurrency, and persistence. In discussing these concepts, we’ll end up discussing most of the important things an operating system does; hopefully, you’ll also have some fun along the way. Learning new things is fun, right? At least, it should be.

Each major concept is divided into a set of chapters, most of which present a particular problem and then show how to solve it. The chapters are short, and try (as best as possible) to reference the source material where the ideas really came from. One of our goals in writing this book is to make the paths of history as clear as possible, as we think that helps a student understand what is, what was, and what will be more clearly. In this case, seeing how the sausage was made is nearly as important as understanding what the sausage is good for.

There are a couple devices we use throughout the book which are probably worth introducing here. The first is the crux of the problem. Anytime we are trying to solve a problem, we first try to state what the most important issue is; such a crux of the problem is explicitly called out in the text, and hopefully solved via the techniques, algorithms, and ideas presented in the rest of the text.

There are also numerous asides and tips throughout the text, adding a little color to the mainline presentation. Asides tend to discuss something relevant (but perhaps not essential) to the main text; tips tend to be general lessons that can be applied to systems you build. An index at the end of the book lists all of these tips and asides (as well as cruces, the odd plural of crux) for your convenience.

We use one of the oldest didactic methods, the dialogue, throughout the book, as a way of presenting some of the material in a different light. These are used to introduce the major thematic concepts (in a peachy way, as we will see), as well as to review material every now and then. They are also a chance to write in a more

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1Hint: eating! Or if you’re a vegetarian, running away from.
humorous style. Whether you find them useful, or humorous, well, that’s another matter entirely.

At the beginning of each major section, we’ll first present an abstraction that an operating system provides, and then work in subsequent chapters on the mechanisms, policies, and other support needed to provide the abstraction. Abstractions are fundamental to all aspects of Computer Science, so it is perhaps no surprise that they are also essential in operating systems.

Throughout the chapters, we try to use real code (not pseudocode) where possible, so for virtually all examples, you should be able to type them up yourself and run them. Running real code on real systems is the best way to learn about operating systems, so we encourage you to do so when you can.

In various parts of the text, we have sprinkled in a few homeworks to ensure that you are understanding what is going on. Many of these homeworks are little simulations of pieces of the operating system; you should download the homeworks, and run them to quiz yourself. The homework simulators have the following feature: by giving them a different random seed, you can generate a virtually infinite set of problems; the simulators can also be told to solve the problems for you. Thus, you can test and re-test yourself until you have achieved a good level of understanding.

The most important addendum to this book is a set of projects in which you learn about how real systems work by designing, implementing, and testing your own code. All projects (as well as the code examples, mentioned above) are in the C programming language [KR88]; C is a simple and powerful language that underlies most operating systems, and thus worth adding to your tool-chest of languages. Two types of projects are available (see the online appendix for ideas). The first are systems programming projects; these projects are great for those who are new to C and UNIX and want to learn how to do low-level C programming. The second type are based on a real operating system kernel developed at MIT called xv6 [CK+08]; these projects are great for students that already have some C and want to get their hands dirty inside the OS. At Wisconsin, we’ve run the course in three different ways: either all systems programming, all xv6 programming, or a mix of both.
To Educators

If you are an instructor or professor who wishes to use this book, please feel free to do so. As you may have noticed, they are free and available on-line from the following web page:

http://www.ostep.org

You can also purchase a printed copy from lulu.com. Look for it on the web page above.

The (current) proper citation for the book is as follows:

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Remzi H. Arpaci-Dusseau and Andrea C. Arpaci-Dusseau
Arpaci-Dusseau Books, Inc.
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http://www.ostep.org

The course divides fairly well across a 15-week semester, in which you can cover most of the topics within at a reasonable level of depth. Cramming the course into a 10-week quarter probably requires dropping some detail from each of the pieces. There are also a few chapters on virtual machine monitors, which we usually squeeze in sometime during the semester, either right at end of the large section on virtualization, or near the end as an aside.

One slightly unusual aspect of the book is that concurrency, a topic at the front of many OS books, is pushed off herein until the student has built an understanding of virtualization of the CPU and of memory. In our experience in teaching this course for nearly 15 years, students have a hard time understanding how the concurrency problem arises, or why they are trying to solve it, if they don’t yet understand what an address space is, what a process is, or why context switches can occur at arbitrary points in time. Once they do understand these concepts, however, introducing the notion of threads and the problems that arise due to them becomes rather easy, or at least, easier.

You may have noticed there are no slides that go hand-in-hand with the book. The major reason for this omission is that we believe in the most old-fashioned of teaching methods: chalk and a blackboard. Thus, when we teach the course, we come to class with a few major ideas and examples in mind and use the board to present them; handouts and live code demos sprinkled are also useful. In our experience, using too many slides encourages students to “check out” of lecture (and log into facebook.com), as they know the material is there for them to digest later; using the blackboard makes lecture a “live” viewing experience and thus (hopefully) more interactive, dynamic, and enjoyable for the students in your class.

If you’d like a copy of the notes we use in preparation for class, please drop us an email. We have already shared them with many others around the world.

One last request: if you use the free online chapters, please just link to them, instead of making a local copy. This helps us track usage (over 1 million chapters downloaded in the past few years!) and also ensures students get the latest and greatest version.
To Students

If you are a student reading this book, thank you! It is an honor for us to provide some material to help you in your pursuit of knowledge about operating systems. We both think back fondly towards some textbooks of our undergraduate days (e.g., Hennessy and Patterson [HP90], the classic book on computer architecture) and hope this book will become one of those positive memories for you.

You may have noticed this book is free and available online. There is one major reason for this: textbooks are generally too expensive. This book, we hope, is the first of a new wave of free materials to help those in pursuit of their education, regardless of which part of the world they come from or how much they are willing to spend for a book. Failing that, it is one free book, which is better than none.

We also hope, where possible, to point you to the original sources of much of the material in the book: the great papers and persons who have shaped the field of operating systems over the years. Ideas are not pulled out of the air; they come from smart and hard-working people (including numerous Turing-award winners2), and thus we should strive to celebrate those ideas and people where possible. In doing so, we hopefully can better understand the revolutions that have taken place, instead of writing texts as if those thoughts have always been present [K62]. Further, perhaps such references will encourage you to dig deeper on your own; reading the famous papers of our field is certainly one of the best ways to learn.

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2The Turing Award is the highest award in Computer Science; it is like the Nobel Prize, except that you have never heard of it.
Acknowledgments

This section will contain thanks to those who helped us put the book together. The important thing for now: your name could go here! But, you have to help. So send us some feedback and help debug this book. And you could be famous! Or, at least, have your name in some book.

The people who have helped so far include: Abhirami Senthilkumaran*, Adam Drescher, Adam Eggum, Ahmed Fikri*, Ajaykrishna Raghavan, Alex Wyler, Anand Mundada, B. Brahmananda Reddy (Minnesota), Bala Subrahmanyan Kambala, Benita Bose, Biswajit Mazumder (Clemson), Bobby Jack, Björn Lindberg, Brennan Payne, Brian Kroth, Cara Lauritzen, Charlotte Kissinger, Chien-Chung Shen (Delaware)*, Cody Hanson, Dan Soendergaard (U. Aarhus), David Hanle (Grinnell), Deepika Muthukumaran, Dorian Arnold (New Mexico), Dustin Metzler, Dustin Passofaro, Emily Jacobson, Emmett Witchel (Texas), Ernst Biersack (France), Finn Kuusisto*, Guilherme Baptista, Hamid Reza Ghasemi, Henry Abbey, Hrishikesh Amur, Huanchen Zhang*, Jake Gillberg, James Perry (U. Michigan-Dearborn)*, Jay Lim, Jerod Weinman (Grinnell), Joel Sommers (Colgate), Jonathan Perry (MIT), Jun He, Karl Wallinger, Kaushik Kannan, Kevin Liu*, Lei Tian (U. Nebraska-Lincoln), Leslie Schultz, Lihao Wang, Martha Ferris, Masashi Kishikawa (Sony), Matt Rechoff, Matty Williams, Meng Huang, Mike Griepentrog, Ming Chen (Stonybrook), Mohammed Alali (Delaware), Murugan Kandaswamy, Natasha Elber, Nathan Dipiazza, Nathan Sullivan, Neeraj Badlani (N.C. State), Nelson Gomez, Nghia Huynh (Texas), Patricio Jara, Radford Smith, Ripudaman Singh, Ross Aiken, Ruslan Kiselev, Ryland Herrick, Samer Al-Kiswany, Sandeep Ummadigari, Shreepa Chennuri, Shreepad Naik, Shreevatsa R., Sivaraman Sivaraman*, Srinivasan Thirunaranayanan*, Suriyaprabha Balaram Sankari, Sy Jin Cheah, Thomas Griebel, Tongxin Zheng, Tony Adkins, Torin Rudeen (Princeton), Tuow Wang, Varun Vats, Xiang Peng, Xu Di, Yue Zhuo (Texas A&M), Yufui Ren, Zef RosnBrick, Zuyu Zhang. Special thanks to those marked with an asterisk above, who have gone above and beyond in their suggestions for improvement.

Special thanks to Professor Joe Meehean (Lynchburg) for his detailed notes on each chapter, to Professor Jerod Weinman (Grinnell) and his entire class for their incredible booklets, and to Professor Chien-Chung Shen (Delaware) for his invaluable and detailed reading and comments about the book. All three have helped these authors immeasurably in the refinement of the materials herein.

Also, many thanks to the hundreds of students who have taken 537 over the years. In particular, the Fall ’08 class who encouraged the first written form of these notes (they were sick of not having any kind of textbook to read – pushy students!), and then praised them enough for us to keep going (including one hilarious “ZOMG! You should totally write a textbook!” comment in our course evaluations that year).

A great debt of thanks is also owed to the brave few who took the xv6 project lab course, much of which is now incorporated into the main 537 course. From Spring ’09: Justin Cherniak, Patrick Deline, Matt Czech, Tony Gregerson, Michael Griepentrog, Tyler Harter, Ryan Kroiss, Eric Radzikowski, Wesley Reardon, Rajiv Vaidyanathan, and Christopher Waclawik. From Fall ’09: Nick Bearson, Aaron Brown, Alex Bird, David Capel, Keith Gould, Tom Grim, Jeffrey Hugo, Brandon Johnson, John Kjell, Boyan Li, James Loethen, Will McCordell, Ryan Szaroletta, Si-
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Although they do not directly help with the book, our graduate students have taught us much of what we know about systems. We talk with them regularly while they are at Wisconsin, but they do all the real work – and by telling us about what they are doing, we learn new things every week. This list includes the following collection of current and former students with whom we published papers; an asterisk marks those who received a Ph.D. under our guidance: Abhishek Rajimwale, Ao Ma, Brian Forney, Chris Dragga, Deepak Ramamurthi, Florentina Popovici*, Haryadi S. Gunawi*, James Nugent, John Bent*, Lanyue Lu, Lakshmi Bairavasundaram*, Laxman Visampalli, Leo Arulraj, Meenali Rungta, Muthian Sivathanu*, Nathan Burnett*, Nitin Agrawal*, Sriram Subramanian*, Stephen Todd Jones*, Swaminathan Sundararaman*, Swetha Krishnan, Thanh Do, Thanumalayan S. Pillai, Timothy Denehy*, Tyler Harter, Venkat Venkataramani, Vijay Chidambaram, Vijayan Prabhakaran*, Yiyeng Zhang*, Yupu Zhang*, Zev Weiss.

A final debt of gratitude is also owed to Aaron Brown, who first took this course many years ago (Spring ’09), then took the xv6 lab course (Fall ’09), and finally was a graduate teaching assistant for the course for two years or so (Fall ’10 through Spring ’12). His tireless work has vastly improved the state of the projects (particularly those in xv6 land) and thus has helped better the learning experience for countless undergraduates and graduates here at Wisconsin. As Aaron would say (in his usual succinct manner): “Thx.”
Final Words

Yeats famously said “Education is not the filling of a pail but the lighting of a fire.” He was right but wrong at the same time\(^3\). You do have to “fill the pail” a bit, and these notes are certainly here to help with that part of your education; after all, when you go to interview at Google, and they ask you a trick question about how to use semaphores, it might be good to actually know what a semaphore is, right?

But Yeats’s larger point is obviously on the mark: the real point of education is to get you interested in something, to learn something more about the subject matter on your own and not just what you have to digest to get a good grade in some class. As one of our fathers (Remzi’s dad, Vedat Arpaci) used to say, “Learn beyond the classroom”.

We created these notes to spark your interest in operating systems, to read more about the topic on your own, to talk to your professor about all the exciting research that is going on in the field, and even to get involved with that research. It is a great field(!), full of exciting and wonderful ideas that have shaped computing history in profound and important ways. And while we understand this fire won’t light for all of you, we hope it does for many, or even a few. Because once that fire is lit, well, that is when you truly become capable of doing something great. And thus the real point of the educational process: to go forth, to study many new and fascinating topics, to learn, to mature, and most importantly, to find something that lights a fire for you.

Andrea and Remzi
Married couple
Professors of Computer Science at the University of Wisconsin
Chief Lighters of Fires, hopefully\(^4\)

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\(^3\)If he actually said this; as with many famous quotes, the history of this gem is murky.

\(^4\)If this sounds like we are admitting some past history as arsonists, you are probably missing the point. Probably. If this sounds cheesy, well, that’s because it is, but you’ll just have to forgive us for that.
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[CK+08] “The xv6 Operating System”
Russ Cox, Frans Kaashoek, Robert Morris, Nickolai Zeldovich
xv6 was developed as a part of the original UNIX version 6 and represents a beautiful, clean, and simple way to understand a modern operating system.

Richard P. Feynman
Basic Books, 1996
This book reprints the six easiest chapters of Feynman’s Lectures on Physics, from 1963. If you like Physics, it is a fantastic read.

[HP90] “Computer Architecture a Quantitative Approach” (1st ed.)
David A. Patterson and John L. Hennessy
Morgan-Kaufman, 1990
A book that encouraged each of us at our undergraduate institutions to pursue graduate studies; we later both had the pleasure of working with Patterson, who greatly shaped the foundations of our research careers.

Brian Kernighan and Dennis Ritchie
Prentice-Hall, April 1988
The C programming reference that everyone should have, by the people who invented the language.

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A great and famous read about the fundamentals of the scientific process. Mop-up work, anomaly, crisis, and revolution. We are mostly destined to do mop-up work, alas.
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A Dialogue on the Book

Professor: Welcome to this book! It’s called *Operating Systems in Three Easy Pieces*, and I am here to teach you the things you need to know about operating systems. I am called “Professor”; who are you?

Student: Hi Professor! I am called “Student”, as you might have guessed. And I am here and ready to learn!

Professor: Sounds good. Any questions?

Student: Sure! Why is it called “Three Easy Pieces”?

Professor: That’s an easy one. Well, you see, there are these great lectures on Physics by Richard Feynman...

Student: Oh! The guy who wrote “Surely You’re Joking, Mr. Feynman”, right? Great book! Is this going to be hilarious like that book was?

Professor: Um... well, no. That book was great, and I’m glad you’ve read it. Hopefully this book is more like his notes on Physics. Some of the basics were summed up in a book called “Six Easy Pieces”. He was talking about Physics; we’re going to do Three Easy Pieces on the fine topic of Operating Systems. This is appropriate, as Operating Systems are about half as hard as Physics.

Student: Well, I liked physics, so that is probably good. What are those pieces?

Professor: They are the three key ideas we’re going to learn about: **virtualization**, **concurrency**, and **persistence**. In learning about these ideas, we’ll learn all about how an operating system works, including how it decides what program to run next on a CPU, how it handles memory overload in a virtual memory system, how virtual machine monitors work, how to manage information on disks, and even a little about how to build a distributed system that works when parts have failed. That sort of stuff.

Student: I have no idea what you’re talking about, really.

Professor: Good! That means you are in the right class.

Student: I have another question: what’s the best way to learn this stuff?

Professor: Excellent query! Well, each person needs to figure this out on their
own, of course, but here is what I would do: go to class, to hear the professor introduce the material. Then, say at the end of every week, read these notes, to help the ideas sink into your head a bit better. Of course, some time later (hint: before the exam!), read the notes again to firm up your knowledge. Of course, your professor will no doubt assign some homeworks and projects, so you should do those; in particular, doing projects where you write real code to solve real problems is the best way to put the ideas within these notes into action. As Confucius said...

Student: Oh, I know! 'I hear and I forget. I see and I remember. I do and I understand.' Or something like that.

Professor: (surprised) How did you know what I was going to say?!

Student: It seemed to follow. Also, I am a big fan of Confucius.

Professor: Well, I think we are going to get along just fine! Just fine indeed.

Student: Professor – just one more question, if I may. What are these dialogues for? I mean, isn’t this just supposed to be a book? Why not present the material directly?

Professor: Ah, good question, good question! Well, I think it is sometimes useful to pull yourself outside of a narrative and think a bit; these dialogues are those times. So you and I are going to work together to make sense of all of these pretty complex ideas. Are you up for it?

Student: So we have to think? Well, I’m up for that. I mean, what else do I have to do anyhow? It’s not like I have much of a life outside of this book.

Professor: Me neither, sadly. So let’s get to work!
If you are taking an undergraduate operating systems course, you should already have some idea of what a computer program does when it runs. If not, this book (and the corresponding course) is going to be difficult – so you should probably stop reading this book, or run to the nearest bookstore and quickly consume the necessary background material before continuing (both Patt/Patel [PP03] and particularly Bryant/O’Hallaron [BOH10] are pretty great books).

So what happens when a program runs?

Well, a running program does one very simple thing: it executes instructions. Many millions (and these days, even billions) of times every second, the processor fetches an instruction from memory, decodes it (i.e., figures out which instruction this is), and executes it (i.e., it does the thing that it is supposed to do, like add two numbers together, access memory, check a condition, jump to a function, and so forth). After it is done with this instruction, the processor moves on to the next instruction, and so on, and so on, until the program finally completes 1.

Thus, we have just described the basics of the Von Neumann model of computing 2. Sounds simple, right? But in this class, we will be learning that while a program runs, a lot of other wild things are going on with the primary goal of making the system easy to use.

There is a body of software, in fact, that is responsible for making it easy to run programs (even allowing you to seemingly run many at the same time), allowing programs to share memory, enabling programs to interact with devices, and other fun stuff like that. That body of software

---

1 Of course, modern processors do many bizarre and frightening things underneath the hood to make programs run faster, e.g., executing multiple instructions at once, and even issuing and completing them out of order! But that is not our concern here; we are just concerned with the simple model most programs assume: that instructions seemingly execute one at a time, in an orderly and sequential fashion.

2 Von Neumann was one of the early pioneers of computing systems. He also did pioneering work on game theory and atomic bombs, and played in the NBA for six years. OK, one of those things isn't true.
THE CRUX OF THE PROBLEM: HOW TO VIRTUALIZE RESOURCES

One central question we will answer in this book is quite simple: how does the operating system virtualize resources? This is the crux of our problem. Why the OS does this is not the main question, as the answer should be obvious: it makes the system easier to use. Thus, we focus on the how: what mechanisms and policies are implemented by the OS to attain virtualization? How does the OS do so efficiently? What hardware support is needed?

We will use the “crux of the problem”, in shaded boxes such as this one, as a way to call out specific problems we are trying to solve in building an operating system. Thus, within a note on a particular topic, you may find one or more cruces (yes, this is the proper plural) which highlight the problem. The details within the chapter, of course, present the solution, or at least the basic parameters of a solution.

is called the operating system (OS), as it is in charge of making sure the system operates correctly and efficiently in an easy-to-use manner.

The primary way the OS does this is through a general technique that we call virtualization. That is, the OS takes a physical resource (such as the processor, or memory, or a disk) and transforms it into a more general, powerful, and easy-to-use virtual form of itself. Thus, we sometimes refer to the operating system as a virtual machine.

Of course, in order to allow users to tell the OS what to do and thus make use of the features of the virtual machine (such as running a program, or allocating memory, or accessing a file), the OS also provides some interfaces (APIs) that you can call. A typical OS, in fact, exports a few hundred system calls that are available to applications. Because the OS provides these calls to run programs, access memory and devices, and other related actions, we also sometimes say that the OS provides a standard library to applications.

Finally, because virtualization allows many programs to run (thus sharing the CPU), and many programs to concurrently access their own instructions and data (thus sharing memory), and many programs to access devices (thus sharing disks and so forth), the OS is sometimes known as a resource manager. Each of the CPU, memory, and disk is a resource of the system; it is thus the operating system’s role to manage those resources, doing so efficiently or fairly or indeed with many other possible goals in mind. To understand the role of the OS a little bit better, let’s take a look at some examples.

---

Another early name for the OS was the supervisor or even the master control program. Apparently, the latter sounded a little overzealous (see the movie Tron for details) and thus, thankfully, “operating system” caught on instead.
Figure 2.1: Simple Example: Code That Loops and Prints

2.1 Virtualizing the CPU

Figure 2.1 depicts our first program. It doesn’t do much. In fact, all it does is call Spin(), a function that repeatedly checks the time and returns once it has run for a second. Then, it prints out the string that the user passed in on the command line, and repeats, forever.

Let’s say we save this file as cpu.c and decide to compile and run it on a system with a single processor (or CPU as we will sometimes call it). Here is what we will see:

```
prompt> gcc -o cpu cpu.c -Wall
prompt> ./cpu "A"
A
A
A
^C
prompt>
```

Not too interesting of a run – the system begins running the program, which repeatedly checks the time until a second has elapsed. Once a second has passed, the code prints the input string passed in by the user (in this example, the letter “A”), and continues. Note the program will run forever; only by pressing “Control-c” (which on UNIX-based systems will terminate the program running in the foreground) can we halt the program.

Now, let’s do the same thing, but this time, let’s run many different instances of this same program. Figure 2.2 shows the results of this slightly more complicated example.

```
#include <stdio.h>
#include <stdlib.h>
#include <sys/time.h>
#include <assert.h>
#include "common.h"

int main(int argc, char *argv[])
{
    if (argc != 2) {
        fprintf(stderr, "usage: cpu <string>\n");
        exit(1);
    }
    char *str = argv[1];
    while (1) {
        Spin(1);
        printf("%s\n", str);
    }
    return 0;
}
```
Well, now things are getting a little more interesting. Even though we have only one processor, somehow all four of these programs seem to be running at the same time! How does this magic happen?\(^4\)

It turns out that the operating system, with some help from the hardware, is in charge of this illusion, i.e., the illusion that the system has a very large number of virtual CPUs. Turning a single CPU (or small set of them) into a seemingly infinite number of CPUs and thus allowing many programs to seemingly run at once is what we call virtualizing the CPU, the focus of the first major part of this book.

Of course, to run programs, and stop them, and otherwise tell the OS which programs to run, there need to be some interfaces (APIs) that you can use to communicate your desires to the OS. We’ll talk about these APIs throughout this book; indeed, they are the major way in which most users interact with operating systems.

You might also notice that the ability to run multiple programs at once raises all sorts of new questions. For example, if two programs want to run at a particular time, which should run? This question is answered by a policy of the OS; policies are used in many different places within an OS to answer these types of questions, and thus we will study them as we learn about the basic mechanisms that operating systems implement (such as the ability to run multiple programs at once). Hence the role of the OS as a resource manager.

---

\(^4\)Note how we ran four processes at the same time, by using the \& symbol. Doing so runs a job in the background in the tcsh shell, which means that the user is able to immediately issue their next command, which in this case is another program to run. The semi-colon between commands allows us to run multiple programs at the same time in tcsh. If you’re using a different shell (e.g., bash), it works slightly differently; read documentation online for details.
#include <unistd.h>
#include <stdio.h>
#include <stdlib.h>
#include "common.h"

int main(int argc, char *argv[]) {
    int *p = malloc(sizeof(int)); // a1
    assert(p != NULL);
    printf("(%d) address of p: %08x\n", getpid(), (unsigned) p); // a2
    *p = 0; // a3
    while (1) {
        Spin(1);
        *p = *p + 1;
        printf("(%d) p: %d\n", getpid(), *p); // a4
    }
    return 0;
}

Figure 2.3: A Program that Accesses Memory

2.2 Virtualizing Memory

Now let’s consider memory. The model of physical memory presented by modern machines is very simple. Memory is just an array of bytes; to read memory, one must specify an address to be able to access the data stored there; to write (or update) memory, one must also specify the data to be written to the given address.

Memory is accessed all the time when a program is running. A program keeps all of its data structures in memory, and accesses them through various instructions, like loads and stores or other explicit instructions that access memory in doing their work. Don’t forget that each instruction of the program is in memory too; thus memory is accessed on each instruction fetch.

Let’s take a look at a program (in Figure 2.3) that allocates some memory by calling malloc(). The output of this program can be found here:

prompt> ./mem
(2134) memory address of p: 00200000
(2134) p: 1
(2134) p: 2
(2134) p: 3
(2134) p: 4
(2134) p: 5
^C

The program does a couple of things. First, it allocates some memory (line a1). Then, it prints out the address of the memory (a2), and then puts the number zero into the first slot of the newly allocated memory (a3). Finally, it loops, delaying for a second and incrementing the value stored at the address held in p. With every print statement, it also prints out what is called the process identifier (the PID) of the running program. This PID is unique per running process.

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prompt> ./mem &; ./mem &
[1] 24113
[2] 24114
(24113) memory address of p: 00200000
(24114) memory address of p: 00200000
(24113) p: 1
(24114) p: 1
(24113) p: 2
(24114) p: 2
(24113) p: 3
(24114) p: 3
(24113) p: 4
(24114) p: 4
...

Figure 2.4: Running The Memory Program Multiple Times

Again, this first result is not too interesting. The newly allocated memory is at address 00200000. As the program runs, it slowly updates the value and prints out the result.

Now, we again run multiple instances of this same program to see what happens (Figure 2.4). We see from the example that each running program has allocated memory at the same address (00200000), and yet each seems to be updating the value at 00200000 independently! It is as if each running program has its own private memory, instead of sharing the same physical memory with other running programs. Indeed, that is exactly what is happening here as the OS is virtualizing memory. Each process accesses its own private virtual address space (sometimes just called its address space), which the OS somehow maps onto the physical memory of the machine. A memory reference within one running program does not affect the address space of other processes (or the OS itself); as far as the running program is concerned, it has physical memory all to itself. The reality, however, is that physical memory is a shared resource, managed by the operating system. Exactly how all of this is accomplished is also the subject of the first part of this book, on the topic of virtualization.

2.3 Concurrency

Another main theme of this book is concurrency. We use this conceptual term to refer to a host of problems that arise, and must be addressed, when working on many things at once (i.e., concurrently) in the same program. The problems of concurrency arose first within the operating system itself; as you can see in the examples above on virtualization, the OS is juggling many things at once, first running one process, then another, and so forth. As it turns out, doing so leads to some deep and interesting problems.

---

5For this example to work, you need to make sure address-space randomization is disabled; randomization, as it turns out, can be a good defense against certain kinds of security flaws. Read more about it on your own, especially if you want to learn how to break into computer systems via stack-smashing attacks. Not that we would recommend such a thing...
#include <stdio.h>
#include <stdlib.h>
#include "common.h"

volatile int counter = 0;
int loops;

void *worker(void *arg) {
    int i;
    for (i = 0; i < loops; i++) {
        counter++;
    }
    return NULL;
}

int main(int argc, char *argv[]) {
    if (argc != 2) {
        fprintf(stderr, "usage: threads <value>\n");
        exit(1);
    }
    loops = atoi(argv[1]);
    pthread_t p1, p2;
    printf("Initial value : %d\n", counter);
    Pthread_create(&p1, NULL, worker, NULL);
    Pthread_create(&p2, NULL, worker, NULL);
    Pthread_join(p1, NULL);
    Pthread_join(p2, NULL);
    printf("Final value : %d\n", counter);
    return 0;
}

Figure 2.5: A Multi-threaded Program

Unfortunately, the problems of concurrency are no longer limited just to the OS itself. Indeed, modern multi-threaded programs exhibit the same problems. Let us demonstrate with an example of a multi-threaded program (Figure 2.5).

Although you might not understand this example fully at the moment (and we’ll learn a lot more about it in later chapters, in the section of the book on concurrency), the basic idea is simple. The main program creates two threads using Pthread_create(). You can think of a thread as a function running within the same memory space as other functions, with more than one of them active at a time. In this example, each thread starts running in a routine called worker(), in which it simply increments a counter in a loop for loops number of times.

Below is a transcript of what happens when we run this program with the input value for the variable loops set to 1000. The value of loops

The actual call should be to lower-case pthread_create(); the upper-case version is our own wrapper that calls pthread_create() and makes sure that the return code indicates that the call succeeded. See the code for details.
THE CRUX OF THE PROBLEM:
HOW TO BUILD CORRECT CONCURRENT PROGRAMS

When there are many concurrently executing threads within the same memory space, how can we build a correctly working program? What primitives are needed from the OS? What mechanisms should be provided by the hardware? How can we use them to solve the problems of concurrency?

determines how many times each of the two workers will increment the shared counter in a loop. When the program is run with the value of loops set to 1000, what do you expect the final value of counter to be?

prompt> gcc -o thread thread.c -Wall -pthread
prompt> ./thread 1000
Initial value : 0
Final value : 2000

As you probably guessed, when the two threads are finished, the final value of the counter is 2000, as each thread incremented the counter 1000 times. Indeed, when the input value of loops is set to $N$, we would expect the final output of the program to be $2N$. But life is not so simple, as it turns out. Let’s run the same program, but with higher values for loops, and see what happens:

prompt> ./thread 100000
Initial value : 0
Final value : 143012 // huh??
prompt> ./thread 100000
Initial value : 0
Final value : 137298 // what the??

In this run, when we gave an input value of 100,000, instead of getting a final value of 200,000, we instead first get 143,012. Then, when we run the program a second time, we not only again get the wrong value, but also a different value than the last time. In fact, if you run the program over and over with high values of loops, you may find that sometimes you even get the right answer! So why is this happening?

As it turns out, the reason for these odd and unusual outcomes relate to how instructions are executed, which is one at a time. Unfortunately, a key part of the program above, where the shared counter is incremented, takes three instructions: one to load the value of the counter from memory into a register, one to increment it, and one to store it back into memory. Because these three instructions do not execute atomically (all at once), strange things can happen. It is this problem of concurrency that we will address in great detail in the second part of this book.
#include <stdio.h>
#include <unistd.h>
#include <assert.h>
#include <fcntl.h>
#include <sys/types.h>

int main(int argc, char *argv[])
{
    int fd = open("/tmp/file", O_WRONLY | O_CREAT | O_TRUNC, S_IRWXU);
    assert(fd > -1);
    int rc = write(fd, "hello world\n", 13);
    assert(rc == 13);
    close(fd);
    return 0;
}

Figure 2.6: A Program That Does I/O

2.4 Persistence

The third major theme of the course is persistence. In system memory, data can be easily lost, as devices such as DRAM store values in a volatile manner; when power goes away or the system crashes, any data in memory is lost. Thus, we need hardware and software to be able to store data persistently; such storage is thus critical to any system as users care a great deal about their data.

The hardware comes in the form of some kind of input/output or I/O device; in modern systems, a hard drive is a common repository for long-lived information, although solid-state drives (SSDs) are making headway in this arena as well.

The software in the operating system that usually manages the disk is called the file system; it is thus responsible for storing any files the user creates in a reliable and efficient manner on the disks of the system.

Unlike the abstractions provided by the OS for the CPU and memory, the OS does not create a private, virtualized disk for each application. Rather, it is assumed that often times, users will want to share information that is in files. For example, when writing a C program, you might first use an editor (e.g., Emacs) to create and edit the C file (emacs -nw main.c). Once done, you might use the compiler to turn the source code into an executable (e.g., gcc -o main main.c). When you’re finished, you might run the new executable (e.g., ./main). Thus, you can see how files are shared across different processes. First, Emacs creates a file that serves as input to the compiler; the compiler uses that input file to create a new executable file (in many steps – take a compiler course for details); finally, the new executable is then run. And thus a new program is born!

To understand this better, let’s look at some code. Figure 2.6 presents code to create a file (/tmp/file) that contains the string “hello world”.

7You should be using Emacs. If you are using vi, there is probably something wrong with you. If you are using something that is not a real code editor, that is even worse.
The file system is the part of the OS in charge of managing persistent data. What techniques are needed to do so correctly? What mechanisms and policies are required to do so with high performance? How is reliability achieved, in the face of failures in hardware and software?

To accomplish this task, the program makes three calls into the operating system. The first, a call to `open()`, opens the file and creates it; the second, `write()`, writes some data to the file; the third, `close()`, simply closes the file thus indicating the program won’t be writing any more data to it. These system calls are routed to the part of the operating system called the file system, which then handles the requests and returns some kind of error code to the user.

You might be wondering what the OS does in order to actually write to disk. We would show you but you’d have to promise to close your eyes first; it is that unpleasant. The file system has to do a fair bit of work: first figuring out where on disk this new data will reside, and then keeping track of it in various structures the file system maintains. Doing so requires issuing I/O requests to the underlying storage device, to either read existing structures or update (write) them. As anyone who has written a device driver\(^8\) knows, getting a device to do something on your behalf is an intricate and detailed process. It requires a deep knowledge of the low-level device interface and its exact semantics. Fortunately, the OS provides a standard and simple way to access devices through its system calls. Thus, the OS is sometimes seen as a standard library.

Of course, there are many more details in how devices are accessed, and how file systems manage data persistently atop said devices. For performance reasons, most file systems first delay such writes for a while, hoping to batch them into larger groups. To handle the problems of system crashes during writes, most file systems incorporate some kind of intricate write protocol, such as journaling or copy-on-write, carefully ordering writes to disk to ensure that if a failure occurs during the write sequence, the system can recover to reasonable state afterwards. To make different common operations efficient, file systems employ many different data structures and access methods, from simple lists to complex b-trees. If all of this doesn’t make sense yet, good! We’ll be talking about all of this quite a bit more in the third part of this book on persistence, where we’ll discuss devices and I/O in general, and then disks, RAIDs, and file systems in great detail.

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\(^8\)A device driver is some code in the operating system that knows how to deal with a specific device. We will talk more about devices and device drivers later.
2.5 Design Goals

So now you have some idea of what an OS actually does: it takes physical resources, such as a CPU, memory, or disk, and virtualizes them. It handles tough and tricky issues related to concurrency. And it stores files persistently, thus making them safe over the long-term. Given that we want to build such a system, we want to have some goals in mind to help focus our design and implementation and make trade-offs as necessary; finding the right set of trade-offs is a key to building systems.

One of the most basic goals is to build up some abstractions in order to make the system convenient and easy to use. Abstractions are fundamental to everything we do in computer science. Abstraction makes it possible to write a large program by dividing it into small and understandable pieces, to write such a program in a high-level language like C\(^9\) without thinking about assembly, to write code in assembly without thinking about logic gates, and to build a processor out of gates without thinking too much about transistors. Abstraction is so fundamental that sometimes we forget its importance, but we won’t here; thus, in each section, we’ll discuss some of the major abstractions that have developed over time, giving you a way to think about pieces of the OS.

One goal in designing and implementing an operating system is to provide high performance; another way to say this is our goal is to minimize the overheads of the OS. Virtualization and making the system easy to use are well worth it, but not at any cost; thus, we must strive to provide virtualization and other OS features without excessive overheads. These overheads arise in a number of forms: extra time (more instructions) and extra space (in memory or on disk). We’ll seek solutions that minimize one or the other or both, if possible. Perfection, however, is not always attainable, something we will learn to notice and (where appropriate) tolerate.

Another goal will be to provide protection between applications, as well as between the OS and applications. Because we wish to allow many programs to run at the same time, we want to make sure that the malicious or accidental bad behavior of one does not harm others; we certainly don’t want an application to be able to harm the OS itself (as that would affect all programs running on the system). Protection is at the heart of one of the main principles underlying an operating system, which is that of isolation; isolating processes from one another is the key to protection and thus underlies much of what an OS must do.

The operating system must also run non-stop; when it fails, all applications running on the system fail as well. Because of this dependence, operating systems often strive to provide a high degree of reliability. As operating systems grow evermore complex (sometimes containing millions of lines of code), building a reliable operating system is quite a chal-

\(^{9}\)Some of you might object to calling C a high-level language. Remember this is an OS course, though, where we’re simply happy not to have to code in assembly all the time!
lenge – and indeed, much of the on-going research in the field (including some of our own work [BS+09, SS+10]) focuses on this exact problem.

Other goals make sense: energy-efficiency is important in our increasingly green world; security (an extension of protection, really) against malicious applications is critical, especially in these highly-networked times; mobility is increasingly important as OSes are run on smaller and smaller devices. Depending in how the system is used, the OS will have different goals and thus likely be implemented in at least slightly different ways. However, as we will see, many of the principles we will present on how to build operating systems are useful in the range of different devices.

2.6 Some History

Before closing this introduction, let us present a brief history of how operating systems developed. Like any system built by humans, good ideas accumulated in operating systems over time, as engineers learned what was important in their design. Here, we discuss a few major developments. For a richer treatment, see Brinch Hansen’s excellent history of operating systems [BH00].

Early Operating Systems: Just Libraries

In the beginning, the operating system didn’t do too much. Basically, it was just a set of libraries of commonly-used functions; for example, instead of having each programmer of the system write low-level I/O handling code, the “OS” would provide such APIs, and thus make life easier for the developer.

Usually, on these old mainframe systems, one program ran at a time, as controlled by a human operator. Much of what you think a modern OS would do (e.g., deciding what order to run jobs in) was performed by this operator. If you were a smart developer, you would be nice to this operator, so that they might move your job to the front of the queue.

This mode of computing was known as batch processing, as a number of jobs were set up and then run in a “batch” by the operator. Computers, as of that point, were not used in an interactive manner, because of cost: it was simply too costly to let a user sit in front of the computer and use it, as most of the time it would just sit idle then, costing the facility hundreds of thousands of dollars per hour [BH00].

Beyond Libraries: Protection

In moving beyond being a simple library of commonly-used services, operating systems took on a more central role in managing machines. One important aspect of this was the realization that code run on behalf of the OS was special; it had control of devices and thus should be treated dif-
ferently than normal application code. Why is this? Well, imagine if you allowed any application to read from anywhere on the disk; the notion of privacy goes out the window, as any program could read any file. Thus, implementing a file system (to manage your files) as a library makes little sense. Instead, something else was needed.

Thus, the idea of a system call was invented, pioneered by the Atlas computing system [K+61,L78]. Instead of providing OS routines as a library (where you just make a procedure call to access them), the idea here was to add a special pair of hardware instructions and hardware state to make the transition into the OS a more formal, controlled process.

The key difference between a system call and a procedure call is that a system call transfers control (i.e., jumps) into the OS while simultaneously raising the hardware privilege level. User applications run in what is referred to as user mode which means the hardware restricts what applications can do; for example, an application running in user mode can’t typically initiate an I/O request to the disk, access any physical memory page, or send a packet on the network. When a system call is initiated (usually through a special hardware instruction called a trap), the hardware transfers control to a pre-specified trap handler (that the OS set up previously) and simultaneously raises the privilege level to kernel mode. In kernel mode, the OS has full access to the hardware of the system and thus can do things like initiate an I/O request or make more memory available to a program. When the OS is done servicing the request, it passes control back to the user via a special return-from-trap instruction, which reverts to user mode while simultaneously passing control back to where the application left off.

The Era of Multiprogramming

Where operating systems really took off was in the era of computing beyond the mainframe, that of the minicomputer. Classic machines like the PDP family from Digital Equipment made computers hugely more affordable; thus, instead of having one mainframe per large organization, now a smaller collection of people within an organization could likely have their own computer. Not surprisingly, one of the major impacts of this drop in cost was an increase in developer activity; more smart people got their hands on computers and thus made computer systems do more interesting and beautiful things.

In particular, multiprogramming became commonplace due to the desire to make better use of machine resources. Instead of just running one job at a time, the OS would load a number of jobs into memory and switch rapidly between them, thus improving CPU utilization. This switching was particularly important because I/O devices were slow; having a program wait on the CPU while its I/O was being serviced was a waste of CPU time. Instead, why not switch to another job and run it for a while?

The desire to support multiprogramming and overlap in the presence of I/O and interrupts forced innovation in the conceptual development of
operating systems along a number of directions. Issues such as memory protection became important; we wouldn’t want one program to be able to access the memory of another program. Understanding how to deal with the concurrency issues introduced by multiprogramming was also critical; making sure the OS was behaving correctly despite the presence of interrupts is a great challenge. We will study these issues and related topics later in the book.

One of the major practical advances of the time was the introduction of the UNIX operating system, primarily thanks to Ken Thompson (and Dennis Ritchie) at Bell Labs (yes, the phone company). UNIX took many good ideas from different operating systems (particularly from Multics [O72], and some from systems like TENEX [B+72] and the Berkeley Time-Sharing System [S+68]), but made them simpler and easier to use. Soon this team was shipping tapes containing UNIX source code to people around the world, many of whom then got involved and added to the system themselves; see the Aside (next page) for more detail10.

The Modern Era

Beyond the minicomputer came a new type of machine, cheaper, faster, and for the masses: the personal computer, or PC as we call it today. Led by Apple’s early machines (e.g., the Apple II) and the IBM PC, this new breed of machine would soon become the dominant force in computing, as their low-cost enabled one machine per desktop instead of a shared minicomputer per workgroup.

Unfortunately, for operating systems, the PC at first represented a great leap backwards, as early systems forgot (or never knew of) the lessons learned in the era of minicomputers. For example, early operating systems such as DOS (the Disk Operating System, from Microsoft) didn’t think memory protection was important; thus, a malicious (or perhaps just a poorly-programmed) application could scribble all over memory. The first generations of the Mac OS (v9 and earlier) took a cooperative approach to job scheduling; thus, a thread that accidentally got stuck in an infinite loop could take over the entire system, forcing a reboot. The painful list of OS features missing in this generation of systems is long, too long for a full discussion here.

Fortunately, after some years of suffering, the old features of minicomputer operating systems started to find their way onto the desktop. For example, Mac OS X has UNIX at its core, including all of the features one would expect from such a mature system. Windows has similarly adopted many of the great ideas in computing history, starting in particular with Windows NT, a great leap forward in Microsoft OS technology. Even today’s cell phones run operating systems (such as Linux) that are

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10We’ll use asides and other related text boxes to call attention to various items that don’t quite fit the main flow of the text. Sometimes, we’ll even use them just to make a joke, because why not have a little fun along the way? Yes, many of the jokes are bad.
ASIDE: The Importance of Unix

It is difficult to overstate the importance of Unix in the history of operating systems. Influenced by earlier systems (in particular, the famous Multics system from MIT), Unix brought together many great ideas and made a system that was both simple and powerful.

Underlying the original “Bell Labs” Unix was the unifying principle of building small powerful programs that could be connected together to form larger workflows. The shell, where you type commands, provided primitives such as pipes to enable such meta-level programming, and thus it became easy to string together programs to accomplish a bigger task. For example, to find lines of a text file that have the word “foo” in them, and then to count how many such lines exist, you would type: `grep foo file.txt | wc -l`, thus using the `grep` and `wc` (word count) programs to achieve your task.

The Unix environment was friendly for programmers and developers alike, also providing a compiler for the new C programming language. Making it easy for programmers to write their own programs, as well as share them, made Unix enormously popular. And it probably helped a lot that the authors gave out copies for free to anyone who asked, an early form of open-source software.

Also of critical importance was the accessibility and readability of the code. Having a beautiful, small kernel written in C invited others to play with the kernel, adding new and cool features. For example, an enterprising group at Berkeley, led by Bill Joy, made a wonderful distribution (the Berkeley Systems Distribution, or BSD) which had some advanced virtual memory, file system, and networking subsystems. Joy later co-founded Sun Microsystems.

Unfortunately, the spread of Unix was slowed a bit as companies tried to assert ownership and profit from it, an unfortunate (but common) result of lawyers getting involved. Many companies had their own variants: SunOS from Sun Microsystems, AIX from IBM, HPUX (a.k.a. “H-Pucks”) from HP, and IRIX from SGI. The legal wrangling among AT&T/Bell Labs and these other players cast a dark cloud over Unix, and many wondered if it would survive, especially as Windows was introduced and took over much of the PC market...

much more like what a minicomputer ran in the 1970s than what a PC ran in the 1980s (thank goodness); it is good to see that the good ideas developed in the heyday of OS development have found their way into the modern world. Even better is that these ideas continue to develop, providing more features and making modern systems even better for users and applications.
Fortunately for UNIX, a young Finnish hacker named Linus Torvalds decided to write his own version of UNIX which borrowed heavily on the principles and ideas behind the original system, but not from the code base, thus avoiding issues of legality. He enlisted help from many others around the world, and soon Linux was born (as well as the modern open-source software movement).

As the internet era came into place, most companies (such as Google, Amazon, Facebook, and others) chose to run Linux, as it was free and could be readily modified to suit their needs; indeed, it is hard to imagine the success of these new companies had such a system not existed. As smart phones became a dominant user-facing platform, Linux found a stronghold there too (via Android), for many of the same reasons. And Steve Jobs took his UNIX-based NeXTStep operating environment with him to Apple, thus making UNIX popular on desktops (though many users of Apple technology are probably not even aware of this fact). And thus UNIX lives on, more important today than ever before. The computing gods, if you believe in them, should be thanked for this wonderful outcome.

2.7 Summary

Thus, we have an introduction to the OS. Today’s operating systems make systems relatively easy to use, and virtually all operating systems you use today have been influenced by the developments we will discuss throughout the book.

Unfortunately, due to time constraints, there are a number of parts of the OS we won’t cover in the book. For example, there is a lot of networking code in the operating system; we leave it to you to take the networking class to learn more about that. Similarly, graphics devices are particularly important; take the graphics course to expand your knowledge in that direction. Finally, some operating system books talk a great deal about security; we will do so in the sense that the OS must provide protection between running programs and give users the ability to protect their files, but we won’t delve into deeper security issues that one might find in a security course.

However, there are many important topics that we will cover, including the basics of virtualization of the CPU and memory, concurrency, and persistence via devices and file systems. Don’t worry! While there is a lot of ground to cover, most of it is quite cool, and at the end of the road, you’ll have a new appreciation for how computer systems really work. Now get to work!
References

[BS+09] “Tolerating File-System Mistakes with EnvyFS”
Lakshmi N. Bairavasundaram, Swaminathan Sundararaman, Andrea C. Arpaci-Dusseau, Remzi H. Arpaci-Dusseau
USENIX ’09, San Diego, CA, June 2009
A fun paper about using multiple file systems at once to tolerate a mistake in any one of them.

[BH00] “The Evolution of Operating Systems”
P. Brinch Hansen
In Classic Operating Systems: From Batch Processing to Distributed Systems
Springer-Verlag, New York, 2000
This essay provides an intro to a wonderful collection of papers about historically significant systems.

[B+72] “TENEX, A Paged Time Sharing System for the PDP-10”
Daniel G. Bobrow, Jerry D. Burchfiel, Daniel L. Murphy, Raymond S. Tomlinson
CACM, Volume 15, Number 3, March 1972
TENEX has much of the machinery found in modern operating systems; read more about it to see how much innovation was already in place in the early 1970’s.

[B75] “The Mythical Man-Month”
Fred Brooks
Addison-Wesley, 1975
A classic text on software engineering; well worth the read.

Randal E. Bryant and David R. O’Hallaron
Addison-Wesley, 2010
Another great intro to how computer systems work. Has a little bit of overlap with this book – so if you’d like, you can skip the last few chapters of that book, or simply read them to get a different perspective on some of the same material. After all, one good way to build up your own knowledge is to hear as many other perspectives as possible, and then develop your own opinion and thoughts on the matter. You know, by thinking!

[K+61] “One-Level Storage System”
IRE Transactions on Electronic Computers, April 1962
The Atlas pioneered much of what you see in modern systems. However, this paper is not the best read. If you were to only read one, you might try the historical perspective below [L78].

S. H. Lavington
Communications of the ACM archive
Volume 21, Issue 1 (January 1978), pages 4-12
A nice piece of history on the early development of computer systems and the pioneering efforts of the Atlas. Of course, one could go back and read the Atlas papers themselves, but this paper provides a great overview and adds some historical perspective.

Elliott Organick, 1972
A great overview of Multics. So many good ideas, and yet it was an over-designed system, shooting for too much, and thus never really worked as expected. A classic example of what Fred Brooks would call the “second-system effect” [B75].
[PP03] “Introduction to Computing Systems: From Bits and Gates to C and Beyond”
Yale N. Patt and Sanjay J. Patel
One of our favorite intro to computing systems books. Starts at transistors and gets you all the way up to C; the early material is particularly great.

[RT74] “The UNIX Time-Sharing System”
Dennis M. Ritchie and Ken Thompson
CACM, Volume 17, Number 7, July 1974, pages 365-375
A great summary of UNIX written as it was taking over the world of computing, by the people who wrote it.

[S68] “SDS 940 Time-Sharing System”
Scientific Data Systems Inc.
TECHNICAL MANUAL, SDS 90 11168 August 1968
Available: http://goo.gl/EN0Zrn
Yes, a technical manual was the best we could find. But it is fascinating to read these old system documents, and see how much was already in place in the late 1960’s. One of the minds behind the Berkeley Time-Sharing System (which eventually became the SDS system) was Butler Lampson, who later won a Turing award for his contributions in systems.

Swaminathan Sundararaman, Sriram Subramanian, Abhishek Rajimwale, Andrea C. Arpaci-Dusseau, Remzi H. Arpaci-Dusseau, Michael M. Swift
FAST ’10, San Jose, CA, February 2010
The great thing about writing your own class notes: you can advertise your own research. But this paper is actually pretty neat – when a file system hits a bug and crashes, Membrane auto-magically restarts it, all without applications or the rest of the system being affected.
Part I

Virtualization
A Dialogue on Virtualization

Professor: And thus we reach the first of our three pieces on operating systems: virtualization.

Student: But what is virtualization, oh noble professor?

Professor: Imagine we have a peach.

Student: A peach? (incredulous)

Professor: Yes, a peach. Let us call that the physical peach. But we have many eaters who would like to eat this peach. What we would like to present to each eater is their own peach, so that they can be happy. We call the peach we give eaters virtual peaches; we somehow create many of these virtual peaches out of the one physical peach. And the important thing: in this illusion, it looks to each eater like they have a physical peach, but in reality they don’t.

Student: So you are sharing the peach, but you don’t even know it?

Professor: Right! Exactly.

Student: But there’s only one peach.

Professor: Yes. And...?

Student: Well, if I was sharing a peach with somebody else, I think I would notice.

Professor: Ah yes! Good point. But that is the thing with many eaters; most of the time they are napping or doing something else, and thus, you can snatch that peach away and give it to someone else for a while. And thus we create the illusion of many virtual peaches, one peach for each person!

Student: Sounds like a bad campaign slogan. You are talking about computers, right Professor?

Professor: Ah, young grasshopper, you wish to have a more concrete example. Good idea! Let us take the most basic of resources, the CPU. Assume there is one physical CPU in a system (though now there are often two or four or more). What virtualization does is take that single CPU and make it look like many virtual CPUs to the applications running on the system. Thus, while each applications
thinks it has its own CPU to use, there is really only one. And thus the OS has created a beautiful illusion: it has virtualized the CPU.

**Student:** Wow! That sounds like magic. Tell me more! How does that work?

**Professor:** In time, young student, in good time. Sounds like you are ready to begin.

**Student:** I am! Well, sort of. I must admit, I’m a little worried you are going to start talking about peaches again.

**Professor:** Don’t worry too much; I don’t even like peaches. And thus we begin...
In this note, we discuss one of the most fundamental abstractions that the OS provides to users: the **process**. The definition of a process, informally, is quite simple: it is a **running program** [V+65,B70]. The program itself is a lifeless thing: it just sits there on the disk, a bunch of instructions (and maybe some static data), waiting to spring into action. It is the operating system that takes these bytes and gets them running, transforming the program into something useful.

It turns out that one often wants to run more than one program at once; for example, consider your desktop or laptop where you might like to run a web browser, mail program, a game, a music player, and so forth. In fact, a typical system may be seemingly running tens or even hundreds of processes at the same time. Doing so makes the system easy to use, as one never need be concerned with whether a CPU is available; one simply runs programs. Hence our challenge:

**THE CRUX OF THE PROBLEM:**
**HOW TO PROVIDE THE ILLUSION OF MANY CPUs?**
Although there are only a few physical CPUs available, how can the OS provide the illusion of a nearly-endless supply of said CPUs?

The OS creates this illusion by **virtualizing** the CPU. By running one process, then stopping it and running another, and so forth, the OS can promote the illusion that many virtual CPUs exist when in fact there is only one physical CPU (or a few). This basic technique, known as **time sharing** of the CPU, allows users to run as many concurrent processes as they would like; the potential cost is performance, as each will run more slowly if the CPU(s) must be shared.

To implement virtualization of the CPU, and to implement it well, the OS will need both some low-level machinery as well as some high-level intelligence. We call the low-level machinery **mechanisms**; mechanisms are low-level methods or protocols that implement a needed piece of
Tip: Use Time Sharing (and Space Sharing)

Time sharing is one of the most basic techniques used by an OS to share a resource. By allowing the resource to be used for a little while by one entity, and then a little while by another, and so forth, the resource in question (e.g., the CPU, or a network link) can be shared by many. The natural counterpart of time sharing is space sharing, where a resource is divided (in space) among those who wish to use it. For example, disk space is naturally a space-shared resource, as once a block is assigned to a file, it is not likely to be assigned to another file until the user deletes it.

functionality. For example, we’ll learn below how to implement a context switch, which gives the OS the ability to stop running one program and start running another on a given CPU; this time-sharing mechanism is employed by all modern OSes.

On top of these mechanisms resides some of the intelligence in the OS, in the form of policies. Policies are algorithms for making some kind of decision within the OS. For example, given a number of possible programs to run on a CPU, which program should the OS run? A scheduling policy in the OS will make this decision, likely using historical information (e.g., which program has run more over the last minute?), workload knowledge (e.g., what types of programs are run), and performance metrics (e.g., is the system optimizing for interactive performance, or throughput?) to make its decision.

4.1 The Abstraction: A Process

The abstraction provided by the OS of a running program is something we will call a process. As we said above, a process is simply a running program; at any instant in time, we can summarize a process by taking an inventory of the different pieces of the system it accesses or affects during the course of its execution.

To understand what constitutes a process, we thus have to understand its machine state: what a program can read or update when it is running. At any given time, what parts of the machine are important to the execution of this program?

One obvious component of machine state that comprises a process is its memory. Instructions lie in memory; the data that the running program reads and writes sits in memory as well. Thus the memory that the process can address (called its address space) is part of the process.

Also part of the process’s machine state are registers; many instructions explicitly read or update registers and thus clearly they are important to the execution of the process.

Note that there are some particularly special registers that form part of this machine state. For example, the program counter (PC) (sometimes called the instruction pointer or IP) tells us which instruction of the pro-
In many operating systems, a common design paradigm is to separate high-level policies from their low-level mechanisms [L+75]. You can think of the mechanism as providing the answer to a *how* question about a system; for example, *how* does an operating system perform a context switch? The policy provides the answer to a *which* question; for example, *which* process should the operating system run right now? Separating the two allows one easily to change policies without having to rethink the mechanism and is thus a form of *modularity*, a general software design principle.

For example, a program is currently being executed; similarly a **stack pointer** and associated **frame pointer** are used to manage the stack for function parameters, local variables, and return addresses.

Finally, programs often access persistent storage devices too. Such **I/O information** might include a list of the files the process currently has open.

### 4.2 Process API

Though we defer discussion of a real process API until a subsequent chapter, here we first give some idea of what must be included in any interface of an operating system. These APIs, in some form, are available on any modern operating system.

- **Create**: An operating system must include some method to create new processes. When you type a command into the shell, or double-click on an application icon, the OS is invoked to create a new process to run the program you have indicated.
- **Destroy**: As there is an interface for process creation, systems also provide an interface to destroy processes forcefully. Of course, many processes will run and just exit by themselves when complete; when they don’t, however, the user may wish to kill them, and thus an interface to halt a runaway process is quite useful.
- **Wait**: Sometimes it is useful to wait for a process to stop running; thus some kind of waiting interface is often provided.
- **Miscellaneous Control**: Other than killing or waiting for a process, there are sometimes other controls that are possible. For example, most operating systems provide some kind of method to suspend a process (stop it from running for a while) and then resume it (continue it running).
- **Status**: There are usually interfaces to get some status information about a process as well, such as how long it has run for, or what state it is in.
4.3 Process Creation: A Little More Detail

One mystery that we should unmask a bit is how programs are transformed into processes. Specifically, how does the OS get a program up and running? How does process creation actually work?

The first thing that the OS must do to run a program is to load its code and any static data (e.g., initialized variables) into memory, into the address space of the process. Programs initially reside on disk (or, in some modern systems, flash-based SSDs) in some kind of executable format; thus, the process of loading a program and static data into memory requires the OS to read those bytes from disk and place them in memory somewhere (as shown in Figure 4.1).

In early (or simple) operating systems, the loading process is done eagerly, i.e., all at once before running the program; modern OSes perform the process lazily, i.e., by loading pieces of code or data only as they are needed during program execution. To truly understand how lazy loading of pieces of code and data works, you’ll have to understand more about the machinery of paging and swapping, topics we’ll cover in the future when we discuss the virtualization of memory. For now, just remember that before running anything, the OS clearly must do some work to get the important program bits from disk into memory.
Once the code and static data are loaded into memory, there are a few other things the OS needs to do before running the process. Some memory must be allocated for the program’s run-time stack (or just stack). As you should likely already know, C programs use the stack for local variables, function parameters, and return addresses; the OS allocates this memory and gives it to the process. The OS will also likely initialize the stack with arguments; specifically, it will fill in the parameters to the main() function, i.e., argc and the argv array.

The OS may also create some initial memory for the program’s heap. In C programs, the heap is used for explicitly requested dynamically-allocated data; programs request such space by calling malloc() and free it explicitly by calling free(). The heap is needed for data structures such as linked lists, hash tables, trees, and other interesting data structures. The heap will be small at first; as the program runs, and requests more memory via the malloc() library API, the OS may get involved and allocate more memory to the process to help satisfy such calls.

The OS will also do some other initialization tasks, particularly as related to input/output (I/O). For example, in UNIX systems, each process by default has three open file descriptors, for standard input, output, and error; these descriptors let programs easily read input from the terminal as well as print output to the screen. We’ll learn more about I/O, file descriptors, and the like in the third part of the book on persistence.

By loading the code and static data into memory, by creating and initializing a stack, and by doing other work as related to I/O setup, the OS has now (finally) set the stage for program execution. It thus has one last task: to start the program running at the entry point, namely main(). By jumping to the main() routine (through a specialized mechanism that we will discuss next chapter), the OS transfers control of the CPU to the newly-created process, and thus the program begins its execution.

4.4 Process States

Now that we have some idea of what a process is (though we will continue to refine this notion), and (roughly) how it is created, let us talk about the different states a process can be in at a given time. The notion that a process can be in one of these states arose in early computer systems [V+65,DV66]. In a simplified view, a process can be in one of three states:

- **Running**: In the running state, a process is running on a processor. This means it is executing instructions.
- **Ready**: In the ready state, a process is ready to run but for some reason the OS has chosen not to run it at this given moment.
- **Blocked**: In the blocked state, a process has performed some kind of operation that makes it not ready to run until some other event takes place. A common example: when a process initiates an I/O request to a disk, it becomes blocked and thus some other process can use the processor.
If we were to map these states to a graph, we would arrive at the diagram in Figure 4.2. As you can see in the diagram, a process can be moved between the ready and running states at the discretion of the OS. Being moved from ready to running means the process has been scheduled; being moved from running to ready means the process has been descheduled. Once a process has become blocked (e.g., by initiating an I/O operation), the OS will keep it as such until some event occurs (e.g., I/O completion); at that point, the process moves to the ready state again (and potentially immediately to running again, if the OS so decides).

4.5 Data Structures

The OS is a program, and like any program, it has some key data structures that track various relevant pieces of information. To track the state of each process, for example, the OS likely will keep some kind of process list for all processes that are ready, as well as some additional information to track which process is currently running. The OS must also track, in some way, blocked processes; when an I/O event completes, the OS should make sure to wake the correct process and ready it to run again.

Figure 4.3 shows what type of information an OS needs to track about each process in the xv6 kernel [CK+08]. Similar process structures exist in “real” operating systems such as Linux, Mac OS X, or Windows; look them up and see how much more complex they are.

From the figure, you can see a couple of important pieces of information the OS tracks about a process. The register context will hold, for a stopped process, the contents of its register state. When a process is stopped, its register state will be saved to this memory location; by restoring these registers (i.e., placing their values back into the actual physical registers), the OS can resume running the process. We’ll learn more about this technique known as a context switch in future chapters.
// the registers xv6 will save and restore
// to stop and subsequently restart a process
struct context {
    int eip;
    int esp;
    int ebx;
    int ecx;
    int edx;
    int esi;
    int edi;
    int ebp;
};

// the different states a process can be in
enum proc_state { UNUSED, EMBRYO, SLEEPING,
                  Runnable, Running, Zombie };

// the information xv6 tracks about each process
// including its register context and state
struct proc {
    char *mem; // Start of process memory
    uint sz; // Size of process memory
    char *kstack; // Bottom of kernel stack
                  // for this process
    enum proc_state state; // Process state
    int pid; // Process ID
    struct proc *parent; // Parent process
    void *chan; // If non-zero, sleeping on chan
    int killed; // If non-zero, have been killed
    struct file *ofile[NOFILE]; // Open files
    struct inode *cwd; // Current directory
    struct context context; // Switch here to run process
    struct trapframe *tf; // Trap frame for the
                         // current interrupt
};

Figure 4.3: The xv6 Proc Structure

You can also see from the figure that there are some other states a process can be in, beyond running, ready, and blocked. Sometimes a system will have an initial state that the process is in when it is being created. Also, a process could be placed in a final state where it has exited but has not yet been cleaned up (in UNIX-based systems, this is called the zombie state1). This final state can be useful as it allows other processes (usually the parent that created the process) to examine the return code of the process and see if it the just-finished process executed successfully (usually, programs return zero in UNIX-based systems when they have accomplished a task successfully, and non-zero otherwise). When finished, the parent will make one final call (e.g., wait()) to wait for the completion of the child, and to also indicate to the OS that it can clean up any relevant data structures that referred to the now-extinct process.

1Yes, the zombie state. Just like real zombies, these zombies are relatively easy to kill. However, different techniques are usually recommended.
4.6 Summary

We have introduced the most basic abstraction of the OS: the process. It is quite simply viewed as a running program. With this conceptual view in mind, we will now move on to the nitty-gritty: the low-level mechanisms needed to implement processes, and the higher-level policies required to schedule them in an intelligent way. By combining mechanisms and policies, we will build up our understanding of how an operating system virtualizes the CPU.
References

[CK+08] “The xv6 Operating System” 
Russ Cox, Frans Kaashoek, Robert Morris, Nickolai Zeldovich 
The coolest real and little OS in the world. Download and play with it to learn more about the details of how operating systems actually work.

[DV66] “Programming Semantics for Multiprogrammed Computations” 
Jack B. Dennis and Earl C. Van Horn 
Communications of the ACM, Volume 9, Number 3, March 1966
This paper defined many of the early terms and concepts around building multiprogrammed systems.

[H70] “The Nucleus of a Multiprogramming System” 
Per Brinch Hansen 
Communications of the ACM, Volume 13, Number 4, April 1970
This paper introduces one of the first microkernels in operating systems history, called Nucleus. The idea of smaller, more minimal systems is a theme that rears its head repeatedly in OS history; it all began with Brinch Hansen’s work described herein.

[L+75] “Policy/mechanism separation in Hydra” 
R. Levin, E. Cohen, W. Corwin, F. Pollack, W. Wulf 
SOSP 1975
An early paper about how to structure operating systems in a research OS known as Hydra. While Hydra never became a mainstream OS, some of its ideas influenced OS designers.

[V+65] “Structure of the Multics Supervisor” 
V.A. Vyssotsky, F. J. Corbato, R. M. Graham 
Fall Joint Computer Conference, 1965
An early paper on Multics, which described many of the basic ideas and terms that we find in modern systems. Some of the vision behind computing as a utility are finally being realized in modern cloud systems.
Interlude: Process API

**Aside: Interludes**
Interludes will cover more practical aspects of systems, including a particular focus on operating system APIs and how to use them. If you don’t like practical things, you could skip these interludes. But you should like practical things, because, well, they are generally useful in real life; companies, for example, don’t usually hire you for your non-practical skills.

In this interlude, we discuss process creation in UNIX systems. UNIX presents one of the most intriguing ways to create a new process with a pair of system calls: `fork()` and `exec()`. A third routine, `wait()`, can be used by a process wishing to wait for a process it has created to complete. We now present these interfaces in more detail, with a few simple examples to motivate us. And thus, our problem:

**Crux: How to Create and Control Processes**
What interfaces should the OS present for process creation and control? How should these interfaces be designed to enable ease of use as well as utility?

5.1 The `fork()` System Call

The `fork()` system call is used to create a new process [C63]. However, be forewarned: it is certainly the strangest routine you will ever call\(^1\). More specifically, you have a running program whose code looks like what you see in Figure 5.1; examine the code, or better yet, type it in and run it yourself!

\(^1\)Well, OK, we admit that we don’t know that for sure; who knows what routines you call when no one is looking? But `fork()` is pretty odd, no matter how unusual your routine-calling patterns are.
When you run this program (called p1.c), you'll see the following:

prompt> ./p1
hello world (pid:29146)
hello, I am parent of 29147 (pid:29146)
hello, I am child (pid:29147)
prompt>

Let us understand what happened in more detail in p1.c. When it first started running, the process prints out a hello world message; included in that message is its process identifier, also known as a PID. The process has a PID of 29146; in UNIX systems, the PID is used to name the process if one wants to do something with the process, such as (for example) stop it from running. So far, so good.

Now the interesting part begins. The process calls the fork() system call, which the OS provides as a way to create a new process. The odd part: the process that is created is an (almost) exact copy of the calling process. That means that to the OS, it now looks like there are two copies of the program p1 running, and both are about to return from the fork() system call. The newly-created process (called the child, in contrast to the creating parent) doesn’t start running at main(), like you might expect (note, the “hello, world” message only got printed out once); rather, it just comes into life as if it had called fork() itself.

You might have noticed: the child isn’t an exact copy. Specifically, although it now has its own copy of the address space (i.e., its own private memory), its own registers, its own PC, and so forth, the value it returns to the caller of fork() is different. Specifically, while the parent receives the PID of the newly-created child, the child is simply returned a 0. This differentiation is useful, because it is simple then to write the code that handles the two different cases (as above).
You might also have noticed: the output is not deterministic. When the child process is created, there are now two active processes in the system that we care about: the parent and the child. Assuming we are running on a system with a single CPU (for simplicity), then either the child or the parent might run at that point. In our example (above), the parent did and thus printed out its message first. In other cases, the opposite might happen, as we show in this output trace:

```
prompt> ./p1
hello world (pid:29146)
hello, I am child (pid:29147)
hello, I am parent of 29147 (pid:29146)
prompt>
```

The CPU scheduler, a topic we’ll discuss in great detail soon, determines which process runs at a given moment in time; because the scheduler is complex, we cannot usually make strong assumptions about what it will choose to do, and hence which process will run first. This non-determinism, as it turns out, leads to some interesting problems, particularly in multi-threaded programs; hence, we’ll see a lot more non-determinism when we study concurrency in the second part of the book.

### 5.2 Adding `wait()` System Call

So far, we haven’t done much: just created a child that prints out a message and exits. Sometimes, as it turns out, it is quite useful for a parent to wait for a child process to finish what it has been doing. This task is accomplished with the `wait()` system call (or its more complete sibling `waitpid()`); see Figure 5.2 for details.

---

**Figure 5.2: p2.c: Calling `fork()` And `wait()`**

```c
#include <stdio.h>
#include <stdlib.h>
#include <unistd.h>
#include <sys/wait.h>

int main(int argc, char *argv[])
{
    printf("hello world (pid:%d)\n", (int) getpid());
    int rc = fork();
    if (rc < 0) { // fork failed; exit
        fprintf(stderr, "fork failed\n");
        exit(1);
    } else if (rc == 0) { // child (new process)
        printf("hello, I am child (pid:%d)\n", (int) getpid());
    } else { // parent goes down this path (main)
        int wc = wait(NULL);
        printf("hello, I am parent of %d (wc:%d) (pid:%d)\n", rc, wc, (int) getpid());
    }
    return 0;
}
```
In this example (p2.c), the parent process calls `wait()` to delay its execution until the child finishes executing. When the child is done, `wait()` returns to the parent.

Adding a `wait()` call to the code above makes the output deterministic. Can you see why? Go ahead, think about it.

*(waiting for you to think .... and done)*

Now that you have thought a bit, here is the output:

```
prompt> ./p2
hello world (pid:29266)
hello, I am child (pid:29267)
hello, I am parent of 29267 (wc:29267) (pid:29266)
prompt>
```

With this code, we now know that the child will always print first. Why do we know that? Well, it might simply run first, as before, and thus print before the parent. However, if the parent does happen to run first, it will immediately call `wait()`; this system call won’t return until the child has run and exited\(^2\). Thus, even when the parent runs first, it politely waits for the child to finish running, then `wait()` returns, and then the parent prints its message.

5.3 Finally, the `exec()` System Call

A final and important piece of the process creation API is the `exec()` system call\(^3\). This system call is useful when you want to run a program that is different from the calling program. For example, calling `fork()` in p2.c is only useful if you want to keep running copies of the same program. However, often you want to run a different program; `exec()` does just that (Figure 5.3).

In this example, the child process calls `execvp()` in order to run the program `wc`, which is the word counting program. In fact, it runs `wc` on the source file `p3.c`, thus telling us how many lines, words, and bytes are found in the file:

```
prompt> ./p3
hello world (pid:29383)
hello, I am child (pid:29384)
29 107 1030 p3.c
hello, I am parent of 29384 (wc:29384) (pid:29383)
prompt>
```

\(^2\)There are a few cases where `wait()` returns before the child exits; read the man page for more details, as always. And beware of any absolute and unqualified statements this book makes, such as “the child will always print first” or “UNIX is the best thing in the world, even better than ice cream.”

\(^3\)Actually, there are six variants of `exec()`: `execl()`, `execle()`, `execlp()`, `execv()`, and `execvp()`. Read the man pages to learn more.
#include <stdio.h>
#include <stdlib.h>
#include <unistd.h>
#include <string.h>
#include <sys/wait.h>

int main(int argc, char *argv[])
{
    printf("hello world (pid:%d)\n", (int) getpid());
    int rc = fork();
    if (rc < 0) { // fork failed; exit
        fprintf(stderr, "fork failed\n");
        exit(1);
    } else if (rc == 0) { // child (new process)
        printf("hello, I am child (pid:%d)\n", (int) getpid());
        char *myargs[3];
        myargs[0] = strdup("wc"); // program: "wc" (word count)
        myargs[1] = strdup("p3.c"); // argument: file to count
        myargs[2] = NULL; // marks end of array
        execvp(myargs[0], myargs); // runs word count
        printf("this shouldn’t print out\n");
    } else { // parent goes down this path (main)
        printf("hello, I am parent of %d (wc:%d) (pid:%d)\n", rc, rc, (int) getpid());
        int wc = wait(NULL);
        printf("hello, I am parent of %d (wc:%d) (pid:%d)\n", wc, rc, (int) getpid());
    }
    return 0;
}

Figure 5.3: p3.c: Calling fork(), wait(), And exec()

If fork() was strange, exec() is not so normal either. What it does: given the name of an executable (e.g., wc), and some arguments (e.g., p3.c), it loads code (and static data) from that executable and overwrites its current code segment (and current static data) with it; the heap and stack and other parts of the memory space of the program are re-initialized. Then the OS simply runs that program, passing in any arguments as the argv of that process. Thus, it does not create a new process; rather, it transforms the currently running program (formerly p3) into a different running program (wc). After the exec() in the child, it is almost as if p3.c never ran; a successful call to exec() never returns.

5.4 Why? Motivating the API

Of course, one big question you might have: why would we build such an odd interface to what should be the simple act of creating a new process? Well, as it turns out, the separation of fork() and exec() is essential in building a UNIX shell, because it lets the shell run code after the call to fork() but before the call to exec(); this code can alter the environment of the about-to-be-run program, and thus enables a variety of interesting features to be readily built.
TIP: GETTING IT RIGHT (LAMPSON’S LAW)

As Lampson states in his well-regarded “Hints for Computer Systems Design” [L83], “Get it right. Neither abstraction nor simplicity is a substitute for getting it right.” Sometimes, you just have to do the right thing, and when you do, it is way better than the alternatives. There are lots of ways to design APIs for process creation; however, the combination of `fork()` and `exec()` are simple and immensely powerful. Here, the UNIX designers simply got it right. And because Lampson so often “got it right”, we name the law in his honor.

The shell is just a user program. It shows you a prompt and then waits for you to type something into it. You then type a command (i.e., the name of an executable program, plus any arguments) into it; in most cases, the shell then figures out where in the file system the executable resides, calls `fork()` to create a new child process to run the command, calls some variant of `exec()` to run the command, and then waits for the command to complete by calling `wait()`. When the child completes, the shell returns from `wait()` and prints out a prompt again, ready for your next command.

The separation of `fork()` and `exec()` allows the shell to do a whole bunch of useful things rather easily. For example:

```
prompt> wc p3.c > newfile.txt
```

In the example above, the output of the program `wc` is redirected into the output file `newfile.txt` (the greater-than sign is how said redirection is indicated). The way the shell accomplishes this task is quite simple: when the child is created, before calling `exec()`, the shell closes standard output and opens the file `newfile.txt`. By doing so, any output from the soon-to-be-running program `wc` are sent to the file instead of the screen.

Figure 5.4 shows a program that does exactly this. The reason this redirection works is due to an assumption about how the operating system manages file descriptors. Specifically, UNIX systems start looking for free file descriptors at zero. In this case, STD_OUT_FILENO will be the first available one and thus get assigned when `open()` is called. Subsequent writes by the child process to the standard output file descriptor, for example by routines such as `printf()`, will then be routed transparently to the newly-opened file instead of the screen.

Here is the output of running the `p4.c` program:

```
prompt> ./p4
prompt> cat p4.output
 32   109   846 p4.c
prompt>
```

And there are lots of shells; `tcsh`, `bash`, and `zsh` to name a few. You should pick one, read its man pages, and learn more about it; all UNIX experts do.
#include <stdio.h>
#include <stdlib.h>
#include <unistd.h>
#include <string.h>
#include <fcntl.h>
#include <sys/wait.h>

int main(int argc, char *argv[]) {
    int rc = fork();
    if (rc < 0) { // fork failed; exit
        fprintf(stderr, "fork failed\n");
        exit(1);
    } else if (rc == 0) { // child: redirect standard output to a file
        close(STDOUT_FILENO);
        open("./p4.output", O_CREAT|O_WRONLY|O_TRUNC, S_IRWXU);
        // now exec "wc"...
        char *myargs[3];
        myargs[0] = strdup("wc"); // program: "wc" (word count)
        myargs[1] = strdup("p4.c"); // argument: file to count
        myargs[2] = NULL; // marks end of array
        execvp(myargs[0], myargs); // runs word count
    } else { // parent goes down this path (main)
        int wc = wait(NULL);
    }
    return 0;
}

Figure 5.4: p4.c: All Of The Above With Redirection

You’ll notice (at least) two interesting tidbits about this output. First, when p4 is run, it looks as if nothing has happened; the shell just prints the command prompt and is immediately ready for your next command. However, that is not the case; the program p4 did indeed call fork() to create a new child, and then run the wc program via a call to execvp(). You don’t see any output printed to the screen because it has been redirected to the file p4.output. Second, you can see that when we cat the output file, all the expected output from running wc is found. Cool, right?

UNIX pipes are implemented in a similar way, but with the pipe() system call. In this case, the output of one process is connected to an in-kernel pipe (i.e., queue), and the input of another process is connected to that same pipe; thus, the output of one process seamlessly is used as input to the next, and long and useful chains of commands can be strung together. As a simple example, consider the looking for a word in a file, and then counting how many times said word occurs; with pipes and the utilities grep and wc, it is easy – just type grep foo file | wc -l into the command prompt and marvel at the result.

Finally, while we just have sketched out the process API at a high level, there is a lot more detail about these calls out there to be learned and digested; we’ll learn more, for example, about file descriptors when we talk about file systems in the third part of the book. For now, suffice it to say that the fork()/exec() combination is a powerful way to create and manipulate processes.

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Many times in this book, when referring to a particular system call or library call, we’ll tell you to read the manual pages, or man pages for short. Man pages are the original form of documentation that exist on UNIX systems; realize that they were created before the thing called the web existed.

Spending some time reading man pages is a key step in the growth of a systems programmer; there are tons of useful tidbits hidden in those pages. Some particularly useful pages to read are the man pages for whichever shell you are using (e.g., tcsh, or bash), and certainly for any system calls your program makes (in order to see what return values and error conditions exist).

Finally, reading the man pages can save you some embarrassment. When you ask colleagues about some intricacy of fork(), they may simply reply: “RTFM.” This is your colleagues’ way of gently urging you to Read The Man pages. The F in RTFM just adds a little color to the phrase...

5.5 Other Parts of the API

Beyond fork(), exec(), and wait(), there are a lot of other interfaces for interacting with processes in UNIX systems. For example, the kill() system call is used to send signals to a process, including directives to go to sleep, die, and other useful imperatives. In fact, the entire signals subsystem provides a rich infrastructure to deliver external events to processes, including ways to receive and process those signals.

There are many command-line tools that are useful as well. For example, using the ps command allows you to see which processes are running; read the man pages for some useful flags to pass to ps. The tool top is also quite helpful, as it displays the processes of the system and how much CPU and other resources they are eating up. Humorously, many times when you run it, top claims it is the top resource hog; perhaps it is a bit of an egomaniac. Finally, there are many different kinds of CPU meters you can use to get a quick glance understanding of the load on your system; for example, we always keep MenuMeters (from Raging Menace software) running on our Macintosh toolbars, so we can see how much CPU is being utilized at any moment in time. In general, the more information about what is going on, the better.

5.6 Summary

We have introduced some of the APIs dealing with UNIX process creation: fork(), exec(), and wait(). However, we have just skimmed the surface. For more detail, read Stevens and Rago [SR05], of course, particularly the chapters on Process Control, Process Relationships, and Signals. There is much to extract from the wisdom therein.
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Lampson’s famous hints on how to design computer systems. You should read it at some point in your life, and probably at many points in your life.

[SR05] “Advanced Programming in the UNIX Environment”
W. Richard Stevens and Stephen A. Rago
Addison-Wesley, 2005
All nuances and subtleties of using UNIX APIs are found herein. Buy this book! Read it! And most importantly, live it.
Mechanism: Limited Direct Execution

In order to virtualize the CPU, the operating system needs to somehow share the physical CPU among many jobs running seemingly at the same time. The basic idea is simple: run one process for a little while, then run another one, and so forth. By time sharing the CPU in this manner, virtualization is achieved.

There are a few challenges, however, in building such virtualization machinery. The first is performance: how can we implement virtualization without adding excessive overhead to the system? The second is control: how can we run processes efficiently while retaining control over the CPU? Control is particularly important to the OS, as it is in charge of resources; without control, a process could simply run forever and take over the machine, or access information that it should not be allowed to access. Attaining performance while maintaining control is thus one of the central challenges in building an operating system.

The Crux:
How To Efficiently Virtualize The CPU With Control
The OS must virtualize the CPU in an efficient manner, but while retaining control over the system. To do so, both hardware and operating systems support will be required. The OS will often use a judicious bit of hardware support in order to accomplish its work effectively.

6.1 Basic Technique: Limited Direct Execution

To make a program run as fast as one might expect, not surprisingly OS developers came up with a technique, which we call limited direct execution. The “direct execution” part of the idea is simple: just run the program directly on the CPU. Thus, when the OS wishes to start a program running, it creates a process entry for it in a process list, allocates some memory pages for it, loads the program code into memory (from
**Table 6.1: Direction Execution Protocol (Without Limits)**

<table>
<thead>
<tr>
<th>OS</th>
<th>Program</th>
</tr>
</thead>
<tbody>
<tr>
<td>Create entry for process list</td>
<td>Run main()</td>
</tr>
<tr>
<td>Allocate memory for program</td>
<td>Execute return from main</td>
</tr>
<tr>
<td>Load program into memory</td>
<td></td>
</tr>
<tr>
<td>Set up stack with argc/argv</td>
<td></td>
</tr>
<tr>
<td>Clear registers</td>
<td></td>
</tr>
<tr>
<td>Execute call main()</td>
<td></td>
</tr>
<tr>
<td>Free memory of process</td>
<td></td>
</tr>
<tr>
<td>Remove from process list</td>
<td></td>
</tr>
</tbody>
</table>

disk), locates its entry point (i.e., the main() routine or something similar), jumps to it, and starts running the user’s code. Table 6.1 shows this basic direct execution protocol (without any limits, yet), using a normal call and return to jump to the program’s main() and later to get back into the kernel.

Sounds simple, no? But this approach gives rise to a few problems in our quest to virtualize the CPU. The first is simple: if we just run a program, how can the OS make sure the program doesn’t do anything that we don’t want it to do, while still running it efficiently? The second: when we are running a process, how does the operating system stop it from running and switch to another process, thus implementing the **time sharing** we require to virtualize the CPU?

In answering these questions below, we’ll get a much better sense of what is needed to virtualize the CPU. In developing these techniques, we’ll also see where the “limited” part of the name arises from; without limits on running programs, the OS wouldn’t be in control of anything and thus would be “just a library” – a very sad state of affairs for an aspiring operating system!

### 6.2 Problem #1: Restricted Operations

Direct execution has the obvious advantage of being fast; the program runs natively on the hardware CPU and thus executes as quickly as one would expect. But running on the CPU introduces a problem: what if the process wishes to perform some kind of restricted operation, such as issuing an I/O request to a disk, or gaining access to more system resources such as CPU or memory?

**THE CRUX: HOW TO PERFORM RESTRICTED OPERATIONS**

A process must be able to perform I/O and some other restricted operations, but without giving the process complete control over the system. How can the OS and hardware work together to do so?
One approach would simply be to let any process do whatever it wants in terms of I/O and other related operations. However, doing so would prevent the construction of many kinds of systems that are desirable. For example, if we wish to build a file system that checks permissions before granting access to a file, we can’t simply let any user process issue I/Os to the disk; if we did, a process could simply read or write the entire disk and thus all protections would be lost.

Thus, the approach we take is to introduce a new processor mode, known as \textit{user mode}; code that runs in user mode is restricted in what it can do. For example, when running in user mode, a process can’t issue I/O requests; doing so would result in the processor raising an exception; the OS would then likely kill the process.

In contrast to user mode is \textit{kernel mode}, which the operating system (or kernel) runs in. In this mode, code that runs can do what it likes, including privileged operations such as issuing I/O requests and executing all types of restricted instructions.

We are still left with a challenge, however: what should a user process do when it wishes to perform some kind of privileged operation, such as reading from disk? To enable this, virtually all modern hardware provides the ability for user programs to perform a \textit{system call}. Pioneered on ancient machines such as the Atlas [K+61,L78], system calls allow the kernel to carefully expose certain key pieces of functionality to user programs, such as accessing the file system, creating and destroying processes, communicating with other processes, and allocating more memory. Most operating systems provide a few hundred calls (see the POSIX standard for details [P10]); early Unix systems exposed a more concise subset of around twenty calls.

To execute a system call, a program must execute a special \textit{trap} instruction. This instruction simultaneously jumps into the kernel and raises the privilege level to kernel mode; once in the kernel, the system can now perform whatever privileged operations are needed (if allowed), and thus do the required work for the calling process. When finished, the OS calls a special \textit{return-from-trap} instruction, which, as you might expect, returns into the calling user program while simultaneously reducing the privilege level back to user mode.

The hardware needs to be a bit careful when executing a trap, in that it must make sure to save enough of the caller’s register state in order
Aside: Why System Calls Look Like Procedure Calls

You may wonder why a call to a system call, such as `open()` or `read()`, looks exactly like a typical procedure call in C; that is, if it looks just like a procedure call, how does the system know it’s a system call, and do all the right stuff? The simple reason: it is a procedure call, but hidden inside that procedure call is the famous trap instruction. More specifically, when you call `open()` (for example), you are executing a procedure call into the C library. Therein, whether for `open()` or any of the other system calls provided, the library uses an agreed-upon calling convention with the kernel to put the arguments to open in well-known locations (e.g., on the stack, or in specific registers), puts the system-call number into a well-known location as well (again, onto the stack or a register), and then executes the aforementioned trap instruction. The code in the library after the trap unpacks return values and returns control to the program that issued the system call. Thus, the parts of the C library that make system calls are hand-coded in assembly, as they need to carefully follow convention in order to process arguments and return values correctly, as well as execute the hardware-specific trap instruction. And now you know why you personally don’t have to write assembly code to trap into an OS; somebody has already written that assembly for you.

to be able to return correctly when the OS issues the return-from-trap instruction. On x86, for example, the processor will push the program counter, flags, and a few other registers onto a per-process kernel stack; the return-from-trap will pop these values off the stack and resume execution of the user-mode program (see the Intel systems manuals [I11] for details). Other hardware systems use different conventions, but the basic concepts are similar across platforms.

There is one important detail left out of this discussion: how does the trap know which code to run inside the OS? Clearly, the calling process can’t specify an address to jump to (as you would when making a procedure call); doing so would allow programs to jump anywhere into the kernel which clearly is a bad idea (imagine jumping into code to access a file, but just after a permission check; in fact, it is likely such ability would enable a wily programmer to get the kernel to run arbitrary code sequences [S07]). Thus the kernel must carefully control what code executes upon a trap.

The kernel does so by setting up a trap table at boot time. When the machine boots up, it does so in privileged (kernel) mode, and thus is free to configure machine hardware as need be. One of the first things the OS thus does is to tell the hardware what code to run when certain exceptional events occur. For example, what code should run when a hard-disk interrupt takes place, when a keyboard interrupt occurs, or when program makes a system call? The OS informs the hardware of the locations of these trap handlers, usually with some kind of special

Operating Systems

[Version 0.80] WWW.OSTEP.ORG
## Mechanism: Limited Direct Execution

<table>
<thead>
<tr>
<th>OS @ boot (kernel mode)</th>
<th>Hardware</th>
<th>Program (user mode)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>initialize trap table</strong></td>
<td>remember address of... syscall handler</td>
<td></td>
</tr>
</tbody>
</table>

### OS @ run (kernel mode)

<table>
<thead>
<tr>
<th>Hardware</th>
<th>Program (user mode)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Create entry for process list</td>
<td>Run main()</td>
</tr>
</tbody>
</table>
| Allocate memory for program | ...
| Load program into memory | Call system call trap into OS |
| Setup user stack with argv | save regs to kernel stack |
| Fill kernel stack with reg/PC | move to kernel mode |
| **return-from-trap** | jump to trap handler |

<table>
<thead>
<tr>
<th>Hardware</th>
<th>Program (user mode)</th>
</tr>
</thead>
</table>
| Handle trap | ...
| Do work of syscall | return from main trap (via `exit()`) |
| **return-from-trap** | |

Free memory of process

Remove from process list

| Table 6.2: Limited Direction Execution Protocol |

Instruction. Once the hardware is informed, it remembers the location of these handlers until the machine is next rebooted, and thus the hardware knows what to do (i.e., what code to jump to) when system calls and other exceptional events take place.

One last aside: being able to execute the instruction to tell the hardware where the trap tables are is a very powerful capability. Thus, as you might have guessed, it is also a privileged operation. If you try to execute this instruction in user mode, the hardware won’t let you, and you can probably guess what will happen (hint: adios, offending program). Point to ponder: what horrible things could you do to a system if you could install your own trap table? Could you take over the machine?

The timeline (with time increasing downward, in Table 6.2) summarizes the protocol. We assume each process has a kernel stack where registers (including general purpose registers and the program counter) are saved to and restored from (by the hardware) when transitioning into and out of the kernel.
There are two phases in the LDE protocol. In the first (at boot time), the kernel initializes the trap table, and the CPU remembers its location for subsequent use. The kernel does so via a privileged instruction (all privileged instructions are highlighted in bold).

In the second (when running a process), the kernel sets up a few things (e.g., allocating a node on the process list, allocating memory) before using a return-from-trap instruction to start the execution of the process; this switches the CPU to user mode and begins running the process. When the process wishes to issue a system call, it traps back into the OS, which handles it and once again returns control via a return-from-trap to the process. The process then completes its work, and returns from main(); this usually will return into some stub code which will properly exit the program (say, by calling the exit() system call, which traps into the OS). At this point, the OS cleans up and we are done.

6.3 Problem #2: Switching Between Processes

The next problem with direct execution is achieving a switch between processes. Switching between processes should be simple, right? The OS should just decide to stop one process and start another. What’s the big deal? But it actually is a little bit tricky: specifically, if a process is running on the CPU, this by definition means the OS is not running. If the OS is not running, how can it do anything at all? (hint: it can’t) While this sounds almost philosophical, it is a real problem: there is clearly no way for the OS to take an action if it is not running on the CPU. Thus we arrive at the crux of the problem.

**THE CRUX: HOW TO REGAIN CONTROL OF THE CPU**

How can the operating system regain control of the CPU so that it can switch between processes?

A Cooperative Approach: Wait For System Calls

One approach that some systems have taken in the past (for example, early versions of the Macintosh operating system [M11], or the old Xerox Alto system [A79]) is known as the cooperative approach. In this style, the OS trusts the processes of the system to behave reasonably. Processes that run for too long are assumed to periodically give up the CPU so that the OS can decide to run some other task.

Thus, you might ask, how does a friendly process give up the CPU in this utopian world? Most processes, as it turns out, transfer control of the CPU to the OS quite frequently by making system calls, for example, to open a file and subsequently read it, or to send a message to another machine, or to create a new process. Systems like this often include an
Operating systems often have to deal with misbehaving processes, those that either through design (maliciousness) or accident (bugs) attempt to do something that they shouldn’t. In modern systems, the way the OS tries to handle such malfeasance is to simply terminate the offender. One strike and you’re out! Perhaps brutal, but what else should the OS do when you try to access memory illegally or execute an illegal instruction?

The OS can regain control of the CPU by waiting for a system call or an illegal operation of some kind to take place. You might also be thinking: isn’t this passive approach less than ideal? What happens, for example, if a process (whether malicious, or just full of bugs) ends up in an infinite loop, and never makes a system call? What can the OS do then?

A Non-Cooperative Approach: The OS Takes Control

Without some additional help from the hardware, it turns out the OS can’t do much at all when a process refuses to make system calls (or mistakes) and thus return control to the OS. In fact, in the cooperative approach, your only recourse when a process gets stuck in an infinite loop is to resort to the age-old solution to all problems in computer systems: reboot the machine. Thus, we again arrive at a subproblem of our general quest to gain control of the CPU.

The answer turns out to be simple and was discovered by a number of people building computer systems many years ago: a timer interrupt [M+63]. A timer device can be programmed to raise an interrupt every so many milliseconds; when the interrupt is raised, the currently running process is halted, and a pre-configured interrupt handler in the OS runs. At this point, the OS has regained control of the CPU, and thus can do what it pleases: stop the current process, and start a different one.
The addition of a timer interrupt gives the OS the ability to run again on a CPU even if processes act in a non-cooperative fashion. Thus, this hardware feature is essential in helping the OS maintain control of the machine.

As we discussed before with system calls, the OS must inform the hardware of which code to run when the timer interrupt occurs; thus, at boot time, the OS does exactly that. Second, also during the boot sequence, the OS must start the timer, which is of course a privileged operation. Once the timer has begun, the OS can thus feel safe in that control will eventually be returned to it, and thus the OS is free to run user programs. The timer can also be turned off (also a privileged operation), something we will discuss later when we understand concurrency in more detail.

Note that the hardware has some responsibility when an interrupt occurs, in particular to save enough of the state of the program that was running when the interrupt occurred such that a subsequent return-from-trap instruction will be able to resume the running program correctly. This set of actions is quite similar to the behavior of the hardware during an explicit system-call trap into the kernel, with various registers thus getting saved (e.g., onto a kernel stack) and thus easily restored by the return-from-trap instruction.

### Saving and Restoring Context

Now that the OS has regained control, whether cooperatively via a system call, or more forcefully via a timer interrupt, a decision has to be made: whether to continue running the currently-running process, or switch to a different one. This decision is made by a part of the operating system known as the scheduler; we will discuss scheduling policies in great detail in the next few chapters.

If the decision is made to switch, the OS then executes a low-level piece of code which we refer to as a context switch. A context switch is conceptually simple: all the OS has to do is save a few register values for the currently-executing process (onto its kernel stack, for example) and restore a few for the soon-to-be-executing process (from its kernel stack). By doing so, the OS thus ensures that when the return-from-trap instruction is finally executed, instead of returning to the process that was running, the system resumes execution of another process.

To save the context of the currently-running process, the OS will execute some low-level assembly code to save the general purpose registers, PC, as well as the kernel stack pointer of the currently-running process, and then restore said registers, PC, and switch to the kernel stack for the soon-to-be-executing process. By switching stacks, the kernel enters the
OS @ boot (kernel mode) | Hardware
---|---
initialize trap table | remember addresses of...
syscall handler
timer handler
start interrupt timer | start timer
interrupt CPU in X ms

OS @ run (kernel mode) | Hardware | Program (user mode)
---|---|---
timer interrupt | save regs(A) to k-stack(A)
move to kernel mode
jump to trap handler | Process A
...
Handle the trap
Call switch() routine
save regs(A) to proc-struct(A)
restore regs(B) from proc-struct(B)
switch to k-stack(B)
return-from-trap (into B)
restore regs(B) from k-stack(B)
move to user mode
jump to B’s PC | Process B
...

Table 6.3: Limited Direction Execution Protocol (Timer Interrupt)

call to the switch code in the context of one process (the one that was interrupted) and returns in the context of another (the soon-to-be-executing one). When the OS then finally executes a return-from-trap instruction, the soon-to-be-executing process becomes the currently-running process. And thus the context switch is complete.

A timeline of the entire process is shown in Table 6.3. In this example, Process A is running and then is interrupted by the timer interrupt. The hardware saves its state (onto its kernel stack) and enters the kernel (switching to kernel mode). In the timer interrupt handler, the OS decides to switch from running Process A to Process B. At that point, it calls the switch() routine, which carefully saves current register values (into the process structure of A), restores the registers of Process B (from its process structure entry), and then switches contexts, specifically by changing the stack pointer to use B’s kernel stack (and not A’s). Finally, the OS returns-from-trap, which restores B’s register state and starts running it.

Note that there are two types of register saves/restores that happen during this protocol. The first is when the timer interrupt occurs; in this case, the user register state of the running process is implicitly saved by the hardware, using the kernel stack of that process. The second is when the OS decides to switch from A to B; in this case, the kernel register state...
is explicitly saved by the software (i.e., the OS), but this time into memory in the process structure of the process. The latter action moves the system from running as if it just trapped into the kernel from A to as if it just trapped into the kernel from B.

To give you a better sense of how such a switch is enacted, Figure 6.1 shows the context switch code for xv6. See if you can make sense of it (you’ll have to know a bit of x86, as well as some xv6, to do so). The context structures old and new are found the old and new process’s process structures, respectively.

6.4 Worried About Concurrency?

Some of you, as attentive and thoughtful readers, may be now thinking: “Hmm... what happens when, during a system call, a timer interrupt occurs?” or “What happens when you’re handling one interrupt and another one happens? Doesn’t that get hard to handle in the kernel?” Good questions – we really have some hope for you yet!

The answer is yes, the OS does indeed need to be concerned as to what happens if, during interrupt or trap handling, another interrupt occurs. This, in fact, is the exact topic of the entire second piece of this book, on concurrency; we’ll defer a detailed discussion until then.
 Aside: How Long Context Switches Take

A natural question you might have is: how long does something like a context switch take? Or even a system call? For those of you that are curious, there is a tool called lmbench [MS96] that measures exactly those things, as well as a few other performance measures that might be relevant.

Results have improved quite a bit over time, roughly tracking processor performance. For example, in 1996 running Linux 1.3.37 on a 200-MHz P6 CPU, system calls took roughly 4 microseconds, and a context switch roughly 6 microseconds [MS96]. Modern systems perform almost an order of magnitude better, with sub-microsecond results on systems with 2- or 3-GHz processors.

It should be noted that not all operating-system actions track CPU performance. As Ousterhout observed, many OS operations are memory intensive, and memory bandwidth has not improved as dramatically as processor speed over time [O90]. Thus, depending on your workload, buying the latest and greatest processor may not speed up your OS as much as you might hope.

To whet your appetite, we’ll just sketch some basics of how the OS handles these tricky situations. One simple thing an OS might do is disable interrupts during interrupt processing; doing so ensures that when one interrupt is being handled, no other one will be delivered to the CPU. Of course, the OS has to be careful in doing so; disabling interrupts for too long could lead to lost interrupts, which is (in technical terms) bad.

Operating systems also have developed a number of sophisticated locking schemes to protect concurrent access to internal data structures. This enables multiple activities to be on-going within the kernel at the same time, particularly useful on multiprocessors. As we’ll see in the next piece of this book on concurrency, though, such locking can be complicated and lead to a variety of interesting and hard-to-find bugs.

6.5 Summary

We have described some key low-level mechanisms to implement CPU virtualization, a set of techniques which we collectively refer to as limited direct execution. The basic idea is straightforward: just run the program you want to run on the CPU, but first make sure to set up the hardware so as to limit what the process can do without OS assistance.

This general approach is taken in real life as well. For example, those of you who have children, or, at least, have heard of children, may be familiar with the concept of baby proofing a room: locking cabinets containing dangerous stuff and covering electrical sockets. When the room is thus readied, you can let your baby roam freely, secure in the knowledge that the most dangerous aspects of the room have been restricted.
Tip: Reboot Is Useful

Earlier on, we noted that the only solution to infinite loops (and similar behaviors) under cooperative preemption is to **reboot** the machine. While you may scoff at this hack, researchers have shown that reboot (or in general, starting over some piece of software) can be a hugely useful tool in building robust systems [C+04].

Specifically, reboot is useful because it moves software back to a known and likely more tested state. Reboots also reclaim stale or leaked resources (e.g., memory) which may otherwise be hard to handle. Finally, reboots are easy to automate. For all of these reasons, it is not uncommon in large-scale cluster Internet services for system management software to periodically reboot sets of machines in order to reset them and thus obtain the advantages listed above.

Thus, next time you reboot, you are not just enacting some ugly hack. Rather, you are using a time-tested approach to improving the behavior of a computer system. Well done!

In an analogous manner, the OS “baby proofs” the CPU, by first (during boot time) setting up the trap handlers and starting an interrupt timer, and then by only running processes in a restricted mode. By doing so, the OS can feel quite assured that processes can run efficiently, only requiring OS intervention to perform privileged operations or when they have monopolized the CPU for too long and thus need to be switched out.

We thus have the basic mechanisms for virtualizing the CPU in place. But a major question is left unanswered: which process should we run at a given time? It is this question that the scheduler must answer, and thus the next topic of our study.
References

Xerox Palo Alto Research Center, September 1979
An amazing system, way ahead of its time. Became famous because Steve Jobs visited, took notes, and built Lisa and eventually Mac.

[C+04] “Microreboot – A Technique for Cheap Recovery”
George Candea, Shinichi Kawamoto, Yuichi Fujiki, Greg Friedman, Armando Fox
OSDI ’04, San Francisco, CA, December 2004
An excellent paper pointing out how far one can go with reboot in building more robust systems.

Volume 3A and 3B: System Programming Guide
Intel Corporation, January 2011

[K+61] “One-Level Storage System”
IRE Transactions on Electronic Computers, April 1962
The Atlas pioneered much of what you see in modern systems. However, this paper is not the best one to read. If you were to only read one, you might try the historical perspective below [L78].

S. H. Lavington
Communications of the ACM, 21:1, January 1978
A history of the early development of computers and the pioneering efforts of Atlas.

J. McCarthy, S. Boilen, E. Fredkin, J. C. R. Licklider
AFIPS ’63 (Spring), May, 1963, New York, USA
An early paper about time-sharing that refers to using a timer interrupt; the quote that discusses it:
“The basic task of the channel 17 clock routine is to decide whether to remove the current user from core and if so to decide which user program to swap in as he goes out.”

[MS96] “lmbench: Portable tools for performance analysis”
Larry McVoy and Carl Staelin
USENIX Annual Technical Conference, January 1996
A fun paper about how to measure a number of different things about your OS and its performance. Download lmbench and give it a try.

[M11] “Mac OS 9”
January 2011
Available: http://en.wikipedia.org/wiki/Mac_OS_9

[O90] “Why Aren’t Operating Systems Getting Faster as Fast as Hardware?”
J. Ousterhout
USENIX Summer Conference, June 1990
A classic paper on the nature of operating system performance.

The Open Group, May 2010
This is hard and painful to read, so probably avoid it if you can.

[S07] “The Geometry of Innocent Flesh on the Bone: Return-into-libc without Function Calls (on the x86)”
Hovav Shacham
CCS ’07, October 2007
One of those awesome, mind-blowing ideas that you’ll see in research from time to time. The author shows that if you can jump into code arbitrarily, you can essentially stitch together any code sequence you like (given a large code base) – read the paper for the details. The technique makes it even harder to defend against malicious attacks, alas.
Homework (Measurement)

**Aside: Measurement Homeworks**

Measurement homeworks are small exercises where you write code to run on a real machine, in order to measure some aspect of OS or hardware performance. The idea behind such homeworks is to give you a little bit of hands-on experience with a real operating system.

In this homework, you’ll measure the costs of a system call and context switch. Measuring the cost of a system call is relatively easy. For example, you could repeatedly call a simple system call (e.g., performing a 0-byte read), and time how long it takes; dividing the time by the number of iterations gives you an estimate of the cost of a system call.

One thing you’ll have to take into account is the precision and accuracy of your timer. A typical timer that you can use is `gettimeofday()`; read the man page for details. What you’ll see there is that `gettimeofday()` returns the time in microseconds since 1970; however, this does not mean that the timer is precise to the microsecond. Measure back-to-back calls to `gettimeofday()` to learn something about how precise the timer really is; this will tell you how many iterations of your null system-call test you’ll have to run in order to get a good measurement result. If `gettimeofday()` is not precise enough for you, you might look into using the `rdtsc` instruction available on x86 machines.

Measuring the cost of a context switch is a little trickier. The lmbench benchmark does so by running two processes on a single CPU, and setting up two UNIX pipes between them; a pipe is just one of many ways processes in a UNIX system can communicate with one another. The first process then issues a write to the first pipe, and waits for a read on the second; upon seeing the first process waiting for something to read from the second pipe, the OS puts the first process in the blocked state, and switches to the other process, which reads from the first pipe and then writes to the second. When the second process tries to read from the first pipe again, it blocks, and thus the back-and-forth cycle of communication continues. By measuring the cost of communicating like this repeatedly, lmbench can make a good estimate of the cost of a context switch. You can try to re-create something similar here, using pipes, or perhaps some other communication mechanism such as UNIX sockets.

One difficulty in measuring context-switch cost arises in systems with more than one CPU; what you need to do on such a system is ensure that your context-switching processes are located on the same processor. Fortunately, most operating systems have calls to bind a process to a particular processor; on Linux, for example, the `sched_setaffinity()` call is what you’re looking for. By ensuring both processes are on the same processor, you are making sure to measure the cost of the OS stopping one process and restoring another on the same CPU.
Scheduling: Introduction

By now low-level mechanisms of running processes (e.g., context switching) should be clear; if they are not, go back a chapter or two, and read the description of how that stuff works again. However, we have yet to understand the high-level policies that an OS scheduler employs. We will now do just that, presenting a series of scheduling policies (sometimes called disciplines) that various smart and hard-working people have developed over the years.

The origins of scheduling, in fact, predate computer systems; early approaches were taken from the field of operations management and applied to computers. This reality should be no surprise: assembly lines and many other human endeavors also require scheduling, and many of the same concerns exist therein, including a laser-like desire for efficiency.

And thus, our problem:

**THE CRUX: HOW TO DEVELOP SCHEDULING POLICY**

How should we develop a basic framework for thinking about scheduling policies? What are the key assumptions? What metrics are important? What basic approaches have been used in the earliest of computer systems?

7.1 Workload Assumptions

Before getting into the range of possible policies, let us first make a number of simplifying assumptions about the processes running in the system, sometimes collectively called the workload. Determining the workload is a critical part of building policies, and the more you know about workload, the more fine-tuned your policy can be.

The workload assumptions we make here are clearly unrealistic, but that is alright (for now), because we will relax them as we go, and eventually develop what we will refer to as ... (dramatic pause) ...
a **fully-operational scheduling discipline**\(^1\).

We will make the following assumptions about the processes, sometimes called **jobs**, that are running in the system:

1. Each job runs for the same amount of time.
2. All jobs arrive at the same time.
3. All jobs only use the CPU (i.e., they perform no I/O)
4. The run-time of each job is known.

We said all of these assumptions were unrealistic, but just as some animals are more equal than others in Orwell’s *Animal Farm* [O45], some assumptions are more unrealistic than others in this chapter. In particular, it might bother you that the run-time of each job is known: this would make the scheduler omniscient, which, although it would be great (probably), is not likely to happen anytime soon.

### 7.2 Scheduling Metrics

Beyond making workload assumptions, we also need one more thing to enable us to compare different scheduling policies: a **scheduling metric**. A metric is just something that we use to **measure** something, and there are a number of different metrics that make sense in scheduling.

For now, however, let us also simplify our life by simply having a single metric: **turnaround time**. The turnaround time of a job is defined as the time at which the job completes minus the time at which the job arrived in the system. More formally, the turnaround time \( T_{\text{turnaround}} \) is:

\[
T_{\text{turnaround}} = T_{\text{completion}} - T_{\text{arrival}} \tag{7.1}
\]

Because we have assumed that all jobs arrive at the same time, for now \( T_{\text{arrival}} = 0 \) and hence \( T_{\text{turnaround}} = T_{\text{completion}} \). This fact will change as we relax the aforementioned assumptions.

You should note that turnaround time is a **performance metric**, which will be our primary focus this chapter. Another metric of interest is **fairness**, as measured (for example) by *Jain’s Fairness Index* [J91]. Performance and fairness are often at odds in scheduling; a scheduler, for example, may optimize performance but at the cost of preventing a few jobs from running, thus decreasing fairness. This conundrum shows us that life isn’t always perfect.

### 7.3 First In, First Out (FIFO)

The most basic algorithm a scheduler can implement is known as **First In, First Out (FIFO)** scheduling or sometimes **First Come, First Served**

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\(^1\)Said in the same way you would say “A fully-operational Death Star.”
FIFO has a number of positive properties: it is clearly very simple and thus easy to implement. Given our assumptions, it works pretty well. Let’s do a quick example together. Imagine three jobs arrive in the system, A, B, and C, at roughly the same time \( T_{arrival} = 0 \). Because FIFO has to put some job first, let’s assume that while they all arrived simultaneously, A arrived just a hair before B which arrived just a hair before C. Assume also that each job runs for 10 seconds. What will the average turnaround time be for these jobs?

From Figure 7.1, you can see that A finished at 10, B at 20, and C at 30. Thus, the average turnaround time for the three jobs is simply \( \frac{10+20+30}{3} = 20 \). Computing turnaround time is as easy as that.

Now let’s relax one of our assumptions. In particular, let’s relax assumption 1, and thus no longer assume that each job runs for the same amount of time. How does FIFO perform now? What kind of workload could you construct to make FIFO perform poorly?

Presumably you’ve figured this out by now, but just in case, let’s do an example to show how jobs of different lengths can lead to trouble for FIFO scheduling. In particular, let’s again assume three jobs (A, B, and C), but this time A runs for 100 seconds while B and C run for 10 each.

As you can see in Figure 7.2, Job A runs first for the full 100 seconds before B or C even get a chance to run. Thus, the average turnaround time for the system is high: a painful 110 seconds \( \frac{100+110+120}{3} = 110 \).

This problem is generally referred to as the convoy effect [B+79], where a number of relatively-short potential consumers of a resource get queued behind a heavyweight resource consumer. This scheduling scenario might remind you of a single line at a grocery store and what you feel like when

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Three Easy Pieces
you see the person in front of you with three carts full of provisions and a checkbook out; it’s going to be a while. 

So what should we do? How can we develop a better algorithm to deal with our new reality of jobs that run for different amounts of time? Think about it first; then read on.

7.4 Shortest Job First (SJF)

It turns out that a very simple approach solves this problem; in fact it is an idea stolen from operations research [C54,PV56] and applied to scheduling of jobs in computer systems. This new scheduling discipline is known as **Shortest Job First (SJF)**, and the name should be easy to remember because it describes the policy quite completely: it runs the shortest job first, then the next shortest, and so on.

![Figure 7.3: SJF Simple Example](image)

Let’s take our example above but with SJF as our scheduling policy. Figure 7.3 shows the results of running A, B, and C. Hopefully the diagram makes it clear why SJF performs much better with regards to average turnaround time. Simply by running B and C before A, SJF reduces average turnaround from 110 seconds to 50 (\( \frac{10 + 20 + 120}{3} = 50 \)), more than a factor of two improvement.

In fact, given our assumptions about jobs all arriving at the same time, we could prove that SJF is indeed an **optimal** scheduling algorithm. How-

---

2Recommended action in this case: either quickly switch to a different line, or take a long, deep, and relaxing breath. That’s right, breathe in, breathe out. It will be OK, don’t worry.
In the old days of batch computing, a number of non-preemptive schedulers were developed; such systems would run each job to completion before considering whether to run a new job. Virtually all modern schedulers are preemptive, and quite willing to stop one process from running in order to run another. This implies that the scheduler employs the mechanisms we learned about previously; in particular, the scheduler can perform a context switch, stopping one running process temporarily and resuming (or starting) another.

ever, you are in a systems class, not theory or operations research; no proofs are allowed.

Thus we arrive upon a good approach to scheduling with SJF, but our assumptions are still fairly unrealistic. Let’s relax another. In particular, we can target assumption 2, and now assume that jobs can arrive at any time instead of all at once. What problems does this lead to?

(Another pause to think ... are you thinking? Come on, you can do it)

Here we can illustrate the problem again with an example. This time, assume A arrives at $t = 0$ and needs to run for 100 seconds, whereas B and C arrive at $t = 10$ and each need to run for 10 seconds. With pure SJF, we’d get the schedule seen in Figure 7.4.

As you can see from the figure, even though B and C arrived shortly after A, they still are forced to wait until A has completed, and thus suffer the same convoy problem. Average turnaround time for these three jobs is $103.33$ seconds ($\frac{100 + (110 - 10) + (120 - 10)}{3}$). What can a scheduler do?

7.5 Shortest Time-to-Completion First (STCF)

As you might have guessed, given our previous discussion about mechanisms such as timer interrupts and context switching, the scheduler can certainly do something else when B and C arrive: it can preempt job A and decide to run another job, perhaps continuing A later. SJF by our definition is a non-preemptive scheduler, and thus suffers from the problems described above.
Fortunately, there is a scheduler which does exactly that: add preemption to SJF, known as the Shortest Time-to-Completion First (STCF) or Preemptive Shortest Job First (PSJF) scheduler [CK68]. Any time a new job enters the system, it determines of the remaining jobs and new job, which has the least time left, and then schedules that one. Thus, in our example, STCF would preempt A and run B and C to completion; only when they are finished would A’s remaining time be scheduled. Figure 7.5 shows an example.

The result is a much-improved average turnaround time: 50 seconds ((120−0)+(20−10)+(30−10))/3). And as before, given our new assumptions, STCF is provably optimal; given that SJF is optimal if all jobs arrive at the same time, you should probably be able to see the intuition behind the optimality of STCF.

Thus, if we knew that job lengths, and jobs only used the CPU, and our only metric was turnaround time, STCF would be a great policy. In fact, for a number of early batch computing systems, these types of scheduling algorithms made some sense. However, the introduction of time-shared machines changed all that. Now users would sit at a terminal and demand interactive performance from the system as well. And thus, a new metric was born: response time.

Response time is defined as the time from when the job arrives in a system to the first time it is scheduled. More formally:

\[ T_{response} = T_{firstrun} - T_{arrival} \]  

For example, if we had the schedule above (with A arriving at time 0, and B and C at time 10), the response time of each job is as follows: 0 for job A, 0 for B, and 10 for C (average: 3.33).

As you might be thinking, STCF and related disciplines are not particularly good for response time. If three jobs arrive at the same time, for example, the third job has to wait for the previous two jobs to run in their entirety before being scheduled just once. While great for turnaround time, this approach is quite bad for response time and interactivity. Indeed, imagine sitting at a terminal, typing, and having to wait 10 seconds to see a response from the system just because some other job got scheduled in front of yours: not too pleasant.

Thus, we are left with another problem: how can we build a scheduler that is sensitive to response time?
7.6 Round Robin

To solve this problem, we will introduce a new scheduling algorithm. This approach is classically known as Round-Robin (RR) scheduling [K64]. The basic idea is simple: instead of running jobs to completion, RR runs a job for a time slice (sometimes called a scheduling quantum) and then switches to the next job in the run queue. It repeatedly does so until the jobs are finished. For this reason, RR is sometimes called time-slicing. Note that the length of a time slice must be a multiple of the timer-interrupt period; thus if the timer interrupts every 10 milliseconds, the time slice could be 10, 20, or any other multiple of 10 ms.

To understand RR in more detail, let’s look at an example. Assume three jobs A, B, and C arrive at the same time in the system, and that they each wish to run for 5 seconds. An SJF scheduler runs each job to completion before running another (Figure 7.6). In contrast, RR with a time-slice of 1 second would cycle through the jobs quickly (Figure 7.7).

The average response time of RR is: \( \frac{0+1+2}{3} = 1 \); for SJF, average response time is: \( \frac{0+5+10}{3} = 5 \).

As you can see, the length of the time slice is critical for RR. The shorter it is, the better the performance of RR under the response-time metric. However, making the time slice too short is problematic: suddenly the cost of context switching will dominate overall performance. Thus, deciding on the length of the time slice presents a trade-off to a system designer, making it long enough to amortize the cost of switching without making it so long that the system is no longer responsive.

Note that the cost of context switching does not arise solely from the OS actions of saving and restoring a few registers. When programs run,
The general technique of amortization is commonly used in systems when there is a fixed cost to some operation. By incurring that cost less often (i.e., by performing the operation fewer times), the total cost to the system is reduced. For example, if the time slice is set to 10 ms, and the context-switch cost is 1 ms, roughly 10% of time is spent context switching and is thus wasted. If we want to amortize this cost, we can increase the time slice, e.g., to 100 ms. In this case, less than 1% of time is spent context switching, and thus the cost of time-slicing has been amortized.

they build up a great deal of state in CPU caches, TLBs, branch predictors, and other on-chip hardware. Switching to another job causes this state to be flushed and new state relevant to the currently-running job to be brought in, which may exact a noticeable performance cost [MB91].

RR, with a reasonable time slice, is thus an excellent scheduler if response time is our only metric. But what about our old friend turnaround time? Let’s look at our example above again. A, B, and C, each with running times of 5 seconds, arrive at the same time, and RR is the scheduler with a (long) 1-second time slice. We can see from the picture above that A finishes at 13, B at 14, and C at 15, for an average of 14. Pretty awful!

It is not surprising, then, that RR is indeed one of the worst policies if turnaround time is our metric. Intuitively, this should make sense: what RR is doing is stretching out each job as long as it can, by only running each job for a short bit before moving to the next. Because turnaround time only cares about when jobs finish, RR is nearly pessimal, even worse than simple FIFO in many cases.

More generally, any policy (such as RR) that is fair, i.e., that evenly divides the CPU among active processes on a small time scale, will perform poorly on metrics such as turnaround time. Indeed, this is an inherent trade-off: if you are willing to be unfair, you can run shorter jobs to completion, but at the cost of response time; if you instead value fairness, response time is lowered, but at the cost of turnaround time. This type of trade-off is common in systems; you can’t have your cake and eat it too.

We have developed two types of schedulers. The first type (SJF, STCF) optimizes turnaround time, but is bad for response time. The second type (RR) optimizes response time but is bad for turnaround. And we still have two assumptions which need to be relaxed: assumption 3 (that jobs do no I/O), and assumption 4 (that the run-time of each job is known). Let’s tackle those assumptions next.

7.7 Incorporating I/O

First we will relax assumption 3 – of course all programs perform I/O. Imagine a program that didn’t take any input: it would produce the same output each time. Imagine one without output: it is the proverbial tree
falling in the forest, with no one to see it; it doesn’t matter that it ran.

A scheduler clearly has a decision to make when a job initiates an I/O request, because the currently-running job won’t be using the CPU during the I/O; it is **blocked** waiting for I/O completion. If the I/O is sent to a hard disk drive, the process might be blocked for a few milliseconds or longer, depending on the current I/O load of the drive. Thus, the scheduler should probably schedule another job on the CPU at that time.

The scheduler also has to make a decision when the I/O completes. When that occurs, an interrupt is raised, and the OS runs and moves the process that issued the I/O from blocked back to the ready state. Of course, it could even decide to run the job at that point. How should the OS treat each job?

To understand this issue better, let us assume we have two jobs, A and B, which each need 50 ms of CPU time. However, there is one obvious difference: A runs for 10 ms and then issues an I/O request (assume here that I/Os each take 10 ms), whereas B simply uses the CPU for 50 ms and performs no I/O. The scheduler runs A first, then B after (Figure 7.8).

![Figure 7.8: Poor Use of Resources](image)

Assume we are trying to build a STCF scheduler. How should such a scheduler account for the fact that A is broken up into 5 10-ms sub-jobs, whereas B is just a single 50-ms CPU demand? Clearly, just running one job and then the other without considering how to take I/O into account makes little sense.

![Figure 7.9: Overlap Allows Better Use of Resources](image)

A common approach is to treat each 10-ms sub-job of A as an independent job. Thus, when the system starts, its choice is whether to schedule a 10-ms A or a 50-ms B. With STCF, the choice is clear: choose the shorter one, in this case A. Then, when the first sub-job of A has completed, only B is left, and it begins running. Then a new sub-job of A is submitted, and it preempts B and runs for 10 ms. Doing so allows for **overlap**, with
TIP: OVERLAP ENABLES HIGHER UTILIZATION
When possible, overlap operations to maximize the utilization of systems. Overlap is useful in many different domains, including when performing disk I/O or sending messages to remote machines; in either case, starting the operation and then switching to other work is a good idea, and improved the overall utilization and efficiency of the system.

the CPU being used by one process while waiting for the I/O of another process to complete; the system is thus better utilized (see Figure 7.9).

And thus we see how a scheduler might incorporate I/O. By treating each CPU burst as a job, the scheduler makes sure processes that are “interactive” get run frequently. While those interactive jobs are performing I/O, other CPU-intensive jobs run, thus better utilizing the processor.

7.8 No More Oracle

With a basic approach to I/O in place, we come to our final assumption: that the scheduler knows the length of each job. As we said before, this is likely the worst assumption we could make. In fact, in a general-purpose OS (like the ones we care about), the OS usually knows very little about the length of each job. Thus, how can we build an approach that behaves like SJF/STCF without such a priori knowledge? Further, how can we incorporate some of the ideas we have seen with the RR scheduler so that response time is also quite good?

7.9 Summary

We have introduced the basic ideas behind scheduling and developed two families of approaches. The first runs the shortest job remaining and thus optimizes turnaround time; the second alternates between all jobs and thus optimizes response time. Both are bad where the other is good, alas, an inherent trade-off common in systems. We have also seen how we might incorporate I/O into the picture, but have still not solved the problem of the fundamental inability of the OS to see into the future. Shortly, we will see how to overcome this problem, by building a scheduler that uses the recent past to predict the future. This scheduler is known as the multi-level feedback queue, and it is the topic of the next chapter.
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Follow-on work that generalizes the SJF approach to machine repair from Cobham’s original work; also postulates the utility of an STCF approach in such an environment. Specifically, “There are certain types of repair work, ... involving much dismantling and covering the floor with nuts and bolts, which certainly should not be interrupted once undertaken; in other cases it would be inadvisable to continue work on a long job if one or more short ones became available (p.81).”

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A nice study on how cache performance can be affected by context switching; less of an issue in today’s systems where processors issue billions of instructions per second but context-switches still happen in the millisecond time range.
Homework

**Aside: Simulation Homeworks**

Simulation homeworks come in the form of simulators you run to make sure you understand some piece of the material. The simulators are generally python programs that enable you both to generate different problems (using different random seeds) as well as to have the program solve the problem for you (with the `--c` flag) so that you can check your answers. Running any simulator with a `--h` or `--help` flag will provide with more information as to all the options the simulator gives you.

This program, `scheduler.py`, allows you to see how different schedulers perform under scheduling metrics such as response time, turnaround time, and total wait time. See the README for details.

**Questions**

1. Compute the response time and turnaround time when running three jobs of length 200 with the SJF and FIFO schedulers.
2. Now do the same but with jobs of different lengths: 100, 200, and 300.
3. Now do the same, but also with the RR scheduler and a time-slice of 1.
4. For what types of workloads does SJF deliver the same turnaround times as FIFO?
5. For what types of workloads and quantum lengths does SJF deliver the same response times as RR?
6. What happens to response time with SJF as job lengths increase? Can you use the simulator to demonstrate the trend?
7. What happens to response time with RR as quantum lengths increase? Can you write an equation that gives the worst-case response time, given $N$ jobs?
In this chapter, we’ll tackle the problem of developing one of the most well-known approaches to scheduling, known as the Multi-level Feedback Queue (MLFQ). The Multi-level Feedback Queue (MLFQ) scheduler was first described by Corbato et al. in 1962 [C+62] in a system known as the Compatible Time-Sharing System (CTSS), and this work, along with later work on Multics, led the ACM to award Corbato its highest honor, the Turing Award. The scheduler has subsequently been refined throughout the years to the implementations you will encounter in some modern systems.

The fundamental problem MLFQ tries to address is two-fold. First, it would like to optimize turnaround time, which, as we saw in the previous note, is done by running shorter jobs first; unfortunately, the OS doesn’t generally know how long a job will run for, exactly the knowledge that algorithms like SJF (or STCF) require. Second, MLFQ would like to make a system feel responsive to interactive users (i.e., users sitting and staring at the screen, waiting for a process to finish), and thus minimize response time; unfortunately, algorithms like Round Robin reduce response time but are terrible for turnaround time. Thus, our problem: given that we in general do not know anything about a process, how can we build a scheduler to achieve these goals? How can the scheduler learn, as the system runs, the characteristics of the jobs it is running, and thus make better scheduling decisions?

The Crux:
How To Schedule Without Perfect Knowledge?
How can we design a scheduler that both minimizes response time for interactive jobs while also minimizing turnaround time without a priori knowledge of job length?
The multi-level feedback queue is an excellent example of a system that learns from the past to predict the future. Such approaches are common in operating systems (and many other places in Computer Science, including hardware branch predictors and caching algorithms). Such approaches work when jobs have phases of behavior and are thus predictable; of course, one must be careful with such techniques, as they can easily be wrong and drive a system to make worse decisions than they would have with no knowledge at all.

8.1 MLFQ: Basic Rules

To build such a scheduler, in this chapter we will describe the basic algorithms behind a multi-level feedback queue; although the specifics of many implemented MLFQs differ [E95], most approaches are similar.

In our treatment, the MLFQ has a number of distinct queues, each assigned a different priority level. At any given time, a job that is ready to run is on a single queue. MLFQ uses priorities to decide which job should run at a given time: a job with higher priority (i.e., a job on a higher queue) is chosen to run.

Of course, more than one job may be on a given queue, and thus have the same priority. In this case, we will just use round-robin scheduling among those jobs.

Thus, the key to MLFQ scheduling lies in how the scheduler sets priorities. Rather than giving a fixed priority to each job, MLFQ varies the priority of a job based on its observed behavior. If, for example, a job repeatedly relinquishes the CPU while waiting for input from the keyboard, MLFQ will keep its priority high, as this is how an interactive process might behave. If, instead, a job uses the CPU intensively for long periods of time, MLFQ will reduce its priority. In this way, MLFQ will try to learn about processes as they run, and thus use the history of the job to predict its future behavior.

Thus, we arrive at the first two basic rules for MLFQ:

- **Rule 1**: If Priority(A) > Priority(B), A runs (B doesn’t).
- **Rule 2**: If Priority(A) = Priority(B), A & B run in RR.

If we were to put forth a picture of what the queues might look like at a given instant, we might see something like the following (Figure 8.1). In the figure, two jobs (A and B) are at the highest priority level, while job C is in the middle and job D is at the lowest priority. Given our current knowledge of how MLFQ works, the scheduler would just alternate time slices between A and B because they are the highest priority jobs in the system; poor jobs C and D would never even get to run – an outrage!

Of course, just showing a static snapshot of some queues does not really give you an idea of how MLFQ works. What we need is to under-
stand how job priority changes over time. And that, in a surprise only to those who are reading a chapter from this book for the first time, is exactly what we will do next.

### 8.2 Attempt #1: How to Change Priority

We now must decide how MLFQ is going to change the priority level of a job (and thus which queue it is on) over the lifetime of a job. To do this, we must keep in mind our workload: a mix of interactive jobs that are short-running (and may frequently relinquish the CPU), and some longer-running “CPU-bound” jobs that need a lot of CPU time but where response time isn’t important. Here is our first attempt at a priority-adjustment algorithm:

- **Rule 3:** When a job enters the system, it is placed at the highest priority (the topmost queue).
- **Rule 4a:** If a job uses up an entire time slice while running, its priority is reduced (i.e., it moves down one queue).
- **Rule 4b:** If a job gives up the CPU before the time slice is up, it stays at the same priority level.

**Example 1: A Single Long-Running Job**

Let’s look at some examples. First, we’ll look at what happens when there has been a long running job in the system. Figure 8.2 shows what happens to this job over time in a three-queue scheduler.
As you can see in the example, the job enters at the highest priority (Q2). After a single time-slice of 10 ms, the scheduler reduces the job’s priority by one, and thus the job is on Q1. After running at Q1 for a time slice, the job is finally lowered to the lowest priority in the system (Q0), where it remains. Pretty simple, no?

**Example 2: Along Came A Short Job**

Now let’s look at a more complicated example, and hopefully see how MLFQ tries to approximate SJF. In this example, there are two jobs: A, which is a long-running CPU-intensive job, and B, which is a short-running interactive job. Assume A has been running for some time, and then B arrives. What will happen? Will MLFQ approximate SJF for B?

Figure 8.3 plots the results of this scenario. A (shown in black) is running along in the lowest-priority queue (as would any long-running CPU-intensive jobs); B (shown in gray) arrives at time \( T = 100 \), and thus is
SCHEDULING:
THE MULTI-LEVEL FEEDBACK QUEUE

Figure 8.4: A Mixed I/O-intensive and CPU-intensive Workload
inserted into the highest queue; as its run-time is short (only 20 ms), B completes before reaching the bottom queue, in two time slices; then A resumes running (at low priority).

From this example, you can hopefully understand one of the major goals of the algorithm: because it doesn’t know whether a job will be a short job or a long-running job, it first assumes it might be a short job, thus giving the job high priority. If it actually is a short job, it will run quickly and complete; if it is not a short job, it will slowly move down the queues, and thus soon prove itself to be a long-running more batch-like process. In this manner, MLFQ approximates SJF.

Example 3: What About I/O?

Let’s now look at an example with some I/O. As Rule 4b states above, if a process gives up the processor before using up its time slice, we keep it at the same priority level. The intent of this rule is simple: if an interactive job, for example, is doing a lot of I/O (say by waiting for user input from the keyboard or mouse), it will relinquish the CPU before its time slice is complete; in such case, we don’t wish to penalize the job and thus simply keep it at the same level.

Figure 8.4 shows an example of how this works, with an interactive job B (shown in gray) that needs the CPU only for 1 ms before performing an I/O competing for the CPU with a long-running batch job A (shown in black). The MLFQ approach keeps B at the highest priority because B keeps releasing the CPU; if B is an interactive job, MLFQ further achieves its goal of running interactive jobs quickly.

Problems With Our Current MLFQ

We thus have a basic MLFQ. It seems to do a fairly good job, sharing the CPU fairly between long-running jobs, and letting short or I/O-intensive interactive jobs run quickly. Unfortunately, the approach we have developed thus far contains serious flaws. Can you think of any?

(This is where you pause and think as deviously as you can)
First, there is the problem of starvation: if there are “too many” interactive jobs in the system, they will combine to consume all CPU time, and thus long-running jobs will never receive any CPU time (they starve). We’d like to make some progress on these jobs even in this scenario.

Second, a smart user could rewrite their program to game the scheduler. Gaming the scheduler generally refers to the idea of doing something sneaky to trick the scheduler into giving you more than your fair share of the resource. The algorithm we have described is susceptible to the following attack: before the timeslice is over, issue an I/O operation (to some file you don’t care about) and thus relinquish the CPU; doing so allows you to remain in the same queue, and thus gain a higher percentage of CPU time. When done right (e.g., by running for 99% of a time slice before relinquishing the CPU), a job could nearly monopolize the CPU.

Finally, a program may change its behavior over time; what was CPU-bound may transition to a phase of interactivity. With our current approach, such a job would be out of luck and not be treated like the other interactive jobs in the system.

8.3 Attempt #2: The Priority Boost

Let’s try to change the rules and see if we can avoid the problem of starvation. What could we do in order to guarantee that CPU-bound jobs will make some progress (even if it is not much?).

The simple idea here is to periodically boost the priority of all the jobs in system. There are many ways to achieve this, but let’s just do something simple: throw them all in the topmost queue; hence, a new rule:

- **Rule 5**: After some time period $S$, move all the jobs in the system to the topmost queue.

Our new rule solves two problems at once. First, processes are guaranteed not to starve: by sitting in the top queue, a job will share the CPU
with other high-priority jobs in a round-robin fashion, and thus eventually receive service. Second, if a CPU-bound job has become interactive, the scheduler treats it properly once it has received the priority boost.

Let's see an example. In this scenario, we just show the behavior of a long-running job when competing for the CPU with two short-running interactive jobs. Two graphs are shown in Figure 8.5. On the left, there is no priority boost, and thus the long-running job gets starved once the two short jobs arrive; on the right, there is a priority boost every 50 ms (which is likely too small of a value, but used here for the example), and thus we at least guarantee that the long-running job will make some progress, getting boosted to the highest priority every 50 ms and thus getting to run periodically.

Of course, the addition of the time period $S$ leads to the obvious question: what should $S$ be set to? John Ousterhout, a well-regarded systems researcher [O11], used to call such values in systems voo-doo constants, because they seemed to require some form of black magic to set them correctly. Unfortunately, $S$ has that flavor. If it is set too high, long-running jobs could starve; too low, and interactive jobs may not get a proper share of the CPU.

### 8.4 Attempt #3: Better Accounting

We now have one more problem to solve: how to prevent gaming of our scheduler? The real culprit here, as you might have guessed, are Rules 4a and 4b, which let a job retain its priority by relinquishing the CPU before the time slice expires. So what should we do?

The solution here is to perform better accounting of CPU time at each level of the MLFQ. Instead of forgetting how much of a time slice a process used at a given level, the scheduler should keep track; once a process has used its allotment, it is demoted to the next priority queue. Whether
it uses the time slice in one long burst or many small ones does not matter. We thus rewrite Rules 4a and 4b to the following single rule:

- **Rule 4:** Once a job uses up its time allotment at a given level (regardless of how many times it has given up the CPU), its priority is reduced (i.e., it moves down one queue).

Let’s look at an example. Figure 8.6 shows what happens when a workload tries to game the scheduler with the old Rules 4a and 4b (on the left) as well the new anti-gaming Rule 4. Without any protection from gaming, a process can issue an I/O just before a time slice ends and thus dominate CPU time. With such protections in place, regardless of the I/O behavior of the process, it slowly moves down the queues, and thus cannot gain an unfair share of the CPU.

### 8.5 Tuning MLFQ And Other Issues

A few other issues arise with MLFQ scheduling. One big question is how to parameterize such a scheduler. For example, how many queues should there be? How big should the time slice be per queue? How often should priority be boosted in order to avoid starvation and account for changes in behavior? There are no easy answers to these questions, and thus only some experience with workloads and subsequent tuning of the scheduler will lead to a satisfactory balance.

For example, most MLFQ variants allow for varying time-slice length across different queues. The high-priority queues are usually given short time slices; they are comprised of interactive jobs, after all, and thus quickly alternating between them makes sense (e.g., 10 or fewer milliseconds). The low-priority queues, in contrast, contain long-running jobs that are CPU-bound; hence, longer time slices work well (e.g., 100s of ms). Figure 8.7 shows an example in which two long-running jobs run for 10 ms at the highest queue, 20 in the middle, and 40 at the lowest.
Avoiding voo-doo constants is a good idea whenever possible. Unfortunately, as in the example above, it is often difficult. One could try to make the system learn a good value, but that too is not straightforward. The frequent result: a configuration file filled with default parameter values that a seasoned administrator can tweak when something isn’t quite working correctly. As you can imagine, these are often left unmodified, and thus we are left to hope that the defaults work well in the field. This tip brought to you by our old OS professor, John Ousterhout, and hence we call it Ousterhout’s Law.

The Solaris MLFQ implementation – the Time-Sharing scheduling class, or TS – is particularly easy to configure; it provides a set of tables that determine exactly how the priority of a process is altered throughout its lifetime, how long each time slice is, and how often to boost the priority of a job [AD00]; an administrator can muck with this table in order to make the scheduler behave in different ways. Default values for the table are 60 queues, with slowly increasing time-slice lengths from 20 milliseconds (highest priority) to a few hundred milliseconds (lowest), and priorities boosted around every 1 second or so.

Other MLFQ schedulers don’t use a table or the exact rules described in this chapter; rather they adjust priorities using mathematical formulae. For example, the FreeBSD scheduler (version 4.3) uses a formula to calculate the current priority level of a job, basing it on how much CPU the process has used [LM+89]; in addition, usage is decayed over time, providing the desired priority boost in a different manner than described herein. See [E95] for an excellent overview of such decay-usage algorithms and their properties.

Finally, many schedulers have a few other features that you might encounter. For example, some schedulers reserve the highest priority levels for operating system work; thus typical user jobs can never obtain the highest levels of priority in the system. Some systems also allow some user advice to help set priorities; for example, by using the command-line utility nice you can increase or decrease the priority of a job (somewhat) and thus increase or decrease its chances of running at any given time. See the man page for more.

8.6 MLFQ: Summary

We have described a scheduling approach known as the Multi-Level Feedback Queue (MLFQ). Hopefully you can now see why it is called that: it has multiple levels of queues, and uses feedback to determine the priority of a given job. History is its guide: pay attention to how jobs behave over time and treat them accordingly.
The refined set of MLFQ rules, spread throughout the chapter, are reproduced here for your viewing pleasure:

- **Rule 1:** If Priority(A) > Priority(B), A runs (B doesn’t).
- **Rule 2:** If Priority(A) = Priority(B), A & B run in RR.
- **Rule 3:** When a job enters the system, it is placed at the highest priority (the topmost queue).
- **Rule 4:** Once a job uses up its time allotment at a given level (regardless of how many times it has given up the CPU), its priority is reduced (i.e., it moves down one queue).
- **Rule 5:** After some time period $S$, move all the jobs in the system to the topmost queue.

MLFQ is interesting because instead of demanding *a priori* knowledge of the nature of a job, it instead observes the execution of a job and prioritizes it accordingly. In this way, it manages to achieve the best of both worlds: it can deliver excellent overall performance (similar to SJF/STCF) for short-running interactive jobs, and is fair and makes progress for long-running CPU-intensive workloads. For this reason, many systems, including BSD UNIX derivatives [LM+89, B86], Solaris [M06], and Windows NT and subsequent Windows operating systems [CS97] use a form of MLFQ as their base scheduler.
References

[AD00] “Multilevel Feedback Queue Scheduling in Solaris”
Andrea Arpaci-Dusseau
A great short set of notes by one of the authors on the details of the Solaris scheduler. OK, we are probably biased in this description, but the notes are pretty darn good.

[B86] “The Design of the UNIX Operating System”
M.J. Bach
Prentice-Hall, 1986
One of the classic old books on how a real UNIX operating system is built; a definite must-read for kernel hackers.

[C+62] “An Experimental Time-Sharing System”
F. J. Corbato, M. M. Daggett, R. C. Daley
IFIPS 1962
A bit hard to read, but the source of many of the first ideas in multi-level feedback scheduling. Much of this later went into Multics, which one could argue was the most influential operating system of all time.

[CS97] “Inside Windows NT”
Helen Custer and David A. Solomon
Microsoft Press, 1997
The NT book, if you want to learn about something other than UNIX. Of course, why would you? OK, we’re kidding; you might actually work for Microsoft some day you know.

[E95] “An Analysis of Decay-Usage Scheduling in Multiprocessors”
D.H.J. Epema
SIGMETRICS ’95
A nice paper on the state of the art of scheduling back in the mid 1990s, including a good overview of the basic approach behind decay-usage schedulers.

[LM+89] “The Design and Implementation of the 4.3BSD UNIX Operating System”
S.J. Leffler, M.K. McKusick, M.J. Karels, J.S. Quarterman
Addison-Wesley, 1989
Another OS classic, written by four of the main people behind BSD. The later versions of this book, while more up to date, don’t quite match the beauty of this one.

[M06] “Solaris Internals: Solaris 10 and OpenSolaris Kernel Architecture”
Richard McDougall
Prentice-Hall, 2006
A good book about Solaris and how it works.

John Ousterhout
The home page of the famous Professor Ousterhout. The two co-authors of this book had the pleasure of taking graduate operating systems from Ousterhout while in graduate school; indeed, this is where the two co-authors got to know each other, eventually leading to marriage, kids, and even this book. Thus, you really can blame Ousterhout for this entire mess you’re in.

[P+95] “Informed Prefetching and Caching”
R.H. Patterson, G.A. Gibson, E. Ginting, D. Stodolsky, J. Zelenka
SOSP ’95
A fun paper about some very cool ideas in file systems, including how applications can give the OS advice about what files it is accessing and how it plans to access them.
Homework

This program, mlfq.py, allows you to see how the MLFQ scheduler presented in this chapter behaves. See the README for details.

Questions

1. Run a few randomly-generated problems with just two jobs and two queues; compute the MLFQ execution trace for each. Make your life easier by limiting the length of each job and turning off I/Os.
2. How would you run the scheduler to reproduce each of the examples in the chapter?
3. How would you configure the scheduler parameters to behave just like a round-robin scheduler?
4. Craft a workload with two jobs and scheduler parameters so that one job takes advantage of the older Rules 4a and 4b (turned on with the -S flag) to game the scheduler and obtain 99% of the CPU over a particular time interval.
5. Given a system with a quantum length of 10 ms in its highest queue, how often would you have to boost jobs back to the highest priority level (with the -B flag) in order to guarantee that a single long-running (and potentially-starving) job gets at least 5% of the CPU?
In this chapter, we’ll examine a different type of scheduler known as a proportional-share scheduler, also sometimes referred to as a fair-share scheduler. Proportional-share is based around a simple concept: instead of optimizing for turnaround or response time, a scheduler might instead try to guarantee that each job obtain a certain percentage of CPU time.

An excellent modern example of proportional-share scheduling is found in research by Waldspurger and Weihl [WW94], and is known as lottery scheduling; however, the idea is certainly much older [KL88]. The basic idea is quite simple: every so often, hold a lottery to determine which process should get to run next; processes that should run more often should be given more chances to win the lottery. Easy, no? Now, onto the details! But not before our crux:

**Crux: How To Share The CPU Proportionally**

How can we design a scheduler to share the CPU in a proportional manner? What are the key mechanisms for doing so? How effective are they?

9.1 Basic Concept: Tickets Represent Your Share

Underlying lottery scheduling is one very basic concept: tickets, which are used to represent the share of a resource that a process (or user or whatever) should receive. The percent of tickets that a process has represents its share of the system resource in question.

Let’s look at an example. Imagine two processes, A and B, and further that A has 75 tickets while B has only 25. Thus, what we would like is for A to receive 75% of the CPU and B the remaining 25%.

Lottery scheduling achieves this probabilistically (but not deterministically) by holding a lottery every so often (say, every time slice). Holding a lottery is straightforward: the scheduler must know how many total tickets there are (in our example, there are 100). The scheduler then picks...
One of the most beautiful aspects of lottery scheduling is its use of randomness. When you have to make a decision, using such a randomized approach is often a robust and simple way of doing so.

Random approaches have at least three advantages over more traditional decisions. First, random often avoids strange corner-case behaviors that a more traditional algorithm may have trouble handling. For example, consider LRU page replacement (studied in more detail in a future chapter on virtual memory); while often a good replacement algorithm, LRU performs pessimistically for some cyclic-sequential workloads. Random, on the other hand, has no such worst case.

Second, random also is lightweight, requiring little state to track alternatives. In a traditional fair-share scheduling algorithm, tracking how much CPU each process has received requires per-process accounting, which must be updated after running each process. Doing so randomly necessitates only the most minimal of per-process state (e.g., the number of tickets each has).

Finally, random can be quite fast. As long as generating a random number is quick, making the decision is also, and thus random can be used in a number of places where speed is required. Of course, the faster the need, the more random tends towards pseudo-random.

A winning ticket, which is a number from 0 to 99. Assuming A holds tickets 0 through 74 and B 75 through 99, the winning ticket simply determines whether A or B runs. The scheduler then loads the state of that winning process and runs it.

Here is an example output of a lottery scheduler’s winning tickets:

63 85 70 39 76 17 29 41 36 39 10 99 68 83 63 62 43 0 49 49

Here is the resulting schedule:

A B A A B A A A A A A B A B A A A A A A

As you can see from the example, the use of randomness in lottery scheduling leads to a probabilistic correctness in meeting the desired proportion, but no guarantee. In our example above, B only gets to run 4 out of 20 time slices (20%), instead of the desired 25% allocation. However, the longer these two jobs compete, the more likely they are to achieve the desired percentages.
One of the most powerful (and basic) mechanisms in the design of lottery (and stride) scheduling is that of the ticket. The ticket is used to represent a process’s share of the CPU in these examples, but can be applied much more broadly. For example, in more recent work on virtual memory management for hypervisors, Waldspurger shows how tickets can be used to represent a guest operating system’s share of memory [W02]. Thus, if you are ever in need of a mechanism to represent a proportion of ownership, this concept just might be ... (wait for it) ... the ticket.

9.2 Ticket Mechanisms

Lottery scheduling also provides a number of mechanisms to manipulate tickets in different and sometimes useful ways. One way is with the concept of ticket currency. Currency allows a user with a set of tickets to allocate tickets among their own jobs in whatever currency they would like; the system then automatically converts said currency into the correct global value.

For example, assume users A and B have each been given 100 tickets. User A is running two jobs, A1 and A2, and gives them each 500 tickets (out of 1000 total) in User A’s own currency. User B is running only 1 job and gives it 10 tickets (out of 10 total). The system will convert A1’s and A2’s allocation from 500 each in A’s currency to 50 each in the global currency; similarly, B1’s 10 tickets will be converted to 100 tickets. The lottery will then be held over the global ticket currency (200 total) to determine which job runs.

User A -> 500 (A’s currency) to A1 -> 50 (global currency)
-> 500 (A’s currency) to A2 -> 50 (global currency)
User B -> 10 (B’s currency) to B1 -> 100 (global currency)

Another useful mechanism is ticket transfer. With transfers, a process can temporarily hand off its tickets to another process. This ability is especially useful in a client/server setting, where a client process sends a message to a server asking it to do some work on the client’s behalf. To speed up the work, the client can pass the tickets to the server and thus try to maximize the performance of the server while the server is handling the client’s request. When finished, the server then transfers the tickets back to the client and all is as before.

Finally, ticket inflation can sometimes be a useful technique. With inflation, a process can temporarily raise or lower the number of tickets it owns. Of course, in a competitive scenario with processes that do not trust one another, this makes little sense; one greedy process could give itself a vast number of tickets and take over the machine. Rather, inflation can be applied in an environment where a group of processes trust one another; in such a case, if any one process knows it needs more CPU time, it can boost its ticket value as a way to reflect that need to the system, all without communicating with any other processes.
// counter: used to track if we’ve found the winner yet
int counter = 0;

// winner: use some call to a random number generator to
// get a value, between 0 and the total # of tickets
int winner = getrandom(0, totaltickets);

// current: use this to walk through the list of jobs
node_t *current = head;

// loop until the sum of ticket values is > the winner
while (current) {
    counter = counter + current->tickets;
    if (counter > winner)
        break; // found the winner
    current = current->next;
}

// 'current’ is the winner: schedule it...

Figure 9.1: Lottery Scheduling Decision Code

9.3 Implementation

Probably the most amazing thing about lottery scheduling is the simplicity of its implementation. All you need is a good random number generator to pick the winning ticket, a data structure to track the processes of the system (e.g., a list), and the total number of tickets.

Let’s assume we keep the processes in a list. Here is an example comprised of three processes, A, B, and C, each with some number of tickets.

\[
\text{head} \rightarrow \text{Job:A Tix:100} \rightarrow \text{Job:B Tix:50} \rightarrow \text{Job:C Tix:250} \rightarrow \text{NULL}
\]

To make a scheduling decision, we first have to pick a random number (the winner) from the total number of tickets (400)\(^2\) Let’s say we pick the number 300. Then, we simply traverse the list, with a simple counter used to help us find the winner (Figure 9.1).

The code walks the list of processes, adding each ticket value to \(\text{counter}\) until the value exceeds \(\text{winner}\). Once that is the case, the current list element is the winner. With our example of the winning ticket being 300, the following takes place. First, \(\text{counter}\) is incremented to 100 to account for A’s tickets; because 100 is less than 300, the loop continues. Then \(\text{counter}\) would be updated to 150 (B’s tickets), still less than 300 and thus again we continue. Finally, \(\text{counter}\) is updated to 400 (clearly greater than 300), and thus we break out of the loop with \(\text{current}\) pointing at C (the winner).

To make this process most efficient, it might generally be best to organize the list in sorted order, from the highest number of tickets to the

\(^2\)Surprisingly, as pointed out by Björn Lindberg, this can be challenging to do correctly; for more details, see http://stackoverflow.com/questions/2509679/how-to-generate-a-random-number-from-within-a-range.
lowest. The ordering does not affect the correctness of the algorithm; however, it does ensure in general that the fewest number of list iterations are taken, especially if there are a few processes that possess most of the tickets.

9.4 An Example

To make the dynamics of lottery scheduling more understandable, we now perform a brief study of the completion time of two jobs competing against one another, each with the same number of tickets (100) and same run time ($R$, which we will vary).

In this scenario, we’d like for each job to finish at roughly the same time, but due to the randomness of lottery scheduling, sometimes one job finishes before the other. To quantify this difference, we define a simple unfairness metric, $U$ which is simply the time the first job completes divided by the time that the second job completes. For example, if $R = 10$, and the first job finishes at time 10 (and the second job at 20), $U = \frac{10}{20} = 0.5$. When both jobs finish at nearly the same time, $U$ will be quite close to 1. In this scenario, that is our goal: a perfectly fair scheduler would achieve $U = 1$.

Figure 9.2 plots the average unfairness as the length of the two jobs ($R$) is varied from 1 to 1000 over thirty trials (results are generated via the simulator provided at the end of the chapter). As you can see from the graph, when the job length is not very long, average unfairness can be quite severe. Only as the jobs run for a significant number of time slices does the lottery scheduler approach the desired outcome.
9.5 How To Assign Tickets?

One problem we have not addressed with lottery scheduling is: how to assign tickets to jobs? This problem is a tough one, because of course how the system behaves is strongly dependent on how tickets are allocated. One approach is to assume that the users know best; in such a case, each user is handed some number of tickets, and a user can allocate tickets to any jobs they run as desired. However, this solution is a non-solution: it really doesn’t tell you what to do. Thus, given a set of jobs, the “ticket-assignment problem” remains open.

9.6 Why Not Deterministic?

You might also be wondering: why use randomness at all? As we saw above, while randomness gets us a simple (and approximately correct) scheduler, it occasionally will not deliver the exact right proportions, especially over short time scales. For this reason, Waldspurger invented **stride scheduling**, a deterministic fair-share scheduler [W95].

Stride scheduling is also straightforward. Each job in the system has a stride, which is inverse in proportion to the number of tickets it has. In our example above, with jobs A, B, and C, with 100, 50, and 250 tickets, respectively, we can compute the stride of each by dividing some large number by the number of tickets each process has been assigned. For example, if we divide 10,000 by each of those ticket values, we obtain the following stride values for A, B, and C: 100, 200, and 40. We call this value the stride of each process; every time a process runs, we will increment a counter for it (called its pass value) by its stride to track its global progress.

The scheduler then uses the stride and pass to determine which process should run next. The basic idea is simple: at any given time, pick the process to run that has the lowest pass value so far; when you run a process, increment its pass counter by its stride. A pseudocode implementation is provided by Waldspurger [W95]:

```c
current = remove_min(queue); // pick client with minimum pass
schedule(current); // use resource for quantum
current->pass += current->stride; // compute next pass using stride
insert(queue, current); // put back into the queue
```

In our example, we start with three processes (A, B, and C), with stride values of 100, 200, and 40, and all with pass values initially at 0. Thus, at first, any of the processes might run, as their pass values are equally low. Assume we pick A (arbitrarily; any of the processes with equal low pass values can be chosen). A runs; when finished with the time slice, we update its pass value to 100. Then we run B, whose pass value is then set to 200. Finally, we run C, whose pass value is incremented to 40. At this point, the algorithm will pick the lowest pass value, which is C’s, and run it, updating its pass to 80 (C’s stride is 40, as you recall). Then C will
run again (still the lowest pass value), raising its pass to 120. A will run now, updating its pass to 200 (now equal to B’s). Then C will run twice more, updating its pass to 160 then 200. At this point, all pass values are equal again, and the process will repeat, ad infinitum. Table 9.1 traces the behavior of the scheduler over time.

As we can see from the table, C ran five times, A twice, and B just once, exactly in proportion to their ticket values of 250, 100, and 50. Lottery scheduling achieves the proportions probabilistically over time; stride scheduling gets them exactly right at the end of each scheduling cycle.

So you might be wondering: given the precision of stride scheduling, why use lottery scheduling at all? Well, lottery scheduling has one nice property that stride scheduling does not: no global state. Imagine a new job enters in the middle of our stride scheduling example above; what should its pass value be? Should it be set to 0? If so, it will monopolize the CPU. With lottery scheduling, there is no global state per process; we simply add a new process with whatever tickets it has, update the single global variable to track how many total tickets we have, and go from there. In this way, lottery makes it much easier to incorporate new processes in a sensible manner.

### Table 9.1: Stride Scheduling: A Trace

<table>
<thead>
<tr>
<th>Pass(A) (stride=100)</th>
<th>Pass(B) (stride=200)</th>
<th>Pass(C) (stride=40)</th>
<th>Who Runs?</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
<td>A</td>
</tr>
<tr>
<td>100</td>
<td>0</td>
<td>0</td>
<td>B</td>
</tr>
<tr>
<td>100</td>
<td>200</td>
<td>0</td>
<td>C</td>
</tr>
<tr>
<td>100</td>
<td>200</td>
<td>40</td>
<td>C</td>
</tr>
<tr>
<td>100</td>
<td>200</td>
<td>80</td>
<td>C</td>
</tr>
<tr>
<td>100</td>
<td>200</td>
<td>120</td>
<td>A</td>
</tr>
<tr>
<td>200</td>
<td>200</td>
<td>120</td>
<td>C</td>
</tr>
<tr>
<td>200</td>
<td>200</td>
<td>160</td>
<td>C</td>
</tr>
<tr>
<td>200</td>
<td>200</td>
<td>200</td>
<td>...</td>
</tr>
</tbody>
</table>

9.7 Summary

We have introduced the concept of proportional-share scheduling and briefly discussed two implementations: lottery and stride scheduling. Lottery uses randomness in a clever way to achieve proportional share; stride does so deterministically. Although both are conceptually interesting, they have not achieved wide-spread adoption as CPU schedulers for a variety of reasons. One is that such approaches do not particularly mesh well with I/O [AC97]; another is that they leave open the hard problem of ticket assignment, i.e., how do you know how many tickets your browser should be allocated? General-purpose schedulers (such as the MLFQ we discussed previously, and other similar Linux schedulers) do so more gracefully and thus are more widely deployed.
As a result, proportional-share schedulers are more useful in domains where some of these problems (such as assignment of shares) are relatively easy to solve. For example, in a virtualized data center, where you might like to assign one-quarter of your CPU cycles to the Windows VM and the rest to your base Linux installation, proportional sharing can be simple and effective. See Waldspurger [W02] for further details on how such a scheme is used to proportionally share memory in VMWare’s ESX Server.
References

[AC97] “Extending Proportional-Share Scheduling to a Network of Workstations”
Andrea C. Arpaci-Dusseau and David E. Culler
PDP TA’97, June 1997
A paper by one of the authors on how to extend proportional-share scheduling to work better in a clustered environment.

[D82] “Why Numbering Should Start At Zero”
Edsger Dijkstra, August 1982
http://www.cs.utexas.edu/users/EWD/EWD831.PDF
A short note from E. Dijkstra, one of the pioneers of computer science. We’ll be hearing much more on this guy in the section on Concurrency. In the meanwhile, enjoy this note, which includes this motivating quote: “One of my colleagues – not a computing scientist – accused a number of younger computing scientists of ‘pedantry’ because they started numbering at zero.” The note explains why doing so is logical.

[KL88] “A Fair Share Scheduler”
J. Kay and P. Lauder
CACM, Volume 31 Issue 1, January 1988
An early reference to a fair-share scheduler.

Carl A. Waldspurger and William E. Weihl
OSDI ’94, November 1994
The landmark paper on lottery scheduling that got the systems community re-energized about scheduling, fair sharing, and the power of simple randomized algorithms.

Carl A. Waldspurger
Ph.D. Thesis, MIT, 1995
The award-winning thesis of Waldspurger’s that outlines lottery and stride scheduling. If you’re thinking of writing a Ph.D. dissertation at some point, you should always have a good example around, to give you something to strive for: this is such a good one.

[W02] “Memory Resource Management in VMware ESX Server”
Carl A. Waldspurger
OSDI ’02, Boston, Massachusetts
The paper to read about memory management in VMMs (a.k.a., hypervisors). In addition to being relatively easy to read, the paper contains numerous cool ideas about this new type of VMM-level memory management.
Homework

This program, `lottery.py`, allows you to see how a lottery scheduler works. See the README for details.

Questions

1. Compute the solutions for simulations with 3 jobs and random seeds of 1, 2, and 3.
2. Now run with two specific jobs: each of length 10, but one (job 0) with just 1 ticket and the other (job 1) with 100 (e.g., `-l 10:1,10:100`). What happens when the number of tickets is so imbalanced? Will job 0 ever run before job 1 completes? How often? In general, what does such a ticket imbalance do to the behavior of lottery scheduling?
3. When running with two jobs of length 100 and equal ticket allocations of 100 (`-l 100:100,100:100`), how unfair is the scheduler? Run with some different random seeds to determine the (probabilistic) answer; let unfairness be determined by how much earlier one job finishes than the other.
4. How does your answer to the previous question change as the quantum size (`-q`) gets larger?
5. Can you make a version of the graph that is found in the chapter? What else would be worth exploring? How would the graph look with a stride scheduler?
This chapter will introduce the basics of **multiprocessor scheduling**. As this topic is relatively advanced, it may be best to cover it *after* you have studied the topic of concurrency in some detail (i.e., the second major “easy piece” of the book).

After years of existence only in the high-end of the computing spectrum, **multiprocessor** systems are increasingly commonplace, and have found their way into desktop machines, laptops, and even mobile devices. The rise of the **multicore** processor, in which multiple CPU cores are packed onto a single chip, is the source of this proliferation; these chips have become popular as computer architects have had a difficult time making a single CPU much faster without using (way) too much power. And thus we all now have a few CPUs available to us, which is a good thing, right?

Of course, there are many difficulties that arise with the arrival of more than a single CPU. A primary one is that a typical application (i.e., some C program you wrote) only uses a single CPU; adding more CPUs does not make that single application run faster. To remedy this problem, you’ll have to rewrite your application to run in **parallel**, perhaps using **threads** (as discussed in great detail in the second piece of this book). Multithreaded applications can spread work across multiple CPUs and thus run faster when given more CPU resources.

**Aside: Advanced Chapters**

Advanced chapters require material from a broad swath of the book to truly understand, while logically fitting into a section that is earlier than said set of prerequisite materials. For example, this chapter on multiprocessor scheduling makes much more sense if you’ve first read the middle piece on concurrency; however, it logically fits into the part of the book on virtualization (generally) and CPU scheduling (specifically). Thus, it is recommended such chapters be covered out of order; in this case, after the second piece of the book.
Beyond applications, a new problem that arises for the operating system is (not surprisingly!) that of multiprocessor scheduling. Thus far we’ve discussed a number of principles behind single-processor scheduling; how can we extend those ideas to work on multiple CPUs? What new problems must we overcome? And thus, our problem:

**Crux: How to Schedule Jobs on Multiple CPUs**
How should the OS schedule jobs on multiple CPUs? What new problems arise? Do the same old techniques work, or are new ideas required?

### 10.1 Background: Multiprocessor Architecture

To understand the new issues surrounding multiprocessor scheduling, we have to understand a new and fundamental difference between single-CPU hardware and multi-CPU hardware. This difference centers around the use of hardware caches (e.g., Figure 10.1), and exactly how data is shared across multiple processors. We now discuss this issue further, at a high level. Details are available elsewhere [CSG99], in particular in an upper-level or perhaps graduate computer architecture course.

In a system with a single CPU, there are a hierarchy of **hardware caches** that in general help the processor run programs faster. Caches are small, fast memories that (in general) hold copies of popular data that is found in the main memory of the system. Main memory, in contrast, holds all of the data, but access to this larger memory is slower. By keeping frequently accessed data in a cache, the system can make the large, slow memory appear to be a fast one.

As an example, consider a program that issues an explicit load instruction to fetch a value from memory, and a simple system with only a single CPU; the CPU has a small cache (say 64 KB) and a large main memory.
The first time a program issues this load, the data resides in main memory, and thus takes a long time to fetch (perhaps in the tens of nanoseconds, or even hundreds). The processor, anticipating that the data may be reused, puts a copy of the loaded data into the CPU cache. If the program later fetches this same data item again, the CPU first checks for it in the cache; because it finds it there, the data is fetched much more quickly (say, just a few nanoseconds), and thus the program runs faster.

Caches are thus based on the notion of locality, of which there are two kinds: temporal locality and spatial locality. The idea behind temporal locality is that when a piece of data is accessed, it is likely to be accessed again in the near future; imagine variables or even instructions themselves being accessed over and over again in a loop. The idea behind spatial locality is that if a program accesses a data item at address $x$, it is likely to access data items near $x$ as well; here, think of a program streaming through an array, or instructions being executed one after the other. Because locality of these types exist in many programs, hardware systems can make good guesses about which data to put in a cache and thus work well.

Now for the tricky part: what happens when you have multiple processors in a single system, with a single shared main memory, as we see in Figure 10.2?

As it turns out, caching with multiple CPUs is much more complicated. Imagine, for example, that a program running on CPU 1 reads a data item (with value $D$) at address $A$; because the data is not in the cache on CPU 1, the system fetches it from main memory, and gets the value $D$. The program then modifies the value at address $A$, just updating its cache with the new value $D'$; writing the data through all the way to main memory is slow, so the system will (usually) do that later. Then assume the OS decides to stop running the program and move it to CPU 2. The program then re-reads the value at address $A$; there is no such data
CPU 2’s cache, and thus the system fetches the value from main memory, and gets the old value \( D \) instead of the correct value \( D’ \). Oops!

This general problem is called the problem of cache coherence, and there is a vast research literature that describes many different subtleties involved with solving the problem [SHW11]. Here, we will skip all of the nuance and make some major points; take a computer architecture class (or three) to learn more.

The basic solution is provided by the hardware: by monitoring memory accesses, hardware can ensure that basically the “right thing” happens and that the view of a single shared memory is preserved. One way to do this on a bus-based system (as described above) is to use an old technique known as bus snooping [G83]; each cache pays attention to memory updates by observing the bus that connects them to main memory. When a CPU then sees an update for a data item it holds in its cache, it will notice the change and either invalidate its copy (i.e., remove it from its own cache) or update it (i.e., put the new value into its cache too). Write-back caches, as hinted at above, make this more complicated (because the write to main memory isn’t visible until later), but you can imagine how the basic scheme might work.

10.2 Don’t Forget Synchronization

Given that the caches do all of this work to provide coherence, do programs (or the OS itself) have to worry about anything when they access shared data? The answer, unfortunately, is yes, and is documented in great detail in the second piece of this book on the topic of concurrency. While we won’t get into the details here, we’ll sketch/review some of the basic ideas here (assuming you’re familiar with concurrency).

When accessing (and in particular, updating) shared data items or structures across CPUs, mutual exclusion primitives (such as locks) should likely be used to guarantee correctness (other approaches, such as building lock-free data structures, are complex and only used on occasion; see the chapter on deadlock in the piece on concurrency for details). For example, assume we have a shared queue being accessed on multiple CPUs concurrently. Without locks, adding or removing elements from the queue concurrently will not work as expected, even with the underlying coherence protocols; one needs locks to atomically update the data structure to its new state.

To make this more concrete, imagine this code sequence, which is used to remove an element from a shared linked list, as we see in Figure 10.3. Imagine if threads on two CPUs enter this routine at the same time. If Thread 1 executes the first line, it will have the current value of head stored in its tmp variable; if Thread 2 then executes the first line as well, it also will have the same value of head stored in its own private tmp variable (tmp is allocated on the stack, and thus each thread will have its own private storage for it). Thus, instead of each thread removing an element from the head of the list, each thread will try to remove the
typedef struct __Node_t {
    int value;
    struct __Node_t *next;
} Node_t;

int List_Pop() {
    Node_t *tmp = head; // remember old head ...
    int value = head->value; // ... and its value
    head = head->next; // advance head to next pointer
    free(tmp); // free old head
    return value; // return value at head
}

Figure 10.3: Simple List Delete Code

same head element, leading to all sorts of problems (such as an attempted
double free of the head element at line 4, as well as potentially returning
the same data value twice).

The solution, of course, is to make such routines correct via locking. In this case, allocating a simple mutex (e.g., pthread_mutex_t m;) and then adding a lock(&m) at the beginning of the routine and
an unlock(&m) at the end will solve the problem, ensuring that the code
will execute as desired. Unfortunately, as we will see, such an approach is
not without problems, in particular with regards to performance. Specifi-
cally, as the number of CPUs grows, access to a synchronized shared data
structure becomes quite slow.

10.3 One Final Issue: Cache Affinity

One final issue arises in building a multiprocessor cache scheduler,
known as cache affinity. This notion is simple: a process, when run on a
particular CPU, builds up a fair bit of state in the caches (and TLBs) of the
CPU. The next time the process runs, it is often advantageous to run it on
the same CPU, as it will run faster if some of its state is already present in
the caches on that CPU. If, instead, one runs a process on a different CPU
each time, the performance of the process will be worse, as it will have to
reload the state each time it runs (note it will run correctly on a different
CPU thanks to the cache coherence protocols of the hardware). Thus, a
multiprocessor scheduler should consider cache affinity when making its
scheduling decisions, perhaps preferring to keep a process on the same
CPU if at all possible.

10.4 Single-Queue Scheduling

With this background in place, we now discuss how to build a sched-
uler for a multiprocessor system. The most basic approach is to simply
reuse the basic framework for single processor scheduling, by putting all
jobs that need to be scheduled into a single queue; we call this single-
queue multiprocessor scheduling or SQMS for short. This approach
has the advantage of simplicity; it does not require much work to take an
existing policy that picks the best job to run next and adapt it to work on
more than one CPU (where it might pick the best two jobs to run, if there
are two CPUs, for example).
However, SQMS has obvious shortcomings. The first problem is a lack of **scalability**. To ensure the scheduler works correctly on multiple CPUs, the developers will have inserted some form of **locking** into the code, as described above. Locks ensure that when SQMS code accesses the single queue (say, to find the next job to run), the proper outcome arises.

Locks, unfortunately, can greatly reduce performance, particularly as the number of CPUs in the systems grows [A91]. As contention for such a single lock increases, the system spends more and more time in lock overhead and less time doing the work the system should be doing (note: it would be great to include a real measurement of this in here someday).

The second main problem with SQMS is cache affinity. For example, let us assume we have five jobs to run \((A, B, C, D, E)\) and four processors. Our scheduling queue thus looks like this:

```
Queue: A -> B -> C -> D -> E -> NULL
```

Over time, assuming each job runs for a time slice and then another job is chosen, here is a possible job schedule across CPUs:

```
CPU 0: A -> E -> D -> C -> B ... (repeat) ...
CPU 1: B -> A -> E -> D -> C ... (repeat) ...
CPU 2: C -> B -> A -> E -> D ... (repeat) ...
CPU 3: D -> C -> B -> A -> E ... (repeat) ...
```

Because each CPU simply picks the next job to run from the globally-shared queue, each job ends up bouncing around from CPU to CPU, thus doing exactly the opposite of what would make sense from the standpoint of cache affinity.

To handle this problem, most SQMS schedulers include some kind of affinity mechanism to try to make it more likely that process will continue to run on the same CPU if possible. Specifically, one might provide affinity for some jobs, but move others around to balance load. For example, imagine the same five jobs scheduled as follows:

```
CPU 0: A -> E -> A -> A ... (repeat) ...
CPU 1: B -> B -> E -> B -> B ... (repeat) ...
CPU 2: C -> C -> C -> E -> C ... (repeat) ...
CPU 3: D -> D -> D -> D -> E ... (repeat) ...
```
In this arrangement, jobs $A$ through $D$ are not moved across processors, with only job $E$ migrating from CPU to CPU, thus preserving affinity for most. You could then decide to migrate a different job the next time through, thus achieving some kind of affinity fairness as well. Implementing such a scheme, however, can be complex.

Thus, we can see the SQMS approach has its strengths and weaknesses. It is straightforward to implement given an existing single-CPU scheduler, which by definition has only a single queue. However, it does not scale well (due to synchronization overheads), and it does not readily preserve cache affinity.

### 10.5 Multi-Queue Scheduling

Because of the problems caused in single-queue schedulers, some systems opt for multiple queues, e.g., one per CPU. We call this approach multi-queue multiprocessor scheduling (or MQMS).

In MQMS, our basic scheduling framework consists of multiple scheduling queues. Each queue will likely follow a particular scheduling discipline, such as round robin, though of course any algorithm can be used. When a job enters the system, it is placed on exactly one scheduling queue, according to some heuristic (e.g., random, or picking one with fewer jobs than others). Then it is scheduled essentially independently, thus avoiding the problems of information sharing and synchronization found in the single-queue approach.

For example, assume we have a system where there are just two CPUs (labeled CPU 0 and CPU 1), and some number of jobs enter the system: $A$, $B$, $C$, and $D$ for example. Given that each CPU has a scheduling queue now, the OS has to decide into which queue to place each job. It might do something like this:

```
Q0 → A → C
Q1 → B → D
```

Depending on the queue scheduling policy, each CPU now has two jobs to choose from when deciding what should run. For example, with round robin, the system might produce a schedule that looks like this:

<table>
<thead>
<tr>
<th>CPU 0</th>
<th>A</th>
<th>A</th>
<th>C</th>
<th>C</th>
<th>A</th>
<th>A</th>
<th>C</th>
<th>C</th>
<th>A</th>
<th>A</th>
<th>C</th>
<th>C</th>
</tr>
</thead>
<tbody>
<tr>
<td>CPU 1</td>
<td>B</td>
<td>B</td>
<td>D</td>
<td>D</td>
<td>B</td>
<td>B</td>
<td>D</td>
<td>D</td>
<td>B</td>
<td>B</td>
<td>D</td>
<td>D</td>
</tr>
</tbody>
</table>

MQMS has a distinct advantage of SQMS in that it should be inherently more scalable. As the number of CPUs grows, so too does the number of queues, and thus lock and cache contention should not become a central problem. In addition, MQMS intrinsically provides cache affinity;
jobs stay on the same CPU and thus reap the advantage of reusing cached contents therein.

But, if you’ve been paying attention, you might see that we have a new problem, which is fundamental in the multi-queue based approach: **load imbalance**. Let’s assume we have the same set up as above (four jobs, two CPUs), but then one of the jobs (say *C*) finishes. We now have the following scheduling queues:

$$
\begin{array}{c}
Q0 & \rightarrow & A \\
Q1 & \rightarrow & B \\
\end{array}
$$

If we then run our round-robin policy on each queue of the system, we will see this resulting schedule:

<table>
<thead>
<tr>
<th>CPU 0</th>
<th>A</th>
<th>A</th>
<th>A</th>
<th>A</th>
<th>A</th>
<th>A</th>
<th>A</th>
<th>A</th>
<th>A</th>
<th>A</th>
<th>...</th>
</tr>
</thead>
<tbody>
<tr>
<td>CPU 1</td>
<td>B</td>
<td>B</td>
<td>D</td>
<td>D</td>
<td>B</td>
<td>B</td>
<td>D</td>
<td>B</td>
<td>B</td>
<td>D</td>
<td>D</td>
</tr>
</tbody>
</table>

As you can see from this diagram, *A* gets twice as much CPU as *B* and *D*, which is not the desired outcome. Even worse, let’s imagine that both *A* and *C* finish, leaving just jobs *B* and *D* in the system. The scheduling queues will look like this:

$$
\begin{array}{c}
Q0 & \rightarrow & \ \\
Q1 & \rightarrow & B \\
\end{array}
$$

As a result, CPU 0 will be left idle! *(insert dramatic and sinister music here)*

And hence our CPU usage timeline looks sad:

<table>
<thead>
<tr>
<th>CPU 0</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>CPU 1</td>
<td>B</td>
</tr>
</tbody>
</table>

So what should a poor multi-queue multiprocessor scheduler do? How can we overcome the insidious problem of load imbalance and defeat the evil forces of ... the Decepticons\(^1\)? How do we stop asking questions that are hardly relevant to this otherwise wonderful book?

---

\(^1\)Little known fact is that the home planet of Cybertron was destroyed by bad CPU scheduling decisions. And now let that be the first and last reference to Transformers in this book, for which we sincerely apologize.
The obvious answer to this query is to move jobs around, a technique which we (once again) refer to as migration. By migrating a job from one CPU to another, true load balance can be achieved.

Let’s look at a couple of examples to add some clarity. Once again, we have a situation where one CPU is idle and the other has some jobs.

In this case, the desired migration is easy to understand: the OS should simply move one of B or D to CPU 0. The result of this single job migration is evenly balanced load and everyone is happy.

A more tricky case arises in our earlier example, where A was left alone on CPU 0 and B and D were alternating on CPU 1:

In this case, a single migration does not solve the problem. What would you do in this case? The answer, alas, is continuous migration of one or more jobs. One possible solution is to keep switching jobs, as we see in the following timeline. In the figure, first A is alone on CPU 0, and B and D alternate on CPU 1. After a few time slices, B is moved to compete with A on CPU 0, while D enjoys a few time slices alone on CPU 1. And thus load is balanced:

Of course, many other possible migration patterns exist. But now for the tricky part: how should the system decide to enact such a migration?

One basic approach is to use a technique known as work stealing [FLR98]. With a work-stealing approach, a (source) queue that is low on jobs will occasionally peek at another (target) queue, to see how full it is. If the target queue is (notably) more full than the source queue, the source will “steal” one or more jobs from the target to help balance load.

Of course, there is a natural tension in such an approach. If you look around at other queues too often, you will suffer from high overhead and have trouble scaling, which was the entire purpose of implementing the
multiple queue scheduling in the first place! If, on the other hand, you don’t look at other queues very often, you are in danger of suffering from severe load balances. Finding the right threshold remains, as is common in system policy design, a black art.

10.6 Linux Multiprocessor Schedulers

Interestingly, in the Linux community, no common solution has approached to building a multiprocessor scheduler. Over time, three different schedulers arose: the O(1) scheduler, the Completely Fair Scheduler (CFS), and the BF Scheduler (BFS). See Meehean’s dissertation for an excellent overview of the strengths and weaknesses of said schedulers [M11]; here we just summarize a few of the basics.

Both O(1) and CFS uses multiple queues, whereas BFS uses a single queue, showing that both approaches can be successful. Of course, there are many other details which separate these schedulers. For example, the O(1) scheduler is a priority-based scheduler (similar to the MLFQ discussed before), changing a process’s priority over time and then scheduling those with highest priority in order to meet various scheduling objectives; interactivity is a particular focus. CFS, in contrast, is a deterministic proportional-share approach (more like Stride scheduling, as discussed earlier). BFS, the only single-queue approach among the three, is also proportional-share, but based on a more complicated scheme known as Earliest Eligible Virtual Deadline First (EEVDF) [SA96]. Read more about these modern algorithms on your own; you should be able to understand how they work now!

10.7 Summary

We have seen various approaches to multiprocessor scheduling. The single-queue approach (SQMS) is rather straightforward to build and balances load well but inherently has difficulty with scaling to many processors and cache affinity. The multiple-queue approach (MQMS) scales better and handles cache affinity well, but has trouble with load imbalance and is more complicated. Whichever approach you take, there is no simple answer: building a general purpose scheduler remains a daunting task, as small code changes can lead to large behavioral differences. Only undertake such an exercise if you know exactly what you are doing, or, at least, are getting paid a large amount of money to do so.

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2Look up what BF stands for on your own; be forewarned, it is not for the faint of heart.
References

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Thomas E. Anderson
IEEE TPDS Volume 1:1, January 1990
A classic paper on how different locking alternatives do and don’t scale. By Tom Anderson, very well
known researcher in both systems and networking. And author of a very fine OS textbook, we must say.

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Matteo Frigo, Charles E. Leiserson, Keith Randall
PLDI ’98, Montreal, Canada, June 1998
Cilk is a lightweight language and runtime for writing parallel programs, and an excellent example of
the work-stealing paradigm.

[G83] “Using Cache Memory To Reduce Processor-Memory Traffic”
James R. Goodman
ISCA ’83, Stockholm, Sweden, June 1983
The pioneering paper on how to use bus snooping, i.e., paying attention to requests you see on the bus, to
build a cache coherence protocol. Goodman’s research over many years at Wisconsin is full of cleverness,
this being but one example.

[M11] “Towards Transparent CPU Scheduling”
Joseph T. Meehean
Doctoral Dissertation at University of Wisconsin–Madison, 2011
A dissertation that covers a lot of the details of how modern Linux multiprocessor scheduling works.
Pretty awesome! But, as co-advisors of Joe’s, we may be a bit biased here.

[SHW11] “A Primer on Memory Consistency and Cache Coherence”
Daniel J. Sorin, Mark D. Hill, and David A. Wood
Synthesis Lectures in Computer Architecture
A definitive overview of memory consistency and multiprocessor caching. Required reading for anyone
who likes to know way too much about a given topic.

[SA96] “Earliest Eligible Virtual Deadline First: A Flexible and Accurate Mechanism for Pro-
portional Share Resource Allocation”
Ion Stoica and Hussein Abdel-Wahab
Technical Report TR-95-22, Old Dominion University, 1996
A tech report on this cool scheduling idea, from Ion Stoica, now a professor at U.C. Berkeley and world
expert in networking, distributed systems, and many other things.
Summary Dialogue on CPU Virtualization

Professor: So, Student, did you learn anything?

Student: Well, Professor, that seems like a loaded question. I think you only want me to say “yes.”

Professor: That’s true. But it’s also still an honest question. Come on, give a professor a break, will you?

Student: OK, OK. I think I did learn a few things. First, I learned a little about how the OS virtualizes the CPU. There are a bunch of important mechanisms that I had to understand to make sense of this: traps and trap handlers, timer interrupts, and how the OS and the hardware have to carefully save and restore state when switching between processes.

Professor: Good, good!

Student: All those interactions do seem a little complicated though; how can I learn more?

Professor: Well, that’s a good question. I think there is no substitute for doing; just reading about these things doesn’t quite give you the proper sense. Do the class projects and I bet by the end it will all kind of make sense.

Student: Sounds good. What else can I tell you?

Professor: Well, did you get some sense of the philosophy of the OS in your quest to understand its basic machinery?

Student: Hmm... I think so. It seems like the OS is fairly paranoid. It wants to make sure it stays in charge of the machine. While it wants a program to run as efficiently as possible (and hence the whole reasoning behind limited direct execution), the OS also wants to be able to say “Ah! Not so fast my friend” in case of an errant or malicious process. Paranoia rules the day, and certainly keeps the OS in charge of the machine. Perhaps that is why we think of the OS as a resource manager.

Professor: Yes indeed – sounds like you are starting to put it together! Nice.

Student: Thanks.
Professor: And what about the policies on top of those mechanisms – any interesting lessons there?
Student: Some lessons to be learned there for sure. Perhaps a little obvious, but obvious can be good. Like the notion of bumping short jobs to the front of the queue – I knew that was a good idea ever since the one time I was buying some gum at the store, and the guy in front of me had a credit card that wouldn’t work. He was no short job, let me tell you.
Professor: That sounds oddly rude to that poor fellow. What else?
Student: Well, that you can build a smart scheduler that tries to be like SJF and RR all at once – that MLFQ was pretty neat. Building up a real scheduler seems difficult.
Professor: Indeed it is. That’s why there is still controversy to this day over which scheduler to use; see the Linux battles between CFS, BFS, and the O(1) scheduler, for example. And no, I will not spell out the full name of BFS.
Student: And I won’t ask you to! These policy battles seem like they could rage forever; is there really a right answer?
Professor: Probably not. After all, even our own metrics are at odds: if your scheduler is good at turnaround time, it’s bad at response time, and vice versa. As Lampson said, perhaps the goal isn’t to find the best solution, but rather to avoid disaster.
Student: That’s a little depressing.
Professor: Good engineering can be that way. And it can also be uplifting! It’s just your perspective on it, really. I personally think being pragmatic is a good thing, and pragmatists realize that not all problems have clean and easy solutions. Anything else that caught your fancy?
Student: I really liked the notion of gaming the scheduler; it seems like that might be something to look into when I’m next running a job on Amazon’s EC2 service. Maybe I can steal some cycles from some other unsuspecting (and more importantly, OS-ignorant) customer!
Professor: It looks like I might have created a monster! Professor Frankenstein is not what I’d like to be called, you know.
Student: But isn’t that the idea? To get us excited about something, so much so that we look into it on our own? Lighting fires and all that?
Professor: I guess so. But I didn’t think it would work!
A Dialogue on Memory Virtualization

**Student:** So, are we done with virtualization?

**Professor:** No!

**Student:** Hey, no reason to get so excited; I was just asking a question. Students are supposed to do that, right?

**Professor:** Well, professors do always say that, but really they mean this: ask questions, if they are good questions, and you have actually put a little thought into them.

**Student:** Well, that sure takes the wind out of my sails.

**Professor:** Mission accomplished. In any case, we are not nearly done with virtualization! Rather, you have just seen how to virtualize the CPU, but really there is a big monster waiting in the closet: memory. Virtualizing memory is complicated and requires us to understand many more intricate details about how the hardware and OS interact.

**Student:** That sounds cool. Why is it so hard?

**Professor:** Well, there are a lot of details, and you have to keep them straight in your head to really develop a mental model of what is going on. We’ll start simple, with very basic techniques like base/bounds, and slowly add complexity to tackle new challenges, including fun topics like TLBs and multi-level page tables. Eventually, we’ll be able to describe the workings of a fully-functional modern virtual memory manager.

**Student:** Neat! Any tips for the poor student, inundated with all of this information and generally sleep-deprived?

**Professor:** For the sleep deprivation, that’s easy: sleep more (and party less). For understanding virtual memory, start with this: every address generated by a user program is a virtual address. The OS is just providing an illusion to each process, specifically that it has its own large and private memory; with some hardware help, the OS will turn these pretend virtual addresses into real physical addresses, and thus be able to locate the desired information.
Student: OK, I think I can remember that... (to self) every address from a user program is virtual, every address from a user program is virtual, every ...

Professor: What are you mumbling about?

Student: Oh nothing.... (awkward pause) ... Anyway, why does the OS want to provide this illusion again?

Professor: Mostly ease of use: the OS will give each program the view that it has a large contiguous address space to put its code and data into; thus, as a programmer, you never have to worry about things like “where should I store this variable?” because the virtual address space of the program is large and has lots of room for that sort of thing. Life, for a programmer, becomes much more tricky if you have to worry about fitting all of your code data into a small, crowded memory.

Student: Why else?

Professor: Well, isolation and protection are big deals, too. We don’t want one errant program to be able to read, or worse, overwrite, some other program’s memory, do we?

Student: Probably not. Unless it’s a program written by someone you don’t like.

Professor: Hmmm.... I think we might need to add a class on morals and ethics to your schedule for next semester. Perhaps OS class isn’t getting the right message across.

Student: Maybe we should. But remember, it’s not me who taught us that the proper OS response to errant process behavior is to kill the offending process!
The Abstraction: Address Spaces

In the early days, building computer systems was easy. Why, you ask? Because users didn’t expect much. It is those darned users with their expectations of “ease of use”, “high performance”, “reliability”, etc., that really have led to all these headaches. Next time you meet one of those computer users, thank them for all the problems they have caused.

13.1 Early Systems

From the perspective of memory, early machines didn’t provide much of an abstraction to users. Basically, the physical memory of the machine looked something like what you see in Figure 13.1.

The OS was a set of routines (a library, really) that sat in memory (starting at physical address 0 in this example), and there would be one running program (a process) that currently sat in physical memory (starting at physical address 64k in this example) and used the rest of memory. There were few illusions here, and the user didn’t expect much from the OS. Life was sure easy for OS developers in those days, wasn’t it?

![Figure 13.1: Operating Systems: The Early Days](image-url)
13.2 Multiprogramming and Time Sharing

After a time, because machines were expensive, people began to share machines more effectively. Thus the era of multiprogramming was born [DV66], in which multiple processes were ready to run at a given time, and the OS would switch between them, for example when one decided to perform an I/O. Doing so increased the effective utilization of the CPU. Such increases in efficiency were particularly important in those days where each machine cost hundreds of thousands or even millions of dollars (and you thought your Mac was expensive!).

Soon enough, however, people began demanding more of machines, and the era of time sharing was born [S59, L60, M62, M83]. Specifically, many realized the limitations of batch computing, particularly on programmers themselves [CV65], who were tired of long (and hence ineffective) program-debug cycles. The notion of interactivity became important, as many users might be concurrently using a machine, each waiting for (or hoping for) a timely response from their currently-executing tasks.

One way to implement time sharing would be to run one process for a short while, giving it full access to all memory (as in Figure 13.1), then stop it, save all of its state to some kind of disk (including all of physical memory), load some other process’s state, run it for a while, and thus implement some kind of crude sharing of the machine [M+63]. Unfortunately, this approach has a big problem: it is way too slow, particularly as memory grew. While saving and restoring register-level state (e.g., the PC, general-purpose registers, etc.) is relatively fast, saving the entire contents of memory to disk is brutally non-performant. Thus, what we’d rather do is leave processes in memory while switching between them, allowing the OS to implement time sharing efficiently (Figure 13.2).
THE ABSTRACTION: ADDRESS SPACES

13.3 The Address Space

However, we have to keep those pesky users in mind, and doing so requires the OS to create an easy to use abstraction of physical memory. We call this abstraction the address space, and it is the running program’s view of memory in the system. Understanding this fundamental OS abstraction of memory is key to understanding how memory is virtualized.

The address space of a process contains all of the memory state of the running program. For example, the code of the program (the instructions) have to live in memory somewhere, and thus they are in the address space. The program, while it is running, uses a stack to keep track of where it is in the function call chain as well as to allocate local variables and pass parameters and return values to and from routines. Finally, the heap is used for dynamically-allocated, user-managed memory, such as that you might receive from a call to malloc() in C or new in an object-oriented language such as C++ or Java. Of course, there are other things in there too (e.g., statically-initialized variables), but for now let us just assume those three components: code, stack, and heap.

Figure 13.3: An Example Address Space

In the diagram, there are three processes (A, B, and C) and each of them have a small part of the 512-KB physical memory carved out for them. Assuming a single CPU, the OS chooses to run one of the processes (say A), while the others (B and C) sit in the ready queue waiting to run.

As time sharing became more popular, you can probably guess that new demands were placed on the operating system. In particular, allowing multiple programs to reside concurrently in memory makes protection an important issue; you don’t want a process to be able to read, or worse, write some other process’s memory.
In the example in Figure 13.3, we have a tiny address space (only 16 KB). The program code lives at the top of the address space (starting at 0 in this example, and is packed into the first 1K of the address space). Code is static (and thus easy to place in memory), so we can place it at the top of the address space and know that it won’t need any more space as the program runs.

Next, we have the two regions of the address space that may grow (and shrink) while the program runs. Those are the heap (at the top) and the stack (at the bottom). We place them like this because each wishes to be able to grow, and by putting them at opposite ends of the address space, we can allow such growth: they just have to grow in opposite directions. The heap thus starts just after the code (at 1KB) and grows downward (say when a user requests more memory via `malloc()`); the stack starts at 16KB and grows upward (say when a user makes a procedure call). However, this placement of stack and heap is just a convention; you could arrange the address space in a different way if you’d like (as we’ll see later, when multiple threads co-exist in an address space, no nice way to divide the address space like this works anymore, alas).

Of course, when we describe the address space, what we are describing is the abstraction that the OS is providing to the running program. The program really isn’t in memory at physical addresses 0 through 16KB; rather it is loaded at some arbitrary physical address(es). Examine processes A, B, and C in Figure 13.2; there you can see how each process is loaded into memory at a different address. And hence the problem:

**THE CRUX: HOW TO VIRTUALIZE MEMORY**

How can the OS build this abstraction of a private, potentially large address space for multiple running processes (all sharing memory) on top of a single, physical memory?

When the OS does this, we say the OS is virtualizing memory, because the running program thinks it is loaded into memory at a particular address (say 0) and has a potentially very large address space (say 32-bits or 64-bits); the reality is quite different.

When, for example, process A in Figure 13.2 tries to perform a load at address 0 (which we will call a virtual address), somehow the OS, in tandem with some hardware support, will have to make sure the load doesn’t actually go to physical address 0 but rather to physical address 320KB (where A is loaded into memory). This is the key to virtualization of memory, which underlies every modern computer system in the world.

---

1We will often use small examples like this because it is a pain to represent a 32-bit address space and the numbers start to become hard to handle.
TIP: THE PRINCIPLE OF ISOLATION

Isolation is a key principle in building reliable systems. If two entities are properly isolated from one another, this implies that one can fail without affecting the other. Operating systems strive to isolate processes from each other and in this way prevent one from harming the other. By using memory isolation, the OS further ensures that running programs cannot affect the operation of the underlying OS. Some modern OS’s take isolation even further, by walling off pieces of the OS from other pieces of the OS. Such microkernels [BH70, R+89, S+03] thus may provide greater reliability than typical monolithic kernel designs.

13.4 Goals

Thus we arrive at the job of the OS in this set of notes: to virtualize memory. The OS will not only virtualize memory, though; it will do so with style. To make sure the OS does so, we need some goals to guide us. We have seen these goals before (think of the Introduction), and we’ll see them again, but they are certainly worth repeating.

One major goal of a virtual memory (VM) system is transparency\(^2\). The OS should implement virtual memory in a way that is invisible to the running program. Thus, the program shouldn’t be aware of the fact that memory is virtualized; rather, the program behaves as if it has its own private physical memory. Behind the scenes, the OS (and hardware) does all the work to multiplex memory among many different jobs, and hence implements the illusion.

Another goal of VM is efficiency. The OS should strive to make the virtualization as efficient as possible, both in terms of time (i.e., not making programs run much more slowly) and space (i.e., not using too much memory for structures needed to support virtualization). In implementing time-efficient virtualization, the OS will have to rely on hardware support, including hardware features such as TLBs (which we will learn about in due course).

Finally, a third VM goal is protection. The OS should make sure to protect processes from one another as well as the OS itself from processes. When one process performs a load, a store, or an instruction fetch, it should not be able to access or affect in any way the memory contents of any other process or the OS itself (that is, anything outside its address space). Protection thus enables us to deliver the property of isolation among processes; each process should be running in its own isolated cocoon, safe from the ravages of other faulty or even malicious processes.

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\(^2\)This usage of transparency is sometimes confusing; some students think that “being transparent” means keeping everything out in the open, i.e., what government should be like. Here, it means the opposite: that the illusion provided by the OS should not be visible to applications. Thus, in common usage, a transparent system is one that is hard to notice, not one that responds to requests as stipulated by the Freedom of Information Act.
Aside: Every Address You See Is Virtual

Ever write a C program that prints out a pointer? The value you see (some large number, often printed in hexadecimal), is a virtual address.

Ever wonder where the code of your program is found? You can print that out too, and yes, if you can print it, it also is a virtual address. In fact, any address you can see as a programmer of a user-level program is a virtual address. It’s only the OS, through its tricky techniques of virtualizing memory, that knows where in the physical memory of the machine these instructions and data values lie. So never forget: if you print out an address in a program, it’s a virtual one, an illusion of how things are laid out in memory; only the OS (and the hardware) knows the real truth.

Here’s a little program that prints out the locations of the main() routine (where code lives), the value of a heap-allocated value returned from malloc(), and the location of an integer on the stack:

```c
#include <stdio.h>
#include <stdlib.h>

int main(int argc, char *argv[]) {
    printf("location of code : %p\n", (void *) main);
    printf("location of heap : %p\n", (void *) malloc(1));
    int x = 3;
    printf("location of stack : %p\n", (void *) &x);
    return x;
}
```

When run on a 64-bit Mac OS X machine, we get the following output:

```
location of code : 0x1095afe50
location of heap : 0x1096008c0
location of stack : 0x7fff691aea64
```

From this, you can see that code comes first in the address space, then the heap, and the stack is all the way at the other end of this large virtual space. All of these addresses are virtual, and will be translated by the OS and hardware in order to fetch values from their true physical locations.

In the next chapters, we’ll focus our exploration on the basic mechanisms needed to virtualize memory, including hardware and operating systems support. We’ll also investigate some of the more relevant policies that you’ll encounter in operating systems, including how to manage free space and which pages to kick out of memory when you run low on space. In doing so, we’ll build up your understanding of how a modern virtual memory system really works.³

³Or, we’ll convince you to drop the course. But hold on; if you make it through VM, you’ll likely make it all the way!
13.5 Summary

We have seen the introduction of a major OS subsystem: virtual memory. The VM system is responsible for providing the illusion of a large, sparse, private address space to programs, which hold all of their instructions and data therein. The OS, with some serious hardware help, will take each of these virtual memory references, and turn them into physical addresses, which can be presented to the physical memory in order to fetch the desired information. The OS will do this for many processes at once, making sure to protect programs from one another, as well as protect the OS. The entire approach requires a great deal of mechanism (lots of low-level machinery) as well as some critical policies to work; we’ll start from the bottom up, describing the critical mechanisms first. And thus we proceed!
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J. McCarthy  
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14 Interlude: Memory API

In this interlude, we discuss the memory allocation interfaces in UNIX systems. The interfaces provided are quite simple, and hence the chapter is short and to the point. The main problem we address is this:

**CRUX: HOW TO ALLOCATE AND MANAGE MEMORY**
In UNIX/C programs, understanding how to allocate and manage memory is critical in building robust and reliable software. What interfaces are commonly used? What mistakes should be avoided?

14.1 Types of Memory

In running a C program, there are two types of memory that are allocated. The first is called stack memory, and allocations and deallocations of it are managed implicitly by the compiler for you, the programmer; for this reason it is sometimes called automatic memory.

Declaring memory on the stack in C is easy. For example, let’s say you need some space in a function `func()` for an integer, called `x`. To declare such a piece of memory, you just do something like this:

```c
void func() {
    int x; // declares an integer on the stack
    ...
}
```

The compiler does the rest, making sure to make space on the stack when you call into `func()`. When your return from the function, the compiler deallocates the memory for you; thus, if you want some information to live beyond the call invocation, you had better not leave that information on the stack.

It is this need for long-lived memory that gets us to the second type of memory, called heap memory, where all allocations and deallocations

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1Indeed, we hope all chapters are! But this one is shorter and pointier, we think.
are *explicitly* handled by you, the programmer. A heavy responsibility, no doubt! And certainly the cause of many bugs. But if you are careful and pay attention, you will use such interfaces correctly and without too much trouble. Here is an example of how one might allocate a pointer to an integer on the heap:

```c
void func() {
    int *x = (int *) malloc(sizeof(int));
    ...
}
```

A couple of notes about this small code snippet. First, you might notice that both stack and heap allocation occur on this line: first the compiler knows to make room for a pointer to an integer when it sees your declaration of said pointer (`int *x`); subsequently, when the program calls `malloc()`, it requests space for an integer on the heap; the routine returns the address of such an integer (upon success, or `NULL` on failure), which is then stored on the stack for use by the program.

Because of its explicit nature, and because of its more varied usage, heap memory presents more challenges to both users and systems. Thus, it is the focus of the remainder of our discussion.

### 14.2 The malloc() Call

The `malloc()` call is quite simple: you pass it a size asking for some room on the heap, and it either succeeds and gives you back a pointer to the newly-allocated space, or fails and returns `NULL`\(^2\).

The manual page shows what you need to do to use `malloc`; type `man malloc` at the command line and you will see:

```c
#include <stdlib.h>
...
void *malloc(size_t size);
```

From this information, you can see that all you need to do is include the header file `stdlib.h` to use `malloc`. In fact, you don’t really need to even do this, as the C library, which all C programs link with by default, has the code for `malloc()` inside of it; adding the header just lets the compiler check whether you are calling `malloc()` correctly (e.g., passing the right number of arguments to it, of the right type).

The single parameter `malloc()` takes is of type `size_t`, which simply describes how many bytes you need. However, most programmers do not type in a number here directly (such as 10); indeed, it would be considered poor form to do so. Instead, various routines and macros are utilized. For example, to allocate space for a double-precision floating point value, you simply do this:

```c
double *d = (double *) malloc(sizeof(double));
```

\(^2\)Note that `NULL` in C isn’t really anything special at all, just a macro for the value zero.
If you aren’t sure how some routine or operator you are using behaves, there is no substitute for simply trying it out and making sure it behaves as you expect. While reading the manual pages or other documentation is useful, how it works in practice is what matters. Write some code and test it! That is no doubt the best way to make sure your code behaves as you desire. Indeed, that is what we did to double-check the things we were saying about `sizeof()` were actually true!

Wow, that’s lot of double-ing! This invocation of `malloc()` uses the `sizeof()` operator to request the right amount of space; in C, this is generally thought of as a compile-time operator, meaning that the actual size is known at compile time and thus a number (in this case, 8, for a double) is substituted as the argument to `malloc()`. For this reason, `sizeof()` is correctly thought of as an operator and not a function call (a function call would take place at run time).

You can also pass in the name of a variable (and not just a type) to `sizeof()`, but in some cases you may not get the desired results, so be careful. For example, let’s look at the following code snippet:

```c
int *x = malloc(10 * sizeof(int));
printf(“%d
”, sizeof(x));
```

In the first line, we’ve declared space for an array of 10 integers, which is fine and dandy. However, when we use `sizeof()` in the next line, it returns a small value, such as 4 (on 32-bit machines) or 8 (on 64-bit machines). The reason is that in this case, `sizeof()` thinks we are simply asking how big a pointer to an integer is, not how much memory we have dynamically allocated. However, sometimes `sizeof()` does work as you might expect:

```c
int x[10];
printf(“%d
”, sizeof(x));
```

In this case, there is enough static information for the compiler to know that 40 bytes have been allocated.

Another place to be careful is with strings. When declaring space for a string, use the following idiom: `malloc(strlen(s) + 1)`, which gets the length of the string using the function `strlen()`, and adds 1 to it in order to make room for the end-of-string character. Using `sizeof()` may lead to trouble here.

You might also notice that `malloc()` returns a pointer to type `void`. Doing so is just the way in C to pass back an address and let the programmer decide what to do with it. The programmer further helps out by using what is called a cast; in our example above, the programmer casts the return type of `malloc()` to a pointer to a `double`. Casting doesn’t really accomplish anything, other than tell the compiler and other
programmers who might be reading your code: “yeah, I know what I’m doing.” By casting the result of malloc(), the programmer is just giving some reassurance; the cast is not needed for the correctness.

14.3 The free() Call

As it turns out, allocating memory is the easy part of the equation; knowing when, how, and even if to free memory is the hard part. To free heap memory that is no longer in use, programmers simply call free():

```c
int *x = malloc(10 * sizeof(int));
...
free(x);
```

The routine takes one argument, a pointer that was returned by malloc(). Thus, you might notice, the size of the allocated region is not passed in by the user, and must be tracked by the memory-allocation library itself.

14.4 Common Errors

There are a number of common errors that arise in the use of malloc() and free(). Here are some we’ve seen over and over again in teaching the undergraduate operating systems course. All of these examples compile and run with nary a peep from the compiler; while compiling a C program is necessary to build a correct C program, it is far from sufficient, as you will learn (often in the hard way).

Correct memory management has been such a problem, in fact, that many newer languages have support for automatic memory management. In such languages, while you call something akin to malloc() to allocate memory (usually new or something similar to allocate a new object), you never have to call something to free space; rather, a garbage collector runs and figures out what memory you no longer have references to and frees it for you.

Forgetting To Allocate Memory

Many routines expect memory to be allocated before you call them. For example, the routine strcpy(dst, src) copies a string from a source pointer to a destination pointer. However, if you are not careful, you might do this:

```c
char *src = "hello";
char *dst; // oops! unallocated
strcpy(dst, src); // segfault and die
```

When you run this code, it will likely lead to a segmentation fault, which is a fancy term for YOU DID SOMETHING WRONG WITH MEMORY YOU FOOLISH PROGRAMMER AND I AM ANGRY.

3Although it sounds arcane, you will soon learn why such an illegal memory access is called a segmentation fault; if that isn’t incentive to read on, what is?
In this case, the proper code might instead look like this:

```c
char *src = "hello";
char *dst = (char *) malloc(strlen(src) + 1);
strcpy(dst, src); // work properly
```

Alternately, you could use `strdup()` and make your life even easier. Read the `strdup` man page for more information.

**Not Allocating Enough Memory**

A related error is not allocating enough memory, sometimes called a buffer overflow. In the example above, a common error is to make *almost* enough room for the destination buffer.

```c
char *src = "hello";
char *dst = (char *) malloc(strlen(src)) + 1); // too small!
strcpy(dst, src); // work properly
```

Oddly enough, depending on how malloc is implemented and many other details, this program will often run seemingly correctly. In some cases, when the string copy executes, it writes one byte too far past the end of the allocated space, but in some cases this is harmless, perhaps overwriting a variable that isn’t used anymore. In some cases, these overflows can be incredibly harmful, and in fact are the source of many security vulnerabilities in systems [W06]. In other cases, the malloc library allocated a little extra space anyhow, and thus your program actually doesn’t scribble on some other variable’s value and works quite fine. In even other cases, the program will indeed fault and crash. And thus we learn another valuable lesson: even though it ran correctly once, doesn’t mean it’s correct.

**Forgetting to Initialize Allocated Memory**

With this error, you call `malloc()` properly, but forget to fill in some values into your newly-allocated data type. Don’t do this! If you do forget, your program will eventually encounter an uninitialized read, where it
reads from the heap some data of unknown value. Who knows what might be in there? If you’re lucky, some value such that the program still works (e.g., zero). If you’re not lucky, something random and harmful.

**Forgetting To Free Memory**

Another common error is known as a **memory leak**, and it occurs when you forget to free memory. In long-running applications or systems (such as the OS itself), this is a huge problem, as slowly leaking memory eventually leads one to run out of memory, at which point a restart is required. Thus, in general, when you are done with a chunk of memory, you should make sure to free it. Note that using a garbage-collected language doesn’t help here: if you still have a reference to some chunk of memory, no garbage collector will ever free it, and thus memory leaks remain a problem even in more modern languages.

Note that not all memory need be freed, at least, in certain cases. For example, when you write a short-lived program, you might allocate some space using `malloc()`. The program runs and is about to complete: is there need to call `free()` a bunch of times just before exiting? While it seems wrong not to, it is in this case quite fine to simply exit. After all, when your program exits, the OS will clean up everything about this process, including any memory it has allocated. Calling `free()` a bunch of times and then exiting is thus pointless, and, if you do so incorrectly, will cause the program to crash. Just call `exit` and be happy instead.

**Freeing Memory Before You Are Done With It**

Sometimes a program will free memory before it is finished using it; such a mistake is called a **dangling pointer**, and it, as you can guess, is also a bad thing. The subsequent use can crash the program, or overwrite valid memory (e.g., you called `free()`, but then called `malloc()` again to allocate something else, which then recycles the errantly-freed memory).

**Freeing Memory Repeatedly**

Programs also sometimes free memory more than once; this is known as the **double free**. The result of doing so is undefined. As you can imagine, the memory-allocation library might get confused and do all sorts of weird things; crashes are a common outcome.

**Calling free () Incorrectly**

One last problem we discuss is the call of `free()` incorrectly. After all, `free()` expects you only to pass to it one of the pointers you received from `malloc()` earlier. When you pass in some other value, bad things can (and do) happen. Thus, such **invalid frees** are dangerous and of course should also be avoided.
Summary

As you can see, there are lots of ways to abuse memory. Because of frequent errors with memory, a whole ecosphere of tools have developed to help find such problems in your code. Check out both purify [HJ92] and valgrind [SN05]; both are excellent at helping you locate the source of your memory-related problems. Once you become accustomed to using these powerful tools, you will wonder how you survived without them.

14.5 Underlying OS Support

You might have noticed that we haven’t been talking about system calls when discussing malloc() and free(). The reason for this is simple: they are not system calls, but rather library calls. Thus the malloc library manages space within your virtual address space, but itself is built on top of some system calls which call into the OS to ask for more memory or release some back to the system.

One such system call is called brk, which is used to change the location of the program’s break: the location of the end of the heap. It takes one argument (the address of the new break), and thus either increases or decreases the size of the heap based on whether the new break is larger or smaller than the current break. An additional call sbrk is passed an increment but otherwise serves a similar purpose.

Note that you should never directly call either brk or sbrk. They are used by the memory-allocation library; if you try to use them, you will likely make something go (horribly) wrong. Stick to malloc() and free() instead.

Finally, you can also obtain memory from the operating system via the mmap() call. By passing in the correct arguments, mmap() can create an anonymous memory region within your program – a region which is not associated with any particular file but rather with swap space, something we’ll discuss in detail later on in virtual memory. This memory can then also be treated like a heap and managed as such. Read the manual page of mmap() for more details.

14.6 Other Calls

There are a few other calls that the memory-allocation library supports. For example, calloc() allocates memory and also zeroes it before returning; this prevents some errors where you assume that memory is zeroed and forget to initialize it yourself (see the paragraph on “uninitialized reads” above). The routine realloc() can also be useful, when you’ve allocated space for something (say, an array), and then need to add something to it: realloc() makes a new larger region of memory, copies the old region into it, and returns the pointer to the new region.
14.7 Summary

We have introduced some of the APIs dealing with memory allocation. As always, we have just covered the basics; more details are available elsewhere. Read the C book [KR88] and Stevens [SR05] (Chapter 7) for more information. For a cool modern paper on how to detect and correct many of these problems automatically, see Novark et al. [N+07]; this paper also contains a nice summary of common problems and some neat ideas on how to find and fix them.
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[W06] “Survey on Buffer Overflow Attacks and Countermeasures”
Tim Werthman
Available: www.nds.rub.de/lehre/seminar/SS06/Werthmann_BufferOverflow.pdf
A nice survey of buffer overflows and some of the security problems they cause. Refers to many of the famous exploits.
Mechanism: Address Translation

In developing the virtualization of the CPU, we focused on a general mechanism known as **limited direct execution** (or LDE). The idea behind LDE is simple: for the most part, let the program run directly on the hardware; however, at certain key points in time (such as when a process issues a system call, or a timer interrupt occurs), arrange so that the OS gets involved and makes sure the “right” thing happens. Thus, the OS, with a little hardware support, tries its best to get out of the way of the running program, to deliver an efficient virtualization; however, by **interposing** at those critical points in time, the OS ensures that it maintains control over the hardware. Efficiency and control together are two of the main goals of any modern operating system.

In virtualizing memory, we will pursue a similar strategy, attaining both efficiency and control while providing the desired virtualization. Efficiency dictates that we make use of hardware support, which at first will be quite rudimentary (e.g., just a few registers) but will grow to be fairly complex (e.g., TLBs, page-table support, and so forth, as you will see). Control implies that the OS ensures that no application is allowed to access any memory but its own; thus, to protect applications from one another, and the OS from applications, we will need help from the hardware here too. Finally, we will need a little more from the VM system, in terms of flexibility; specifically, we’d like for programs to be able to use their address spaces in whatever way they would like, thus making the system easier to program. And thus arrive at the refined crux:

**THE CRUX:**

**How to Efficiently and Flexibly Virtualize Memory**

How can we build an efficient virtualization of memory? How do we provide the flexibility needed by applications? How do we maintain control over which memory locations an application can access, and thus ensure that application memory accesses are properly restricted? How do we do all of this efficiently?
The generic technique we will use, which you can consider an addition to our general approach of limited direct execution, is something that is referred to as **hardware-based address translation**, or just **address translation** for short. With address translation, the hardware transforms each memory access (e.g., an instruction fetch, load, or store), changing the virtual address provided by the instruction to a physical address where the desired information is actually located. Thus, on each and every memory reference, an address translation is performed by the hardware to redirect application memory references to their actual locations in memory.

Of course, the hardware alone cannot virtualize memory, as it just provides the low-level mechanism for doing so efficiently. The OS must get involved at key points to set up the hardware so that the correct translations take place; it must thus **manage memory**, keeping track of which locations are free and which are in use, and judiciously intervening to maintain control over how memory is used.

Once again the goal of all of this work is to create a beautiful **illusion**: that the program has its own private memory, where its own code and data reside. Behind that virtual reality lies the ugly physical truth: that many programs are actually sharing memory at the same time, as the CPU (or CPUs) switches between running one program and the next. Through virtualization, the OS (with the hardware’s help) turns the ugly machine reality into something that is a useful, powerful, and easy to use abstraction.

### 15.1 Assumptions

Our first attempts at virtualizing memory will be very simple, almost laughably so. Go ahead, laugh all you want; pretty soon it will be the OS laughing at you, when you try to understand the ins and outs of TLBs, multi-level page tables, and other technical wonders. Don’t like the idea of the OS laughing at you? Well, you may be out of luck then; that’s just how the OS rolls.

Specifically, we will assume for now that the user’s address space must be placed **contiguously** in physical memory. We will also assume, for simplicity, that the size of the address space is not too big; specifically, that it is **less than the size of physical memory**. Finally, we will also assume that each address space is exactly the **same size**. Don’t worry if these assumptions sound unrealistic; we will relax them as we go, thus achieving a realistic virtualization of memory.

### 15.2 An Example

To understand better what we need to do to implement address translation, and why we need such a mechanism, let’s look at a simple example. Imagine there is a process whose address space as indicated in Figure 15.1. What we are going to examine here is a short code sequence that
Interposition is a generic and powerful technique that is often used to great effect in computer systems. In virtualizing memory, the hardware will interpose on each memory access, and translate each virtual address issued by the process to a physical address where the desired information is actually stored. However, the general technique of interposition is much more broadly applicable; indeed, almost any well-defined interface can be interposed upon, to add new functionality or improve some other aspect of the system. One of the usual benefits of such an approach is transparency; the interposition often is done without changing the client of the interface, thus requiring no changes to said client.

loads a value from memory, increments it by three, and then stores the value back into memory. You can imagine the C-language representation of this code might look like this:

```c
void func()
    int x;
    ...
    x = x + 3; // this is the line of code we are interested in
```

The compiler turns this line of code into assembly, which might look something like this (in x86 assembly). Use `objdump` on Linux or `otool` on Mac OS X to disassemble it:

```
128: movl 0x0(%ebx), %eax ;load 0+ebx into eax
132: addl $0x03, %eax ;add 3 to eax register
135: movl %eax, 0x0(%ebx) ;store eax back to mem
```

This code snippet is relatively straightforward; it presupposes that the address of `x` has been placed in the register `ebx`, and then loads the value at that address into the general-purpose register `eax` using the `movl` instruction (for “longword” move). The next instruction adds 3 to `eax`, and the final instruction stores the value in `eax` back into memory at that same location.

In Figure 15.1, you can see how both the code and data are laid out in the process’s address space; the three-instruction code sequence is located at address 128 (in the code section near the top), and the value of the variable `x` at address 15 KB (in the stack near the bottom). In the figure, the initial value of `x` is 3000, as shown in its location on the stack.

When these instructions run, from the perspective of the process, the following memory accesses take place.

- Fetch instruction at address 128
- Execute this instruction (load from address 15 KB)
- Fetch instruction at address 132
- Execute this instruction (no memory reference)
- Fetch the instruction at address 135
- Execute this instruction (store to address 15 KB)
From the program’s perspective, its address space starts at address 0 and grows to a maximum of 16 KB; all memory references it generates should be within these bounds. However, to virtualize memory, the OS wants to place the process somewhere else in physical memory, not necessarily at address 0. Thus, we have the problem: how can we relocate this process in memory in a way that is transparent to the process? How can provide the illusion of a virtual address space starting at 0, when in reality the address space is located at some other physical address?
An example of what physical memory might look like once this process’s address space has been placed in memory is found in Figure 15.2. In the figure, you can see the OS using the first slot of physical memory for itself, and that it has relocated the process from the example above into the slot starting at physical memory address 32 KB. The other two slots are free (16 KB-32 KB and 48 KB-64 KB).

15.3 Dynamic (Hardware-based) Relocation

To gain some understanding of hardware-based address translation, we’ll first discuss its first incarnation. Introduced in the first time-sharing machines of the late 1950’s is a simple idea referred to as base and bounds (the technique is also referred to as dynamic relocation; we’ll use both terms interchangeably) [SS74].

Specifically, we’ll need two hardware registers within each CPU: one is called the base register, and the other the bounds (sometimes called a limit register). This base-and-bounds pair is going to allow us to place the address space anywhere we’d like in physical memory, and do so while ensuring that the process can only access its own address space.

In this setup, each program is written and compiled as if it is loaded at address zero. However, when a program starts running, the OS decides where in physical memory it should be loaded and sets the base register to that value. In the example above, the OS decides to load the process at physical address 32 KB and thus sets the base register to this value.

Interesting things start to happen when the process is running. Now, when any memory reference is generated by the process, it is translated by the processor in the following manner:

\[
\text{physical address} = \text{virtual address} + \text{base}
\]
ASIDE: SOFTWARE-BASED RELOCATION

In the early days, before hardware support arose, some systems performed a crude form of relocation purely via software methods. The basic technique is referred to as static relocation, in which a piece of software known as the loader takes an executable that is about to be run and rewrites its addresses to the desired offset in physical memory.

For example, if an instruction was a load from address 1000 into a register (e.g., movl 1000, %eax), and the address space of the program was loaded starting at address 3000 (and not 0, as the program thinks), the loader would rewrite the instruction to offset each address by 3000 (e.g., movl 4000, %eax). In this way, a simple static relocation of the process’s address space is achieved.

However, static relocation has numerous problems. First and most importantly, it does not provide protection, as processes can generate bad addresses and thus illegally access other process’s or even OS memory; in general, hardware support is likely needed for true protection [WL+93]. A smaller negative is that once placed, it is difficult to later relocate an address space to another location [M65].

Each memory reference generated by the process is a virtual address; the hardware in turn adds the contents of the base register to this address and the result is a physical address that can be issued to the memory system.

To understand this better, let’s trace through what happens when a single instruction is executed. Specifically, let’s look at one instruction from our earlier sequence:

128: movl 0x0(%ebx), %eax

The program counter (PC) is set to 128; when the hardware needs to fetch this instruction, it first adds the value to the the base register value of 32 KB (32768) to get a physical address of 32896; the hardware then fetches the instruction from that physical address. Next, the processor begins executing the instruction. At some point, the process then issues the load from virtual address 15 KB, which the processor takes and again adds to the base register (32 KB), getting the final physical address of 47 KB and thus the desired contents.

Transforming a virtual address into a physical address is exactly the technique we refer to as address translation; that is, the hardware takes a virtual address the process thinks it is referencing and transforms it into a physical address which is where the data actually resides. Because this relocation of the address happens at runtime, and because we can move address spaces even after the process has started running, the technique is often referred to as dynamic relocation [M65].
**MECHANISM: ADDRESS TRANSLATION**

**TIP: HARDWARE-BASED DYNAMIC RELOCATION**

With dynamic relocation, we can see how a little hardware goes a long way. Namely, a base register is used to transform virtual addresses (generated by the program) into physical addresses. A bounds (or limit) register ensures that such addresses are within the confines of the address space. Together, they combine to provide a simple and efficient virtualization of memory.

Now you might be asking: what happened to that bounds (limit) register? After all, isn’t this supposed to be the base-and-bounds approach? Indeed, it is. And as you might have guessed, the bounds register is there to help with protection. Specifically, the processor will first check that the memory reference is within bounds to make sure it is legal; in the simple example above, the bounds register would always be set to 16 KB. If a process generates a virtual address that is greater than the bounds, or one that is negative, the CPU will raise an exception, and the process will likely be terminated. The point of the bounds is thus to make sure that all addresses generated by the process are legal and within the “bounds” of the process.

We should note that the base and bounds registers are hardware structures kept on the chip (one pair per CPU). Sometimes people call the part of the processor that helps with address translation the memory management unit (MMU); as we develop more sophisticated memory-management techniques, we will be adding more circuitry to the MMU.

A small aside about bound registers, which can be defined in one of two ways. In one way (as above), it holds the size of the address space, and thus the hardware checks the virtual address against it first before adding the base. In the second way, it holds the physical address of the end of the address space, and thus the hardware first adds the base and then makes sure the address is within bounds. Both methods are logically equivalent; for simplicity, we’ll usually assume that the bounds register holds the size of the address space.

**Example Translations**

To understand address translation via base-and-bounds in more detail, let’s take a look at an example. Imagine a process with an address space of size 4 KB (yes, unrealistically small) has been loaded at physical address 16 KB. Here are the results of a number of address translations:

- Virtual Address 0 → Physical Address 16 KB
- VA 1 KB → PA 17 KB
- VA 3000 → PA 19384
- VA 4400 → Fault (out of bounds)
The OS must track which parts of free memory are not in use, so as to be able to allocate memory to processes. Many different data structures can of course be used for such a task; the simplest (which we will assume here) is a **free list**, which simply is a list of the ranges of the physical memory which are not currently in use.

As you can see from the example, it is easy for you to simply add the base address to the virtual address (which can rightly be viewed as an **offset** into the address space) to get the resulting physical address. Only if the virtual address is “too big” or negative will the result be a fault (e.g., 4400 is greater than the 4 KB bounds), causing an exception to be raised and the process to be terminated.

### 15.4 OS Issues

There are a number of new OS issues that arise when using base and bounds to implement a simple virtual memory. Specifically, there are three critical junctures where the OS must take action to implement this base-and-bounds approach to virtualizing memory.

First, the OS must take action when a process is created, finding space for its address space in memory. Fortunately, given our assumptions that each address space is (a) smaller than the size of physical memory and (b) the same size, this is quite easy for the OS; it can simply view physical memory as an array of slots, and track whether each one is free or in use. When a new process is created, the OS will have to search a data structure (often called a **free list**) to find room for the new address space and then mark it used.

An example of what physical memory might look like can be found in Figure 15.2. In the figure, you can see the OS using the first slot of physical memory for itself, and that it has relocated the process from the example above into the slot starting at physical memory address 32 KB. The other two slots are free (16 KB-32 KB and 48 KB-64 KB); thus, the **free list** should consist of these two entries.

Second, the OS must take action when a process is terminated, reclaiming all of its memory for use in other processes or the OS. Upon termination of a process, the OS thus puts its memory back on the free list, and cleans up any associated data structures as need be.

Third, the OS must also take action when a context switch occurs. There is only one base and bounds register on each CPU, after all, and their values differ for each running program, as each program is loaded at a different physical address in memory. Thus, the OS must save and restore the base-and-bounds pair when it switches between processes. Specifically, when the OS decides to stop running a process, it must save the
values of the base and bounds registers to memory, in some per-process structure such as the process structure or process control block (PCB). Similarly, when the OS resumes a running process (or runs it the first time), it must set the values of the base and bounds on the CPU to the correct values for this process.

We should note that when a process is stopped (i.e., not running), it is possible for the OS to move an address space from one location in memory to another rather easily. To move a process’s address space, the OS first deschedules the process; then, the OS copies the address space from the current location to the new location; finally, the OS updates the saved base register (in the process structure) to point to the new location. When the process is resumed, its (new) base register is restored, and it begins running again, oblivious that its instructions and data are now in a completely new spot in memory!

We should also note that access to the base and bounds registers is obviously privileged. Special hardware instructions are required to access base-and-bounds registers; if a process, running in user mode, attempts to do so, the CPU will raise an exception and the OS will likely terminate the process. Only in kernel (or privileged) mode can such registers be modified. Imagine the havoc a user process could wreak\(^1\) if it could arbitrarily change the base register while running. Imagine it! And then quickly flush such dark thoughts from your mind, as they are the ghastly stuff of which nightmares are made.

15.5 Summary

In this chapter, we have extended the concept of limited direct execution with a specific mechanism used in virtual memory, known as address translation. With address translation, the OS can control each and every memory access from a process, ensuring the accesses stay within the bounds of the address space. Key to the efficiency of this technique is hardware support, which performs the translation quickly for each access, turning virtual addresses (the process’s view of memory) into physical ones (the actual view). All of this is performed in a way that is transparent to the process that has been relocated; the process has no idea its memory references are being translated, making for a wonderful illusion.

We have also seen one particular form of virtualization, known as base and bounds or dynamic relocation. Base-and-bounds virtualization is quite efficient, as only a little more hardware logic is required to add a base register to the virtual address and check that the address generated by the process is in bounds. Base-and-bounds also offers protection; the OS and hardware combine to ensure no process can generate memory references outside its own address space. Protection is certainly one of the most important goals of the OS; without it, the OS could not control

\(^1\)Is there anything other than “havoc” that can be “wreaked”? 

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THREE EASY PIECES
the machine (if processes were free to overwrite memory, they could easily do nasty things like overwrite the trap table and take over the system).

Unfortunately, this simple technique of dynamic relocation does have its inefficiencies. For example, as you can see in Figure 15.2 (back a few pages), the relocated process is using physical memory from 32 KB to 48 KB; however, because the process stack and heap are not too big, all of the space between the two is simply wasted. This type of waste is usually called internal fragmentation, as the space inside the allocated unit is not all used (i.e., is fragmented) and thus wasted. In our current approach, although there might be enough physical memory for more processes, we are currently restricted to placing an address space in a fixed-sized slot and thus internal fragmentation can arise. Thus, we are going to need more sophisticated machinery, to try to better utilize physical memory and avoid internal fragmentation. Our first attempt will be a slight generalization of base and bounds known as segmentation, which we will discuss next.

\[A\] different solution might instead place a fixed-sized stack within the address space, just below the code region, and a growing heap below that. However, this limits flexibility by making recursion and deeply-nested function calls challenging, and thus is something we hope to avoid.
References

[M65] “On Dynamic Program Relocation”  
W.C. McGee  
IBM Systems Journal  
Volume 4, Number 3, 1965, pages 184–199  
This paper is a nice summary of early work on dynamic relocation, as well as some basics on static relocation.

[P90] “Relocating loader for MS-DOS .EXE executable files”  
Kenneth D. A. Pillay  
Microprocessors & Microsystems archive  
Volume 14, Issue 7 (September 1990)  
An example of a relocating loader for MS-DOS. Not the first one, but just a relatively modern example of how such a system works.

[SS74] “The Protection of Information in Computer Systems”  
J. Saltzer and M. Schroeder  
CACM, July 1974  
From this paper: “The concepts of base-and-bound register and hardware-interpreted descriptors appeared, apparently independently, between 1957 and 1959 on three projects with diverse goals. At M.I.T., McCarthy suggested the base-and-bound idea as part of the memory protection system necessary to make time-sharing feasible. IBM independently developed the base-and-bound register as a mechanism to permit reliable multiprogramming of the Stretch (7030) computer system. At Burroughs, R. Barton suggested that hardware-interpreted descriptors would provide direct support for the naming scope rules of higher level languages in the B5000 computer system.” We found this quote on Mark Smotherman’s cool history pages [S04]; see them for more information.

[S04] “System Call Support”  
Mark Smotherman, May 2004  
http://people.cs.clemson.edu/~mark/syscall.html  
A neat history of system call support. Smotherman has also collected some early history on items like interrupts and other fun aspects of computing history. See his web pages for more details.

Robert Wahbe, Steven Lucco, Thomas E. Anderson, Susan L. Graham  
SOSP ’93  
A terrific paper about how you can use compiler support to bound memory references from a program, without hardware support. The paper sparked renewed interest in software techniques for isolation of memory references.
Homework

The program `relocation.py` allows you to see how address translations are performed in a system with base and bounds registers. See the README for details.

Questions

- Run with seeds 1, 2, and 3, and compute whether each virtual address generated by the process is in or out of bounds. If in bounds, compute the translation.
- Run with these flags: `-s 0 -n 10`. What value do you have set `-l` (the bounds register) to in order to ensure that all the generated virtual addresses are within bounds?
- Run with these flags: `-s 1 -n 10 -l 100`. What is the maximum value that bounds can be set to, such that the address space still fits into physical memory in its entirety?
- Run some of the same problems above, but with larger address spaces (`-a`) and physical memories (`-p`).
- What fraction of randomly-generated virtual addresses are valid, as a function of the value of the bounds register? Make a graph from running with different random seeds, with limit values ranging from 0 up to the maximum size of the address space.
Segmentation

So far we have been putting the entire address space of each process in memory. With the base and bounds registers, the OS can easily relocate processes to different parts of physical memory. However, you might have noticed something interesting about these address spaces of ours: there is a big chunk of “free” space right in the middle, between the stack and the heap.

As you can imagine from Figure 16.1, although the space between the stack and heap is not being used by the process, it is still taking up physical memory when we relocate the entire address space somewhere in physical memory; thus, the simple approach of using a base and bounds register pair to virtualize memory is wasteful. It also makes it quite hard to run a program when the entire address space doesn’t fit into memory; thus, base and bounds is not as flexible as we would like. And thus:

**THE CRUX: HOW TO SUPPORT A LARGE ADDRESS SPACE**

How do we support a large address space with (potentially) a lot of free space between the stack and the heap? Note that in our examples, with tiny (pretend) address spaces, the waste doesn’t seem too bad. Imagine, however, a 32-bit address space (4 GB in size); a typical program will only use megabytes of memory, but still would demand that the entire address space be resident in memory.

16.1 Segmentation: Generalized Base/Bounds

To solve this problem, an idea was born, and it is called **segmentation**. It is quite an old idea, going at least as far back as the very early 1960’s [H61, G62]. The idea is simple: instead of having just one base and bounds pair in our MMU, why not have a base and bounds pair per logical segment of the address space? A segment is just a contiguous portion of the address space of a particular length, and in our canonical
address space, we have three logically-different segments: code, stack, and heap. What segmentation allows the OS to do is to place each one of those segments in different parts of physical memory, and thus avoid filling physical memory with unused virtual address space.

Let’s look at an example. Assume we want to place the address space from Figure 16.1 into physical memory. With a base and bounds pair per segment, we can place each segment independently in physical memory. For example, see Figure 16.2; there you see a 64-KB physical memory with those three segments within it (and 16KB reserved for the OS).

Figure 16.1: *An Address Space (Again)*
As you can see in the diagram, only used memory is allocated space in physical memory, and thus large address spaces with large amounts of unused address space (which we sometimes call sparse address spaces) can be accommodated.

The hardware structure in our MMU required to support segmentation is just what you’d expect: in this case, a set of three base and bounds register pairs. Table 16.1 below shows the register values for the example above; each bounds register holds the size of a segment.

<table>
<thead>
<tr>
<th>Segment</th>
<th>Base</th>
<th>Size</th>
</tr>
</thead>
<tbody>
<tr>
<td>Code</td>
<td>32K</td>
<td>2K</td>
</tr>
<tr>
<td>Heap</td>
<td>34K</td>
<td>2K</td>
</tr>
<tr>
<td>Stack</td>
<td>28K</td>
<td>2K</td>
</tr>
</tbody>
</table>

Table 16.1: Segment Register Values

You can see from the table that the code segment is placed at physical address 32KB and has a size of 2KB and the heap segment is placed at 34KB and also has a size of 2KB.

Let’s do an example translation, using the address space in Figure 16.1. Assume a reference is made to virtual address 100 (which is in the code segment). When the reference takes place (say, on an instruction fetch), the hardware will add the base value to the offset into this segment (100 in this case) to arrive at the desired physical address: 100 + 32KB, or 32868. It will then check that the address is within bounds (100 is less than 2KB), find that it is, and issue the reference to physical memory address 32868.
The term segmentation fault or violation arises from a memory access on a segmented machine to an illegal address. Humorously, the term persists, even on machines with no support for segmentation at all. Or not so humorously, if you can’t figure why your code keeps faulting.

Now let’s look at an address in the heap, virtual address 4200 (again refer to Figure 16.1). If we just add the virtual address 4200 to the base of the heap (34KB), we get a physical address of 39016, which is not the correct physical address. What we need to first do is extract the offset into the heap, i.e., which byte(s) in this segment the address refers to. Because the heap starts at virtual address 4KB (4096), the offset of 4200 is actually 4200 – 4096 or 104. We then take this offset (104) and add it to the base register physical address (34K or 34816) to get the desired result: 34920.

What if we tried to refer to an illegal address, such as 7KB which is beyond the end of the heap? You can imagine what will happen: the hardware detects that the address is out of bounds, traps into the OS, likely leading to the termination of the offending process. And now you know the origin of the famous term that all C programmers learn to dread: the segmentation violation or segmentation fault.

16.2 Which Segment Are We Referring To?

The hardware uses segment registers during translation. How does it know the offset into a segment, and to which segment an address refers?

One common approach, sometimes referred to as an explicit approach, is to chop up the address space into segments based on the top few bits of the virtual address; this technique was used in the VAX/VMS system [LL82]. In our example above, we have three segments; thus we need two bits to accomplish our task. If we use the top two bits of our 14-bit virtual address to select the segment, our virtual address looks like this:

```
13 12 11 10  9  8  7  6  5  4  3  2  1  0
Segment  Offset
```

In our example, then, if the top two bits are 00, the hardware knows the virtual address is in the code segment, and thus uses the code base and bounds pair to relocate the address to the correct physical location. If the top two bits are 01, the hardware knows the address is in the heap, and thus uses the heap base and bounds. Let’s take our example heap virtual address from above (4200) and translate it, just to make sure this is clear. The virtual address 4200, in binary form, can be seen here:

```
13 12 11 10  9  8  7  6  5  4  3  2  1  0
Segment  Offset
```

```
0 1 0 0 0 0 0 1 1 0 1 0 0 0
```
As you can see from the picture, the top two bits (01) tell the hardware which segment we are referring to. The bottom 12 bits are the offset into the segment: 0000 0110 1000, or hex 0x068, or 104 in decimal. Thus, the hardware simply takes the first two bits to determine which segment register to use, and then takes the next 12 bits as the offset into the segment. By adding the base register to the offset, the hardware arrives at the final physical address. Note the offset eases the bounds check too: we can simply check if the offset is less than the bounds; if not, the address is illegal. Thus, if base and bounds were arrays (with one entry per segment), the hardware would be doing something like this to obtain the desired physical address:

```plaintext
1 // get top 2 bits of 14-bit VA
2 Segment = (VirtualAddress & SEG_MASK) >> SEG_SHIFT
3 // now get offset
4 Offset = VirtualAddress & OFFSET_MASK
5 if (Offset >= Bounds[Segment])
6    RaiseException(PROTECTION_FAULT)
7 else
8    PhysAddr = Base[Segment] + Offset
9    Register = AccessMemory(PhysAddr)
```

In our running example, we can fill in values for the constants above. Specifically, SEG_MASK would be set to 0x3000, SEG_SHIFT to 12, and OFFSET_MASK to 0xFFF.

You may also have noticed that when we use the top two bits, and we only have three segments (code, heap, stack), one segment of the address space goes unused. Thus, some systems put code in the same segment as the heap and thus use only one bit to select which segment to use [LL82].

There are other ways for the hardware to determine which segment a particular address is in. In the implicit approach, the hardware determines the segment by noticing how the address was formed. If, for example, the address was generated from the program counter (i.e., it was an instruction fetch), then the address is within the code segment; if the address is based off of the stack or base pointer, it must be in the stack segment; any other address must be in the heap.

### 16.3 What About The Stack?

Thus far, we’ve left out one important component of the address space: the stack. The stack has been relocated to physical address 28KB in the diagram above, but with one critical difference: it grows backwards – in physical memory, it starts at 28KB and grows back to 26KB, corresponding to virtual addresses 16KB to 14KB; translation has to proceed differently.

The first thing we need is a little extra hardware support. Instead of just base and bounds values, the hardware also needs to know which way the segment grows (a bit, for example, that is set to 1 when the segment grows in the positive direction, and 0 for negative). Our updated view of what the hardware tracks is seen in Table 16.2.
146 S
E G M E N T A T I O N
Table 16.2: Segmentation (With Negative-Growth Support)

<table>
<thead>
<tr>
<th>Segment</th>
<th>Base</th>
<th>Size</th>
<th>Grows Positive?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Code</td>
<td>32K</td>
<td>2K</td>
<td>1</td>
</tr>
<tr>
<td>Heap</td>
<td>34K</td>
<td>2K</td>
<td>1</td>
</tr>
<tr>
<td>Stack</td>
<td>28K</td>
<td>2K</td>
<td>0</td>
</tr>
</tbody>
</table>

With the hardware understanding that segments can grow in the negative direction, the hardware must now translate such virtual addresses slightly differently. Let’s take an example stack virtual address and translate it to understand the process.

In this example, assume we wish to access virtual address 15KB, which should map to physical address 27KB. Our virtual address, in binary form, thus looks like this: 11 1100 0000 0000 (hex 0x3C00). The hardware uses the top two bits (11) to designate the segment, but then we are left with an offset of 3KB. To obtain the correct negative offset, we must subtract the maximum segment size from 3KB: in this example, a segment can be 4KB, and thus the correct negative offset is 3KB - 4KB which equals -1KB. We simply add the negative offset (-1KB) to the base (28KB) to arrive at the correct physical address: 27KB. The bounds check can be calculated by ensuring the absolute value of the negative offset is less than the segment’s size.

16.4 Support for Sharing

As support for segmentation grew, system designers soon realized that they could realize new types of efficiencies with a little more hardware support. Specifically, to save memory, sometimes it is useful to share certain memory segments between address spaces. In particular, code sharing is common and still in use in systems today.

To support sharing, we need a little extra support from the hardware, in the form of protection bits. Basic support adds a few bits per segment, indicating whether or not a program can read or write a segment, or perhaps execute code that lies within the segment. By setting a code segment to read-only, the same code can be shared across multiple processes, without worry of harming isolation; while each process still thinks that it is accessing its own private memory, the OS is secretly sharing memory which cannot be modified by the process, and thus the illusion is preserved.

An example of the additional information tracked by the hardware (and OS) is shown in Figure 16.3. As you can see, the code segment is set to read and execute, and thus the same physical segment in memory could be mapped into multiple virtual address spaces.

With protection bits, the hardware algorithm described earlier would also have to change. In addition to checking whether a virtual address is within bounds, the hardware also has to check whether a particular access is permissible. If a user process tries to write to a read-only page, or execute from a non-executable page, the hardware should raise an exception, and thus let the OS deal with the offending process.
Segmentation

<table>
<thead>
<tr>
<th>Segment</th>
<th>Base</th>
<th>Size</th>
<th>Grows Positive?</th>
<th>Protection</th>
</tr>
</thead>
<tbody>
<tr>
<td>Code</td>
<td>32K</td>
<td>2K</td>
<td>1</td>
<td>Read-Execute</td>
</tr>
<tr>
<td>Heap</td>
<td>34K</td>
<td>2K</td>
<td>1</td>
<td>Read-Write</td>
</tr>
<tr>
<td>Stack</td>
<td>28K</td>
<td>2K</td>
<td>0</td>
<td>Read-Write</td>
</tr>
</tbody>
</table>

Table 16.3: Segment Register Values (with Protection)

16.5 Fine-grained vs. Coarse-grained Segmentation

Most of our examples thus far have focused on systems with just a few segments (i.e., code, stack, heap); we can think of this segmentation as coarse-grained, as it chops up the address space into relatively large, coarse chunks. However, some early systems (e.g., Multics [CV65,DD68]) were more flexible and allowed for address spaces to consist of a large number smaller segments, referred to as fine-grained segmentation.

Supporting many segments requires even further hardware support, with a segment table of some kind stored in memory. Such segment tables usually support the creation of a very large number of segments, and thus enable a system to use segments in more flexible ways than we have thus far discussed. For example, early machines like the Burroughs B5000 had support for thousands of segments, and expected a compiler to chop code and data into separate segments which the OS and hardware would then support [RK68]. The thinking at the time was that by having fine-grained segments, the OS could better learn about which segments are in use and which are not and thus utilize main memory more effectively.

16.6 OS Support

You now should have a basic idea as to how segmentation works. Pieces of the address space are relocated into physical memory as the system runs, and thus a huge savings of physical memory is achieved relative to our simpler approach with just a single base/bounds pair for the entire address space. Specifically, all the unused space between the stack and the heap need not be allocated in physical memory, allowing us to fit more address spaces into physical memory.

However, segmentation raises a number of new issues. We’ll first describe the new OS issues that must be addressed. The first is an old one: what should the OS do on a context switch? You should have a good guess by now: the segment registers must be saved and restored. Clearly, each process has its own virtual address space, and the OS must make sure to set up these registers correctly before letting the process run again.

The second, and more important, issue is managing free space in physical memory. When a new address space is created, the OS has to be able to find space in physical memory for its segments. Previously, we assumed that each address space was the same size, and thus physical memory could be thought of as a bunch of slots where processes would fit in. Now, we have a number of segments per process, and each segment might be a different size.
The general problem that arises is that physical memory quickly becomes full of little holes of free space, making it difficult to allocate new segments, or to grow existing ones. We call this problem external fragmentation [R69]; see Figure 16.3 (left).

In the example, a process comes along and wishes to allocate a 20KB segment. In that example, there is 24KB free, but not in one contiguous segment (rather, in three non-contiguous chunks). Thus, the OS cannot satisfy the 20KB request.

One solution to this problem would be to compact physical memory by rearranging the existing segments. For example, the OS could stop whichever processes are running, copy their data to one contiguous region of memory, change their segment register values to point to the new physical locations, and thus have a large free extent of memory with which to work. By doing so, the OS enables the new allocation request to succeed. However, compaction is expensive, as copying segments is memory-intensive and thus would use a fair amount of processor time. See Figure 16.3 (right) for a diagram of compacted physical memory.

A simpler approach is to use a free-list management algorithm that tries to keep large extents of memory available for allocation. There are literally hundreds of approaches that people have taken, including classic algorithms like best-fit (which keeps a list of free spaces and returns the one closest in size that satisfies the desired allocation to the requester), worst-fit, first-fit, and more complex schemes like buddy algorithm [K68]. An excellent survey by Wilson et al. is a good place to start if you want to learn more about such algorithms [W+95], or you can wait until we cover some of the basics ourselves in a later chapter. Unfortunately, though, no matter how smart the algorithm, external fragmentation will still exist; thus, a good algorithm simply attempts to minimize it.
The fact that so many different algorithms exist to try to minimize external fragmentation is indicative of a stronger underlying truth: there is no one “best” way to solve the problem. Thus, we settle for something reasonable and hope it is good enough. The only real solution (as we will see in forthcoming chapters) is to avoid the problem altogether, by never allocating memory in variable-sized chunks.

### 16.7 Summary

Segmentation solves a number of problems, and helps us build a more effective virtualization of memory. Beyond just dynamic relocation, segmentation can better support sparse address spaces, by avoiding the huge potential waste of memory between logical segments of the address space. It is also fast, as doing the arithmetic segmentation requires in hardware is easy and well-suited to hardware; the overheads of translation are minimal. A fringe benefit arises too: code sharing. If code is placed within a separate segment, such a segment could potentially be shared across multiple running programs.

However, as we learned, allocating variable-sized segments in memory leads to some problems that we’d like to overcome. The first, as discussed above, is external fragmentation. Because segments are variable-sized, free memory gets chopped up into odd-sized pieces, and thus satisfying a memory-allocation request can be difficult. One can try to use smart algorithms [W+95] or periodically compact memory, but the problem is fundamental and hard to avoid.

The second and perhaps more important problem is that segmentation still isn’t flexible enough to support our fully generalized, sparse address space. For example, if we have a large but sparsely-used heap all in one logical segment, the entire heap must still reside in memory in order to be accessed. In other words, if our model of how the address space is being used doesn’t exactly match how the underlying segmentation has been designed to support it, segmentation doesn’t work very well. We thus need to find some new solutions. Ready to find them?
References

[CV65] “Introduction and Overview of the Multics System”  
F. J. Corbato and V. A. Vyssotsky  
Fall Joint Computer Conference, 1965  
One of five papers presented on Multics at the Fall Joint Computer Conference; oh to be a fly on the wall in that room that day!

[DD68] “Virtual Memory, Processes, and Sharing in Multics”  
Robert C. Daley and Jack B. Dennis  
An early paper on how to perform dynamic linking in Multics, which was way ahead of its time. Dynamic linking finally found its way back into systems about 20 years later, as the large X-windows libraries demanded it. Some say that these large X11 libraries were MIT’s revenge for removing support for dynamic linking in early versions of UNIX!

[G62] “Fact Segmentation”  
M. N. Greenfield  
Proceedings of the SJCC, Volume 21, May 1962  
Another early paper on segmentation; so early that it has no references to other work.

[H61] “Program Organization and Record Keeping for Dynamic Storage”  
A. W. Holt  
Communications of the ACM, Volume 4, Issue 10, October 1961  
An incredibly early and difficult to read paper about segmentation and some of its uses.

[I09] “Intel 64 and IA-32 Architectures Software Developer’s Manuals”  
Intel, 2009  
Try reading about segmentation in here (Chapter 3 in Volume 3a); it’ll hurt your head, at least a little bit.

Donald Knuth  
Addison-Wesley, 1968  
Knuth is famous not only for his early books on the Art of Computer Programming but for his typesetting system TeX which is still a powerhouse typesetting tool used by professionals today, and indeed to typeset this very book. His tomes on algorithms are a great early reference to many of the algorithms that underly computing systems today.

Butler Lampson  
ACM Operating Systems Review, 15:5, October 1983  
A treasure-trove of sage advice on how to build systems. Hard to read in one sitting; take it in a little at a time, like a fine wine, or a reference manual.

[LL82] “Virtual Memory Management in the VAX/VMS Operating System”  
Henry M. Levy and Peter H. Lipman  
IEEE Computer, Volume 15, Number 3 (March 1982)  
A classic memory management system, with lots of common sense in its design. We’ll study it in more detail in a later chapter.
B. Randell and C.J. Kuehner  
Communications of the ACM  
A nice overview of the differences between paging and segmentation, with some historical discussion of various machines.

[R69] “A note on storage fragmentation and program segmentation”  
Brian Randell  
Communications of the ACM  
Volume 12(7), pages 365-372, July 1969  
One of the earliest papers to discuss fragmentation.

Paul R. Wilson, Mark S. Johnstone, Michael Neely, and David Boles  
In International Workshop on Memory Management  
Scotland, United Kingdom, September 1995  
A great survey paper on memory allocators.
Homework

This program allows you to see how address translations are performed in a system with segmentation. See the README for details.

Questions

- First let’s use a tiny address space to translate some addresses. Here’s a simple set of parameters with a few different random seeds; can you translate the addresses?

  segmentation.py -a 128 -p 512 -b 0 -l 20 -B 512 -L 20 -s 0
  segmentation.py -a 128 -p 512 -b 0 -l 20 -B 512 -L 20 -s 1
  segmentation.py -a 128 -p 512 -b 0 -l 20 -B 512 -L 20 -s 2

- Now, let’s see if we understand this tiny address space we’ve constructed (using the parameters from the question above). What is the highest legal virtual address in segment 0? What about the lowest legal virtual address in segment 1? What are the lowest and highest illegal addresses in this entire address space? Finally, how would you run segmentation.py with the -A flag to test if you are right?

- Let’s say we have a tiny 16-byte address space in a 128-byte physical memory. What base and bounds would you set up so as to get the simulator to generate the following translation results for the specified address stream: valid, valid, violation, ..., violation, valid, valid? Assume the following parameters:

  segmentation.py -a 16 -p 128
  -A 0,1,2,3,4,5,6,7,8,9,10,11,12,13,14,15
  --b0 ? --l0 ? --b1 ? --l1 ?

- Assuming we want to generate a problem where roughly 90% of the randomly-generated virtual addresses are valid (i.e., not segmentation violations). How should you configure the simulator to do so? Which parameters are important?

- Can you run the simulator such that no virtual addresses are valid? How?
Free-Space Management

In this chapter, we take a small detour from our discussion of virtualizing memory to discuss a fundamental aspect of any memory management system, whether it be a malloc library (managing pages of a process’s heap) or the OS itself (managing portions of the address space of a process). Specifically, we will discuss the issues surrounding free-space management.

Let us make the problem more specific. Managing free space can certainly be easy, as we will see when we discuss the concept of paging. It is easy when the space you are managing is divided into fixed-sized units; in such a case, you just keep a list of these fixed-sized units; when a client requests one of them, return the first entry.

Where free-space management becomes more difficult (and interesting) is when the free space you are managing consists of variable-sized units; this arises in a user-level memory-allocation library (as in malloc() and free()) and in an OS managing physical memory when using segmentation to implement virtual memory. In either case, the problem that exists is known as external fragmentation: the free space gets chopped into little pieces of different sizes and is thus fragmented; subsequent requests may fail because there is no single contiguous space that can satisfy the request, even though the total amount of free space exceeds the size of the request.

The figure shows an example of this problem. In this case, the total free space available is 20 bytes; unfortunately, it is fragmented into two chunks of size 10 each. As a result, a request for 15 bytes will fail even though there are 20 bytes free. And thus we arrive at the problem addressed in this chapter.
CRUX: HOW TO MANAGE FREE SPACE
How should free space be managed, when satisfying variable-sized requests? What strategies can be used to minimize fragmentation? What are the time and space overheads of alternate approaches?

17.1 Assumptions

Most of this discussion will focus on the great history of allocators found in user-level memory-allocation libraries. We draw on Wilson's excellent survey [W+95] but encourage interested readers to go to the source document itself for more details.

We assume a basic interface such as that provided by malloc() and free(). Specifically, void *malloc(size_t size) takes a single parameter, size, which is the number of bytes requested by the application; it hands back a pointer (of no particular type, or a void pointer in C lingo) to a region of that size (or greater). The complementary routine void free(void *ptr) takes a pointer and frees the corresponding chunk. Note the implication of the interface: the user, when freeing the space, does not inform the library of its size; thus, the library must be able to figure out how big a chunk of memory is when handed just a pointer to it. We’ll discuss how to do this a bit later on in the chapter.

The space that this library manages is known historically as the heap, and the generic data structure used to manage free space in the heap is some kind of free list. This structure contains references to all of the free chunks of space in the managed region of memory. Of course, this data structure need not be a list per se, but just some kind of data structure to track free space.

We further assume that primarily we are concerned with external fragmentation, as described above. Allocators could of course also have the problem of internal fragmentation; if an allocator hands out chunks of memory bigger than that requested, any unasked for (and thus unused) space in such a chunk is considered internal fragmentation (because the waste occurs inside the allocated unit) and is another example of space waste. However, for the sake of simplicity, and because it is the more interesting of the two types of fragmentation, we’ll mostly focus on external fragmentation.

We’ll also assume that once memory is handed out to a client, it cannot be relocated to another location in memory. For example, if a program calls malloc() and is given a pointer to some space within the heap, that memory region is essentially “owned” by the program (and cannot be moved by the library) until the program returns it via a corresponding call to free(). Thus, no compaction of free space is possible, which

1It is nearly 80 pages long; thus, you really have to be interested!
would be useful to combat fragmentation\(^2\). Compaction could, however, be used in the OS to deal with fragmentation when implementing segmentation; see the chapter on segmentation for details.

Finally, we’ll assume that the allocator manages a contiguous region of bytes. In some cases, an allocator could ask for that region to grow; for example, a user-level memory-allocation library might call into the kernel to grow the heap (via a system call such as `sbrk`) when it runs out of space. However, for simplicity, we’ll just assume that the region is a single fixed size throughout its life.

### 17.2 Low-level Mechanisms

Before delving into some policy details, we’ll first cover some common mechanisms used in most allocators. First, we’ll discuss the basics of splitting and coalescing, common techniques in most any allocator. Second, we’ll show how one can track the size of allocated regions quickly and with relative ease. Finally, we’ll discuss how to build a simple list inside the free space to keep track of what is free and what isn’t.

#### Splitting and Coalescing

A free list contains a set of elements that describe the free space still remaining in the heap. Thus, assume the following 30-byte heap:

\[
\begin{array}{ccc}
\text{free} & \text{used} & \text{free} \\
0 & 10 & 20 & 30
\end{array}
\]

The free list for this heap would have two elements on it. One entry describes the first 10-byte free segment (bytes 0-9), and one entry describes the other free segment (bytes 20-29):

```
head → addr:0 len:10 → addr:20 len:10 → NULL
```

As described above, a request for anything greater than 10 bytes will fail (returning NULL); there just isn’t a single contiguous chunk of memory of that size available. A request for exactly that size (10 bytes) could be satisfied easily by either of the free chunks. But what happens if the request is for something smaller than 10 bytes?

Assume we have a request for just a single byte of memory. In this case, the allocator will perform an action known as splitting: it will find

\(^2\)Once you hand a pointer to a chunk of memory to a C program, it is generally difficult to determine all references (pointers) to that region, which may be stored in other variables or even in registers at a given point in execution. This may not be the case in more strongly-typed, garbage-collected languages, which would thus enable compaction as a technique to combat fragmentation.
a free chunk of memory that can satisfy the request and split it into two. The first chunk it will return to the caller; the second chunk will remain on the list. Thus, in our example above, if a request for 1 byte were made, and the allocator decided to use the second of the two elements on the list to satisfy the request, the call to malloc() would return 20 (the address of the 1-byte allocated region) and the list would end up looking like this:

```
head -> addr:0 len:10 -> addr:21 len:9 -> NULL
```

In the picture, you can see the list basically stays intact; the only change is that the free region now starts at 21 instead of 20, and the length of that free region is now just 9\(^3\). Thus, the split is commonly used in allocators when requests are smaller than the size of any particular free chunk.

A corollary mechanism found in many allocators is known as **coalescing** of free space. Take our example from above once more (free 10 bytes, used 10 bytes, and another free 10 bytes).

Given this (tiny) heap, what happens when an application calls free(10), thus returning the space in the middle of the heap? If we simply add this free space back into our list without too much thinking, we might end up with a list that looks like this:

```
head -> addr:10 len:10 -> addr:0 len:10 -> addr:20 len:10 -> NULL
```

Note the problem: while the entire heap is now free, it is seemingly divided into three chunks of 10 bytes each. Thus, if a user requests 20 bytes, a simple list traversal will not find such a free chunk, and return failure.

What allocators do in order to avoid this problem is coalesce free space when a chunk of memory is freed. The idea is simple: when returning a free chunk in memory, look carefully at the addresses of the chunk you are returning as well as the nearby chunks of free space; if the newly-freed space sits right next to one (or two, as in this example) existing free chunks, merge them into a single larger free chunk. Thus, with coalescing, our final list should look like this:

```
head -> addr:0 len:30 -> NULL
```

Indeed, this is what the heap list looked like at first, before any allocations were made. With coalescing, an allocator can better ensure that large free extents are available for the application.

---

\(^3\)This discussion assumes that there are no headers, an unrealistic but simplifying assumption we make for now.
The header used by malloc library
The 20 bytes returned to caller

Figure 17.1: An Allocated Region Plus Header

ptr

The 20 bytes returned to caller

Figure 17.2: Specific Contents Of The Header

Tracking The Size Of Allocated Regions

You might have noticed that the interface to `free(void *ptr)` does not take a size parameter; thus it is assumed that given a pointer, the malloc library can quickly determine the size of the region of memory being freed and thus incorporate the space back into the free list.

To accomplish this task, most allocators store a little bit of extra information in a header block which is kept in memory, usually just before the handed-out chunk of memory. Let’s look at an example again (Figure 17.1). In this example, we are examining an allocated block of size 20 bytes, pointed to by `ptr`; imagine the user called `malloc()` and stored the results in `ptr`, e.g., `ptr = malloc(20);`.

The header minimally contains the size of the allocated region (in this case, 20); it may also contain additional pointers to speed up deallocation, a magic number to provide additional integrity checking, and other information. Let’s assume a simple header which contains the size of the region and a magic number, like this:

```c
typedef struct __header_t {
    int size;
    int magic;
} header_t;
```

The example above would look like what you see in Figure 17.2. When
the user calls `free(ptr)`, the library then uses simple pointer arithmetic to figure out where the header begins:

```c
void free(void *ptr) {
    header_t *hptr = (void *)ptr - sizeof(header_t);
}
```

After obtaining such a pointer to the header, the library can easily determine whether the magic number matches the expected value as a sanity check (`assert(hptr->magic == 1234567)`) and calculate the total size of the newly-freed region via simple math (i.e., adding the size of the header to size of the region). Note the small but critical detail in the last sentence: the size of the free region is the size of the header plus the size of the space allocated to the user. Thus, when a user requests $N$ bytes of memory, the library does not search for a free chunk of size $N$; rather, it searches for a free chunk of size $N$ plus the size of the header.

**Embedding A Free List**

Thus far we have treated our simple free list as a conceptual entity; it is just a list describing the free chunks of memory in the heap. But how do we build such a list inside the free space itself?

In a more typical list, when allocating a new node, you would just call `malloc()` when you need space for the node. Unfortunately, within the memory-allocation library, you can’t do this! Instead, you need to build the list `inside` the free space itself. Don’t worry if this sounds a little weird; it is, but not so weird that you can’t do it!

Assume we have a 4096-byte chunk of memory to manage (i.e., the heap is 4KB). To manage this as a free list, we first have to initialize said list; initially, the list should have one entry, of size 4096 (minus the header size). Here is the description of a node of the list:

```c
typedef struct __node_t {
    int size;
    struct __node_t *next;
} node_t;
```

Now let’s look at some code that initializes the heap and puts the first element of the free list inside that space. We are assuming that the heap is built within some free space acquired via a call to the system call `mmap()`; this is not the only way to build such a heap but serves us well in this example. Here is the code:

```c
// mmap() returns a pointer to a chunk of free space
node_t *head = mmap(NULL, 4096, PROT_READ|PROT_WRITE, MAP_ANON|MAP_PRIVATE, -1, 0);
head->size = 4096 - sizeof(node_t);
head->next = NULL;
```

After running this code, the status of the list is that it has a single entry, of size 4088. Yes, this is a tiny heap, but it serves as a fine example for us.
here. The head pointer contains the beginning address of this range; let’s assume it is 16KB (though any virtual address would be fine). Visually, the heap thus looks like what you see in Figure 17.3.

Now, let’s imagine that a chunk of memory is requested, say of size 100 bytes. To service this request, the library will first find a chunk that is large enough to accommodate the request; because there is only one free chunk (size: 4088), this chunk will be chosen. Then, the chunk will be split into two: one chunk big enough to service the request (and header, as described above), and the remaining free chunk. Assuming an 8-byte header (an integer size and an integer magic number), the space in the heap now looks like what you see in Figure 17.4.

Thus, upon the request for 100 bytes, the library allocated 108 bytes out of the existing one free chunk, returns a pointer (marked ptr in the figure above) to it, stashes the header information immediately before the
allocated space for later use upon `free()`, and shrinks the one free node in the list to 3980 bytes (4088 minus 108).

Now let’s look at the heap when there are three allocated regions, each of 100 bytes (or 108 including the header). A visualization of this heap is shown in Figure 17.5.

As you can see therein, the first 324 bytes of the heap are now allocated, and thus we see three headers in that space as well as three 100-byte regions being used by the calling program. The free list remains uninteresting: just a single node (pointed to by `head`), but now only 3764 bytes in size after the three splits. But what happens when the calling program returns some memory via `free()`?
In this example, the application returns the middle chunk of allocated memory, by calling `free(16500)` (the value 16500 is arrived upon by adding the start of the memory region, 16384, to the 108 of the previous chunk and the 8 bytes of the header for this chunk). This value is shown in the previous diagram by the pointer `sptr`.

The library immediately figures out the size of the free region, and then adds the free chunk back onto the free list. Assuming we insert at the head of the free list, the space now looks like this (Figure 17.6).

And now we have a list that starts with a small free chunk (100 bytes, pointed to by the head of the list) and a large free chunk (3764 bytes).
Our list finally has more than one element on it! And yes, the free space is fragmented, an unfortunate but common occurrence.

One last example: let’s assume now that the last two in-use chunks are freed. Without coalescing, you might end up with a free list that is highly fragmented (see Figure 17.7).

As you can see from the figure, we now have a big mess! Why? Simple, we forgot to **coalesce** the list. Although all of the memory is free, it is chopped up into pieces, thus appearing as a fragmented memory despite not being one. The solution is simple: go through the list and **merge** neighboring chunks; when finished, the heap will be whole again.

---

**Figure 17.7: A Non-Coalesced Free List**
Growing The Heap

We should discuss one last mechanism found within many allocation libraries. Specifically, what should you do if the heap runs out of space? The simplest approach is just to fail. In some cases this is the only option, and thus returning NULL is an honorable approach. Don’t feel bad! You tried, and though you failed, you fought the good fight.

Most traditional allocators start with a small-sized heap and then request more memory from the OS when they run out. Typically, this means they make some kind of system call (e.g., sbrk in most UNIX systems) to grow the heap, and then allocate the new chunks from there. To service the sbrk request, the OS finds free physical pages, maps them into the address space of the requesting process, and then returns the value of the end of the new heap; at that point, a larger heap is available, and the request can be successfully serviced.

17.3 Basic Strategies

Now that we have some machinery under our belt, let’s go over some basic strategies for managing free space. These approaches are mostly based on pretty simple policies that you could think up yourself; try it before reading and see if you come up with all of the alternatives (or maybe some new ones!).

The ideal allocator is both fast and minimizes fragmentation. Unfortunately, because the stream of allocation and free requests can be arbitrary (after all, they are determined by the programmer), any particular strategy can do quite badly given the wrong set of inputs. Thus, we will not describe a “best” approach, but rather talk about some basics and discuss their pros and cons.

Best Fit

The best fit strategy is quite simple: first, search through the free list and find chunks of free memory that are as big or bigger than the requested size. Then, return the one that is the smallest in that group of candidates; this is the so called best-fit chunk (it could be called smallest fit too). One pass through the free list is enough to find the correct block to return.

The intuition behind best fit is simple: by returning a block that is close to what the user asks, best fit tries to reduce wasted space. However, there is a cost; naive implementations pay a heavy performance penalty when performing an exhaustive search for the correct free block.

Worst Fit

The worst fit approach is the opposite of best fit; find the largest chunk and return the requested amount; keep the remaining (large) chunk on the free list. Worst fit tries to thus leave big chunks free instead of lots of
small chunks that can arise from a best-fit approach. Once again, however, a full search of free space is required, and thus this approach can be costly. Worse, most studies show that it performs badly, leading to excess fragmentation while still having high overheads.

First Fit

The first fit method simply finds the first block that is big enough and returns the requested amount to the user. As before, the remaining free space is kept free for subsequent requests.

First fit has the advantage of speed – no exhaustive search of all the free spaces are necessary – but sometimes pollutes the beginning of the free list with a small objects. Thus, how the allocator manages the free list’s order becomes an issue. One approach is to use address-based ordering; by keeping the list ordered by the address of the free space, coalescing becomes easier, and fragmentation tends to be reduced.

Next Fit

Instead of always beginning the first-fit search at the beginning of the list, the next fit algorithm keeps an extra pointer to the location within the list where one was looking last. The idea is to spread the searches for free space throughout the list more uniformly, thus avoiding splintering of the beginning of the list. The performance of such an approach is quite similar to first fit, as an exhaustive search is once again avoided.

Examples

Here are a few examples of the above strategies. Envision a free list with three elements on it, of sizes 10, 30, and 20 (we’ll ignore headers and other details here, instead just focusing on how strategies operate):

```
head → 10 → 30 → 20 → NULL
```

Assume an allocation request of size 15. A best-fit approach would search the entire list and find that 20 was the best fit, as it is the smallest free space that can accommodate the request. The resulting free list:

```
head → 10 → 30 → 5 → NULL
```

As happens in this example, and often happens with a best-fit approach, a small free chunk is now left over. A worst-fit approach is similar but instead finds the largest chunk, in this example 30. The resulting list:

```
head → 10 → 15 → 20 → NULL
```
The first-fit strategy, in this example, does the same thing as worst-fit, also finding the first free block that can satisfy the request. The difference is in the search cost; both best-fit and worst-fit look through the entire list; first-fit only examines free chunks until it finds one that fits, thus reducing search cost.

These examples just scratch the surface of allocation policies. More detailed analysis with real workloads and more complex allocator behaviors (e.g., coalescing) are required for a deeper understanding. Perhaps something for a homework section, you say?

17.4 Other Approaches

Beyond the basic approaches described above, there have been a host of suggested techniques and algorithms to improve memory allocation in some way. We list a few of them here for your consideration (i.e., to make you think about a little more than just best-fit allocation).

Segregated Lists

One interesting approach that has been around for some time is the use of segregated lists. The basic idea is simple: if a particular application has one (or a few) popular-sized request that it makes, keep a separate list just to manage objects of that size; all other requests are forwarded to a more general memory allocator.

The benefits of such an approach are obvious. By having a chunk of memory dedicated for one particular size of requests, fragmentation is much less of a concern; moreover, allocation and free requests can be served quite quickly when they are of the right size, as no complicated search of a list is required.

Just like any good idea, this approach introduces new complications into a system as well. For example, how much memory should one dedicate to the pool of memory that serves specialized requests of a given size, as opposed to the general pool? One particular allocator, the slab allocator by uber-engineer Jeff Bonwick (which was designed for use in the Solaris kernel), handles this issue in a rather nice way [B94].

Specifically, when the kernel boots up, it allocates a number of object caches for kernel objects that are likely to be requested frequently (such as locks, file-system inodes, etc.); the object caches thus are each segregated free lists of a given size and serve memory allocation and free requests quickly. When a given cache is running low on free space, it requests some slabs of memory from a more general memory allocator (the total amount requested being a multiple of the page size and the object in question). Conversely, when the reference counts of the objects within a given slab all go to zero, the general allocator can reclaim them from the specialized allocator, which is often done when the VM system needs more memory.
**ASIDE: GREAT ENGINEERS ARE REALLY GREAT**

Engineers like Jeff Bonwick (who not only wrote the slab allocator mentioned herein but also was the lead of an amazing file system, ZFS) are the heart of Silicon Valley. Behind almost any great product or technology is a human (or small group of humans) who are way above average in their talents, abilities, and dedication. As Mark Zuckerberg (of Facebook) says: “Someone who is exceptional in their role is not just a little better than someone who is pretty good. They are 100 times better.” This is why, still today, one or two people can start a company that changes the face of the world forever (think Google, Apple, or Facebook). Work hard and you might become such a “100x” person as well. Failing that, work with such a person; you’ll learn more in a day than most learn in a month. Failing that, feel sad.

The slab allocator also goes beyond most segregated list approaches by keeping free objects on the lists in a pre-initialized state. Bonwick shows that initialization and destruction of data structures is costly [B94]; by keeping freed objects in a particular list in their initialized state, the slab allocator thus avoids frequent initialization and destruction cycles per object and thus lowers overheads noticeably.

**Buddy Allocation**

Because coalescing is critical for an allocator, some approaches have been designed around making coalescing simple. One good example is found in the **binary buddy allocator** [K65].

In such a system, free memory is first conceptually thought of as one big space of size $2^N$. When a request for memory is made, the search for free space recursively divides free space by two until a block that is big enough to accommodate the request is found (and a further split into two would result in a space that is too small). At this point, the requested block is returned to the user. Here is an example of a 64KB free space getting divided in the search for a 7KB block:
In the example, the leftmost 8KB block is allocated (as indicated by the darker shade of gray) and returned to the user; note that this scheme can suffer from internal fragmentation, as you are only allowed to give out power-of-two-sized blocks.

The beauty of buddy allocation is found in what happens when that block is freed. When returning the 8KB block to the free list, the allocator checks whether the “buddy” 8KB is free; if so, it coalesces the two blocks into a 16KB block. The allocator then checks if the buddy of the 16KB block is still free; if so, it coalesces those two blocks. This recursive coalescing process continues up the tree, either restoring the entire free space or stopping when a buddy is found to be in use.

The reason buddy allocation works so well is that it is simple to determine the buddy of a particular block. How, you ask? Think about the addresses of the blocks in the free space above. If you think carefully enough, you’ll see that the address of each buddy pair only differs by a single bit; which bit is determined by the level in the buddy tree. And thus you have a basic idea of how binary buddy allocation schemes work. For more detail, as always, see the Wilson survey [W+95].

Other Ideas

One major problem with many of the approaches described above is their lack of scaling. Specifically, searching lists can be quite slow. Thus, advanced allocators use more complex data structures to address these costs, trading simplicity for performance. Examples include balanced binary trees, splay trees, or partially-ordered trees [W+95].

Given that modern systems often have multiple processors and run multi-threaded workloads (something you’ll learn about in great detail in the section of the book on Concurrency), it is not surprising that a lot of effort has been spent making allocators work well on multiprocessor-based systems. Two wonderful examples are found in Berger et al. [B+00] and Evans [E06]; check them out for the details.

These are but two of the thousands of ideas people have had over time about memory allocators. Read on your own if you are curious.

17.5 Summary

In this chapter, we’ve discussed the most rudimentary forms of memory allocators. Such allocators exist everywhere, linked into every C program you write, as well as in the underlying OS which is managing memory for its own data structures. As with many systems, there are many trade-offs to be made in building such a system, and the more you know about the exact workload presented to an allocator, the more you could do to tune it to work better for that workload. Making a fast, space-efficient, scalable allocator that works well for a broad range of workloads remains an on-going challenge in modern computer systems.

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References

[B+00] “Hoard: A Scalable Memory Allocator for Multithreaded Applications”
Emery D. Berger, Kathryn S. McKinley, Robert D. Blumofe, and Paul R. Wilson
ASPLOS-IX, November 2000
Berger and company’s excellent allocator for multiprocessor systems. Beyond just being a fun paper, also used in practice!

Jeff Bonwick
USENIX ’94
A cool paper about how to build an allocator for an operating system kernel, and a great example of how to specialize for particular common object sizes.

[E06] “A Scalable Concurrent malloc(3) Implementation for FreeBSD”
Jason Evans
April 2006
A detailed look at how to build a real modern allocator for use in multiprocessors. The “jemalloc” allocator is in widespread use today, within FreeBSD, NetBSD, Mozilla Firefox, and within Facebook.

[K65] “A Fast Storage Allocator”
Kenneth C. Knowlton
Communications of the ACM, Volume 8, Number 10, October 1965
The common reference for buddy allocation. Random strange fact: Knuth gives credit for the idea to not to Knouelon but to Harry Markowitz, a Nobel-prize winning economist. Another strange fact: Knuth communicates all of his emails via a secretary; he doesn’t send email himself, rather he tells his secretary what email to send and then the secretary does the work of emailing. Last Knuth fact: he created TeX, the tool used to typeset this book. It is an amazing piece of software.

Paul R. Wilson, Mark S. Johnstone, Michael Neely, David Boles
International Workshop on Memory Management
Kinross, Scotland, September 1995
An excellent and far-reaching survey of many facets of memory allocation. Far too much detail to go into in this tiny chapter!

Actually we use LaTeX, which is based on Lamport’s additions to TeX, but close enough.
Remember our goal: to virtualize memory. Segmentation (a generalization of dynamic relocation) helped us do this, but has some problems; in particular, managing free space becomes quite a pain as memory becomes fragmented and segmentation is not as flexible as we might like. Is there a better solution?

**THE CRUX:**
**HOW TO VIRTUALIZE MEMORY WITHOUT SEGMENTS**

How can we virtualize memory in a way as to avoid the problems of segmentation? What are the basic techniques? How do we make those techniques work well?

Thus comes along the idea of **paging**, which goes back to the earliest of computer systems, namely the Atlas [KE+62,L78]. Instead of splitting up our address space into three logical segments (each of variable size), we split up our address space into fixed-sized units we call a **page**. Here in Figure 18.1 an example of a tiny address space, only 64 bytes total in size, with 16 byte pages (real address spaces are much bigger, of course, commonly 32 bits and thus 4-GB of address space, or even 64 bits). We’ll use tiny examples to make them easier to digest (at first).

![Figure 18.1: A Simple 64-byte Address Space](image)

(page 0 of the address space)

(page 1)

(page 2)

(page 3)
Thus, we have an address space that is split into four pages (0 through 3). With paging, physical memory is also split into some number of pages as well; we sometimes will call each page of physical memory a page frame. For an example, let’s examine Figure 18.2.

Paging, as we will see, has a number of advantages over our previous approaches. Probably the most important improvement will be flexibility: with a fully-developed paging approach, the system will be able to support the abstraction of an address space effectively, regardless of how the processes uses the address space; we won’t, for example, have to make assumptions about how the heap and stack grow and how they are used.

Another advantage is the simplicity of free-space management that paging affords. For example, when the OS wishes to place our tiny 64-byte address space from above into our 8-page physical memory, it simply finds four free pages; perhaps the OS keeps a free list of all free pages for this, and just grabs the first four free pages off of this list. In the example above, the OS has placed virtual page 0 of the address space (AS) in physical page 3, virtual page 1 of the AS on physical page 7, page 2 on page 5, and page 3 on page 2.

To record where each virtual page of the address space is placed in physical memory, the operating system keeps a per-process data structure known as a page table. The major role of the page table is to store address translations for each of the virtual pages of the address space, thus letting us know where in physical memory they live. For our simple example above (Figure 18.2), the page table would thus have the following entries: (Virtual Page 0 → Physical Frame 3), (VP 1 → PF 7), (VP 2 → PF 5), and (VP 3 → PF 2).

It is important to remember that this page table is a per-process data structure (most page table structures we discuss are per-process structures; an exception we’ll touch on is the inverted page table). If another

---

**Figure 18.2: 64-Byte Address Space Placed In Physical Memory**

<table>
<thead>
<tr>
<th>0</th>
<th>reserved for OS</th>
</tr>
</thead>
<tbody>
<tr>
<td>16</td>
<td>page frame 0 of physical memory</td>
</tr>
<tr>
<td>32</td>
<td>page frame 1</td>
</tr>
<tr>
<td>48</td>
<td>page frame 2</td>
</tr>
<tr>
<td>64</td>
<td>page 3 of AS</td>
</tr>
<tr>
<td>80</td>
<td>page frame 3</td>
</tr>
<tr>
<td>96</td>
<td>page frame 4</td>
</tr>
<tr>
<td>112</td>
<td>page frame 5</td>
</tr>
<tr>
<td>128</td>
<td>page frame 6</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>160</th>
<th>page 2 of AS</th>
</tr>
</thead>
<tbody>
<tr>
<td>176</td>
<td>page frame 7</td>
</tr>
<tr>
<td>192</td>
<td>page 1 of AS</td>
</tr>
</tbody>
</table>
process were to run in our example above, the OS would have to manage a different page table for it, as its virtual pages obviously map to different physical pages (modulo any sharing going on).

Now, we know enough to perform an address-translation example. Let’s imagine the process with that tiny address space (64 bytes) is performing a memory access:

```assembly
movl <virtual address>, %eax
```

Specifically, let’s pay attention to the explicit load of the data at `<virtual address>` into the register eax (and thus ignore the instruction fetch that must have happened prior).

To translate this virtual address that the process generated, we have to first split it into two components: the virtual page number (VPN), and the offset within the page. For this example, because the virtual address space of the process is 64 bytes, we need 6 bits total for our virtual address \(2^6 = 64\). Thus, our virtual address:

```
| Va5 | Va4 | Va3 | Va2 | Va1 | Va0 |
```

where Va5 is the highest-order bit of the virtual address, and Va0 the lowest order bit. Because we know the page size (16 bytes), we can further divide the virtual address as follows:

```
VPN offset
+----------+----------+
| Va5 | Va4 | Va3 | Va2 | Va1 | Va0 |
```

The page size is 16 bytes in a 64-byte address space; thus we need to be able to select 4 pages, and the top 2 bits of the address do just that. Thus, we have a 2-bit virtual page number (VPN). The remaining bits tell us which byte of the page we are interested in, 4 bits in this case; we call this the offset.

When a process generates a virtual address, the OS and hardware must combine to translate it into a meaningful physical address. For example, let us assume the load above was to virtual address 21:

```assembly
movl 21, %eax
```

Turning “21” into binary form, we get “010101”, and thus we can examine this virtual address and see how it breaks down into a virtual page number (VPN) and offset:

```
VPN offset
+----------+----------+
| 0 | 1 | 0 | 1 | 0 | 1 |
```
Thus, the virtual address “21” is on the 5th (“0101”th) byte of virtual page “01” (or 1). With our virtual page number, we can now index our page table and find which physical page that virtual page 1 resides within. In the page table above the physical page number (PPN) (a.k.a. physical frame number or PFN) is 7 (binary 111). Thus, we can translate this virtual address by replacing the VPN with the PFN and then issue the load to physical memory (Figure 18.3).

Note the offset stays the same (i.e., it is not translated), because the offset just tells us which byte within the page we want. Our final physical address is 1110101 (117 in decimal), and is exactly where we want our load to fetch data from (Figure 18.2).

18.1 Where Are Page Tables Stored?

Page tables can get awfully large, much bigger than the small segment table or base/bounds pair we have discussed previously. For example, imagine a typical 32-bit address space, with 4-KB pages. This virtual address splits into a 20-bit VPN and 12-bit offset (recall that 10 bits would be needed for a 1-KB page size, and just add two more to get to 4 KB).

A 20-bit VPN implies that there are $2^{20}$ translations that the OS would have to manage for each process (that’s roughly a million); assuming we need 4 bytes per page table entry (PTE) to hold the physical translation plus any other useful stuff, we get an immense 4MB of memory needed for each page table! That is pretty big. Now imagine there are 100 processes running: this means the OS would need 400MB of memory just for all those address translations! Even in the modern era, where machines have gigabytes of memory, it seems a little crazy to use a large chunk of if just for translations, no?

Because page tables are so big, we don’t keep any special on-chip hardware in the MMU to store the page table of the currently-running process. Instead, we store the page table for each process in memory somewhere.
Let’s assume for now that the page tables live in physical memory that the OS manages. In Figure 18.4 is a picture of what that might look like.

18.2 What’s Actually In The Page Table?

Let’s talk a little about page table organization. The page table is just a data structure that is used to map virtual addresses (or really, virtual page numbers) to physical addresses (physical page numbers). Thus, any data structure could work. The simplest form is called a linear page table, which is just an array. The OS indexes the array by the VPN, and looks up the page-table entry (PTE) at that index in order to find the desiredPFN. For now, we will assume this simple linear structure; in later chapters, we will make use of more advanced data structures to help solve some problems with paging.

As for the contents of each PTE, we have a number of different bits in there worth understanding at some level. A valid bit is common to indicate whether the particular translation is valid; for example, when a program starts running, it will have code and heap at one end of its address space, and the stack at the other. All the unused space in-between will be marked invalid, and if the process tries to access such memory, it will generate a trap to the OS which will likely terminate the process. Thus, the valid bit is crucial for supporting a sparse address space; by simply marking all the unused pages in the address space invalid, we remove the need to allocate physical frames for those pages and thus save a great deal of memory.

We also might have protection bits, indicating whether the page could be read from, written to, or executed from. Again, accessing a page in a way not allowed by these bits will generate a trap to the OS.
Figure 18.5: An x86 Page Table Entry (PTE)

There are a couple of other bits that are important but we won’t talk about much for now. A present bit indicates whether this page is in physical memory or on disk (swapped out); we will understand this in more detail when we study how to move parts of the address space to disk and back in order to support address spaces that are larger than physical memory and allow for the pages of processes that aren’t actively being run to be swapped out. A dirty bit is also common, indicating whether the page has been modified since it was brought into memory.

A reference bit (a.k.a. accessed bit) is sometimes used to track whether a page has been accessed, and is useful in determining which pages are popular and thus should be kept in memory; such knowledge is critical during page replacement, a topic we will study in great detail in subsequent chapters.

Figure 18.5 shows an example page table entry from the x86 architecture [I09]. It contains a present bit (P); a read/write bit (R/W) which determines if writes are allowed to this page; a user/ supervisor bit (U/S) which determines if user-mode processes can access the page; a few bits (PWT, PCD, PAT, and G) that determine how hardware caching works for these pages; an accessed bit (A) and a dirty bit (D); and finally, the page frame number (PFN) itself.

Read the Intel Architecture Manuals [I09] for more details on x86 paging support. Be forewarned, however; reading manuals such as these, while quite informative (and certainly necessary for those who write code to use such page tables in the OS), can be challenging at first. A little patience, and a lot of desire, is required.

18.3 Paging: Also Too Slow

With page tables in memory, we already know that they might be too big. Turns out they can slow things down too. For example, take our simple instruction:

```
movl 21, %eax
```

Again, let’s just examine the explicit reference to address 21 and not worry about the instruction fetch. In this example, we will assume the hardware performs the translation for us. To fetch the desired data, the system must first translate the virtual address (21) into the correct physical address (117). Thus, before issuing the load to address 117, the system must first fetch the proper page table entry from the process’s page table, perform the translation, and then finally get the desired data from physical memory.
To do so, the hardware must know where the page table is for the currently-running process. Let’s assume for now that a single page-table base register contains the physical address of the starting location of the page table. To find the location of the desired PTE, the hardware will thus perform the following functions:

\[
\text{VPN} = (\text{VirtualAddress} \& \text{VPN}\_\text{MASK}) >> \text{SHIFT} \\
\text{PTEAddr} = \text{PageTableBaseRegister} + (\text{VPN} \times \text{sizeof(PTE)})
\]

In our example, VPN\_MASK would be set to 0x30 (hex 30, or binary 110000) which picks out the VPN bits from the full virtual address; SHIFT is set to 4 (the number of bits in the offset), such that we move the VPN bits down to form the correct integer virtual page number. For example, with virtual address 21 (010101), and masking turns this value into 010000; the shift turns it into 01, or virtual page 1, as desired. We then use this value as an index into the array of PTEs pointed to by the page table base register.

Once this physical address is known, the hardware can fetch the PTE from memory, extract the PFN, and concatenate it with the offset from the virtual address to form the desired physical address. Specifically, you can think of the PFN being left-shifted by SHIFT, and then logically OR’d with the offset to form the final address as follows:

\[
\text{offset} = \text{VirtualAddress} \& \text{OFFSET}\_\text{MASK} \\
\text{PhysAddr} = (\text{PFN} << \text{PFN}\_\text{SHIFT}) | \text{offset}
\]

Figure 18.6: Accessing Memory With Paging

Finally, the hardware can fetch the desired data from memory and put it into register eax. The program has now succeeded at loading a value from memory!
ASIDE: DATA STRUCTURE – THE PAGE TABLE
One of the most important data structures in the memory management subsystem of a modern OS is the **page table**. In general, a page table stores **virtual-to-physical address translations**, thus letting the system know where each page of an address space actually resides in physical memory. Because each address space requires such translations, in general there is one page table per process in the system. The exact structure of the page table is either determined by the hardware (older systems) or can be more flexibly managed by the OS (modern systems).

To summarize, we now describe the initial protocol for what happens on each memory reference. Figure 18.6 shows the basic approach. For every memory reference (whether an instruction fetch or an explicit load or store), paging requires us to perform one extra memory reference in order to first fetch the translation from the page table. That is a lot of work! Extra memory references are costly, and in this case will likely slow down the process by a factor of two or more.

And now you can hopefully see that there are two real problems that we must solve. Without careful design of both hardware and software, page tables will cause the system to run too slowly, as well as take up too much memory. While seemingly a great solution for our memory virtualization needs, these two crucial problems must first be overcome.

18.4 A Memory Trace

Before closing, we now trace through a simple memory access example to demonstrate all of the resulting memory accesses that occur when using paging. The code snippet (in C, in a file called `array.c`) that are interested in is as follows:

```c
int array[1000];
...
for (i = 0; i < 1000; i++)
    array[i] = 0;
```

We could then compile `array.c` and run it with the following commands:

```
prompt> gcc -o array array.c -Wall -O
prompt> ./array
```

Of course, to truly understand what memory accesses this code snippet (which simply initializes an array) will make, we’ll have to know (or assume) a few more things. First, we’ll have to **disassemble** the resulting binary (using `objdump` on Linux, or `otool` on a Mac) to see what assembly instructions are used to initialize the array in a loop. Here it the resulting assembly code:
The code, if you know a little x86, is actually quite easy to understand. The first instruction moves the value zero (shown as $0x0) into the virtual memory address of the location of the array; this address is computed by taking the contents of %edi and adding %eax multiplied by four to it. Thus, %edi holds the base address of the array, whereas %eax holds the array index (i); we multiply by four because the array is an array of integers, each size four bytes (note we are cheating a little bit here, assuming each instruction is four bytes in size for simplicity; in actuality, x86 instructions are variable-sized).

The second instruction increments the array index held in %eax, and the third instruction compares the contents of that register to the hex value 0x03e8, or decimal 1000. If the comparison shows that that two values are not yet equal (which is what the jne instruction tests), the fourth instruction jumps back to the top of the loop.

To understand which memory accesses this instruction sequence makes (at both the virtual and physical levels), we’ll have assume something about where in virtual memory the code snippet and array are found, as well as the contents and location of the page table.

For this example, we assume a virtual address space of size 64 KB (unrealistically small). We also assume a page size of 1 KB.

All we need to know now are the contents of the page table, and its location in physical memory. Let’s assume we have a linear (array-based) page table and that it is located at physical address 1 KB (1024).

As for its contents, there are just a few virtual pages we need to worry about having mapped for this example. First, there is the virtual page the code lives on. Because the page size is 1 KB, virtual address 1024 resides on the the second page of the virtual address space (VPN=1, as VPN=0 is the first page). Let’s assume this virtual page maps to physical frame 4 (VPN 1 → PFN 4).

Next, there is the array itself. Its size is 4000 bytes (1000 integers), and it lives at virtual addresses 40000 through 44000 (not including the last byte). The virtual pages for this decimal range is VPN=39 ... VPN=42. Thus, we need mappings for these pages. Let’s assume these virtual-to-physical mappings for the example: (VPN 39 → PFN 7), (VPN 40 → PFN 8), (VPN 41 → PFN 9), (VPN 42 → PFN 10).

We are now ready to trace the memory references of the program. When it runs, each instruction fetch will generate two memory references: one to the page table to find the physical frame that the instruction resides within, and one to the instruction itself to fetch it to the CPU for processing. In addition, there is one explicit memory reference in the form of the mov instruction; this adds another page table access first (to translate the array virtual address to the correct physical one) and then the array access itself.
The entire process, for the first five loop iterations, is depicted in Figure 18.7. The bottom most graph shows the instruction memory references on the y-axis in black (with virtual addresses on the left, and the actual physical addresses on the right); the middle graph shows array accesses in dark gray (again with virtual on left and physical on right); finally, the topmost graph shows page table memory accesses in light gray (just physical, as the page table in this example resides in physical memory). The x-axis, for the entire trace, shows memory accesses across the first five iterations of the loop (there are 10 memory accesses per loop, which includes four instruction fetches, one explicit update of memory, and five page table accesses to translate those four fetches and one explicit update).

See if you can make sense of the patterns that show up in this visualization. In particular, what will change as the loop continues to run beyond these first five iterations? Which new memory locations will be accessed? Can you figure it out?

This has just been the simplest of examples (only a few lines of C code), and yet you might already be able to sense the complexity of understanding the actual memory behavior of real applications. Don’t worry: it definitely gets worse, because the mechanisms we are about to introduce only complicate this already complex machinery. Sorry!
18.5 Summary

We have introduced the concept of paging as a solution to our challenge of virtualizing memory. Paging has many advantages over previous approaches (such as segmentation). First, it does not lead to external fragmentation, as paging (by design) divides memory into fixed-sized units. Second, it is quite flexible, enabling the sparse use of virtual address spaces.

However, implementing paging support without care will lead to a slower machine (with many extra memory accesses to access the page table) as well as memory waste (with memory filled with page tables instead of useful application data). We’ll thus have to think a little harder to come up with a paging system that not only works, but works well. The next two chapters, fortunately, will show us how to do so.
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T. Kilburn, and D.B.G. Edwards and M.J. Lanigan and F.H. Sumner
IRE Trans. EC-11, 2 (1962), pp. 223-235
The Atlas pioneered the idea of dividing memory into fixed-sized pages and in many senses was an early form of the memory-management ideas we see in modern computer systems.

[I09] “Intel 64 and IA-32 Architectures Software Developer’s Manuals”
Intel, 2009

S. H. Lavington
Communications of the ACM archive
Volume 21, Issue 1 (January 1978), pp. 4-12
Special issue on computer architecture
This paper is a great retrospective of some of the history of the development of some important computer systems. As we sometimes forget in the US, many of these new ideas came from overseas.
Homework

In this homework, you will use a simple program, which is known as paging-linear-translate.py, to see if you understand how simple virtual-to-physical address translation works with linear page tables. See the README for details.

Questions

- Before doing any translations, let’s use the simulator to study how linear page tables change size given different parameters. Compute the size of linear page tables as different parameters change. Some suggested inputs are below; by using the -v flag, you can see how many page-table entries are filled.
  
  - First, to understand how linear page table size changes as the address space grows:
    
    ```
    paging-linear-translate.py -P 1k -a 1m -p 512m -v -n 0
    paging-linear-translate.py -P 1k -a 2m -p 512m -v -n 0
    paging-linear-translate.py -P 1k -a 4m -p 512m -v -n 0
    ```
  
  - Then, to understand how linear page table size changes as page size grows:
    
    ```
    paging-linear-translate.py -P 1k -a 1m -p 512m -v -n 0
    paging-linear-translate.py -P 2k -a 1m -p 512m -v -n 0
    paging-linear-translate.py -P 4k -a 1m -p 512m -v -n 0
    ```

  Before running any of these, try to think about the expected trends. How should page-table size change as the address space grows? As the page size grows? Why shouldn’t we just use really big pages in general?

- Now let’s do some translations. Start with some small examples, and change the number of pages that are allocated to the address space with the -u flag. For example:

  ```
  paging-linear-translate.py -P 1k -a 16k -p 32k -v -u 0
  paging-linear-translate.py -P 1k -a 16k -p 32k -v -u 25
  paging-linear-translate.py -P 1k -a 16k -p 32k -v -u 50
  paging-linear-translate.py -P 1k -a 16k -p 32k -v -u 75
  paging-linear-translate.py -P 1k -a 16k -p 32k -v -u 100
  ```

  What happens as you increase the percentage of pages that are allocated in each address space?

- Now let’s try some different random seeds, and some different (and sometimes quite crazy) address-space parameters, for variety:

  ```
  paging-linear-translate.py -P 8 -a 32 -p 1024 -v -s 1
  paging-linear-translate.py -P 8k -a 32k -p 1m -v -s 2
  paging-linear-translate.py -P 1m -a 256m -p 512m -v -s 3
  ```

  Which of these parameter combinations are unrealistic? Why?
• Use the program to try out some other problems. Can you find the limits of where the program doesn’t work anymore? For example, what happens if the address-space size is bigger than physical memory?
Paging: Faster Translations (TLBs)

Using paging as the core mechanism to support virtual memory can lead to high performance overheads. By chopping the address space into small, fixed-sized units (i.e., pages), paging requires a large amount of mapping information. Because that mapping information is generally stored in physical memory, paging logically requires an extra memory lookup for each virtual address generated by the program. Going to memory for translation information before every instruction fetch or explicit load or store is prohibitively slow. And thus our problem:

**THE CRUX:**
**How To Speed Up Address Translation**

How can we speed up address translation, and generally avoid the extra memory reference that paging seems to require? What hardware support is required? What OS involvement is needed?

When we want to make things fast, the OS usually needs some help. And help often comes from the OS’s old friend: the hardware. To speed address translation, we are going to add what is called (for historical reasons [CP78]) a **translation-lookaside buffer**, or TLB [C68, C95]. A TLB is part of the chip’s **memory-management unit** (MMU), and is simply a hardware **cache** of popular virtual-to-physical address translations; thus, a better name would be an **address-translation cache**. Upon each virtual memory reference, the hardware first checks the TLB to see if the desired translation is held therein; if so, the translation is performed (quickly) **without** having to consult the page table (which has all translations). Because of their tremendous performance impact, TLBs in a real sense make virtual memory possible [C95].

19.1 TLB Basic Algorithm

Figure 19.1 shows a rough sketch of how hardware might handle a virtual address translation, assuming a simple **linear page table** (i.e., the page table is an array) and a **hardware-managed TLB** (i.e., the hardware handles much of the responsibility of page table accesses; we’ll explain more about this below).
The algorithm the hardware follows works like this: first, extract the virtual page number (VPN) from the virtual address (Line 1 in Figure 19.1), and check if the TLB holds the translation for this VPN (Line 2). If it does, we have a TLB hit, which means the TLB holds the translation. Success! We can now extract the page frame number (PFN) from the relevant TLB entry, concatenate that onto the offset from the original virtual address, and form the desired physical address (PA), and access memory (Lines 5–7), assuming protection checks do not fail (Line 4).

If the CPU does not find the translation in the TLB (a TLB miss), we have some more work to do. In this example, the hardware accesses the page table to find the translation (Lines 11–12), and, assuming that the virtual memory reference generated by the process is valid and accessible (Lines 13, 15), updates the TLB with the translation (Line 18). These set of actions are costly, primarily because of the extra memory reference needed to access the page table (Line 12). Finally, once the TLB is updated, the hardware retries the instruction; this time, the translation is found in the TLB, and the memory reference is processed quickly.

The TLB, like all caches, is built on the premise that in the common case, translations are found in the cache (i.e., are hits). If so, little overhead is added, as the TLB is found near the processing core and is designed to be quite fast. When a miss occurs, the high cost of paging is incurred; the page table must be accessed to find the translation, and an extra memory reference (or more, with more complex page tables) results. If this happens often, the program will likely run noticeably more slowly; memory accesses, relative to most CPU instructions, are quite costly, and TLB misses lead to more memory accesses. Thus, it is our hope to avoid TLB misses as much as we can.

```cpp
VPN = (VirtualAddress & VPN_MASK) >> SHIFT
(Success, TlbEntry) = TLB_Lookup(VPN)
if (Success == True) // TLB Hit
    if (CanAccess(TlbEntry.ProtectBits) == True)
        Offset = VirtualAddress & OFFSET_MASK
        PhysAddr = (TlbEntry.PFN << SHIFT) | Offset
        AccessMemory(PhysAddr)
    else
        RaiseException(PROTECTION_FAULT)
else // TLB Miss
    PTEAddr = PTBR + (VPN * sizeof(PTE))
    PTE = AccessMemory(PTEAddr)
    if (PTE.Valid == False)
        RaiseException(SEGMENTATION_FAULT)
    else if (CanAccess(PTE.ProtectBits) == False)
        RaiseException(PROTECTION_FAULT)
    else
        TLB_Insert(VPN, PTE.PFN, PTE.ProtectBits)
    RetryInstruction()
```

Figure 19.1: TLB Control Flow Algorithm
19.2 Example: Accessing An Array

To make clear the operation of a TLB, let’s examine a simple virtual address trace and see how a TLB can improve its performance. In this example, let’s assume we have an array of 10 4-byte integers in memory, starting at virtual address 100. Assume further that we have a small 8-bit virtual address space, with 16-byte pages; thus, a virtual address breaks down into a 4-bit VPN (there are 16 virtual pages) and a 4-bit offset (there are 16 bytes on each of those pages).

Figure 19.2 shows the array laid out on the 16 16-byte pages of the system. As you can see, the array’s first entry (a[0]) begins on (VPN=06, offset=04); only three 4-byte integers fit onto that page. The array continues onto the next page (VPN=07), where the next four entries (a[3] ... a[6]) are found. Finally, the last three entries of the 10-entry array (a[7] ... a[9]) are located on the next page of the address space (VPN=08).

Now let’s consider a simple loop that accesses each array element, something that would look like this in C:

```c
int sum = 0;
for (i = 0; i < 10; i++) {
    sum += a[i];
}
```
For the sake of simplicity, we will pretend that the only memory accesses the loop generates are to the array (ignoring the variables \( i \) and \( \text{sum} \), as well as the instructions themselves). When the first array element (\( a[0] \)) is accessed, the CPU will see a load to virtual address 100. The hardware extracts the VPN from this (VPN=06), and uses that to check the TLB for a valid translation. Assuming this is the first time the program accesses the array, the result will be a TLB miss.

The next access is to \( a[1] \), and there is some good news here: a TLB hit! Because the second element of the array is packed next to the first, it lives on the same page; because we’ve already accessed this page when accessing the first element of the array, the translation is already loaded into the TLB. And hence the reason for our success. Access to \( a[2] \) encounters similar success (another hit), because it too lives on the same page as \( a[0] \) and \( a[1] \).

Unfortunately, when the program accesses \( a[3] \), we encounter another TLB miss. However, once again, the next entries (\( a[4] \) ... \( a[6] \)) will hit in the TLB, as they all reside on the same page in memory.

Finally, access to \( a[7] \) causes one last TLB miss. The hardware once again consults the page table to figure out the location of this virtual page in physical memory, and updates the TLB accordingly. The final two accesses (\( a[8] \) and \( a[9] \)) receive the benefits of this TLB update; when the hardware looks in the TLB for their translations, two more hits result.

Let us summarize TLB activity during our ten accesses to the array: miss, hit, hit, miss, hit, hit, miss, hit, hit. Thus, our TLB hit rate, which is the number of hits divided by the total number of accesses, is 70%. Although this is not too high (indeed, we desire hit rates that approach 100%), it is non-zero, which may be a surprise. Even though this is the first time the program accesses the array, TLB performance gains benefit from spatial locality. The elements of the array are packed tightly into pages (i.e., they are close to one another in space), and thus only the first access to an element on a page yields a TLB miss.

Also note the role that page size plays in this example. If the page size had simply been twice as big (32 bytes, not 16), the array access would suffer even fewer misses. As typical page sizes are more like 4KB, these types of dense, array-based accesses achieve excellent TLB performance, encountering only a single miss per page of accesses.

One last point about TLB performance: if the program, soon after this loop completes, accesses the array again, we’d likely see an even better result, assuming that we have a big enough TLB to cache the needed translations: hit, hit, hit, hit, hit, hit, hit, hit, hit. In this case, the TLB hit rate would be high because of temporal locality, i.e., the quick re-referencing of memory items in time. Like any cache, TLBs rely upon both spatial and temporal locality for success, which are program properties. If the program of interest exhibits such locality (and many programs do), the TLB hit rate will likely be high.
Caching is one of the most fundamental performance techniques in computer systems, one that is used again and again to make the “common-case fast” [HP06]. The idea behind hardware caches is to take advantage of locality in instruction and data references. There are usually two types of locality: temporal locality and spatial locality. With temporal locality, the idea is that an instruction or data item that has been recently accessed will likely be re-accessed soon in the future. Think of loop variables or instructions in a loop; they are accessed repeatedly over time. With spatial locality, the idea is that if a program accesses memory at address $x$, it will likely soon access memory near $x$. Imagine here streaming through an array of some kind, accessing one element and then the next. Of course, these properties depend on the exact nature of the program, and thus are not hard-and-fast laws but more like rules of thumb.

Hardware caches, whether for instructions, data, or address translations (as in our TLB) take advantage of locality by keeping copies of memory in small, fast on-chip memory. Instead of having to go to a (slow) memory to satisfy a request, the processor can first check if a nearby copy exists in a cache; if it does, the processor can access it quickly (i.e., in a few cycles) and avoid spending the costly time it takes to access memory (many nanoseconds).

You might be wondering: if caches (like the TLB) are so great, why don’t we just make bigger caches and keep all of our data in them? Unfortunately, this is where we run into more fundamental laws like those of physics. If you want a fast cache, it has to be small, as issues like the speed-of-light and other physical constraints become relevant. Any large cache by definition is slow, and thus defeats the purpose. Thus, we are stuck with small, fast caches; the question that remains is how to best use them to improve performance.

### 19.3 Who Handles The TLB Miss?

One question that we must answer: who handles a TLB miss? Two answers are possible: the hardware, or the software (OS). In the olden days, the hardware had complex instruction sets (sometimes called CISC, for complex-instruction set computers) and the people who built the hardware didn’t much trust those sneaky OS people. Thus, the hardware would handle the TLB miss entirely. To do this, the hardware has to know exactly where the page tables are located in memory (via a page-table base register, used in Line 11 in Figure 19.1), as well as their exact format; on a miss, the hardware would “walk” the page table, find the correct page-table entry and extract the desired translation, update the TLB with the translation, and retry the instruction. An example of an “older” architecture that has **hardware-managed TLBs** is the Intel x86 architecture, which uses a fixed **multi-level page table** (see the next chapter for details); the current page table is pointed to by the CR3 register [I09].
VPN = (VirtualAddress & VPN_MASK) >> SHIFT
(Success, TlbEntry) = TLB_Lookup(VPN)
if (Success == True) // TLB Hit
  if (CanAccess(TlbEntry.ProtectBits) == True)
    Offset = VirtualAddress & OFFSET_MASK
    PhysAddr = (TlbEntry.PFN << SHIFT) | Offset
    Register = AccessMemory(PhysAddr)
  else
    RaiseException(PROTECTION_FAULT)
else // TLB Miss
  RaiseException(TLB_MISS)

Figure 19.3: TLB Control Flow Algorithm (OS Handled)

More modern architectures (e.g., MIPS R10k [H93] or Sun’s SPARC v9 [WG00], both RISC or reduced-instruction set computers) have what is known as a software-managed TLB. On a TLB miss, the hardware simply raises an exception (line 11 in Figure 19.3), which pauses the current instruction stream, raises the privilege level to kernel mode, and jumps to a trap handler. As you might guess, this trap handler is code within the OS that is written with the express purpose of handling TLB misses. When run, the code will lookup the translation in the page table, use special “privileged” instructions to update the TLB, and return from the trap; at this point, the hardware retries the instruction (resulting in a TLB hit).

Let’s discuss a couple of important details. First, the return-from-trap instruction needs to be a little different than the return-from-trap we saw before when servicing a system call. In the latter case, the return-from-trap should resume execution at the instruction after the trap into the OS, just as a return from a procedure call returns to the instruction immediately following the call into the procedure. In the former case, when returning from a TLB miss-handling trap, the hardware must resume execution at the instruction that caused the trap; this retry thus lets the instruction run again, this time resulting in a TLB hit. Thus, depending on how a trap or exception was caused, the hardware must save a different PC when trapping into the OS, in order to resume properly when the time to do so arrives.

Second, when running the TLB miss-handling code, the OS needs to be extra careful not to cause an infinite chain of TLB misses to occur. Many solutions exist; for example, you could keep TLB miss handlers in physical memory (where they are unmapped and not subject to address translation), or reserve some entries in the TLB for permanently-valid translations and use some of those permanent translation slots for the handler code itself; these wired translations always hit in the TLB.

The primary advantage of the software-managed approach is flexibility: the OS can use any data structure it wants to implement the page table, without necessitating hardware change. Another advantage is simplicity; as you can see in the TLB control flow (line 11 in Figure 19.3, in contrast to lines 11–19 in Figure 19.1), the hardware doesn’t have to do much on a miss; it raises an exception, and the OS TLB miss handler does the rest.
Aside: RISC vs. CISC

In the 1980’s, a great battle took place in the computer architecture community. On one side was the CISC camp, which stood for Complex Instruction Set Computing; on the other side was RISC, for Reduced Instruction Set Computing [PS81]. The RISC side was spear-headed by David Patterson at Berkeley and John Hennessy at Stanford (who are also co-authors of some famous books [HP06]), although later John Cocke was recognized with a Turing award for his earliest work on RISC [CM00].

CISC instruction sets tend to have a lot of instructions in them, and each instruction is relatively powerful. For example, you might see a string copy, which takes two pointers and a length and copies bytes from source to destination. The idea behind CISC was that instructions should be high-level primitives, to make the assembly language itself easier to use, and to make code more compact.

RISC instruction sets are exactly the opposite. A key observation behind RISC is that instruction sets are really compiler targets, and all compilers really want are a few simple primitives that they can use to generate high-performance code. Thus, RISC proponents argued, let’s rip out as much from the hardware as possible (especially the microcode), and make what’s left simple, uniform, and fast.

In the early days, RISC chips made a huge impact, as they were noticeably faster [BC91]; many papers were written; a few companies were formed (e.g., MIPS and Sun). However, as time progressed, CISC manufacturers such as Intel incorporated many RISC techniques into the core of their processors, for example by adding early pipeline stages that transformed complex instructions into micro-instructions which could then be processed in a RISC-like manner. These innovations, plus a growing number of transistors on each chip, allowed CISC to remain competitive. The end result is that the debate died down, and today both types of processors can be made to run fast.

19.4 TLB Contents: What’s In There?

Let’s look at the contents of the hardware TLB in more detail. A typical TLB might have 32, 64, or 128 entries and be what is called fully associative. Basically, this just means that any given translation can be anywhere in the TLB, and that the hardware will search the entire TLB in parallel to find the desired translation. A typical TLB entry might look like this:

```plaintext
VPN | PFN | other bits
```

Note that both the VPN and PFN are present in each entry, as a translation could end up in any of these locations (in hardware terms, the TLB is known as a fully-associative cache). The hardware searches the entries in parallel to see if there is a match.
A common mistake is to confuse the valid bits found in a TLB with those found in a page table. In a page table, when a page-table entry (PTE) is marked invalid, it means that the page has not been allocated by the process, and should not be accessed by a correctly-working program. The usual response when an invalid page is accessed is to trap to the OS, which will respond by killing the process.

A TLB valid bit, in contrast, simply refers to whether a TLB entry has a valid translation within it. When a system boots, for example, a common initial state for each TLB entry is to be set to invalid, because no address translations are yet cached there. Once virtual memory is enabled, and once programs start running and accessing their virtual address spaces, the TLB is slowly populated, and thus valid entries soon fill the TLB.

The TLB valid bit is quite useful when performing a context switch too, as we’ll discuss further below. By setting all TLB entries to invalid, the system can ensure that the about-to-be-run process does not accidentally use a virtual-to-physical translation from a previous process.

More interesting are the “other bits”. For example, the TLB commonly has a valid bit, which says whether the entry has a valid translation or not. Also common are protection bits, which determine how a page can be accessed (as in the page table). For example, code pages might be marked read and execute, whereas heap pages might be marked read and write. There may also be a few other fields, including an address-space identifier, a dirty bit, and so forth; see below for more information.

### 19.5 TLB Issue: Context Switches

With TLBs, some new issues arise when switching between processes (and hence address spaces). Specifically, the TLB contains virtual-to-physical translations that are only valid for the currently running process; these translations are not meaningful for other processes. As a result, when switching from one process to another, the hardware or OS (or both) must be careful to ensure that the about-to-be-run process does not accidentally use translations from some previously run process.

To understand this situation better, let’s look at an example. When one process (P1) is running, it assumes the TLB might be caching translations that are valid for it, i.e., that come from P1’s page table. Assume, for this example, that the 10th virtual page of P1 is mapped to physical frame 100.

In this example, assume another process (P2) exists, and the OS soon might decide to perform a context switch and run it. Assume here that the 10th virtual page of P2 is mapped to physical frame 170. If entries for both processes were in the TLB, the contents of the TLB would be:
In the TLB above, we clearly have a problem: VPN 10 translates to either PFN 100 (P1) or PFN 170 (P2), but the hardware can’t distinguish which entry is meant for which process. Thus, we need to do some more work in order for the TLB to correctly and efficiently support virtualization across multiple processes. And thus, a crux:

**The Crux:**

**How To Manage TLB Contents On A Context Switch**

When context-switching between processes, the translations in the TLB for the last process are not meaningful to the about-to-be-run process. What should the hardware or OS do in order to solve this problem?

There are a number of possible solutions to this problem. One approach is to simply flush the TLB on context switches, thus emptying it before running the next process. On a software-based system, this can be accomplished with an explicit (and privileged) hardware instruction; with a hardware-managed TLB, the flush could be enacted when the page-table base register is changed (note the OS must change the PTBR on a context switch anyhow). In either case, the flush operation simply sets all valid bits to 0, essentially clearing the contents of the TLB.

By flushing the TLB on each context switch, we now have a working solution, as a process will never accidentally encounter the wrong translations in the TLB. However, there is a cost: each time a process runs, it must incur TLB misses as it touches its data and code pages. If the OS switches between processes frequently, this cost may be high.

To reduce this overhead, some systems add hardware support to enable sharing of the TLB across context switches. In particular, some hardware systems provide an address space identifier (ASID) field in the TLB. You can think of the ASID as a process identifier (PID), but usually it has fewer bits (e.g., 8 bits for the ASID versus 32 bits for a PID).

If we take our example TLB from above and add ASIDs, it is clear processes can readily share the TLB: only the ASID field is needed to differentiate otherwise identical translations. Here is a depiction of a TLB with the added ASID field:

<table>
<thead>
<tr>
<th>VPN</th>
<th>PFN</th>
<th>valid</th>
<th>prot</th>
<th>ASID</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>100</td>
<td>1</td>
<td>rwx</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>170</td>
<td>1</td>
<td>rwx</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

In the TLB above, we clearly have a problem: VPN 10 translates to either PFN 100 (P1) or PFN 170 (P2), but the hardware can’t distinguish which entry is meant for which process. Thus, we need to do some more work in order for the TLB to correctly and efficiently support virtualization across multiple processes. And thus, a crux:
Thus, with address-space identifiers, the TLB can hold translations from different processes at the same time without any confusion. Of course, the hardware also needs to know which process is currently running in order to perform translations, and thus the OS must, on a context switch, set some privileged register to the ASID of the current process.

As an aside, you may also have thought of another case where two entries of the TLB are remarkably similar. In this example, there are two entries for two different processes with two different VPNs that point to the same physical page:

<table>
<thead>
<tr>
<th>VPN</th>
<th>PFN</th>
<th>valid</th>
<th>prot</th>
<th>ASID</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>101</td>
<td>1</td>
<td>r-x</td>
<td>1</td>
</tr>
<tr>
<td>—</td>
<td>—</td>
<td>0</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>50</td>
<td>101</td>
<td>1</td>
<td>r-x</td>
<td>2</td>
</tr>
<tr>
<td>—</td>
<td>—</td>
<td>0</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

This situation might arise, for example, when two processes share a page (a code page, for example). In the example above, Process 1 is sharing physical page 101 with Process 2; P1 maps this page into the 10th page of its address space, whereas P2 maps it to the 50th page of its address space. Sharing of code pages (in binaries, or shared libraries) is useful as it reduces the number of physical pages in use, thus reducing memory overheads.

19.6 Issue: Replacement Policy

As with any cache, and thus also with the TLB, one more issue that we must consider is cache replacement. Specifically, when we are installing a new entry in the TLB, we have to replace an old one, and thus the question: which one to replace?

**The Crux: How To Design TLB Replacement Policy**

Which TLB entry should be replaced when we add a new TLB entry? The goal, of course, being to minimize the miss rate (or increase hit rate) and thus improve performance.

We will study such policies in some detail when we tackle the problem of swapping pages to disk in a virtual memory system; here we’ll just highlight a few of typical policies. One common approach is to evict the least-recently-used or LRU entry. The idea here is to take advantage of locality in the memory-reference stream; thus, it is likely that an entry that has not recently been used is a good candidate for eviction as (perhaps) it won’t soon be referenced again. Another typical approach is to use a random policy. Randomness sometimes makes a bad decision but has the nice property that there are not any weird corner case behaviors that can cause pessimal behavior, e.g., think of a loop accessing \( n + 1 \) pages, a TLB of size \( n \), and an LRU replacement policy.
19.7 A Real TLB Entry

Finally, let’s briefly look at a real TLB. This example is from the MIPS R4000 [H93], a modern system that uses software-managed TLBs. All 64 bits of this TLB entry can be seen in Figure 19.4.

The MIPS R4000 supports a 32-bit address space with 4KB pages. Thus, we would expect a 20-bit VPN and 12-bit offset in our typical virtual address. However, as you can see in the TLB, there are only 19 bits for the VPN; as it turns out, user addresses will only come from half the address space (the rest reserved for the kernel) and hence only 19 bits of VPN are needed. The VPN translates to up to a 24-bit physical frame number (PFN), and hence can support systems with up to 64GB of (physical) main memory ($2^{24}$ 4KB pages).

There are a few other interesting bits in the MIPS TLB. We see a global bit (G), which is used for pages that are globally-shared among processes. Thus, if the global bit is set, the ASID is ignored. We also see the 8-bit ASID, which the OS can use to distinguish between address spaces (as described above). One question for you: what should the OS do if there are more than 256 ($2^8$) processes running at a time? Finally, we see 3 Coherence (C) bits, which determine how a page is cached by the hardware (a bit beyond the scope of these notes); a dirty bit which is marked when the page has been written to (we’ll see the use of this later); a valid bit which tells the hardware if there is a valid translation present in the entry. There is also a page mask field (not shown), which supports multiple page sizes; we’ll see later why having larger pages might be useful. Finally, some of the 64 bits are unused (shaded gray in the diagram).

MIPS TLBs usually have 32 or 64 of these entries, most of which are used by user processes as they run. However, a few are reserved for the OS. A wired register can be set by the OS to tell the hardware how many slots of the TLB to reserve for the OS; the OS uses these reserved mappings for code and data that it wants to access during critical times, where a TLB miss would be problematic (e.g., in the TLB miss handler).

Because the MIPS TLB is software managed, there needs to be instructions to update the TLB. The MIPS provides four such instructions: TLBP, which probes the TLB to see if a particular translation is in there; TLBR, which reads the contents of a TLB entry into registers; TLBWI, which replaces a specific TLB entry; and TLBWR, which replaces a random TLB entry. The OS uses these instructions to manage the TLB’s contents. It is of course critical that these instructions are privileged; imagine what a user process could do if it could modify the contents of the TLB (hint: just about anything, including take over the machine, run its own malicious “OS”, or even make the Sun disappear).
The term random-access memory, or RAM, implies that you can access any part of RAM just as quickly as another. While it is generally good to think of RAM in this way, because of hardware/OS features such as the TLB, accessing a particular page of memory may be costly, particularly if that page isn’t currently mapped by your TLB. Thus, it is always good to remember the implementation tip: RAM isn’t always RAM. Sometimes randomly accessing your address space, particular if the number of pages accessed exceeds the TLB coverage, can lead to severe performance penalties. Because one of our advisors, David Culler, used to always point to the TLB as the source of many performance problems, we name this law in his honor: Culler’s Law.

19.8 Summary

We have seen how hardware can help us make address translation faster. By providing a small, dedicated on-chip TLB as an address-translation cache, most memory references will hopefully be handled without having to access the page table in main memory. Thus, in the common case, the performance of the program will be almost as if memory isn’t being virtualized at all, an excellent achievement for an operating system, and certainly essential to the use of paging in modern systems.

However, TLBs do not make the world rosy for every program that exists. In particular, if the number of pages a program accesses in a short period of time exceeds the number of pages that fit into the TLB, the program will generate a large number of TLB misses, and thus run quite a bit more slowly. We refer to this phenomenon as exceeding the TLB coverage, and it can be quite a problem for certain programs. One solution, as we’ll discuss in the next chapter, is to include support for larger page sizes; by mapping key data structures into regions of the program’s address space that are mapped by larger pages, the effective coverage of the TLB can be increased. Support for large pages is often exploited by programs such as a database management system (a DBMS), which have certain data structures that are both large and randomly-accessed.

One other TLB issue worth mentioning: TLB access can easily become a bottleneck in the CPU pipeline, in particular with what is called a physically-indexed cache. With such a cache, address translation has to take place before the cache is accessed, which can slow things down quite a bit. Because of this potential problem, people have looked into all sorts of clever ways to access caches with virtual addresses, thus avoiding the expensive step of translation in the case of a cache hit. Such a virtually-indexed cache solves some performance problems, but introduces new issues into hardware design as well. See Wiggins’s fine survey for more details [W03].
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Rafael H. Saavedra-Barrera
EECS Department, University of California, Berkeley
A great dissertation about how to predict execution time of applications by breaking them down into constituent pieces and knowing the cost of each piece. Probably the most interesting part that comes out of this work is the tool to measure details of the cache hierarchy (described in Chapter 5). Make sure to check out the wonderful diagrams therein.

Adam Wiggins
University of New South Wales TR UNSW-CSE-TR-0321, August, 2003
An excellent survey of how TLBs interact with other parts of the CPU pipeline, namely hardware caches.

David L. Weaver and Tom Germond, September 2000
SPARC International, San Jose, California
Homework (Measurement)

In this homework, you are to measure the size and cost of accessing a TLB. The idea is based on work by Saavedra-Barrera [SB92], who developed a simple but beautiful method to measure numerous aspects of cache hierarchies, all with a very simple user-level program. Read his work for more details.

The basic idea is to access some number of pages within large data structure (e.g., an array) and to time those accesses. For example, let’s say the TLB size of a machine happens to be 4 (which would be very small, but useful for the purposes of this discussion). If you write a program that touches 4 or fewer pages, each access should be a TLB hit, and thus relatively fast. However, once you touch 5 pages or more, repeatedly in a loop, each access will suddenly jump in cost, to that of a TLB miss.

The basic code to loop through an array once should look like this:

```c
int jump = PAGESIZE / sizeof(int);
for (i = 0; i < NUMPAGES * jump; i += jump) {
    a[i] += 1;
}
```

In this loop, one integer per page of the the array `a` is updated, up to the number of pages specified by `NUMPAGES`. By timing such a loop repeatedly (say, a few hundred million times in another loop around this one, or however many loops are needed to run for a few seconds), you can time how long each access takes (on average). By looking for jumps in cost as `NUMPAGES` increases, you can roughly determine how big the first-level TLB is, determine whether a second-level TLB exists (and how big it is if it does), and in general get a good sense of how TLB hits and misses can affect performance.

Here is an example graph:

As you can see in the graph, when just a few pages are accessed (8 or fewer), the average access time is roughly 5 nanoseconds. When 16 or more pages are accessed, there is a sudden jump to about 20 nanoseconds per access. A final jump in cost occurs at around 1024 pages, at which point each access takes around 70 nanoseconds. From this data, we can conclude that there is a two-level TLB hierarchy; the first is quite small (probably holding between 8 and 16 entries); the second is larger but slower (holding roughly 512 entries). The overall difference between hits in the first-level TLB and misses is quite large, roughly a factor of fourteen. TLB performance matters!
Questions

• For timing, you’ll need to use a timer such as that made available by gettimeofday(). How precise is such a timer? How long does an operation have to take in order for you to time it precisely? (this will help determine how many times, in a loop, you’ll have to repeat a page access in order to time it successfully)

• Write the program, called tlb.c, that can roughly measure the cost of accessing each page. Inputs to the program should be: the number of pages to touch and the number of trials.

• Now write a script in your favorite scripting language (csh, python, etc.) to run this program, while varying the number of pages accessed from 1 up to a few thousand, perhaps incrementing by a factor of two per iteration. Run the script on different machines and gather some data. How many trials are needed to get reliable measurements?

• Next, graph the results, making a graph that looks similar to the one above. Use a good tool like ploticus. Visualization usually makes the data much easier to digest; why do you think that is?

• One thing to watch out for is compiler optimization. Compilers do all sorts of clever things, including removing loops which increment values that no other part of the program subsequently uses. How can you ensure the compiler does not remove the main loop above from your TLB size estimator?

• Another thing to watch out for is the fact that most systems today ship with multiple CPUs, and each CPU, of course, has its own TLB hierarchy. To really get good measurements, you have to run your
code on just one CPU, instead of letting the scheduler bounce it from one CPU to the next. How can you do that? (hint: look up “pinning a thread” on Google for some clues) What will happen if you don’t do this, and the code moves from one CPU to the other?

• Another issue that might arise relates to initialization. If you don’t initialize the array \( a \) above before accessing it, the first time you access it will be very expensive, due to initial access costs such as demand zeroing. Will this affect your code and its timing? What can you do to counterbalance these potential costs?
Paging: Smaller Tables

We now tackle the second problem that paging introduces: page tables are too big and thus consume too much memory. Let’s start out with a linear page table. As you might recall\(^1\), linear page tables get pretty big. Assume again a 32-bit address space \((2^{32}\) bytes\)), with 4KB \((2^{12}\) byte\)) pages and a 4-byte page-table entry. An address space thus has roughly one million virtual pages in it \((2^{32})\); multiply by the page-table size and you see that our page table is 4MB in size. Recall also: we usually have one page table for every process in the system! With a hundred active processes (not uncommon on a modern system), we will be allocating hundreds of megabytes of memory just for page tables! As a result, we are in search of some techniques to reduce this heavy burden. There are a lot of them, so let’s get going. But not before our crux:

**Crux: How to make page tables smaller?**

Simple array-based page tables (usually called linear page tables) are too big, taking up far too much memory on typical systems. How can we make page tables smaller? What are the key ideas? What inefficiencies arise as a result of these new data structures?

20.1 Simple Solution: Bigger Pages

We could reduce the size of the page table in one simple way: use bigger pages. Take our 32-bit address space again, but this time assume 16KB pages. We would thus have an 18-bit VPN plus a 14-bit offset. Assuming the same size for each PTE (4 bytes), we now have \(2^{18}\) entries in our linear page table and thus a total size of 1MB per page table, a factor

\(^1\) Or indeed, you might not; this paging thing is getting out of control, no? That said, always make sure you understand the problem you are solving before moving onto the solution; indeed, if you understand the problem, you can often derive the solution yourself. Here, the problem should be clear: simple linear (array-based) page tables are too big.
As an aside, do note that many architectures (e.g., MIPS, SPARC, x86-64) now support multiple page sizes. Usually, a small (4KB or 8KB) page size is used. However, if a “smart” application requests it, a single large page (e.g., of size 4MB) can be used for a specific portion of the address space, enabling such applications to place a frequently-used (and large) data structure in such a space while consuming only a single TLB entry. This type of large page usage is common in database management systems and other high-end commercial applications. The main reason for multiple page sizes is not to save page table space, however; it is to reduce pressure on the TLB, enabling a program to access more of its address space without suffering from too many TLB misses. However, as researchers have shown \[N+02\], using multiple page sizes makes the OS virtual memory manager notably more complex, and thus large pages are sometimes most easily used simply by exporting a new interface to applications to request large pages directly.

The major problem with this approach, however, is that big pages lead to waste within each page, a problem known as internal fragmentation (as the waste is internal to the unit of allocation). Applications thus end up allocating pages but only using little bits and pieces of each, and memory quickly fills up with these overly-large pages. Thus, most systems use relatively small page sizes in the common case: 4KB (as in x86) or 8KB (as in SPARCv9). Our problem will not be solved so simply, alas.

### 20.2 Hybrid Approach: Paging and Segments

Whenever you have two reasonable but different approaches to something in life, you should always examine the combination of the two to see if you can obtain the best of both worlds. We call such a combination a **hybrid**. For example, why eat just chocolate or plain peanut butter when you can instead combine the two in a lovely hybrid known as the Reese’s Peanut Butter Cup [M28]?

Years ago, the creators of Multics (in particular Jack Dennis) chanced upon such an idea in the construction of the Multics virtual memory system [M07]. Specifically, Dennis had the idea of combining paging and segmentation in order to reduce the memory overhead of page tables. We can see why this might work by examining a typical linear page table in more detail. Assume we have an address space in which the used portions of the heap and stack are small. For the example, we use a tiny 16KB address space with 1KB pages (Figure 20.1); the page table for this address space is in Table 20.1.
This example assumes the single code page (VPN 0) is mapped to physical page 10, the single heap page (VPN 4) to physical page 23, and the two stack pages at the other end of the address space (VPNs 14 and 15) are mapped to physical pages 28 and 4, respectively. As you can see from the picture, most of the page table is unused, full of invalid entries. What a waste! And this is for a tiny 16KB address space. Imagine the page table of a 32-bit address space and all the potential wasted space in there! Actually, don’t imagine such a thing; it’s far too gruesome.

<table>
<thead>
<tr>
<th>PFN</th>
<th>valid</th>
<th>prot</th>
<th>present</th>
<th>dirty</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>1</td>
<td>r-x</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>23</td>
<td>1</td>
<td>rw-</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>28</td>
<td>1</td>
<td>rw-</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>4</td>
<td>1</td>
<td>rw-</td>
<td>1</td>
<td>1</td>
</tr>
</tbody>
</table>

Table 20.1: A Page Table For 16-KB Address Space
Thus, our hybrid approach: instead of having a single page table for the entire address space of the process, why not have one per logical segment? In this example, we might thus have three page tables, one for the code, heap, and stack parts of the address space.

Now, remember with segmentation, we had a base register that told us where each segment lived in physical memory, and a bound or limit register that told us the size of said segment. In our hybrid, we still have those structures in the MMU; here, we use the base not to point to the segment itself but rather to hold the physical address of the page table of that segment. The bounds register is used to indicate the end of the page table (i.e., how many valid pages it has).

Let’s do a simple example to clarify. Assume a 32-bit virtual address space with 4KB pages, and an address space split into four segments. We’ll only use three segments for this example: one for code, one for heap, and one for stack.

To determine which segment an address refers to, we’ll use the top two bits of the address space. Let’s assume 00 is the unused segment, with 01 for code, 10 for the heap, and 11 for the stack. Thus, a virtual address looks like this:

```
31 30 29 28 27 26 25 24 23 22 21 20 19 18 17 16 15 14 13 12 11 10 9 8 7 6 5 4 3 2 1 0
Seg VPN Offset
```

In the hardware, assume that there are thus three base/bounds pairs, one each for code, heap, and stack. When a process is running, the base register for each of these segments contains the physical address of a linear page table for that segment; thus, each process in the system now has three page tables associated with it. On a context switch, these registers must be changed to reflect the location of the page tables of the newly-running process.

On a TLB miss (assuming a hardware-managed TLB, i.e., where the hardware is responsible for handling TLB misses), the hardware uses the segment bits (SN) to determine which base and bounds pair to use. The hardware then takes the physical address therein and combines it with the VPN as follows to form the address of the page table entry (PTE):

```
SN = (VirtualAddress & SEG_MASK) >> SN_SHIFT
VPN = (VirtualAddress & VPN_MASK) >> VPN_SHIFT
AddressOfPTE = Base[SN] + (VPN * sizeof(PTE))
```

This sequence should look familiar; it is virtually identical to what we saw before with linear page tables. The only difference, of course, is the use of one of three segment base registers instead of the single page table base register.

The critical difference in our hybrid scheme is the presence of a bounds register per segment; each bounds register holds the value of the maximum valid page in the segment. For example, if the code segment is using its first three pages (0, 1, and 2), the code segment page table will only have three entries allocated to it and the bounds register will be set...
When you have two good and seemingly opposing ideas, you should always see if you can combine them into a hybrid that manages to achieve the best of both worlds. Hybrid corn species, for example, are known to be more robust than any naturally-occurring species. Of course, not all hybrids are a good idea; see the Zeedonk (or Zonkey), which is a cross of a Zebra and a Donkey. If you don’t believe such a creature exists, look it up, and prepare to be amazed.

However, as you might notice, this approach is not without problems. First, it still requires us to use segmentation; as we discussed before, segmentation is not quite as flexible as we would like, as it assumes a certain usage pattern of the address space; if we have a large but sparsely-used heap, for example, we can still end up with a lot of page table waste. Second, this hybrid causes external fragmentation to arise again. While most of memory is managed in page-sized units, page tables now can be of arbitrary size (in multiples of PTEs). Thus, finding free space for them in memory is more complicated. For these reasons, people continued to look for better approaches to implementing smaller page tables.

20.3 Multi-level Page Tables

A different approach doesn’t rely on segmentation but attacks the same problem: how to get rid of all those invalid regions in the page table instead of keeping them all in memory? We call this approach a multi-level page table, as it turns the linear page table into something like a tree. This approach is so effective that many modern systems employ it (e.g., x86 [BOH10]). We now describe this approach in detail.

The basic idea behind a multi-level page table is simple. First, chop up the page table into page-sized units; then, if an entire page of page-table entries (PTEs) is invalid, don’t allocate that page of the page table at all. To track whether a page of the page table is valid (and if valid, where it is in memory), use a new structure, called the page directory. The page directory thus either can be used to tell you where a page of the page table is, or that the entire page of the page table contains no valid pages.

Figure 20.2 shows an example. On the left of the figure is the classic linear page table; even though most of the middle regions of the address space are not valid, we still have to have page-table space allocated for those regions (i.e., the middle two pages of the page table). On the right is a multi-level page table. The page directory marks just two pages of
Figure 20.2: Linear (Left) And Multi-Level (Right) Page Tables

The page table as valid (the first and last); thus, just those two pages of the page table reside in memory. And thus you can see one way to visualize what a multi-level table is doing: it just makes parts of the linear page table disappear (freeing those frames for other uses), and tracks which pages of the page table are allocated with the page directory.

The page directory, in a simple two-level table, contains one entry per page of the page table. It consists of a number of page directory entries (PDE). A PDE (minimally) has a valid bit and a page frame number (PFN), similar to a PTE. However, as hinted at above, the meaning of this valid bit is slightly different: if the PDE entry is valid, it means that at least one of the pages of the page table that the entry points to (via the PFN) is valid, i.e., in at least one PTE on that page pointed to by this PDE, the valid bit in that PTE is set to one. If the PDE entry is not valid (i.e., equal to zero), the rest of the PDE is not defined.

Multi-level page tables have some obvious advantages over approaches we’ve seen thus far. First, and perhaps most obviously, the multi-level table only allocates page-table space in proportion to the amount of address space you are using; thus it is generally compact and supports sparse address spaces.

Second, if carefully constructed, each portion of the page table fits neatly within a page, making it easier to manage memory; the OS can simply grab the next free page when it needs to allocate or grow a page table. Contrast this to a simple (non-paged) linear page table\(^2\), which is just an array of PTEs indexed by VPN; with such a structure, the entire linear page table must reside contiguously in physical memory. For a large page table (say 4MB), finding such a large chunk of unused contiguous free physical memory can be quite a challenge. With a multi-level

\(^2\)We are making some assumptions here, i.e., that all page tables reside in their entirety in physical memory (i.e., they are not swapped to disk); we’ll soon relax this assumption.
When building a data structure, one should always consider time-space trade-offs in its construction. Usually, if you wish to make access to a particular data structure faster, you will have to pay a space-usage penalty for the structure.

structure, we add a level of indirection through use of the page directory, which points to pieces of the page table; that indirection allows us to place page-table pages wherever we would like in physical memory.

It should be noted that there is a cost to multi-level tables; on a TLB miss, two loads from memory will be required to get the right translation information from the page table (one for the page directory, and one for the PTE itself), in contrast to just one load with a linear page table. Thus, the multi-level table is a small example of a time-space trade-off. We wanted smaller tables (and got them), but not for free; although in the common case (TLB hit), performance is obviously identical, a TLB miss suffers from a higher cost with this smaller table.

Another obvious negative is complexity. Whether it is the hardware or OS handling the page-table lookup (on a TLB miss), doing so is undoubtedly more involved than a simple linear page-table lookup. Often we are willing to increase complexity in order to improve performance or reduce overheads; in the case of a multi-level table, we make page-table lookups more complicated in order to save valuable memory.

A Detailed Multi-Level Example

To understand the idea behind multi-level page tables better, let’s do an example. Imagine a small address space of size 16 KB, with 64-byte pages. Thus, we have a 14-bit virtual address space, with 8 bits for the VPN and 6 bits for the offset. A linear page table would have $2^8$ (256) entries, even if only a small portion of the address space is in use. Figure 20.3 presents one example of such an address space.

![Figure 20.3: A 16-KB Address Space With 64-byte Pages](image-url)
System designers should be wary of adding complexity into their system. What a good systems builder does is implement the least complex system that achieves the task at hand. For example, if disk space is abundant, you shouldn’t design a file system that works hard to use as few bytes as possible; similarly, if processors are fast, it is better to write a clean and understandable module within the OS than perhaps the most CPU-optimized, hand-assembled code for the task at hand. Be wary of needless complexity, in prematurely-optimized code or other forms; such approaches make systems harder to understand, maintain, and debug.

As Antoine de Saint-Exupery famously wrote: “Perfection is finally attained not when there is no longer anything to add, but when there is no longer anything to take away.” What he didn’t write: “It’s a lot easier to say something about perfection than to actually achieve it.”

In this example, virtual pages 0 and 1 are for code, virtual pages 4 and 5 for the heap, and virtual pages 254 and 255 for the stack; the rest of the pages of the address space are unused.

To build a two-level page table for this address space, we start with our full linear page table and break it up into page-sized units. Recall our full table (in this example) has 256 entries; assume each PTE is 4 bytes in size. Thus, our page table is 1KB (256 × 4 bytes) in size. Given that we have 64-byte pages, the 1-KB page table can be divided into 16 64-byte pages; each page can hold 16 PTEs.

What we need to understand now is how to take a VPN and use it to index first into the page directory and then into the page of the page table. Remember that each is an array of entries; thus, all we need to figure out is how to construct the index for each from pieces of the VPN.

Let’s first index into the page directory. Our page table in this example is small: 256 entries, spread across 16 pages. The page directory needs one entry per page of the page table; thus, it has 16 entries. As a result, we need four bits of the VPN to index into the directory; we use the top four bits of the VPN, as follows:

<table>
<thead>
<tr>
<th>VPN Offset</th>
</tr>
</thead>
<tbody>
<tr>
<td>13 12 11 10</td>
</tr>
<tr>
<td>9 8 7 6 5 4 3 2 1 0</td>
</tr>
</tbody>
</table>

Once we extract the page-directory index (PDIIndex for short) from the VPN, we can use it to find the address of the page-directory entry (PDE) with a simple calculation: PDEAddr = PageDirBase + (PDIIndex * sizeof(PDE)). This results in our page directory, which we now examine to make further progress in our translation.

If the page-directory entry is marked invalid, we know that the access is invalid, and thus raise an exception. If, however, the PDE is valid,
we have more work to do. Specifically, we now have to fetch the page-
table entry (PTE) from the page of the page table pointed to by this page-
directory entry. To find this PTE, we have to index into the portion of the
page table using the remaining bits of the VPN:

\[
\begin{array}{c|c|c}
\text{VPN} & \text{Page Directory Index} & \text{Page Table Index} \\
\hline
13 & 12 & 11 & 10 & 9 & 8 & 7 & 6 & 5 & 4 & 3 & 2 & 1 & 0 \\
\end{array}
\]

This page-table index (PTIndex for short) can then be used to index
into the page table itself, giving us the address of our PTE:

\[
PTE\text{Addr} = (\text{PDE.PFN } \ll \text{SHIFT}) + (\text{PTIndex} \times \text{sizeof(PTE)})
\]

Note that the page-frame number (PFN) obtained from the page-directory
entry must be left-shifted into place before combining it with the page-
table index to form the address of the PTE.

To see if this all makes sense, we’ll now fill in a multi-level page ta-
ble with some actual values, and translate a single virtual address. Let’s
begin with the page directory for this example (left side of Table 20.2).

In the figure, you can see that each page directory entry (PDE) de-
scribes something about a page of the page table for the address space.
In this example, we have two valid regions in the address space (at the
beginning and end), and a number of invalid mappings in-between.

In physical page 100 (the physical frame number of the 0th page of the
page table), we have the first page of 16 page table entries for the first 16
VPNs in the address space. See Table 20.2 (middle part) for the contents
of this portion of the page table.

<table>
<thead>
<tr>
<th>Page Directory</th>
<th>Page of PT (@PFN:100)</th>
<th>Page of PT (@PFN:101)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PFN</td>
<td>valid?</td>
<td>PFN</td>
</tr>
<tr>
<td>100</td>
<td>1</td>
<td>10</td>
</tr>
<tr>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>---</td>
<td>0</td>
<td>23</td>
</tr>
<tr>
<td>---</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>---</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>---</td>
<td>0</td>
<td>80</td>
</tr>
<tr>
<td>---</td>
<td>0</td>
<td>59</td>
</tr>
<tr>
<td>---</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>---</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>---</td>
<td>0</td>
<td>0</td>
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<td>0</td>
<td>0</td>
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<tr>
<td>---</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>---</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>---</td>
<td>0</td>
<td>55</td>
</tr>
<tr>
<td>101</td>
<td>1</td>
<td>0</td>
</tr>
</tbody>
</table>

Table 20.2: A Page Directory, And Pieces Of Page Table
This page of the page table contains the mappings for the first 16 VPNs; in our example, VPNs 0 and 1 are valid (the code segment), as are 4 and 5 (the heap). Thus, the table has mapping information for each of those pages. The rest of the entries are marked invalid.

The other valid page of page table is found inside PFN 101. This page contains mappings for the last 16 VPNs of the address space; see Table 20.2 (right) for details.

In the example, VPNs 254 and 255 (the stack) have valid mappings. Hopefully, what we can see from this example is how much space savings are possible with a multi-level indexed structure. In this example, instead of allocating the full sixteen pages for a linear page table, we allocate only three: one for the page directory, and two for the chunks of the page table that have valid mappings. The savings for large (32-bit or 64-bit) address spaces could obviously be much greater.

Finally, let’s use this information in order to perform a translation. Here is an address that refers to the 0th byte of VPN 254: 0x3F80, or 11 1111 1000 0000 in binary.

Recall that we will use the top 4 bits of the VPN to index into the page directory. Thus, 1111 will choose the last (15th, if you start at the 0th) entry of the page directory above. This points us to a valid page of the page table located at address 101. We then use the next 4 bits of the VPN (1110) to index into that page of the page table and find the desired PTE. 1110 is the next-to-last (14th) entry on the page, and tells us that page 254 of our virtual address space is mapped at physical page 55. By concatenating PFN=55 (or hex 0x37) with offset=000000, we can thus form our desired physical address and issue the request to the memory system: PhysAddr = (PTE.PFN << SHIFT) + offset = 00 1101 1100 0000 = 0x0DC0.

You should now have some idea of how to construct a two-level page table, using a page directory which points to pages of the page table. Unfortunately, however, our work is not done. As we’ll now discuss, sometimes two levels of page table is not enough!

More Than Two Levels

In our example thus far, we’ve assumed that multi-level page tables only have two levels: a page directory and then pieces of the page table. In some cases, a deeper tree is possible (and indeed, needed).

Let’s take a simple example and use it to show why a deeper multi-level table can be useful. In this example, assume we have a 30-bit virtual address space, and a small (512 byte) page. Thus our virtual address has a 21-bit virtual page number component and a 9-bit offset.

Remember our goal in constructing a multi-level page table: to make each piece of the page table fit within a single page. Thus far, we’ve only considered the page table itself; however, what if the page directory gets too big?
To determine how many levels are needed in a multi-level table to make all pieces of the page table fit within a page, we start by determining how many page-table entries fit within a page. Given our page size of 512 bytes, and assuming a PTE size of 4 bytes, you should see that you can fit 128 PTEs on a single page. When we index into a page of the page table, we can thus conclude we’ll need the least significant 7 bits ($\log_2 128$) of the VPN as an index:

![Virtual Address Diagram]

What you also might notice from the diagram above is how many bits are left into the (large) page directory: 14. If our page directory has $2^{14}$ entries, it spans not one page but 128, and thus our goal of making every piece of the multi-level page table fit into a page vanishes.

To remedy this problem, we build a further level of the tree, by splitting the page directory itself into multiple pages, and then adding another page directory on top of that, to point to the pages of the page directory. We can thus split up our virtual address as follows:

![Virtual Address Diagram 2]

Now, when indexing the upper-level page directory, we use the very top bits of the virtual address (PD Index 0 in the diagram); this index can be used to fetch the page-directory entry from the top-level page directory. If valid, the second level of the page directory is consulted by combining the physical frame number from the top-level PDE and the next part of the VPN (PD Index 1). Finally, if valid, the PTE address can be formed by using the page-table index combined with the address from the second-level PDE. Whew! That’s a lot of work. And all just to look something up in a multi-level table.

**The Translation Process: Remember the TLB**

To summarize the entire process of address translation using a two-level page table, we once again present the control flow in algorithmic form (Figure 20.4). The figure shows what happens in hardware (assuming a hardware-managed TLB) upon every memory reference.

As you can see from the figure, before any of the complicated multi-level page table access occurs, the hardware first checks the TLB; upon
VPN = (VirtualAddress & VPN_MASK) >> SHIFT
(Success, TlbEntry) = TLB_Lookup(VPN)
if (Success == True) // TLB Hit
  if (CanAccess(TlbEntry.ProtectBits) == True)
    Offset = VirtualAddress & OFFSET_MASK
    PhysAddr = (TlbEntry.PFN << SHIFT) | Offset
    Register = AccessMemory(PhysAddr)
  else
    RaiseException(PROTECTION_FAULT)
else
  // first, get page directory entry
  PDIndex = (VPN & PD_MASK) >> PD_SHIFT
  PDEAddr = PDBR + (PDIndex * sizeof(PDE))
  PDE = AccessMemory(PDEAddr)
  if (PDE.Valid == False)
    RaiseException(SEGMENTATION_FAULT)
  else
    // PDE is valid: now fetch PTE from page table
    PTIndex = (VPN & PT_MASK) >> PT_SHIFT
    PTEAddr = (PDE.PFN << SHIFT) + (PTIndex * sizeof(PTE))
    PTE = AccessMemory(PTEAddr)
    if (PTE.Valid == False)
      RaiseException(SEGMENTATION_FAULT)
    else if (CanAccess(PTE.ProtectBits) == False)
      RaiseException(PROTECTION_FAULT)
    else
      TLB_Insert(VPN, PTE.PFN, PTE.ProtectBits)
      RetryInstruction()

Figure 20.4: Multi-level Page Table Control Flow

a hit, the physical address is formed directly without accessing the page
table at all, as before. Only upon a TLB miss does the hardware need to
perform the full multi-level lookup. On this path, you can see the cost of
our traditional two-level page table: two additional memory accesses to
look up a valid translation.

20.4 Inverted Page Tables

An even more extreme space savings in the world of page tables is
found with inverted page tables. Here, instead of having many page
tables (one per process of the system), we keep a single page table that
has an entry for each physical page of the system. The entry tells us which
process is using this page, and which virtual page of that process maps to
this physical page.

Finding the correct entry is now a matter of searching through this
data structure. A linear scan would be expensive, and thus a hash table is
often built over the base structure to speed lookups. The PowerPC is one
example of such an architecture [JM98].

More generally, inverted page tables illustrate what we’ve said from
the beginning: page tables are just data structures. You can do lots of
crazy things with data structures, making them smaller or bigger, making
them slower or faster. Multi-level and inverted page tables are just two
eXamples of the many things one could do.
20.5 Swapping the Page Tables to Disk

Finally, we discuss the relaxation of one final assumption. Thus far, we have assumed that page tables reside in kernel-owned physical memory. Even with our many tricks to reduce the size of page tables, it is still possible, however, that they may be too big to fit into memory all at once. Thus, some systems place such page tables in kernel virtual memory, thereby allowing the system to swap some of these page tables to disk when memory pressure gets a little tight. We’ll talk more about this in a future chapter (namely, the case study on VAX/VMS), once we understand how to move pages in and out of memory in more detail.

20.6 Summary

We have now seen how real page tables are built; not necessarily just as linear arrays but as more complex data structures. The trade-offs such tables present are in time and space – the bigger the table, the faster a TLB miss can be serviced, as well as the converse – and thus the right choice of structure depends strongly on the constraints of the given environment.

In a memory-constrained system (like many older systems), small structures make sense; in a system with a reasonable amount of memory and with workloads that actively use a large number of pages, a bigger table that speeds up TLB misses might be the right choice. With software-managed TLBs, the entire space of data structures opens up to the delight of the operating system innovator (hint: that’s you). What new structures can you come up with? What problems do they solve? Think of these questions as you fall asleep, and dream the big dreams that only operating-system developers can dream.
References

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Addison-Wesley, 2010
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Hank Levy and P. Lipman
A terrific paper about a real virtual memory manager in a classic operating system, VMS. So terrific, in fact, that we’ll use it to review everything we’ve learned about virtual memory thus far a few chapters from now.

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Mars Candy Corporation.
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[N+02] “Practical, Transparent Operating System Support for Superpages”
Juan Navarro, Sitaram Iyer, Peter Druschel, Alan Cox
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A nice paper showing all the details you have to get right to incorporate large pages, or superpages, into a modern OS. Not as easy as you might think, alas.

[M07] “Multics: History”
Available: http://www.multicians.org/history.html
This amazing web site provides a huge amount of history on the Multics system, certainly one of the most influential systems in OS history. The quote from therein: “Jack Dennis of MIT contributed influential architectural ideas to the beginning of Multics, especially the idea of combining paging and segmentation.” (from Section 1.2.1)
Homework

This fun little homework tests if you understand how a multi-level page table works. And yes, there is some debate over the use of the term “fun” in the previous sentence. The program is called, perhaps unsurprisingly: paging-multilevel-translate.py; see the README for details.

Questions

• With a linear page table, you need a single register to locate the page table, assuming that hardware does the lookup upon a TLB miss. How many registers do you need to locate a two-level page table? A three-level table?
• Use the simulator to perform translations given random seeds 0, 1, and 2, and check your answers using the -c flag. How many memory references are needed to perform each lookup?
• Given your understanding of how cache memory works, how do you think memory references to the page table will behave in the cache? Will they lead to lots of cache hits (and thus fast accesses?) Or lots of misses (and thus slow accesses)?
Thus far, we’ve assumed that an address space is unrealistically small and fits into physical memory. In fact, we’ve been assuming that every address space of every running process fits into memory. We will now relax these big assumptions, and assume that we wish to support many concurrently-running large address spaces.

To do so, we require an additional level in the memory hierarchy. Thus far, we have assumed that all pages reside in physical memory. However, to support large address spaces, the OS will need a place to stash away portions of address spaces that currently aren’t in great demand. In general, the characteristics of such a location are that it should have more capacity than memory; as a result, it is generally slower (if it were faster, we would just use it as memory, no?). In modern systems, this role is usually served by a hard disk drive. Thus, in our memory hierarchy, big and slow hard drives sit at the bottom, with memory just above. And thus we arrive at the crux of the problem:

**The Crux: How To Go Beyond Physical Memory**

How can the OS make use of a larger, slower device to transparently provide the illusion of a large virtual address space?

One question you might have: why do we want to support a single large address space for a process? Once again, the answer is convenience and ease of use. With a large address space, you don’t have to worry about if there is room enough in memory for your program’s data structures; rather, you just write the program naturally, allocating memory as needed. It is a powerful illusion that the OS provides, and makes your life vastly simpler. You’re welcome! A contrast is found in older systems that used memory overlays, which required programmers to manually move pieces of code or data in and out of memory as they were needed [D97]. Try imagining what this would be like: before calling a function or accessing some data, you need to first arrange for the code or data to be in memory; yuck!
Beyond just a single process, the addition of swap space allows the OS to support the illusion of a large virtual memory for multiple concurrently-running processes. The invention of multiprogramming (running multiple programs “at once”, to better utilize the machine) almost demanded the ability to swap out some pages, as early machines clearly could not hold all the pages needed by all processes at once. Thus, the combination of multiprogramming and ease-of-use leads us to want to support using more memory than is physically available. It is something that all modern VM systems do; it is now something we will learn more about.

21.1 Swap Space

The first thing we will need to do is to reserve some space on the disk for moving pages back and forth. In operating systems, we generally refer to such space as swap space, because we swap pages out of memory to it and swap pages into memory from it. Thus, we will simply assume that the OS can read from and write to the swap space, in page-sized units. To do so, the OS will need to remember the disk address of a given page.

The size of the swap space is important, as ultimately it determines the maximum number of memory pages that can be in use by a system at a given time. Let us assume for simplicity that it is very large for now.

In the tiny example (Figure 21.1), you can see a little example of a 4-page physical memory and an 8-page swap space. In the example, three processes (Proc 0, Proc 1, and Proc 2) are actively sharing physical memory; each of the three, however, only have some of their valid pages in memory, with the rest located in swap space on disk. A fourth process (Proc 3) has all of its pages swapped out to disk, and thus clearly isn’t currently running. One block of swap remains free. Even from this tiny example, hopefully you can see how using swap space allows the system to pretend that memory is larger than it actually is.

We should note that swap space is not the only on-disk location for swapping traffic. For example, assume you are running a program binary (e.g., ls, or your own compiled main program). The code pages from this binary are initially found on disk, and when the program runs, they are loaded into memory (either all at once when the program starts execution,
or, as in modern systems, one page at a time when needed). However, if the system needs to make room in physical memory for other needs, it can safely re-use the memory space for these code pages, knowing that it can later swap them in again from the on-disk binary in the file system.

21.2 The Present Bit

Now that we have some space on the disk, we need to add some machinery higher up in the system in order to support swapping pages to and from the disk. Let us assume, for simplicity, that we have a system with a hardware-managed TLB.

Recall first what happens on a memory reference. The running process generates virtual memory references (for instruction fetches, or data accesses), and, in this case, the hardware translates them into physical addresses before fetching the desired data from memory.

Remember that the hardware first extracts the VPN from the virtual address, checks the TLB for a match (a TLB hit), and if a hit, produces the resulting physical address and fetches it from memory. This is hopefully the common case, as it is fast (requiring no additional memory accesses).

If the VPN is not found in the TLB (i.e., a TLB miss), the hardware locates the page table in memory (using the page table base register) and looks up the page table entry (PTE) for this page using the VPN as an index. If the page is valid and present in physical memory, the hardware extracts the PFN from the PTE, installs it in the TLB, and retries the instruction, this time generating a TLB hit; so far, so good.

If we wish to allow pages to be swapped to disk, however, we must add even more machinery. Specifically, when the hardware looks in the PTE, it may find that the page is not present in physical memory. The way the hardware (or the OS, in a software-managed TLB approach) determines this is through a new piece of information in each page-table entry, known as the present bit. If the present bit is set to one, it means the page is present in physical memory and everything proceeds as above; if it is set to zero, the page is not in memory but rather on disk somewhere. The act of accessing a page that is not in physical memory is commonly referred to as a page fault.
Terminology in virtual memory systems can be a little confusing and variable across machines and operating systems. For example, a page fault more generally could refer to any reference to a page table that generates a fault of some kind: this could include the type of fault we are discussing here, i.e., a page-not-present fault, but sometimes can refer to illegal memory accesses. Indeed, it is odd that we call what is definitely a legal access (to a page mapped into the virtual address space of a process, but simply not in physical memory at the time) a “fault” at all; really, it should be called a page miss. But often, when people say a program is “page faulting”, they mean that it is accessing parts of its virtual address space that the OS has swapped out to disk.

We suspect the reason that this behavior became known as a “fault” relates to the machinery in the operating system to handle it. When something unusual happens, i.e., when something the hardware doesn’t know how to handle occurs, the hardware simply transfers control to the OS, hoping it can make things better. In this case, a page that a process wants to access is missing from memory; the hardware does the only thing it can, which is raise an exception, and the OS takes over from there. As this is identical to what happens when a process does something illegal, it is perhaps not surprising that we term the activity a “fault.”

Upon a page fault, the OS is invoked to service the page fault. A particular piece of code, known as a page-fault handler, runs, and must service the page fault, as we now describe.

21.3 The Page Fault

Recall that with TLB misses, we have two types of systems: hardware-managed TLBs (where the hardware looks in the page table to find the desired translation) and software-managed TLBs (where the OS does). In either type of system, if a page is not present, the OS is put in charge to handle the page fault. The appropriately-named OS page-fault handler runs to determine what to do. Virtually all systems handle page faults in software; even with a hardware-managed TLB, the hardware trusts the OS to manage this important duty.

If a page is not present and has been swapped to disk, the OS will need to swap the page into memory in order to service the page fault. Thus, a question arises: how will the OS know where to find the desired page? In many systems, the page table is a natural place to store such information. Thus, the OS could use the bits in the PTE normally used for data such as thePFN of the page for a disk address. When the OS receives a page fault for a page, it looks in the PTE to find the address, and issues the request to disk to fetch the page into memory.
We know from our experience with the TLB that hardware designers are loathe to trust the OS to do much of anything. So why do they trust the OS to handle a page fault? There are a few main reasons. First, page faults to disk are slow; even if the OS takes a long time to handle a fault, executing tons of instructions, the disk operation itself is traditionally so slow that the extra overheads of running software are minimal. Second, to be able to handle a page fault, the hardware would have to understand swap space, how to issue I/Os to the disk, and a lot of other details which it currently doesn’t know much about. Thus, for both reasons of performance and simplicity, the OS handles page faults, and even hardware types can be happy.

When the disk I/O completes, the OS will then update the page table to mark the page as present, update the PFN field of the page-table entry (PTE) to record the in-memory location of the newly-fetched page, and retry the instruction. This next attempt may generate a TLB miss, which would then be serviced and update the TLB with the translation (one could alternately update the TLB upon when servicing the page fault, to avoid this step). Finally, a last restart would find the translation in the TLB and thus proceed to fetch the desired data or instruction from memory at the translated physical address.

Note that while the I/O is in flight, the process will be in the blocked state. Thus, the OS will be free to run other ready processes while the page fault is being serviced. Because I/O is expensive, this overlap of the I/O (page fault) of one process and the execution of another is yet another way a multiprogrammed system can make the most effective use of its hardware.

21.4 What If Memory Is Full?

In the process described above, you may notice that we assumed there is plenty of free memory in which to page in a page from swap space. Of course, this may not be the case; memory may be full (or close to it). Thus, the OS might like to first page out one or more pages to make room for the new page(s) the OS is about to bring in. The process of picking a page to kick out, or replace is known as the page-replacement policy.

As it turns out, a lot of thought has been put into creating a good page-replacement policy, as kicking out the wrong page can exact a great cost on program performance. Making the wrong decision can cause a program to run at disk-like speeds instead of memory-like speeds; in current technology that means a program could run 10,000 or 100,000 times slower. Thus, such a policy is something we should study in some detail; indeed, that is exactly what we will do in the next chapter. For now, it is good enough to understand that such a policy exists, built on top of the mechanisms described here.
Figure 21.2: Page-Fault Control Flow Algorithm (Hardware)

21.5 Page Fault Control Flow

With all of this knowledge in place, we can now roughly sketch the complete control flow of memory access. In other words, when somebody asks you “what happens when a program fetches some data from memory?”, you should have a pretty good idea of all the different possibilities. See the control flow in Figures 21.2 and 21.3 for more details; the first figure shows what the hardware does during translation, and the second what the OS does upon a page fault.

From the hardware control flow diagram in Figure 21.2, notice that there are now three important cases to understand when a TLB miss occurs. First, that the page was both present and valid (Lines 18–21); in this case, the TLB miss handler can simply grab thePFN from the PTE, retry the instruction (this time resulting in a TLB hit), and thus continue as described (many times) before. In the second case (Lines 22–23), the page fault handler must be run; although this was a legitimate page for the process to access (it is valid, after all), it is not present in physical memory. Third (and finally), the access could be to an invalid page, due for example to a bug in the program (Lines 13–14). In this case, no other bits in the PTE really matter; the hardware traps this invalid access, and the OS trap handler runs, likely terminating the offending process.

From the software control flow in Figure 21.3, we can see what the OS roughly must do in order to service the page fault. First, the OS must find a physical frame for the soon-to-be-faulted-in page to reside within; if there is no such page, we’ll have to wait for the replacement algorithm to run and kick some pages out of memory, thus freeing them for use here.
PFN = FindFreePhysicalPage()
if (PFN == -1) // no free page found
PFN = EvictPage() // run replacement algorithm
DiskRead(PTE.DiskAddr, pfn) // sleep (waiting for I/O)
PTE.present = True // update page table with present
PTE.PFN = PFN // bit and translation (PFN)
RetryInstruction() // retry instruction

Figure 21.3: Page-Fault Control Flow Algorithm (Software)

With a physical frame in hand, the handler then issues the I/O request to read in the page from swap space. Finally, when that slow operation completes, the OS updates the page table and retries the instruction. The retry will result in a TLB miss, and then, upon another retry, a TLB hit, at which point the hardware will be able to access the desired item.

21.6 When Replacements Really Occur

Thus far, the way we’ve described how replacements occur assumes that the OS waits until memory is entirely full, and only then replaces (evicts) a page to make room for some other page. As you can imagine, this is a little bit unrealistic, and there are many reasons for the OS to keep a small portion of memory free more proactively.

To keep a small amount of memory free, most operating systems thus have some kind of high watermark (HW) and low watermark (LW) to help decide when to start evicting pages from memory. How this works is as follows: when the OS notices that there are fewer than LW pages available, a background thread that is responsible for freeing memory runs. The thread evicts pages until there are HW pages available. The background thread, sometimes called the swap daemon or page daemon¹, then goes to sleep, happy that it has freed some memory for running processes and the OS to use.

By performing a number of replacements at once, new performance optimizations become possible. For example, many systems will cluster or group a number of pages and write them out at once to the swap partition, thus increasing the efficiency of the disk [LL82]; as we will see later when we discuss disks in more detail, such clustering reduces seek and rotational overheads of a disk and thus increases performance noticeably.

To work with the background paging thread, the control flow in Figure 21.3 should be modified slightly; instead of performing a replacement directly, the algorithm would instead simply check if there are any free pages available. If not, it would signal that the background paging thread that free pages are needed; when the thread frees up some pages, it would re-awaken the original thread, which could then page in the desired page and go about its work.

¹The word “daemon”, usually pronounced “demon”, is an old term for a background thread or process that does something useful. Turns out (once again!) that the source of the term is Multics [CS94].
21.7 Summary

In this brief chapter, we have introduced the notion of accessing more memory than is physically present within a system. To do so requires more complexity in page-table structures, as a present bit (of some kind) must be included to tell us whether the page is present in memory or not. When not, the operating system page-fault handler runs to service the page fault, and thus arranges for the transfer of the desired page from disk to memory, perhaps first replacing some pages in memory to make room for those soon to be swapped in.

Recall, importantly (and amazingly!), that these actions all take place transparently to the process. As far as the process is concerned, it is just accessing its own private, contiguous virtual memory. Behind the scenes, pages are placed in arbitrary (non-contiguous) locations in physical memory, and sometimes they are not even present in memory, requiring a fetch from disk. While we hope that in the common case a memory access is fast, in some cases it will take multiple disk operations to service it; something as simple as performing a single instruction can, in the worst case, take many milliseconds to complete.
References

[CS94] “Take Our Word For It”  
F. Corbato and R. Steinberg  
Richard Steinberg writes: “Someone has asked me the origin of the word daemon as it applies to computing. Best I can tell based on my research, the word was first used by people on your team at Project MAC using the IBM 7094 in 1963.” Professor Corbato replies: “Our use of the word daemon was inspired by the Maxwell’s daemon of physics and thermodynamics (my background is in physics). Maxwell’s daemon was an imaginary agent which helped sort molecules of different speeds and worked tirelessly in the background. We fancifully began to use the word daemon to describe background processes which worked tirelessly to perform system chores.”

[D97] “Before Memory Was Virtual”  
Peter Denning  
From In the Beginning: Recollections of Software Pioneers, Wiley, November 1997  
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[G+95] “Idleness is not sloth”  
Richard Golding, Peter Bosch, Carl Staelin, Tim Sullivan, John Wilkes  
USENIX ATC ’95, New Orleans, Louisiana  
A fun and easy-to-read discussion of how idle time can be better used in systems, with lots of good examples.

[LL82] “Virtual Memory Management in the VAX/VMS Operating System”  
Hank Levy and P. Lipman  
Not the first place where such clustering was used, but a clear and simple explanation of how such a mechanism works.
Beyond Physical Memory: Policies

In a virtual memory manager, life is easy when you have a lot of free memory. A page fault occurs, you find a free page on the free-page list, and assign it to the faulting page. Hey, Operating System, congratulations! You did it again.

Unfortunately, things get a little more interesting when little memory is free. In such a case, this memory pressure forces the OS to start paging out pages to make room for actively-used pages. Deciding which page (or pages) to evict is encapsulated within the replacement policy of the OS; historically, it was one of the most important decisions the early virtual memory systems made, as older systems had little physical memory. Minimally, it is an interesting set of policies worth knowing a little more about. And thus our problem:

THE CRUX: HOW TO DECIDE WHICH PAGE TO EVICT
How can the OS decide which page (or pages) to evict from memory? This decision is made by the replacement policy of the system, which usually follows some general principles (discussed below) but also includes certain tweaks to avoid corner-case behaviors.

22.1 Cache Management

Before diving into policies, we first describe the problem we are trying to solve in more detail. Given that main memory holds some subset of all the pages in the system, it can rightly be viewed as a cache for virtual memory pages in the system. Thus, our goal in picking a replacement policy for this cache is to minimize the number of cache misses; that is, to minimize the number of times that we have to go to disk to fetch the desired page. Alternately, one can view our goal as maximizing the number of cache hits, i.e., the number of times a page that is read or written is found in memory.
Knowing the number of cache hits and misses let us calculate the average memory access time (AMAT) for a program (a metric computer architects compute for hardware caches [HP06]). Specifically, given these values, we can compute the AMAT of a program as follows:

\[
AMAT = (\text{Hit}\% \cdot T_M) + (\text{Miss}\% \cdot T_D)
\]

(22.1)

where \(T_M\) represents the cost of accessing memory, and represents \(T_D\) the cost of accessing disk.

For example, let us imagine a machine with a (tiny) address space: 4KB, with 256-byte pages. Thus, a virtual address has two components: a 4-bit VPN (the most-significant bits) and an 8-bit offset (the least-significant bits). Thus, a process in this example can access \(2^4\) or 16 total virtual pages. In this example, the process generates the following memory references (i.e., virtual addresses): 0x000, 0x100, 0x200, 0x300, 0x400, 0x500, 0x600, 0x700, 0x800, 0x900. These virtual addresses refer to the first byte of each of the first ten pages of the address space (the page number being the first hex digit of each virtual address).

Let us further assume that every page except virtual page 3 are already in memory. Thus, our sequence of memory references will encounter the following behavior: hit, hit, hit, miss, hit, hit, hit, hit, hit, hit. We can compute the hit rate (the percent of references found in memory): 90%, as 9 out of 10 references are in memory. The miss rate is obviously 10%.

To calculate AMAT, we simply need to know the cost of accessing memory and the cost of accessing disk. Assuming the cost of accessing memory \((T_M)\) is around 100 nanoseconds, and the cost of accessing disk \((T_D)\) is about 10 milliseconds, we have the following AMAT: 0.9 · 100ns + 0.1 · 10ms, which is 90ns + 1ms, or 1.00009 ms, or about 1 millisecond. If our hit rate had instead been 99.9%, the result is quite different: AMAT is 10.1 microseconds, or roughly 100 times faster. As the hit rate approaches 100%, AMAT approaches 100 nanoseconds.

Unfortunately, as you can see in this example, the cost of disk access is so high in modern systems that even a tiny miss rate will quickly dominate the overall AMAT of running programs. Clearly, we need to avoid as many misses as possible or run slowly, at the rate of the disk. One way to help with this is to carefully develop a smart policy, as we now do.

### 22.2 The Optimal Replacement Policy

To better understand how a particular replacement policy works, it would be nice to compare it to the best possible replacement policy. As it turns out, such an optimal policy was developed by Belady many years ago [B66] (he originally called it MIN). The optimal replacement policy leads to the fewest number of misses overall. Belady showed that a simple (but, unfortunately, difficult to implement!) approach that replaces the page that will be accessed furtherest in the future is the optimal policy, resulting in the fewest-possible cache misses.
Although optimal is not very practical as a real policy, it is incredibly useful as a comparison point in simulation or other studies. Saying that your fancy new algorithm has a 80% hit rate isn’t meaningful in isolation; saying that optimal achieves an 82% hit rate (and thus your new approach is quite close to optimal) makes the result more meaningful and gives it context. Thus, in any study you perform, knowing what the optimal is lets you perform a better comparison, showing how much improvement is still possible, and also when you can stop making your policy better, because it is close enough to the ideal [AD03].

Hopefully, the intuition behind the optimal policy makes sense. Think about it like this: if you have to throw out some page, why not throw out the one that is needed the furthest from now? By doing so, you are essentially saying that all the other pages in the cache are more important than the one furthest out. The reason this is true is simple: you will refer to the other pages before you refer to the one furthest out.

Let’s trace through a simple example to understand the decisions the optimal policy makes. Assume a program accesses the following stream of virtual pages: 0, 1, 2, 0, 1, 3, 0, 3, 1, 2, 1. Table 22.1 shows the behavior of optimal, assuming a cache that fits three pages.

In the table, you can see the following actions. Not surprisingly, the first three accesses are misses, as the cache begins in an empty state; such a miss is sometimes referred to as a cold-start miss (or compulsory miss). Then we refer again to pages 0 and 1, which both hit in the cache. Finally, we reach another miss (to page 3), but this time the cache is full; a replacement must take place! Which begs the question: which page should we replace? With the optimal policy, we examine the future for each page currently in the cache (0, 1, and 2), and see that 0 is accessed almost immediately, 1 is accessed a little later, and 2 is accessed furthest in the future. Thus the optimal policy has an easy choice: evict page 2, resulting in pages 0, 1, and 3 in the cache. The next three references are hits, but then

<table>
<thead>
<tr>
<th>Access</th>
<th>Hit/Miss?</th>
<th>Evict</th>
<th>Cache State</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Miss</td>
<td></td>
<td>0</td>
</tr>
<tr>
<td>1</td>
<td>Miss</td>
<td>0, 1</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Miss</td>
<td>0, 1, 2</td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>Hit</td>
<td>0, 1, 2</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>Hit</td>
<td>0, 1, 2</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Miss</td>
<td>2</td>
<td>0, 1, 3</td>
</tr>
<tr>
<td>0</td>
<td>Hit</td>
<td>0, 1, 3</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Hit</td>
<td>0, 1, 3</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>Hit</td>
<td>0, 1, 3</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Miss</td>
<td>3</td>
<td>0, 1, 2</td>
</tr>
<tr>
<td>1</td>
<td>Hit</td>
<td>0, 1, 2</td>
<td></td>
</tr>
</tbody>
</table>

Table 22.1: Tracing the Optimal Policy
In the computer architecture world, architects sometimes find it useful to characterize misses by type, into one of three categories: compulsory, capacity, and conflict misses, sometimes called the Three C’s [H87]. A **compulsory miss** (or **cold-start miss** [EF78]) occurs because the cache is empty to begin with and this is the first reference to the item; in contrast, a **capacity miss** occurs because the cache ran out of space and had to evict an item to bring a new item into the cache. The third type of miss (a **conflict miss**) arises in hardware because of limits on where an item can be placed in a hardware cache, due to something known as **set-associativity**; it does not arise in the OS page cache because such caches are always **fully-associative**, i.e., there are no restrictions on where in memory a page can be placed. See H&P for details [HP06].

we get to page 2, which we evicted long ago, and suffer another miss. Here the optimal policy again examines the future for each page in the cache (0, 1, and 3), and sees that as long as it doesn’t evict page 1 (which is about to be accessed), we’ll be OK. The example shows page 3 getting evicted, although 0 would have been a fine choice too. Finally, we hit on page 1 and the trace completes.

We can also calculate the hit rate for the cache: with 6 hits and 5 misses, the hit rate is 
\[ \frac{\text{Hits}}{\text{Hits} + \text{Misses}} \] which is \( \frac{6}{6+5} \) or 54.6%. You can also compute the hit rate modulo compulsory misses (i.e., ignore the first miss to a given page), resulting in a 85.7% hit rate.

Unfortunately, as we saw before in the development of scheduling policies, the future is not generally known; you can’t build the optimal policy for a general-purpose operating system\(^1\). Thus, in developing a real, deployable policy, we will focus on approaches that find some other way to decide which page to evict. The optimal policy will thus serve only as a comparison point, to know how close we are to “perfect”.

### 22.3 A Simple Policy: FIFO

Many early systems avoided the complexity of trying to approach optimal and employed very simple replacement policies. For example, some systems used **FIFO** (first-in, first-out) replacement, where pages were simply placed in a queue when they enter the system; when a replacement occurs, the page on the tail of the queue (the “first-in” page) is evicted. FIFO has one great strength: it is quite simple to implement.

Let’s examine how FIFO does on our example reference stream (Table 22.2). We again begin our trace with three compulsory misses to pages 0, 1, and 2, and then hit on both 0 and 1. Next, page 3 is referenced, causing a miss; the replacement decision is easy with FIFO: pick the page that

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\(^1\)If you can, let us know! We can become rich together. Or, like the scientists who “discovered” cold fusion, widely scorned and mocked.
was the “first one” in (the cache state in the table is kept in FIFO order, with the first-in page on the left), which is page 0. Unfortunately, our next access is to page 0, causing another miss and replacement (of page 1). We then hit on page 3, but miss on 1 and 2, and finally hit on 3.

Comparing FIFO to optimal, FIFO does notably worse: a 36.4% hit rate (or 57.1% excluding compulsory misses). FIFO simply can’t determine the importance of blocks: even though page 0 had been accessed a number of times, FIFO still kicks it out, simply because it was the first one brought into memory.

Table 22.2: Tracing the FIFO Policy

<table>
<thead>
<tr>
<th>Access</th>
<th>Hit/Miss?</th>
<th>Evict</th>
<th>Resulting Cache State</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Miss</td>
<td>First-in→</td>
<td>0</td>
</tr>
<tr>
<td>1</td>
<td>Miss</td>
<td>First-in→</td>
<td>0, 1</td>
</tr>
<tr>
<td>2</td>
<td>Miss</td>
<td>First-in→</td>
<td>0, 1, 2</td>
</tr>
<tr>
<td>0</td>
<td>Hit</td>
<td>First-in→</td>
<td>0, 1, 2</td>
</tr>
<tr>
<td>1</td>
<td>Hit</td>
<td>First-in→</td>
<td>0, 1, 2</td>
</tr>
<tr>
<td>3</td>
<td>Miss</td>
<td>First-in→</td>
<td>1, 2, 3</td>
</tr>
<tr>
<td>0</td>
<td>Miss</td>
<td>First-in→</td>
<td>2, 3, 0</td>
</tr>
<tr>
<td>3</td>
<td>Hit</td>
<td>First-in→</td>
<td>2, 3, 0</td>
</tr>
<tr>
<td>1</td>
<td>Miss</td>
<td>First-in→</td>
<td>3, 0, 1</td>
</tr>
<tr>
<td>2</td>
<td>Miss</td>
<td>First-in→</td>
<td>0, 1, 2</td>
</tr>
<tr>
<td>1</td>
<td>Hit</td>
<td>First-in→</td>
<td>0, 1, 2</td>
</tr>
</tbody>
</table>

ASIDE: Belady’s Anomaly
Belady (of the optimal policy) and colleagues found an interesting reference stream that behaved a little unexpectedly [BNS69]. The memory-reference stream: 1, 2, 3, 4, 1, 2, 5, 1, 2, 3, 4, 5. The replacement policy they were studying was FIFO. The interesting part: how the cache hit rate changed when moving from a cache size of 3 to 4 pages.

In general, you would expect the cache hit rate to increase (get better) when the cache gets larger. But in this case, with FIFO, it gets worse! Calculate the hits and misses yourself and see. This odd behavior is generally referred to as Belady’s Anomaly (to the chagrin of his co-authors).

Some other policies, such as LRU, don’t suffer from this problem. Can you guess why? As it turns out, LRU has what is known as a stack property [M+70]. For algorithms with this property, a cache of size $N + 1$ naturally includes the contents of a cache of size $N$. Thus, when increasing the cache size, hit rate will either stay the same or improve. FIFO and Random (among others) clearly do not obey the stack property, and thus are susceptible to anomalous behavior.
Another similar replacement policy is Random, which simply picks a random page to replace under memory pressure. Random has properties similar to FIFO; it is simple to implement, but it doesn’t really try to be too intelligent in picking which blocks to evict. Let’s look at how Random does on our famous example reference stream (see Table 22.3).

Of course, how Random does depends entirely upon how lucky (or unlucky) Random gets in its choices. In the example above, Random does a little better than FIFO, and a little worse than optimal. In fact, we can run the Random experiment thousands of times and determine how it does in general. Figure 22.1 shows how many hits Random achieves over 10,000 trials, each with a different random seed. As you can see, sometimes (just over 40% of the time), Random is as good as optimal, achieving 6 hits on the example trace; sometimes it does much worse, achieving 2 hits or fewer. How Random does depends on the luck of the draw.

<table>
<thead>
<tr>
<th>Access</th>
<th>Hit/Miss?</th>
<th>Evict Cache State</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Miss</td>
<td>0</td>
</tr>
<tr>
<td>1</td>
<td>Miss</td>
<td>0, 1</td>
</tr>
<tr>
<td>2</td>
<td>Miss</td>
<td>0, 1, 2</td>
</tr>
<tr>
<td>0</td>
<td>Hit</td>
<td>0, 1, 2</td>
</tr>
<tr>
<td>1</td>
<td>Hit</td>
<td>0, 1, 2</td>
</tr>
<tr>
<td>3</td>
<td>Miss</td>
<td>0, 1, 2, 3</td>
</tr>
<tr>
<td>0</td>
<td>Miss</td>
<td>1, 2, 3</td>
</tr>
<tr>
<td>3</td>
<td>Hit</td>
<td>2, 3, 0</td>
</tr>
<tr>
<td>1</td>
<td>Miss</td>
<td>2, 0, 1</td>
</tr>
<tr>
<td>2</td>
<td>Hit</td>
<td>2, 0, 1</td>
</tr>
<tr>
<td>1</td>
<td>Hit</td>
<td>2, 0, 1</td>
</tr>
</tbody>
</table>

Table 22.3: Tracing the Random Policy

![Figure 22.1: Random Performance over 10,000 Trials](image-url)
### Table 22.4: Tracing the LRU Policy

<table>
<thead>
<tr>
<th>Access</th>
<th>Hit/Miss?</th>
<th>Evict</th>
<th>Resulting Cache State</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Miss</td>
<td>LRU→ 0</td>
<td>0</td>
</tr>
<tr>
<td>1</td>
<td>Miss</td>
<td>LRU→ 0, 1</td>
<td>0, 1</td>
</tr>
<tr>
<td>2</td>
<td>Miss</td>
<td>LRU→ 0, 1, 2</td>
<td>0, 1, 2</td>
</tr>
<tr>
<td>0</td>
<td>Hit</td>
<td>LRU→ 1, 2, 0</td>
<td>1, 2, 0</td>
</tr>
<tr>
<td>1</td>
<td>Hit</td>
<td>LRU→ 2, 0, 1</td>
<td>2, 0, 1</td>
</tr>
<tr>
<td>3</td>
<td>Miss</td>
<td>LRU→ 0, 1, 3</td>
<td>0, 1, 3</td>
</tr>
<tr>
<td>0</td>
<td>Hit</td>
<td>LRU→ 1, 3, 0</td>
<td>1, 3, 0</td>
</tr>
<tr>
<td>3</td>
<td>Hit</td>
<td>LRU→ 1, 0, 3</td>
<td>1, 0, 3</td>
</tr>
<tr>
<td>1</td>
<td>Hit</td>
<td>LRU→ 0, 3, 1</td>
<td>0, 3, 1</td>
</tr>
<tr>
<td>2</td>
<td>Miss</td>
<td>LRU→ 3, 1, 2</td>
<td>3, 1, 2</td>
</tr>
<tr>
<td>1</td>
<td>Hit</td>
<td>LRU→ 3, 2, 1</td>
<td>3, 2, 1</td>
</tr>
</tbody>
</table>

#### 22.5 Using History: LRU

Unfortunately, any policy as simple as FIFO or Random is likely to have a common problem: it might kick out an important page, one that is about to be referenced again. FIFO kicks out the page that was first brought in; if this happens to be a page with important code or data structures upon it, it gets thrown out anyhow, even though it will soon be paged back in. Thus, FIFO, Random, and similar policies are not likely to approach optimal; something smarter is needed.

As we did with scheduling policy, to improve our guess at the future, we once again lean on the past and use history as our guide. For example, if a program has accessed a page in the near past, it is likely to access it again in the near future.

One type of historical information a page-replacement policy could use is frequency; if a page has been accessed many times, perhaps it should not be replaced as it clearly has some value. A more commonly-used property of a page is its recency of access; the more recently a page has been accessed, perhaps the more likely it will be accessed again.

This family of policies is based on what people refer to as the principle of locality [D70], which basically is just an observation about programs and their behavior. What this principle says, quite simply, is that programs tend to access certain code sequences (e.g., in a loop) and data structures (e.g., an array accessed by the loop) quite frequently; we should thus try to use history to figure out which pages are important, and keep those pages in memory when it comes to eviction time.

And thus, a family of simple historically-based algorithms are born. The Least-Frequently-Used (LFU) policy replaces the least-frequently-used page when an eviction must take place. Similarly, the Least-Recently-Used (LRU) policy replaces the least-recently-used page. These algorithms are easy to remember: once you know the name, you know exactly what it does, which is an excellent property for a name.

To better understand LRU, let’s examine how LRU does on our exam-
There are two types of locality that programs tend to exhibit. The first is known as **spatial locality**, which states that if a page \( P \) is accessed, it is likely the pages around it (say \( P - 1 \) or \( P + 1 \)) will also likely be accessed. The second is **temporal locality**, which states that pages that have been accessed in the near past are likely to be accessed again in the near future. The assumption of the presence of these types of locality plays a large role in the caching hierarchies of hardware systems, which deploy many levels of instruction, data, and address-translation caching to help programs run fast when such locality exists.

Of course, the **principle of locality**, as it is often called, is no hard-and-fast rule that all programs must obey. Indeed, some programs access memory (or disk) in rather random fashion and don’t exhibit much or any locality in their access streams. Thus, while locality is a good thing to keep in mind while designing caches of any kind (hardware or software), it does not guarantee success. Rather, it is a heuristic that often proves useful in the design of computer systems.

**Aside: Types of Locality**

Of course, the principle of locality, as it is often called, is no hard-and-fast rule that all programs must obey. Indeed, some programs access memory (or disk) in rather random fashion and don’t exhibit much or any locality in their access streams. Thus, while locality is a good thing to keep in mind while designing caches of any kind (hardware or software), it does not guarantee success. Rather, it is a heuristic that often proves useful in the design of computer systems.

Let’s look at a few more examples in order to better understand how some of these policies behave. We’ll look at more complex workloads instead just a small trace of references. However, even these workloads are greatly simplified; a real study would include application traces.

Our first workload has no locality, which means that each reference is to a random page within the set of accessed pages. In this simple example, the workload accesses 100 unique pages over time, choosing the next page to refer to at random; overall, 10,000 pages are accessed. In the experiment, we vary the cache size from very small (1 page) to enough to hold all the unique pages (100 page), in order to see how each policy behaves over the range of cache sizes.

**22.6 Workload Examples**

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Figure 22.2 plots the results of the experiment for optimal, LRU, Random, and FIFO. The y-axis of the figure shows the hit rate that each policy achieves; the x-axis varies the cache size as described above.

We can draw a number of conclusions from the graph. First, when there is no locality in the workload, it doesn’t matter much which realistic policy you are using; LRU, FIFO, and Random all perform the same, with the hit rate exactly determined by the size of the cache. Second, when the cache is large enough to fit the entire workload, it also doesn’t matter which policy you use; all policies (even optimal) converge to a 100% hit rate when all the referenced blocks fit in cache. Finally, you can see that optimal performs noticeably better than the realistic policies; peeking into the future, if it were possible, does a much better job of replacement.

The next workload we examine is called the “80-20” workload, which exhibits locality: 80% of the references are made to 20% of the pages (the “hot” pages); the remaining 20% of the references are made to the remaining 80% of the pages (the “cold” pages). In our workload, there are a total 100 unique pages again; thus, “hot” pages are referred to most of the time, and “cold” pages the remainder. Figure 22.3 shows how the policies perform with this workload.

As you can see from the figure, while both random and FIFO do reasonably well, LRU does better, as it is more likely to hold onto the hot pages; as those pages have been referred to frequently in the past, they are likely to be referred to again in the near future. Optimal once again does better, showing that LRU’s historical information is not perfect.
You might now be wondering: is LRU’s improvement over Random and FIFO really that big of a deal? The answer, as usual, is “it depends.” If each miss is very costly (not uncommon), then even a small increase in hit rate (reduction in miss rate) can make a huge difference on performance. If misses are not so costly, then of course the benefits possible with LRU are not nearly as important.

Let’s look at one final workload. We call this one the “looping sequential” workload, as in it, we refer to 50 pages in sequence, starting at 0, then 1, ..., up to page 49, and then we loop, repeating those accesses, for a total of 10,000 accesses to 50 unique pages. The last graph in Figure 22.4 shows the behavior of the policies under this workload.

This workload, common in many applications (including important commercial applications such as databases [CD85]), represents a worst-case for both LRU and FIFO. These algorithms, under a looping-sequential workload, kick out older pages; unfortunately, due to the looping nature of the workload, these older pages are going to be accessed sooner than the pages that the policies prefer to keep in cache. Indeed, even with a cache of size 49, a looping-sequential workload of 50 pages results in a 0% hit rate. Interestingly, Random fares notably better, not quite approaching optimal, but at least achieving a non-zero hit rate. Turns out that random has some nice properties; one such property is not having weird corner-case behaviors.
22.7 Implementing Historical Algorithms

As you can see, an algorithm such as LRU can generally do a better job than simpler policies like FIFO or Random, which may throw out important pages. Unfortunately, historical policies present us with a new challenge: how do we implement them?

Let’s take, for example, LRU. To implement it perfectly, we need to do a lot of work. Specifically, upon each page access (i.e., each memory access, whether an instruction fetch or a load or store), we must update some data structure to move this page to the front of the list (i.e., the MRU side). Contrast this to FIFO, where the FIFO list of pages is only accessed when a page is evicted (by removing the first-in page) or when a new page is added to the list (to the last-in side). To keep track of which pages have been least- and most-recently used, the system has to do some accounting work on every memory reference. Clearly, without great care, such accounting could greatly reduce performance.

One method that could help speed this up is to add a little bit of hardware support. For example, a machine could update, on each page access, a time field in memory (for example, this could be in the per-process page table, or just in some separate array in memory, with one entry per physical page of the system). Thus, when a page is accessed, the time field would be set, by hardware, to the current time. Then, when replacing a page, the OS could simply scan all the time fields in the system to find the least-recently-used page.
Unfortunately, as the number of pages in a system grows, scanning a huge array of times just to find the absolute least-recently-used page is prohibitively expensive. Imagine a modern machine with 4GB of memory, chopped into 4KB pages. This machine has 1 million pages, and thus finding the LRU page will take a long time, even at modern CPU speeds. Which begs the question: do we really need to find the absolute oldest page to replace? Can we instead survive with an approximation?

**Crux: How To Implement An LRU Replacement Policy**
Given that it will be expensive to implement perfect LRU, can we approximate it in some way, and still obtain the desired behavior?

### 22.8 Approximating LRU

As it turns out, the answer is yes: approximating LRU is more feasible from a computational-overhead standpoint, and indeed it is what many modern systems do. The idea requires some hardware support, in the form of a use bit (sometimes called the reference bit), the first of which was implemented in the first system with paging, the Atlas one-level store [KE+62]. There is one use bit per page of the system, and the use bits live in memory somewhere (they could be in the per-process page tables, for example, or just in an array somewhere). Whenever a page is referenced (i.e., read or written), the use bit is set by hardware to 1. The hardware never clears the bit, though (i.e., sets it to 0); that is the responsibility of the OS.

How does the OS employ the use bit to approximate LRU? Well, there could be a lot of ways, but with the clock algorithm [C69], one simple approach was suggested. Imagine all the pages of the system arranged in a circular list. A clock hand points to some particular page to begin with (it doesn’t really matter which). When a replacement must occur, the OS checks if the currently-pointed to page $P$ has a use bit of 1 or 0. If 1, this implies that page $P$ was recently used and thus is not a good candidate for replacement. Thus, the clock hand is incremented to the next page $P + 1$, and the use bit for $P$ set to 0 (cleared). The algorithm continues until it finds a use bit that is set to 0, implying this page has not been recently used (or, in the worst case, that all pages have been and that we have now searched through the entire set of pages, clearing all the bits).

Note that this approach is not the only way to employ a use bit to approximate LRU. Indeed, any approach which periodically clears the use bits and then differentiates between which pages have use bits of 1 versus 0 to decide which to replace would be fine. The clock algorithm of Corbato’s was just one early approach which met with some success, and had the nice property of not repeatedly scanning through all of memory looking for an unused page.
The behavior of a clock algorithm variant is shown in Figure 22.5. This variant randomly scans pages when doing a replacement; when it encounters a page with a reference bit set to 1, it clears the bit (i.e., sets it to 0); when it finds a page with the reference bit set to 0, it chooses it as its victim. As you can see, although it doesn’t do quite as well as perfect LRU, it does better than approaches that don’t consider history at all.

### 22.9 Considering Dirty Pages

One small modification to the clock algorithm (also originally suggested by Corbato [C69]) that is commonly made is the additional consideration of whether a page has been modified or not while in memory. The reason for this: if a page has been modified and is thus dirty, it must be written back to disk to evict it, which is expensive. If it has not been modified (and is thus clean), the eviction is free; the physical frame can simply be reused for other purposes without additional I/O. Thus, some VM systems prefer to evict clean pages over dirty pages.

To support this behavior, the hardware should include a modified bit (a.k.a. dirty bit). This bit is set any time a page is written, and thus can be incorporated into the page-replacement algorithm. The clock algorithm, for example, could be changed to scan for pages that are both unused and clean to evict first; failing to find those, then for unused pages that are dirty; etc.
22.10 Other VM Policies

Page replacement is not the only policy the VM subsystem employs (though it may be the most important). For example, the OS also has to decide when to bring a page into memory. This policy, sometimes called the page selection policy (as it was called by Denning [D70]), presents the OS with some different options.

For most pages, the OS simply uses demand paging, which means the OS brings the page into memory when it is accessed, “on demand” as it were. Of course, the OS could guess that a page is about to be used, and thus bring it in ahead of time; this behavior is known as prefetching and should only be done when there is reasonable chance of success. For example, some systems will assume that if a code page $P$ is brought into memory, that code page $P+1$ will likely soon be accessed and thus should be brought into memory too.

Another policy determines how the OS writes pages out to disk. Of course, they could simply be written out one at a time; however, many systems instead collect a number of pending writes together in memory and write them to disk in one (more efficient) write. This behavior is usually called clustering or simply grouping of writes, and is effective because of the nature of disk drives, which perform a single large write more efficiently than many small ones.

22.11 Thrashing

Before closing, we address one final question: what should the OS do when memory is simply oversubscribed, and the memory demands of the set of running processes simply exceed the available physical memory? In this case, the system will constantly be paging, a condition sometimes referred to as thrashing [D70].

Some earlier operating systems had a fairly sophisticated set of mechanisms to both detect and cope with thrashing when it took place. For example, given a set of processes, a system could decide not to run a subset of processes, with the hope that the reduced set of processes working sets (the pages that they are using actively) fit in memory and thus can make progress. This approach, generally known as admission control, states that it is sometimes better to do less work well than to try to do everything at once poorly, a situation we often encounter in real life as well as in modern computer systems (sadly).

Some current systems take a more draconian approach to memory overload. For example, some versions of Linux run an out-of-memory killer when memory is oversubscribed; this daemon chooses a memory-intensive process and kills it, thus reducing memory in a none-too-subtle manner. While successful at reducing memory pressure, this approach can have problems, if, for example, it kills the X server and thus renders any applications requiring the display unusable.
22.12 Summary

We have seen the introduction of a number of page-replacement (and other) policies, which are part of the VM subsystem of all modern operating systems. Modern systems add some tweaks to straightforward LRU approximations like clock; for example, scan resistance is an important part of many modern algorithms, such as ARC [MM03]. Scan-resistant algorithms are usually LRU-like but also try to avoid the worst-case behavior of LRU, which we saw with the looping-sequential workload. Thus, the evolution of page-replacement algorithms continues.

However, in many cases the importance of said algorithms has decreased, as the discrepancy between memory-access and disk-access times has increased. Because paging to disk is so expensive, the cost of frequent paging is prohibitive. Thus, the best solution to excessive paging is often a simple (if intellectually dissatisfying) one: buy more memory.
References

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Remzi H. Arpaci-Dusseau
ACM TOCS, 21:1, February 2003
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Laszlo A. Belady
The paper that introduces the simple way to compute the optimal behavior of a policy (the MIN algorithm).

L. A. Belady and R. A. Nelson and G. S. Shedler
Communications of the ACM, 12:6, June 1969
Introduction of the little sequence of memory references known as Belady’s Anomaly. How do Nelson and Shedler feel about this name, we wonder?

Hong-Tai Chou and David J. DeWitt
VLDB ’85, Stockholm, Sweden, August 1985
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F. J. Corbato
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[D70] “Virtual Memory”
Peter J. Denning
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Malcolm C. Easton and Ronald Fagin
Communications of the ACM, 21:10, October 1978
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John Hennessy and David Patterson
Morgan-Kaufmann, 2006
A great and marvelous book about computer architecture. Read it!

[H87] “Aspects of Cache Memory and Instruction Buffer Performance”
Mark D. Hill
Mark Hill, in his dissertation work, introduced the Three C’s, which later gained wide popularity with its inclusion in H&P [HP06]. The quote from therein: “I have found it useful to partition misses ... into three components intuitively based on the cause of the misses (page 49).”
BEYOND PHYSICAL MEMORY: POLICIES

[KE+62] “One-level Storage System”  
T. Kilburn, and D.B.G. Edwards and M.J. Lanigan and F.H. Sumner  
IRE Trans. EC-11:2, 1962  
Although Atlas had a use bit, it only had a very small number of pages, and thus the scanning of the use bits in large memories was not a problem the authors solved.

[M+70] “Evaluation Techniques for Storage Hierarchies”  
R. L. Mattson, J. Gecsei, D. R. Slutz, I. L. Traiger  
IBM Systems Journal, Volume 9:2, 1970  
A paper that is mostly about how to simulate cache hierarchies efficiently; certainly a classic in that regard, as well for its excellent discussion of some of the properties of various replacement algorithms. Can you figure out why the stack property might be useful for simulating a lot of different-sized caches at once?

[MM03] “ARC: A Self-Tuning, Low Overhead Replacement Cache”  
Nimrod Megiddo and Dharmendra S. Modha  
FAST 2003, February 2003, San Jose, California  
An excellent modern paper about replacement algorithms, which includes a new policy, ARC, that is now used in some systems. Recognized in 2014 as a “Test of Time” award winner by the storage systems community at the FAST ’14 conference.
Homework

This simulator, `paging-policy.py`, allows you to play around with different page-replacement policies. See the README for details.

Questions

- Generate random addresses with the following arguments: `-s 0 -n 10`, `-s 1 -n 10`, and `-s 2 -n 10`. Change the policy from FIFO, to LRU, to OPT. Compute whether each access in said address traces are hits or misses.
- For a cache of size 5, generate worst-case address reference streams for each of the following policies: FIFO, LRU, and MRU (worst-case reference streams cause the most misses possible. For the worst case reference streams, how much bigger of a cache is needed to improve performance dramatically and approach OPT?
- Generate a random trace (use python or perl). How would you expect the different policies to perform on such a trace?
- Now generate a trace with some locality. How can you generate such a trace? How does LRU perform on it? How much better than RAND is LRU? How does CLOCK do? How about CLOCK with different numbers of clock bits?
- Use a program like `valgrind` to instrument a real application and generate a virtual page reference stream. For example, running `valgrind --tool=lackey --trace-mem=yes ls` will output a nearly-complete reference trace of every instruction and data reference made by the program `ls`. To make this useful for the simulator above, you'll have to first transform each virtual memory reference into a virtual page-number reference (done by masking off the offset and shifting the resulting bits downward). How big of a cache is needed for your application trace in order to satisfy a large fraction of requests? Plot a graph of its working set as the size of the cache increases.
The VAX/VMS Virtual Memory System

Before we end our study of virtual memory, let us take a closer look at one particularly clean and well done virtual memory manager, that found in the VAX/VMS operating system [LL82]. In this note, we will discuss the system to illustrate how some of the concepts brought forth in earlier chapters together in a complete memory manager.

23.1 Background

The VAX-11 minicomputer architecture was introduced in the late 1970's by Digital Equipment Corporation (DEC). DEC was a massive player in the computer industry during the era of the mini-computer; unfortunately, a series of bad decisions and the advent of the PC slowly (but surely) led to their demise [C03]. The architecture was realized in a number of implementations, including the VAX-11/780 and the less powerful VAX-11/750.

The OS for the system was known as VAX/VMS (or just plain VMS), one of whose primary architects was Dave Cutler, who later led the effort to develop Microsoft's Windows NT [C93]. VMS had the general problem that it would be run on a broad range of machines, including very inexpensive VAXen (yes, that is the proper plural) to extremely high-end and powerful machines in the same architecture family. Thus, the OS had to have mechanisms and policies that worked (and worked well) across this huge range of systems.

THE CRUX: HOW TO AVOID THE CURSE OF GENERALITY

Operating systems often have a problem known as “the curse of generality”, where they are tasked with general support for a broad class of applications and systems. The fundamental result of the curse is that the OS is not likely to support any one installation very well. In the case of VMS, the curse was very real, as the VAX-11 architecture was realized in a number of different implementations. Thus, how can an OS be built so as to run effectively on a wide range of systems?
As an additional issue, VMS is an excellent example of software innovations used to hide some of the inherent flaws of the architecture. Although the OS often relies on the hardware to build efficient abstractions and illusions, sometimes the hardware designers don’t quite get everything right; in the VAX hardware, we’ll see a few examples of this, and what the VMS operating system does to build an effective, working system despite these hardware flaws.

23.2 Memory Management Hardware

The VAX-11 provided a 32-bit virtual address space per process, divided into 512-byte pages. Thus, a virtual address consisted of a 23-bit VPN and a 9-bit offset. Further, the upper two bits of the VPN were used to differentiate which segment the page resided within; thus, the system was a hybrid of paging and segmentation, as we saw previously.

The lower-half of the address space was known as “process space” and is unique to each process. In the first half of process space (known as P0), the user program is found, as well as a heap which grows downward. In the second half of process space (P1), we find the stack, which grows upwards. The upper-half of the address space is known as system space (S), although only half of it is used. Protected OS code and data reside here, and the OS is in this way shared across processes.

One major concern of the VMS designers was the incredibly small size of pages in the VAX hardware (512 bytes). This size, chosen for historical reasons, has the fundamental problem of making simple linear page tables excessively large. Thus, one of the first goals of the VMS designers was to make sure that VMS would not overwhelm memory with page tables.

The system reduced the pressure page tables place on memory in two ways. First, by segmenting the user address space into two, the VAX-11 provides a page table for each of these regions (P0 and P1) per process; thus, no page-table space is needed for the unused portion of the address space between the stack and the heap. The base and bounds registers are used as you would expect; a base register holds the address of the page table for that segment, and the bounds holds its size (i.e., number of page-table entries).

Second, the OS reduces memory pressure even further by placing user page tables (for P0 and P1, thus two per process) in kernel virtual memory. Thus, when allocating or growing a page table, the kernel allocates space out of its own virtual memory, in segment S. If memory comes under severe pressure, the kernel can swap pages of these page tables out to disk, thus making physical memory available for other uses.

Putting page tables in kernel virtual memory means that address translation is even further complicated. For example, to translate a virtual address in P0 or P1, the hardware has to first try to look up the page-table entry for that page in its page table (the P0 or P1 page table for that pro-
cess); in doing so, however, the hardware may first have to consult the system page table (which lives in physical memory); with that translation complete, the hardware can learn the address of the page of the page table, and then finally learn the address of the desired memory access. All of this, fortunately, is made faster by the VAX’s hardware-managed TLBs, which usually (hopefully) circumvent this laborious lookup.

23.3 A Real Address Space

One neat aspect of studying VMS is that we can see how a real address space is constructed (Figure 23.1. Thus far, we have assumed a simple address space of just user code, user data, and user heap, but as we can see above, a real address space is notably more complex.
Aside: Why Null Pointer Accesses Cause Seg Faults

You should now have a good understanding of exactly what happens on a null-pointer dereference. A process generates a virtual address of 0, by doing something like this:

```c
int *p = NULL; // set p = 0
*p = 10; // try to store value 10 to virtual address 0
```

The hardware tries to look up the VPN (also 0 here) in the TLB, and suffers a TLB miss. The page table is consulted, and the entry for VPN 0 is found to be marked invalid. Thus, we have an invalid access, which transfers control to the OS, which likely terminates the process (on UNIX systems, processes are sent a signal which allows them to react to such a fault; if uncaught, however, the process is killed).

For example, the code segment never begins at page 0. This page, instead, is marked inaccessible, in order to provide some support for detecting null-pointer accesses. Thus, one concern when designing an address space is support for debugging, which the inaccessible zero page provides here in some form.

Perhaps more importantly, the kernel virtual address space (i.e., its data structures and code) is a part of each user address space. On a context switch, the OS changes the P0 and P1 registers to point to the appropriate page tables of the soon-to-be-run process; however, it does not change the S base and bound registers, and as a result the “same” kernel structures are mapped into each user address space.

The kernel is mapped into each address space for a number of reasons. This construction makes life easier for the kernel; when, for example, the OS is handed a pointer from a user program (e.g., on a write() system call), it is easy to copy data from that pointer to its own structures. The OS is naturally written and compiled, without worry of where the data it is accessing comes from. If in contrast the kernel were located entirely in physical memory, it would be quite hard to do things like swap pages of the page table to disk; if the kernel were given its own address space, moving data between user applications and the kernel would again be complicated and painful. With this construction (now used widely), the kernel appears almost as a library to applications, albeit a protected one.

One last point about this address space relates to protection. Clearly, the OS does not want user applications reading or writing OS data or code. Thus, the hardware must support different protection levels for pages to enable this. The VAX did so by specifying, in protection bits in the page table, what privilege level the CPU must be at in order to access a particular page. Thus, system data and code are set to a higher level of protection than user data and code; an attempted access to such information from user code will generate a trap into the OS, and (you guessed it) the likely termination of the offending process.
23.4 Page Replacement

The page table entry (PTE) in VAX contains the following bits: a valid bit, a protection field (4 bits), a modify (or dirty) bit, a field reserved for OS use (5 bits), and finally a physical frame number (PFN) to store the location of the page in physical memory. The astute reader might note: no reference bit! Thus, the VMS replacement algorithm must make do without hardware support for determining which pages are active.

The developers were also concerned about memory hogs, programs that use a lot of memory and make it hard for other programs to run. Most of the policies we have looked at thus far are susceptible to such hogging; for example, LRU is a global policy that doesn’t share memory fairly among processes.

Segmented FIFO

To address these two problems, the developers came up with the segmented FIFO replacement policy [RL81]. The idea is simple: each process has a maximum number of pages it can keep in memory, known as its resident set size (RSS). Each of these pages is kept on a FIFO list; when a process exceeds its RSS, the “first-in” page is evicted. FIFO clearly does not need any support from the hardware, and is thus easy to implement.

Of course, pure FIFO does not perform particularly well, as we saw earlier. To improve FIFO’s performance, VMS introduced two second-chance lists where pages are placed before getting evicted from memory, specifically a global clean-page free list and dirty-page list. When a process \( P \) exceeds its RSS, a page is removed from its per-process FIFO; if clean (not modified), it is placed on the end of the clean-page list; if dirty (modified), it is placed on the end of the dirty-page list.

If another process \( Q \) needs a free page, it takes the first free page off of the global clean list. However, if the original process \( P \) faults on that page before it is reclaimed, \( P \) reclaims it from the free (or dirty) list, thus avoiding a costly disk access. The bigger these global second-chance lists are, the closer the segmented FIFO algorithm performs to LRU [RL81].

Page Clustering

Another optimization used in VMS also helps overcome the small page size in VMS. Specifically, with such small pages, disk I/O during swapping could be highly inefficient, as disks do better with large transfers. To make swapping I/O more efficient, VMS adds a number of optimizations, but most important is clustering. With clustering, VMS groups large batches of pages together from the global dirty list, and writes them to disk in one fell swoop (thus making them clean). Clustering is used in most modern systems, as the freedom to place pages anywhere within swap space lets the OS group pages, perform fewer and bigger writes, and thus improve performance.
ASIDE: EMULATING REFERENCE BITS
As it turns out, you don’t need a hardware reference bit in order to get some notion of which pages are in use in a system. In fact, in the early 1980’s, Babaoglu and Joy showed that protection bits on the VAX can be used to emulate reference bits [BJ81]. The basic idea: if you want to gain some understanding of which pages are actively being used in a system, mark all of the pages in the page table as inaccessible (but keep around the information as to which pages are really accessible by the process, perhaps in the “reserved OS field” portion of the page table entry). When a process accesses a page, it will generate a trap into the OS; the OS will then check if the page really should be accessible, and if so, revert the page to its normal protections (e.g., read-only, or read-write). At the time of a replacement, the OS can check which pages remain marked inaccessible, and thus get an idea of which pages have not been recently used.

The key to this “emulation” of reference bits is reducing overhead while still obtaining a good idea of page usage. The OS must not be too aggressive in marking pages inaccessible, or overhead would be too high. The OS also must not be too passive in such marking, or all pages will end up referenced; the OS will again have no good idea which page to evict.

23.5 Other Neat VM Tricks

VMS had two other now-standard tricks: demand zeroing and copy-on-write. We now describe these lazy optimizations.

One form of laziness in VMS (and most modern systems) is demand zeroing of pages. To understand this better, let’s consider the example of adding a page to your address space, say in your heap. In a naive implementation, the OS responds to a request to add a page to your heap by finding a page in physical memory, zeroing it (required for security; otherwise you’d be able to see what was on the page from when some other process used it!), and then mapping it into your address space (i.e., setting up the page table to refer to that physical page as desired). But the naive implementation can be costly, particularly if the page does not get used by the process.

With demand zeroing, the OS instead does very little work when the page is added to your address space; it puts an entry in the page table that marks the page inaccessible. If the process then reads or writes the page, a trap into the OS takes place. When handling the trap, the OS notices (usually through some bits marked in the “reserved for OS” portion of the page table entry) that this is actually a demand-zero page; at this point, the OS then does the needed work of finding a physical page, zeroing it, and mapping it into the process’s address space. If the process never accesses the page, all of this work is avoided, and thus the virtue of demand zeroing.
TIP: BE LAZY
Being lazy can be a virtue in both life as well as in operating systems. Laziness can put off work until later, which is beneficial within an OS for a number of reasons. First, putting off work might reduce the latency of the current operation, thus improving responsiveness; for example, operating systems often report that writes to a file succeeded immediately, and only write them to disk later in the background. Second, and more importantly, laziness sometimes obviates the need to do the work at all; for example, delaying a write until the file is deleted removes the need to do the write at all. Laziness is also good in life: for example, by putting off your OS project, you may find that the project specification bugs are worked out by your fellow classmates; however, the class project is unlikely to get canceled, so being too lazy may be problematic, leading to a late project, bad grade, and a sad professor. Don’t make professors sad!

Another cool optimization found in VMS (and again, in virtually every modern OS) is **copy-on-write** (COW for short). The idea, which goes at least back to the TENEX operating system [BB+72], is simple: when the OS needs to copy a page from one address space to another, instead of copying it, it can map it into the target address space and mark it read-only in both address spaces. If both address spaces only read the page, no further action is taken, and thus the OS has affected a fast copy without actually moving any data.

If, however, one of the address spaces does indeed try to write to the page, it will trap into the OS. The OS will then notice that the page is a COW page, and thus (lazily) allocate a new page, fill it with the data, and map this new page into the address space of the faulting process. The process then continues and now has its own private copy of the page.

COW is useful for a number of reasons. Certainly any sort of shared library can be mapped copy-on-write into the address spaces of many processes, saving valuable memory space. In UNIX systems, COW is even more critical, due to the semantics of `fork()` and `exec()`. As you might recall, `fork()` creates an exact copy of the address space of the caller; with a large address space, making such a copy is slow and data intensive. Even worse, most of the address space is immediately over-written by a subsequent call to `exec()`, which overlays the calling process’s address space with that of the soon-to-be-exec’d program. By instead performing a copy-on-write `fork()`, the OS avoids much of the needless copying and thus retains the correct semantics while improving performance.
23.6 Summary

You have now seen a top-to-bottom review of an entire virtual memory system. Hopefully, most of the details were easy to follow, as you should have already had a good understanding of most of the basic mechanisms and policies. More detail is available in the excellent (and short) paper by Levy and Lipman [LL82]; we encourage you to read it, a great way to see what the source material behind these chapters is like.

You should also learn more about the state of the art by reading about Linux and other modern systems when possible. There is a lot of source material out there, including some reasonable books [BC05]. One thing that will amaze you: how classic ideas, found in old papers such as this one on VAX/VMS, still influence how modern operating systems are built.
References

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Daniel G. Bobrow, Jerry D. Burchfiel, Daniel L. Murphy, Raymond S. Tomlinson
Communications of the ACM, Volume 15, March 1972
An early time-sharing OS where a number of good ideas came from. Copy-on-write was just one of those; inspiration for many other aspects of modern systems, including process management, virtual memory, and file systems are found herein.

[BJ81] “Converting a Swap-Based System to do Paging in an Architecture Lacking Page-Reference Bits”
Ozalp Babaoglu and William N. Joy
SOSP ’81, December 1981, Pacific Grove, California
A clever idea paper on how to exploit existing protection machinery within a machine in order to emulate reference bits. The idea came from the group at Berkeley working on their own version of UNIX, known as the Berkeley Systems Distribution, or BSD. The group was heavily influential in the development of UNIX, in virtual memory, file systems, and networking.

[BC05] “Understanding the Linux Kernel (Third Edition)”
Daniel P. Bovet and Marco Cesati
O’Reilly Media, November 2005
One of the many books you can find on Linux. They go out of date quickly, but many of the basics remain and are worth reading about.

[C03] “The Innovator’s Dilemma”
Clayton M. Christenson
Harper Paperbacks, January 2003
A fantastic book about the disk-drive industry and how new innovations disrupt existing ones. A good read for business majors and computer scientists alike. Provides insight on how large and successful companies completely fail.

[C93] “Inside Windows NT”
Helen Custer and David Solomon
Microsoft Press, 1993
The book about Windows NT that explains the system top to bottom, in more detail than you might like. But seriously, a pretty good book.

[LL82] “Virtual Memory Management in the VAX/VMS Operating System”
Henry M. Levy, Peter H. Lipman
IEEE Computer, Volume 15, Number 3 (March 1982) Read the original source of most of this material; it is a concise and easy read. Particularly important if you wish to go to graduate school, where all you do is read papers, work, read some more papers, work more, eventually write a paper, and then work some more. But it is fun!

[RL81] “Segmented FIFO Page Replacement”
Rollins Turner and Henry Levy
SIGMETRICS ’81
A short paper that shows for some workloads, segmented FIFO can approach the performance of LRU.
Summary Dialogue on Memory Virtualization

Student: (Gulps) Wow, that was a lot of material.
Professor: Yes, and?
Student: Well, how am I supposed to remember it all? You know, for the exam?
Professor: Goodness, I hope that’s not why you are trying to remember it.
Student: Why should I then?
Professor: Come on, I thought you knew better. You’re trying to learn something here, so that when you go off into the world, you’ll understand how systems actually work.
Student: Hmm... can you give an example?
Professor: Sure! One time back in graduate school, my friends and I were measuring how long memory accesses took, and once in a while the numbers were way higher than we expected; we thought all the data was fitting nicely into the second-level hardware cache, you see, and thus should have been really fast to access.
Student: (nods)
Professor: We couldn’t figure out what was going on. So what do you do in such a case? Easy, ask a professor! So we went and asked one of our professors, who looked at the graph we had produced, and simply said “TLB”. Aha! Of course, TLB misses! Why didn’t we think of that? Having a good model of how virtual memory works helps diagnose all sorts of interesting performance problems.
Student: I think I see. I’m trying to build these mental models of how things work, so that when I’m out there working on my own, I won’t be surprised when a system doesn’t quite behave as expected. I should even be able to anticipate how the system will work just by thinking about it.
Professor: Exactly. So what have you learned? What’s in your mental model of how virtual memory works?
Student: Well, I think I now have a pretty good idea of what happens when memory is referenced by a process, which, as you’ve said many times, happens
on each instruction fetch as well as explicit loads and stores.

**Professor:** Sounds good – tell me more.

**Student:** Well, one thing I’ll always remember is that the addresses we see in a user program, written in C for example...

**Professor:** What other language is there?

**Student:** (continuing) ... Yes, I know you like C. So do I! Anyhow, as I was saying, I now really know that all addresses that we can observe within a program are virtual addresses; that I, as a programmer, am just given this illusion of where data and code are in memory. I used to think it was cool that I could print the address of a pointer, but now I find it frustrating – it’s just a virtual address! I can’t see the real physical address where the data lives.

**Professor:** Nope, the OS definitely hides that from you. What else?

**Student:** Well, I think the TLB is a really key piece, providing the system with a small hardware cache of address translations. Page tables are usually quite large and hence live in big and slow memories. Without that TLB, programs would certainly run a great deal more slowly. Seems like the TLB truly makes virtualizing memory possible. I couldn’t imagine building a system without one! And I shudder at the thought of a program with a working set that exceeds the coverage of the TLB: with all those TLB misses, it would be hard to watch.

**Professor:** Yes, cover the eyes of the children! Beyond the TLB, what did you learn?

**Student:** I also now understand that the page table is one of those data structures you need to know about; it’s just a data structure, though, and that means almost any structure could be used. We started with simple structures, like arrays (a.k.a. linear page tables), and advanced all the way up to multi-level tables (which look like trees), and even crazier things like pageable page tables in kernel virtual memory. All to save a little space in memory!

**Professor:** Indeed.

**Student:** And here’s one more important thing: I learned that the address translation structures need to be flexible enough to support what programmers want to do with their address spaces. Structures like the multi-level table are perfect in this sense; they only create table space when the user needs a portion of the address space, and thus there is little waste. Earlier attempts, like the simple base and bounds register, just weren’t flexible enough; the structures need to match what users expect and want out of their virtual memory system.

**Professor:** That’s a nice perspective. What about all of the stuff we learned about swapping to disk?

**Student:** Well, it’s certainly fun to study, and good to know how page replacement works. Some of the basic policies are kind of obvious (like LRU, for example), but building a real virtual memory system seems more interesting, like we saw in the VMS case study. But somehow, I found the mechanisms more interesting, and the policies less so.
Professor: Oh, why is that?

Student: Well, as you said, in the end the best solution to policy problems is simple: buy more memory. But the mechanisms you need to understand to know how stuff really works. Speaking of which...

Professor: Yes?

Student: Well, my machine is running a little slowly these days... and memory certainly doesn’t cost that much...

Professor: Oh fine, fine! Here’s a few bucks. Go and get yourself some DRAM, cheapskate.

Student: Thanks professor! I’ll never swap to disk again – or, if I do, at least I’ll know what’s actually going on!
Part II

Concurrency
A Dialogue on Concurrency

Professor: And thus we reach the second of our three pillars of operating systems: concurrency.

Student: I thought there were four pillars...?

Professor: Nope, that was in an older version of the book.

Student: Umm... OK. So what is concurrency, oh wonderful professor?

Professor: Well, imagine we have a peach –

Student: (interrupting) Peaches again! What is it with you and peaches?

Professor: Ever read T.S. Eliot? The Love Song of J. Alfred Prufrock, “Do I dare to eat a peach”, and all that fun stuff?

Student: Oh yes! In English class in high school. Great stuff! I really liked the part where –

Professor: (interrupting) This has nothing to do with that – I just like peaches. Anyhow, imagine there are a lot of peaches on a table, and a lot of people who wish to eat them. Let’s say we did it this way: each eater first identifies a peach visually, and then tries to grab it and eat it. What is wrong with this approach?

Student: Hmmm... seems like you might see a peach that somebody else also sees. If they get there first, when you reach out, no peach for you!

Professor: Exactly! So what should we do about it?

Student: Well, probably develop a better way of going about this. Maybe form a line, and when you get to the front, grab a peach and get on with it.

Professor: Good! But what’s wrong with your approach?

Student: Sheesh, do I have to do all the work?

Professor: Yes.

Student: OK, let me think. Well, we used to have many people grabbing for peaches all at once, which is faster. But in my way, we just go one at a time, which is correct, but quite a bit slower. The best kind of approach would be fast and correct, probably.
Professor: You are really starting to impress. In fact, you just told us everything we need to know about concurrency! Well done.

Student: I did? I thought we were just talking about peaches. Remember, this is usually a part where you make it about computers again.

Professor: Indeed. My apologies! One must never forget the concrete. Well, as it turns out, there are certain types of programs that we call multi-threaded applications; each thread is kind of like an independent agent running around in this program, doing things on the program’s behalf. But these threads access memory, and for them, each spot of memory is kind of like one of those peaches. If we don’t coordinate access to memory between threads, the program won’t work as expected. Make sense?

Student: Kind of. But why do we talk about this in an OS class? Isn’t that just application programming?

Professor: Good question! A few reasons, actually. First, the OS must support multi-threaded applications with primitives such as locks and condition variables, which we’ll talk about soon. Second, the OS itself was the first concurrent program – it must access its own memory very carefully or many strange and terrible things will happen. Really, it can get quite grisly.

Student: I see. Sounds interesting. There are more details, I imagine?

Professor: Indeed there are...
Thus far, we have seen the development of the basic abstractions that the OS performs. We have seen how to take a single physical CPU and turn it into multiple virtual CPUs, thus enabling the illusion of multiple programs running at the same time. We have also seen how to create the illusion of a large, private virtual memory for each process; this abstraction of the address space enables each program to behave as if it has its own memory when indeed the OS is secretly multiplexing address spaces across physical memory (and sometimes, disk).

In this note, we introduce a new abstraction for a single running process: that of a thread. Instead of our classic view of a single point of execution within a program (i.e., a single PC where instructions are being fetched from and executed), a multi-threaded program has more than one point of execution (i.e., multiple PCs, each of which is being fetched and executed from). Perhaps another way to think of this is that each thread is very much like a separate process, except for one difference: they share the same address space and thus can access the same data.

The state of a single thread is thus very similar to that of a process. It has a program counter (PC) that tracks where the program is fetching instructions from. Each thread has its own private set of registers it uses for computation; thus, if there are two threads that are running on a single processor, when switching from running one (T1) to running the other (T2), a context switch must take place. The context switch between threads is quite similar to the context switch between processes, as the register state of T1 must be saved and the register state of T2 restored before running T2. With processes, we saved state to a process control block (PCB); now, we’ll need one or more thread control blocks (TCBs) to store the state of each thread of a process. There is one major difference, though, in the context switch we perform between threads as compared to processes: the address space remains the same (i.e., there is no need to switch which page table we are using).

One other major difference between threads and processes concerns the stack. In our simple model of the address space of a classic process (which we can now call a single-threaded process), there is a single stack, usually residing at the bottom of the address space (Figure 26.1, left).
However, in a multi-threaded process, each thread runs independently and of course may call into various routines to do whatever work it is doing. Instead of a single stack in the address space, there will be one per thread. Let’s say we have a multi-threaded process that has two threads in it; the resulting address space looks different (Figure 26.1, right).

In this figure, you can see two stacks spread throughout the address space of the process. Thus, any stack-allocated variables, parameters, return values, and other things that we put on the stack will be placed in what is sometimes called thread-local storage, i.e., the stack of the relevant thread.

You might also notice how this ruins our beautiful address space layout. Before, the stack and heap could grow independently and trouble only arose when you ran out of room in the address space. Here, we no longer have such a nice situation. Fortunately, this is usually OK, as stacks do not generally have to be very large (the exception being in programs that make heavy use of recursion).

26.1 An Example: Thread Creation

Let’s say we wanted to run a program that created two threads, each of which was doing some independent work, in this case printing “A” or “B”. The code is shown in Figure 26.2.

The main program creates two threads, each of which will run the function mythread(), though with different arguments (the string A or B). Once a thread is created, it may start running right away (depending on the whims of the scheduler); alternately, it may be put in a “ready” but not “running” state and thus not run yet. After creating the two threads (T1 and T2), the main thread calls pthread_join(), which waits for a particular thread to complete.
Let us examine the possible execution ordering of this little program. In the execution diagram (Table 26.1), time increases in the downwards direction, and each column shows when a different thread (the main one, or Thread 1, or Thread 2) is running.

Note, however, that this ordering is not the only possible ordering. In fact, given a sequence of instructions, there are quite a few, depending on which thread the scheduler decides to run at a given point. For example, once a thread is created, it may run immediately, which would lead to the execution shown in Table 26.2.

We also could even see “B” printed before “A”, if, say, the scheduler decided to run Thread 2 first even though Thread 1 was created earlier; there is no reason to assume that a thread that is created first will run first. Table 26.3 shows this final execution ordering, with Thread 2 getting to strut its stuff before Thread 1.

As you might be able to see, one way to think about thread creation is that it is a bit like making a function call; however, instead of first executing the function and then returning to the caller, the system instead creates a new thread of execution for the routine that is being called, and it runs independently of the caller, perhaps before returning from the create, but perhaps much later.

As you also might be able to tell from this example, threads make life complicated: it is already hard to tell what will run when! Computers are hard enough to understand without concurrency. Unfortunately, with concurrency, it gets worse. Much worse.
main
starts running
prints “main: begin”
creates Thread 1
creates Thread 2
waits for T1
prints “A”
returns
waits for T2
returns
prints “main: end”

Table 26.1: Thread Trace (1)

main
starts running
prints “main: begin”
creates Thread 1
runs
prints “A”
returns
creates Thread 2
runs
prints “B”
returns
waits for T1
returns immediately; T1 is done
waits for T2
returns immediately; T2 is done
prints “main: end”

Table 26.2: Thread Trace (2)

main
starts running
prints “main: begin”
creates Thread 1
creates Thread 2
waits for T1
runs
prints “B”
returns
waits for T2
returns immediately; T2 is done
prints “A”
returns
waits for T2
returns immediately; T2 is done
prints “main: end”

Table 26.3: Thread Trace (3)
#include <stdio.h>
#include <pthread.h>
#include "mythreads.h"

static volatile int counter = 0;

// mythread()
// Simply adds 1 to counter repeatedly, in a loop
// No, this is not how you would add 10,000,000 to
// a counter, but it shows the problem nicely.
void *
mythread(void *arg)
{
    printf("%s: begin\n", (char *) arg);
    int i;
    for (i = 0; i < 1e7; i++) {
        counter = counter + 1;
    }
    printf("%s: done\n", (char *) arg);
    return NULL;
}

// main()
// Just launches two threads (pthread_create)
// and then waits for them (pthread_join)
int
main(int argc, char *argv[])
{
    pthread_t p1, p2;
    printf("main: begin (counter = %d)\n", counter);
    Pthread_create(&p1, NULL, mythread, "A");
    Pthread_create(&p2, NULL, mythread, "B");
    // join waits for the threads to finish
    Pthread_join(p1, NULL);
    Pthread_join(p2, NULL);
    printf("main: done with both (counter = %d)\n", counter);
    return 0;
}

Figure 26.3: Sharing Data: Oh Oh (t2)

26.2 Why It Gets Worse: Shared Data

The simple thread example we showed above was useful in showing how threads are created and how they can run in different orders depending on how the scheduler decides to run them. What it doesn’t show you, though, is how threads interact when they access shared data.
Let us imagine a simple example where two threads wish to update a global shared variable. The code we’ll study is in Figure 26.3.

Here are a few notes about the code. First, as Stevens suggests [SR05], we wrap the thread creation and join routines to simply exit on failure; for a program as simple as this one, we want to at least notice an error occurred (if it did), but not do anything very smart about it (e.g., just exit). Thus, `Pthread_create()` simply calls `pthread_create()` and makes sure the return code is 0; if it isn’t, `Pthread_create()` just prints a message and exits.

Second, instead of using two separate function bodies for the worker threads, we just use a single piece of code, and pass the thread an argument (in this case, a string) so we can have each thread print a different letter before its messages.

Finally, and most importantly, we can now look at what each worker is trying to do: add a number to the shared variable `counter`, and do so 10 million times (1e7) in a loop. Thus, the desired final result is: 20,000,000.

We now compile and run the program, to see how it behaves. Sometimes, everything works how we might expect:

```
prompt> gcc -o main main.c -Wall -pthread
prompt> ./main
main: begin (counter = 0)
A: begin
B: begin
A: done
B: done
main: done with both (counter = 20000000)
```

Unfortunately, when we run this code, even on a single processor, we don’t necessarily get the desired result. Sometimes, we get:

```
prompt> ./main
main: begin (counter = 0)
A: begin
B: begin
A: done
B: done
main: done with both (counter = 19345221)
```

Let’s try it one more time, just to see if we’ve gone crazy. After all, aren’t computers supposed to produce deterministic results, as you have been taught?! Perhaps your professors have been lying to you? (gasp)

```
prompt> ./main
main: begin (counter = 0)
A: begin
B: begin
A: done
B: done
main: done with both (counter = 19221041)
```

Not only is each run wrong, but also yields a different result! A big question remains: why does this happen?
You should always learn new tools that help you write, debug, and understand computer systems. Here, we use a neat tool called a disassembler. When you run a disassembler on an executable, it shows you what assembly instructions make up the program. For example, if we wish to understand the low-level code to update a counter (as in our example), we run `objdump` (Linux) to see the assembly code:

```
prompt> objdump -d main
```

Doing so produces a long listing of all the instructions in the program, neatly labeled (particularly if you compiled with the `-g` flag), which includes symbol information in the program. The `objdump` program is just one of many tools you should learn how to use; a debugger like `gdb`, memory profilers like `valgrind` or `purify`, and of course the compiler itself are others that you should spend time to learn more about; the better you are at using your tools, the better systems you’ll be able to build.

26.3 The Heart of the Problem: Uncontrolled Scheduling

To understand why this happens, we must understand the code sequence that the compiler generates for the update to `counter`. In this case, we wish to simply add a number (1) to `counter`. Thus, the code sequence for doing so might look something like this (in x86);

```
mov 0x8049a1c, %eax
add $0x1, %eax
mov %eax, 0x8049a1c
```

This example assumes that the variable `counter` is located at address 0x8049a1c. In this three-instruction sequence, the x86 `mov` instruction is used first to get the memory value at the address and put it into register `eax`. Then, the add is performed, adding 1 (0x1) to the contents of the `eax` register, and finally, the contents of `eax` are stored back into memory at the same address.

Let us imagine one of our two threads (Thread 1) enters this region of code, and is thus about to increment `counter` by one. It loads the value of `counter` (let’s say it’s 50 to begin with) into its register `eax`. Thus, `eax=50` for Thread 1. Then it adds one to the register; thus `eax=51`. Now, something unfortunate happens: a timer interrupt goes off; thus, the OS saves the state of the currently running thread (its PC, its registers including `eax`, etc.) to the thread’s TCB.

Now something worse happens: Thread 2 is chosen to run, and it enters this same piece of code. It also executes the first instruction, getting the value of `counter` and putting it into its `eax` (remember: each thread when running has its own private registers; the registers are virtualized by the context-switch code that saves and restores them). The value of
The Problem: Up Close and Personal

Table 26.4: The Problem: Up Close and Personal

<table>
<thead>
<tr>
<th>OS</th>
<th>Thread 1</th>
<th>Thread 2</th>
<th>(after instruction)</th>
<th>PC</th>
<th>%eax counter</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>before critical section</td>
<td></td>
<td></td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>mov 0x8049a1c, %eax</td>
<td></td>
<td></td>
<td>105</td>
<td>50</td>
</tr>
<tr>
<td></td>
<td>add $0x1, %eax</td>
<td></td>
<td></td>
<td>108</td>
<td>51</td>
</tr>
<tr>
<td>interrupt</td>
<td>save T1's state</td>
<td></td>
<td></td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>restore T2's state</td>
<td></td>
<td></td>
<td>105</td>
<td>50</td>
</tr>
<tr>
<td></td>
<td>mov 0x8049a1c, %eax</td>
<td></td>
<td></td>
<td>108</td>
<td>51</td>
</tr>
<tr>
<td></td>
<td>add $0x1, %eax</td>
<td></td>
<td></td>
<td>113</td>
<td>51</td>
</tr>
<tr>
<td></td>
<td>mov %eax, 0x8049a1c</td>
<td></td>
<td></td>
<td>108</td>
<td>51</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>113</td>
<td>51</td>
</tr>
</tbody>
</table>

**counter** is still 50 at this point, and thus Thread 2 has \( \text{eax}=50 \). Let’s then assume that Thread 2 executes the next two instructions, incrementing \( \text{eax} \) by 1 (thus \( \text{eax}=51 \)), and then saving the contents of \( \text{eax} \) into **counter** (address 0x8049a1c). Thus, the global variable **counter** now has the value 51.

Finally, another context switch occurs, and Thread 1 resumes running. Recall that it had just executed the **mov** and **add**, and is now about to perform the final **mov** instruction. Recall also that \( \text{eax}=51 \). Thus, the final **mov** instruction executes, and saves the value to memory; the counter is set to 51 again.

Put simply, what has happened is this: the code to increment **counter** has been run twice, but **counter**, which started at 50, is now only equal to 51. A “correct” version of this program should have resulted in **counter** equal to 52.

Here is a pictorial depiction of what happened and when in the example above. Assume, for this depiction, that the above code is loaded at address 100 in memory, like the following sequence (note for those of you used to nice, RISC-like instruction sets: x86 has variable-length instructions; the **mov** instructions here take up 5 bytes of memory, whereas the **add** takes only 3):

```
100 mov 0x8049a1c, %eax
105 add $0x1, %eax
108 mov %eax, 0x8049a1c
```

With these assumptions, what happens is seen in Table 26.4. Assume the counter starts at value 50, and trace through this example to make sure you understand what is going on.

What we have demonstrated here is called a **race condition**: the results depend on the timing execution of the code. With some bad luck (i.e., context switches that occur at untimely points in the execution), we get the wrong result. In fact, we may get a different result each time; thus, instead of a nice **deterministic** computation (which we are used to from computers), we call this result **indeterminate**, where it is not known what the output will be and it is indeed likely to be different across runs.
Because multiple threads executing this code can result in a race condition, we call this code a **critical section**. A critical section is a piece of code that accesses a shared variable (or more generally, a shared resource) and must not be concurrently executed by more than one thread.

What we really want for this code is what we call **mutual exclusion**. This property guarantees that if one thread is executing within the critical section, the others will be prevented from doing so.

Virtually all of these terms, by the way, were coined by Edsger Dijkstra, who was a pioneer in the field and indeed won the Turing Award because of this and other work; see his 1968 paper on “Cooperating Sequential Processes” [D68] for an amazingly clear description of the problem. We’ll be hearing more about Dijkstra in this section of the book.

### 26.4 The Wish For Atomicity

One way to solve this problem would be to have more powerful instructions that, in a single step, did exactly whatever we needed done and thus removed the possibility of an untimely interrupt. For example, what if we had a super instruction that looked like this?

\[
\text{memory-add 0x8049a1c, } 0x1
\]

Assume this instruction adds a value to a memory location, and the hardware guarantees that it executes **atomically**; when the instruction executed, it would perform the update as desired. It could not be interrupted mid-instruction, because that is precisely the guarantee we receive from the hardware: when an interrupt occurs, either the instruction has not run at all, or it has run to completion; there is no in-between state. Hardware can be a beautiful thing, no?

Atomically, in this context, means “as a unit”, which sometimes we take as “all or none.” What we’d like is to execute the three instruction sequence atomically:

\[
\begin{align*}
\text{mov} & \ 0x8049a1c, \ %eax \\
\text{add} & \ 0x1, \ %eax \\
\text{mov} & \ %eax, \ 0x8049a1c
\end{align*}
\]

As we said, if we had a single instruction to do this, we could just issue that instruction and be done. But in the general case, we won’t have such an instruction. Imagine we were building a concurrent B-tree, and wished to update it; would we really want the hardware to support an “atomic update of B-tree” instruction? Probably not, at least in a sane instruction set.

Thus, what we will instead do is ask the hardware for a few useful instructions upon which we can build a general set of what we call **synchronization primitives**. By using these hardware synchronization primitives, in combination with some help from the operating system, we will be able to build multi-threaded code that accesses critical sections in a
These four terms are so central to concurrent code that we thought it worth while to call them out explicitly. See some of Dijkstra’s early work [D65,D68] for more details.

- A **critical section** is a piece of code that accesses a *shared* resource, usually a variable or data structure.

- A **race condition** arises if multiple threads of execution enter the critical section at roughly the same time; both attempt to update the shared data structure, leading to a surprising (and perhaps undesirable) outcome.

- An **indeterminate** program consists of one or more race conditions; the output of the program varies from run to run, depending on which threads ran when. The outcome is thus not **deterministic**, something we usually expect from computer systems.

- To avoid these problems, threads should use some kind of **mutual exclusion** primitives; doing so guarantees that only a single thread ever enters a critical section, thus avoiding races, and resulting in deterministic program outputs.

synchronized and controlled manner, and thus reliably produces the correct result despite the challenging nature of concurrent execution. Pretty awesome, right?

This is the problem we will study in this section of the book. It is a wonderful and hard problem, and should make your mind hurt (a bit). If it doesn’t, then you don’t understand! Keep working until your head hurts; you then know you’re headed in the right direction. At that point, take a break; we don’t want your head hurting too much.

**THE CRUX:**

**HOW TO PROVIDE SUPPORT FOR SYNCHRONIZATION**

What support do we need from the hardware in order to build useful synchronization primitives? What support do we need from the OS? How can we build these primitives correctly and efficiently? How can programs use them to get the desired results?
26.5 One More Problem: Waiting For Another

This chapter has set up the problem of concurrency as if only one type of interaction occurs between threads, that of accessing shared variables and the need to support atomicity for critical sections. As it turns out, there is another common interaction that arises, where one thread must wait for another to complete some action before it continues. This interaction arises, for example, when a process performs a disk I/O and is put to sleep; when the I/O completes, the process needs to be roused from its slumber so it can continue.

Thus, in the coming chapters, we’ll be not only studying how to build support for synchronization primitives to support atomicity but also for mechanisms to support this type of sleeping/waking interaction that is common in multi-threaded programs. If this doesn’t make sense right now, that is OK! It will soon enough, when you read the chapter on condition variables. If it doesn’t by then, well, then it is less OK, and you should read that chapter again (and again) until it does make sense.

26.6 Summary: Why in OS Class?

Before wrapping up, one question that you might have is: why are we studying this in OS class? “History” is the one-word answer; the OS was the first concurrent program, and many techniques were created for use within the OS. Later, with multi-threaded processes, application programmers also had to consider such things.

For example, imagine the case where there are two processes running. Assume they both call write() to write to the file, and both wish to append the data to the file (i.e., add the data to the end of the file, thus increasing its length). To do so, both must allocate a new block, record in the inode of the file where this block lives, and change the size of the file to reflect the new larger size (among other things; we’ll learn more about files in the third part of the book). Because an interrupt may occur at any time, the code that updates to these shared structures (e.g., a bitmap for allocation, or the file’s inode) are critical sections; thus, OS designers, from the very beginning of the introduction of the interrupt, had to worry about how the OS updates internal structures. An untimely interrupt causes all of the problems described above. Not surprisingly, page tables, process lists, file system structures, and virtually every kernel data structure has to be carefully accessed, with the proper synchronization primitives, to work correctly.
Atomic operations are one of the most powerful underlying techniques in building computer systems, from the computer architecture, to concurrent code (what we are studying here), to file systems (which we’ll study soon enough), database management systems, and even distributed systems [L+93].

The idea behind making a series of actions atomic is simply expressed with the phrase “all or nothing”; it should either appear as if all of the actions you wish to group together occurred, or that none of them occurred, with no in-between state visible. Sometimes, the grouping of many actions into a single atomic action is called a transaction, an idea developed in great detail in the world of databases and transaction processing [GR92].

In our theme of exploring concurrency, we’ll be using synchronization primitives to turn short sequences of instructions into atomic blocks of execution, but the idea of atomicity is much bigger than that, as we will see. For example, file systems use techniques such as journaling or copy-on-write in order to atomically transition their on-disk state, critical for operating correctly in the face of system failures. If that doesn’t make sense, don’t worry – it will, in some future chapter.
References

[D65] “Solution of a problem in concurrent programming control”
E. W. Dijkstra
Communications of the ACM, 8(9):569, September 1965
Pointed to as the first paper of Dijkstra’s where he outlines the mutual exclusion problem and a solution.
The solution, however, is not widely used; advanced hardware and OS support is needed, as we will see
in the coming chapters.

[D68] “Cooperating sequential processes”
Edsger W. Dijkstra, 1968
Available: http://www.cs.utexas.edu/users/EWD/ewd01xx/EWD123.PDF
Dijkstra has an amazing number of his old papers, notes, and thoughts recorded (for posterity) on this
website at the last place he worked, the University of Texas. Much of his foundational work, however,
was done years earlier while he was at the Technische Hochshule of Eindhoven (THE), including this
famous paper on “cooperating sequential processes”, which basically outlines all of the thinking that
has to go into writing multi-threaded programs. Dijkstra discovered much of this while working on an
operating system named after his school: the “THE” operating system (said “T”, “H”, “E”, and not
like the word “the”).

Jim Gray and Andreas Reuter
Morgan Kaufmann, September 1992
This book is the bible of transaction processing, written by one of the legends of the field, Jim Gray. It is,
for this reason, also considered Jim Gray’s “brain dump”, in which he wrote down everything he knows
about how database management systems work. Sadly, Gray passed away tragically a few years back,
and many of us lost a friend and great mentor, including the co-authors of said book, who were lucky
enough to interact with Gray during their graduate school years.

[L+93] “Atomic Transactions”
Nancy Lynch, Michael Merritt, William Weihl, Alan Fekete
Morgan Kaufmann, August 1993
A nice text on some of the theory and practice of atomic transactions for distributed systems. Perhaps a
bit formal for some, but lots of good material is found herein.

[SR05] ”Advanced Programming in the UNIX Environment”
W. Richard Stevens and Stephen A. Rago
Addison-Wesley, 2005
As we’ve said many times, buy this book, and read it, in little chunks, preferably before going to bed.
This way, you will actually fall asleep more quickly; more importantly, you learn a little more about
how to become a serious UNIX programmer.
Homework

This program, *x86.py*, allows you to see how different thread interleaveings either cause or avoid race conditions. See the README for details on how the program works and its basic inputs, then answer the questions below.

Questions

1. To start, let’s examine a simple program, “loop.s”. First, just look at the program, and see if you can understand it: `cat loop.s`. Then, run it with these arguments:

   ```
   ./x86.py -p loop.s -t 1 -i 100 -R dx
   ```

   This specifies a single thread, an interrupt every 100 instructions, and tracing of register `%dx`. Can you figure out what the value of `%dx` will be during the run? Once you have, run the same above and use the `-c` flag to check your answers; note the answers, on the left, show the value of the register (or memory value) *after* the instruction on the right has run.

2. Now run the same code but with these flags:

   ```
   ./x86.py -p loop.s -t 2 -i 100 -a dx=3,dx=3 -R dx
   ```

   This specifies two threads, and initializes each `%dx` register to 3. What values will `%dx` see? Run with the `-c` flag to see the answers. Does the presence of multiple threads affect anything about your calculations? Is there a race condition in this code?

3. Now run the following:

   ```
   ./x86.py -p loop.s -t 2 -i 3 -r -a dx=3,dx=3 -R dx
   ```

   This makes the interrupt interval quite small and random; use different seeds with `-s` to see different interleavings. Does the frequency of interruption change anything about this program?

4. Next we’ll examine a different program (*looping-race-nolock.s*). This program accesses a shared variable located at memory address 2000; we’ll call this variable *x* for simplicity. Run it with a single thread and make sure you understand what it does, like this:

   ```
   ./x86.py -p looping-race-nolock.s -t 1 -M 2000
   ```

   What value is found in *x* (i.e., at memory address 2000) throughout the run? Use `-c` to check your answer.
5. Now run with multiple iterations and threads:

```
./x86.py -p looping-race-nolock.s -t 2 -a bx=3 -M 2000
```

Do you understand why the code in each thread loops three times? What will the final value of x be?

6. Now run with random interrupt intervals:

```
./x86.py -p looping-race-nolock.s -t 2 -M 2000 -i 4 -r -s 0
```

Then change the random seed, setting -s 1, then -s 2, etc. Can you tell, just by looking at the thread interleaving, what the final value of x will be? Does the exact location of the interrupt matter? Where can it safely occur? Where does an interrupt cause trouble? In other words, where is the critical section exactly?

7. Now use a fixed interrupt interval to explore the program further. Run:

```
./x86.py -p looping-race-nolock.s -a bx=1 -t 2 -M 2000 -i 1
```

See if you can guess what the final value of the shared variable x will be. What about when you change -i 2, -i 3, etc.? For which interrupt intervals does the program give the “correct” final answer?

8. Now run the same code for more loops (e.g., set -a bx=100). What interrupt intervals, set with the -i flag, lead to a “correct” outcome? Which intervals lead to surprising results?

9. We’ll examine one last program in this homework (wait-for-me.s). Run the code like this:

```
./x86.py -p wait-for-me.s -a ax=1,ax=0 -R ax -M 2000
```

This sets the %ax register to 1 for thread 0, and 0 for thread 1, and watches the value of %ax and memory location 2000 throughout the run. How should the code behave? How is the value at location 2000 being used by the threads? What will its final value be?

10. Now switch the inputs:

```
./x86.py -p wait-for-me.s -a ax=0,ax=1 -R ax -M 2000
```

How do the threads behave? What is thread 0 doing? How would changing the interrupt interval (e.g., -i 1000, or perhaps to use random intervals) change the trace outcome? Is the program efficiently using the CPU?
This chapter briefly covers the main portions of the thread API. Each part will be explained further in the subsequent chapters, as we show how to use the API. More details can be found in various books and online sources [B97, B+96, K+96]. We should note that the subsequent chapters introduce the concepts of locks and condition variables more slowly, with many examples; this chapter is thus better used as a reference.

CRUX: HOW TO CREATE AND CONTROL THREADS
What interfaces should the OS present for thread creation and control? How should these interfaces be designed to enable ease of use as well as utility?

27.1 Thread Creation

The first thing you have to be able to do to write a multi-threaded program is to create new threads, and thus some kind of thread creation interface must exist. In POSIX, it is easy:

```
#include <pthread.h>
int
pthread_create(   pthread_t *   thread,
                 const pthread_attr_t * attr,
                 void *           (*start_routine)(void*),
                 void *            arg);
```

This declaration might look a little complex (particularly if you haven’t used function pointers in C), but actually it’s not too bad. There are four arguments: thread, attr, start_routine, and arg. The first, thread, is a pointer to a structure of type pthread_t; we’ll use this structure to interact with this thread, and thus we need to pass it to pthread_create() in order to initialize it.
The second argument, `attr`, is used to specify any attributes this thread might have. Some examples include setting the stack size or perhaps information about the scheduling priority of the thread. An attribute is initialized with a separate call to `pthread_attr_init()`; see the manual page for details. However, in most cases, the defaults will be fine; in this case, we will simply pass the value `NULL` in.

The third argument is the most complex, but is really just asking: which function should this thread start running in? In C, we call this a function pointer, and this one tells us the following is expected: a function name (`start_routine`), which is passed a single argument of type `void *` (as indicated in the parentheses after `start_routine`), and which returns a value of type `void *` (i.e., a void pointer).

If this routine instead required an integer argument, instead of a void pointer, the declaration would look like this:

```c
int pthread_create(..., // first two args are the same
  void * (*start_routine)(int),
  int arg);
```

If instead the routine took a void pointer as an argument, but returned an integer, it would look like this:

```c
int pthread_create(..., // first two args are the same
  int (*start_routine)(void *),
  void * arg);
```

Finally, the fourth argument, `arg`, is exactly the argument to be passed to the function where the thread begins execution. You might ask: why do we need these void pointers? Well, the answer is quite simple: having a void pointer as an argument to the function `start_routine` allows us to pass in any type of argument; having it as a return value allows the thread to return any type of result.

Let’s look at an example in Figure 27.1. Here we just create a thread that is passed two arguments, packaged into a single type we define ourselves (`myarg_t`). The thread, once created, can simply cast its argument to the type it expects and thus unpack the arguments as desired.

And there it is! Once you create a thread, you really have another live executing entity, complete with its own call stack, running within the same address space as all the currently existing threads in the program. The fun thus begins!

### 27.2 Thread Completion

The example above shows how to create a thread. However, what happens if you want to wait for a thread to complete? You need to do something special in order to wait for completion; in particular, you must call the routine `pthread_join()`.

```c
int pthread_join(pthread_t thread, void **value_ptr);
```
### Interlude: Thread API

```c
#include <pthread.h>

typedef struct __myarg_t {
    int a;
    int b;
} myarg_t;

void *mythread(void *arg) {
    myarg_t *m = (myarg_t *) arg;
    printf("%d %d\n", m->a, m->b);
    return NULL;
}

int main(int argc, char *argv[]) {
    pthread_t p;
    int rc;
    myarg_t args;
    args.a = 10;
    args.b = 20;
    rc = pthread_create(&p, NULL, mythread, &args);
    ...
    return rc;
}
```

Figure 27.1: Creating a Thread

This routine takes only two arguments. The first is of type `pthread_t`, and is used to specify which thread to wait for. This value is exactly what you passed into the thread library during creation; if you held onto it, you can now use it to wait for the thread to stop running.

The second argument is a pointer to the return value you expect to get back. Because the routine can return anything, it is defined to return a pointer to void; because the `pthread_join()` routine changes the value of the passed in argument, you need to pass in a pointer to that value, not just the value itself.

Let’s look at another example (Figure 27.2). In the code, a single thread is again created, and passed a couple of arguments via the `myarg_t` structure. To return values, the `myret_t` type is used. Once the thread is finished running, the main thread, which has been waiting inside of the `pthread_join()` routine\(^1\), then returns, and we can access the values returned from the thread, namely whatever is in `myret_t`.

A few things to note about this example. First, often times we don’t have to do all of this painful packing and unpacking of arguments. For example, if we just create a thread with no arguments, we can pass `NULL` in as an argument when the thread is created. Similarly, we can pass `NULL` into `pthread_join()` if we don’t care about the return value.

Second, if we are just passing in a single value (e.g., an int), we don’t have to package it up as an argument. Figure 27.3 shows an example. In

---

\(^1\) Note we use wrapper functions here; specifically, we call `Malloc()`, `PthreadJoin()`, and `PthreadCreate()`, which just call their similarly-named lower-case versions and make sure the routines did not return anything unexpected.
#include <stdio.h>
#include <pthread.h>
#include <assert.h>
#include <stdlib.h>

typedef struct __myarg_t {
    int a;
    int b;
} myarg_t;

typedef struct __myret_t {
    int x;
    int y;
} myret_t;

void *mythread(void *arg) {
    myarg_t *m = (myarg_t *) arg;
    printf("%d %d\n", m->a, m->b);
    myret_t r = Malloc(sizeof(myret_t));
    r->x = 1;
    r->y = 2;
    return (void *) r;
}

int main(int argc, char *argv[]) {
    int rc;
    pthread_t p;
    myret_t *m;

    myarg_t args;
    args.a = 10;
    args.b = 20;

    Pthread_create(&p, NULL, mythread, &args);
    Pthread_join(p, (void **) &m);
    printf("returned %d %d\n", m->x, m->y);
    return 0;
}

Figure 27.2: Waiting for Thread Completion

this case, life is a bit simpler, as we don’t have to package arguments and return values inside of structures.

Third, we should note that one has to be extremely careful with how values are returned from a thread. In particular, never return a pointer which refers to something allocated on the thread’s call stack. If you do, what do you think will happen? (think about it!) Here is an example of a dangerous piece of code, modified from the example in Figure 27.2.

void *mythread(void *arg) {
    myarg_t *m = (myarg_t *) arg;
    printf("%d %d\n", m->a, m->b);
    myret_t r; // ALLOCATED ON STACK: BAD!
    r.x = 1;
    r.y = 2;
    return (void *) &r;
}
void *mythread(void *arg) {
    int m = (int) arg;
    printf("%d\n", m);
    return (void*) (arg + 1);
}

int main(int argc, char *argv[]) {
    pthread_t p;
    int rc, m;
    pthread_create(&p, NULL, mythread, (void*) 100);
    pthread_join(p, (void **) &m);
    printf("returned %d\n", m);
    return 0;
}

Figure 27.3: Simpler Argument Passing to a Thread

In this case, the variable r is allocated on the stack of mythread. However, when it returns, the value is automatically deallocated (that’s why the stack is so easy to use, after all!), and thus, passing back a pointer to a now deallocated variable will lead to all sorts of bad results. Certainly, when you print out the values you think you returned, you’ll probably (but not necessarily!) be surprised. Try it and find out for yourself2!

Finally, you might notice that the use of pthread_create() to create a thread, followed by an immediate call to pthread_join(), is a pretty strange way to create a thread. In fact, there is an easier way to accomplish this exact task; it’s called a procedure call. Clearly, we’ll usually be creating more than just one thread and waiting for it to complete, otherwise there is not much purpose to using threads at all.

We should note that not all code that is multi-threaded uses the join routine. For example, a multi-threaded web server might create a number of worker threads, and then use the main thread to accept requests and pass them to the workers, indefinitely. Such long-lived programs thus may not need to join. However, a parallel program that creates threads to execute a particular task (in parallel) will likely use join to make sure all such work completes before exiting or moving onto the next stage of computation.

27.3 Locks

Beyond thread creation and join, probably the next most useful set of functions provided by the POSIX threads library are those for providing mutual exclusion to a critical section via locks. The most basic pair of routines to use for this purpose is provided by this pair of routines:

```c
int pthread_mutex_lock(pthread_mutex_t *mutex);
int pthread_mutex_unlock(pthread_mutex_t *mutex);
```

Fortunately the compiler gcc will likely complain when you write code like this, which is yet another reason to pay attention to compiler warnings.
The routines should be easy to understand and use. When you have a region of code you realize is a critical section, and thus needs to be protected by locks in order to operate as desired. You can probably imagine what the code looks like:

```c
pthread_mutex_t lock;
pthread_mutex_lock(&lock);
x = x + 1; // or whatever your critical section is
pthread_mutex_unlock(&lock);
```

The intent of the code is as follows: if no other thread holds the lock when `pthread_mutex_lock()` is called, the thread will acquire the lock and enter the critical section. If another thread does indeed hold the lock, the thread trying to grab the lock will not return from the call until it has acquired the lock (implying that the thread holding the lock has released it via the unlock call). Of course, many threads may be stuck waiting inside the lock acquisition function at a given time; only the thread with the lock acquired, however, should call unlock.

Unfortunately, this code is broken, in two important ways. The first problem is a lack of proper initialization. All locks must be properly initialized in order to guarantee that they have the correct values to begin with and thus work as desired when lock and unlock are called.

With POSIX threads, there are two ways to initialize locks. One way to do this is to use `PTHREAD_MUTEX_INITIALIZER`, as follows:

```c
pthread_mutex_t lock = PTHREAD_MUTEX_INITIALIZER;
```

Doing so sets the lock to the default values and thus makes the lock usable. The dynamic way to do it (i.e., at run time) is to make a call to `pthread_mutex_init()`, as follows:

```c
int rc = pthread_mutex_init(&lock, NULL);
assert(rc == 0); // always check success!
```

The first argument to this routine is the address of the lock itself, whereas the second is an optional set of attributes. Read more about the attributes yourself; passing `NULL` in simply uses the defaults. Either way works, but we usually use the dynamic (latter) method. Note that a corresponding call to `pthread_cond_destroy()` should also be made, when you are done with the lock; see the manual page for all of details.

The second problem with the code above is that it fails to check errors code when calling lock and unlock. Just like virtually any library routine you call in a UNIX system, these routines can also fail! If your code doesn’t properly check error codes, the failure will happen silently, which in this case could allow multiple threads into a critical section. Minimally, use wrappers, which assert that the routine succeeded (e.g., as in Figure 27.4); more sophisticated (non-toy) programs, which can’t simply exit when something goes wrong, should check for failure and do something appropriate when the lock or unlock does not succeed.
INTERLUDE: THREAD API

// Use this to keep your code clean but check for failures
// Only use if exiting program is OK upon failure
void Pthread_mutex_lock(pthread_mutex_t*mutex) {
    int rc = pthread_mutex_lock(mutex);
    assert(rc == 0);
}

Figure 27.4: An Example Wrapper

The lock and unlock routines are not the only routines that pthreads
has to interact with locks. In particular, here are two more routines which
may be of interest:

int pthread_mutex_trylock(pthread_mutex_t*mutex);
int pthread_mutex_timedlock(pthread_mutex_t *mutex,
    struct timespec *abs_timeout);

These two calls are used in lock acquisition. The trylock version re-
turns failure if the lock is already held; the timedlock version of acquir-
ing a lock returns after a timeout or after acquiring the lock, whichever
happens first. Thus, the timedlock with a timeout of zero degenerates
to the trylock case. Both of these versions should generally be avoided;
however, there are a few cases where avoiding getting stuck (perhaps in-
definitely) in a lock acquisition routine can be useful, as we’ll see in future
chapters (e.g., when we study deadlock).

27.4 Condition Variables

The other major component of any threads library, and certainly the
case with POSIX threads, is the presence of a condition variable. Con-
dition variables are useful when some kind of signaling must take place
between threads, if one thread is waiting for another to do something be-
fore it can continue. Two primary routines are used by programs wishing
to interact in this way:

int pthread_cond_wait(pthread_cond_t*cond, pthread_mutex_t *mutex);
int pthread_cond_signal(pthread_cond_t *cond);

To use a condition variable, one has to in addition have a lock that is
associated with this condition. When calling either of the above routines,
this lock should be held.

The first routine, pthread_cond_wait(), puts the calling thread to
sleep, and thus waits for some other thread to signal it, usually when
something in the program has changed that the now-sleeping thread might
care about. For example, a typical usage looks like this:

pthread_mutex_t lock = PTHREAD_MUTEX_INITIALIZER;
pthread_cond_t init = PTHREAD_COND_INITIALIZER;

Pthread_mutex_lock(&lock);
while (initialized == 0)
    Pthread_cond_wait(&init, &lock);
Pthread_mutex_unlock(&lock);
In this code, after initialization of the relevant lock and condition, a thread checks to see if the variable `initialized` has yet been set to something other than zero. If not, the thread simply calls the wait routine in order to sleep until some other thread wakes it.

The code to wake a thread, which would run in some other thread, looks like this:

```c
Pthread_mutex_lock(&lock);
initialized = 1;
Pthread_cond_signal(&init);
Pthread_mutex_unlock(&lock);
```

A few things to note about this code sequence. First, when signaling (as well as when modifying the global variable `initialized`), we always make sure to have the lock held. This ensures that we don’t accidentally introduce a race condition into our code.

Second, you might notice that the wait call takes a lock as its second parameter, whereas the signal call only takes a condition. The reason for this difference is that the wait call, in addition to putting the calling thread to sleep, releases the lock when putting said caller to sleep. Imagine if it did not: how could the other thread acquire the lock and signal it to wake up? However, before returning after being woken, the `pthread_cond_wait()` re-acquires the lock, thus ensuring that any time the waiting thread is running between the lock acquire at the beginning of the wait sequence, and the lock release at the end, it holds the lock.

One last oddity: the waiting thread re-checks the condition in a while loop, instead of a simple if statement. We’ll discuss this issue in detail when we study condition variables in a future chapter, but in general, using a while loop is the simple and safe thing to do. Although it rechecks the condition (perhaps adding a little overhead), there are some `pthread` implementations that could spuriously wake up a waiting thread; in such a case, without rechecking, the waiting thread will continue thinking that the condition has changed even though it has not. It is safer thus to view waking up as a hint that something might have changed, rather than an absolute fact.

Note that sometimes it is tempting to use a simple flag to signal between two threads, instead of a condition variable and associated lock. For example, we could rewrite the waiting code above to look more like this in the waiting code:

```c
while (initialized == 0)
    ; // spin
```

The associated signaling code would look like this:

```c
initialized = 1;
```

---

3Note that one could use `pthread_cond_init()` (and corresponding the `pthread_cond_destroy()` call) instead of the static initializer `PTHREAD_COND_INITIALIZER`. Sound like more work? It is.
Don’t ever do this, for the following reasons. First, it performs poorly in many cases (spinning for a long time just wastes CPU cycles). Second, it is error prone. As recent research shows [X+10], it is surprisingly easy to make mistakes when using flags (as above) to synchronize between threads; roughly half the uses of these ad hoc synchronizations were buggy! Don’t be lazy; use condition variables even when you think you can get away without doing so.

27.5 Compiling and Running

All of the code examples in this chapter are relatively easy to get up and running. To compile them, you must include the header pthread.h in your code. On the link line, you must also explicitly link with the pthreads library, by adding the -pthread flag.

For example, to compile a simple multi-threaded program, all you have to do is the following:

```
prompt> gcc -o main main.c -Wall -pthread
```

As long as main.c includes the pthreads header, you have now successfully compiled a concurrent program. Whether it works or not, as usual, is a different matter entirely.

27.6 Summary

We have introduced the basics of the pthread library, including thread creation, building mutual exclusion via locks, and signaling and waiting via condition variables. You don’t need much else to write robust and efficient multi-threaded code, except patience and a great deal of care!

We now end the chapter with a set of tips that might be useful to you when you write multi-threaded code (see the aside on the following page for details). There are other aspects of the API that are interesting; if you want more information, type man -k pthread on a Linux system to see over one hundred APIs that make up the entire interface. However, the basics discussed herein should enable you to build sophisticated (and hopefully, correct and performant) multi-threaded programs. The hard part with threads is not the APIs, but rather the tricky logic of how you build concurrent programs. Read on to learn more.
ASIDE: THREAD API GUIDELINES

There are a number of small but important things to remember when you use the POSIX thread library (or really, any thread library) to build a multi-threaded program. They are:

- **Keep it simple.** Above all else, any code to lock or signal between threads should be as simple as possible. Tricky thread interactions lead to bugs.

- **Minimize thread interactions.** Try to keep the number of ways in which threads interact to a minimum. Each interaction should be carefully thought out and constructed with tried and true approaches (many of which we will learn about in the coming chapters).

- **Initialize locks and condition variables.** Failure to do so will lead to code that sometimes works and sometimes fails in very strange ways.

- **Check your return codes.** Of course, in any C and UNIX programming you do, you should be checking each and every return code, and it’s true here as well. Failure to do so will lead to bizarre and hard to understand behavior, making you likely to (a) scream, (b) pull some of your hair out, or (c) both.

- **Be careful with how you pass arguments to, and return values from, threads.** In particular, any time you are passing a reference to a variable allocated on the stack, you are probably doing something wrong.

- **Each thread has its own stack.** As related to the point above, please remember that each thread has its own stack. Thus, if you have a locally-allocated variable inside of some function a thread is executing, it is essentially private to that thread; no other thread can (easily) access it. To share data between threads, the values must be in the heap or otherwise some locale that is globally accessible.

- **Always use condition variables to signal between threads.** While it is often tempting to use a simple flag, don’t do it.

- **Use the manual pages.** On Linux, in particular, the pthread man pages are highly informative and discuss much of the nuances presented here, often in even more detail. Read them carefully!
References

[B97] “Programming with POSIX Threads”
David R. Butenhof
Addison-Wesley, May 1997
Another one of these books on threads.

Dick Buttlar, Jacqueline Farrell, Bradford Nichols
O’Reilly, September 1996
A reasonable book from the excellent, practical publishing house O’Reilly. Our bookshelves certainly contain a great deal of books from this company, including some excellent offerings on Perl, Python, and Javascript (particularly Crockford’s “Javascript: The Good Parts”.)

[K+96] “Programming With Threads”
Steve Kleiman, Devang Shah, Bart Smaalders
Prentice Hall, January 1996
Probably one of the better books in this space. Get it at your local library.

[X+10] “Ad Hoc Synchronization Considered Harmful”
Weiwei Xiong, Soyeon Park, Jiaqi Zhang, Yuanyuan Zhou, Zhiqiang Ma
OSDI 2010, Vancouver, Canada
This paper shows how seemingly simple synchronization code can lead to a surprising number of bugs. Use condition variables and do the signaling correctly!
Locks

From the introduction to concurrency, we saw one of the fundamental problems in concurrent programming: we would like to execute a series of instructions atomically, but due to the presence of interrupts on a single processor (or multiple threads executing on multiple processors concurrently), we couldn’t. In this chapter, we thus attack this problem directly, with the introduction of something referred to as a lock. Programmers annotate source code with locks, putting them around critical sections, and thus ensure that any such critical section executes as if it were a single atomic instruction.

28.1 Locks: The Basic Idea

As an example, assume our critical section looks like this, the canonical update of a shared variable:

```c
balance = balance + 1;
```

Of course, other critical sections are possible, such as adding an element to a linked list or other more complex updates to shared structures, but we’ll just keep to this simple example for now. To use a lock, we add some code around the critical section like this:

```c
1 lock_t mutex; // some globally-allocated lock ‘mutex’
2 ...
3 lock(&mutex);
4 balance = balance + 1;
5 unlock(&mutex);
```

A lock is just a variable, and thus to use one, you must declare a lock variable of some kind (such as mutex above). This lock variable (or just “lock” for short) holds the state of the lock at any instant in time. It is either available (or unlocked or free) and thus no thread holds the lock, or acquired (or locked or held), and thus exactly one thread holds the lock and presumably is in a critical section. We could store other information
in the data type as well, such as which thread holds the lock, or a queue for ordering lock acquisition, but information like that is hidden from the user of the lock.

The semantics of the lock() and unlock() routines are simple. Calling the routine lock() tries to acquire the lock; if no other thread holds the lock (i.e., it is free), the thread will acquire the lock and enter the critical section; this thread is sometimes said to be the owner of the lock. If another thread then calls lock() on that same lock variable (mutex in this example), it will not return while the lock is held by another thread; in this way, other threads are prevented from entering the critical section while the first thread that holds the lock is in there.

Once the owner of the lock calls unlock(), the lock is now available (free) again. If no other threads are waiting for the lock (i.e., no other thread has called lock() and is stuck therein), the state of the lock is simply changed to free. If there are waiting threads (stuck in lock()), one of them will (eventually) notice (or be informed of) this change of the lock’s state, acquire the lock, and enter the critical section.

Locks provide some minimal amount of control over scheduling to programmers. In general, we view threads as entities created by the programmer but scheduled by the OS, in any fashion that the OS chooses. Locks yield some of that control back to the programmer; by putting a lock around a section of code, the programmer can guarantee that no more than a single thread can ever be active within that code. Thus locks help transform the chaos that is traditional OS scheduling into a more controlled activity.

28.2 Pthread Locks

The name that the POSIX library uses for a lock is a mutex, as it is used to provide mutual exclusion between threads, i.e., if one thread is in the critical section, it excludes the others from entering until it has completed the section. Thus, when you see the following POSIX threads code, you should understand that it is doing the same thing as above (we again use our wrappers that check for errors upon lock and unlock):

```c
1 pthread_mutex_t lock = PTHREAD_MUTEX_INITIALIZER;
2
3 Pthread_mutex_lock(&lock); // wrapper for pthread_mutex_lock()
4 balance = balance + 1;
5 Pthread_mutex_unlock(&lock);
```

You might also notice here that the POSIX version passes a variable to lock and unlock, as we may be using different locks to protect different variables. Doing so can increase concurrency: instead of one big lock that is used any time any critical section is accessed (a coarse-grained locking strategy), one will often protect different data and data structures with different locks, thus allowing more threads to be in locked code at once (a more fine-grained approach).
28.3 Building A Lock

By now, you should have some understanding of how a lock works, from the perspective of a programmer. But how should we build a lock? What hardware support is needed? What OS support? It is this set of questions we address in the rest of this chapter.

The Crux: How To Build A Lock

How can we build an efficient lock? Efficient locks provided mutual exclusion at low cost, and also might attain a few other properties we discuss below. What hardware support is needed? What OS support?

To build a working lock, we will need some help from our old friend, the hardware, as well as our good pal, the OS. Over the years, a number of different hardware primitives have been added to the instruction sets of various computer architectures; while we won’t study how these instructions are implemented (that, after all, is the topic of a computer architecture class), we will study how to use them in order to build a mutual exclusion primitive like a lock. We will also study how the OS gets involved to complete the picture and enable us to build a sophisticated locking library.

28.4 Evaluating Locks

Before building any locks, we should first understand what our goals are, and thus we ask how to evaluate the efficacy of a particular lock implementation. To evaluate whether a lock works (and works well), we should first establish some basic criteria. The first is whether the lock does its basic task, which is to provide mutual exclusion. Basically, does the lock work, preventing multiple threads from entering a critical section?

The second is fairness. Does each thread contending for the lock get a fair shot at acquiring it once it is free? Another way to look at this is by examining the more extreme case: does any thread contending for the lock starve while doing so, thus never obtaining it?

The final criterion is performance, specifically the time overheads added by using the lock. There are a few different cases that are worth considering here. One is the case of no contention; when a single thread is running and grabs and releases the lock, what is the overhead of doing so? Another is the case where multiple threads are contending for the lock on a single CPU; in this case, are there performance concerns? Finally, how does the lock perform when there are multiple CPUs involved, and threads on each contending for the lock? By comparing these different scenarios, we can better understand the performance impact of using various locking techniques, as described below.
28.5 Controlling Interrupts

One of the earliest solutions used to provide mutual exclusion was to disable interrupts for critical sections; this solution was invented for single-processor systems. The code would look like this:

```c
void lock() {
    DisableInterrupts();
}

void unlock() {
    EnableInterrupts();
}
```

Assume we are running on such a single-processor system. By turning off interrupts (using some kind of special hardware instruction) before entering a critical section, we ensure that the code inside the critical section will not be interrupted, and thus will execute as if it were atomic. When we are finished, we re-enable interrupts (again, via a hardware instruction) and thus the program proceeds as usual.

The main positive of this approach is its simplicity. You certainly don’t have to scratch your head too hard to figure out why this works. Without interruption, a thread can be sure that the code it executes will execute and that no other thread will interfere with it.

The negatives, unfortunately, are many. First, this approach requires us to allow any calling thread to perform a privileged operation (turning interrupts on and off), and thus trust that this facility is not abused. As you already know, any time we are required to trust an arbitrary program, we are probably in trouble. Here, the trouble manifests in numerous ways: a greedy program could call `lock()` at the beginning of its execution and thus monopolize the processor; worse, an errant or malicious program could call `lock()` and go into an endless loop. In this latter case, the OS never regains control of the system, and there is only one recourse: restart the system. Using interrupt disabling as a general-purpose synchronization solution requires too much trust in applications.

Second, the approach does not work on multiprocessors. If multiple threads are running on different CPUs, and each try to enter the same critical section, it does not matter whether interrupts are disabled; threads will be able to run on other processors, and thus could enter the critical section. As multiprocessors are now commonplace, our general solution will have to do better than this.

Third, and probably least important, this approach can be inefficient. Compared to normal instruction execution, code that masks or unmasks interrupts tends to be executed slowly by modern CPUs.

For these reasons, turning off interrupts is only used in limited contexts as a mutual-exclusion primitive. For example, in some cases an operating system itself will sometimes use interrupt masking to guarantee atomicity when accessing its own data structures, or at least to prevent certain messy interrupt handling situations from arising. This usage makes sense, as the trust issue disappears inside the OS, which always trusts itself to perform privileged operations anyhow.
ASIDE: DEKKER’S AND PETERSON’S ALGORITHMS

In the 1960’s, Dijkstra posed the concurrency problem to his friends, and one of them, a mathematician named Theodorus Jozef Dekker, came up with a solution [D68]. Unlike the solutions we discuss here, which use special hardware instructions and even OS support, Dekker’s approach uses just loads and stores (assuming they are atomic with respect to each other, which was true on early hardware).

Dekker’s approach was later refined by Peterson [P81] (and thus “Peterson’s algorithm”), shown here. Once again, just loads and stores are used, and the idea is to ensure that two threads never enter a critical section at the same time. Here is Peterson’s algorithm (for two threads); see if you can understand it.

```c
int flag[2];
int turn;

void init() {
    flag[0] = flag[1] = 0;  // 1->thread wants to grab lock
    turn = 0;  // whose turn? (thread 0 or 1?)
}

void lock() {
    flag[1-self] = 1;  // self: thread ID of caller
    turn = 1 - self;  // make it other thread’s turn
    while ((flag[1-self] == 1) && (turn == 1 - self))
        ;  // spin-wait
}

void unlock() {
    flag[1-self] = 0;  // simply undo your intent
}
```

For some reason, developing locks that work without special hardware support became all the rage for a while, giving theory-types a lot of problems to work on. Of course, this all became quite useless when people realized it is much easier to assume a little hardware support (and indeed that support had been around from the very earliest days of multiprocessing). Further, algorithms like the ones above don’t work on modern hardware (due to relaxed memory consistency models), thus making them even less useful than they were before. Yet more research relegated to the dustbin of history...

28.6 Test And Set (Atomic Exchange)

Because disabling interrupts does not work on multiple processors, system designers started to invent hardware support for locking. The earliest multiprocessor systems, such as the Burroughs B5000 in the early 1960’s [M82], had such support; today all systems provide this type of support, even for single CPU systems.
typedef struct __lock_t { int flag; } lock_t;

void init(lock_t *mutex) {
    // 0 -> lock is available, 1 -> held
    mutex->flag = 0;
}

void lock(lock_t *mutex) {
    while (mutex->flag == 1) // TEST the flag
        ; // spin-wait (do nothing)
    mutex->flag = 1; // now SET it!
}

void unlock(lock_t *mutex) {
    mutex->flag = 0;
}

Figure 28.1: First Attempt: A Simple Flag

The simplest bit of hardware support to understand is what is known as a **test-and-set instruction**, also known as **atomic exchange**. To understand how test-and-set works, let’s first try to build a simple lock without it. In this failed attempt, we use a simple flag variable to denote whether the lock is held or not.

In this first attempt (Figure 28.1), the idea is quite simple: use a simple variable to indicate whether some thread has possession of a lock. The first thread that enters the critical section will call `lock()`, which **tests** whether the flag is equal to 1 (in this case, it is not), and then **sets** the flag to 1 to indicate that the thread now **holds** the lock. When finished with the critical section, the thread calls `unlock()` and clears the flag, thus indicating that the lock is no longer held.

If another thread happens to call `lock()` while that first thread is in the critical section, it will simply **spin-wait** in the while loop for that thread to call `unlock()` and clear the flag. Once that first thread does so, the waiting thread will fall out of the while loop, set the flag to 1 for itself, and proceed into the critical section.

Unfortunately, the code has two problems: one of correctness, and another of performance. The correctness problem is simple to see once you get used to thinking about concurrent programming. Imagine the code interleaving in Table 28.1 (assume `flag=0` to begin).

<table>
<thead>
<tr>
<th>Thread 1</th>
<th>Thread 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>call lock()</td>
<td>call lock()</td>
</tr>
<tr>
<td>while (flag == 1)</td>
<td>while (flag == 1)</td>
</tr>
<tr>
<td>interrupt: switch to Thread 2</td>
<td>flag = 1;</td>
</tr>
<tr>
<td></td>
<td>interrupt: switch to Thread 1</td>
</tr>
</tbody>
</table>
| flag = 1;              | / / set flag to 1 (too!) | Table 28.1: Trace: No Mutual Exclusion
As you can see from this interleaving, with timely (untimely?) interrupts, we can easily produce a case where both threads set their flags to 1 and both threads are thus able to enter the critical section. This is bad! We have obviously failed to provide the most basic requirement: providing mutual exclusion.

The performance problem, which we will address more later on, is the fact that the way a thread waits to acquire a lock that is already held: it endlessly checks the value of flag, a technique known as spin-waiting. Spin-waiting wastes time waiting for another thread to release a lock. The waste is exceptionally high on a uniprocessor, where the thread that the waiter is waiting for cannot even run (at least, until a context switch occurs)! Thus, as we move forward and develop more sophisticated solutions, we should also consider ways to avoid this kind of waste.

28.7 Building A Working Spin Lock

While the idea behind the example above is a good one, it is not possible to implement without some support from the hardware. Fortunately, some systems provide an instruction to support the creation of simple locks based on this concept. This more powerful instruction has different names – on SPARC, it is the load/store unsigned byte instruction (ldstub), whereas on x86, it is the atomic exchange instruction (xchg) – but basically does the same thing across platforms, and is usually generally referred to as test-and-set. We define what the test-and-set instruction does with the following C code snippet:

```c
int TestAndSet(int *ptr, int new) {
    int old = *ptr; // fetch old value at ptr
    *ptr = new;     // store 'new' into ptr
    return old;    // return the old value
}
```

What the test-and-set instruction does is as follows. It returns the old value pointed to by the `ptr`, and simultaneously updates said value to `new`. The key, of course, is that this sequence of operations is performed atomically. The reason it is called “test and set” is that it enables you to “test” the old value (which is what is returned) while simultaneously
typedef struct __lock_t {
    int flag;
} lock_t;

void init(lock_t *lock) {
    // 0 indicates that lock is available, 1 that it is held
    lock->flag = 0;
}

void lock(lock_t *lock) {
    while (TestAndSet(&lock->flag, 1) == 1) // spin-wait (do nothing)
}

void unlock(lock_t *lock) {
    lock->flag = 0;
}

Figure 28.2: A Simple Spin Lock Using Test-and-set

“setting” the memory location to a new value; as it turns out, this slightly more powerful instruction is enough to build a simple spin lock, as we now examine in Figure 28.2.

Let’s make sure we understand why this works. Imagine first the case where a thread calls lock() and no other thread currently holds the lock; thus, flag should be 0. When the thread then calls TestAndSet(flag, 1), the routine will return the old value of flag, which is 0; thus, the calling thread, which is testing the value of flag, will not get caught spinning in the while loop and will acquire the lock. The thread will also atomically set the value to 1, thus indicating that the lock is now held. When the thread is finished with its critical section, it calls unlock() to set the flag back to zero.

The second case we can imagine arises when one thread already has the lock held (i.e., flag is 1). In this case, this thread will call lock() and then call TestAndSet(flag, 1) as well. This time, TestAndSet() will return the old value at flag, which is 1 (because the lock is held), while simultaneously setting it to 1 again. As long as the lock is held by another thread, TestAndSet() will repeatedly return 1, and thus this thread will spin and spin until the lock is finally released. When the flag is finally set to 0 by some other thread, this thread will call TestAndSet() again, which will now return 0 while atomically setting the value to 1 and thus acquire the lock and enter the critical section.

By making both the test (of the old lock value) and set (of the new value) a single atomic operation, we ensure that only one thread acquires the lock. And that’s how to build a working mutual exclusion primitive!

You may also now understand why this type of lock is usually referred to as a spin lock. It is the simplest type of lock to build, and simply spins, using CPU cycles, until the lock becomes available. To work correctly on a single processor, it requires a preemptive scheduler (i.e., one that will interrupt a thread via a timer, in order to run a different thread, from time to time). Without preemption, spin locks don’t make much sense on a single CPU, as a thread spinning on a CPU will never relinquish it.
28.8 Evaluating Spin Locks

Given our basic spin lock, we can now evaluate how effective it is along our previously described axes. The most important aspect of a lock is correctness: does it provide mutual exclusion? The answer here is obviously yes: the spin lock only allows a single thread to enter the critical section at a time. Thus, we have a correct lock.

The next axis is fairness. How fair is a spin lock to a waiting thread? Can you guarantee that a waiting thread will ever enter the critical section? The answer here, unfortunately, is bad news: spin locks don’t provide any fairness guarantees. Indeed, a thread spinning may spin forever, under contention. Spin locks are not fair and may lead to starvation.

The final axis is performance. What are the costs of using a spin lock? To analyze this more carefully, we suggest thinking about a few different cases. In the first, imagine threads competing for the lock on a single processor; in the second, consider the threads as spread out across many processors.

For spin locks, in the single CPU case, performance overheads can be quite painful; imagine the case where the thread holding the lock is pre-empted within a critical section. The scheduler might then run every other thread (imagine there are \( N - 1 \) others), each of which tries to acquire the lock. In this case, each of those threads will spin for the duration of a time slice before giving up the CPU, a waste of CPU cycles.

However, on multiple CPUs, spin locks work reasonably well (if the number of threads roughly equals the number of CPUs). The thinking goes as follows: imagine Thread A on CPU 1 and Thread B on CPU 2, both contending for a lock. If Thread A (CPU 1) grabs the lock, and then Thread B tries to, B will spin (on CPU 2). However, presumably the critical section is short, and thus soon the lock becomes available, and is acquired by Thread B. Spinning to wait for a lock held on another processor doesn’t waste many cycles in this case, and thus can be quite effective.

28.9 Compare-And-Swap

Another hardware primitive that some systems provide is known as the compare-and-swap instruction (as it is called on SPARC, for example), or compare-and-exchange (as it called on x86). The C pseudocode for this single instruction is found in Figure 28.3.

The basic idea is for compare-and-swap to test whether the value at the

```c
int CompareAndSwap(int *ptr, int expected, int new) {
    int actual = *ptr;
    if (actual == expected) {
        *ptr = new;
    }
    return actual;
}
```

Figure 28.3: Compare-and-swap
address specified by \texttt{ptr} is equal to \texttt{expected}; if so, update the memory location pointed to by \texttt{ptr} with the new value. If not, do nothing. In either case, return the actual value at that memory location, thus allowing the code calling compare-and-swap to know whether it succeeded or not.

With the compare-and-swap instruction, we can build a lock in a manner quite similar to that with test-and-set. For example, we could just replace the \texttt{lock()} routine above with the following:

```c
void lock(lock_t *lock) {
    while (CompareAndSwap(&lock->flag, 0, 1) == 1) // spin
}
```

The rest of the code is the same as the test-and-set example above. This code works quite similarly; it simply checks if the flag is 0 and if so, atomically swaps in a 1 thus acquiring the lock. Threads that try to acquire the lock while it is held will get stuck spinning until the lock is finally released.

If you want to see how to really make a C-callable x86-version of compare-and-swap, this code sequence might be useful (from [S05]):

```c
char CompareAndSwap(int *ptr, int old, int new) {
    unsigned char ret;

    __asm__ __volatile__ ("lock
        cmpxchgl %2,%1
        sete %0
": "=q" (ret), "=m" (*ptr)
: "r" (new), "m" (*ptr), "a" (old)
: "memory");

    return ret;
}
```

Finally, as you may have sensed, compare-and-swap is a more powerful instruction than test-and-set. We will make some use of this power in the future when we briefly delve into \textit{wait-free synchronization} [H91]. However, if we just build a simple spin lock with it, its behavior is identical to the spin lock we analyzed above.

### 28.10 Load-Linked and Store-Conditional

Some platforms provide a pair of instructions that work in concert to help build critical sections. On the MIPS architecture [H93], for example, the \texttt{load-linked} and \texttt{store-conditional} instructions can be used in tandem to build locks and other concurrent structures. The C pseudocode for these instructions is as found in Figure 28.4. Alpha, PowerPC, and ARM provide similar instructions [W09].
int LoadLinked(int *ptr) {
    return *ptr;
}

int StoreConditional(int *ptr, int value) {
    if (no one has updated *ptr since the LoadLinked to this address) {
        *ptr = value;
        return 1; // success!
    } else {
        return 0; // failed to update
    }
}

Figure 28.4: Load-linked And Store-conditional

void lock(lock_t *lock) {
    while (1) {
        while (LoadLinked(&lock->flag) == 1)
            ; // spin until it’s zero
        if (StoreConditional(&lock->flag, 1) == 1)
            return; // if set-it-to-1 was a success: all done
        // otherwise: try it all over again
    }
}

void unlock(lock_t *lock) {
    lock->flag = 0;
}

Figure 28.5: Using LL/SC To Build A Lock

The load-linked operates much like a typical load instruction, and simply fetches a value from memory and places it in a register. The key difference comes with the store-conditional, which only succeeds (and updates the value stored at the address just load-linked from) if no intermittent store to the address has taken place. In the case of success, the store-conditional returns 1 and updates the value at ptr to value; if it fails, the value at ptr is not updated and 0 is returned.

As a challenge to yourself, try thinking about how to build a lock using load-linked and store-conditional. Then, when you are finished, look at the code below which provides one simple solution. Do it! The solution is in Figure 28.5.

The lock() code is the only interesting piece. First, a thread spins waiting for the flag to be set to 0 (and thus indicate the lock is not held). Once so, the thread tries to acquire the lock via the store-conditional; if it succeeds, the thread has atomically changed the flag’s value to 1 and thus can proceed into the critical section.

Note how failure of the store-conditional might arise. One thread calls lock() and executes the load-linked, returning 0 as the lock is not held. Before it can attempt the store-conditional, it is interrupted and another thread enters the lock code, also executing the load-linked instruction, and also getting a 0 and continuing. At this point, two threads have each executed the load-linked and each are about to attempt the store-conditional. The key feature of these instructions is that only one of these threads will succeed in updating the flag to 1 and thus acquire the lock;
Programmers tend to brag about how much code they wrote to do something. Doing so is fundamentally broken. What one should brag about, rather, is how little code one wrote to accomplish a given task. Short, concise code is always preferred; it is likely easier to understand and has fewer bugs. As Hugh Lauer said, when discussing the construction of the Pilot operating system: “If the same people had twice as much time, they could produce as good of a system in half the code.” [L81] We’ll call this Lauer’s Law, and it is well worth remembering. So next time you’re bragging about how much code you wrote to finish the assignment, think again, or better yet, go back, rewrite, and make the code as clear and concise as possible.

the second thread to attempt the store-conditional will fail (because the other thread updated the value of flag between its load-linked and store-conditional) and thus have to try to acquire the lock again.

In class a few years ago, undergraduate student David Capels suggested a more concise form of the above, for those of you who enjoy short-circuiting boolean conditionals. See if you can figure out why it is equivalent. It certainly is shorter!

```
1  void lock(lock_t *lock) {
2      while (LoadLinked(&lock->flag)||!StoreConditional(&lock->flag, 1))
3           ; // spin
4  }
```

28.11 Fetch-And-Add

One final hardware primitive is the fetch-and-add instruction, which atomically increments a value while returning the old value at a particular address. The C pseudocode for the fetch-and-add instruction looks like this:

```
1  int FetchAndAdd(int *ptr) {
2      int old = *ptr;
3      *ptr = old + 1;
4      return old;
5  }
```

In this example, we’ll use fetch-and-add to build a more interesting ticket lock, as introduced by Mellor-Crummey and Scott [MS91]. The lock and unlock code looks like what you see in Figure 28.6.

Instead of a single value, this solution uses a ticket and turn variable in combination to build a lock. The basic operation is pretty simple: when a thread wishes to acquire a lock, it first does an atomic fetch-and-add on the ticket value; that value is now considered this thread’s “turn” (myturn). The globally shared lock->turn is then used to determine which thread’s turn it is; when (myturn == turn) for a given thread,
typedef struct __lock_t {
    int ticket;
    int turn;
} lock_t;

void lock_init(lock_t *lock) {
    lock->ticket = 0;
    lock->turn = 0;
}

void lock(lock_t *lock) {
    int myturn = FetchAndAdd(&lock->ticket);
    while (lock->turn != myturn)
        ; // spin
}

void unlock(lock_t *lock) {
    FetchAndAdd(&lock->turn);
}

Figure 28.6: Ticket Locks

it is that thread’s turn to enter the critical section. Unlock is accomplished simply by incrementing the turn such that the next waiting thread (if there is one) can now enter the critical section.

Note one important difference with this solution versus our previous attempts: it ensures progress for all threads. Once a thread is assigned its ticket value, it will be scheduled at some point in the future (once those in front of it have passed through the critical section and released the lock). In our previous attempts, no such guarantee existed; a thread spinning on test-and-set (for example) could spin forever even as other threads acquire and release the lock.

28.12 Summary: So Much Spinning

Our simple hardware-based locks are simple (only a few lines of code) and they work (you could even prove that if you’d like to, by writing some code), which are two excellent properties of any system or code. However, in some cases, these solutions can be quite inefficient. Imagine you are running two threads on a single processor. Now imagine that one thread (thread 0) is in a critical section and thus has a lock held, and unfortunately gets interrupted. The second thread (thread 1) now tries to acquire the lock, but finds that it is held. Thus, it begins to spin. And spin. Then it spins some more. And finally, a timer interrupt goes off, thread 0 is run again, which releases the lock, and finally (the next time it runs, say), thread 1 won’t have to spin so much and will be able to acquire the lock. Thus, any time a thread gets caught spinning in a situation like this, it wastes an entire time slice doing nothing but checking a value that isn’t going to change! The problem gets worse with $N$ threads contending for a lock; $N - 1$ time slices may be wasted in a similar manner, simply spinning and waiting for a single thread to release the lock. And thus, our next problem:
THE CRUX: HOW TO AVOID SPINNING
How can we develop a lock that doesn’t needlessly waste time spinning on the CPU?

Hardware support alone cannot solve the problem. We’ll need OS support too! Let’s now figure out just how that might work.

28.13 A Simple Approach: Just Yield, Baby

Hardware support got us pretty far: working locks, and even (as with the case of the ticket lock) fairness in lock acquisition. However, we still have a problem: what to do when a context switch occurs in a critical section, and threads start to spin endlessly, waiting for the interrupt (lock-holding) thread to be run again?

Our first try is a simple and friendly approach: when you are going to spin, instead give up the CPU to another thread. Or, as Al Davis might say, “just yield, baby!” [D91]. Figure 28.7 presents the approach.

In this approach, we assume an operating system primitive `yield()` which a thread can call when it wants to give up the CPU and let another thread run. Because a thread can be in one of three states (running, ready, or blocked), you can think of this as an OS system call that moves the caller from the running state to the ready state, and thus promotes another thread to running.

Think about the example with two threads on one CPU; in this case, our yield-based approach works quite well. If a thread happens to call `lock()` and find a lock held, it will simply yield the CPU, and thus the other thread will run and finish its critical section. In this simple case, the yielding approach works well.

Let us now consider the case where there are many threads (say 100) contending for a lock repeatedly. In this case, if one thread acquires the lock and is preempted before releasing it, the other 99 will each call

```c
void init() {
    flag = 0;
}

void lock() {
    while (TestAndSet(&flag, 1) == 1)
        yield(); // give up the CPU
}

void unlock() {
    flag = 0;
}
```

Figure 28.7: Lock With Test-and-set And Yield
lock(), find the lock held, and yield the CPU. Assuming some kind of round-robin scheduler, each of the 99 will execute this run-and-yield pattern before the thread holding the lock gets to run again. While better than our spinning approach (which would waste 99 time slices spinning), this approach is still costly; the cost of a context switch can be substantial, and there is thus plenty of waste.

Worse, we have not tackled the starvation problem at all. A thread may get caught in an endless yield loop while other threads repeatedly enter and exit the critical section. We clearly will need an approach that addresses this problem directly.

28.14 Using Queues: Sleeping Instead Of Spinning

The real problem with our previous approaches is that they leave too much to chance. The scheduler determines which thread runs next; if the scheduler makes a bad choice, a thread runs that must either spin waiting for the lock (our first approach), or yield the CPU immediately (our second approach). Either way, there is potential for waste and no prevention of starvation.

Thus, we must explicitly exert some control over who gets to acquire the lock next after the current holder releases it. To do this, we will need a little more OS support, as well as a queue to keep track of which threads are waiting to enter the lock.

For simplicity, we will use the support provided by Solaris, in terms of two calls: park() to put a calling thread to sleep, and unpark(threadID) to wake a particular thread as designated by threadID. These two routines can be used in tandem to build a lock that puts a caller to sleep if it tries to acquire a held lock and wakes it when the lock is free. Let’s look at the code in Figure 28.8 to understand one possible use of such primitives.

We do a couple of interesting things in this example. First, we combine the old test-and-set idea with an explicit queue of lock waiters to make a more efficient lock. Second, we use a queue to help control who gets the lock next and thus avoid starvation.

You might notice how the guard is used, basically as a spin-lock around the flag and queue manipulations the lock is using. This approach thus doesn’t avoid spin-waiting entirely; a thread might be interrupted while acquiring or releasing the lock, and thus cause other threads to spin-wait for this one to run again. However, the time spent spinning is quite limited (just a few instructions inside the lock and unlock code, instead of the user-defined critical section), and thus this approach may be reasonable.

Second, you might notice that in lock(), when a thread can not acquire the lock (it is already held), we are careful to add ourselves to a queue (by calling the getpid() call to get the thread ID of the current thread), set guard to 0, and yield the CPU. A question for the reader: What would happen if the release of the guard lock came after the park(), and not before? Hint: something bad.
typedef struct __lock_t {
   int flag;
   int guard;
   queue_t *q;
} lock_t;

void lock_init(lock_t *m) {
   m->flag = 0;
   m->guard = 0;
   queue_init(m->q);
}

void lock(lock_t *m) {
   while (TestAndSet(&m->guard, 1) == 1) ; //acquire guard lock by spinning
   if (m->flag == 0) {
      m->flag = 1; // lock is acquired
      m->guard = 0;
   } else {
      queue_add(m->q, gettid());
      m->guard = 0;
      park();
   }
}

void unlock(lock_t *m) {
   while (TestAndSet(&m->guard, 1) == 1) ; //acquire guard lock by spinning
   if (queue_empty(m->q))
      m->flag = 0; // let go of lock; no one wants it
   else
      unpark(queue_remove(m->q)); // hold lock (for next thread!)
   m->guard = 0;
}

Figure 28.8: Lock With Queues, Test-and-set, Yield, And Wakeup

You might also notice the interesting fact that the flag does not get set back to 0 when another thread gets woken up. Why is this? Well, it is not an error, but rather a necessity! When a thread is woken up, it will be as if it is returning from park(); however, it does not hold the guard at that point in the code and thus cannot even try to set the flag to 1. Thus, we just pass the lock directly from the thread releasing the lock to the next thread acquiring it; flag is not set to 0 in-between.

Finally, you might notice the perceived race condition in the solution, just before the call to park(). With just the wrong timing, a thread will be about to park, assuming that it should sleep until the lock is no longer held. A switch at that time to another thread (say, a thread holding the lock) could lead to trouble, for example, if that thread then released the lock. The subsequent park by the first thread would then sleep forever (potentially). This problem is sometimes called the wakeup/waiting race; to avoid it, we need to do some extra work.

Solaris solves this problem by adding a third system call: setpark(). By calling this routine, a thread can indicate it is about to park. If it then happens to be interrupted and another thread calls unpark before park is
actually called, the subsequent park returns immediately instead of sleeping. The code modification, inside of `lock()`, is quite small:

```c
queue_add(m->q, gettid());
setpark(); // new code
m->guard = 0;
```

A different solution could pass the guard into the kernel. In that case, the kernel could take precautions to atomically release the lock and dequeue the running thread.

### 28.15 Different OS, Different Support

We have thus far seen one type of support that an OS can provide in order to build a more efficient lock in a thread library. Other OS’s provide similar support; the details vary.

For example, Linux provides something called a `futex` which is similar to the Solaris interface but provides a bit more in-kernel functionality. Specifically, each futex has associated with it a specific physical memory location; associated with each such memory location is an in-kernel queue. Callers can use futex calls (described below) to sleep and wake as need be.

Specifically, two calls are available. The call to `futex_wait(address, expected)` puts the calling thread to sleep, assuming the value at `address` is equal to `expected`. If it is not equal, the call returns immediately. The call to the routine `futex_wake(address)` wakes one thread that is waiting on the queue. The usage of these in Linux is as found in 28.9.

This code snippet from `lowlevellock.h` in the nptl library (part of the gnu libc library) [L09] is pretty interesting. Basically, it uses a single integer to track both whether the lock is held or not (the high bit of the integer) and the number of waiters on the lock (all the other bits). Thus, if the lock is negative, it is held (because the high bit is set and that bit determines the sign of the integer). The code is also interesting because it shows how to optimize for the common case where there is no contention: with only one thread acquiring and releasing a lock, very little work is done (the atomic bit test-and-set to lock and an atomic add to release the lock). See if you can puzzle through the rest of this “real-world” lock to see how it works.

### 28.16 Two-Phase Locks

One final note: the Linux approach has the flavor of an old approach that has been used on and off for years, going at least as far back to Dahm Locks in the early 1960’s [M82], and is now referred to as a **two-phase lock**. A two-phase lock realizes that spinning can be useful, particularly if the lock is about to be released. So in the first phase, the lock spins for a while, hoping that it can acquire the lock.
void mutex_lock (int *mutex) {
    int v;
    /* Bit 31 was clear, we got the mutex (this is the fastpath) */
    if (atomic_bit_test_set (mutex, 31) == 0)
        return;
    atomic_increment (mutex);
    while (1) {
        if (atomic_bit_test_set (mutex, 31) == 0) {
            atomic_decrement (mutex);
            return;
        }
        /* We have to wait now. First make sure the futex value
         * we are monitoring is truly negative (i.e. locked). */
        v = *mutex;
        if (v >= 0)
            continue;
        futex_wait (mutex, v);
    }
}

void mutex_unlock (int *mutex) {
    /* Adding 0x80000000 to the counter results in 0 if and only if
     * there are not other interested threads */
    if (atomic_add_zero (mutex, 0x80000000))
        return;
    /* There are other threads waiting for this mutex,
     * wake one of them up. */
    futex_wake (mutex);
}

Figure 28.9: Linux-based Futex Locks

However, if the lock is not acquired during the first spin phase, a second phase is entered, where the caller is put to sleep, and only woken up when the lock becomes free later. The Linux lock above is a form of such a lock, but it only spins once; a generalization of this could spin in a loop for a fixed amount of time before using futex support to sleep.

Two-phase locks are yet another instance of a hybrid approach, where combining two good ideas may indeed yield a better one. Of course, whether it does depends strongly on many things, including the hardware environment, number of threads, and other workload details. As always, making a single general-purpose lock, good for all possible use cases, is quite a challenge.

28.17 Summary

The above approach shows how real locks are built these days: some hardware support (in the form of a more powerful instruction) plus some operating system support (e.g., in the form of park() and unpark() primitives on Solaris, or futex on Linux). Of course, the details differ, and the exact code to perform such locking is usually highly tuned. Check out the Solaris or Linux open source code bases if you want to see more details; they are a fascinating read [L09, S09].
References

[D91] “Just Win, Baby: Al Davis and His Raiders”
Glenn Dickey, Harcourt 1991
There is even an undoubtedly bad book about Al Davis and his famous “just win” quote. Or, we suppose, the book is more about Al Davis and the Raiders, and maybe not just the quote. Read the book to find out?

[D68] “Cooperating sequential processes”
Edsger W. Dijkstra, 1968
Available: http://www.cs.utexas.edu/users/EWD/EWD123.PDF
One of the early seminal papers in the area. Discusses how Dijkstra posed the original concurrency problem, and Dekker’s solution.

Joe Heinrich, Prentice-Hall, June 1993

[H91] “Wait-free Synchronization”
Maurice Herlihy
ACM Transactions on Programming Languages and Systems (TOPLAS)
Volume 13, Issue 1, January 1991
A landmark paper introducing a different approach to building concurrent data structures. However, because of the complexity involved, many of these ideas have been slow to gain acceptance in deployed systems.

Hugh Lauer
SOSP ’81
A must-read retrospective about the development of the Pilot OS, an early PC operating system. Fun and full of insights.

[L09] “glibc 2.9 (include Linux pthreads implementation)”
In particular, take a look at the nptl subdirectory where you will find most of the pthread support in Linux today.

20 Years Later and Still Ahead of the Times?”
Alastair J.W. Mayer, 1982
www.ajwm.net/amayer/papers/B5000.html
From the paper: “One particularly useful instruction is the RDLK (read-lock). It is an indivisible operation which reads from and writes into a memory location.” RDLK is thus an early test-and-set primitive, if not the earliest. Some credit here goes to an engineer named Dave Dahm, who apparently invented a number of these things for the Burroughs systems, including a form of spin locks (called “Buzz Locks” as well as a two-phase lock eponymously called “Dahm Locks.”)

[MS91] “Algorithms for Scalable Synchronization on Shared-Memory Multiprocessors”
John M. Mellor-Crummey and M. L. Scott
ACM TOCS, February 1991
An excellent survey on different locking algorithms. However, no OS support is used, just fancy hardware instructions.

[P81] “Myths About the Mutual Exclusion Problem”
G.L. Peterson
Information Processing Letters. 12(3) 1981, 115–116
Peterson’s algorithm introduced here.
“Guide to porting from Solaris to Linux on x86”
Ajay Sood, April 29, 2005

“OpenSolaris Thread Library”
usr/src/lib/libc/port/threads/synch.c
This is also pretty interesting to look at, though who knows what will happen to it now that Oracle owns
Sun. Thanks to Mike Swift for the pointer to the code.

“Load-Link, Store-Conditional”
Wikipedia entry on said topic, as of October 22, 2009
http://en.wikipedia.org/wiki/Load-Link/Store-Conditional
Can you believe we referenced wikipedia? Pretty shabby. But, we found the information there first,
and it felt wrong not to cite it. Further, they even listed the instructions for the different architectures:
ldl/l/stl_c and ldg/l/stq_c (Alpha), lwarx/stwcx (PowerPC), ll/sc (MIPS), and
ldrex/strex (ARM version 6 and above).

David L. Weaver and Tom Germond, September 2000
SPARC International, San Jose, California
Also see: http://developers.sun.com/solaris/articles/atomic_sparc/ for some
more details on Sparc atomic operations.
Lock-based Concurrent Data Structures

Before moving beyond locks, we’ll first describe how to use locks in some common data structures. Adding locks to a data structure to make it usable by threads makes the structure thread safe. Of course, exactly how such locks are added determines both the correctness and performance of the data structure. And thus, our challenge:

**CRUX: HOW TO ADD LOCKS TO DATA STRUCTURES**

When given a particular data structure, how should we add locks to it, in order to make it work correctly? Further, how do we add locks such that the data structure yields high performance, enabling many threads to access the structure at once, i.e., concurrently?

Of course, we will be hard pressed to cover all data structures or all methods for adding concurrency, as this is a topic that has been studied for years, with (literally) thousands of research papers published about it. Thus, we hope to provide a sufficient introduction to the type of thinking required, and refer you to some good sources of material for further inquiry on your own. We found Moir and Shavit’s survey to be a great source of information [MS04].

29.1 Concurrent Counters

One of the simplest data structures is a counter. It is a structure that is commonly used and has a simple interface. We define a simple non-concurrent counter in Figure 29.1.

**Simple But Not Scalable**

As you can see, the non-synchronized counter is a trivial data structure, requiring a tiny amount of code to implement. We now have our next challenge: how can we make this code thread safe? Figure 29.2 shows how we do so.
typedef struct __counter_t {
    int value;
} counter_t;

void init(counter_t *c) {
    c->value = 0;
}

void increment(counter_t *c) {
    c->value++;
}

void decrement(counter_t *c) {
    c->value--;
}

int get(counter_t *c) {
    return c->value;
}

Figure 29.1: A Counter Without Locks

typedef struct __counter_t {
    int value;
    pthread_lock_t lock;
} counter_t;

void init(counter_t *c) {
    c->value = 0;
    Pthread_mutex_init(&c->lock, NULL);
}

void increment(counter_t *c) {
    Pthread_mutex_lock(&c->lock);
    c->value++;
    Pthread_mutex_unlock(&c->lock);
}

void decrement(counter_t *c) {
    Pthread_mutex_lock(&c->lock);
    c->value--;
    Pthread_mutex_unlock(&c->lock);
}

int get(counter_t *c) {
    Pthread_mutex_lock(&c->lock);
    int rc = c->value;
    Pthread_mutex_unlock(&c->lock);
    return rc;
}

Figure 29.2: A Counter With Locks

This concurrent counter is simple and works correctly. In fact, it follows a design pattern common to the simplest and most basic concurrent data structures: it simply adds a single lock, which is acquired when calling a routine that manipulates the data structure, and is released when returning from the call. In this manner, it is similar to a data structure built with monitors [BH73], where locks are acquired and released automatically as you call and return from object methods.
At this point, you have a working concurrent data structure. The problem you might have is performance. If your data structure is too slow, you’ll have to do more than just add a single lock; such optimizations, if needed, are thus the topic of the rest of the chapter. Note that if the data structure is not too slow, you are done! No need to do something fancy if something simple will work.

To understand the performance costs of the simple approach, we run a benchmark in which each thread updates a single shared counter a fixed number of times; we then vary the number of threads. Figure 29.3 shows the total time taken, with one to four threads active; each thread updates the counter one million times. This experiment was run upon an iMac with four Intel 2.7 GHz i5 CPUs; with more CPUs active, we hope to get more total work done per unit time.

From the top line in the figure (labeled precise), you can see that the performance of the synchronized counter scales poorly. Whereas a single thread can complete the million counter updates in a tiny amount of time (roughly 0.03 seconds), having two threads each update the counter one million times concurrently leads to a massive slowdown (taking over 5 seconds!). It only gets worse with more threads.

Ideally, you’d like to see the threads complete just as quickly on multiple processors as the single thread does on one. Achieving this end is called perfect scaling; even though more work is done, it is done in parallel, and hence the time taken to complete the task is not increased.

Scalable Counting

Amazingly, researchers have studied how to build more scalable counters for years [MS04]. Even more amazing is the fact that scalable counters matter, as recent work in operating system performance analysis has shown [B+10]; without scalable counting, some workloads running on
Linux suffer from serious scalability problems on multicore machines.

Though many techniques have been developed to attack this problem, we’ll now describe one particular approach. The idea, introduced in recent research [B+10], is known as a sloppy counter.

The sloppy counter works by representing a single logical counter via numerous local physical counters, one per CPU core, as well as a single global counter. Specifically, on a machine with four CPUs, there are four local counters and one global one. In addition to these counters, there are also locks: one for each local counter, and one for the global counter.

The basic idea of sloppy counting is as follows. When a thread running on a given core wishes to increment the counter, it increments its local counter; access to this local counter is synchronized via the corresponding local lock. Because each CPU has its own local counter, threads across CPUs can update local counters without contention, and thus counter updates are scalable.

However, to keep the global counter up to date (in case a thread wishes to read its value), the local values are periodically transferred to the global counter, by acquiring the global lock and incrementing it by the local counter’s value; the local counter is then reset to zero.

How often this local-to-global transfer occurs is determined by a threshold, which we call $S$ here (for sloppiness). The smaller $S$ is, the more the counter behaves like the non-scalable counter above; the bigger $S$ is, the more scalable the counter, but the further off the global value might be from the actual count. One could simply acquire all the local locks and the global lock (in a specified order, to avoid deadlock) to get an exact value, but that is not scalable.

To make this clear, let’s look at an example (Table 29.1). In this example, the threshold $S$ is set to 5, and there are threads on each of four CPUs updating their local counters $L_1$ ... $L_4$. The global counter value ($G$) is also shown in the trace, with time increasing downward. At each time step, a local counter may be incremented; if the local value reaches the threshold $S$, the local value is transferred to the global counter and the local counter is reset.

The lower line in Figure 29.3 (labeled sloppy) shows the performance of sloppy counters with a threshold $S$ of 1024. Performance is excellent; the time taken to update the counter four million times on four processors is hardly higher than the time taken to update it one million times on one processor.

**Table 29.1: Tracing the Sloppy Counters**

<table>
<thead>
<tr>
<th>Time</th>
<th>$L_1$</th>
<th>$L_2$</th>
<th>$L_3$</th>
<th>$L_4$</th>
<th>$G$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>1</td>
<td>0</td>
<td>2</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>3</td>
<td>2</td>
<td>0</td>
<td>3</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>4</td>
<td>3</td>
<td>0</td>
<td>3</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>5</td>
<td>4</td>
<td>1</td>
<td>3</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>6</td>
<td>5 → 0</td>
<td>1</td>
<td>3</td>
<td>4</td>
<td>5 (from $L_1$)</td>
</tr>
<tr>
<td>7</td>
<td>0</td>
<td>2</td>
<td>4</td>
<td>5 → 0</td>
<td>10 (from $L_4$)</td>
</tr>
</tbody>
</table>
typedef struct __counter_t {
    int global; // global count
    pthread_mutex_t glock; // global lock
    int local[NUMCPUS]; // local count (per cpu)
    pthread_mutex_t llock[NUMCPUS]; // ... and locks
    int threshold; // update frequency
} counter_t;

// init: record threshold, init locks, init values
// of all local counts and global count
void init(counter_t *c, int threshold) {
    c->threshold = threshold;
    c->global = 0;
    pthread_mutex_init(&c->glock, NULL);
    int i;
    for (i = 0; i < NUMCPUS; i++) {
        c->local[i] = 0;
        pthread_mutex_init(&c->llock[i], NULL);
    }
}

// update: usually, just grab local lock and update local amount
// once local count has risen by ‘threshold’, grab global
// lock and transfer local values to it
void update(counter_t *c, int threadID, int amt) {
    pthread_mutex_lock(&c->llock[threadID]);
    c->local[threadID] += amt; // assumes amt > 0
    if (c->local[threadID] >= c->threshold) { // transfer to global
        pthread_mutex_lock(&c->glock);
        c->global += c->local[threadID];
        pthread_mutex_unlock(&c->glock);
        c->local[threadID] = 0;
    }
    pthread_mutex_unlock(&c->llock[threadID]);
}

// get: just return global amount (which may not be perfect)
int get(counter_t *c) {
    pthread_mutex_lock(&c->glock);
    int val = c->global;
    pthread_mutex_unlock(&c->glock);
    return val; // only approximate!
}

Figure 29.4: Sloppy Counter Implementation

Figure 29.5 shows the importance of the threshold value $S$, with four threads each incrementing the counter 1 million times on four CPUs. If $S$ is low, performance is poor (but the global count is always quite accurate); if $S$ is high, performance is excellent, but the global count lags (by the number of CPUs multiplied by $S$). This accuracy/performance trade-off is what sloppy counters enables.

A rough version of such a sloppy counter is found in Figure 29.4. Read it, or better yet, run it yourself in some experiments to better understand how it works.
29.2 Concurrent Linked Lists

We next examine a more complicated structure, the linked list. Let’s start with a basic approach once again. For simplicity, we’ll omit some of the obvious routines that such a list would have and just focus on concurrent insert; we’ll leave it to the reader to think about lookup, delete, and so forth. Figure 29.6 shows the code for this rudimentary data structure.

As you can see in the code, the code simply acquires a lock in the insert routine upon entry, and releases it upon exit. One small tricky issue arises if malloc() happens to fail (a rare case); in this case, the code must also release the lock before failing the insert.

This kind of exceptional control flow has been shown to be quite error prone; a recent study of Linux kernel patches found that a huge fraction of bugs (nearly 40%) are found on such rarely-taken code paths (indeed, this observation sparked some of our own research, in which we removed all memory-failing paths from a Linux file system, resulting in a more robust system [S+11]).

Thus, a challenge: can we rewrite the insert and lookup routines to remain correct under concurrent insert but avoid the case where the failure path also requires us to add the call to unlock?

The answer, in this case, is yes. Specifically, we can rearrange the code a bit so that the lock and release only surround the actual critical section in the insert code, and that a common exit path is used in the lookup code. The former works because part of the lookup actually need not be locked; assuming that malloc() itself is thread-safe, each thread can call into it without worry of race conditions or other concurrency bugs. Only when updating the shared list does a lock need to be held. See Figure 29.7 for the details of these modifications.
LOCK-BASED CONCURRENT DATA STRUCTURES

1 // basic node structure
2 typedef struct __node_t {
3     int key;
4     struct __node_t *next;
5 } node_t;
6
7 // basic list structure (one used per list)
8 typedef struct __list_t {
9     node_t *head;
10     pthread_mutex_t lock;
11 } list_t;
12
13 void List_Init(list_t *L) {
14     L->head = NULL;
15     pthread_mutex_init(&L->lock, NULL);
16 }
17
18 int List_Insert(list_t *L, int key) {
19     pthread_mutex_lock(&L->lock);
20     node_t *new = malloc(sizeof(node_t));
21     if (new == NULL) {
22         perror("malloc");
23         pthread_mutex_unlock(&L->lock);
24         return -1; // fail
25     }
26     new->key = key;
27     new->next = L->head;
28     L->head = new;
29     pthread_mutex_unlock(&L->lock);
30     return 0; // success
31 }
32
33 int List_Lookup(list_t *L, int key) {
34     pthread_mutex_lock(&L->lock);
35     node_t *curr = L->head;
36     while (curr) {
37         if (curr->key == key) {
38             pthread_mutex_unlock(&L->lock);
39             return 0; // success
40         }
41         curr = curr->next;
42     }
43     pthread_mutex_unlock(&L->lock);
44     return -1; // failure
45 }

Figure 29.6: Concurrent Linked List

As for the lookup routine, it is a simple code transformation to jump out of the main search loop to a single return path. Doing so again reduces the number of lock acquire/release points in the code, and thus decreases the chances of accidentally introducing bugs (such as forgetting to unlock before returning) into the code.

Scaling Linked Lists

Though we again have a basic concurrent linked list, once again we are in a situation where it does not scale particularly well. One technique that researchers have explored to enable more concurrency within a list is
Figure 29.7: Concurrent Linked List: Rewritten

something called hand-over-hand locking (a.k.a. lock coupling) [MS04].

The idea is pretty simple. Instead of having a single lock for the entire list, you instead add a lock per node of the list. When traversing the list, the code first grabs the next node’s lock and then releases the current node’s lock (which inspires the name hand-over-hand).

Conceptually, a hand-over-hand linked list makes some sense; it enables a high degree of concurrency in list operations. However, in practice, it is hard to make such a structure faster than the simple single lock approach, as the overheads of acquiring and releasing locks for each node of a list traversal is prohibitive. Even with very large lists, and a large number of threads, the concurrency enabled by allowing multiple ongoing traversals is unlikely to be faster than simply grabbing a single lock, performing an operation, and releasing it. Perhaps some kind of hybrid (where you grab a new lock every so many nodes) would be worth investigating.
Tip: More Concurrency Isn’t Necessarily Faster

If the scheme you design adds a lot of overhead (for example, by acquiring and releasing locks frequently, instead of once), the fact that it is more concurrent may not be important. Simple schemes tend to work well, especially if they use costly routines rarely. Adding more locks and complexity can be your downfall. All of that said, there is one way to really know: build both alternatives (simple but less concurrent, and complex but more concurrent) and measure how they do. In the end, you can’t cheat on performance; your idea is either faster, or it isn’t.

Tip: Be Wary Of Locks and Control Flow

A general design tip, which is useful in concurrent code as well as elsewhere, is to be wary of control flow changes that lead to function returns, exits, or other similar error conditions that halt the execution of a function. Because many functions will begin by acquiring a lock, allocating some memory, or doing other similar stateful operations, when errors arise, the code has to undo all of the state before returning, which is error-prone. Thus, it is best to structure code to minimize this pattern.

29.3 Concurrent Queues

As you know by now, there is always a standard method to make a concurrent data structure: add a big lock. For a queue, we’ll skip that approach, assuming you can figure it out.

Instead, we’ll take a look at a slightly more concurrent queue designed by Michael and Scott [MS98]. The data structures and code used for this queue are found in Figure 29.8 on the following page.

If you study this code carefully, you’ll notice that there are two locks, one for the head of the queue, and one for the tail. The goal of these two locks is to enable concurrency of enqueue and dequeue operations. In the common case, the enqueue routine will only access the tail lock, and dequeue only the head lock.

One trick used by the Michael and Scott is to add a dummy node (allocated in the queue initialization code); this dummy enables the separation of head and tail operations. Study the code, or better yet, type it in, run it, and measure it, to understand how it works deeply.

Queues are commonly used in multi-threaded applications. However, the type of queue used here (with just locks) often does not completely meet the needs of such programs. A more fully developed bounded queue, that enables a thread to wait if the queue is either empty or overly full, is the subject of our intense study in the next chapter on condition variables. Watch for it!
typedef struct __node_t {
  int value;
  struct __node_t *next;
} node_t;

typedef struct __queue_t {
  node_t *head;
  node_t *tail;
  pthread_mutex_t headLock;
  pthread_mutex_t tailLock;
} queue_t;

void Queue_Init(queue_t *q) {
  node_t *tmp = malloc(sizeof(node_t));
  tmp->next = NULL;
  q->head = q->tail = tmp;
  pthread_mutex_init(&q->headLock, NULL);
  pthread_mutex_init(&q->tailLock, NULL);
}

void Queue_Enqueue(queue_t *q, int value) {
  node_t *tmp = malloc(sizeof(node_t));
  assert(tmp != NULL);
  tmp->value = value;
  tmp->next = NULL;
  pthread_mutex_lock(&q->tailLock);
  q->tail->next = tmp;
  q->tail = tmp;
  pthread_mutex_unlock(&q->tailLock);
}

int Queue_Dequeue(queue_t *q, int *value) {
  pthread_mutex_lock(&q->headLock);
  node_t *tmp = q->head;
  node_t *newHead = tmp->next;
  if (newHead == NULL) {
    pthread_mutex_unlock(&q->headLock);
    return -1; // queue was empty
  }
  *value = newHead->value;
  q->head = newHead;
  pthread_mutex_unlock(&q->headLock);
  free(tmp);
  return 0;
}

Figure 29.8: Michael and Scott Concurrent Queue

29.4 Concurrent Hash Table

We end our discussion with a simple and widely applicable concurrent data structure, the hash table. We’ll focus on a simple hash table that does not resize; a little more work is required to handle resizing, which we leave as an exercise for the reader (sorry!).

This concurrent hash table is straightforward, is built using the concurrent lists we developed earlier, and works incredibly well. The reason
#define BUCKETS (101)

typedef struct __hash_t {
    list_t lists[BUCKETS];
} hash_t;

void Hash_Init(hash_t *H) {
    int i;
    for (i = 0; i < BUCKETS; i++) {
        List_Init(&H->lists[i]);
    }
}

int Hash_Insert(hash_t *H, int key) {
    int bucket = key % BUCKETS;
    return List_Insert(&H->lists[bucket], key);
}

int Hash_Lookup(hash_t *H, int key) {
    int bucket = key % BUCKETS;
    return List_Lookup(&H->lists[bucket], key);
}

Figure 29.9: A Concurrent Hash Table

for its good performance is that instead of having a single lock for the entire structure, it uses a lock per hash bucket (each of which is represented by a list). Doing so enables many concurrent operations to take place.

Figure 29.10 shows the performance of the hash table under concurrent updates (from 10,000 to 50,000 concurrent updates from each of four threads, on the same iMac with four CPUs). Also shown, for the sake of comparison, is the performance of a linked list (with a single lock). As you can see from the graph, this simple concurrent hash table scales magnificently; the linked list, in contrast, does not.
When building a concurrent data structure, start with the most basic approach, which is to add a single big lock to provide synchronized access. By doing so, you are likely to build a correct lock; if you then find that it suffers from performance problems, you can refine it, thus only making it fast if need be. As Knuth famously stated, “Premature optimization is the root of all evil.”

Many operating systems added a single lock when transitioning to multi-processors, including Sun OS and Linux. In the latter, it even had a name, the big kernel lock (BKL), and was the source of performance problems for many years until it was finally removed in 2011. In SunOS (which was a BSD variant), the notion of removing the single lock protecting the kernel was so painful that the Sun engineers decided on a different route: building the entirely new Solaris operating system, which was multi-threaded from day one. Read the Linux and Solaris kernel books for more information [BC05, MM00].

29.5 Summary

We have introduced a sampling of concurrent data structures, from counters, to lists and queues, and finally to the ubiquitous and heavily-used hash table. We have learned a few important lessons along the way: to be careful with acquisition and release of locks around control flow changes; that enabling more concurrency does not necessarily increase performance; that performance problems should only be remedied once they exist. This last point, of avoiding premature optimization, is central to any performance-minded developer; there is no value in making something faster if doing so will not improve the overall performance of the application.

Of course, we have just scratched the surface of high performance structures. See Moir and Shavit’s excellent survey for more information, as well as links to other sources [MS04]. In particular, you might be interested in other structures (such as B-trees); for this knowledge, a database class is your best bet. You also might be interested in techniques that don’t use traditional locks at all; such non-blocking data structures are something we’ll get a taste of in the chapter on common concurrency bugs, but frankly this topic is an entire area of knowledge requiring more study than is possible in this humble book. Find out more on your own if you are interested (as always!).
References

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Robert Morris, Nickolai Zeldovich
OSDI ’10, Vancouver, Canada, October 2010
A great study of how Linux performs on multicore machines, as well as some simple solutions.

[BH73] “Operating System Principles”
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One of the first books on operating systems; certainly ahead of its time. Introduced monitors as a
concurrency primitive.

[BC05] “Understanding the Linux Kernel (Third Edition)”
Daniel P. Bovet and Marco Cesati
O’Reilly Media, November 2005
The classic book on the Linux kernel. You should read it.

Lanyue Lu, Andrea C. Arpaci-Dusseau, Remzi H. Arpaci-Dusseau, Shan Lu
FAST ’13, San Jose, CA, February 2013
Our paper that studies every patch to Linux file systems over nearly a decade. Lots of fun findings in
there; read it to see! The work was painful to do though; the poor graduate student, Lanyue Lu, had to
look through every single patch by hand in order to understand what they did.

[MS98] “Nonblocking Algorithms and Preemption-safe Locking on Multiprogrammed Shared-
memory Multiprocessors”
M. Michael and M. Scott
Professor Scott and his students have been at the forefront of concurrent algorithms and data structures
for many years; check out his web page, numerous papers, or books to find out more.

[MS04] “Concurrent Data Structures”
Mark Moir and Nir Shavit
In Handbook of Data Structures and Applications
(Editors D. Metha and S. Sahni)
Chapman and Hall/CRC Press, 2004
A short but relatively comprehensive reference on concurrent data structures. Though it is missing
some of the latest works in the area (due to its age), it remains an incredibly useful reference.

[MM00] “Solaris Internals: Core Kernel Architecture”
Jim Mauro and Richard McDougall
Prentice Hall, October 2000
The Solaris book. You should also read this, if you want to learn in great detail about something other
than Linux.

[S+11] “Making the Common Case the Only Case with Anticipatory Memory Allocation”
Swaminathan Sundararaman, Yupu Zhang, Srim Subramanian,
Andrea C. Arpaci-Dusseau, Remzi H. Arpaci-Dusseau
FAST ’11, San Jose, CA, February 2011
Our work on removing possibly-failing calls to malloc from kernel code paths. The idea is to allocate all
potentially needed memory before doing any of the work, thus avoiding failure deep down in the storage
stack.
Thus far we have developed the notion of a lock and seen how one can be properly built with the right combination of hardware and OS support. Unfortunately, locks are not the only primitives that are needed to build concurrent programs.

In particular, there are many cases where a thread wishes to check whether a condition is true before continuing its execution. For example, a parent thread might wish to check whether a child thread has completed before continuing (this is often called a `join()`); how should such a wait be implemented? Let’s look at Figure 30.1.

```c
void *child(void *arg) {
    printf("child\n");
    // XXX how to indicate we are done?
    return NULL;
}

int main(int argc, char *argv[]) {
    printf("parent: begin\n");
    pthread_t c;
    pthread_create(&c, NULL, child, NULL); // create child
    // XXX how to wait for child?
    printf("parent: end\n");
    return 0;
}
```

**Figure 30.1: A Parent Waiting For Its Child**

What we would like to see here is the following output:

```
parent: begin
child
parent: end
```

We could try using a shared variable, as you see in Figure 30.2. This solution will generally work, but it is hugely inefficient as the parent spins and wastes CPU time. What we would like here instead is some way to put the parent to sleep until the condition we are waiting for (e.g., the child is done executing) comes true.
volatile int done = 0;

void *child(void *arg) {
    printf("child\n");
    done = 1;
    return NULL;
}

int main(int argc, char *argv[]) {
    printf("parent: begin\n");
    pthread_t c;
    Pthread_create(&c, NULL, child, NULL); // create child
    while (done == 0)
        ; // spin
    printf("parent: end\n");
    return 0;
}

Figure 30.2: Parent Waiting For Child: Spin-based Approach

THE CRUX: HOW TO WAIT FOR A CONDITION

In multi-threaded programs, it is often useful for a thread to wait for some condition to become true before proceeding. The simple approach, of just spinning until the condition becomes true, is grossly inefficient and wastes CPU cycles, and in some cases, can be incorrect. Thus, how should a thread wait for a condition?

30.1 Definition and Routines

To wait for a condition to become true, a thread can make use of what is known as a condition variable. A condition variable is an explicit queue that threads can put themselves on when some state of execution (i.e., some condition) is not as desired (by waiting on the condition); some other thread, when it changes said state, can then wake one (or more) of those waiting threads and thus allow them to continue (by signaling on the condition). The idea goes back to Dijkstra’s use of “private semaphores” [D68]; a similar idea was later named a “condition variable” by Hoare in his work on monitors [H74].

To declare such a condition variable, one simply writes something like this: `pthread_cond_t c;`, which declares `c` as a condition variable (note: proper initialization is also required). A condition variable has two operations associated with it: `wait()` and `signal()`. The `wait()` call is executed when a thread wishes to put itself to sleep; the `signal()` call is executed when a thread has changed something in the program and thus wants to wake a sleeping thread waiting on this condition. Specifically, the POSIX calls look like this:

```c
pthread_cond_wait(pthread_cond_t *c, pthread_mutex_t *m);
pthread_cond_signal(pthread_cond_t *c);
```
Figure 30.3: Parent Waiting For Child: Use A Condition Variable

We will often refer to these as wait() and signal() for simplicity. One thing you might notice about the wait() call is that it also takes a mutex as a parameter; it assumes that this mutex is locked when wait() is called. The responsibility of wait() is to release the lock and put the calling thread to sleep (atomically); when the thread wakes up (after some other thread has signaled it), it must re-acquire the lock before returning to the caller. This complexity stems from the desire to prevent certain race conditions from occurring when a thread is trying to put itself to sleep. Let’s take a look at the solution to the join problem (Figure 30.3) to understand this better.

There are two cases to consider. In the first, the parent creates the child thread but continues running itself (assume we have only a single processor) and thus immediately calls into thr_join() to wait for the child thread to complete. In this case, it will acquire the lock, check if the child is done (it is not), and put itself to sleep by calling wait() (hence releasing the lock). The child will eventually run, print the message “child”, and call thr_exit() to wake the parent thread; this code just grabs the lock, sets the state variable done, and signals the parent thus waking it. Finally, the parent will run (returning from wait() with the lock held), unlock the lock, and print the final message “parent: end”.
In the second case, the child runs immediately upon creation, sets done to 1, calls signal to wake a sleeping thread (but there is none, so it just returns), and is done. The parent then runs, calls thr_join(), sees that done is 1, and thus does not wait and returns.

One last note: you might observe the parent uses a while loop instead of just an if statement when deciding whether to wait on the condition. While this does not seem strictly necessary per the logic of the program, it is always a good idea, as we will see below.

To make sure you understand the importance of each piece of the thr_exit() and thr_join() code, let's try a few alternate implementations. First, you might be wondering if we need the state variable done. What if the code looked like the example below? Would this work?

```c
void thr_exit() {
    Pthread_mutex_lock(&m);
    Pthread_cond_signal(&c);
    Pthread_mutex_unlock(&m);
}

void thr_join() {
    Pthread_mutex_lock(&m);
    Pthread_cond_wait(&c, &m);
    Pthread_mutex_unlock(&m);
}
```

Unfortunately this approach is broken. Imagine the case where the child runs immediately and calls thr_exit() immediately; in this case, the child will signal, but there is no thread asleep on the condition. When the parent runs, it will simply call wait and be stuck; no thread will ever wake it. From this example, you should appreciate the importance of the state variable done; it records the value the threads are interested in knowing. The sleeping, waking, and locking all are built around it.

Here is another poor implementation. In this example, we imagine that one does not need to hold a lock in order to signal and wait. What problem could occur here? Think about it!

```c
void thr_exit() {
    done = 1;
    Pthread_cond_signal(&c);
}

void thr_join() {
    if (done == 0)
        Pthread_cond_wait(&c);
}
```

The issue here is a subtle race condition. Specifically, if the parent calls thr_join() and then checks the value of done, it will see that it is 0 and thus try to go to sleep. But just before it calls wait to go to sleep, the parent is interrupted, and the child runs. The child changes the state variable done to 1 and signals, but no thread is waiting and thus no thread is woken. When the parent runs again, it sleeps forever, which is sad.
Although it is strictly not necessary in all cases, it is likely simplest and best to hold the lock while signaling when using condition variables. The example above shows a case where you \textit{must} hold the lock for correctness; however, there are some other cases where it is likely OK not to, but probably is something you should avoid. Thus, for simplicity, \textbf{hold the lock when calling signal.}

The converse of this tip, i.e., hold the lock when calling \textit{wait}, is not just a tip, but rather mandated by the semantics of \textit{wait}, because \textit{wait} always (a) assumes the lock is held when you call it, (b) releases said lock when putting the caller to sleep, and (c) re-acquires the lock just before returning. Thus, the generalization of this tip is correct: \textbf{hold the lock when calling signal or wait}, and you will always be in good shape.

Hopefully, from this simple join example, you can see some of the basic requirements of using condition variables properly. To make sure you understand, we now go through a more complicated example: the \textbf{producer/consumer} or \textbf{bounded-buffer} problem.

\section{The Producer/Consumer (Bound Buffer) Problem}

The next synchronization problem we will confront in this chapter is known as the \textbf{producer/consumer} problem, or sometimes as the \textbf{bounded buffer} problem, which was first posed by Dijkstra \cite{D72}. Indeed, it was this very producer/consumer problem that led Dijkstra and his co-workers to invent the generalized semaphore (which can be used as either a lock or a condition variable) \cite{D01}; we will learn more about semaphores later.

Imagine one or more producer threads and one or more consumer threads. Producers produce data items and wish to place them in a buffer; consumers grab data items out of the buffer consume them in some way. This arrangement occurs in many real systems. For example, in a multi-threaded web server, a producer puts HTTP requests into a work queue (i.e., the bounded buffer); consumer threads take requests out of this queue and process them.

A bounded buffer is also used when you pipe the output of one program into another, e.g., `\texttt{grep foo file.txt | wc -l}`. This example runs two processes concurrently; \texttt{grep} writes lines from \texttt{file.txt} with the string \texttt{foo} in them to what it thinks is standard output; the UNIX shell redirects the output to what is called a UNIX pipe (created by the \texttt{pipe} system call). The other end of this pipe is connected to the standard input of the process \texttt{wc}, which simply counts the number of lines in the input stream and prints out the result. Thus, the \texttt{grep} process is the producer; the \texttt{wc} process is the consumer; between them is an in-kernel bounded buffer; you, in this example, are just the happy user.
int buffer;
int count = 0; // initially, empty

void put(int value) {
    assert(count == 0);
    count = 1;
    buffer = value;
}

int get() {
    assert(count == 1);
    count = 0;
    return buffer;
}

Figure 30.4: The Put and Get Routines (Version 1)

void *producer(void *arg) {
    int i;
    int loops = (int) arg;
    for (i = 0; i < loops; i++) {
        put(i);
    }
}

void *consumer(void *arg) {
    int i;
    while (1) {
        int tmp = get();
        printf("%d\n", tmp);
    }
}

Figure 30.5: Producer/Consumer Threads (Version 1)

Because the bounded buffer is a shared resource, we must of course require synchronized access to it, lest a race condition arise. To begin to understand this problem better, let us examine some actual code.

The first thing we need is a shared buffer, into which a producer puts data, and out of which a consumer takes data. Let's just use a single integer for simplicity (you can certainly imagine placing a pointer to a data structure into this slot instead), and the two inner routines to put a value into the shared buffer, and to get a value out of the buffer. See Figure 30.4 for details.

Pretty simple, no? The put() routine assumes the buffer is empty (and checks this with an assertion), and then simply puts a value into the shared buffer and marks it full by setting count to 1. The get() routine does the opposite, setting the buffer to empty (i.e., setting count to 0) and returning the value. Don’t worry that this shared buffer has just a single entry; later, we’ll generalize it to a queue that can hold multiple entries, which will be even more fun than it sounds.

Now we need to write some routines that know when it is OK to access the buffer to either put data into it or get data out of it. The conditions for

---

1This is where we drop some serious Old English on you, and the subjunctive form.
this should be obvious: only put data into the buffer when count is zero (i.e., when the buffer is empty), and only get data from the buffer when count is one (i.e., when the buffer is full). If we write the synchronization code such that a producer puts data into a full buffer, or a consumer gets data from an empty one, we have done something wrong (and in this code, an assertion will fire).

This work is going to be done by two types of threads, one set of which we’ll call the producer threads, and the other set which we’ll call consumer threads. Figure 30.5 shows the code for a producer that puts an integer into the shared buffer loops number of times, and a consumer that gets the data out of that shared buffer (forever), each time printing out the data item it pulled from the shared buffer.

A Broken Solution

Now imagine that we have just a single producer and a single consumer. Obviously the put() and get() routines have critical sections within them, as put() updates the buffer, and get() reads from it. However, putting a lock around the code doesn’t work; we need something more. Not surprisingly, that something more is some condition variables. In this (broken) first try (Figure 30.6), we have a single condition variable cond and associated lock mutex.

```c
cond_t cond;
mutex_t mutex;

void *producer(void *arg) {
    int i;
    for (i = 0; i < loops; i++) {
        Pthread_mutex_lock(&mutex); // p1
        if (count == 1) // p2
            Pthread_cond_wait(&cond, &mutex); // p3
        put(i); // p4
        Pthread_cond_signal(&cond); // p5
        Pthread_mutex_unlock(&mutex); // p6
    }
}

void *consumer(void *arg) {
    int i;
    for (i = 0; i < loops; i++) {
        Pthread_mutex_lock(&mutex); // c1
        if (count == 0) // c2
            Pthread_cond_wait(&cond, &mutex); // c3
        int tmp = get(); // c4
        Pthread_cond_signal(&cond); // c5
        Pthread_mutex_unlock(&mutex); // c6
        printf("%d
", tmp);
    }
}
```

Figure 30.6: Producer/Consumer: Single CV and If Statement
Let’s examine the signaling logic between producers and consumers. When a producer wants to fill the buffer, it waits for it to be empty (p1–p3). The consumer has the exact same logic, but waits for a different condition: fullness (c1–c3).

With just a single producer and a single consumer, the code in Figure 30.6 works. However, if we have more than one of these threads (e.g., two consumers), the solution has two critical problems. What are they?

... (pause here to think) ...

Let’s understand the first problem, which has to do with the if statement before the wait. Assume there are two consumers (T\textsubscript{c1} and T\textsubscript{c2}) and one producer (T\textsubscript{p}). First, a consumer (T\textsubscript{c1}) runs; it acquires the lock (c1), checks if any buffers are ready for consumption (c2), and finding that none are, waits (c3) (which releases the lock).

Then the producer (T\textsubscript{p}) runs. It acquires the lock (p1), checks if all buffers are full (p2), and finding that not to be the case, goes ahead and fills the buffer (p4). The producer then signals that a buffer has been filled (p5). Critically, this moves the first consumer (T\textsubscript{c1}) from sleeping on a condition variable to the ready queue; T\textsubscript{c1} is now able to run (but not yet running). The producer then continues until realizing the buffer is full, at which point it sleeps (p6, p1–p3).

Here is where the problem occurs: another consumer (T\textsubscript{c2}) sneaks in and consumes the one existing value in the buffer (c1, c2, c4, c5, c6, skipping the wait at c3 because the buffer is full). Now assume T\textsubscript{c1} runs; just before returning from the wait, it re-acquires the lock and then returns. It then calls get() (c4), but there are no buffers to consume! An assertion triggers, and the code has not functioned as desired. Clearly, we should have somehow prevented T\textsubscript{c1} from trying to consume because T\textsubscript{c2} snuck in and consumed the one value in the buffer that had been produced. Table 30.1 shows the action each thread takes, as well as its scheduler state (Ready, Running, or Sleeping) over time.

<table>
<thead>
<tr>
<th>T\textsubscript{c1} State</th>
<th>T\textsubscript{c2} State</th>
<th>T\textsubscript{p} State</th>
<th>Count</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Running</td>
<td>Ready</td>
<td>Ready</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Running</td>
<td>Ready</td>
<td>Ready</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Sleep</td>
<td>Ready</td>
<td>Ready</td>
<td>0</td>
<td>Nothing to get</td>
</tr>
<tr>
<td>Sleep</td>
<td>Ready</td>
<td>p1 Running</td>
<td>0</td>
<td>Buffer now full</td>
</tr>
<tr>
<td>Sleep</td>
<td>Ready</td>
<td>p2 Running</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Sleep</td>
<td>Ready</td>
<td>p4 Running</td>
<td>1</td>
<td>T\textsubscript{c1} awakened</td>
</tr>
<tr>
<td>Ready</td>
<td>Ready</td>
<td>p5 Running</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Ready</td>
<td>Ready</td>
<td>p6 Running</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Ready</td>
<td>Ready</td>
<td>p1 Running</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Ready</td>
<td>Ready</td>
<td>p2 Running</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Ready</td>
<td>Ready</td>
<td>p3 Sleep</td>
<td>1</td>
<td>Buffer full; sleep</td>
</tr>
<tr>
<td>Ready</td>
<td>c1 Running</td>
<td>Sleep</td>
<td>1</td>
<td>T\textsubscript{c2} sneaks in ...</td>
</tr>
<tr>
<td>Ready</td>
<td>c2 Running</td>
<td>Sleep</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Ready</td>
<td>c4 Running</td>
<td>Sleep</td>
<td>0</td>
<td>... and grabs data</td>
</tr>
<tr>
<td>Ready</td>
<td>c5 Running</td>
<td>Ready</td>
<td>0</td>
<td>T\textsubscript{p} awakened</td>
</tr>
<tr>
<td>Ready</td>
<td>c6 Running</td>
<td>Ready</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>c4 Running</td>
<td>Ready</td>
<td>Ready</td>
<td>0</td>
<td>Oh oh! No data</td>
</tr>
</tbody>
</table>

Table 30.1: Thread Trace: Broken Solution (Version 1)
Figure 30.7: Producer/Consumer: Single CV and While

The problem arises for a simple reason: after the producer woke \( T_c_1 \), but before \( T_c_1 \) ever ran, the state of the bounded buffer changed (thanks to \( T_c_2 \)). Signaling a thread only wakes them up; it is thus a hint that the state of the world has changed (in this case, that a value has been placed in the buffer), but there is no guarantee that when the woken thread runs, the state will still be as desired. This interpretation of what a signal means is often referred to as Mesa semantics, after the first research that built a condition variable in such a manner [LR80]; the contrast, referred to as Hoare semantics, is harder to build but provides a stronger guarantee that the woken thread will run immediately upon being woken [H74]. Virtually every system ever built employs Mesa semantics.

Better, But Still Broken: While, Not If

Fortunately, this fix is easy (Figure 30.7): change the if to a while. Think about why this works; now consumer \( T_c_1 \) wakes up and (with the lock held) immediately re-checks the state of the shared variable (c2). If the buffer is empty at that point, the consumer simply goes back to sleep (c3). The corollary if is also changed to a while in the producer (p2).

Thanks to Mesa semantics, a simple rule to remember with condition variables is to always use while loops. Sometimes you don’t have to re-check the condition, but it is always safe to do so; just do it and be happy.

```c
cond_t cond;
mutex_t mutex;

void *producer(void *arg) {
    int i;
    for (i = 0; i < loops; i++) {
        Pthread_mutex_lock(&mutex); // p1
        while (count == 1) // p2
            Pthread_cond_wait(&cond, &mutex); // p3
        put(i); // p4
        Pthread_cond_signal(&cond); // p5
        Pthread_mutex_unlock(&mutex); // p6
    }
}

void *consumer(void *arg) {
    int i;
    for (i = 0; i < loops; i++) {
        Pthread_mutex_lock(&mutex); // c1
        while (count == 0) // c2
            Pthread_cond_wait(&cond, &mutex); // c3
        int tmp = get(); // c4
        Pthread_cond_signal(&cond); // c5
        Pthread_mutex_unlock(&mutex); // c6
        printf("%d\n", tmp);
    }
}
However, this code still has a bug, the second of two problems mentioned above. Can you see it? It has something to do with the fact that there is only one condition variable. Try to figure out what the problem is, before reading ahead. DO IT!

Let’s confirm you figured it out correctly, or perhaps let’s confirm that you are now awake and reading this part of the book. The problem occurs when two consumers run first (Tc1 and Tc2), and both go to sleep (c3). Then, a producer runs, put a value in the buffer, wakes one of the consumers (say Tc1), and goes back to sleep. Now we have one consumer ready to run (Tc1), and two threads sleeping on a condition (Tc2 andTp). And we are about to cause a problem to occur: things are getting exciting!

The consumer Tc1 then wakes by returning from wait() (c3), re-checks the condition (c2), and finding the buffer full, consumes the value (c4). This consumer then, critically, signals on the condition (c5), waking one thread that is sleeping. However, which thread should it wake?

Because the consumer has emptied the buffer, it clearly should wake the producer. However, if it wakes the consumer Tc2 (which is definitely possible, depending on how the wait queue is managed), we have a problem. Specifically, the consumer Tc2 will wake up and find the buffer empty (c2), and go back to sleep (c3). The producer Tp, which has a value to put into the buffer, is left sleeping. The other consumer thread, Tc1, also goes back to sleep. All three threads are left sleeping, a clear bug; see Table 30.2 for the brutal step-by-step of this terrible calamity.

Signaling is clearly needed, but must be more directed. A consumer should not wake other consumers, only producers, and vice-versa.

### Table 30.2: Thread Trace: Broken Solution (Version 2)

<table>
<thead>
<tr>
<th>Tc1 State</th>
<th>Tc2 State</th>
<th>Tp State</th>
<th>Count</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Running</td>
<td>Ready</td>
<td>Ready</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Running</td>
<td>Ready</td>
<td>Ready</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Sleep</td>
<td>Ready</td>
<td>Ready</td>
<td>0</td>
<td>Nothing to get</td>
</tr>
<tr>
<td>Sleep</td>
<td>c1</td>
<td>Running</td>
<td>Ready</td>
<td>0</td>
</tr>
<tr>
<td>Sleep</td>
<td>c2</td>
<td>Running</td>
<td>Ready</td>
<td>0</td>
</tr>
<tr>
<td>Sleep</td>
<td>c3</td>
<td>Sleep</td>
<td>Ready</td>
<td>0</td>
</tr>
<tr>
<td>Sleep</td>
<td>Sleep</td>
<td>p1</td>
<td>Running</td>
<td>0</td>
</tr>
<tr>
<td>Sleep</td>
<td>Sleep</td>
<td>p2</td>
<td>Running</td>
<td>0</td>
</tr>
<tr>
<td>Sleep</td>
<td>Sleep</td>
<td>p4</td>
<td>Running</td>
<td>1</td>
</tr>
<tr>
<td>Ready</td>
<td>Sleep</td>
<td>p5</td>
<td>Running</td>
<td>1</td>
</tr>
<tr>
<td>Ready</td>
<td>Sleep</td>
<td>p6</td>
<td>Running</td>
<td>1</td>
</tr>
<tr>
<td>Ready</td>
<td>Sleep</td>
<td>p1</td>
<td>Running</td>
<td>1</td>
</tr>
<tr>
<td>Ready</td>
<td>Sleep</td>
<td>p2</td>
<td>Running</td>
<td>1</td>
</tr>
<tr>
<td>Ready</td>
<td>Sleep</td>
<td>p3</td>
<td>Sleep</td>
<td>1 Must sleep (full)</td>
</tr>
<tr>
<td>Running</td>
<td>Sleep</td>
<td>p1</td>
<td>Sleep</td>
<td>1 Recheck condition</td>
</tr>
<tr>
<td>Running</td>
<td>Sleep</td>
<td>Sleep</td>
<td>0</td>
<td>Tc1 grabs data</td>
</tr>
<tr>
<td>Running</td>
<td>Ready</td>
<td>Ready</td>
<td>0</td>
<td>Oops! Woke Tc2</td>
</tr>
<tr>
<td>Running</td>
<td>Ready</td>
<td>Sleep</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Running</td>
<td>Ready</td>
<td>Sleep</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Sleep</td>
<td>Ready</td>
<td>Sleep</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Sleep</td>
<td>c2</td>
<td>Running</td>
<td>Sleep</td>
<td>0</td>
</tr>
<tr>
<td>Sleep</td>
<td>c3</td>
<td>Sleep</td>
<td>Sleep</td>
<td>0</td>
</tr>
</tbody>
</table>

Operating Systems
[Version 0.80]  WWW.OSTEP.ORG
The Single Buffer Producer/Consumer Solution

The solution here is once again a small one: use two condition variables, instead of one, in order to properly signal which type of thread should wake up when the state of the system changes. Figure 30.8 shows the resulting code.

In the code above, producer threads wait on the condition empty, and signals fill. Conversely, consumer threads wait on fill and signal empty. By doing so, the second problem above is avoided by design: a consumer can never accidentally wake a consumer, and a producer can never accidentally wake a producer.

The Final Producer/Consumer Solution

We now have a working producer/consumer solution, albeit not a fully general one. The last change we make is to enable more concurrency and efficiency; specifically, we add more buffer slots, so that multiple values can be produced before sleeping, and similarly multiple values can be consumed before sleeping. With just a single producer and consumer, this approach is more efficient as it reduces context switches; with multiple producers or consumers (or both), it even allows concurrent producing or consuming to take place, thus increasing concurrency. Fortunately, it is a small change from our current solution.
```c
int buffer[MAX];
int fill = 0;
int use = 0;
int count = 0;

void put(int value) {
    buffer[fill] = value;
    fill = (fill + 1) % MAX;
    count++;
}

int get() {
    int tmp = buffer[use];
    use = (use + 1) % MAX;
    count--;
    return tmp;
}
```

Figure 30.9: The Final Put and Get Routines

```c
cond_t empty, fill;
mutex_t mutex;

void *producer(void *arg) {
    int i;
    for (i = 0; i < loops; i++) {
        Pthread_mutex_lock(&mutex); // p1
        while (count == MAX) // p2
            Pthread_cond_wait(&empty, &mutex); // p3
        put(i); // p4
        Pthread_mutex_unlock(&mutex); // p6
    }
}

void *consumer(void *arg) {
    int i;
    for (i = 0; i < loops; i++) {
        Pthread_mutex_lock(&mutex); // c1
        while (count == 0) // c2
            Pthread_cond_wait(&fill, &mutex); // c3
        int tmp = get(); // c4
        Pthread_mutex_unlock(&mutex); // c5
        Pthread_mutex_unlock(&mutex); // c6
        printf("%d\n", tmp);
    }
}
```

Figure 30.10: The Final Working Solution

The first change for this final solution is within the buffer structure itself and the corresponding `put()` and `get()` (Figure 30.9). We also slightly change the conditions that producers and consumers check in order to determine whether to sleep or not. Figure 30.10 shows the final waiting and signaling logic. A producer only sleeps if all buffers are currently filled (p2); similarly, a consumer only sleeps if all buffers are currently empty (c2). And thus we solve the producer/consumer problem.
When checking for a condition in a multi-threaded program, using a `while` loop is always correct; using an `if` statement only might be, depending on the semantics of signaling. Thus, always use `while` and your code will behave as expected.

Using while loops around conditional checks also handles the case where spurious wakeups occur. In some thread packages, due to details of the implementation, it is possible that two threads get woken up though just a single signal has taken place [L11]. Spurious wakeups are further reason to re-check the condition a thread is waiting on.

### 30.3 Covering Conditions

We’ll now look at one more example of how condition variables can be used. This code study is drawn from Lampson and Redell’s paper on Pilot [LR80], the same group who first implemented the Mesa semantics described above (the language they used was Mesa, hence the name).

The problem they ran into is best shown via simple example, in this case in a simple multi-threaded memory allocation library. Figure 30.11 shows a code snippet which demonstrates the issue.

As you might see in the code, when a thread calls into the memory allocation code, it might have to wait in order for more memory to become free. Conversely, when a thread frees memory, it signals that more memory is free. However, our code above has a problem: which waiting thread (there can be more than one) should be woken up?

Consider the following scenario. Assume there are zero bytes free; thread $T_a$ calls `allocate(100)`, followed by thread $T_b$ which asks for less memory by calling `allocate(10)`. Both $T_a$ and $T_b$ thus wait on the condition and go to sleep; there aren’t enough free bytes to satisfy either of these requests.

At that point, assume a third thread, $T_c$, calls `free(50)`. Unfortunately, when it calls signal to wake a waiting thread, it might not wake the correct waiting thread, $T_b$, which is waiting for only 10 bytes to be freed; $T_a$ should remain waiting, as not enough memory is yet free. Thus, the code in the figure does not work, as the thread waking other threads does not know which thread (or threads) to wake up.

The solution suggested by Lampson and Redell is straightforward: replace the `pthread_cond_signal()` call in the code above with a call to `pthread_cond_broadcast()`, which wakes up all waiting threads. By doing so, we guarantee that any threads that should be woken are. The downside, of course, can be a negative performance impact, as we might needlessly wake up many other waiting threads that shouldn’t (yet) be awake. Those threads will simply wake up, re-check the condition, and then go immediately back to sleep.
```c
// how many bytes of the heap are free?
int bytesLeft = MAX_HEAP_SIZE;

// need lock and condition too
cond_t  c;
mutex_t m;

void *
allocate(int size) {
    Pthread_mutex_lock(&m);
    while (bytesLeft < size)
        Pthread_cond_wait(&c, &m);
    void *ptr = ...; // get mem from heap
    bytesLeft -= size;
    Pthread_mutex_unlock(&m);
    return ptr;
}

void free(void *ptr, int size) {
    Pthread_mutex_lock(&m);
    bytesLeft += size;
    Pthread_cond_signal(&c); // whom to signal??
    Pthread_mutex_unlock(&m);
}
```

Figure 30.11: Covering Conditions: An Example

Lampson and Redell call such a condition a covering condition, as it covers all the cases where a thread needs to wake up (conservatively); the cost, as we’ve discussed, is that too many threads might be woken. The astute reader might also have noticed we could have used this approach earlier (see the producer/consumer problem with only a single condition variable). However, in that case, a better solution was available to us, and thus we used it. In general, if you find that your program only works when you change your signals to broadcasts (but you don’t think it should need to), you probably have a bug; fix it! But in cases like the memory allocator above, broadcast may be the most straightforward solution available.

30.4 Summary

We have seen the introduction of another important synchronization primitive beyond locks: condition variables. By allowing threads to sleep when some program state is not as desired, CVs enable us to neatly solve a number of important synchronization problems, including the famous (and still important) producer/consumer problem, as well as covering conditions. A more dramatic concluding sentence would go here, such as “He loved Big Brother” [O49].
References

[D72] “Information Streams Sharing a Finite Buffer”
E.W. Dijkstra
Information Processing Letters 1: 179180, 1972
Available: http://www.cs.utexas.edu/users/EWD/ewd03xx/EWD329.PDF
The famous paper that introduced the producer/consumer problem.

[D01] “My recollections of operating system design”
E.W. Dijkstra
April, 2001
Available: http://www.cs.utexas.edu/users/EWD/ewd13xx/EWD1303.PDF
A fascinating read for those of you interested in how the pioneers of our field came up with some very basic and fundamental concepts, including ideas like “interrupts” and even “a stack”!

[H74] “Monitors: An Operating System Structuring Concept”
C.A.R. Hoare
Communications of the ACM, 17:10, pages 549–557, October 1974
Hoare did a fair amount of theoretical work in concurrency. However, he is still probably most known for his work on Quicksort, the coolest sorting algorithm in the world, at least according to these authors.

Available: http://linux.die.net/man/3/pthread_cond_signal
March, 2011
The Linux man page shows a nice simple example of why a thread might get a spurious wakeup, due to race conditions within the signal/wakeup code.

[LR80] “Experience with Processes and Monitors in Mesa”
B.W. Lampson, D.R. Redell
Communications of the ACM. 23:2, pages 105-117, February 1980
A terrific paper about how to actually implement signaling and condition variables in a real system, leading to the term “Mesa” semantics for what it means to be woken up; the older semantics, developed by Tony Hoare [H74], then became known as “Hoare” semantics, which is hard to say out loud in class with a straight face.

[O49] “1984”
George Orwell, 1949, Secker and Warburg
A little heavy-handed, but of course a must read. That said, we kind of gave away the ending by quoting the last sentence. Sorry! And if the government is reading this, let us just say that we think that the government is “double plus good”. Hear that, our pals at the NSA?
31

Semaphores

As we know now, one needs both locks and condition variables to solve a broad range of relevant and interesting concurrency problems. One of the first people to realize this years ago was Edsger Dijkstra (though it is hard to know the exact history [GR92]), known among other things for his famous “shortest paths” algorithm in graph theory [D59], an early polemic on structured programming entitled “Goto Statements Considered Harmful” [D68a] (what a great title!), and, in the case we will study here, the introduction of a synchronization primitive called the semaphore [D68b,D72]. Indeed, Dijkstra and colleagues invented the semaphore as a single primitive for all things related to synchronization; as you will see, one can use semaphores as both locks and condition variables.

THE CRUX: HOW TO USE SEMAPHORES
How can we use semaphores instead of locks and condition variables? What is the definition of a semaphore? What is a binary semaphore? Is it straightforward to build a semaphore out of locks and condition variables? What about building locks and condition variables out of semaphores?

31.1 Semaphores: A Definition

A semaphore is as an object with an integer value that we can manipulate with two routines; in the POSIX standard, these routines are sem_wait() and sem_post(). Because the initial value of the semaphore determines its behavior, before calling any other routine to interact with the semaphore, we must first initialize it to some value, as the code in Figure 31.1 does.

1Historically, sem_wait() was first called P() by Dijkstra (for the Dutch word “to probe”) and sem_post() was called V() (for the Dutch word “to test”). Sometimes, people call them down and up, too. Use the Dutch versions to impress your friends.
#include <semaphore.h>

sem_t s;
sem_init(&s, 0, 1);

Figure 31.1: Initializing A Semaphore

In the figure, we declare a semaphore s and initialize it to the value 1 by passing 1 in as the third argument. The second argument to sem_init() will be set to 0 in all of the examples we’ll see; this indicates that the semaphore is shared between threads in the same process. See the man page for details on other usages of semaphores (namely, how they can be used to synchronize access across different processes), which require a different value for that second argument.

After a semaphore is initialized, we can call one of two functions to interact with it, sem_wait() or sem_post(). The behavior of these two functions is seen in Figure 31.2.

For now, we are not concerned with the implementation of these routines, which clearly requires some care; with multiple threads calling into sem_wait() and sem_post(), there is the obvious need for managing these critical sections. We will now focus on how to use these primitives; later we may discuss how they are built.

We should discuss a few salient aspects of the interfaces here. First, we can see that sem_wait() will either return right away (because the value of the semaphore was one or higher when we called sem_wait()), or it will cause the caller to suspend execution waiting for a subsequent post. Of course, multiple calling threads may call into sem_wait(), and thus all be queued waiting to be woken.

Second, we can see that sem_post() does not wait for some particular condition to hold like sem_wait() does. Rather, it simply increments the value of the semaphore and then, if there is a thread waiting to be woken, wakes one of them up.

Third, the value of the semaphore, when negative, is equal to the number of waiting threads [D68b]. Though the value generally isn’t seen by users of the semaphores, this invariant is worth knowing and perhaps can help you remember how a semaphore functions.

Don’t worry (yet) about the seeming race conditions possible within the semaphore; assume that the actions they make are performed atomically. We will soon use locks and condition variables to do just this.

1 int sem_wait(sem_t *s) {
  2     decrement the value of semaphore s by one
  3     wait if value of semaphore s is negative
  4 }
1 5
6 int sem_post(sem_t *s) {
  7     increment the value of semaphore s by one
  8     if there are one or more threads waiting, wake one
  9 }
We are now ready to use a semaphore. Our first use will be one with which we are already familiar: using a semaphore as a lock. See Figure 31.3 for a code snippet; therein, you’ll see that we simply surround the critical section of interest with a `sem_wait()`/`sem_post()` pair. Critical to making this work, though, is the initial value of the semaphore \( m \) (initialized to \( X \) in the figure). What should \( X \) be?

... (Try thinking about it before going on) ...

Looking back at definition of the `sem_wait()` and `sem_post()` routines above, we can see that the initial value should be 1.

To make this clear, let’s imagine a scenario with two threads. The first thread (Thread 0) calls `sem_wait()`; it will first decrement the value of the semaphore, changing it to 0. Then, it will wait only if the value is not greater than or equal to 0; because the value is 0, the calling thread will simply return and continue; Thread 0 is now free to enter the critical section. If no other thread tries to acquire the lock while Thread 0 is inside the critical section, when it calls `sem_post()`, it will simply restore the value of the semaphore to 1 (and not wake any waiting thread, because there are none). Table 31.1 shows a trace of this scenario.

A more interesting case arises when Thread 0 “holds the lock” (i.e., it has called `sem_wait()` but not yet called `sem_post()`), and another thread (Thread 1) tries to enter the critical section by calling `sem_wait()`. In this case, Thread 1 will decrement the value of the semaphore to -1, and thus wait (putting itself to sleep and relinquishing the processor). When Thread 0 runs again, it will eventually call `sem_post()`, incrementing the value of the semaphore back to zero, and then wake the waiting thread (Thread 1), which will then be able to acquire the lock for itself. When Thread 1 finishes, it will again increment the value of the semaphore, restoring it to 1 again.

<table>
<thead>
<tr>
<th>Value of Semaphore</th>
<th>Thread 0</th>
<th>Thread 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>call <code>sem_wait()</code></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td><code>sem_wait()</code> returns</td>
<td></td>
</tr>
<tr>
<td>0 (crit sect)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>call <code>sem_post()</code></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td><code>sem_post()</code> returns</td>
<td></td>
</tr>
</tbody>
</table>

Table 31.1: Thread Trace: Single Thread Using A Semaphore
Table 31.2: Thread Trace: Two Threads Using A Semaphore

Table 31.2 shows a trace of this example. In addition to thread actions, the table shows the scheduler state of each thread: Running, Ready (i.e., runnable but not running), and Sleeping. Note in particular that Thread 1 goes into the sleeping state when it tries to acquire the already-held lock; only when Thread 0 runs again can Thread 1 be awoken and potentially run again.

If you want to work through your own example, try a scenario where multiple threads queue up waiting for a lock. What would the value of the semaphore be during such a trace?

Thus we are able to use semaphores as locks. Because locks only have two states (held and not held), this usage is sometimes known as a binary semaphore and in fact can be implemented in a more simplified manner than discussed here; we instead use the generalized semaphore as a lock.

31.3 Semaphores As Condition Variables

Semaphores are also useful when a thread wants to halt its progress waiting for a condition to become true. For example, a thread may wish to wait for a list to become non-empty, so it can delete an element from it. In this pattern of usage, we often find a thread waiting for something to happen, and a different thread making that something happen and then signaling that it has happened, thus waking the waiting thread. Because the waiting thread (or threads) is waiting for some condition in the program to change, we are using the semaphore as a condition variable.
A simple example is as follows. Imagine a thread creates another thread and then wants to wait for it to complete its execution (Figure 31.4). When this program runs, we would like to see the following:

parent: begin
child
parent: end

The question, then, is how to use a semaphore to achieve this effect, and as it turns out, it is relatively easy to understand. As you can see in the code, the parent simply calls `sem_wait()` and the child `sem_post()` to wait for the condition of the child finishing its execution to become true. However, this raises the question: what should the initial value of this semaphore be?

(Again, think about it here, instead of reading ahead)

The answer, of course, is that the value of the semaphore should be set to 0. There are two cases to consider. First, let us assume that the parent creates the child but the child has not run yet (i.e., it is sitting in a ready queue but not running). In this case (Table 31.3), the parent will call `sem_wait()` before the child has called `sem_post()`; we'd like the parent to wait for the child to run. The only way this will happen is if the value of the semaphore is not greater than 0; hence, 0 is the initial value. The parent runs, decrements the semaphore (to -1), then waits (sleeping). When the child finally runs, it will call `sem_post()`, increment the value of the semaphore to 0, and wake the parent, which will then return from `sem_wait()` and finish the program.

The second case (Table 31.4) occurs when the child runs to completion before the parent gets a chance to call `sem_wait()`. In this case, the child will first call `sem_post()`, thus incrementing the value of the semaphore from 0 to 1. When the parent then gets a chance to run, it will call `sem_wait()` and find the value of the semaphore to be 1; the parent will then decrement the value (to 0) and return from `sem_wait()` without waiting, also achieving the desired effect.
Table 31.3: Thread Trace: Parent Waiting For Child (Case 1)

<table>
<thead>
<tr>
<th>Value</th>
<th>Parent</th>
<th>State</th>
<th>Child</th>
<th>State</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>create(Child)</td>
<td>Running</td>
<td>(Child exists; is runnable)</td>
<td>Ready</td>
</tr>
<tr>
<td>0</td>
<td>call sem_wait()</td>
<td>Running</td>
<td></td>
<td>Ready</td>
</tr>
<tr>
<td>-1</td>
<td>decrement sem</td>
<td>Running</td>
<td></td>
<td>Ready</td>
</tr>
<tr>
<td>-1</td>
<td>(sem&lt;0)→sleep</td>
<td>Sleeping</td>
<td>child runs</td>
<td>Running</td>
</tr>
<tr>
<td>-1</td>
<td>Switch→Child</td>
<td>Sleeping</td>
<td>call sem_post()</td>
<td>Running</td>
</tr>
<tr>
<td>0</td>
<td>Sleep</td>
<td>Running</td>
<td>increment sem</td>
<td>Running</td>
</tr>
<tr>
<td>0</td>
<td>Ready</td>
<td>Running</td>
<td>wake(Parent)</td>
<td>Running</td>
</tr>
<tr>
<td>0</td>
<td>Ready</td>
<td>Running</td>
<td>sem_post() returns</td>
<td>Running</td>
</tr>
<tr>
<td>0</td>
<td>Ready</td>
<td>Running</td>
<td>Interrupt; Switch→Parent</td>
<td>Ready</td>
</tr>
<tr>
<td>0</td>
<td>sem_wait() returns</td>
<td>Ready</td>
<td></td>
<td>Ready</td>
</tr>
</tbody>
</table>

Table 31.4: Thread Trace: Parent Waiting For Child (Case 2)

31.4 The Producer/Consumer (Bounded-Buffer) Problem

The next problem we will confront in this chapter is known as the producer/consumer problem, or sometimes as the bounded buffer problem [D72]. This problem is described in detail in the previous chapter on condition variables; see there for details.

First Attempt

Our first attempt at solving the problem introduces two semaphores, empty and full, which the threads will use to indicate when a buffer entry has been emptied or filled, respectively. The code for the put and get routines is in Figure 31.5, and our attempt at solving the producer and consumer problem is in Figure 31.6.

In this example, the producer first waits for a buffer to become empty in order to put data into it, and the consumer similarly waits for a buffer to become filled before using it. Let us first imagine that MAX=1 (there is only one buffer in the array), and see if this works.

Imagine again there are two threads, a producer and a consumer. Let us examine a specific scenario on a single CPU. Assume the consumer gets to run first. Thus, the consumer will hit line c1 in the figure above, calling sem_wait(&full). Because full was initialized to the value 0,
int buffer[MAX];
int fill = 0;
int use = 0;

void put(int value) {
    buffer[fill] = value; // line f1
    fill = (fill + 1) % MAX; // line f2
}

int get() {
    int tmp = buffer[use]; // line g1
    use = (use + 1) % MAX; // line g2
    return tmp;
}

Figure 31.5: The Put and Get Routines

void *producer(void *arg) {
    int i;
    for (i = 0; i < loops; i++) {
        sem_wait(&empty); // line P1
        put(i); // line P2
        sem_post(&full); // line P3
    }
}

void *consumer(void *arg) {
    int i, tmp = 0;
    while (tmp != -1) {
        sem_wait(&full); // line C1
        tmp = get(); // line C2
        sem_post(&empty); // line C3
        printf("%d\n", tmp);
    }
}

int main(int argc, char *argv[]) {
    // ...
    sem_init(&empty, 0, MAX); // MAX buffers are empty to begin with...
    sem_init(&full, 0, 0); // ... and 0 are full
    // ...
}

Figure 31.6: Adding the Full and Empty Conditions

the call will decrement full (to -1), block the consumer, and wait for another thread to call sem_post() on full, as desired.

Assume the producer then runs. It will hit line P1, thus calling the sem_wait(&empty) routine. Unlike the consumer, the producer will continue through this line, because empty was initialized to the value MAX (in this case, 1). Thus, empty will be decremented to 0 and the producer will put a data value into the first entry of buffer (line P2). The producer will then continue on to P3 and call sem_post(&full), changing the value of the full semaphore from -1 to 0 and waking the consumer (e.g., move it from blocked to ready).
In this case, one of two things could happen. If the producer continues to run, it will loop around and hit line P1 again. This time, however, it would block, as the empty semaphore’s value is 0. If the producer instead was interrupted and the consumer began to run, it would call `sem_wait(&full)` (line c1) and find that the buffer was indeed full and thus consume it. In either case, we achieve the desired behavior.

You can try this same example with more threads (e.g., multiple producers, and multiple consumers). It should still work.

Let us now imagine that MAX is greater than 1 (say $\text{MAX} = 10$). For this example, let us assume that there are multiple producers and multiple consumers. We now have a problem: a race condition. Do you see where it occurs? (take some time and look for it) If you can’t see it, here’s a hint: look more closely at the put() and get() code.

OK, let’s understand the issue. Imagine two producers (Pa and Pb) both calling into put() at roughly the same time. Assume producer Pa gets to run first, and just starts to fill the first buffer entry (fill = 0 at line f1). Before Pa gets a chance to increment the fill counter to 1, it is interrupted. Producer Pb starts to run, and at line f1 it also puts its data into the 0th element of buffer, which means that the old data there is overwritten! This is a no-no; we don’t want any data from the producer to be lost.

**A Solution: Adding Mutual Exclusion**

As you can see, what we’ve forgotten here is *mutual exclusion*. The filling of a buffer and incrementing of the index into the buffer is a critical section, and thus must be guarded carefully. So let’s use our friend the binary semaphore and add some locks. Figure 31.7 shows our attempt.

Now we’ve added some locks around the entire put()/get() parts of the code, as indicated by the NEW LINE comments. That seems like the right idea, but it also doesn’t work. Why? Deadlock. Why does deadlock occur? Take a moment to consider it; try to find a case where deadlock arises. What sequence of steps must happen for the program to deadlock?

**Avoiding Deadlock**

OK, now that you figured it out, here is the answer. Imagine two threads, one producer and one consumer. The consumer gets to run first. It acquires the mutex (line c0), and then calls `sem_wait()` on the full semaphore (line c1); because there is no data yet, this call causes the consumer to block and thus yield the CPU; importantly, though, the consumer still holds the lock.

A producer then runs. It has data to produce and if it were able to run, it would be able to wake the consumer thread and all would be good. Unfortunately, the first thing it does is call `sem_wait()` on the binary mutex semaphore (line p0). The lock is already held. Hence, the producer is now stuck waiting too.
```c
1 #include <semaphore.h>

2 void *producer(void *arg) {  
3     int i;
4     for (i = 0; i < loops; i++) {
5         sem_wait(&mutex); // line p0 (NEW LINE)
6         sem_wait(&empty); // line p1
7         put(i); // line p2
8         sem_post(&full); // line p3
9         sem_post(&mutex); // line p4 (NEW LINE)
10     }
11 }
12 
13 void *consumer(void *arg) {
14     int i;
15     for (i = 0; i < loops; i++) {
16         sem_wait(&mutex); // line c0 (NEW LINE)
17         sem_wait(&full); // line c1
18         int tmp = get(); // line c2
19         sem_post(&empty); // line c3
20         sem_post(&mutex); // line c4 (NEW LINE)
21         printf("%d\n", tmp);
22     }
23 }
24
25 int main(int argc, char *argv[]) {
26 // ...
27     sem_init(&empty, 0, MAX); // MAX buffers are empty to begin with...
28     sem_init(&full, 0, 0); // ... and 0 are full
29     sem_init(&mutex, 0, 1); // mutex=1 because it is a lock (NEW LINE)
30     // ...
31 }
```

Figure 31.7: Adding Mutual Exclusion (Incorrectly)

There is a simple cycle here. The consumer holds the mutex and is waiting for someone to signal full. The producer could signal full but is waiting for the mutex. Thus, the producer and consumer are each stuck waiting for each other: a classic deadlock.

Finally, A Working Solution

To solve this problem, we simply must reduce the scope of the lock. Figure 31.8 shows the final working solution. As you can see, we simply move the mutex acquire and release to be just around the critical section; the full and empty wait and signal code is left outside. The result is a simple and working bounded buffer, a commonly-used pattern in multithreaded programs. Understand it now; use it later. You will thank us for years to come. Or at least, you will thank us when the same question is asked on the final exam.
void *producer(void *arg) {
    int i;
    for (i = 0; i < loops; i++) {
        sem_wait(&empty); // line p1
        sem_wait(&mutex); // line p1.5 (MOVED MUTEX HERE...)
        put(i); // line p2
        sem_post(&mutex); // line p2.5 (... AND HERE)
        sem_post(&full); // line p3
    }
}

void *consumer(void *arg) {
    int i;
    for (i = 0; i < loops; i++) {
        sem_wait(&full); // line c1
        sem_wait(&mutex); // line c1.5 (MOVED MUTEX HERE...)
        int tmp = get(); // line c2
        sem_post(&mutex); // line c2.5 (... AND HERE)
        sem_post(&empty); // line c3
        printf("%d\n", tmp);
    }
}

int main(int argc, char *argv[]) {
    sem_init(&empty, 0, MAX); // MAX buffers are empty to begin with...
    sem_init(&full, 0, 0); // ... and 0 are full
    sem_init(&mutex, 0, 1); // mutex=1 because it is a lock
    // ...
}

Figure 31.8: Adding Mutual Exclusion (Correctly)

31.5 Reader-Writer Locks

Another classic problem stems from the desire for a more flexible locking primitive that admits that different data structure accesses might require different kinds of locking. For example, imagine a number of concurrent list operations, including inserts and simple lookups. While inserts change the state of the list (and thus a traditional critical section makes sense), lookups simply read the data structure; as long as we can guarantee that no insert is on-going, we can allow many lookups to proceed concurrently. The special type of lock we will now develop to support this type of operation is known as a reader-writer lock [CHP71]. The code for such a lock is available in Figure 31.9.

The code is pretty simple. If some thread wants to update the data structure in question, it should call the new pair of synchronization operations: rwlock_acquire_writelock(), to acquire a write lock, and rwlock_release_writelock(), to release it. Internally, these simply use the writelock semaphore to ensure that only a single writer can ac-
typedef struct _rwlock_t {
    sem_t lock; // binary semaphore (basic lock)
    sem_t writelock; // used to allow ONE writer or MANY readers
    int readers; // count of readers reading in critical section
} rwlock_t;

void rwlock_init(rwlock_t *rw) {
    rw->readers = 0;
    sem_init(&rw->lock, 0, 1);
    sem_init(&rw->writelock, 0, 1);
}

void rwlock_acquire_readlock(rwlock_t *rw) {
    sem_wait(&rw->lock);
    rw->readers++;
    if (rw->readers == 1)
        sem_wait(&rw->writelock); // first reader acquires writelock
    sem_post(&rw->lock);
}

void rwlock_release_readlock(rwlock_t *rw) {
    sem_wait(&rw->lock);
    rw->readers--;
    if (rw->readers == 0)
        sem_post(&rw->writelock); // last reader releases writelock
    sem_post(&rw->lock);
}

void rwlock_acquire_writelock(rwlock_t *rw) {
    sem_wait(&rw->writelock);
}

void rwlock_release_writelock(rwlock_t *rw) {
    sem_post(&rw->writelock);
}

Figure 31.9: A Simple Reader-Writer Lock

quire the lock and thus enter the critical section to update the data structure in question.

More interesting is the pair of routines to acquire and release read locks. When acquiring a read lock, the reader first acquires lock and then increments the readers variable to track how many readers are currently inside the data structure. The important step then taken within rwlock_acquire_readlock() occurs when the first reader acquires the lock; in that case, the reader also acquires the write lock by calling sem_wait() on the writelock semaphore, and then finally releasing the lock by calling sem_post().

Thus, once a reader has acquired a read lock, more readers will be allowed to acquire the read lock too; however, any thread that wishes to acquire the write lock will have to wait until all readers are finished; the last one to exit the critical section calls sem_post() on “writelock” and thus enables a waiting writer to acquire the lock.

This approach works (as desired), but does have some negatives, espe-
TIP: SIMPLE AND DUMB CAN BE BETTER (HILL’S LAW)
You should never underestimate the notion that the simple and dumb approach can be the best one. With locking, sometimes a simple spin lock works best, because it is easy to implement and fast. Although something like reader/writer locks sounds cool, they are complex, and complex can mean slow. Thus, always try the simple and dumb approach first. This idea, of appealing to simplicity, is found in many places. One early source is Mark Hill’s dissertation [H87], which studied how to design caches for CPUs. Hill found that simple direct-mapped caches worked better than fancy set-associative designs (one reason is that in caching, simpler designs enable faster lookups). As Hill succinctly summarized his work: “Big and dumb is better.” And thus we call this similar advice Hill’s Law.

Socially when it comes to fairness. In particular, it would be relatively easy for readers to starve writers. More sophisticated solutions to this problem exist; perhaps you can think of a better implementation? Hint: think about what you would need to do to prevent more readers from entering the lock once a writer is waiting.

Finally, it should be noted that reader-writer locks should be used with some caution. They often add more overhead (especially with more sophisticated implementations), and thus do not end up speeding up performance as compared to just using simple and fast locking primitives [CB08]. Either way, they showcase once again how we can use semaphores in an interesting and useful way.

31.6 The Dining Philosophers

One of the most famous concurrency problems posed, and solved, by Dijkstra, is known as the dining philosopher’s problem [DHO71]. The problem is famous because it is fun and somewhat intellectually interesting; however, its practical utility is low. However, its fame forces its inclusion here; indeed, you might be asked about it on some interview, and you’d really hate your OS professor if you miss that question and don’t get the job. Conversely, if you get the job, please feel free to send your OS professor a nice note, or some stock options.

The basic setup for the problem is this (as shown in Figure 31.10): assume there are five “philosophers” sitting around a table. Between each pair of philosophers is a single fork (and thus, five total). The philosophers each have times where they think, and don’t need any forks, and times where they eat. In order to eat, a philosopher needs two forks, both the one on their left and the one on their right. The contention for these forks, and the synchronization problems that ensue, are what makes this a problem we study in concurrent programming.
Here is the basic loop of each philosopher:

```c
while (1) {
    think();
    getforks();
    eat();
    putforks();
}
```

The key challenge, then, is to write the routines `getforks()` and `putforks()` such that there is no deadlock, no philosopher starves and never gets to eat, and concurrency is high (i.e., as many philosophers can eat at the same time as possible).

Following Downey’s solutions [D08], we’ll use a few helper functions to get us towards a solution. They are:

```c
int left(int p) { return p; }
int right(int p) { return (p + 1) % 5; }
```

When philosopher \( p \) wishes to refer to the fork on their left, they simply call `left(p)`. Similarly, the fork on the right of a philosopher \( p \) is referred to by calling `right(p)`; the modulo operator therein handles the one case where the last philosopher (\( p=4 \)) tries to grab the fork on their right, which is fork 0.

We’ll also need some semaphores to solve this problem. Let us assume we have five, one for each fork: `sem_t forks[5]`. 

void getforks() {
    sem_wait(forks[left(p)]);
    sem_wait(forks[right(p)]);
}

void putforks() {
    sem_post(forks[left(p)]);
    sem_post(forks[right(p)]);
}

Figure 31.11: The getforks() and putforks() Routines

Broken Solution

We attempt our first solution to the problem. Assume we initialize each semaphore (in the forks array) to a value of 1. Assume also that each philosopher knows its own number (p). We can thus write the getforks() and putforks() routine as shown in Figure 31.11.

The intuition behind this (broken) solution is as follows. To acquire the forks, we simply grab a “lock” on each one: first the one on the left, and then the one on the right. When we are done eating, we release them. Simple, no? Unfortunately, in this case, simple means broken. Can you see the problem that arises? Think about it.

The problem is deadlock. If each philosopher happens to grab the fork on their left before any philosopher can grab the fork on their right, each will be stuck holding one fork and waiting for another, forever. Specifically, philosopher 0 grabs fork 0, philosopher 1 grabs fork 1, philosopher 2 grabs fork 2, philosopher 3 grabs fork 3, and philosopher 4 grabs fork 4; all the forks are acquired, and all the philosophers are stuck waiting for a fork that another philosopher possesses. We’ll study deadlock in more detail soon; for now, it is safe to say that this is not a working solution.

A Solution: Breaking The Dependency

The simplest way to attack this problem is to change how forks are acquired by at least one of the philosophers; indeed, this is how Dijkstra himself solved the problem. Specifically, let’s assume that philosopher 4 (the highest numbered one) acquires the forks in a different order. The code to do so is as follows:

Because the last philosopher tries to grab right before left, there is no situation where each philosopher grabs one fork and is stuck waiting for another; the cycle of waiting is broken. Think through the ramifications of this solution, and convince yourself that it works.
typedef struct __Zem_t {
    int value;
    pthread_cond_t cond;
    pthread_mutex_t lock;
} Zem_t;

// only one thread can call this
void Zem_init(Zem_t *s, int value) {
    s->value = value;
    Cond_init(&s->cond);
    Mutex_init(&s->lock);
}

void Zem_wait(Zem_t *s) {
    Mutex_lock(&s->lock);
    while (s->value <= 0)
        Cond_wait(&s->cond, &s->lock);
    s->value--;
    Mutex_unlock(&s->lock);
}

void Zem_post(Zem_t *s) {
    Mutex_lock(&s->lock);
    s->value++;
    Cond_signal(&s->cond);
    Mutex_unlock(&s->lock);
}

Figure 31.12: Implementing Zemaphores with Locks and CVs

There are other “famous” problems like this one, e.g., the cigarette smoker’s problem or the sleeping barber problem. Most of them are just excuses to think about concurrency; some of them have fascinating names. Look them up if you are interested in learning more, or just getting more practice thinking in a concurrent manner [D08].

31.7 How To Implement Semaphores

Finally, let’s use our low-level synchronization primitives, locks and condition variables, to build our own version of semaphores called ... (drum roll here) ... Zemaphores. This task is fairly straightforward, as you can see in Figure 31.12.

As you can see from the figure, we use just one lock and one condition variable, plus a state variable to track the value of the semaphore. Study the code for yourself until you really understand it. Do it!

One subtle difference between our Zemaphore and pure semaphores as defined by Dijkstra is that we don’t maintain the invariant that the value of the semaphore, when negative, reflects the number of waiting threads; indeed, the value will never be lower than zero. This behavior is easier to implement and matches the current Linux implementation.
The abstract technique of generalization can thus be quite useful in systems design, where one good idea can be made slightly broader and thus solve a larger class of problems. However, be careful when generalizing; as Lampson warns us “Don’t generalize; generalizations are generally wrong” [L83].

One could view semaphores as a generalization of locks and condition variables; however, is such a generalization needed? And, given the difficulty of realizing a condition variable on top of a semaphore, perhaps this generalization is not as general as you might think.

Curiously, building locks and condition variables out of semaphores is a much trickier proposition. Some highly experienced concurrent programmers tried to do this in the Windows environment, and many different bugs ensued [B04]. Try it yourself, and see if you can figure out why building condition variables out of semaphores is more challenging than it might appear.

31.8 Summary

Semaphores are a powerful and flexible primitive for writing concurrent programs. Some programmers use them exclusively, shunning locks and condition variables, due to their simplicity and utility.

In this chapter, we have presented just a few classic problems and solutions. If you are interested in finding out more, there are many other materials you can reference. One great (and free reference) is Allen Downey’s book on concurrency and programming with semaphores [D08]. This book has lots of puzzles you can work on to improve your understanding of both semaphores in specific and concurrency in general. Becoming a real concurrency expert takes years of effort; going beyond what you learn in this class is undoubtedly the key to mastering such a topic.
References

[B04] “Implementing Condition Variables with Semaphores”  
Andrew Birrell  
December 2004  
An interesting read on how difficult implementing CVs on top of semaphores really is, and the mistakes the author and co-workers made along the way. Particularly relevant because the group had done a ton of concurrent programming; Birrell, for example, is known for (among other things) writing various thread-programming guides.

[CB08] “Real-world Concurrency”  
Bryan Cantrill and Jeff Bonwick  
ACM Queue. Volume 6, No. 5. September 2008  
A nice article by some kernel hackers from a company formerly known as Sun on the real problems faced in concurrent code.

[CHP71] “Concurrent Control with Readers and Writers”  
P.J. Courtosis, F. Heymans, D.L. Parnas  
Communications of the ACM, 14:10, October 1971  
The introduction of the reader-writer problem, and a simple solution. Later work introduced more complex solutions, skipped here because, well, they are pretty complex.

[D59] “A Note on Two Problems in Connexion with Graphs”  
E. W. Dijkstra  
Numerische Mathematik 1, 269-271, 1959  
Can you believe people worked on algorithms in 1959? We can’t. Even before computers were any fun to use, these people had a sense that they would transform the world...

[D68a] “Go-to Statement Considered Harmful”  
E.W. Dijkstra  
Sometimes thought as the beginning of the field of software engineering.

[D68b] “The Structure of the THE Multiprogramming System”  
E.W. Dijkstra  
One of the earliest papers to point out that systems work in computer science is an engaging intellectual endeavor. Also argues strongly for modularity in the form of layered systems.

[D72] “Information Streams Sharing a Finite Buffer”  
E.W. Dijkstra  
Information Processing Letters 1: 179-180, 1972  
Did Dijkstra invent everything? No, but maybe close. He certainly was the first to clearly write down what the problems were in concurrent code. However, it is true that practitioners in operating system design knew of many of the problems described by Dijkstra, so perhaps giving him too much credit would be a misrepresentation of history.

[D08] “The Little Book of Semaphores”  
A.B. Downey  
A nice (and free!) book about semaphores. Lots of fun problems to solve, if you like that sort of thing.
“Hierarchical ordering of sequential processes” E.W. Dijkstra
Available: http://www.cs.utexas.edu/users/EWD/ewd03xx/EWD310.PDF
Presents numerous concurrency problems, including the Dining Philosophers. The wikipedia page about this problem is also quite informative.

Jim Gray and Andreas Reuter
Morgan Kaufmann, September 1992
The exact quote that we find particularly humorous is found on page 485, at the top of Section 8.8: “The first multiprocessors, circa 1960, had test and set instructions ... presumably the OS implementors worked out the appropriate algorithms, although Dijkstra is generally credited with inventing semaphores many years later.”

[H87] “Aspects of Cache Memory and Instruction Buffer Performance”
Mark D. Hill
Hill’s dissertation work, for those obsessed with caching in early systems. A great example of a quantitative dissertation.
Common Concurrency Problems

Researchers have spent a great deal of time and effort looking into concurrency bugs over many years. Much of the early work focused on deadlock, a topic which we’ve touched on in the past chapters but will now dive into deeply [C+71]. More recent work focuses on studying other types of common concurrency bugs (i.e., non-deadlock bugs). In this chapter, we take a brief look at some example concurrency problems found in real code bases, to better understand what problems to look out for. And thus our problem:

**CRUX: HOW TO HANDLE COMMON CONCURRENCY BUGS**
Concurrency bugs tend to come in a variety of common patterns. Knowing which ones to look out for is the first step to writing more robust, correct concurrent code.

32.1 What Types Of Bugs Exist?

The first, and most obvious, question is this: what types of concurrency bugs manifest in complex, concurrent programs? This question is difficult to answer in general, but fortunately, some others have done the work for us. Specifically, we rely upon a study by Lu et al. [L+08], which analyzes a number of popular concurrent applications in great detail to understand what types of bugs arise in practice.

The study focuses on four major and important open-source applications: MySQL (a popular database management system), Apache (a well-known web server), Mozilla (the famous web browser), and OpenOffice (a free version of the MS Office suite, which some people actually use). In the study, the authors examine concurrency bugs that have been found and fixed in each of these code bases, turning the developers’ work into a quantitative bug analysis; understanding these results can help you understand what types of problems actually occur in mature code bases.
Table 32.1: Bugs In Modern Applications

Table 32.1 shows a summary of the bugs Lu and colleagues studied. From the table, you can see that there were 105 total bugs, most of which were not deadlock (74); the remaining 31 were deadlock bugs. Further, you can see that the number of bugs studied from each application; while OpenOffice only had 8 total concurrency bugs, Mozilla had nearly 60.

We now dive into these different classes of bugs (non-deadlock, deadlock) a bit more deeply. For the first class of non-deadlock bugs, we use examples from the study to drive our discussion. For the second class of deadlock bugs, we discuss the long line of work that has been done in either preventing, avoiding, or handling deadlock.

32.2 Non-Deadlock Bugs

Non-deadlock bugs make up a majority of concurrency bugs, according to Lu’s study. But what types of bugs are these? How do they arise? How can we fix them? We now discuss the two major types of non-deadlock bugs found by Lu et al.: atomicity violation bugs and order violation bugs.

Atomicity-Violation Bugs

The first type of problem encountered is referred to as an atomicity violation. Here is a simple example, found in MySQL. Before reading the explanation, try figuring out what the bug is. Do it!

```c
1 Thread 1::
2   if (thd->proc_info) {
3     ...
4     fputs(thd->proc_info, ...);
5     ...
6   }
7
8 Thread 2::
9   thd->proc_info = NULL;
```

In the example, two different threads access the field `proc_info` in the structure `thd`. The first thread checks if the value is non-NULL and then prints its value; the second thread sets it to NULL. Clearly, if the first thread performs the check but then is interrupted before the call to `fputs`, the second thread could run in-between, thus setting the pointer to NULL; when the first thread resumes, it will crash, as a NULL pointer will be dereferenced by `fputs`.
The more formal definition of an atomicity violation, according to Lu et al, is this: "The desired serializability among multiple memory accesses is violated (i.e. a code region is intended to be atomic, but the atomicity is not enforced during execution)." In our example above, the code has an atomicity assumption (in Lu’s words) about the check for non-NULL of proc_info and the usage of proc_info in the fputs() call; when assumption is broken, the code will not work as desired.

Finding a fix for this type of problem is often (but not always) straightforward. Can you think of how to fix the code above?

In this solution, we simply add locks around the shared-variable references, ensuring that when either thread accesses the proc_info field, it has a lock held. Of course (not shown), any other code that accesses the structure should also acquire this lock before doing so.

```
1 pthread_mutex_t lock = PTHREAD_MUTEX_INITIALIZER;
2
3 Thread 1::
4    pthread_mutex_lock(&lock);
5    if (thd->proc_info) {
6      ...
7      fputs(thd->proc_info, ...);
8      ...
9    }
10    pthread_mutex_unlock(&lock);
11
12 Thread 2::
13    pthread_mutex_lock(&lock);
14    thd->proc_info = NULL;
15    pthread_mutex_unlock(&lock);
```

Order-Violation Bugs

Another common type of non-deadlock bug found by Lu et al. is known as an order violation. Here is another simple example; once again, see if you can figure out why the code below has a bug in it.

```
1 Thread 1::
2    void init() {
3      ...
4      mThread = PR_CreateThread(mMain, ...);
5      ...
6    }
7
8 Thread 2::
9    void mMain(...) {
10      ...
11      mState = mThread->State;
12      ...
13    }
```

As you probably figured out, the code in Thread 2 seems to assume that the variable mThread has already been initialized (and is not NULL); however, if Thread 1 does not happen to run first, we are out of luck, and Thread 2 will likely crash with a NULL pointer dereference (assuming
that the value of mThread is initially NULL; if not, even stranger things could happen as arbitrary memory locations are read through the dereference in Thread 2).

The more formal definition of an order violation is this: “The desired order between two (groups of) memory accesses is flipped (i.e., A should always be executed before B, but the order is not enforced during execution).” [L+08]

The fix to this type of bug is generally to enforce ordering. As we discussed in detail previously, using condition variables is an easy and robust way to add this style of synchronization into modern code bases.

In the example above, we could thus rewrite the code as follows:

```c
pthread_mutex_t mtLock = PTHREAD_MUTEX_INITIALIZER;
pthread_cond_t mtCond = PTHREAD_COND_INITIALIZER;
int mtInit = 0;

Thread 1::
void init() {
    mThread = PR_CreateThread(mMain, ...);
    pthread_mutex_lock(&mtLock);
    mtInit = 1;
    pthread_cond_signal(&mtCond);
    pthread_mutex_unlock(&mtLock);
    ...
}

Thread 2::
void mMain(...) {
    // wait for the thread to be initialized...
    pthread_mutex_lock(&mtLock);
    while (mtInit == 0)
        pthread_cond_wait(&mtCond, &mtLock);
    pthread_mutex_unlock(&mtLock);
    mState = mThread->State;
    ...
}
```

In this fixed-up code sequence, we have added a lock (mtLock) and corresponding condition variable (mtCond), as well as a state variable (mtInit). When the initialization code runs, it sets the state of mtInit to 1 and signals that it has done so. If Thread 2 had run before this point, it will be waiting for this signal and corresponding state change; if it runs later, it will check the state and see that the initialization has already occurred (i.e., mtInit is set to 1), and thus continue as is proper. Note that we could likely use mThread as the state variable itself, but do not do so for the sake of simplicity here. When ordering matters between threads, condition variables (or semaphores) can come to the rescue.
Non-Deadlock Bugs: Summary

A large fraction (97%) of non-deadlock bugs studied by Lu et al. are either atomicity or order violations. Thus, by carefully thinking about these types of bug patterns, programmers can likely do a better job of avoiding them. Moreover, as more automated code-checking tools develop, they should likely focus on these two types of bugs as they constitute such a large fraction of non-deadlock bugs found in deployment.

Unfortunately, not all bugs are as easily fixable as the examples we looked at above. Some require a deeper understanding of what the program is doing, or a larger amount of code or data structure reorganization to fix. Read Lu et al.’s excellent (and readable) paper for more details.

32.3 Deadlock Bugs

Beyond the concurrency bugs mentioned above, a classic problem that arises in many concurrent systems with complex locking protocols is known as deadlock. Deadlock occurs, for example, when a thread (say Thread 1) is holding a lock (L1) and waiting for another one (L2); unfortunately, the thread (Thread 2) that holds lock L2 is waiting for L1 to be released.

Here is a code snippet that demonstrates such a potential deadlock:

Thread 1:
lock(L1);
lock(L2);
Thread 2:
lock(L2);
lock(L1);

Note that if this code runs, deadlock does not necessarily occur; rather, it may occur, if, for example, Thread 1 grabs lock L1 and then a context switch occurs to Thread 2. At that point, Thread 2 grabs L2, and tries to acquire L1. Thus we have a deadlock, as each thread is waiting for the other and neither can run. See Figure 32.1 for details; the presence of a cycle in the graph is indicative of the deadlock.

The figure should make clear the problem. How should programmers write code so as to handle deadlock in some way?

CRUX: HOW TO DEAL WITH DEADLOCK

How should we build systems to prevent, avoid, or at least detect and recover from deadlock? Is this a real problem in systems today?

Why Do Deadlocks Occur?

As you may be thinking, simple deadlocks such as the one above seem readily avoidable. For example, if Thread 1 and 2 both made sure to grab locks in the same order, the deadlock would never arise. So why do deadlocks happen?
One reason is that in large code bases, complex dependencies arise between components. Take the operating system, for example. The virtual memory system might need to access the file system in order to page in a block from disk; the file system might subsequently require a page of memory to read the block into and thus contact the virtual memory system. Thus, the design of locking strategies in large systems must be carefully done to avoid deadlock in the case of circular dependencies that may occur naturally in the code.

Another reason is due to the nature of encapsulation. As software developers, we are taught to hide details of implementations and thus make software easier to build in a modular way. Unfortunately, such modularity does not mesh well with locking. As Jula et al. point out [J+08], some seemingly innocuous interfaces almost invite you to deadlock. For example, take the Java Vector class and the method `AddAll()`. This routine would be called as follows:

```java
Vector v1, v2;
v1.addAll(v2);
```

Internally, because the method needs to be multi-thread safe, locks for both the vector being added to (v1) and the parameter (v2) need to be acquired. The routine acquires said locks in some arbitrary order (say v1 then v2) in order to add the contents of v2 to v1. If some other thread calls `v2.addAll(v1)` at nearly the same time, we have the potential for deadlock, all in a way that is quite hidden from the calling application.
Conditions for Deadlock

Four conditions need to hold for a deadlock to occur [C+71]:

- **Mutual exclusion**: Threads claim exclusive control of resources that they require (e.g., a thread grabs a lock).
- **Hold-and-wait**: Threads hold resources allocated to them (e.g., locks that they have already acquired) while waiting for additional resources (e.g., locks that they wish to acquire).
- **No preemption**: Resources (e.g., locks) cannot be forcibly removed from threads that are holding them.
- **Circular wait**: There exists a circular chain of threads such that each thread holds one more resources (e.g., locks) that are being requested by the next thread in the chain.

If any of these four conditions are not met, deadlock cannot occur. Thus, we first explore techniques to **prevent** deadlock; each of these strategies seeks to prevent one of the above conditions from arising and thus is one approach to handling the deadlock problem.

Prevention

Circular Wait

Probably the most practical prevention technique (and certainly one that is used frequently) is to write your locking code such that you never induce a circular wait. The way to do that is to provide a **total ordering** on lock acquisition. For example, if there are only two locks in the system (L1 and L2), we can prevent deadlock by always acquiring L1 before L2. Such strict ordering ensures that no cyclical wait arises; hence, no deadlock.

As you can imagine, this approach requires careful design of global locking strategies and must be done with great care. Further, it is just a convention, and a sloppy programmer can easily ignore the locking protocol and potentially cause deadlock. Finally, it requires a deep understanding of the code base, and how various routines are called; just one mistake could result in the wrong ordering of lock acquisition, and hence deadlock.

Hold-and-wait

The hold-and-wait requirement for deadlock can be avoided by acquiring all locks at once, atomically. In practice, this could be achieved as follows:

```cpp
1  lock(prevention);
2  lock(L1);
3  lock(L2);
4  ...
5  unlock(prevention);
```
By first grabbing the lock prevention, this code guarantees that no untimely thread switch can occur in the midst of lock acquisition and thus deadlock can once again be avoided. Of course, it requires that any time any thread grabs a lock, it first acquires the global prevention lock. For example, if another thread was trying to grab locks L1 and L2 in a different order, it would be OK, because it would be holding the prevention lock while doing so.

Note that the solution is problematic for a number of reasons. As before, encapsulation works against us: this approach requires us to know when calling a routine exactly which locks must be held and to acquire them ahead of time. Further, the approach likely decreases concurrency as all locks must be acquired early on (at once) instead of when they are truly needed.

No Preemption

Because we generally view locks as held until unlock is called, multiple lock acquisition often gets us into trouble because when waiting for one lock we are holding another. Many thread libraries provide a more flexible set of interfaces to help avoid this situation. Specifically, a trylock() routine will grab the lock (if it is available) or return -1 indicating that the lock is held right now and that you should try again later if you want to grab that lock.

Such an interface could be used as follows to build a deadlock-free, ordering-robust lock acquisition protocol:

```
1  top:
2     lock(L1);
3     if (trylock(L2) == -1) {
4         unlock(L1);
5     goto top;
6 }
```

Note that another thread could follow the same protocol but grab the locks in the other order (L2 then L1) and the program would still be deadlock free. One new problem does arise, however: livelock. It is possible (though perhaps unlikely) that two threads could both be repeatedly attempting this sequence and repeatedly failing to acquire both locks. In this case, both systems are running through this code sequence over and over again (and thus it is not a deadlock), but progress is not being made, hence the name livelock. There are solutions to the livelock problem, too: for example, one could add a random delay before looping back and trying the entire thing over again, thus decreasing the odds of repeated interference among competing threads.

One final point about this solution: it skirts around the hard parts of using a trylock approach. The first problem that would likely exist again arises due to encapsulation: if one of these locks is buried in some routine that is getting called, the jump back to the beginning becomes more complex to implement. If the code had acquired some resources (other than
L1) along the way, it must make sure to carefully release them as well; for example, if after acquiring L1, the code had allocated some memory, it would have to release that memory upon failure to acquire L2, before jumping back to the top to try the entire sequence again. However, in limited circumstances (e.g., the Java vector method above), this type of approach could work well.

**Mutual Exclusion**

The final prevention technique would be to avoid the need for mutual exclusion at all. In general, we know this is difficult, because the code we wish to run does indeed have critical sections. So what can we do?

Herlihy had the idea that one could design various data structures to be wait-free [H91]. The idea here is simple: using powerful hardware instructions, you can build data structures in a manner that does not require explicit locking.

As a simple example, let us assume we have a compare-and-swap instruction, which as you may recall is an atomic instruction provided by the hardware that does the following:

```c
1 int CompareAndSwap(int *address, int expected, int new) {
2   if (*address == expected) {
3     *address = new;
4     return 1; // success
5   }
6   return 0; // failure
7 }
```

Imagine we now wanted to atomically increment a value by a certain amount. We could do it as follows:

```c
1 void AtomicIncrement(int *value, int amount) {
2   do {
3     int old = *value;
4   } while (CompareAndSwap(value, old, old + amount) == 0);
5 }
```

Instead of acquiring a lock, doing the update, and then releasing it, we have instead built an approach that repeatedly tries to update the value to the new amount and uses the compare-and-swap to do so. In this manner, no lock is acquired, and no deadlock can arise (though livelock is still a possibility).

Let us consider a slightly more complex example: list insertion. Here is code that inserts at the head of a list:

```c
1 void insert(int value) {
2   node_t *n = malloc(sizeof(node_t));
3   assert(n != NULL);
4   n->value = value;
5   n->next = head;
6   head = n;
7 }
```
This code performs a simple insertion, but if called by multiple threads at the “same time”, has a race condition (see if you can figure out why). Of course, we could solve this by surrounding this code with a lock acquire and release:

```c
void insert(int value) {
  node_t *n = malloc(sizeof(node_t));
  assert(n != NULL);
  n->value = value;
  lock(listlock); // begin critical section
  n->next = head;
  head = n;
  unlock(listlock); // end of critical section
}
```

In this solution, we are using locks in the traditional manner. Instead, let us try to perform this insertion in a wait-free manner simply using the compare-and-swap instruction. Here is one possible approach:

```c
void insert(int value) {
  node_t *n = malloc(sizeof(node_t));
  assert(n != NULL);
  n->value = value;
  do {
    n->next = head;
  } while (CompareAndSwap(&head, n->next, n));
}
```

The code here updates the next pointer to point to the current head, and then tries to swap the newly-created node into position as the new head of the list. However, this will fail if some other thread successfully swapped in a new head in the meanwhile, causing this thread to retry again with the new head.

Of course, building a useful list requires more than just a list insert, and not surprisingly building a list that you can insert into, delete from, and perform lookups on in a wait-free manner is non-trivial. Read the rich literature on wait-free synchronization if you find this interesting.

### Deadlock Avoidance via Scheduling

Instead of deadlock prevention, in some scenarios deadlock avoidance is preferable. Avoidance requires some global knowledge of which locks various threads might grab during their execution, and subsequently schedules said threads in a way as to guarantee no deadlock can occur.

For example, assume we have two processors and four threads which must be scheduled upon them. Assume further we know that Thread 1 (T1) grabs locks L1 and L2 (in some order, at some point during its execution), T2 grabs L1 and L2 as well, T3 grabs just L2, and T4 grabs no

---

1The astute reader might be asking why we grabbed the lock so late, instead of right when entering the `insert()` routine; can you, astute reader, figure out why that is likely OK?
locks at all. We can show these lock acquisition demands of the threads in tabular form:

<table>
<thead>
<tr>
<th></th>
<th>T1</th>
<th>T2</th>
<th>T3</th>
<th>T4</th>
</tr>
</thead>
<tbody>
<tr>
<td>L1</td>
<td>yes</td>
<td>yes</td>
<td>no</td>
<td>no</td>
</tr>
<tr>
<td>L2</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>no</td>
</tr>
</tbody>
</table>

A smart scheduler could thus compute that as long as T1 and T2 are not run at the same time, no deadlock could ever arise. Here is one such schedule:

CPU 1: T3 T4
CPU 2: T1 T2

Note that it is OK for (T3 and T1) or (T3 and T2) to overlap. Even though T3 grabs lock L2, it can never cause a deadlock by running concurrently with other threads because it only grabs one lock.

Let’s look at one more example. In this one, there is more contention for the same resources (again, locks L1 and L2), as indicated by the following contention table:

<table>
<thead>
<tr>
<th></th>
<th>T1</th>
<th>T2</th>
<th>T3</th>
<th>T4</th>
</tr>
</thead>
<tbody>
<tr>
<td>L1</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>no</td>
</tr>
<tr>
<td>L2</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>no</td>
</tr>
</tbody>
</table>

In particular, threads T1, T2, and T3 all need to grab both locks L1 and L2 at some point during their execution. Here is a possible schedule that guarantees that no deadlock could ever occur:

CPU 1: T4
CPU 2: T1 T2 T3

As you can see, static scheduling leads to a conservative approach where T1, T2, and T3 are all run on the same processor, and thus the total time to complete the jobs is lengthened considerably. Though it may have been possible to run these tasks concurrently, the fear of deadlock prevents us from doing so, and the cost is performance.

One famous example of an approach like this is Dijkstra’s Banker’s Algorithm [D64], and many similar approaches have been described in the literature. Unfortunately, they are only useful in very limited environments, for example, in an embedded system where one has full knowledge of the entire set of tasks that must be run and the locks that they need. Further, such approaches can limit concurrency, as we saw in the second example above. Thus, avoidance of deadlock via scheduling is not a widely-used general-purpose solution.
TIP: DON’T ALWAYS DO IT PERFECTLY (TOM WEST’S LAW)
Tom West, famous as the subject of the classic computer-industry book “Soul of a New Machine” [K81], says famously: “Not everything worth doing is worth doing well”, which is a terrific engineering maxim. If a bad thing happens rarely, certainly one should not spend a great deal of effort to prevent it, particularly if the cost of the bad thing occurring is small.

Detect and Recover
One final general strategy is to allow deadlocks to occasionally occur, and then take some action once such a deadlock has been detected. For example, if an OS froze once a year, you would just reboot it and get happily (or grumpily) on with your work. If deadlocks are rare, such a non-solution is indeed quite pragmatic.

Many database systems employ deadlock detection and recovery techniques. A deadlock detector runs periodically, building a resource graph and checking it for cycles. In the event of a cycle (deadlock), the system needs to be restarted. If more intricate repair of data structures is first required, a human being may be involved to ease the process.

32.4 Summary
In this chapter, we have studied the types of bugs that occur in concurrent programs. The first type, non-deadlock bugs, are surprisingly common, but often are easier to fix. They include atomicity violations, in which a sequence of instructions that should have been executed together was not, and order violations, in which the needed order between two threads was not enforced.

We have also briefly discussed deadlock: why it occurs, and what can be done about it. The problem is as old as concurrency itself, and many hundreds of papers have been written about the topic. The best solution in practice is to be careful, develop a lock acquisition total order, and thus prevent deadlock from occurring in the first place. Wait-free approaches also have promise, as some wait-free data structures are now finding their way into commonly-used libraries and critical systems, including Linux. However, their lack of generality and the complexity to develop a new wait-free data structure will likely limit the overall utility of this approach. Perhaps the best solution is to develop new concurrent programming models: in systems such as MapReduce (from Google) [GD02], programmers can describe certain types of parallel computations without any locks whatsoever. Locks are problematic by their very nature; perhaps we should seek to avoid using them unless we truly must.
References

[C+71] “System Deadlocks”
E.G. Coffman, M.J. Elphick, A. Shoshani
ACM Computing Surveys, 3:2, June 1971
The classic paper outlining the conditions for deadlock and how you might go about dealing with it. There are certainly some earlier papers on this topic; see the references within this paper for details.

[D64] “Een algorithme ter voorkoming van de dodelijke omarming”
Circulated privately, around 1964
Available: http://www.cs.utexas.edu/users/EWD/ewd01xx/EWD108.PDF
Indeed, not only did Dijkstra come up with a number of solutions to the deadlock problem, he was the first to note its existence, at least in written form. However, he called it the “deadly embrace”, which (thankfully) did not catch on.

[GD02] “MapReduce: Simplified Data Processing on Large Clusters”
Sanjay Ghemawat and Jeff Dean
OSDI ’04, San Francisco, CA, October 2004
The MapReduce paper ushered in the era of large-scale data processing, and proposes a framework for performing such computations on clusters of generally unreliable machines.

[H91] “Wait-free Synchronization”
Maurice Herlihy
ACM TOPLAS, 13(1), pages 124-149, January 1991
Herlihy’s work pioneers the ideas behind wait-free approaches to writing concurrent programs. These approaches tend to be complex and hard, often more difficult than using locks correctly, probably limiting their success in the real world.

[J+08] “Deadlock Immunity: Enabling Systems To Defend Against Deadlocks”
Horatiu Jula, Daniel Tralamazza, Cristian Zamfir, George Candea
OSDI ’08, San Diego, CA, December 2008
An excellent recent paper on deadlocks and how to avoid getting caught in the same ones over and over again in a particular system.

[K81] “Soul of a New Machine”
Tracy Kidder, 1980
A must-read for any systems builder or engineer, detailing the early days of how a team inside Data General (DG), led by Tom West, worked to produce a “new machine.” Kidder’s other book are also excellent, in particular, “Mountains beyond Mountains”. Or maybe you don’t agree with me, comma?

[L+08] “Learning from Mistakes – A Comprehensive Study on Real World Concurrency Bug Characteristics”
Shan Lu, Soyeon Park, Eunsoo Seo, Yuanyuan Zhou
ASPLOS ’08, March 2008, Seattle, Washington
The first in-depth study of concurrency bugs in real software, and the basis for this chapter. Look at Y.Y. Zhou’s or Shan Lu’s web pages for many more interesting papers on bugs.
Thus far, we’ve written about concurrency as if the only way to build concurrent applications is to use threads. Like many things in life, this is not completely true. Specifically, a different style of concurrent programming is often used in both GUI-based applications [O96] as well as some types of internet servers [PDZ99]. This style, known as event-based concurrency, has become popular in some modern systems, including server-side frameworks such as node.js [N13], but its roots are found in C/UNIX systems that we’ll discuss below.

The problem that event-based concurrency addresses is two-fold. The first is that managing concurrency correctly in multi-threaded applications can be challenging; as we’ve discussed, missing locks, deadlock, and other nasty problems can arise. The second is that in a multi-threaded application, the developer has little or no control over what is scheduled at a given moment in time; rather, the programmer simply creates threads and then hopes that the underlying OS schedules them in a reasonable manner across available CPUs. Given the difficulty of building a general-purpose scheduler that works well in all cases for all workloads, sometimes the OS will schedule work in a manner that is less than optimal.

The crux:

**The Crux:**

**How To Build Concurrent Servers Without Threads**

How can we build a concurrent server without using threads, and thus retain control over concurrency as well as avoid some of the problems that seem to plague multi-threaded applications?

33.1 The Basic Idea: An Event Loop

The basic approach we’ll use, as stated above, is called event-based concurrency. The approach is quite simple: you simply wait for something (i.e., an “event”) to occur; when it does, you check what type of
event it is and do the small amount of work it requires (which may include issuing I/O requests, or scheduling other events for future handling, etc.). That’s it!

Before getting into the details, let’s first examine what a canonical event-based server looks like. Such applications are based around a simple construct known as the event loop. Pseudocode for an event loop looks like this:

```c
while (1) {
    events = getEvents();
    for (e in events)
        processEvent(e);
}
```

It’s really that simple. The main loop simply waits for something to do (by calling `getEvents()` in the code above) and then, for each event returned, processes them, one at a time; the code that processes each event is known as an event handler. Importantly, when a handler processes an event, it is the only activity taking place in the system; thus, deciding which event to handle next is equivalent to scheduling. This explicit control over scheduling is one of the fundamental advantages of the event-based approach.

But this discussion leaves us with a bigger question: how exactly does an event-based server determine which events are taking place, in particular with regards to network and disk I/O? Specifically, how can an event server tell if a message has arrived for it?

### 33.2 An Important API: `select()` (or `poll()`)

With that basic event loop in mind, we next must address the question of how to receive events. In most systems, a basic API is available, via either the `select()` or `poll()` system calls.

What these interfaces enable a program to do is simple: check whether there is any incoming I/O that should be attended to. For example, imagine that a network application (such as a web server) wishes to check whether any network packets have arrived, in order to service them. These system calls let you do exactly that.

Take `select()` for example. The manual page (on Mac OS X) describes the API in this manner:

```c
int select(int nfds,
           fd_set *restrict readfds,
           fd_set *restrict writefds,
           fd_set *restrict errorfds,
           struct timeval *restrict timeout);
```

The actual description from the man page: `select()` examines the I/O descriptor sets whose addresses are passed in readfds, writefds, and errorfds to see if some of their descriptors are ready for reading, are ready for writing, or have
ASIDE: Blocking vs. Non-blocking Interfaces

Blocking (or synchronous) interfaces do all of their work before returning to the caller; non-blocking (or asynchronous) interfaces begin some work but return immediately, thus letting whatever work that needs to be done get done in the background.

The usual culprit in blocking calls is I/O of some kind. For example, if a call must read from disk in order to complete, it might block, waiting for the I/O request that has been sent to the disk to return.

Non-blocking interfaces can be used in any style of programming (e.g., with threads), but are essential in the event-based approach, as a call that blocks will halt all progress.

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Non-blocking interfaces can be used in any style of programming (e.g., with threads), but are essential in the event-based approach, as a call that blocks will halt all progress.
```c
#include <stdio.h>
#include <stdlib.h>
#include <sys/time.h>
#include <sys/types.h>
#include <unistd.h>

int main(void) {
    // open and set up a bunch of sockets (not shown)
    // main loop
    while (1) {
        // initialize the fd_set to all zero
        fd_set readFDs;
        FD_ZERO(&readFDs);
        // now set the bits for the descriptors
        // this server is interested in
        // (for simplicity, all of them from min to max)
        int fd;
        for (fd = minFD; fd < maxFD; fd++)
            FD_SET(fd, &readFDs);
        // do the select
        int rc = select(maxFD+1, &readFDs, NULL, NULL, NULL);
        // check which actually have data using FD_ISSET()
        int fd;
        for (fd = minFD; fd < maxFD; fd++)
            if (FD_ISSET(fd, &readFDs))
                processFD(fd);
    }

Figure 33.1: Simple Code using select()
33.5 A Problem: Blocking System Calls

Thus far, event-based programming sounds great, right? You program a simple loop, and handle events as they arise. You don’t even need to think about locking! But there is an issue: what if an event requires that you issue a system call that might block?

For example, imagine a request comes from a client into a server to read a file from disk and return its contents to the requesting client (much like a simple HTTP request). To service such a request, some event handler will eventually have to issue an `open()` system call to open the file, followed by a series of `read()` calls to read the file. When the file is read into memory, the server will likely start sending the results to the client.

Both the `open()` and `read()` calls may issue I/O requests to the storage system (when the needed metadata or data is not in memory already), and thus may take a long time to service. With a thread-based server, this is no issue: while the thread issuing the I/O request suspends (waiting for the I/O to complete), other threads can run, thus enabling the server to make progress. Indeed, this natural overlap of I/O and other computation is what makes thread-based programming quite natural and straightforward.

With an event-based approach, however, there are no other threads to run: just the main event loop. And this implies that if an event handler issues a call that blocks, the entire server will do just that: block until the call completes. When the event loop blocks, the system sits idle, and thus is a huge potential waste of resources. We thus have a rule that must be obeyed in event-based systems: no blocking calls are allowed.

33.6 A Solution: Asynchronous I/O

To overcome this limit, many modern operating systems have introduced new ways to issue I/O requests to the disk system, referred to generically as asynchronous I/O. These interfaces enable an application to issue an I/O request and return control immediately to the caller, before the I/O has completed; additional interfaces enable an application to determine whether various I/Os have completed.

For example, let us examine the interface provided on Mac OS X (other systems have similar APIs). The APIs revolve around a basic structure,
the **struct aiocb** or **AIO control block** in common terminology. A simplified version of the structure looks like this (see the manual pages for more information):

```c
struct aiocb {
    int  aio_fildes;  /* File descriptor */
    off_t aio_offset; /* File offset */
    volatile void *aio_buf; /* Location of buffer */
    size_t aio_nbytes; /* Length of transfer */
};
```

To issue an asynchronous read to a file, an application should first fill in this structure with the relevant information: the file descriptor of the file to be read (**aio_fildes**), the offset within the file (**aio_offset**) as well as the length of the request (**aio_nbytes**), and finally the target memory location into which the results of the read should be copied (**aio_buf**).

After this structure is filled in, the application must issue the asynchronous call to read the file; on Mac OS X, this API is simply the **asynchronous read** API:

```c
int aio_read(struct aiocb *aiocbp);
```

This call tries to issue the I/O; if successful, it simply returns right away and the application (i.e., the event-based server) can continue with its work.

There is one last piece of the puzzle we must solve, however. How can we tell when an I/O is complete, and thus that the buffer (pointed to by **aio_buf**) now has the requested data within it?

One last API is needed. On Mac OS X, it is referred to (somewhat confusingly) as **aio_error()**. The API looks like this:

```c
int aio_error(const struct aiocb *aiocbp);
```

This system call checks whether the request referred to by **aiocbp** has completed. If it has, the routine returns success (indicated by a zero); if not, EINPROGRESS is returned. Thus, for every outstanding asynchronous I/O, an application can periodically poll the system via a call to **aio_error()** to determine whether said I/O has yet completed.

One thing you might have noticed is that it is painful to check whether an I/O has completed; if a program has tens or hundreds of I/Os issued at a given point in time, should it simply keep checking each of them repeatedly, or wait a little while first, or ... ?

To remedy this issue, some systems provide an approach based on the **interrupt**. This method uses **UNIX signals** to inform applications when an asynchronous I/O completes, thus removing the need to repeatedly ask the system. This polling vs. interrupts issue is seen in devices too, as you will see (or already have seen) in the chapter on I/O devices.
**ASIDE: UNIX SIGNALS**

A huge and fascinating infrastructure known as signals is present in all modern UNIX variants. At its simplest, signals provide a way to communicate with a process. Specifically, a signal can be delivered to an application; doing so stops the application from whatever it is doing to run a **signal handler**, i.e., some code in the application to handle that signal. When finished, the process just resumes its previous behavior.

Each signal has a name, such as **HUP** (hang up), **INT** (interrupt), **SEGV** (segmentation violation), etc; see the manual page for details. Interestingly, sometimes it is the kernel itself that does the signaling. For example, when your program encounters a segmentation violation, the OS sends it a **SIGSEGV** (prepending SIG to signal names is common); if your program is configured to catch that signal, you can actually run some code in response to this erroneous program behavior (which can be useful for debugging). When a signal is sent to a process not configured to handle that signal, some default behavior is enacted; for SEGV, the process is killed.

Here is a simple program that goes into an infinite loop, but has first set up a signal handler to catch SIGHUP:

```c
#include <stdio.h>
#include <signal.h>

void handle(int arg) {
    printf("stop wakin' me up...\n");
}

int main(int argc, char *argv[]) {
    signal(SIGHUP, handle);
    while (1)
        ; // doin' nothin' except catchin' some sigs
    return 0;
}
```

You can send signals to it with the `kill` command line tool (yes, this is an odd and aggressive name). Doing so will interrupt the main while loop in the program and run the handler code `handle()`:

```
prompt> ./main &
[3] 36705
prompt> kill -HUP 36705
stop wakin' me up...
prompt> kill -HUP 36705
stop wakin' me up...
prompt> kill -HUP 36705
stop wakin' me up...
prompt> kill -HUP 36705
stop wakin' me up...
```

There is a lot more to learn about signals, so much that a single page, much less a single chapter, does not nearly suffice. As always, there is one great source: Stevens and Rago [SR05]. Read more if interested.
In systems without asynchronous I/O, the pure event-based approach cannot be implemented. However, clever researchers have derived methods that work fairly well in their place. For example, Pai et al. [PDZ99] describe a hybrid approach in which events are used to process network packets, and a thread pool is used to manage outstanding I/Os. Read their paper for details.

33.7 Another Problem: State Management

Another issue with the event-based approach is that such code is generally more complicated to write than traditional thread-based code. The reason is as follows: when an event handler issues an asynchronous I/O, it must package up some program state for the next event handler to use when the I/O finally completes; this additional work is not needed in thread-based programs, as the state the program needs is on the stack of the thread. Adya et al. call this work manual stack management, and it is fundamental to event-based programming [A+02].

To make this point more concrete, let’s look at a simple example in which a thread-based server needs to read from a file descriptor (fd) and, once complete, write the data that it read from the file to a network socket descriptor (sd). The code (ignoring error checking) looks like this:

```c
int rc = read(fd, buffer, size);
rc = write(sd, buffer, size);
```

As you can see, in a multi-threaded program, doing this kind of work is trivial; when the read() finally returns, the code immediately knows which socket to write to because that information is on the stack of the thread (in the variable sd).

In an event-based system, life is not so easy. To perform the same task, we’d first issue the read asynchronously, using the AIO calls described above. Let’s say we then periodically check for completion of the read using the aio_error() call; when that call informs us that the read is complete, how does the event-based server know what to do?

The solution, as described by Adya et al. [A+02], is to use an old programming language construct known as a continuation [FHK84]. Though it sounds complicated, the idea is rather simple: basically, record the needed information to finish processing this event in some data structure; when the event happens (i.e., when the disk I/O completes), look up the needed information and process the event.

In this specific case, the solution would be to record the socket descriptor (sd) in some kind of data structure (e.g., a hash table), indexed by the file descriptor (fd). When the disk I/O completes, the event handler would use the file descriptor to look up the continuation, which will return the value of the socket descriptor to the caller. At this point (finally), the server can then do the last bit of work to write the data to the socket.
33.8 What Is Still Difficult With Events

There are a few other difficulties with the event-based approach that we should mention. For example, when systems moved from a single CPU to multiple CPUs, some of the simplicity of the event-based approach disappeared. Specifically, in order to utilize more than one CPU, the event server has to run multiple event handlers in parallel; when doing so, the usual synchronization problems (e.g., critical sections) arise, and the usual solutions (e.g., locks) must be employed. Thus, on modern multicore systems, simple event handling without locks is no longer possible.

Another problem with the event-based approach is that it does not integrate well with certain kinds of systems activity, such as paging. For example, if an event-handler page faults, it will block, and thus the server will not make progress until the page fault completes. Even though the server has been structured to avoid explicit blocking, this type of implicit blocking due to page faults is hard to avoid and thus can lead to large performance problems when prevalent.

A third issue is that event-based code can be hard to manage over time, as the exact semantics of various routines changes [A+02]. For example, if a routine changes from non-blocking to blocking, the event handler that calls that routine must also change to accommodate its new nature, by ripping itself into two pieces. Because blocking is so disastrous for event-based servers, a programmer must always be on the lookout for such changes in the semantics of the APIs each event uses.

Finally, though asynchronous disk I/O is now possible on most platforms, it has taken a long time to get there [PDZ99], and it never quite integrates with asynchronous network I/O in as simple and uniform a manner as you might think. For example, while one would simply like to use the select() interface to manage all outstanding I/Os, usually some combination of select() for networking and the AIO calls for disk I/O are required.

33.9 Summary

We’ve presented a bare bones introduction to a different style of concurrency based on events. Event-based servers give control of scheduling to the application itself, but do so at some cost in complexity and difficulty of integration with other aspects of modern systems (e.g., paging). Because of these challenges, no single approach has emerged as best; thus, both threads and events are likely to persist as two different approaches to the same concurrency problem for many years to come. Read some research papers (e.g., [A+02, PDZ99, vB+03, WCB01]) or better yet, write some event-based code, to learn more.
References

Atul Adya, Jon Howell, Marvin Theimer, William J. Bolosky, John R. Douceur
USENIX ATC ’02, Monterey, CA, June 2002
This gem of a paper is the first to clearly articulate some of the difficulties of event-based concurrency,
and suggests some simple solutions, as well explores the even crazier idea of combining the two types of
concurrency management into a single application!

[FHK84] “Programming With Continuations”
Daniel P. Friedman, Christopher T. Haynes, Eugene E. Kohlbecker
In Program Transformation and Programming Environments, Springer Verlag, 1984
The classic reference to this old idea from the world of programming languages. Now increasingly
popular in some modern languages.

[N13] “Node.js Documentation”
By the folks who build node.js
Available: http://nodejs.org/api/
One of the many cool new frameworks that help you readily build web services and applications. Every
modern systems hacker should be proficient in frameworks such as this one (and likely, more than one).
Spend the time and do some development in one of these worlds and become an expert.

[O96] “Why Threads Are A Bad Idea (for most purposes)”
John Ousterhout
Invited Talk at USENIX ‘96, San Diego, CA, January 1996
A great talk about how threads aren’t a great match for GUI-based applications (but the ideas are more
general). Ousterhout formed many of these opinions while he was developing Tcl/Tk, a cool scripting
language and toolkit that made it 100x easier to develop GUI-based applications than the state of the
art at the time. While the Tk GUI toolkit lives on (in Python for example), Tcl seems to be slowly dying
(unfortunately).

[PDZ99] “Flash: An Efficient and Portable Web Server”
Vivek S. Pai, Peter Druschel, Willy Zwaenepoel
USENIX ’99, Monterey, CA, June 1999
A pioneering paper on how to structure web servers in the then-burgeoning Internet era. Read it to
understand the basics as well as to see the authors’ ideas on how to build hybrids when support for
asynchronous I/O is lacking.

[SR05] “Advanced Programming in the UNIX Environment”
W. Richard Stevens and Stephen A. Rago
Addison-Wesley, 2005
Once again, we refer to the classic must-have-on-your-bookshelf book of UNIX systems programming.
If there is some detail you need to know, it is in here.

[vB+03] “Capriccio: Scalable Threads for Internet Services”
Rob von Behren, Jeremy Condit, Feng Zhou, George C. Necula, Eric Brewer
SOSP ’03, Lake George, New York, October 2003
A paper about how to make threads work at extreme scale; a counter to all the event-based work ongoing
at the time.

[WCB01] “SEDA: An Architecture for Well-Conditioned, Scalable Internet Services”
Matt Welsh, David Culler, and Eric Brewer
SOSP ’01, Banff, Canada, October 2001
A nice twist on event-based serving that combines threads, queues, and event-based handling into one
streamlined whole. Some of these ideas have found their way into the infrastructures of companies such
as Google, Amazon, and elsewhere.
Summary Dialogue on Concurrency

**Professor:** So, does your head hurt now?

**Student:** (taking two Motrin tablets) Well, some. It’s hard to think about all the ways threads can interleave.

**Professor:** Indeed it is. I am always amazed at how so few line of code, when concurrent execution is involved, can become nearly impossible to understand.

**Student:** Me too! It’s kind of embarrassing, as a Computer Scientist, not to be able to make sense of five lines of code.

**Professor:** Oh, don’t feel too badly. If you look through the first papers on concurrent algorithms, they are sometimes wrong! And the authors often professors!

**Student:** (gasps) Professors can be ... umm... wrong?

**Professor:** Yes, it is true. Though don’t tell anybody – it’s one of our trade secrets.

**Student:** I am sworn to secrecy. Though don’t tell anybody – it’s one of our trade secrets.

**Professor:** Well that is the real question, isn’t it? I think it starts with a few simple things. First, keep it simple! Avoid complex interactions between threads, and use well-known and tried-and-true ways to manage thread interactions.

**Student:** Like simple locking, and maybe a producer-consumer queue?

**Professor:** Exactly! Those are common paradigms, and you should be able to produce the working solutions given what you’ve learned. Second, only use concurrency when absolutely needed; avoid it if at all possible. There is nothing worse than premature optimization of a program.

**Student:** I see – why add threads if you don’t need them?

**Professor:** Exactly. Third, if you really need parallelism, seek it in other simplified forms. For example, the Map-Reduce method for writing parallel data analysis code is an excellent example of achieving parallelism without having to handle any of the horrific complexities of locks, condition variables, and the other nasty things we’ve talked about.
Student: Map-Reduce, huh? Sounds interesting – I’ll have to read more about it on my own.

Professor: Good! You should. In the end, you’ll have to do a lot of that, as what we learn together can only serve as the barest introduction to the wealth of knowledge that is out there. Read, read, and read some more! And then try things out, write some code, and then write some more too. As Gladwell talks about in his book “Outliers”, you need to put roughly 10,000 hours into something in order to become a real expert. You can’t do that all inside of class time!

Student: Wow, I’m not sure if that is depressing, or uplifting. But I’ll assume the latter, and get to work! Time to write some more concurrent code...
Part III

Persistence
A Dialogue on Persistence

Professor: And thus we reach the third of our four ... err... three pillars of operating systems: persistence.

Student: Did you say there were three pillars, or four? What is the fourth?

Professor: No. Just three, young student, just three. Trying to keep it simple here.

Student: OK, fine. But what is persistence, oh fine and noble professor?

Professor: Actually, you probably know what it means in the traditional sense, right? As the dictionary would say: “a firm or obstinate continuance in a course of action in spite of difficulty or opposition.”

Student: It’s kind of like taking your class: some obstinance required.

Professor: Ha! Yes. But persistence here means something else. Let me explain.

Imagine you are outside, in a field, and you pick a –

Student: (interrupting) I know! A peach! From a peach tree!

Professor: I was going to say apple, from an apple tree. Oh well; we’ll do it your way, I guess.

Student: (stares blankly)

Professor: Anyhow, you pick a peach; in fact, you pick many many peaches, but you want to make them last for a long time. Winter is hard and cruel in Wisconsin, after all. What do you do?

Student: Well, I think there are some different things you can do. You can pickle it! Or bake a pie. Or make a jam of some kind. Lots of fun!

Professor: Fun? Well, maybe. Certainly, you have to do a lot more work to make the peach persist. And so it is with information as well; making information persist, despite computer crashes, disk failures, or power outages is a tough and interesting challenge.

Student: Nice segue; you’re getting quite good at that.

Professor: Thanks! A professor can always use a few kind words, you know.
**Student:** I’ll try to remember that. I guess it’s time to stop talking peaches, and start talking computers?

**Professor:** Yes, it is that time...
Before delving into the main content of this part of the book (on persistence), we first introduce the concept of an input/output (I/O) device and show how the operating system might interact with such an entity. I/O is quite critical to computer systems, of course; imagine a program without any input (it produces the same result each time); now imagine a program with no output (what was the purpose of it running?). Clearly, for computer systems to be interesting, both input and output are required. And thus, our general problem:

**Crux: How To Integrate I/O Into Systems**

How should I/O be integrated into systems? What are the general mechanisms? How can we make them efficient?

### 36.1 System Architecture

To begin our discussion, let’s look at the structure of a typical system (Figure 36.1). The picture shows a single CPU attached to the main memory of the system via some kind of memory bus or interconnect. Some devices are connected to the system via a general I/O bus, which in many modern systems would be PCI (or one of its many derivatives); graphics and some other higher-performance I/O devices might be found here. Finally, even lower down are one or more of what we call a peripheral bus, such as **SCSI**, **SATA**, or **USB**. These connect the slowest devices to the system, including disks, mice, and other similar components.

One question you might ask is: why do we need a hierarchical structure like this? Put simply: physics, and cost. The faster a bus is, the shorter it must be; thus, a high-performance memory bus does not have much room to plug devices and such into it. In addition, engineering a bus for high performance is quite costly. Thus, system designers have adopted this hierarchical approach, where components that demand high performance (such as the graphics card) are nearer the CPU. Lower per-
formance components are further away. The benefits of placing disks and other slow devices on a peripheral bus are manifold; in particular, you can place a large number of devices on it.

36.2 A Canonical Device

Let us now look at a canonical device (not a real one), and use this device to drive our understanding of some of the machinery required to make device interaction efficient. From Figure 36.2, we can see that a device has two important components. The first is the hardware interface it presents to the rest of the system. Just like a piece of software, hardware must also present some kind of interface that allows the system software to control its operation. Thus, all devices have some specified interface and protocol for typical interaction.

The second part of any device is its internal structure. This part of the device is implementation specific and is responsible for implementing the abstraction the device presents to the system. Very simple devices will have one or a few hardware chips to implement their functionality; more complex devices will include a simple CPU, some general purpose memory, and other device-specific chips to get their job done. For example, modern RAID controllers might consist of hundreds of thousands of lines of firmware (i.e., software within a hardware device) to implement its functionality.
### 36.3 The Canonical Protocol

In the picture above, the (simplified) device interface is comprised of three registers: a **status** register, which can be read to see the current status of the device; a **command** register, to tell the device to perform a certain task; and a **data** register to pass data to the device, or get data from the device. By reading and writing these registers, the operating system can control device behavior.

Let us now describe a typical interaction that the OS might have with the device in order to get the device to do something on its behalf. The protocol is as follows:

```plaintext
While (STATUS == BUSY)
    ; // wait until device is not busy
Write data to DATA register
Write command to COMMAND register
    (Doing so starts the device and executes the command)
While (STATUS == BUSY)
    ; // wait until device is done with your request
```

The protocol has four steps. In the first, the OS waits until the device is ready to receive a command by repeatedly reading the status register; we call this polling the device (basically, just asking it what is going on). Second, the OS sends some data down to the data register; one can imagine that if this were a disk, for example, that multiple writes would need to take place to transfer a disk block (say 4KB) to the device. When the main CPU is involved with the data movement (as in this example protocol), we refer to it as **programmed I/O (PIO)**. Third, the OS writes a command to the command register; doing so implicitly lets the device know that both the data is present and that it should begin working on the command. Finally, the OS waits for the device to finish by again polling it in a loop, waiting to see if it is finished (it may then get an error code to indicate success or failure).

This basic protocol has the positive aspect of being simple and working. However, there are some inefficiencies and inconveniences involved. The first problem you might notice in the protocol is that polling seems inefficient; specifically, it wastes a great deal of CPU time just waiting for the (potentially slow) device to complete its activity, instead of switching to another ready process and thus better utilizing the CPU.
36.4 Lowering CPU Overhead With Interrupts

The invention that many engineers came upon years ago to improve this interaction is something we’ve seen already: the interrupt. Instead of polling the device repeatedly, the OS can issue a request, put the calling process to sleep, and context switch to another task. When the device is finally finished with the operation, it will raise a hardware interrupt, causing the CPU to jump into the OS at a pre-determined interrupt service routine (ISR) or more simply an interrupt handler. The handler is just a piece of operating system code that will finish the request (for example, by reading data and perhaps an error code from the device) and wake the process waiting for the I/O, which can then proceed as desired.

Interrupts thus allow for overlap of computation and I/O, which is key for improved utilization. This timeline shows the problem:

```
CPU: 1 1 1 1 1 p p p p 1 1 1 1
Disk: 1 1 1 1
```

In the diagram, Process 1 runs on the CPU for some time (indicated by a repeated 1 on the CPU line), and then issues an I/O request to the disk to read some data. Without interrupts, the system simply spins, polling the status of the device repeatedly until the I/O is complete (indicated by a p). The disk services the request and finally Process 1 can run again.

If instead we utilize interrupts and allow for overlap, the OS can do something else while waiting for the disk:

```
CPU: 1 1 1 1 1 2 2 2 2 1 1 1 1
Disk: 1 1 1 1
```

In this example, the OS runs Process 2 on the CPU while the disk services Process 1’s request. When the disk request is finished, an interrupt occurs, and the OS wakes up Process 1 and runs it again. Thus, both the CPU and the disk are properly utilized during the middle stretch of time.

Note that using interrupts is not always the best solution. For example, imagine a device that performs its tasks very quickly: the first poll usually finds the device to be done with task. Using an interrupt in this case will actually slow down the system: switching to another process, handling the interrupt, and switching back to the issuing process is expensive. Thus, if a device is fast, it may be best to poll; if it is slow, interrupts, which allow
Although interrupts allow for overlap of computation and I/O, they only really make sense for slow devices. Otherwise, the cost of interrupt handling and context switching may outweigh the benefits interrupts provide. There are also cases where a flood of interrupts may overload a system and lead it to livelock [MR96]; in such cases, polling provides more control to the OS in its scheduling and thus is again useful.

Overlap, are best. If the speed of the device is not known, or sometimes fast and sometimes slow, it may be best to use a **hybrid** that polls for a little while and then, if the device is not yet finished, uses interrupts. This **two-phased** approach may achieve the best of both worlds.

Another reason not to use interrupts arises in networks [MR96]. When a huge stream of incoming packets each generate an interrupt, it is possible for the OS to **livelock**, that is, find itself only processing interrupts and never allowing a user-level process to run and actually service the requests. For example, imagine a web server that suddenly experiences a high load due to the “slashdot effect”. In this case, it is better to occasionally use polling to better control what is happening in the system and allow the web server to service some requests before going back to the device to check for more packet arrivals.

Another interrupt-based optimization is **coalescing**. In such a setup, a device which needs to raise an interrupt first waits for a bit before delivering the interrupt to the CPU. While waiting, other requests may soon complete, and thus multiple interrupts can be coalesced into a single interrupt delivery, thus lowering the overhead of interrupt processing. Of course, waiting too long will increase the latency of a request, a common trade-off in systems. See Ahmad et al. [A+11] for an excellent summary.

### 36.5 More Efficient Data Movement With DMA

Unfortunately, there is one other aspect of our canonical protocol that requires our attention. In particular, when using programmed I/O (PIO) to transfer a large chunk of data to a device, the CPU is once again overburdened with a rather trivial task, and thus wastes a lot of time and effort that could better be spent running other processes. This timeline illustrates the problem:

```
CPU | 1 | 1 | 1 | 1 | c | c | c | 2 | 2 | 2 | 2 | 1 | 1
Disk| 1 | 1 | 1 | 1 | 1 |
```

In the timeline, Process 1 is running and then wishes to write some data to the disk. It then initiates the I/O, which must copy the data from memory to the device explicitly, one word at a time (marked c in the diagram). When the copy is complete, the I/O begins on the disk and the CPU can finally be used for something else.
THE CRUX: HOW TO LOWER PIO OVERHEADS

With PIO, the CPU spends too much time moving data to and from devices by hand. How can we offload this work and thus allow the CPU to be more effectively utilized?

The solution to this problem is something we refer to as **Direct Memory Access (DMA)**. A DMA engine is essentially a very specific device within a system that can orchestrate transfers between devices and main memory without much CPU intervention.

DMA works as follows. To transfer data to the device, for example, the OS would program the DMA engine by telling it where the data lives in memory, how much data to copy, and which device to send it to. At that point, the OS is done with the transfer and can proceed with other work. When the DMA is complete, the DMA controller raises an interrupt, and the OS thus knows the transfer is complete. The revised timeline:

```
CPU: 1 1 1 1 2 2 2 2 2 2 2 2 1 1
DMA: c c c
Disk: 1 1 1 1 1
```

From the timeline, you can see that the copying of data is now handled by the DMA controller. Because the CPU is free during that time, the OS can do something else, here choosing to run Process 2. Process 2 thus gets to use more CPU before Process 1 runs again.

36.6 Methods Of Device Interaction

Now that we have some sense of the efficiency issues involved with performing I/O, there are a few other problems we need to handle to incorporate devices into modern systems. One problem you may have noticed thus far: we have not really said anything about how the OS actually communicates with the device! Thus, the problem:

THE CRUX: HOW TO COMMUNICATE WITH DEVICES

How should the hardware communicate with a device? Should there be explicit instructions? Or are there other ways to do it?

Over time, two primary methods of device communication have developed. The first, oldest method (used by IBM mainframes for many years) is to have explicit **I/O instructions**. These instructions specify a way for the OS to send data to specific device registers and thus allow the construction of the protocols described above.
For example, on x86, the `in` and `out` instructions can be used to communicate with devices. For example, to send data to a device, the caller specifies a register with the data in it, and a specific `port` which names the device. Executing the instruction leads to the desired behavior.

Such instructions are usually **privileged**. The OS controls devices, and the OS thus is the only entity allowed to directly communicate with them. Imagine if any program could read or write the disk, for example: total chaos (as always), as any user program could use such a loophole to gain complete control over the machine.

The second method to interact with devices is known as **memory-mapped I/O**. With this approach, the hardware makes device registers available as if they were memory locations. To access a particular register, the OS issues a load (to read) or store (to write) the address; the hardware then routes the load/store to the device instead of main memory.

There is not some great advantage to one approach or the other. The memory-mapped approach is nice in that no new instructions are needed to support it, but both approaches are still in use today.

### 36.7 Fitting Into The OS: The Device Driver

One final problem we will discuss: how to fit devices, each of which have very specific interfaces, into the OS, which we would like to keep as general as possible. For example, consider a file system. We’d like to build a file system that worked on top of SCSI disks, IDE disks, USB keychain drives, and so forth, and we’d like the file system to be relatively oblivious to all of the details of how to issue a read or write request to these difference types of drives. Thus, our problem:

**The Crux: How To Build A Device-neutral OS**

How can we keep most of the OS device-neutral, thus hiding the details of device interactions from major OS subsystems?

The problem is solved through the age-old technique of **abstraction**. At the lowest level, a piece of software in the OS must know in detail how a device works. We call this piece of software a **device driver**, and any specifics of device interaction are encapsulated within.

Let us see how this abstraction might help OS design and implementation by examining the Linux file system software stack. Figure 36.3 is a rough and approximate depiction of the Linux software organization. As you can see from the diagram, a file system (and certainly, an application above) is completely oblivious to the specifics of which disk class it is using; it simply issues block read and write requests to the generic block layer, which routes them to the appropriate device driver, which handles the details of issuing the specific request. Although simplified, the diagram shows how such detail can be hidden from most of the OS.
For example, if there is a device that has many special capabilities, but has to present a generic interface to the rest of the kernel, those special capabilities will go unused. This situation arises, for example, in Linux with SCSI devices, which have very rich error reporting; because other block devices (e.g., ATA/IDE) have much simpler error handling, all that higher levels of software ever receive is a generic EIO (generic IO error) error code; any extra detail that SCSI may have provided is thus lost to the file system [G08].

Interestingly, because device drivers are needed for any device you might plug into your system, over time they have come to represent a huge percentage of kernel code. Studies of the Linux kernel reveal that over 70% of OS code is found in device drivers [C01]; for Windows-based systems, it is likely quite high as well. Thus, when people tell you that the OS has millions of lines of code, what they are really saying is that the OS has millions of lines of device-driver code. Of course, for any given installation, most of that code may not be active (i.e., only a few devices are connected to the system at a time). Perhaps more depressingly, as drivers are often written by “amateurs” (instead of full-time kernel developers), they tend to have many more bugs and thus are a primary contributor to kernel crashes [S03].

36.8 Case Study: A Simple IDE Disk Driver

To dig a little deeper here, let’s take a quick look at an actual device: an IDE disk drive [L94]. We summarize the protocol as described in this reference [W10]; we’ll also peek at the xv6 source code for a simple example of a working IDE driver [CK+08].

An IDE disk presents a simple interface to the system, consisting of four types of register: control, command block, status, and error. These registers are available by reading or writing to specific “I/O addresses” (such as 0x3F6 below) using (on x86) the in and out I/O instructions.
Control Register:
Address 0x3F6 = 0x80 (0000 1RE0): R=reset, E=0 means "enable interrupt"

Command Block Registers:
Address 0x1F0 = Data Port
Address 0x1F1 = Error
Address 0x1F2 = Sector Count
Address 0x1F3 = LBA low byte
Address 0x1F4 = LBA mid byte
Address 0x1F5 = LBA hi byte
Address 0x1F6 = 1B1D TOP4LBA: B=LBA, D=drive
Address 0x1F7 = Command/status

Status Register (Address 0x1F7):
7 6 5 4 3 2 1 0
BUSY READY FAULT SEEK DRQ CORR IDDEX ERROR

Error Register (Address 0x1F1): (check when Status ERROR==1)
7 6 5 4 3 2 1 0
BBK = Bad Block
UNC = Uncorrectable data error
MC = Media Changed
IDNF = ID mark Not Found
MCR = Media Change Requested
ABRT = Command aborted
TONF = Track 0 Not Found
AMNF = Address Mark Not Found

Figure 36.4: The IDE Interface
The basic protocol to interact with the device is as follows, assuming it has already been initialized.

- **Wait for drive to be ready.** Read Status Register (0x1F7) until drive is not busy and READY.
- **Write parameters to command registers.** Write the sector count, logical block address (LBA) of the sectors to be accessed, and drive number (master=0x00 or slave=0x10, as IDE permits just two drives) to command registers (0x1F2-0x1F6).
- **Start the I/O.** by issuing read/write to command register. Write READ—WRITE command to command register (0x1F7).
- **Data transfer (for writes):** Wait until drive status is READY and DRQ (drive request for data); write data to data port.
- **Handle interrupts.** In the simplest case, handle an interrupt for each sector transferred; more complex approaches allow batching and thus one final interrupt when the entire transfer is complete.
- **Error handling.** After each operation, read the status register. If the ERROR bit is on, read the error register for details.

Most of this protocol is found in the xv6 IDE driver (Figure 36.5), which (after initialization) works through four primary functions. The first is `ide_rw()`, which queues a request (if there are others pending), or issues it directly to the disk (via `ide_start_request()`); in either
static int ide_wait_ready() {
    while (((int r = inb(0x1f7)) & IDE_BSY) || !(r & IDE_DRDY))
        // loop until drive isn’t busy
}

static void ide_start_request(struct buf *b) {
    ide_wait_ready();
    outb(0x3f6, 0); // generate interrupt
    outb(0x1f3, b->sector & 0xff); // LBA goes here ... 
    outb(0x1f4, (b->sector >> 8) & 0xff); // ... and here 
    outb(0x1f5, (b->sector >> 16) & 0xff); // ... and here!
    outb(0x1f6, 0xe0 | ((b->dev&1)<<4) | ((b->sector>>24)&0x0f));
    if(b->flags & B_DIRTY){
        outb(0x1f7, IDE_CMD_WRITE); // this is a WRITE
        insl(0x1f0, b->data, 512/4); // transfer data too!
    } else {
        outb(0x1f7, IDE_CMD_READ); // this is a READ (no data)
    }
}

void ide_rw(struct buf *b) {
    acquire(&ide_lock);
    for (struct buf **pp = &ide_queue; *pp; pp=&(*pp)->qnext)
        // walk queue
    *pp = b; // add request to end
    if (ide_queue == b) // if q is empty
        ide_start_request(b); // send req to disk
    while ((b->flags & (B_VALID|B_DIRTY)) != B_VALID)
        sleep(b, &ide_lock); // wait for completion
    release(&ide_lock);
}

void ide_intr() {
    struct buf *b;
    acquire(&ide_lock);
    if (!((b->flags & B_DIRTY) && ide_wait_ready(1) >= 0))
        insl(0x1f0, b->data, 512/4); // if READ: get data
    b->flags |= B_VALID;
    b->flags ^= B_DIRTY;
    wakeup(b); // wake waiting process
    if ((ide_queue = b->qnext) != 0) // start next request
        ide_start_request(ide_queue); // (if one exists)
    release(&ide_lock);
}

Figure 36.5: The xv6 IDE Disk Driver (Simplified)

case, the routine waits for the request to complete and the calling process is put to sleep. The second is ide_start_request(), which is used to send a request (and perhaps data, in the case of a write) to the disk; the in and out x86 instructions are called to read and write device registers, respectively. The start request routine uses the third function, ide_wait_ready(), to ensure the drive is ready before issuing a request to it. Finally, ide_intr() is invoked when an interrupt takes place; it reads data from the device (if the request is a read, not a write), wakes the process waiting for the I/O to complete, and (if there are more requests in the I/O queue), launches the next I/O via ide_start_request().

Figure 36.5: The xv6 IDE Disk Driver (Simplified)
36.9 Historical Notes

Before ending, we include a brief historical note on the origin of some of these fundamental ideas. If you are interested in learning more, read Smotherman’s excellent summary [S08].

Interrupts are an ancient idea, existing on the earliest of machines. For example, the UNIVAC in the early 1950’s had some form of interrupt vectoring, although it is unclear in exactly which year this feature was available [S08]. Sadly, even in its infancy, we are beginning to lose the origins of computing history.

There is also some debate as to which machine first introduced the idea of DMA. For example, Knuth and others point to the DYSEAC (a “mobile” machine, which at the time meant it could be hauled in a trailer), whereas others think the IBM SAGE may have been the first [S08]. Either way, by the mid 50’s, systems with I/O devices that communicated directly with memory and interrupted the CPU when finished existed.

The history here is difficult to trace because the inventions are tied to real, and sometimes obscure, machines. For example, some think that the Lincoln Labs TX-2 machine was first with vectored interrupts [S08], but this is hardly clear.

Because the ideas are relatively obvious – no Einsteinian leap is required to come up with the idea of letting the CPU do something else while a slow I/O is pending – perhaps our focus on “who first?” is misguided. What is certainly clear: as people built these early machines, it became obvious that I/O support was needed. Interrupts, DMA, and related ideas are all direct outcomes of the nature of fast CPUs and slow devices; if you were there at the time, you might have had similar ideas.

36.10 Summary

You should now have a very basic understanding of how an OS interacts with a device. Two techniques, the interrupt and DMA, have been introduced to help with device efficiency, and two approaches to accessing device registers, explicit I/O instructions and memory-mapped I/O, have been described. Finally, the notion of a device driver has been presented, showing how the OS itself can encapsulate low-level details and thus make it easier to build the rest of the OS in a device-neutral fashion.
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Andy Chou, Junfeng Yang, Benjamin Chelf, Seth Hallem, Dawson Engler
SOSP ’01
One of the first papers to systematically explore how many bugs are in modern operating systems.
Among other neat findings, the authors show that device drivers have something like seven times more bugs than mainline kernel code.

[CK+08] “The xv6 Operating System”
Russ Cox, Frans Kaashoek, Robert Morris, Nickolai Zeldovich
See ide.c for the IDE device driver, with a few more details therein.

[D07] “What Every Programmer Should Know About Memory”
Ulrich Drepper
November, 2007
A fantastic read about modern memory systems, starting at DRAM and going all the way up to virtualization and cache-optimized algorithms.

[G08] “EIO: Error-handling is Occasionally Correct”
Haryadi Gunawi, Cindy Rubio-Gonzalez, Andrea Arpaci-Dusseau, Remzi Arpaci-Dusseau, Ben Liblit
FAST ’08, San Jose, CA, February 2008
Our own work on building a tool to find code in Linux file systems that does not handle error return properly. We found hundreds and hundreds of bugs, many of which have now been fixed.

[L94] “AT Attachment Interface for Disk Drives”
Lawrence J. Lamers, X3T10 Technical Editor

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USENIX ’96, San Diego, CA, January 1996
Mogul and colleagues did a great deal of pioneering work on web server network performance. This paper is but one example.

[S08] “Interrupts”
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Available: http://people.cs.clemson.edu/~mark/interrupts.html
A treasure trove of information on the history of interrupts, DMA, and related early ideas in computing.
[S03] “Improving the Reliability of Commodity Operating Systems”  
Michael M. Swift, Brian N. Bershad, and Henry M. Levy  
SOSP ’03  
Swift’s work revived interest in a more microkernel-like approach to operating systems; minimally, it finally gave some good reasons why address-space based protection could be useful in a modern OS.

[W10] “Hard Disk Driver”  
Washington State Course Homepage  
Available: http://eecs.wsu.edu/~cs460/cs560/HDdriver.html  
A nice summary of a simple IDE disk drive’s interface and how to build a device driver for it.
The last chapter introduced the general concept of an I/O device and showed you how the OS might interact with such a beast. In this chapter, we dive into more detail about one device in particular: the hard disk drive. These drives have been the main form of persistent data storage in computer systems for decades and much of the development of file system technology (coming soon) is predicated on their behavior. Thus, it is worth understanding the details of a disk’s operation before building the file system software that manages it. Many of these details are available in excellent papers by Ruemmler and Wilkes [RW92] and Anderson, Dykes, and Riedel [ADR03].

**Crux: How to Store and Access Data on Disk**

- How do modern hard-disk drives store data? What is the interface?
- How is the data actually laid out and accessed? How does disk scheduling improve performance?

### 37.1 The Interface

Let’s start by understanding the interface to a modern disk drive. The basic interface for all modern drives is straightforward. The drive consists of a large number of sectors (512-byte blocks), each of which can be read or written. The sectors are numbered from 0 to \( n - 1 \) on a disk with \( n \) sectors. Thus, we can view the disk as an array of sectors; 0 to \( n - 1 \) is thus the address space of the drive.

Multi-sector operations are possible; indeed, many file systems will read or write 4KB at a time (or more). However, when updating the disk, the only guarantee drive manufactures make is that a single 512-byte write is atomic (i.e., it will either complete in its entirety or it won’t complete at all); thus, if an untimely power loss occurs, only a portion of a larger write may complete (sometimes called a torn write).
37.2 Basic Geometry

Let’s start to understand some of the components of a modern disk. We start with a **platter**, a circular hard surface on which data is stored persistently by inducing magnetic changes to it. A disk may have one or more platters; each platter has 2 sides, each of which is called a **surface**. These platters are usually made of some hard material (such as aluminum), and then coated with a thin magnetic layer that enables the drive to persistently store bits even when the drive is powered off.

The platters are all bound together around the **spindle**, which is connected to a motor that spins the platters around (while the drive is powered on) at a constant (fixed) rate. The rate of rotation is often measured in **rotations per minute (RPM)**, and typical modern values are in the 7,200 RPM to 15,000 RPM range. Note that we will often be interested in the time of a single rotation, e.g., a drive that rotates at 10,000 RPM means that a single rotation takes about 6 milliseconds (6 ms).

Data is encoded on each surface in concentric circles of sectors; we call one such concentric circle a **track**. A single surface contains many thousands and thousands of tracks, tightly packed together, with hundreds of tracks fitting into the width of a human hair.

To read and write from the surface, we need a mechanism that allows us to either sense (i.e., read) the magnetic patterns on the disk or to induce a change in (i.e., write) them. This process of reading and writing is accomplished by the **disk head**; there is one such head per surface of the drive. The disk head is attached to a single **disk arm**, which moves across the surface to position the head over the desired track.

37.3 A Simple Disk Drive

Let’s understand how disks work by building up a model one track at a time. Assume we have a simple disk with a single track (Figure 37.1).
This track has just 12 sectors, each of which is 512 bytes in size (our typical sector size, recall) and addressed therefore by the numbers 0 through 11. The single platter we have here rotates around the spindle, to which a motor is attached. Of course, the track by itself isn’t too interesting; we want to be able to read or write those sectors, and thus we need a disk head, attached to a disk arm, as we now see (Figure 37.2).

In the figure, the disk head, attached to the end of the arm, is positioned over sector 6, and the surface is rotating counter-clockwise.

**Single-track Latency: The Rotational Delay**

To understand how a request would be processed on our simple, one-track disk, imagine we now receive a request to read block 0. How should the disk service this request?

In our simple disk, the disk doesn’t have to do much. In particular, it must just wait for the desired sector to rotate under the disk head. This wait happens often enough in modern drives, and is an important enough component of I/O service time, that it has a special name: **rotational delay** (sometimes **rotation delay**, though that sounds weird). In the example, if the full rotational delay is $R$, the disk has to incur a rotational delay of about $\frac{R}{2}$ to wait for 0 to come under the read/write head (if we start at 6). A worst-case request on this single track would be to sector 5, causing nearly a full rotational delay in order to service such a request.

**Multiple Tracks: Seek Time**

So far our disk just has a single track, which is not too realistic; modern disks of course have many millions. Let’s thus look at ever-so-slightly more realistic disk surface, this one with three tracks (Figure 37.3, left).

In the figure, the head is currently positioned over the innermost track (which contains sectors 24 through 35); the next track over contains the next set of sectors (12 through 23), and the outermost track contains the first sectors (0 through 11).
To understand how the drive might access a given sector, we now trace what would happen on a request to a distant sector, e.g., a read to sector 11. To service this read, the drive has to first move the disk arm to the correct track (in this case, the outermost one), in a process known as a seek. Seeks, along with rotations, are one of the most costly disk operations.

The seek, it should be noted, has many phases: first an acceleration phase as the disk arm gets moving; then coasting as the arm is moving at full speed, then deceleration as the arm slows down; finally settling as the head is carefully positioned over the correct track. The settling time is often quite significant, e.g., 0.5 to 2 ms, as the drive must be certain to find the right track (imagine if it just got close instead!).

After the seek, the disk arm has positioned the head over the right track. A depiction of the seek is found in Figure 37.3 (right).

As we can see, during the seek, the arm has been moved to the desired track, and the platter of course has rotated, in this case about 3 sectors. Thus, sector 9 is just about to pass under the disk head, and we must only endure a short rotational delay to complete the transfer.

When sector 11 passes under the disk head, the final phase of I/O will take place, known as the transfer, where data is either read from or written to the surface. And thus, we have a complete picture of I/O time: first a seek, then waiting for the rotational delay, and finally the transfer.

**Some Other Details**

Though we won’t spend too much time on it, there are some other interesting details about how hard drives operate. Many drives employ some kind of track skew to make sure that sequential reads can be properly serviced even when crossing track boundaries. In our simple example disk, this might appear as seen in Figure 37.4.
Sectors are often skewed like this because when switching from one track to another, the disk needs time to reposition the head (even to neighboring tracks). Without such skew, the head would be moved to the next track but the desired next block would have already rotated under the head, and thus the drive would have to wait almost the entire rotational delay to access the next block.

Another reality is that outer tracks tend to have more sectors than inner tracks, which is a result of geometry; there is simply more room out there. These tracks are often referred to as multi-zoned disk drives, where the disk is organized into multiple zones, and where a zone is consecutive set of tracks on a surface. Each zone has the same number of sectors per track, and outer zones have more sectors than inner zones.

Finally, an important part of any modern disk drive is its cache, for historical reasons sometimes called a track buffer. This cache is just some small amount of memory (usually around 8 or 16 MB) which the drive can use to hold data read from or written to the disk. For example, when reading a sector from the disk, the drive might decide to read in all of the sectors on that track and cache them in its memory; doing so allows the drive to quickly respond to any subsequent requests to the same track.

On writes, the drive has a choice: should it acknowledge the write has completed when it has put the data in its memory, or after the write has actually been written to disk? The former is called write back caching (or sometimes immediate reporting), and the latter write through. Write back caching sometimes makes the drive appear “faster”, but can be dangerous; if the file system or applications require that data be written to disk in a certain order for correctness, write-back caching can lead to problems (read the chapter on file-system journaling for details).
ASIDE: DIMENSIONAL ANALYSIS

Remember in Chemistry class, how you solved virtually every problem by simply setting up the units such that they canceled out, and somehow the answers popped out as a result? That chemical magic is known by the highfalutin name of **dimensional analysis** and it turns out it is useful in computer systems analysis too.

Let’s do an example to see how dimensional analysis works and why it is useful. In this case, assume you have to figure out how long, in milliseconds, a single rotation of a disk takes. Unfortunately, you are given only the **RPM** of the disk, or rotations per minute. Let’s assume we’re talking about a 10K RPM disk (i.e., it rotates 10,000 times per minute). How do we set up the dimensional analysis so that we get time per rotation in milliseconds?

To do so, we start by putting the desired units on the left; in this case, we wish to obtain the time (in milliseconds) per rotation, so that is exactly what we write down: \( \text{Time (ms)} \quad 1 \text{ Rotation} \). We then write down everything we know, making sure to cancel units where possible. First, we obtain \( \frac{1 \text{ minute}}{10,000 \text{ Rotations}} \) (keeping rotation on the bottom, as that’s where it is on the left), then transform minutes into seconds with \( \frac{60 \text{ seconds}}{1 \text{ minute}} \), and then finally transform seconds in milliseconds with \( \frac{1000 \text{ ms}}{1 \text{ second}} \). The final result is this equation, with units nicely canceled, is:

\[
\frac{\text{Time (ms)}}{1 \text{ Rot.}} = \frac{1 \text{ minute}}{10,000 \text{ Rot.}} \cdot \frac{60 \text{ seconds}}{1 \text{ minute}} \cdot \frac{1000 \text{ ms}}{1 \text{ second}} = \frac{6 \text{ ms}}{1 \text{ Rotation}}
\]

As you can see from this example, dimensional analysis makes what seems obvious into a simple and repeatable process. Beyond the RPM calculation above, it comes in handy with I/O analysis regularly. For example, you will often be given the transfer rate of a disk, e.g., 100 MB/second, and then asked: how long does it take to transfer a 512 KB block (in milliseconds)? With dimensional analysis, it’s easy:

\[
\frac{\text{Time (ms)}}{1 \text{ Request}} = \frac{512 \text{ KB}}{1 \text{ Request}} \cdot \frac{1 \text{ MB}}{1024 \text{ KB}} \cdot \frac{1 \text{ second}}{100 \text{ MB}} \cdot \frac{1000 \text{ ms}}{1 \text{ second}} = \frac{5 \text{ ms}}{1 \text{ Request}}
\]

37.4 I/O Time: Doing The Math

Now that we have an abstract model of the disk, we can use a little analysis to better understand disk performance. In particular, we can now represent I/O time as the sum of three major components:

\[
T_{I/O} = T_{\text{seek}} + T_{\text{rotation}} + T_{\text{transfer}} \quad (37.1)
\]
Table 37.1: **Disk Drive Specs: SCSI Versus SATA**

<table>
<thead>
<tr>
<th></th>
<th>Cheetah 15K.5</th>
<th>Barracuda</th>
</tr>
</thead>
<tbody>
<tr>
<td>Capacity</td>
<td>300 GB</td>
<td>1 TB</td>
</tr>
<tr>
<td>RPM</td>
<td>15,000</td>
<td>7,200</td>
</tr>
<tr>
<td>Average Seek</td>
<td>4 ms</td>
<td>9 ms</td>
</tr>
<tr>
<td>Max Transfer</td>
<td>125 MB/s</td>
<td>105 MB/s</td>
</tr>
<tr>
<td>Platters</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>Cache</td>
<td>16 MB</td>
<td>16/32 MB</td>
</tr>
<tr>
<td>Connects via</td>
<td>SCSI</td>
<td>SATA</td>
</tr>
</tbody>
</table>

Note that the rate of I/O ($R_{I/O}$), which is often more easily used for comparison between drives (as we will do below), is easily computed from the time. Simply divide the size of the transfer by the time it took:

$$R_{I/O} = \frac{\text{Size Transfer}}{T_{I/O}} \tag{37.2}$$

To get a better feel for I/O time, let us perform the following calculation. Assume there are two workloads we are interested in. The first, known as the **random** workload, issues small (e.g., 4KB) reads to random locations on the disk. Random workloads are common in many important applications, including database management systems. The second, known as the **sequential** workload, simply reads a large number of sectors consecutively from the disk, without jumping around. Sequential access patterns are quite common and thus important as well.

To understand the difference in performance between random and sequential workloads, we need to make a few assumptions about the disk drive first. Let’s look at a couple of modern disks from Seagate. The first, known as the Cheetah 15K.5 [S09b], is a high-performance SCSI drive. The second, the Barracuda [S09a], is a drive built for capacity. Details on both are found in Table 37.1.

As you can see, the drives have quite different characteristics, and in many ways nicely summarize two important components of the disk drive market. The first is the “high performance” drive market, where drives are engineered to spin as fast as possible, deliver low seek times, and transfer data quickly. The second is the “capacity” market, where cost per byte is the most important aspect; thus, the drives are slower but pack as many bits as possible into the space available.

From these numbers, we can start to calculate how well the drives would do under our two workloads outlined above. Let’s start by looking at the random workload. Assuming each 4 KB read occurs at a random location on disk, we can calculate how long each such read would take. On the Cheetah:

$$T_{\text{seek}} = 4 \text{ ms}, \ T_{\text{rotation}} = 2 \text{ ms}, \ T_{\text{transfer}} = 30 \text{ microsecs} \tag{37.3}$$
When at all possible, transfer data to and from disks in a sequential manner. If sequential is not possible, at least think about transferring data in large chunks: the bigger, the better. If I/O is done in little random pieces, I/O performance will suffer dramatically. Also, users will suffer. Also, you will suffer, knowing what suffering you have wrought with your careless random I/Os.

The average seek time (4 milliseconds) is just taken as the average time reported by the manufacturer; note that a full seek (from one end of the surface to the other) would likely take two or three times longer. The average rotational delay is calculated from the RPM directly. 15000 RPM is equal to 250 RPS (rotations per second); thus, each rotation takes 4 ms. On average, the disk will encounter a half rotation and thus 2 ms is the average time. Finally, the transfer time is just the size of the transfer over the peak transfer rate; here it is vanishingly small (30 microseconds; note that we need 1000 microseconds just to get 1 millisecond!).

Thus, from our equation above, $T_{I/O}$ for the Cheetah roughly equals 6 ms. To compute the rate of I/O, we just divide the size of the transfer by the average time, and thus arrive at $R_{I/O}$ for the Cheetah under the random workload of about 0.66 MB/s. The same calculation for the Barracuda yields a $T_{I/O}$ of about 13.2 ms, more than twice as slow, and thus a rate of about 0.31 MB/s.

Now let’s look at the sequential workload. Here we can assume there is a single seek and rotation before a very long transfer. For simplicity, assume the size of the transfer is 100 MB. Thus, $T_{I/O}$ for the Barracuda and Cheetah is about 800 ms and 950 ms, respectively. The rates of I/O are thus very nearly the peak transfer rates of 125 MB/s and 105 MB/s, respectively. Table 37.2 summarizes these numbers.

The table shows us a number of important things. First, and most importantly, there is a huge gap in drive performance between random and sequential workloads, almost a factor of 200 or so for the Cheetah and more than a factor 300 difference for the Barracuda. And thus we arrive at the most obvious design tip in the history of computing.

A second, more subtle point: there is a large difference in performance between high-end “performance” drives and low-end “capacity” drives. For this reason (and others), people are often willing to pay top dollar for the former while trying to get the latter as cheaply as possible.

<table>
<thead>
<tr>
<th>Drive</th>
<th>Random</th>
<th>Sequential</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cheetah</td>
<td>0.66 MB/s</td>
<td>125 MB/s</td>
</tr>
<tr>
<td>Barracuda</td>
<td>0.31 MB/s</td>
<td>105 MB/s</td>
</tr>
</tbody>
</table>

Table 37.2: Disk Drive Performance: SCSI Versus SATA
ASIDE: Computing The “Average” Seek

In many books and papers, you will see average disk-seek time cited as being roughly one-third of the full seek time. Where does this come from?

Turns out it arises from a simple calculation based on average seek distance, not time. Imagine the disk as a set of tracks, from 0 to \( N \). The seek distance between any two tracks \( x \) and \( y \) is thus computed as the absolute value of the difference between them: \( |x - y| \).

To compute the average seek distance, all you need to do is to first add up all possible seek distances:

\[
\sum_{x=0}^{N} \sum_{y=0}^{N} |x - y|.
\]

(37.4)

Then, divide this by the number of different possible seeks: \( N^2 \). To compute the sum, we’ll just use the integral form:

\[
\int_{x=0}^{N} \int_{y=0}^{N} |x - y| \, dy \, dx.
\]

(37.5)

To compute the inner integral, let’s break it out the absolute value:

\[
\int_{y=0}^{x} (x - y) \, dy + \int_{y=x}^{N} (y - x) \, dy.
\]

(37.6)

Solving this leads to \((xy - \frac{1}{2}y^2)|_0^x + (\frac{1}{2}y^2 - xy)|_x^N\) which can be simplified to \((x^2 - Nx + \frac{1}{2}N^2)\). Now we have to compute the outer integral:

\[
\int_{x=0}^{N} (x^2 - Nx + \frac{1}{2}N^2) \, dx,
\]

(37.7)

which results in:

\[
\left(\frac{1}{3}x^3 - \frac{N}{2}x^2 + \frac{N^2}{2}x\right)|_0^N = \frac{N^3}{3}.
\]

(37.8)

Remember that we still have to divide by the total number of seeks \( (N^2) \) to compute the average seek distance: \( \frac{\frac{N^3}{3}}{N^2} = \frac{1}{3}N \). Thus the average seek distance on a disk, over all possible seeks, is one-third the full distance. And now when you hear that an average seek is one-third of a full seek, you’ll know where it came from.
Because of the high cost of I/O, the OS has historically played a role in deciding the order of I/Os issued to the disk. More specifically, given a set of I/O requests, the disk scheduler examines the requests and decides which one to schedule next [SCO90, JW91].

Unlike job scheduling, where the length of each job is usually unknown, with disk scheduling, we can make a good guess at how long a “job” (i.e., disk request) will take. By estimating the seek and possible the rotational delay of a request, the disk scheduler can know how long each request will take, and thus (greedily) pick the one that will take the least time to service first. Thus, the disk scheduler will try to follow the principle of SJF (shortest job first) in its operation.

**SSTF: Shortest Seek Time First**

One early disk scheduling approach is known as **shortest-seek-time-first** (SSTF) (also called **shortest-seek-first** or SSF). SSTF orders the queue of I/O requests by track, picking requests on the nearest track to complete first. For example, assuming the current position of the head is over the inner track, and we have requests for sectors 21 (middle track) and 2 (outer track), we would then issue the request to 21 first, wait for it to complete, and then issue the request to 2 (Figure 37.5).

SSTF works well in this example, seeking to the middle track first and then the outer track. However, SSTF is not a panacea, for the following reasons. First, the drive geometry is not available to the host OS; rather, it sees an array of blocks. Fortunately, this problem is rather easily fixed. Instead of SSTF, an OS can simply implement **nearest-block-first** (NBF), which schedules the request with the nearest block address next.
The second problem is more fundamental: starvation. Imagine in our example above if there were a steady stream of requests to the inner track, where the head currently is positioned. Requests to any other tracks would then be ignored completely by a pure SSTF approach. And thus the crux of the problem:

**Crux: How To Handle Disk Starvation**
How can we implement SSTF-like scheduling but avoid starvation?

**Elevator (a.k.a. SCAN or C-SCAN)**

The answer to this query was developed some time ago (see [CKR72] for example), and is relatively straightforward. The algorithm, originally called SCAN, simply moves across the disk servicing requests in order across the tracks. Let us call a single pass across the disk a sweep. Thus, if a request comes for a block on a track that has already been serviced on this sweep of the disk, it is not handled immediately, but rather queued until the next sweep.

SCAN has a number of variants, all of which do about the same thing. For example, Coffman et al. introduced F-SCAN, which freezes the queue to be serviced when it is doing a sweep [CKR72]; this action places requests that come in during the sweep into a queue to be serviced later. Doing so avoids starvation of far-away requests, by delaying the servicing of late-arriving (but nearer by) requests.

C-SCAN is another common variant, short for Circular SCAN. Instead of sweeping in one direction across the disk, the algorithm sweeps from outer-to-inner, and then inner-to-outer, etc.

For reasons that should now be obvious, this algorithm (and its variants) is sometimes referred to as the elevator algorithm, because it behaves like an elevator which is either going up or down and not just servicing requests to floors based on which floor is closer. Imagine how annoying it would be if you were going down from floor 10 to 1, and somebody got on at 3 and pressed 4, and the elevator went up to 4 because it was “closer” than 1! As you can see, the elevator algorithm, when used in real life, prevents fights from taking place on elevators. In disks, it just prevents starvation.

Unfortunately, SCAN and its cousins do not represent the best scheduling technology. In particular, SCAN (or SSTF even) do not actually adhere as closely to the principle of SJF as they could. In particular, they ignore rotation. And thus, another crux:

**Crux: How To Account For Disk Rotation Costs**
How can we implement an algorithm that more closely approximates SJF by taking both seek and rotation into account?
SPTF: Shortest Positioning Time First

Before discussing shortest positioning time first or SPTF scheduling (sometimes also called shortest access time first or SATF), which is the solution to our problem, let us make sure we understand the problem in more detail. Figure 37.6 presents an example.

In the example, the head is currently positioned over sector 30 on the inner track. The scheduler thus has to decide: should it schedule sector 16 (on the middle track) or sector 8 (on the outer track) for its next request. So which should it service next?

The answer, of course, is “it depends”. In engineering, it turns out “it depends” is almost always the answer, reflecting that trade-offs are part of the life of the engineer; such maxims are also good in a pinch, e.g., when you don’t know an answer to your boss’s question, you might want to try this gem. However, it is almost always better to know why it depends, which is what we discuss here.

What it depends on here is the relative time of seeking as compared to rotation. If, in our example, seek time is much higher than rotational delay, then SSTF (and variants) are just fine. However, imagine if seek is quite a bit faster than rotation. Then, in our example, it would make more sense to seek further to service request 8 on the outer track than it would to perform the shorter seek to the middle track to service 16, which has to rotate all the way around before passing under the disk head.

On modern drives, as we saw above, both seek and rotation are roughly equivalent (depending, of course, on the exact requests), and thus SPTF is useful and improves performance. However, it is even more difficult to implement in an OS, which generally does not have a good idea where track boundaries are or where the disk head currently is (in a rotational sense). Thus, SPTF is usually performed inside a drive, described below.
Almost any question can be answered with “it depends”, as our colleague Miron Livny always says. However, use with caution, as if you answer too many questions this way, people will stop asking you questions altogether. For example, somebody asks: “want to go to lunch?” You reply: “it depends, are you coming along?”

Other Scheduling Issues

There are many other issues we do not discuss in this brief description of basic disk operation, scheduling, and related topics. One such issue is this: where is disk scheduling performed on modern systems? In older systems, the operating system did all the scheduling; after looking through the set of pending requests, the OS would pick the best one, and issue it to the disk. When that request completed, the next one would be chosen, and so forth. Disks were simpler then, and so was life.

In modern systems, disks can accommodate multiple outstanding requests, and have sophisticated internal schedulers themselves (which can implement SPTF accurately; inside the disk controller, all relevant details are available, including exact head position). Thus, the OS scheduler usually picks what it thinks the best few requests are (say 16) and issues them all to disk; the disk then uses its internal knowledge of head position and detailed track layout information to service said requests in the best possible (SPTF) order.

Another important related task performed by disk schedulers is I/O merging. For example, imagine a series of requests to read blocks 33, then 8, then 34, as in Figure 37.6. In this case, the scheduler should merge the requests for blocks 33 and 34 into a single two-block request; any reordering that the scheduler does is performed upon the merged requests. Merging is particularly important at the OS level, as it reduces the number of requests sent to the disk and thus lowers overheads.

One final problem that modern schedulers address is this: how long should the system wait before issuing an I/O to disk? One might naively think that the disk, once it has even a single I/O, should immediately issue the request to the drive; this approach is called work-conserving, as the disk will never be idle if there are requests to serve. However, research on anticipatory disk scheduling has shown that sometimes it is better to wait for a bit [ID01], in what is called a non-work-conserving approach. By waiting, a new and “better” request may arrive at the disk, and thus overall efficiency is increased. Of course, deciding when to wait, and for how long, can be tricky; see the research paper for details, or check out the Linux kernel implementation to see how such ideas are transitioned into practice (if you are the ambitious sort).
37.6 Summary

We have presented a summary of how disks work. The summary is actually a detailed functional model; it does not describe the amazing physics, electronics, and material science that goes into actual drive design. For those interested in even more details of that nature, we suggest a different major (or perhaps minor); for those that are happy with this model, good! We can now proceed to using the model to build more interesting systems on top of these incredible devices.
References

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A cool paper showing how waiting can improve disk scheduling: better requests may be on their way!

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A more modern take on disk scheduling. It remains a technical report (and not a published paper) because the authors were scooped by Seltzer et al. [SCO90].

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C. Ruemmler, J. Wilkes
A terrific introduction to the basics of disk operation. Some pieces are out of date, but most of the basics remain.

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[SG04] “MEMS-based storage devices and standard disk interfaces: A square peg in a round hole?”
Steven W. Schlosser, Gregory R. Ganger
FAST’04, pp. 87-100, 2004
While the MEMS aspect of this paper hasn’t yet made an impact, the discussion of the contract between file systems and disks is wonderful and a lasting contribution.

[S09a] “Barracuda ES.2 data sheet”

[S09b] “Cheetah 15K.5”

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This homework uses disk.py to familiarize you with how a modern hard drive works. It has a lot of different options, and unlike most of the other simulations, has a graphical animator to show you exactly what happens when the disk is in action. See the README for details.

1. Compute the seek, rotation, and transfer times for the following sets of requests: -a 0, -a 6, -a 30, -a 7, 30, 8, and finally -a 10, 11, 12, 13.
2. Do the same requests above, but change the seek rate to different values: -S 2, -S 4, -S 8, -S 10, -S 40, -S 0.1. How do the times change?
3. Do the same requests above, but change the rotation rate: -R 0.1, -R 0.5, -R 0.01. How do the times change?
4. You might have noticed that some request streams would be better served with a policy better than FIFO. For example, with the request stream -a 7, 30, 8, what order should the requests be processed in? Now run the shortest seek-time first (SSTF) scheduler (-p SSTF) on the same workload; how long should it take (seek, rotation, transfer) for each request to be served?
5. Now do the same thing, but using the shortest access-time first (SATF) scheduler (-p SATF). Does it make any difference for the set of requests as specified by -a 7, 30, 8? Find a set of requests where SATF does noticeably better than SSTF; what are the conditions for a noticeable difference to arise?
6. You might have noticed that the request stream -a 10, 11, 12, 13 wasn’t particularly well handled by the disk. Why is that? Can you introduce a track skew to address this problem (-o skew, where skew is a non-negative integer)? Given the default seek rate, what should the skew be to minimize the total time for this set of requests? What about for different seek rates (e.g., -S 2, -S 4)? In general, could you write a formula to figure out the skew, given the seek rate and sector layout information?
7. Multi-zone disks pack more sectors into the outer tracks. To configure this disk in such a way, run with the -z flag. Specifically, try running some requests against a disk run with -z 10, 20, 30 (the numbers specify the angular space occupied by a sector, per track; in this example, the outer track will be packed with a sector every 10 degrees, the middle track every 20 degrees, and the inner track with a sector every 30 degrees). Run some random requests (e.g., -a -1 -A 5, -1, 0, which specifies that random requests should be used via the -a -1 flag and that five requests ranging from 0 to the max be generated), and see if you can compute the seek, rotation, and transfer times. Use different random seeds (-s 1, -s 2, etc.). What is the bandwidth (in sectors per unit time) on the outer, middle, and inner tracks?
8. Scheduling windows determine how many sector requests a disk can examine at once in order to determine which sector to serve next. Generate some random workloads of a lot of requests (e.g., \(-A \ 1000, \ -1, \ 0\), with different seeds perhaps) and see how long the SATF scheduler takes when the scheduling window is changed from 1 up to the number of requests (e.g., \(-w \ 1 \) up to \(-w \ 1000\), and some values in between). How big of a scheduling window is needed to approach the best possible performance? Make a graph and see. Hint: use the \(-c\) flag and don’t turn on graphics with \(-G\) to run these more quickly. When the scheduling window is set to 1, does it matter which policy you are using?

9. Avoiding starvation is important in a scheduler. Can you think of a series of requests such that a particular sector is delayed for a very long time given a policy such as SATF? Given that sequence, how does it perform if you use a bounded SATF or BSATF scheduling approach? In this approach, you specify the scheduling window (e.g., \(-w \ 4\)) as well as the BSATF policy (\(-p \ \text{BSATF}\)); the scheduler then will only move onto the next window of requests when all of the requests in the current window have been serviced. Does this solve the starvation problem? How does it perform, as compared to SATF? In general, how should a disk make this trade-off between performance and starvation avoidance?

10. All the scheduling policies we have looked at thus far are greedy, in that they simply pick the next best option instead of looking for the optimal schedule over a set of requests. Can you find a set of requests in which this greedy approach is not optimal?
Redundant Arrays of Inexpensive Disks (RAIDs)

When we use a disk, we sometimes wish it to be faster; I/O operations are slow and thus can be the bottleneck for the entire system. When we use a disk, we sometimes wish it to be larger; more and more data is being put online and thus our disks are getting fuller and fuller. When we use a disk, we sometimes wish for it to be more reliable; when a disk fails, if our data isn’t backed up, all that valuable data is gone.

CRUX: HOW TO MAKE A LARGE, FAST, RELIABLE DISK

How can we make a large, fast, and reliable storage system? What are the key techniques? What are trade-offs between different approaches?

In this chapter, we introduce the **Redundant Array of Inexpensive Disks** better known as **RAID** \[P+88\], a technique to use multiple disks in concert to build a faster, bigger, and more reliable disk system. The term was introduced in the late 1980s by a group of researchers at U.C. Berkeley (led by Professors David Patterson and Randy Katz and then student Garth Gibson); it was around this time that many different researchers simultaneously arrived upon the basic idea of using multiple disks to build a better storage system \[BG88, K86, K88, PB86, SG86\].

Externally, a RAID looks like a disk: a group of blocks one can read or write. Internally, the RAID is a complex beast, consisting of multiple disks, memory (both volatile and non-), and one or more processors to manage the system. A hardware RAID is very much like a computer system, specialized for the task of managing a group of disks.

RAIDs offer a number of advantages over a single disk. One advantage is **performance**. Using multiple disks in parallel can greatly speed up I/O times. Another benefit is **capacity**. Large data sets demand large disks. Finally, RAIDs can improve **reliability**; spreading data across multiple disks (without RAID techniques) makes the data vulnerable to the loss of a single disk; with some form of **redundancy**, RAIDs can tolerate the loss of a disk and keep operating as if nothing were wrong.
When considering how to add new functionality to a system, one should always consider whether such functionality can be added transparently, in a way that demands no changes to the rest of the system. Requiring a complete rewrite of the existing software (or radical hardware changes) lessens the chance of impact of an idea. RAID is a perfect example, and certainly its transparency contributed to its success; administrators could install a SCSI-based RAID storage array instead of a SCSI disk, and the rest of the system (host computer, OS, etc.) did not have to change one bit to start using it. By solving this problem of deployment, RAID was made more successful from day one.

Amazingly, RAIDs provide these advantages transparently to systems that use them, i.e., a RAID just looks like a big disk to the host system. The beauty of transparency, of course, is that it enables one to simply replace a disk with a RAID and not change a single line of software; the operating system and client applications continue to operate without modification. In this manner, transparency greatly improves the deployability of RAID, enabling users and administrators to put a RAID to use without worries of software compatibility.

We now discuss some of the important aspects of RAIDs. We begin with the interface, fault model, and then discuss how one can evaluate a RAID design along three important axes: capacity, reliability, and performance. We then discuss a number of other issues that are important to RAID design and implementation.

38.1 Interface And RAID Internals

To a file system above, a RAID looks like a big, (hopefully) fast, and (hopefully) reliable disk. Just as with a single disk, it presents itself as a linear array of blocks, each of which can be read or written by the file system (or other client).

When a file system issues a logical I/O request to the RAID, the RAID internally must calculate which disk (or disks) to access in order to complete the request, and then issue one or more physical I/Os to do so. The exact nature of these physical I/Os depends on the RAID level, as we will discuss in detail below. However, as a simple example, consider a RAID that keeps two copies of each block (each one on a separate disk); when writing to such a mirrored RAID system, the RAID will have to perform two physical I/Os for every one logical I/O it is issued.

A RAID system is often built as a separate hardware box, with a standard connection (e.g., SCSI, or SATA) to a host. Internally, however, RAIDs are fairly complex, consisting of a microcontroller that runs firmware to direct the operation of the RAID, volatile memory such as DRAM to buffer data blocks as they are read and written, and in some cases,
non-volatile memory to buffer writes safely and perhaps even specialized logic to perform parity calculations (useful in some RAID levels, as we will also see below). At a high level, a RAID is very much a specialized computer system: it has a processor, memory, and disks; however, instead of running applications, it runs specialized software designed to operate the RAID.

38.2 Fault Model

To understand RAID and compare different approaches, we must have a fault model in mind. RAIDs are designed to detect and recover from certain kinds of disk faults; thus, knowing exactly which faults to expect is critical in arriving upon a working design.

The first fault model we will assume is quite simple, and has been called the fail-stop fault model [S84]. In this model, a disk can be in exactly one of two states: working or failed. With a working disk, all blocks can be read or written. In contrast, when a disk has failed, we assume it is permanently lost.

One critical aspect of the fail-stop model is what it assumes about fault detection. Specifically, when a disk has failed, we assume that this is easily detected. For example, in a RAID array, we would assume that the RAID controller hardware (or software) can immediately observe when a disk has failed.

Thus, for now, we do not have to worry about more complex “silent” failures such as disk corruption. We also do not have to worry about a single block becoming inaccessible upon an otherwise working disk (sometimes called a latent sector error). We will consider these more complex (and unfortunately, more realistic) disk faults later.

38.3 How To Evaluate A RAID

As we will soon see, there are a number of different approaches to building a RAID. Each of these approaches has different characteristics which are worth evaluating, in order to understand their strengths and weaknesses.

Specifically, we will evaluate each RAID design along three axes. The first axis is capacity; given a set of N disks, how much useful capacity is available to systems that use the RAID? Without redundancy, the answer is obviously N; however, if we have a system that keeps a two copies of each block, we will obtain a useful capacity of N/2. Different schemes (e.g., parity-based ones) tend to fall in between.

The second axis of evaluation is reliability. How many disk faults can the given design tolerate? In alignment with our fault model, we assume only that an entire disk can fail; in later chapters (i.e., on data integrity), we’ll think about how to handle more complex failure modes.

Finally, the third axis is performance. Performance is somewhat chal-
lenging to evaluate, because it depends heavily on the workload presented to the disk array. Thus, before evaluating performance, we will first present a set of typical workloads that one should consider.

We now consider three important RAID designs: RAID Level 0 (striping), RAID Level 1 (mirroring), and RAID Levels 4/5 (parity-based redundancy). The naming of each of these designs as a “level” stems from the pioneering work of Patterson, Gibson, and Katz at Berkeley [P+88].

38.4 RAID Level 0: Striping

The first RAID level is actually not a RAID level at all, in that there is no redundancy. However, RAID level 0, or striping as it is better known, serves as an excellent upper-bound on performance and capacity and thus is worth understanding.

The simplest form of striping will stripe blocks across the disks of the system as follows (assume here a 4-disk array):

<table>
<thead>
<tr>
<th>Disk 0</th>
<th>Disk 1</th>
<th>Disk 2</th>
<th>Disk 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
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<tr>
<td>4</td>
<td>5</td>
<td>6</td>
<td>7</td>
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<tr>
<td>8</td>
<td>9</td>
<td>10</td>
<td>11</td>
</tr>
<tr>
<td>12</td>
<td>13</td>
<td>14</td>
<td>15</td>
</tr>
</tbody>
</table>

Table 38.1: RAID-0: Simple Striping

From Table 38.1, you get the basic idea: spread the blocks of the array across the disks in a round-robin fashion. This approach is designed to extract the most parallelism from the array when requests are made for contiguous chunks of the array (as in a large, sequential read, for example). We call the blocks in the same row a stripe; thus, blocks 0, 1, 2, and 3 are in the same stripe above.

In the example, we have made the simplifying assumption that only 1 block (each of say size 4KB) is placed on each disk before moving on to the next. However, this arrangement need not be the case. For example, we could arrange the blocks across disks as in Table 38.2:

<table>
<thead>
<tr>
<th>Disk 0</th>
<th>Disk 1</th>
<th>Disk 2</th>
<th>Disk 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>2</td>
<td>4</td>
<td>6</td>
</tr>
<tr>
<td>1</td>
<td>3</td>
<td>5</td>
<td>7</td>
</tr>
<tr>
<td>8</td>
<td>10</td>
<td>12</td>
<td>14</td>
</tr>
<tr>
<td>9</td>
<td>11</td>
<td>13</td>
<td>15</td>
</tr>
</tbody>
</table>

Table 38.2: Striping with a Bigger Chunk Size

In this example, we place two 4KB blocks on each disk before moving on to the next disk. Thus, the chunk size of this RAID array is 8KB, and a stripe thus consists of 4 chunks or 32KB of data.
Before studying the capacity, reliability, and performance characteristics of the RAID, we first present an aside on what we call the mapping problem. This problem arises in all RAID arrays; simply put, given a logical block to read or write, how does the RAID know exactly which physical disk and offset to access? For these simple RAID levels, we do not need much sophistication in order to correctly map logical blocks onto their physical locations. Take the first striping example above (chunk size = 1 block = 4KB). In this case, given a logical block address A, the RAID can easily compute the desired disk and offset with two simple equations:

\[
\begin{align*}
\text{Disk} & = A \mod \text{number_of_disks} \\
\text{Offset} & = A / \text{number_of_disks}
\end{align*}
\]

Note that these are all integer operations (e.g., \(4 / 3 = 1\) not \(1.33333\ldots\)).

Let's see how these equations work for a simple example. Imagine in the first RAID above that a request arrives for block 14. Given that there are 4 disks, this would mean that the disk we are interested in is \((14 \mod 4 = 2):\) disk 2. The exact block is calculated as \((14 / 4 = 3):\) block 3. Thus, block 14 should be found on the fourth block (block 3, starting at 0) of the third disk (disk 2, starting at 0), which is exactly where it is.

You can think about how these equations would be modified to support different chunk sizes. Try it! It’s not too hard.

### Chunk Sizes

Chunk size mostly affects performance of the array. For example, a small chunk size implies that many files will get striped across many disks, thus increasing the parallelism of reads and writes to a single file; however, the positioning time to access blocks across multiple disks increases, because the positioning time for the entire request is determined by the maximum of the positioning times of the requests across all drives.

A big chunk size, on the other hand, reduces such intra-file parallelism, and thus relies on multiple concurrent requests to achieve high throughput. However, large chunk sizes reduce positioning time; if, for example, a single file fits within a chunk and thus is placed on a single disk, the positioning time incurred while accessing it will just be the positioning time of a single disk.

Thus, determining the “best” chunk size is hard to do, as it requires a great deal of knowledge about the workload presented to the disk system [CL95]. For the rest of this discussion, we will assume that the array uses a chunk size of a single block (4KB). Most arrays use larger chunk sizes (e.g., 64 KB), but for the issues we discuss below, the exact chunk size does not matter; thus we use a single block for the sake of simplicity.
Back To RAID-0 Analysis

Let us now evaluate the capacity, reliability, and performance of striping. From the perspective of capacity, it is perfect: given N disks, striping delivers N disks worth of useful capacity. From the standpoint of reliability, striping is also perfect, but in the bad way: any disk failure will lead to data loss. Finally, performance is excellent: all disks are utilized, often in parallel, to service user I/O requests.

Evaluating RAID Performance

In analyzing RAID performance, one can consider two different performance metrics. The first is single-request latency. Understanding the latency of a single I/O request to a RAID is useful as it reveals how much parallelism can exist during a single logical I/O operation. The second is steady-state throughput of the RAID, i.e., the total bandwidth of many concurrent requests. Because RAIDs are often used in high-performance environments, the steady-state bandwidth is critical, and thus will be the main focus of our analyses.

To understand throughput in more detail, we need to put forth some workloads of interest. We will assume, for this discussion, that there are two types of workloads: sequential and random. With a sequential workload, we assume that requests to the array come in large contiguous chunks; for example, a request (or series of requests) that accesses 1 MB of data, starting at block (B) and ending at block (B + 1 MB), would be deemed sequential. Sequential workloads are common in many environments (think of searching through a large file for a keyword), and thus are considered important.

For random workloads, we assume that each request is rather small, and that each request is to a different random location on disk. For example, a random stream of requests may first access 4KB at logical address 10, then at logical address 550,000, then at 20,100, and so forth. Some important workloads, such as transactional workloads on a database management system (DBMS), exhibit this type of access pattern, and thus it is considered an important workload.

Of course, real workloads are not so simple, and often have a mix of sequential and random-seeming components as well as behaviors in-between the two. For simplicity, we just consider these two possibilities.

As you can tell, sequential and random workloads will result in widely different performance characteristics from a disk. With sequential access, a disk operates in its most efficient mode, spending little time seeking and waiting for rotation and most of its time transferring data. With random access, just the opposite is true: most time is spent seeking and waiting for rotation and relatively little time is spent transferring data. To capture this difference in our analysis, we will assume that a disk can transfer data at S MB/s under a sequential workload, and R MB/s when under a random workload. In general, S is much greater than R.
To make sure we understand this difference, let’s do a simple exercise. Specifically, let’s calculate $S$ and $R$ given the following disk characteristics. Assume a sequential transfer of size 10 MB on average, and a random transfer of 10 KB on average. Also, assume the following disk characteristics:

- Average seek time: 7 ms
- Average rotational delay: 3 ms
- Transfer rate of disk: 50 MB/s

To compute $S$, we need to first figure out how time is spent in a typical 10 MB transfer. First, we spend 7 ms seeking, and then 3 ms rotating. Finally, transfer begins; 10 MB @ 50 MB/s leads to 1/5th of a second, or 200 ms, spent in transfer. Thus, for each 10 MB request, we spend 210 ms completing the request. To compute $S$, we just need to divide:

$$S = \frac{\text{Amount of Data}}{\text{Time to access}} = \frac{10 \text{ MB}}{210 \text{ ms}} = 47.62 \text{ MB/s}$$

As we can see, because of the large time spent transferring data, $S$ is very near the peak bandwidth of the disk (the seek and rotational costs have been amortized).

We can compute $R$ similarly. Seek and rotation are the same; we then compute the time spent in transfer, which is 10 KB @ 50 MB/s, or 0.195 ms.

$$R = \frac{\text{Amount of Data}}{\text{Time to access}} = \frac{10 \text{ KB}}{0.195 \text{ ms}} = 51.28 \text{ MB/s}$$

As we can see, $R$ is less than 1 MB/s, and $S/R$ is almost 50.

### Back To RAID-0 Analysis, Again

Let’s now evaluate the performance of striping. As we said above, it is generally good. From a latency perspective, for example, the latency of a single-block request should be just about identical to that of a single disk; after all, RAID-0 will simply redirect that request to one of its disks.

From the perspective of steady-state throughput, we’d expect to get the full bandwidth of the system. Thus, throughput equals $N$ (the number of disks) multiplied by $S$ (the sequential bandwidth of a single disk). For a large number of random I/Os, we can again use all of the disks, and thus obtain $N \cdot R$ MB/s. As we will see below, these values are both the simplest to calculate and will serve as an upper bound in comparison with other RAID levels.

### 38.5 RAID Level 1: Mirroring

Our first RAID level beyond striping is known as RAID level 1, or mirroring. With a mirrored system, we simply make more than one copy of each block in the system; each copy should be placed on a separate disk, of course. By doing so, we can tolerate disk failures.
In a typical mirrored system, we will assume that for each logical block, the RAID keeps two physical copies of it. Here is an example:

<table>
<thead>
<tr>
<th>Disk 0</th>
<th>Disk 1</th>
<th>Disk 2</th>
<th>Disk 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>1</td>
<td>1</td>
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<td>2</td>
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<td>4</td>
<td>4</td>
<td>5</td>
<td>5</td>
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<tr>
<td>6</td>
<td>6</td>
<td>7</td>
<td>7</td>
</tr>
</tbody>
</table>

Table 38.3: Simple RAID-1: Mirroring

In the example, disk 0 and disk 1 have identical contents, and disk 2 and disk 3 do as well; the data is striped across these mirror pairs. In fact, you may have noticed that there are a number of different ways to place block copies across the disks. The arrangement above is a common one and is sometimes called RAID-10 or (RAID 1+0) because it uses mirrored pairs (RAID-1) and then stripes (RAID-0) on top of them; another common arrangement is RAID-01 (or RAID 0+1), which contains two large striping (RAID-0) arrays, and then mirrors (RAID-1) on top of them. For now, we will just talk about mirroring assuming the above layout.

When reading a block from a mirrored array, the RAID has a choice: it can read either copy. For example, if a read to logical block 5 is issued to the RAID, it is free to read it from either disk 2 or disk 3. When writing a block, though, no such choice exists: the RAID must update both copies of the data, in order to preserve reliability. Do note, though, that these writes can take place in parallel; for example, a write to logical block 5 could proceed to disks 2 and 3 at the same time.

**RAID-1 Analysis**

Let us assess RAID-1. From a capacity standpoint, RAID-1 is expensive; with the mirroring level = 2, we only obtain half of our peak useful capacity. Thus, with N disks, the useful capacity of mirroring is N/2.

From a reliability standpoint, RAID-1 does well. It can tolerate the failure of any one disk. You may also notice RAID-1 can actually do better than this, with a little luck. Imagine, in the figure above, that disk 0 and disk 2 both failed. In such a situation, there is no data loss! More generally, a mirrored system (with mirroring level of 2) can tolerate 1 disk failure for certain, and up to N/2 failures depending on which disks fail. In practice, we generally don’t like to leave things like this to chance; thus most people consider mirroring to be good for handling a single failure.

Finally, we analyze performance. From the perspective of the latency of a single read request, we can see it is the same as the latency on a single disk; all the RAID-1 does is direct the read to one of its copies. A write is a little different: it requires two physical writes to complete before it is done. These two writes happen in parallel, and thus the time will be roughly equivalent to the time of a single write; however, because the logical write must wait for both physical writes to complete, it suffers the worst-case seek and rotational delay of the two requests, and thus (on average) will be slightly higher than a write to a single disk.
ASIDE: THE RAID CONSISTENT-UPDATE PROBLEM

Before analyzing RAID-1, let us first discuss a problem that arises in any multi-disk RAID system, known as the consistent-update problem [DAA05]. The problem occurs on a write to any RAID that has to update multiple disks during a single logical operation. In this case, let us assume we are considering a mirrored disk array.

Imagine the write is issued to the RAID, and then the RAID decides that it must be written to two disks, disk 0 and disk 1. The RAID then issues the write to disk 0, but just before the RAID can issue the request to disk 1, a power loss (or system crash) occurs. In this unfortunate case, let us assume that the request to disk 0 completed (but clearly the request to disk 1 did not, as it was never issued).

The result of this untimely power loss is that the two copies of the block are now inconsistent; the copy on disk 0 is the new version, and the copy on disk 1 is the old. What we would like to happen is for the state of both disks to change atomically, i.e., either both should end up as the new version or neither.

The general way to solve this problem is to use a write-ahead log of some kind to first record what the RAID is about to do (i.e., update two disks with a certain piece of data) before doing it. By taking this approach, we can ensure that in the presence of a crash, the right thing will happen; by running a recovery procedure that replays all pending transactions to the RAID, we can ensure that no two mirrored copies (in the RAID-1 case) are out of sync.

One last note: because logging to disk on every write is prohibitively expensive, most RAID hardware includes a small amount of non-volatile RAM (e.g., battery-backed) where it performs this type of logging. Thus, consistent update is provided without the high cost of logging to disk.

To analyze steady-state throughput, let us start with the sequential workload. When writing out to disk sequentially, each logical write must result in two physical writes; for example, when we write logical block 0 (in the figure above), the RAID internally would write it to both disk 0 and disk 1. Thus, we can conclude that the maximum bandwidth obtained during sequential writing to a mirrored array is \( \frac{N}{2} \cdot S \), or half the peak bandwidth.

Unfortunately, we obtain the exact same performance during a sequential read. One might think that a sequential read could do better, because it only needs to read one copy of the data, not both. However, let’s use an example to illustrate why this doesn’t help much. Imagine we need to read blocks 0, 1, 2, 3, 4, 5, 6, and 7. Let’s say we issue the read of 0 to disk 0, the read of 1 to disk 2, the read of 2 to disk 1, and the read of 3 to disk 3. We continue by issuing reads to 4, 5, 6, and 7 to disks 0, 2, 1, and 3, respectively. One might naively think that because we are utilizing all disks, we are achieving the full bandwidth of the array.

To see that this is not the case, however, consider the requests a single
First, it gets a request for block 0; then, it gets a request for block 4 (skipping block 2). In fact, each disk receives a request for every other block. While it is rotating over the skipped block, it is not delivering useful bandwidth to the client. Thus, each disk will only deliver half its peak bandwidth. And thus, the sequential read will only obtain a bandwidth of \((\frac{N}{2} \cdot S)\) MB/s.

Random reads are the best case for a mirrored RAID. In this case, we can distribute the reads across all the disks, and thus obtain the full possible bandwidth. Thus, for random reads, RAID-1 delivers \(N \cdot R\) MB/s.

Finally, random writes perform as you might expect: \(\frac{N}{2} \cdot R\) MB/s. Each logical write must turn into two physical writes, and thus while all the disks will be in use, the client will only perceive this as half the available bandwidth. Even though a write to logical block \(X\) turns into two parallel writes to two different physical disks, the bandwidth of many small requests only achieves half of what we saw with striping. As we will soon see, getting half the available bandwidth is actually pretty good!

### 38.6 RAID Level 4: Saving Space With Parity

We now present a different method of adding redundancy to a disk array known as parity. Parity-based approaches attempt to use less capacity and thus overcome the huge space penalty paid by mirrored systems. They do so at a cost, however: performance.

In a five-disk RAID-4 system, we might observe the following layout:

<table>
<thead>
<tr>
<th>Disk 0</th>
<th>Disk 1</th>
<th>Disk 2</th>
<th>Disk 3</th>
<th>Disk 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>P0</td>
</tr>
<tr>
<td>4</td>
<td>5</td>
<td>6</td>
<td>7</td>
<td>P1</td>
</tr>
<tr>
<td>8</td>
<td>9</td>
<td>10</td>
<td>11</td>
<td>P2</td>
</tr>
<tr>
<td>12</td>
<td>13</td>
<td>14</td>
<td>15</td>
<td>P3</td>
</tr>
</tbody>
</table>

As you can see, for each stripe of data, we have added a single parity block that stores the redundant information for that stripe of blocks. For example, parity block P1 has redundant information that it calculated from blocks 4, 5, 6, and 7.

To compute parity, we need to use a mathematical function that enables us to withstand the loss of any one block from our stripe. It turns out the simple function XOR does the trick quite nicely. For a given set of bits, the XOR of all of those bits returns a 0 if there are an even number of 1’s in the bits, and a 1 if there are an odd number of 1’s. For example:

<table>
<thead>
<tr>
<th>C0</th>
<th>C1</th>
<th>C2</th>
<th>C3</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>XOR(0,0,1,1) = 0</td>
</tr>
<tr>
<td>0</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>XOR(0,1,0,0) = 1</td>
</tr>
</tbody>
</table>

In the first row (0,0,1,1), there are two 1’s (C2, C3), and thus XOR of all of those values will be 0 (P); similarly, in the second row there is only
one 1 (C1), and thus the XOR must be 1 (P). You can remember this in a very simple way: that the number of 1’s in any row must be an even (not odd) number; that is the invariant that the RAID must maintain in order for parity to be correct.

From the example above, you might also be able to guess how parity information can be used to recover from a failure. Imagine the column labeled C2 is lost. To figure out what values must have been in the column, we simply have to read in all the other values in that row (including the XOR’d parity bit) and reconstruct the right answer. Specifically, assume the first row’s value in column C2 is lost (it is a 1); by reading the other values in that row (0 from C0, 0 from C1, 1 from C3, and 0 from the parity column P), we get the values 0, 0, 1, and 0. Because we know that XOR keeps an even number of 1’s in each row, we know what the missing data must be: a 1. And that is how reconstruction works in a XOR-based parity scheme! Note also how we compute the reconstructed value: we just XOR the data bits and the parity bits together, in the same way that we calculated the parity in the first place.

Now you might be wondering: we are talking about XORing all of these bits, and yet above we know that the RAID places 4KB (or larger) blocks on each disk; how do we apply XOR to a bunch of blocks to compute the parity? It turns out this is easy as well. Simply perform a bitwise XOR across each bit of the data blocks; put the result of each bitwise XOR into the corresponding bit slot in the parity block. For example, if we had blocks of size 4 bits (yes, this is still quite a bit smaller than a 4KB block, but you get the picture), they might look something like this:

<table>
<thead>
<tr>
<th>Block0</th>
<th>Block1</th>
<th>Block2</th>
<th>Block3</th>
<th>Parity</th>
</tr>
</thead>
<tbody>
<tr>
<td>00</td>
<td>10</td>
<td>11</td>
<td>10</td>
<td>11</td>
</tr>
<tr>
<td>10</td>
<td>01</td>
<td>00</td>
<td>01</td>
<td>10</td>
</tr>
</tbody>
</table>

As you can see from the figure, the parity is computed for each bit of each block and the result placed in the parity block.

**RAID-4 Analysis**

Let us now analyze RAID-4. From a capacity standpoint, RAID-4 uses 1 disk for parity information for every group of disks it is protecting. Thus, our useful capacity for a RAID group is (N-1).

Reliability is also quite easy to understand: RAID-4 tolerates 1 disk failure and no more. If more than one disk is lost, there is simply no way to reconstruct the lost data.

Finally, there is performance. This time, let us start by analyzing steady-state throughput. Sequential read performance can utilize all of the disks except for the parity disk, and thus deliver a peak effective bandwidth of \((N - 1) \cdot 5 \text{ MB/s}\) (an easy case).

To understand the performance of sequential writes, we must first understand how they are done. When writing a big chunk of data to disk,
RAID-4 can perform a simple optimization known as a full-stripe write. For example, imagine the case where the blocks 0, 1, 2, and 3 have been sent to the RAID as part of a write request (Table 38.4).

<table>
<thead>
<tr>
<th>Disk 0</th>
<th>Disk 1</th>
<th>Disk 2</th>
<th>Disk 3</th>
<th>Disk 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>P0</td>
</tr>
<tr>
<td>4</td>
<td>5</td>
<td>6</td>
<td>7</td>
<td>P1</td>
</tr>
<tr>
<td>8</td>
<td>9</td>
<td>10</td>
<td>11</td>
<td>P2</td>
</tr>
<tr>
<td>12</td>
<td>13</td>
<td>14</td>
<td>15</td>
<td>P3</td>
</tr>
</tbody>
</table>

Table 38.4: Full-stripe Writes In RAID-4

In this case, the RAID can simply calculate the new value of P0 (by performing an XOR across the blocks 0, 1, 2, and 3) and then write all of the blocks (including the parity block) to the five disks above in parallel (highlighted in gray in the figure). Thus, full-stripe writes are the most efficient way for RAID-4 to write to disk.

Once we understand the full-stripe write, calculating the performance of sequential writes on RAID-4 is easy; the effective bandwidth is also \((N - 1) \cdot S\) MB/s. Even though the parity disk is constantly in use during the operation, the client does not gain performance advantage from it.

Now let us analyze the performance of random reads. As you can also see from the figure above, a set of 1-block random reads will be spread across the data disks of the system but not the parity disk. Thus, the effective performance is: \((N - 1) \cdot R\) MB/s.

Random writes, which we have saved for last, present the most interesting case for RAID-4. Imagine we wish to overwrite block 1 in the example above. We could just go ahead and overwrite it, but that would leave us with a problem: the parity block P0 would no longer accurately reflect the correct parity value for the stripe. Thus, in this example, P0 must also be updated. But how can we update it both correctly and efficiently?

It turns out there are two methods. The first, known as additive parity, requires us to do the following. To compute the value of the new parity block, read in all of the other data blocks in the stripe in parallel (in the example, blocks 0, 2, and 3) and XOR those with the new block (1). The result is your new parity block. To complete the write, you can then write the new data and new parity to their respective disks, also in parallel.

The problem with this technique is that it scales with the number of disks, and thus in larger RAIDs requires a high number of reads to compute parity. Thus, the subtractive parity method.

For example, imagine this string of bits (4 data bits, one parity):

<table>
<thead>
<tr>
<th>C0</th>
<th>C1</th>
<th>C2</th>
<th>C3</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>XOR(0,0,1,1) = 0</td>
</tr>
</tbody>
</table>

Let’s imagine that we wish to overwrite bit C2 with a new value which we will call C2(new). The subtractive method works in three steps. First, we read in the old data at C2 (C2(old) = 1) and the old parity (P(old) = 0).
0). Then, we compare the old data and the new data; if they are the same (e.g., \( C_2(\text{new}) = C_2(\text{old}) \)), then we know the parity bit will also remain the same (i.e., \( P(\text{new}) = P(\text{old}) \)). If, however, they are different, then we must flip the old parity bit to the opposite of its current state, that is, if \( P(\text{old}) = 1 \), \( P(\text{new}) \) will be set to 0; if \( P(\text{old}) = 0 \), \( P(\text{new}) \) will be set to 1. We can express this whole mess neatly with XOR as it turns out (if you understand XOR, this will now make sense to you):

\[
P(\text{new}) = (C(\text{old}) \oplus C(\text{new})) \oplus P(\text{old})
\]

Because we are dealing with blocks, not bits, we perform this calculation over all the bits in the block (e.g., 4096 bytes in each block multiplied by 8 bits per byte). Thus, in most cases, the new block will be different than the old block and thus the new parity block will too.

You should now be able to figure out when we would use the additive parity calculation and when we would use the subtractive method. Think about how many disks would need to be in the system so that the additive method performs fewer I/Os than the subtractive method; what is the cross-over point?

For this performance analysis, let us assume we are using the subtractive method. Thus, for each write, the RAID has to perform 4 physical I/Os (two reads and two writes). Now imagine there are lots of writes submitted to the RAID; how many can RAID-4 perform in parallel? To understand, let us again look at the RAID-4 layout (Figure 38.5).

<table>
<thead>
<tr>
<th>Disk 0</th>
<th>Disk 1</th>
<th>Disk 2</th>
<th>Disk 3</th>
<th>Disk 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>P0</td>
</tr>
<tr>
<td>+4</td>
<td>5</td>
<td>6</td>
<td>7</td>
<td>+P1</td>
</tr>
<tr>
<td>8</td>
<td>9</td>
<td>10</td>
<td>11</td>
<td>P2</td>
</tr>
<tr>
<td>12</td>
<td>+13</td>
<td>14</td>
<td>15</td>
<td>+P3</td>
</tr>
</tbody>
</table>

Table 38.5: Example: Writes To 4, 13, And Respective Parity Blocks

Now imagine there were 2 small writes submitted to the RAID-4 at about the same time, to blocks 4 and 13 (marked with * in the diagram). The data for those disks is on disks 0 and 1, and thus the read and write to data could happen in parallel, which is good. The problem that arises is with the parity disk; both the requests have to read the related parity blocks for 4 and 13, parity blocks 1 and 3 (marked with +). Hopefully, the issue is now clear: the parity disk is a bottleneck under this type of workload; we sometimes thus call this the small-write problem for parity-based RAIDs. Thus, even though the data disks could be accessed in parallel, the parity disk prevents any parallelism from materializing; all writes to the system will be serialized because of the parity disk. Because the parity disk has to perform two I/Os (one read, one write) per logical I/O, we can compute the performance of small random writes in RAID-4 by computing the parity disk’s performance on those two I/Os, and thus we achieve \((R/2) \text{ MB/s}\). RAID-4 throughput under random small writes is terrible; it does not improve as you add disks to the system.
We conclude by analyzing I/O latency in RAID-4. As you now know, a single read (assuming no failure) is just mapped to a single disk, and thus its latency is equivalent to the latency of a single disk request. The latency of a single write requires two reads and then two writes; the reads can happen in parallel, as can the writes, and thus total latency is about twice that of a single disk (with some differences because we have to wait for both reads to complete and thus get the worst-case positioning time, but then the updates don’t incur seek cost and thus may be a better-than-average positioning cost).

### 38.7 RAID Level 5: Rotating Parity

To address the small-write problem (at least, partially), Patterson, Gibson, and Katz introduced RAID-5. RAID-5 works almost identically to RAID-4, except that it rotates the parity block across drives (Figure 38.6).

#### Table 38.6: RAID-5 With Rotated Parity

<table>
<thead>
<tr>
<th>Disk 0</th>
<th>Disk 1</th>
<th>Disk 2</th>
<th>Disk 3</th>
<th>Disk 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>P0</td>
</tr>
<tr>
<td>5</td>
<td>6</td>
<td>7</td>
<td>P1</td>
<td>4</td>
</tr>
<tr>
<td>10</td>
<td>11</td>
<td>P2</td>
<td>8</td>
<td>9</td>
</tr>
<tr>
<td>15</td>
<td>P3</td>
<td>12</td>
<td>13</td>
<td>14</td>
</tr>
<tr>
<td>P4</td>
<td>16</td>
<td>17</td>
<td>18</td>
<td>19</td>
</tr>
</tbody>
</table>

As you can see, the parity block for each stripe is now rotated across the disks, in order to remove the parity-disk bottleneck for RAID-4.

#### RAID-5 Analysis

Much of the analysis for RAID-5 is identical to RAID-4. For example, the effective capacity and failure tolerance of the two levels are identical. So are sequential read and write performance. The latency of a single request (whether a read or a write) is also the same as RAID-4.

Random read performance is a little better, because we can utilize all of the disks. Finally, random write performance improves noticeably over RAID-4, as it allows for parallelism across requests. Imagine a write to block 1 and a write to block 10; this will turn into requests to disk 1 and disk 4 (for block 1 and its parity) and requests to disk 0 and disk 2 (for block 10 and its parity). Thus, they can proceed in parallel. In fact, we can generally assume that that given a large number of random requests, we will be able to keep all the disks about evenly busy. If that is the case, then our total bandwidth for small writes will be \( N/4 \cdot R \) MB/s. The factor of four loss is due to the fact that each RAID-5 write still generates 4 total I/O operations, which is simply the cost of using parity-based RAID.
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Because RAID-5 is basically identical to RAID-4 except in the few cases where it is better, it has almost completely replaced RAID-4 in the marketplace. The only place where it has not is in systems that know they will never perform anything other than a large write, thus avoiding the small-write problem altogether [HLM94]; in those cases, RAID-4 is sometimes used as it is slightly simpler to build.

### 38.8 RAID Comparison: A Summary

We now summarize our simplified comparison of RAID levels in Table 38.7. Note that we have omitted a number of details to simplify our analysis. For example, when writing in a mirrored system, the average seek time is a little higher than when writing to just a single disk, because the seek time is the max of two seeks (one on each disk). Thus, random write performance to two disks will generally be a little less than random write performance of a single disk. Also, when updating the parity disk in RAID-4/5, the first read of the old parity will likely cause a full seek and rotation, but the second write of the parity will only result in rotation.

However, our comparison does capture the essential differences, and is useful for understanding tradeoffs across RAID levels. We present a summary in the table below; for the latency analysis, we simply use $D$ to represent the time that a request to a single disk would take.

To conclude, if you strictly want performance and do not care about reliability, striping is obviously best. If, however, you want random I/O performance and reliability, mirroring is the best; the cost you pay is in lost capacity. If capacity and reliability are your main goals, then RAID-5 is the winner; the cost you pay is in small-write performance. Finally, if you are always doing sequential I/O and want to maximize capacity, RAID-5 also makes the most sense.

<table>
<thead>
<tr>
<th>Capacity</th>
<th>RAID-0</th>
<th>RAID-1</th>
<th>RAID-4</th>
<th>RAID-5</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$N$</td>
<td>$N/2$</td>
<td>$N-1$</td>
<td>$N-1$</td>
</tr>
<tr>
<td>Reliability</td>
<td>0</td>
<td>1 (for sure)</td>
<td>1</td>
<td>1</td>
</tr>
</tbody>
</table>

\[
\frac{N}{2} \text{ (if lucky)}
\]

<table>
<thead>
<tr>
<th>Throughput</th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Sequential Read</td>
<td>$N \cdot S$</td>
<td>$(N/2) \cdot S$</td>
<td>$(N-1) \cdot S$</td>
<td>$(N-1) \cdot S$</td>
</tr>
<tr>
<td>Sequential Write</td>
<td>$N \cdot S$</td>
<td>$(N/2) \cdot S$</td>
<td>$(N-1) \cdot S$</td>
<td>$(N-1) \cdot S$</td>
</tr>
<tr>
<td>Random Read</td>
<td>$N \cdot R$</td>
<td>$N \cdot R$</td>
<td>$(N-1) \cdot R$</td>
<td>$N \cdot R$</td>
</tr>
<tr>
<td>Random Write</td>
<td>$N \cdot R$</td>
<td>$(N/2) \cdot R$</td>
<td>$\frac{1}{2} \cdot R$</td>
<td>$\frac{N}{T} \cdot R$</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Latency</th>
<th>$D$</th>
<th>$D$</th>
<th>$D$</th>
<th>$D$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Read</td>
<td>$D$</td>
<td>$D$</td>
<td>$2D$</td>
<td>$2D$</td>
</tr>
<tr>
<td>Write</td>
<td>$D$</td>
<td>$D$</td>
<td>$2D$</td>
<td>$2D$</td>
</tr>
</tbody>
</table>

Table 38.7: RAID Capacity, Reliability, and Performance
38.9 Other Interesting RAID Issues

There are a number of other interesting ideas that one could (and perhaps should) discuss when thinking about RAID. Here are some things we might eventually write about.

For example, there are many other RAID designs, including Levels 2 and 3 from the original taxonomy, and Level 6 to tolerate multiple disk faults [C+04]. There is also what the RAID does when a disk fails; sometimes it has a hot spare sitting around to fill in for the failed disk. What happens to performance under failure, and performance during reconstruction of the failed disk? There are also more realistic fault models, to take into account latent sector errors or block corruption [B+08], and lots of techniques to handle such faults (see the data integrity chapter for details). Finally, you can even build raid as a software layer: such software RAID systems are cheaper but have other problems, including the consistent-update problem [DAA05].

38.10 Summary

We have discussed RAID. RAID transforms a number of independent disks into a large, more capacious, and more reliable single entity; importantly, it does so transparently, and thus hardware and software above is relatively oblivious to the change.

There are many possible RAID levels to choose from, and the exact RAID level to use depends heavily on what is important to the end-user. For example, mirrored RAID is simple, reliable, and generally provides good performance but at a high capacity cost. RAID-5, in contrast, is reliable and better from a capacity standpoint, but performs quite poorly when there are small writes in the workload. Picking a RAID and setting its parameters (chunk size, number of disks, etc.) properly for a particular workload is challenging, and remains more of an art than a science.
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References

[B+08] “An Analysis of Data Corruption in the Storage Stack”
Lakshmi N. Bairavasundaram, Garth R. Goodson, Bianca Schroeder, Andrea C. Arpaci-Dusseau, Remzi H. Arpaci-Dusseau
FAST ’08, San Jose, CA, February 2008
Our own work analyzing how often disks actually corrupt your data. Not often, but sometimes! And thus something a reliable storage system must consider.

[BJ88] “Disk Shadowing”
D. Bitton and J. Gray
VLDB 1988
One of the first papers to discuss mirroring, herein called “shadowing”.

[CL95] “Striping in a RAID level 5 disk array”
Peter M. Chen, Edward K. Lee
SIGMETRICS 1995
A nice analysis of some of the important parameters in a RAID-5 disk array.

[C+04] “Row-Diagonal Parity for Double Disk Failure Correction”
P. Corbett, B. English, A. Goel, T. Gracanac, S. Kleiman, J. Leong, S. Sankar
FAST ’04, February 2004
Though not the first paper on a RAID system with two disks for parity, it is a recent and highly-understandable version of said idea. Read it to learn more.

[DAA05] “Journal-guided Resynchronization for Software RAID”
Timothy E. Denehy, A. Arpaci-Dusseau, R. Arpaci-Dusseau
FAST 2005
Our own work on the consistent-update problem. Here we solve it for Software RAID by integrating the journaling machinery of the file system above with the software RAID beneath it.

Dave Hitz, James Lau, Michael Malcolm
USENIX Winter 1994, San Francisco, California, 1994
The sparse paper introducing a landmark product in storage, the write-anywhere file layout or WAFL file system that underlies the NetApp file server.

[K86] “Synchronized Disk Interleaving”
M.Y. Kim.
IEEE Transactions on Computers, Volume C-35: 11, November 1986
Some of the earliest work on RAID is found here.

F. Kurzweil.
Presentation at Spring COMPCON ’88, March 1, 1988, San Francisco, California
Another early RAID reference.

[P+88] “Redundant Arrays of Inexpensive Disks”
D. Patterson, G. Gibson, R. Katz.
SIGMOD 1988
This is considered the RAID paper, written by famous authors Patterson, Gibson, and Katz. The paper has since won many test-of-time awards and ushered in the RAID era, including the name RAID itself!
Another early work on RAID.

And yes, another early RAID work. There are a lot of these, which kind of came out of the woodwork when the RAID paper was published in SIGMOD.

Finally, a paper that is not about RAID! This paper is actually about how systems fail, and how to make something behave in a fail-stop manner.
Homework

This section introduces raid.py, a simple RAID simulator you can use to shore up your knowledge of how RAID systems work. See the README for details.

Questions

1. Use the simulator to perform some basic RAID mapping tests. Run with different levels (0, 1, 4, 5) and see if you can figure out the mappings of a set of requests. For RAID-5, see if you can figure out the difference between left-symmetric and left-asymmetric layouts. Use some different random seeds to generate different problems than above.
2. Do the same as the first problem, but this time vary the chunk size with -C. How does chunk size change the mappings?
3. Do the same as above, but use the -r flag to reverse the nature of each problem.
4. Now use the reverse flag but increase the size of each request with the -S flag. Try specifying sizes of 8k, 12k, and 16k, while varying the RAID level. What happens to the underlying I/O pattern when the size of the request increases? Make sure to try this with the sequential workload too (-W sequential); for what request sizes are RAID-4 and RAID-5 much more I/O efficient?
5. Use the timing mode of the simulator (-t) to estimate the performance of 100 random reads to the RAID, while varying the RAID levels, using 4 disks.
6. Do the same as above, but increase the number of disks. How does the performance of each RAID level scale as the number of disks increases?
7. Do the same as above, but use all writes (-w 100) instead of reads. How does the performance of each RAID level scale now? Can you do a rough estimate of the time it will take to complete the workload of 100 random writes?
8. Run the timing mode one last time, but this time with a sequential workload (-W sequential). How does the performance vary with RAID level, and when doing reads versus writes? How about when varying the size of each request? What size should you write to a RAID when using RAID-4 or RAID-5?
Thus far we have seen the development of two key operating system abstractions: the process, which is a virtualization of the CPU, and the address space, which is a virtualization of memory. In tandem, these two abstractions allow a program to run as if it is in its own private, isolated world; as if it has its own processor (or processors); as if it has its own memory. This illusion makes programming the system much easier and thus is prevalent today not only on desktops and servers but increasingly on all programmable platforms including mobile phones and the like.

In this section, we add one more critical piece to the virtualization puzzle: persistent storage. A persistent-storage device, such as a classic hard disk drive or a more modern solid-state storage device, stores information permanently (or at least, for a long time). Unlike memory, whose contents are lost when there is a power loss, a persistent-storage device keeps such data intact. Thus, the OS must take extra care with such a device: this is where users keep data that they really care about.

**CRUX: HOW TO MANAGE A PERSISTENT DEVICE**

How should the OS manage a persistent device? What are the APIs? What are the important aspects of the implementation?

Thus, in the next few chapters, we will explore critical techniques for managing persistent data, focusing on methods to improve performance and reliability. We begin, however, with an overview of the API: the interfaces you’ll expect to see when interacting with a UNIX file system.

### 39.1 Files and Directories

Two key abstractions have developed over time in the virtualization of storage. The first is the file. A file is simply a linear array of bytes, each of which you can read or write. Each file has some kind of *low-level*
name, usually a number of some kind; often, the user is not aware of this name (as we will see). For historical reasons, the low-level name of a file is often referred to as its inode number. We’ll be learning a lot more about inodes in future chapters; for now, just assume that each file has an inode number associated with it.

In most systems, the OS does not know much about the structure of the file (e.g., whether it is a picture, or a text file, or C code); rather, the responsibility of the file system is simply to store such data persistently on disk and make sure that when you request the data again, you get what you put there in the first place. Doing so is not as simple as it seems!

The second abstraction is that of a directory. A directory, like a file, also has a low-level name (i.e., an inode number), but its contents are quite specific: it contains a list of (user-readable name, low-level name) pairs. For example, let’s say there is a file with the low-level name “10”, and it is referred to by the user-readable name of “foo”. The directory “foo” resides in thus would have an entry (“foo”, “10”) that maps the user-readable name to the low-level name. Each entry in a directory refers to either files or other directories. By placing directories within other directories, users are able to build an arbitrary directory tree (or directory hierarchy), under which all files and directories are stored.

The directory hierarchy starts at a root directory (in UNIX-based systems, the root directory is simply referred to as /) and uses some kind of separator to name subsequent sub-directories until the desired file or directory is named. For example, if a user created a directory foo in the root directory /, and then created a file bar.txt in the directory foo, we could refer to the file by its absolute pathname, which in this case would be /foo/bar.txt. See Figure 39.1 for a more complex directory tree; valid directories in the example are /, /foo, /bar, /bar/bar, /bar/foo and valid files are /foo/bar.txt and /bar/foo/bar.txt. Directories and files can have the same name as long as they are in different locations in the file-system tree (e.g., there are two files named bar.txt in the figure, /foo/bar.txt and /bar/foo/bar.txt).
I N T E R L U D E: F I L E A N D D I R E C T O R I E S

39.2 The File System Interface

Let’s now discuss the file system interface in more detail. We’ll start with the basics of creating, accessing, and deleting files. You may think this straightforward, but along the way we’ll discover the mysterious call that is used to remove files, known as unlink(). Hopefully, by the end of this chapter, this mystery won’t be so mysterious to you!

39.3 Creating Files

We’ll start with the most basic of operations: creating a file. This can be accomplished with the open system call; by calling open() and passing it the O_CREAT flag, a program can create a new file. Here is some example code to create a file called “foo” in the current working directory.

```c
int fd = open("foo", O_CREAT | O_WRONLY | O_TRUNC);
```

The routine open() takes a number of different flags. In this example, the program creates the file (O_CREAT), can only write to that file while opened in this manner (O_WRONLY), and, if the file already exists, first truncate it to a size of zero bytes thus removing any existing content (O_TRUNC).
The older way of creating a file is to call `creat()`, as follows:

```c
int fd = creat("foo");
```

You can think of `creat()` as `open()` with the following flags: `O_CREAT | O_WRONLY | O_TRUNC`. Because `open()` can create a file, the usage of `creat()` has somewhat fallen out of favor (indeed, it could just be implemented as a library call to `open()`); however, it does hold a special place in UNIX lore. Specifically, when Ken Thompson was asked what he would do differently if he were redesigning UNIX, he replied: “I’d spell creat with an e.”

One important aspect of `open()` is what it returns: a file descriptor. A file descriptor is just an integer, private per process, and is used in UNIX systems to access files; thus, once a file is opened, you use the file descriptor to read or write the file, assuming you have permission to do so. In this way, a file descriptor is a capability [L84], i.e., an opaque handle that gives you the power to perform certain operations. Another way to think of a file descriptor is as a pointer to an object of type file; once you have such an object, you can call other “methods” to access the file, like `read()` and `write()`. We’ll see just how a file descriptor is used below.

39.4 Reading and Writing Files

Once we have some files, of course we might like to read or write them. Let’s start by reading an existing file. If we were typing at a command line, we might just use the program `cat` to dump the contents of the file to the screen.

```
prompt> echo hello > foo
prompt> cat foo
hello
prompt>
```

In this code snippet, we redirect the output of the program `echo` to the file `foo`, which then contains the word “hello” in it. We then use `cat` to see the contents of the file. But how does the `cat` program access the file `foo`?

To find this out, we’ll use an incredibly useful tool to trace the system calls made by a program. On Linux, the tool is called `strace`; other systems have similar tools (see `dtruss` on Mac OS X, or `truss` on some older UNIX variants). What `strace` does is trace every system call made by a program while it runs, and dump the trace to the screen for you to see.
TIP: USE STRACE (AND SIMILAR TOOLS)
The strace tool provides an awesome way to see what programs are up to. By running it, you can trace which system calls a program makes, see the arguments and return codes, and generally get a very good idea of what is going on.
The tool also takes some arguments which can be quite useful. For example, -f follows any fork’d children too; -t reports the time of day at each call; -e trace=open,close,read,write only traces calls to those system calls and ignores all others. There are many more powerful flags – read the man pages and find out how to harness this wonderful tool.

Here is an example of using strace to figure out what cat is doing (some calls removed for readability):

```
prompt> strace cat foo
...
open("foo", O_RDONLY|O_LARGEFILE) = 3
read(3, "hello\n", 4096) = 6
write(1, "hello\n", 6) = 6
hello
read(3, "", 4096) = 0
close(3) = 0
...
prompt>
```

The first thing that cat does is open the file for reading. A couple of things we should note about this; first, that the file is only opened for reading (not writing), as indicated by the O_RDONLY flag; second, that the 64-bit offset be used (O_LARGEFILE); third, that the call to open() succeeds and returns a file descriptor, which has the value of 3.

Why does the first call to open() return 3, not 0 or perhaps 1 as you might expect? As it turns out, each running process already has three files open, standard input (which the process can read to receive input), standard output (which the process can write to in order to dump information to the screen), and standard error (which the process can write error messages to). These are represented by file descriptors 0, 1, and 2, respectively. Thus, when you first open another file (as cat does above), it will almost certainly be file descriptor 3.

After the open succeeds, cat uses the read() system call to repeatedly read some bytes from a file. The first argument to read() is the file descriptor, thus telling the file system which file to read; a process can of course have multiple files open at once, and thus the descriptor enables the operating system to know which file a particular read refers to. The second argument points to a buffer where the result of the read() will be placed; in the system-call trace above, strace shows the results of the read in this spot (“hello”). The third argument is the size of the buffer, which
in this case is 4 KB. The call to `read()` returns successfully as well, here returning the number of bytes it read (6, which includes 5 for the letters in the word “hello” and one for an end-of-line marker).

At this point, you see another interesting result of the strace: a single call to the `write()` system call, to the file descriptor 1. As we mentioned above, this descriptor is known as the standard output, and thus is used to write the word “hello” to the screen as the program `cat` is meant to do. But does it call `write()` directly? Maybe (if it is highly optimized). But if not, what `cat` might do is call the library routine `printf()`; internally, `printf()` figures out all the formatting details passed to it, and eventually calls `write()` on the standard output to print the results to the screen.

The `cat` program then tries to read more from the file, but since there are no bytes left in the file, the `read()` returns 0 and the program knows that this means it has read the entire file. Thus, the program calls `close()` to indicate that it is done with the file “foo”, passing in the corresponding file descriptor. The file is thus closed, and the reading of it thus complete.

Writing a file is accomplished via a similar set of steps. First, a file is opened for writing, then the `write()` system call is called, perhaps repeatedly for larger files, and then `close()`. Use `strace` to trace writes to a file, perhaps of a program you wrote yourself, or by tracing the `dd` utility, e.g., `dd if=foo of=bar`.

### 39.5 Reading And Writing, But Not Sequentially

Thus far, we’ve discussed how to read and write files, but all access has been sequential; that is, we have either read a file from the beginning to the end, or written a file out from beginning to end.

Sometimes, however, it is useful to be able to read or write to a specific offset within a file; for example, if you build an index over a text document, and use it to look up a specific word, you may end up reading from some random offsets within the document. To do so, we will use the `lseek()` system call. Here is the function prototype:

```c
off_t lseek(int fildes, off_t offset, int whence);
```

The first argument is familiar (a file descriptor). The second argument is the `offset`, which positions the file offset to a particular location within the file. The third argument, called `whence` for historical reasons, determines exactly how the seek is performed. From the man page:

- If `whence` is SEEK_SET, the offset is set to offset bytes.
- If `whence` is SEEK_CUR, the offset is set to its current location plus offset bytes.
- If `whence` is SEEK_END, the offset is set to the size of the file plus offset bytes.

As you can tell from this description, for each file a process opens, the OS tracks a “current” offset, which determines where the next read or
The poorly-named system call `lseek()` confuses many a student trying to understand disks and how the file systems atop them work. Do not confuse the two! The `lseek()` call simply changes a variable in OS memory that tracks, for a particular process, at which offset to which its next read or write will start. A disk seek occurs when a read or write issued to the disk is not on the same track as the last read or write, and thus necessitates a head movement. Making this even more confusing is the fact that calling `lseek()` to read or write from/to random parts of a file, and then reading/writing to those random parts, will indeed lead to more disk seeks. Thus, calling `lseek()` can certainly lead to a seek in an upcoming read or write, but absolutely does not cause any disk I/O to occur itself.

write will begin reading from or writing to within the file. Thus, part of the abstraction of an open file is that it has a current offset, which is updated in one of two ways. The first is when a read or write of \(N\) bytes takes place, \(N\) is added to the current offset; thus each read or write implicitly updates the offset. The second is explicitly with `lseek()`, which changes the offset as specified above.

Note that this call `lseek()` has nothing to do with the seek operation of a disk, which moves the disk arm. The call to `lseek()` simply changes the value of a variable within the kernel; when the I/O is performed, depending on where the disk head is, the disk may or may not perform an actual seek to fulfill the request.

### 39.6 Writing Immediately with `fsync()`

Most times when a program calls `write()`, it is just telling the file system: please write this data to persistent storage, at some point in the future. The file system, for performance reasons, will buffer such writes in memory for some time (say 5 seconds, or 30); at that later point in time, the write(s) will actually be issued to the storage device. From the perspective of the calling application, writes seem to complete quickly, and only in rare cases (e.g., the machine crashes after the `write()` call but before the write to disk) will data be lost.

However, some applications require something more than this eventual guarantee. For example, in a database management system (DBMS), development of a correct recovery protocol requires the ability to force writes to disk from time to time.

To support these types of applications, most file systems provide some additional control APIs. In the UNIX world, the interface provided to applications is known as `fsync(int fd)`. When a process calls `fsync()` for a particular file descriptor, the file system responds by forcing all dirty (i.e., not yet written) data to disk, for the file referred to by the specified
file descriptor. The \texttt{fsync()} routine returns once all of these writes are complete.

Here is a simple example of how to use \texttt{fsync()}. The code opens the file \texttt{foo}, writes a single chunk of data to it, and then calls \texttt{fsync()} to ensure the writes are forced immediately to disk. Once the \texttt{fsync()} returns, the application can safely move on, knowing that the data has been persisted (if \texttt{fsync()} is correctly implemented, that is).

\begin{verbatim}
int fd = open("foo", O_CREAT | O_WRONLY | O_TRUNC);
assert(fd > -1);
int rc = write(fd, buffer, size);
assert(rc == size);
rc = fsync(fd);
assert(rc == 0);
\end{verbatim}

Interestingly, this sequence does not guarantee everything that you might expect; in some cases, you also need to \texttt{fsync()} the directory that contains the file \texttt{foo}. Adding this step ensures not only that the file itself is on disk, but that the file, if newly created, also is durably a part of the directory. Not surprisingly, this type of detail is often overlooked, leading to many application-level bugs [P+13].

\subsection*{39.7 Renaming Files}

Once we have a file, it is sometimes useful to be able to give a file a different name. When typing at the command line, this is accomplished with \texttt{mv} command; in this example, the file \texttt{foo} is renamed \texttt{bar}:

\begin{verbatim}
prompt> mv foo bar
\end{verbatim}

Using \texttt{strace}, we can see that \texttt{mv} uses the system call \texttt{rename(char *old, char *new)}, which takes precisely two arguments: the original name of the file (old) and the new name (new).

One interesting guarantee provided by the \texttt{rename()} call is that it is (usually) implemented as an atomic call with respect to system crashes; if the system crashes during the renaming, the file will either be named the old name or the new name, and no odd in-between state can arise. Thus, \texttt{rename()} is critical for supporting certain kinds of applications that require an atomic update to file state.

Let’s be a little more specific here. Imagine that you are using a file editor (e.g., emacs), and you insert a line into the middle of a file. The file’s name, for the example, is \texttt{foo.txt}. The way the editor might update the file to guarantee that the new file has the original contents plus the line inserted is as follows (ignoring error-checking for simplicity):

\begin{verbatim}
int fd = open("foo.txt.tmp", O_WRONLY|O_CREAT|O_TRUNC);
write(fd, buffer, size); // write out new version of file
fsync(fd);
close(fd);
rename("foo.txt.tmp", "foo.txt");
\end{verbatim}
What the editor does in this example is simple: write out the new version of the file under temporary name (foot.txt.tmp), force it to disk with \texttt{fsync()}, and then, when the application is certain the new file metadata and contents are on the disk, rename the temporary file to the original file’s name. This last step atomically swaps the new file into place, while concurrently deleting the old version of the file, and thus an atomic file update is achieved.

### 39.8 Getting Information About Files

Beyond file access, we expect the file system to keep a fair amount of information about each file it is storing. We generally call such data about files \textit{metadata}. To see the metadata for a certain file, we can use \texttt{stat()} or \texttt{fstat()} system call – read their man pages for details on how to call them. These calls take a pathname (or file descriptor) to a file and fill in a \texttt{stat} structure as seen here:

```c
struct stat {
  dev_t    st_dev; /* ID of device containing file */
  ino_t    st_ino; /* inode number */
  mode_t   st_mode; /* protection */
  nlink_t  st_nlink; /* number of hard links */
  uid_t    st_uid; /* user ID of owner */
  gid_t    st_gid; /* group ID of owner */
  dev_t    st_rdev; /* device ID (if special file) */
  off_t    st_size; /* total size, in bytes */
  blksize_t st_blksize; /* blocksize for filesystem I/O */
  blkcnt_t st_blocks; /* number of blocks allocated */
  time_t   st_atime; /* time of last access */
  time_t   st_mtime; /* time of last modification */
  time_t   st_ctime; /* time of last status change */
};
```

You can see that there is a lot of information kept about each file, including its size (in bytes), its low-level name (i.e., inode number), some ownership information, and some information about when the file was accessed or modified, among other things. To see this information, you can use the command line tool \texttt{stat}:

```
prompt> echo hello > file
prompt> stat file
File: ‘file’
  Size: 6 Blocks: 8     IO Block: 4096  regular file
Device: 811h/2065d Inode: 67158084   Links: 1
Access: (0640/-rw-r-----) Uid: (30686/ remzi) Gid: (30686/ remzi)
```

As it turns out, each file system usually keeps this type of information...
in a structure called an **inode**\(^1\). We’ll be learning a lot more about inodes when we talk about file system implementation. For now, you should just think of an inode as a persistent data structure kept by the file system that has information like we see above inside of it.

### 39.9 Removing Files

At this point, we know how to create files and access them, either sequentially or not. But how do you delete files? If you’ve used **UNIX**, you probably think you know: just run the program `rm`. But what system call does `rm` use to remove a file?

Let’s use our old friend `strace` again to find out. Here we remove that pesky file “foo”:

```
prompt> strace rm foo
...
unlink("foo") = 0
...
```

We’ve removed a bunch of unrelated cruft from the traced output, leaving just a single call to the mysteriously-named system call `unlink()`. As you can see, `unlink()` just takes the name of the file to be removed, and returns zero upon success. But this leads us to a great puzzle: why is this system call named “unlink”? Why not just “remove” or “delete”?

To understand the answer to this puzzle, we must first understand more than just files, but also directories.

### 39.10 Making Directories

Beyond files, a set of directory-related system calls enable you to make, read, and delete directories. Note you can never write to a directory directly; because the format of the directory is considered file system metadata, you can only update a directory indirectly by, for example, creating files, directories, or other object types within it. In this way, the file system makes sure that the contents of the directory always are as expected.

To create a directory, a single system call, `mkdir()`, is available. The eponymous `mkdir` program can be used to create such a directory. Let’s take a look at what happens when we run the `mkdir` program to make a simple directory called `foo`:

```
prompt> strace mkdir foo
...
mkdir("foo", 0777) = 0
...
prompt>
```

\(^1\)Some file systems call these structures similar, but slightly different, names, such as dnodes; the basic idea is similar however.
The program `rm` provides us with a great example of powerful commands, and how sometimes too much power can be a bad thing. For example, to remove a bunch of files at once, you can type something like:

```
prompt> rm *
```

where the `*` will match all files in the current directory. But sometimes you want to also delete the directories too, and in fact all of their contents. You can do this by telling `rm` to recursively descend into each directory, and remove its contents too:

```
prompt> rm -rf *
```

Where you get into trouble with this small string of characters is when you issue the command, accidentally, from the root directory of a file system, thus removing every file and directory from it. Oops!

Thus, remember the double-edged sword of powerful commands; while they give you the ability to do a lot of work with a small number of keystrokes, they also can quickly and readily do a great deal of harm.

When such a directory is created, it is considered "empty", although it does have a bare minimum of contents. Specifically, an empty directory has two entries: one entry that refers to itself, and one entry that refers to its parent. The former is referred to as the "." (dot) directory, and the latter as ".." (dot-dot). You can see these directories by passing a flag (`-a`) to the program `ls`:

```
prompt> ls -a
./ ../
prompt> ls -al
```

```
total 8
drwxr-x--- 2 remzi remzi 6 Apr 30 16:17 ./
```

```
```

```
drwxr-x--- 26 remzi remzi 4096 Apr 30 16:17 ../
```

### 39.11 Reading Directories

Now that we’ve created a directory, we might wish to read one too. Indeed, that is exactly what the program `ls` does. Let’s write our own little tool like `ls` and see how it is done.

Instead of just opening a directory as if it were a file, we instead use a new set of calls. Below is an example program that prints the contents of a directory. The program uses three calls, `opendir()`, `readdir()`, and `closedir()`, to get the job done, and you can see how simple the interface is; we just use a simple loop to read one directory entry at a time, and print out the name and inode number of each file in the directory.
int main(int argc, char *argv[]) {
    DIR *dp = opendir(".");
    assert(dp != NULL);
    struct dirent *d;
    while ((d = readdir(dp)) != NULL) {
        printf("%d %s\n", (int) d->d_ino, d->d_name);
    }
    closedir(dp);
    return 0;
}

The declaration below shows the information available within each directory entry in the struct dirent data structure:

```
struct dirent {
    char d_name[256]; /* filename */
    ino_t d_ino; /* inode number */
    off_t d_off; /* offset to the next dirent */
    unsigned short d_reclen; /* length of this record */
    unsigned char d_type; /* type of file */
};
```

Because directories are light on information (basically, just mapping the name to the inode number, along with a few other details), a program may want to call stat() on each file to get more information on each, such as its length or other detailed information. Indeed, this is exactly what ls does when you pass it the -l flag; try strace on ls with and without that flag to see for yourself.

### 39.12 Deleting Directories

Finally, you can delete a directory with a call to rmdir() (which is used by the program of the same name, rmdir). Unlike file deletion, however, removing directories is more dangerous, as you could potentially delete a large amount of data with a single command. Thus, rmdir() has the requirement that the directory be empty (i.e., only has “.” and “..” entries) before it is deleted. If you try to delete a non-empty directory, the call to rmdir() simply will fail.

### 39.13 Hard Links

We now come back to the mystery of why removing a file is performed via unlink(), by understanding a new way to make an entry in the file system tree, through a system call known as link(). The link() system call takes two arguments, an old pathname and a new one; when you “link” a new file name to an old one, you essentially create another way to refer to the same file. The command-line program ln is used to do this, as we see in this example:
Here we created a file with the word “hello” in it, and called the file file2. We then create a hard link to that file using the ln program. After this, we can examine the file by either opening file or file2.

The way link works is that it simply creates another name in the directory you are creating the link to, and refers it to the same inode number (i.e., low-level name) of the original file. The file is not copied in any way; rather, you now just have two human names (file and file2) that both refer to the same file. We can even see this in the directory itself, by printing out the inode number of each file:

```
prompt> ls -i file file2
67158084 file
67158084 file2
prompt>
```

By passing the -i flag to ls, it prints out the inode number of each file (as well as the file name). And thus you can see what link really has done: just make a new reference to the same exact inode number (67158084 in this example).

By now you might be starting to see why unlink() is called unlink(). When you create a file, you are really doing two things. First, you are making a structure (the inode) that will track virtually all relevant information about the file, including its size, where its blocks are on disk, and so forth. Second, you are linking a human-readable name to that file, and putting that link into a directory.

After creating a hard link to a file, to the file system, there is no difference between the original file name (file) and the newly created file name (file2); indeed, they are both just links to the underlying metadata about the file, which is found in inode number 67158084.

Thus, to remove a file from the file system, we call unlink(). In the example above, we could for example remove the file named file, and still access the file without difficulty:

```
prompt> rm file
removed 'file'
prompt> cat file2
hello
```

The reason this works is because when the file system unlinks file, it checks a reference count within the inode number. This reference count

---

2Note how creative the authors of this book are. We also used to have a cat named “Cat” (true story). However, she died, and we now have a hamster named “Hammy.”
(sometimes called the **link count**) allows the file system to track how many different file names have been linked to this particular inode. When `unlink()` is called, it removes the “link” between the human-readable name (the file that is being deleted) to the given inode number, and decrements the reference count; only when the reference count reaches zero does the file system also free the inode and related data blocks, and thus truly “delete” the file.

You can see the reference count of a file using `stat()` of course. Let’s see what it is when we create and delete hard links to a file. In this example, we’ll create three links to the same file, and then delete them. Watch the link count!

```
prompt> echo hello > file
prompt> stat file
... Inode: 67158084  Links: 1 ...
prompt> ln file file2
prompt> stat file
... Inode: 67158084  Links: 2 ...
prompt> ln file2 file3
prompt> stat file
... Inode: 67158084  Links: 3 ...
prompt> rm file
prompt> rm file2
... Inode: 67158084  Links: 2 ...
prompt> rm file2
prompt> rm file3
... Inode: 67158084  Links: 1 ...
prompt> rm file3
```

### 39.14 Symbolic Links

There is one other type of link that is really useful, and it is called a **symbolic link** or sometimes a **soft link**. As it turns out, hard links are somewhat limited: you can’t create one to a directory (for fear that you will create a cycle in the directory tree); you can’t hard link to files in other disk partitions (because inode numbers are only unique within a particular file system, not across file systems); etc. Thus, a new type of link called the symbolic link was created.

To create such a link, you can use the same program `ln`, but with the `-s` flag. Here is an example:

```
prompt> echo hello > file
prompt> ln -s file file2
prompt> cat file2
hello
```

As you can see, creating a soft link looks much the same, and the original file can now be accessed through the file name `file` as well as the symbolic link name `file2`. 
However, beyond this surface similarity, symbolic links are actually quite different from hard links. The first difference is that a symbolic link is actually a file itself, of a different type. We’ve already talked about regular files and directories; symbolic links are a third type the file system knows about. A `stat` on the symlink reveals all:

```bash
prompt> stat file
... regular file ...
prompt> stat file2
... symbolic link ...
```

Running `ls` also reveals this fact. If you look closely at the first character of the long-form of the output from `ls`, you can see that the first character in the left-most column is `d` for regular files, `d` for directories, and `l` for soft links. You can also see the size of the symbolic link (4 bytes in this case), as well as what the link points to (the file named `file`).

```bash
prompt> ls -al
```

```
drwxr-x--- 2 remzi remzi 29 May 3 19:10 ./
drwxr-x--- 27 remzi remzi 4096 May 3 15:14 ../
-rw-r----- 1 remzi remzi 6 May 3 19:10 file
lrwxrwxrwx 1 remzi remzi 4 May 3 19:10 file2 -> file
```

The reason that `file2` is 4 bytes is because the way a symbolic link is formed is by holding the pathname of the linked-to file as the data of the link file. Because we’ve linked to a file named `file`, our link file `file2` is small (4 bytes). If we link to a longer pathname, our link file would be bigger:

```bash
prompt> echo hello > alongerfilename
prompt> ln -s alongerfilename file3
prompt> ls -al alongerfilename file3
```

```
-rw-r----- 1 remzi remzi 6 May 3 19:17 alongerfilename
lrwxrwxrwx 1 remzi remzi 15 May 3 19:17 file3 -> alongerfilename
```

Finally, because of the way symbolic links are created, they leave the possibility for what is known as a **dangling reference**:

```bash
prompt> echo hello > file
prompt> ln -s file file2
prompt> cat file2
hello
prompt> rm file
prompt> cat file2
```

```
cat: file2: No such file or directory
```

As you can see in this example, quite unlike hard links, removing the original file named `file` causes the link to point to a pathname that no longer exists.
39.15 Making and Mounting a File System

We’ve now toured the basic interfaces to access files, directories, and certain types of special types of links. But there is one more topic we should discuss: how to assemble a full directory tree from many underlying file systems. This task is accomplished via first making file systems, and then mounting them to make their contents accessible.

To make a file system, most file systems provide a tool, usually referred to as mkfs (pronounced “make Fs”), that performs exactly this task. The idea is as follows: give the tool, as input, a device (such as a disk partition, e.g., /dev/sda1) a file system type (e.g., ext3), and it simply writes an empty file system, starting with a root directory, onto that disk partition. And mkfs said, let there be a file system!

However, once such a file system is created, it needs to be made accessible within the uniform file-system tree. This task is achieved via the mount program (which makes the underlying system call mount() to do the real work). What mount does, quite simply is take an existing directory as a target mount point and essentially paste a new file system onto the directory tree at that point.

An example here might be useful. Imagine we have an unmounted ext3 file system, stored in device partition /dev/sda1, that has the following contents: a root directory which contains two sub-directories, a and b, each of which in turn holds a single file named foo. Let’s say we wish to mount this file system at the mount point /home/users. We would type something like this:

```
prompt> mount -t ext3 /dev/sda1 /home/users
```

If successful, the mount would thus make this new file system available. However, note how the new file system is now accessed. To look at the contents of the root directory, we would use ls like this:

```
prompt> ls /home/users/
 a b
```

As you can see, the pathname /home/users/ now refers to the root of the newly-mounted directory. Similarly, we could access files a and b with the pathnames /home/users/a and /home/users/b. Finally, the files named foo could be accessed via /home/users/a/foo and /home/users/b/foo. And thus the beauty of mount: instead of having a number of separate file systems, mount unifies all file systems into one tree, making naming uniform and convenient.

To see what is mounted on your system, and at which points, simply run the mount program. You’ll see something like this:

```
/dev/sda1 on / type ext3 (rw)
 proc on /proc type proc (rw)
 sysfs on /sys type sysfs (rw)
 /dev/sda5 on /tmp type ext3 (rw)
 /dev/sda7 on /var/vice/cache type ext3 (rw)
 tmpfs on /dev/shm type tmpfs (rw)
 AFS on /afs type afs (rw)
```
This crazy mix shows that a whole number of different file systems, including ext3 (a standard disk-based file system), the proc file system (a file system for accessing information about current processes), tmpfs (a file system just for temporary files), and AFS (a distributed file system) are all glued together onto this one machine’s file-system tree.

39.16 Summary

The file system interface in Unix systems (and indeed, in any system) is seemingly quite rudimentary, but there is a lot to understand if you wish to master it. Nothing is better, of course, than simply using it (a lot). So please do so! Of course, read more; as always, Stevens [SR05] is the place to begin.

We’ve toured the basic interfaces, and hopefully understood a little bit about how they work. Even more interesting is how to implement a file system that meets the needs of the API, a topic we will delve into in great detail next.
References

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pseudo file system. A clever idea that you can still see in modern UNIX systems.

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and Remzi H. Arpaci-Dusseau
HotDep ’13, November 2013
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work correctly only if they are running on a specific file system.

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This tour de force of systems is a must-read for anybody interested in the field. It’s how they teach
systems at MIT. Read it once, and then read it a few more times to let it all soak in.

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We have probably referenced this book a few hundred thousand times. It is that useful to you, if you care
to become an awesome systems programmer.
Homework

In this homework, we’ll just familiarize ourselves with how the APIs described in the chapter work. To do so, you’ll just write a few different programs, mostly based on various UNIX utilities.

Questions

1. **Stat**: Write your own version of the command line program `stat`, which simply calls the `stat()` system call on a given file or directory. Print out file size, number of blocks allocated, reference (link) count, and so forth. What is the link count of a directory, as the number of entries in the directory changes? Useful interfaces: `stat()`

2. **List Files**: Write a program that lists files in the given directory. When called without any arguments, the program should just print the file names. When invoked with the `-l` flag, the program should print out information about each file, such as the owner, group, permissions, and other information obtained from the `stat()` system call. The program should take one additional argument, which is the directory to read, e.g., `myls -l directory`. If no directory is given, the program should just use the current working directory. Useful interfaces: `stat()`, `opendir()`, `readdir()`, `getcwd()`.

3. **Tail**: Write a program that prints out the last few lines of a file. The program should be efficient, in that it seeks to near the end of the file, reads in a block of data, and then goes backwards until it finds the requested number of lines; at this point, it should print out those lines from beginning to the end of the file. To invoke the program, one should type: `mytail -n file`, where `n` is the number of lines at the end of the file to print. Useful interfaces: `stat()`, `lseek()`, `open()`, `read()`, `close()`.

4. **Recursive Search**: Write a program that prints out the names of each file and directory in the file system tree, starting at a given point in the tree. For example, when run without arguments, the program should start with the current working directory and print its contents, as well as the contents of any sub-directories, etc., until the entire tree, root at the CWD, is printed. If given a single argument (of a directory name), use that as the root of the tree instead. Refine your recursive search with more fun options, similar to the powerful `find` command line tool. Useful interfaces: you figure it out.
In this chapter, we introduce a simple file system implementation, known as vsfs (the Very Simple File System). This file system is a simplified version of a typical UNIX file system and thus serves to introduce some of the basic on-disk structures, access methods, and various policies that you will find in many file systems today.

The file system is pure software; unlike our development of CPU and memory virtualization, we will not be adding hardware features to make some aspect of the file system work better (though we will want to pay attention to device characteristics to make sure the file system works well). Because of the great flexibility we have in building a file system, many different ones have been built, literally from AFS (the Andrew File System) [H+88] to ZFS (Sun’s Zettabyte File System) [B07]. All of these file systems have different data structures and do some things better or worse than their peers. Thus, the way we will be learning about file systems is through case studies: first, a simple file system (vsfs) in this chapter to introduce most concepts, and then a series of studies of real file systems to understand how they can differ in practice.

**The Crux: How To Implement A Simple File System**
How can we build a simple file system? What structures are needed on the disk? What do they need to track? How are they accessed?

40.1 The Way To Think

To think about file systems, we usually suggest thinking about two different aspects of them; if you understand both of these aspects, you probably understand how the file system basically works.

The first is the data structures of the file system. In other words, what types of on-disk structures are utilized by the file system to organize its data and metadata? The first file systems we’ll see (including vsfs below) employ simple structures, like arrays of blocks or other objects, whereas
As we've discussed before, mental models are what you are really trying to develop when learning about systems. For file systems, your mental model should eventually include answers to questions like: what on-disk structures store the file system's data and metadata? What happens when a process opens a file? Which on-disk structures are accessed during a read or write? By working on and improving your mental model, you develop an abstract understanding of what is going on, instead of just trying to understand the specifics of some file-system code (though that is also useful, of course!).

more sophisticated file systems, like SGI’s XFS, use more complicated tree-based structures [S+96].

The second aspect of a file system is its access methods. How does it map the calls made by a process, such as `open()`, `read()`, `write()`, etc., onto its structures? Which structures are read during the execution of a particular system call? Which are written? How efficiently are all of these steps performed?

If you understand the data structures and access methods of a file system, you have developed a good mental model of how it truly works, a key part of the systems mindset. Try to work on developing your mental model as we delve into our first implementation.

40.2 Overall Organization

We now develop the overall on-disk organization of the data structures of the vsfs file system. The first thing we’ll need to do is divide the disk into blocks; simple file systems use just one block size, and that’s exactly what we’ll do here. Let’s choose a commonly-used size of 4 KB. Thus, our view of the disk partition where we’re building our file system is simple: a series of blocks, each of size 4 KB. The blocks are addressed from 0 to \( N - 1 \), in a partition of size \( N \) 4-KB blocks. Assume we have a really small disk, with just 64 blocks:

```
  0  1  2  3  4  5  6  7
  8  9 10 11 12 13 14 15
 16 17 18 19 20 21 22 23
 24 25 26 27 28 29 30 31

 32 33 34 35 36 37 38 39
 40 41 42 43 44 45 46 47
 48 49 50 51 52 53 54 55
 56 57 58 59 60 61 62 63
```

Let’s now think about what we need to store in these blocks to build a file system. Of course, the first thing that comes to mind is user data. In fact, most of the space in any file system is (and should be) user data. Let’s call the region of the disk we use for user data the data region, and,
again for simplicity, reserve a fixed portion of the disk for these blocks, say the last 56 of 64 blocks on the disk:

As we learned about (a little) last chapter, the file system has to track information about each file. This information is a key piece of metadata, and tracks things like which data blocks (in the data region) comprise a file, the size of the file, its owner and access rights, access and modify times, and other similar kinds of information. To store this information, file system usually have a structure called an inode (we’ll read more about inodes below).

To accommodate inodes, we’ll need to reserve some space on the disk for them as well. Let’s call this portion of the disk the inode table, which simply holds an array of on-disk inodes. Thus, our on-disk image now looks like this picture, assuming that we use 5 of our 64 blocks for inodes (denoted by I’s in the diagram):

We should note here that inodes are typically not that big, for example 128 or 256 bytes. Assuming 256 bytes per inode, a 4-KB block can hold 16 inodes, and our file system above contains 80 total inodes. In our simple file system, built on a tiny 64-block partition, this number represents the maximum number of files we can have in our file system; however, do note that the same file system, built on a larger disk, could simply allocate a larger inode table and thus accommodate more files.

Our file system thus far has data blocks (D), and inodes (I), but a few things are still missing. One primary component that is still needed, as you might have guessed, is some way to track whether inodes or data blocks are free or allocated. Such allocation structures are thus a requisite element in any file system.

Many allocation-tracking methods are possible, of course. For example, we could use a free list that points to the first free block, which then points to the next free block, and so forth. We instead choose a simple and popular structure known as a bitmap, one for the data region (the data bitmap), and one for the inode table (the inode bitmap). A bitmap is a
simple structure: each bit is used to indicate whether the corresponding object/block is free (0) or in-use (1). And thus our new on-disk layout, with an inode bitmap (i) and a data bitmap (d):

```
   Inodes       Data Region
   0 1 0 1 1 1 1 1 0 0 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
   7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 31
```

You may notice that it is a bit of overkill to use an entire 4-KB block for these bitmaps; such a bitmap can track whether 32K objects are allocated, and yet we only have 80 inodes and 56 data blocks. However, we just use an entire 4-KB block for each of these bitmaps for simplicity.

The careful reader (i.e., the reader who is still awake) may have noticed there is one block left in the design of the on-disk structure of our very simple file system. We reserve this for the superblock, denoted by an S in the diagram below. The superblock contains information about this particular file system, including, for example, how many inodes and data blocks are in the file system (80 and 56, respectively in this instance), where the inode table begins (block 3), and so forth. It will likely also include a magic number of some kind to identify the file system type (in this case, vsfs).

```
   Inodes       Data Region
   S 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
   0 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 31
```

Thus, when mounting a file system, the operating system will read the superblock first, to initialize various parameters, and then attach the volume to the file-system tree. When files within the volume are accessed, the system will thus know exactly where to look for the needed on-disk structures.

40.3 File Organization: The Inode

One of the most important on-disk structures of a file system is the inode; virtually all file systems have a structure similar to this. The name inode is short for index node, the historical name given to it by UNIX inventor Ken Thompson [RT74], used because these nodes were originally arranged in an array, and the array indexed into when accessing a particular inode.
The **inode** is the generic name that is used in many file systems to describe the structure that holds the metadata for a given file, such as its length, permissions, and the location of its constituent blocks. The name goes back at least as far as **UNIX** (and probably further back to Multics if not earlier systems); it is short for **index node**, as the inode number is used to index into an array of on-disk inodes in order to find the inode of that number. As we’ll see, design of the inode is one key part of file system design. Most modern systems have some kind of structure like this for every file they track, but perhaps call them different things (such as dnodes, fnodes, etc.).

Each inode is implicitly referred to by a number (called the **inumber**), which we’ve earlier called the **low-level name** of the file. In vsfs (and other simple file systems), given an i-number, you should directly be able to calculate where on the disk the corresponding inode is located. For example, take the inode table of vsfs as above: 20-KB in size (5 4-KB blocks) and thus consisting of 80 inodes (assuming each inode is 256 bytes); further assume that the inode region starts at 12KB (i.e., the superblock starts at 0KB, the inode bitmap is at address 4KB, the data bitmap at 8 KB, and thus the inode table comes right after). In vsfs, we thus have the following layout for the beginning of the file system partition (in closeup view):

![The Inode Table (Closeup)](image)

To read inode number 32, the file system would first calculate the offset into the inode region (32-**sizeof(inode)**) or 8192, add it to the start address of the inode table on disk (**inodeStartAddr** = 12KB), and thus arrive upon the correct byte address of the desired block of inodes: 20KB. Recall that disks are not byte addressable, but rather consist of a large number of addressable sectors, usually 512 bytes. Thus, to fetch the block of inodes that contains inode 32, the file system would issue a read to sector **20 × 1024 / 512**, or 40, to fetch the desired inode block. More generally, the sector address (**iaddr**) of the inode block can be calculated as follows:

\[
\text{blk} = \frac{(\text{inumber} \times \text{sizeof(inode_t)})}{\text{blockSize}}; \\
\text{sector} = \frac{(\text{blk} \times \text{blockSize}) + \text{inodeStartAddr}}{\text{sectorSize}};
\]

Inside each inode is virtually all of the information you need about a file: its **type** (e.g., regular file, directory, etc.), its **size**, the number of **blocks**...
Table 40.1: The ext2 inode

<table>
<thead>
<tr>
<th>Size</th>
<th>Name</th>
<th>What is this inode field for?</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>mode</td>
<td>can this file be read/written/executed?</td>
</tr>
<tr>
<td>2</td>
<td>uid</td>
<td>who owns this file?</td>
</tr>
<tr>
<td>4</td>
<td>size</td>
<td>how many bytes are in this file?</td>
</tr>
<tr>
<td>4</td>
<td>time</td>
<td>what time was this file last accessed?</td>
</tr>
<tr>
<td>4</td>
<td>ctime</td>
<td>what time was this file created?</td>
</tr>
<tr>
<td>4</td>
<td>mtime</td>
<td>what time was this file last modified?</td>
</tr>
<tr>
<td>4</td>
<td>dtime</td>
<td>what time was this inode deleted?</td>
</tr>
<tr>
<td>2</td>
<td>gid</td>
<td>which group does this file belong to?</td>
</tr>
<tr>
<td>2</td>
<td>links_count</td>
<td>how many hard links are there to this file?</td>
</tr>
<tr>
<td>4</td>
<td>blocks</td>
<td>how many blocks have been allocated to this file?</td>
</tr>
<tr>
<td>4</td>
<td>flags</td>
<td>how should ext2 use this inode?</td>
</tr>
<tr>
<td>4</td>
<td>osd1</td>
<td>an OS-dependent field</td>
</tr>
<tr>
<td>60</td>
<td>block</td>
<td>a set of disk pointers (15 total)</td>
</tr>
<tr>
<td>4</td>
<td>generation</td>
<td>file version (used by NFS)</td>
</tr>
<tr>
<td>4</td>
<td>file_acl</td>
<td>a new permissions model beyond mode bits</td>
</tr>
<tr>
<td>4</td>
<td>dir_acl</td>
<td>called access control lists</td>
</tr>
<tr>
<td>4</td>
<td>faddr</td>
<td>an unsupported field</td>
</tr>
<tr>
<td>12</td>
<td>i_osd2</td>
<td>another OS-dependent field</td>
</tr>
</tbody>
</table>

allocated to it, protection information (such as who owns the file, as well as who can access it), some time information, including when the file was created, modified, or last accessed, as well as information about where its data blocks reside on disk (e.g., pointers of some kind). We refer to all such information about a file as metadata; in fact, any information inside the file system that isn’t pure user data is often referred to as such. An example inode from ext2 [P09] is shown below in Table 40.1.

One of the most important decisions in the design of the inode is how it refers to where data blocks are. One simple approach would be to have one or more direct pointers (disk addresses) inside the inode; each pointer refers to one disk block that belongs to the file. Such an approach is limited: for example, if you want to have a file that is really big (e.g., bigger than the size of a block multiplied by the number of direct pointers), you are out of luck.

The Multi-Level Index

To support bigger files, file system designers have had to introduce different structures within inodes. One common idea is to have a special pointer known as an indirect pointer. Instead of pointing to a block that contains user data, it points to a block that contains more pointers, each of which point to user data. Thus, an inode may have some fixed number of direct pointers (e.g., 12), and a single indirect pointer. If a file grows large enough, an indirect block is allocated (from the data-block region of the disk), and the inode’s slot for an indirect pointer is set to point to it. Assuming that a block is 4KB and 4-byte disk addresses, that adds another 1024 pointers; the file can grow to be \((12 + 1024) \cdot 4K = 4144KB\).
A different approach is to use **extents** instead of pointers. An extent is simply a disk pointer plus a length (in blocks); thus, instead of requiring a pointer for every block of a file, all one needs is a pointer and a length to specify the on-disk location of a file. Just a single extent is limiting, as one may have trouble finding a contiguous chunk of on-disk free space when allocating a file. Thus, extent-based file systems often allow for more than one extent, thus giving more freedom to the file system during file allocation.

In comparing the two approaches, pointer-based approaches are the most flexible but use a large amount of metadata per file (particularly for large files). Extent-based approaches are less flexible but more compact; in particular, they work well when there is enough free space on the disk and files can be laid out contiguously (which is the goal for virtually any file allocation policy anyhow).

Not surprisingly, in such an approach, you might want to support even larger files. To do so, just add another pointer to the inode: the **double indirect pointer**. This pointer refers to a block that contains pointers to indirect blocks, each of which contain pointers to data blocks. A double indirect block thus adds the possibility to grow files with an additional \(1024 \cdot 1024\) or 1-million 4KB blocks, in other words supporting files that are over 4GB in size. You may want even more, though, and we bet you know where this is headed: the **triple indirect pointer**.

Overall, this imbalanced tree is referred to as the **multi-level index** approach to pointing to file blocks. Let’s examine an example with twelve direct pointers, as well as both a single and a double indirect block. Assuming a block size of 4 KB, and 4-byte pointers, this structure can accommodate a file of just over 4 GB in size (i.e., \((12 + 1024 + 1024^2) \times 4 \text{ KB}\)). Can you figure out how big of a file can be handled with the addition of a triple-indirect block? (hint: pretty big)

Many file systems use a multi-level index, including commonly-used file systems such as Linux ext2 [P09] and ext3, NetApp’s WAFL, as well as the original Unix file system. Other file systems, including SGI XFS and Linux ext4, use **extents** instead of simple pointers; see the earlier aside for details on how extent-based schemes work (they are akin to segments in the discussion of virtual memory).

You might be wondering: why use an imbalanced tree like this? Why not a different approach? Well, as it turns out, many researchers have studied file systems and how they are used, and virtually every time they find certain “truths” that hold across the decades. One such finding is that **most files are small**. This imbalanced design reflects such a reality; if most files are indeed small, it makes sense to optimize for this case. Thus, with a small number of direct pointers (12 is a typical number), an inode
ASIDE: LINKED-BASED APPROACHES

Another simpler approach in designing inodes is to use a linked list. Thus, inside an inode, instead of having multiple pointers, you just need one, to point to the first block of the file. To handle larger files, add another pointer at the end of that data block, and so on, and thus you can support large files.

As you might have guessed, linked file allocation performs poorly for some workloads; think about reading the last block of a file, for example, or just doing random access. Thus, to make linked allocation work better, some systems will keep an in-memory table of link information, instead of storing the next pointers with the data blocks themselves. The table is indexed by the address of a data block \( D \); the content of an entry is simply \( D \)'s next pointer, i.e., the address of the next block in a file which follows \( D \). A null-value could be there too (indicating an end-of-file), or some other marker to indicate that a particular block is free. Having such a table of next pointers makes it so that a linked allocation scheme can effectively do random file accesses, simply by first scanning through the (in memory) table to find the desired block, and then accessing (on disk) it directly.

Does such a table sound familiar? What we have described is the basic structure of what is known as the file allocation table, or FAT file system. Yes, this classic old Windows file system, before NTFS [C94], is based on a simple linked-based allocation scheme. There are other differences from a standard UNIX file system too; for example, there are no inodes per se, but rather directory entries which store metadata about a file and refer directly to the first block of said file, which makes creating hard links impossible. See Brouwer [B02] for more of the inelegant details.

can directly point to 48 KB of data, needing one (or more) indirect blocks for larger files. See Agrawal et. al [A+07] for a recent study; Table 40.2 summarizes those results.

Of course, in the space of inode design, many other possibilities exist; after all, the inode is just a data structure, and any data structure that stores the relevant information, and can query it effectively, is sufficient. As file system software is readily changed, you should be willing to explore different designs should workloads or technologies change.

<table>
<thead>
<tr>
<th>Most files are small</th>
<th>Roughly 2K is the most common size</th>
</tr>
</thead>
<tbody>
<tr>
<td>Average file size is growing</td>
<td>Almost 200K is the average</td>
</tr>
<tr>
<td>Most bytes are stored in large files</td>
<td>A few big files use most of the space</td>
</tr>
<tr>
<td>File systems contain lots of files</td>
<td>Almost 100K on average</td>
</tr>
<tr>
<td>File systems are roughly half full</td>
<td>Even as disks grow, file systems remain ~50% full</td>
</tr>
<tr>
<td>Directories are typically small</td>
<td>Many have few entries; most have 20 or fewer</td>
</tr>
</tbody>
</table>

Table 40.2: File System Measurement Summary
40.4 Directory Organization

In vsfs (as in many file systems), directories have a simple organization; a directory basically just contains a list of (entry name, inode number) pairs. For each file or directory in a given directory, there is a string and a number in the data block(s) of the directory. For each string, there may also be a length (assuming variable-sized names).

For example, assume a directory dir (inode number 5) has three files in it (foo, bar, and foobar), and their inode numbers are 12, 13, and 24 respectively. The on-disk data for dir might look like this:

<table>
<thead>
<tr>
<th>inum</th>
<th>reclen</th>
<th>strlen</th>
<th>name</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>4</td>
<td>2</td>
<td>..</td>
</tr>
<tr>
<td>2</td>
<td>4</td>
<td>4</td>
<td>..</td>
</tr>
<tr>
<td>12</td>
<td>4</td>
<td>4</td>
<td>foo</td>
</tr>
<tr>
<td>13</td>
<td>4</td>
<td>4</td>
<td>bar</td>
</tr>
<tr>
<td>24</td>
<td>8</td>
<td>7</td>
<td>foobar</td>
</tr>
</tbody>
</table>

In this example, each entry has an inode number, record length (the total bytes for the name plus any leftover space), string length (the actual length of the name), and finally the name of the entry. Note that each directory has two extra entries, . “dot” and .. “dot-dot”; the dot directory is just the current directory (in this example, dir), whereas dot-dot is the parent directory (in this case, the root).

Deleting a file (e.g., calling unlink()) can leave an empty space in the middle of the directory, and hence there should be some way to mark that as well (e.g., with a reserved inode number such as zero). Such a delete is one reason the record length is used: a new entry may reuse an old, bigger entry and thus have extra space within.

You might be wondering where exactly directories are stored. Often, file systems treat directories as a special type of file. Thus, a directory has an inode, somewhere in the inode table (with the type field of the inode marked as “directory” instead of “regular file”). The directory has data blocks pointed to by the inode (and perhaps, indirect blocks); these data blocks live in the data block region of our simple file system. Our on-disk structure thus remains unchanged.

We should also note again that this simple linear list of directory entries is not the only way to store such information. As before, any data structure is possible. For example, XFS [S+96] stores directories in B-tree form, making file create operations (which have to ensure that a file name has not been used before creating it) faster than systems with simple lists that must be scanned in their entirety.

40.5 Free Space Management

A file system must track which inodes and data blocks are free, and which are not, so that when a new file or directory is allocated, it can find space for it. Thus free space management is important for all file systems. In vsfs, we have two simple bitmaps for this task.
There are many ways to manage free space; bitmaps are just one way. Some early file systems used free lists, where a single pointer in the super block was kept to point to the first free block; inside that block the next free pointer was kept, thus forming a list through the free blocks of the system. When a block was needed, the head block was used and the list updated accordingly.

Modern file systems use more sophisticated data structures. For example, SGI’s XFS [S+96] uses some form of a B-tree to compactly represent which chunks of the disk are free. As with any data structure, different time-space trade-offs are possible.

For example, when we create a file, we will have to allocate an inode for that file. The file system will thus search through the bitmap for an inode that is free, and allocate it to the file; the file system will have to mark the inode as used (with a 1) and eventually update the on-disk bitmap with the correct information. A similar set of activities take place when a data block is allocated.

Some other considerations might also come into play when allocating data blocks for a new file. For example, some Linux file systems, such as ext2 and ext3, will look for a sequence of blocks (say 8) that are free when a new file is created and needs data blocks; by finding such a sequence of free blocks, and then allocating them to the newly-created file, the file system guarantees that a portion of the file will be on the disk and contiguous, thus improving performance. Such a pre-allocation policy is thus a commonly-used heuristic when allocating space for data blocks.

40.6 Access Paths: Reading and Writing

Now that we have some idea of how files and directories are stored on disk, we should be able to follow the flow of operation during the activity of reading or writing a file. Understanding what happens on this access path is thus the second key in developing an understanding of how a file system works; pay attention!

For the following examples, let us assume that the file system has been mounted and thus that the superblock is already in memory. Everything else (i.e., inodes, directories) is still on the disk.

Reading A File From Disk

In this simple example, let us first assume that you want to simply open a file (e.g., /foo/bar, read it, and then close it. For this simple example, let’s assume the file is just 4KB in size (i.e., 1 block).

When you issue an open("/foo/bar", O_RDONLY) call, the file system first needs to find the inode for the file bar, to obtain some basic information about the file (permissions information, file size, etc.). To do so,
the file system must be able to find the inode, but all it has right now is the full pathname. The file system must traverse the pathname and thus locate the desired inode.

All traversals begin at the root of the file system, in the root directory which is simply called /. Thus, the first thing the FS will read from disk is the inode of the root directory. But where is this inode? To find an inode, we must know its i-number. Usually, we find the i-number of a file or directory in its parent directory; the root has no parent (by definition). Thus, the root inode number must be “well known”; the FS must know what it is when the file system is mounted. In most UNIX file systems, the root inode number is 2. Thus, to begin the process, the FS reads in the block that contains inode number 2 (the first inode block).

Once the inode is read in, the FS can look inside of it to find pointers to data blocks, which contain the contents of the root directory. The FS will thus use these on-disk pointers to read through the directory, in this case looking for an entry for foo. By reading in one or more directory data blocks, it will find the entry for foo; once found, the FS will also have found the inode number of foo (say it is 44) which it will need next.

The next step is to recursively traverse the pathname until the desired inode is found. In this example, the FS would next read the block containing the inode of foo and then read in its directory data, finally finding the inode number of bar. The final step of open(), then, is to read its inode into memory; the FS can then do a final permissions check, allocate a file descriptor for this process in the per-process open-file table, and return it to the user.

Once open, the program can then issue a read() system call to read from the file. The first read (at offset 0 unless lseek() has been called) will thus read in the first block of the file, consulting the inode to find the location of such a block; it may also update the inode with a new last-accessed time. The read will further update the in-memory open file table for this file descriptor, updating the file offset such that the next read will read the second file block, etc.
Aside: Reads Don’t Access Allocation Structures

We’ve seen many students get confused by allocation structures such as bitmaps. In particular, many often think that when you are simply reading a file, and not allocating any new blocks, that the bitmap will still be consulted. This is not true! Allocation structures, such as bitmaps, are only accessed when allocation is needed. The inodes, directories, and indirect blocks have all the information they need to complete a read request; there is no need to make sure a block is allocated when the inode already points to it.

At some point, the file will be closed. There is much less work to be done here; clearly, the file descriptor should be deallocated, but for now, that is all the FS really needs to do. No disk I/Os take place.

A depiction of this entire process is found in Figure 40.3 (time increases downward). In the figure, the open causes numerous reads to take place in order to finally locate the inode of the file. Afterwards, reading each block requires the file system to first consult the inode, then read the block, and then update the inode’s last-accessed-time field with a write. Spend some time and try to understand what is going on.

Also note that the amount of I/O generated by the open is proportional to the length of the pathname. For each additional directory in the path, we have to read its inode as well as its data. Making this worse would be the presence of large directories; here, we only have to read one block to get the contents of a directory, whereas with a large directory, we might have to read many data blocks to find the desired entry. Yes, life can get pretty bad when reading a file; as you’re about to find out, writing out a file (and especially, creating a new one) is even worse.

Writing to Disk

Writing to a file is a similar process. First, the file must be opened (as above). Then, the application can issue write() calls to update the file with new contents. Finally, the file is closed.

Unlike reading, writing to the file may also allocate a block (unless the block is being overwritten, for example). When writing out a new file, each write not only has to write data to disk but has to first decide which block to allocate to the file and thus update other structures of the disk accordingly (e.g., the data bitmap). Thus, each write to a file logically generates three I/Os: one to read the data bitmap, which is then updated to mark the newly-allocated block as used, one to write the bitmap (to reflect its new state to disk), and one to write the actual block itself.

The amount of write traffic is even worse when one considers a simple and common operation such as file creation. To create a file, the file system must not only allocate an inode, but also allocate space within the directory containing the new file. The total amount of I/O traffic to do so is quite high: one read to the inode bitmap (to find a free inode),
one write to the inode bitmap (to mark it allocated), one write to the new 
inode itself (to initialize it), one to the data of the directory (to link the 
high-level name of the file to its inode number), and one read and write 
to the directory inode to update it. If the directory needs to grow to ac-
commodate the new entry, additional I/Os (i.e., to the data bitmap, and 
the new directory block) will be needed too. All that just to create a file!

Let’s look at a specific example, where the file /foo/bar is created, 
and three blocks are written to it. Figure 40.4 shows what happens during 
the open() (which creates the file) and during each of three 4KB writes.

In the figure, reads and writes to the disk are grouped under which 
system call caused them to occur, and the rough ordering they might take 
place in goes from top to bottom of the figure. You can see how much 
work it is to create the file: 10 I/Os in this case, to walk the pathname 
and then finally create the file. You can also see that each allocating write 
costs 5 I/Os: a pair to read and update the inode, another pair to read 
and update the data bitmap, and then finally the write of the data itself.
How can a file system accomplish any of this with reasonable efficiency?

### Table 40.4: File Creation Timeline (Time Increasing Downward)

<table>
<thead>
<tr>
<th>System Call</th>
<th>File Name</th>
<th>I/O Operations</th>
</tr>
</thead>
<tbody>
<tr>
<td>create (/foo/bar)</td>
<td></td>
<td>read inode, read data, write bitmap, write inode, read inode, read data, write data, read data[0], read data[1], write data[1]</td>
</tr>
<tr>
<td>write()</td>
<td></td>
<td>read inode, write data, read data[1], write data[1]</td>
</tr>
<tr>
<td>write()</td>
<td></td>
<td>read inode, write data, read data[0], write data[0]</td>
</tr>
<tr>
<td>write()</td>
<td></td>
<td>read inode, write data, read data[1], write data[1]</td>
</tr>
</tbody>
</table>

THE CRUX: HOW TO REDUCE FILE SYSTEM I/O COSTS

Even the simplest of operations like opening, reading, or writing a file 
incurs a huge number of I/O operations, scattered over the disk. What 
can a file system do to reduce the high costs of doing so many I/Os?
40.7 Caching and Buffering

As the examples above show, reading and writing files can be expensive, incurring many I/Os to the (slow) disk. To remedy what would clearly be a huge performance problem, most file systems aggressively use system memory (DRAM) to cache important blocks.

Imagine the open example above: without caching, every file open would require at least two reads for every level in the directory hierarchy (one to read the inode of the directory in question, and at least one to read its data). With a long pathname (e.g., /1/2/3/ ... /100/file.txt), the file system would literally perform hundreds of reads just to open the file!

Early file systems thus introduced a fix-sized cache to hold popular blocks. As in our discussion of virtual memory, strategies such as LRU and different variants would decide which blocks to keep in cache. This fix-sized cache would usually be allocated at boot time to be roughly 10% of total memory. Modern systems integrate virtual memory pages and file system pages into a unified page cache [500]. In this way, memory can be allocated more flexibly across virtual memory and file system, depending on which needs more memory at a given time.

Now imagine the file open example with caching. The first open may generate a lot of I/O traffic to read in directory inode and data, but subsequent file opens of that same file (or files in the same directory) will mostly hit in the cache and thus no I/O is needed.

Let us also consider the effect of caching on writes. Whereas read I/O can be avoided altogether with a sufficiently large cache, write traffic has to go to disk in order to become persistent. Thus, a cache does not serve as the same kind of filter on write traffic that it does for reads. That said, write buffering (as it is sometimes called) certainly has a number of performance benefits. First, by delaying writes, the file system can batch some updates into a smaller set of I/Os; for example, if an inode bitmap is updated when one file is created and then updated moments later as another file is created, the file system saves an I/O by delaying the write after the first update. Second, by buffering a number of writes in memory, the system can then schedule the subsequent I/Os and thus increase performance. Finally, some writes are avoided altogether by delaying them; for example, if an application creates a file and then deletes it, delaying the writes to reflect the file creation to disk avoids them entirely. In this case, laziness (in writing blocks to disk) is a virtue.

For the reasons above, most modern file systems buffer writes in memory for anywhere between five and thirty seconds, representing yet another trade-off: if the system crashes before the updates have been propagated to disk, the updates are lost; however, by keeping writes in memory longer, performance can be improved by batching, scheduling, and even avoiding writes.

Some applications (such as databases) don’t enjoy this trade-off. Thus, to avoid unexpected data loss due to write buffering, they simply force writes to disk, by calling \texttt{fsync()}, by using direct I/O interfaces that
work around the cache, or by using the raw disk interface and avoiding the file system altogether\(^1\). While most applications live with the trade-offs made by the file system, there are enough controls in place to get the system to do what you want it to, should the default not be satisfying.

### 40.8 Summary

We have seen the basic machinery required in building a file system. There needs to be some information about each file (metadata), usually stored in a structure called an inode. Directories are just a specific type of file that store name—inode-number mappings. And other structures are needed too; for example, file systems often use a structure such as a bitmap to track which inodes or data blocks are free or allocated.

The terrific aspect of file system design is its freedom; the file systems we explore in the coming chapters each take advantage of this freedom to optimize some aspect of the file system. There are also clearly many policy decisions we have left unexplored. For example, when a new file is created, where should it be placed on disk? This policy and others will also be the subject of future chapters. Or will they?

\(^1\)Take a database class to learn more about old-school databases and their former insistence on avoiding the OS and controlling everything themselves. But watch out! Those database types are always trying to bad mouth the OS. Shame on you, database people. Shame.
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the trail of file-system analysis papers back to the early 1980s.

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February 1988
A classic distributed file system; we’ll be learning more about it later, don’t worry.

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FREENIX, 2000
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cache. Many other systems do the same type of thing.

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Mike Nishimoto, Geoff Peck
USENIX ’96, January 1996, San Diego, CA
The first attempt to make scalability of operations, including things like having millions of files in a
directory, a central focus. A great example of pushing an idea to the extreme. The key idea behind this
file system: everything is a tree. We should have a chapter on this file system too.

Operating Systems
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Homework

Use this tool, vsfs.py, to study how file system state changes as various operations take place. The file system begins in an empty state, with just a root directory. As the simulation takes place, various operations are performed, thus slowly changing the on-disk state of the file system. See the README for details.

Questions

1. Run the simulator with some different random seeds (say 17, 18, 19, 20), and see if you can figure out which operations must have taken place between each state change.
2. Now do the same, using different random seeds (say 21, 22, 23, 24), except run with the \(-r\) flag, thus making you guess the state change while being shown the operation. What can you conclude about the inode and data-block allocation algorithms, in terms of which blocks they prefer to allocate?
3. Now reduce the number of data blocks in the file system, to very low numbers (say two), and run the simulator for a hundred or so requests. What types of files end up in the file system in this highly-constrained layout? What types of operations would fail?
4. Now do the same, but with inodes. With very few inodes, what types of operations can succeed? Which will usually fail? What is the final state of the file system likely to be?

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Locality and The Fast File System

When the UNIX operating system was first introduced, the UNIX wizard himself Ken Thompson wrote the first file system. We will call that the “old UNIX file system”, and it was really simple. Basically, its data structures looked like this on the disk:

<table>
<thead>
<tr>
<th></th>
<th>Inodes</th>
<th>Data</th>
</tr>
</thead>
<tbody>
<tr>
<td>S</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The super block (S) contained information about the entire file system: how big the volume is, how many inodes there are, a pointer to the head of a free list of blocks, and so forth. The inode region of the disk contained all the inodes for the file system. Finally, most of the disk was taken up by data blocks.

The good thing about the old file system was that it was simple, and supported the basic abstractions the file system was trying to deliver: files and the directory hierarchy. This easy-to-use system was a real step forward from the clumsy, record-based storage systems of the past, and the directory hierarchy a true advance over simpler, one-level hierarchies provided by earlier systems.

41.1 The Problem: Poor Performance

The problem: performance was terrible. As measured by Kirk McKusick and his colleagues at Berkeley [MJLF84], performance started off bad and got worse over time, to the point where the file system was delivering only 2% of overall disk bandwidth!

The main issue was that the old UNIX file system treated the disk like it was a random-access memory; data was spread all over the place without regard to the fact that the medium holding the data was a disk, and thus had real and expensive positioning costs. For example, the data blocks of a file were often very far away from its inode, thus inducing an expensive seek whenever one first read the inode and then the data blocks of a file (a pretty common operation).
Worse, the file system would end up getting quite **fragmented**, as the free space was not carefully managed. The free list would end up pointing to a bunch of blocks spread across the disk, and as files got allocated, they would simply take the next free block. The result was that a logically contiguous file would be accessed by going back and forth across the disk, thus reducing performance dramatically.

For example, imagine the following data block region, which contains four files (A, B, C, and D), each of size 2 blocks:

<table>
<thead>
<tr>
<th>A1</th>
<th>A2</th>
<th>B1</th>
<th>B2</th>
<th>C1</th>
<th>C2</th>
<th>D1</th>
<th>D2</th>
</tr>
</thead>
</table>

If B and D are deleted, the resulting layout is:

<table>
<thead>
<tr>
<th>A1</th>
<th>A2</th>
<th></th>
<th></th>
<th>C1</th>
<th>C2</th>
<th></th>
<th></th>
</tr>
</thead>
</table>

As you can see, the free space is fragmented into two chunks of two blocks, instead of one nice contiguous chunk of four. Let's say we now wish to allocate a file E, of size four blocks:

<table>
<thead>
<tr>
<th>A1</th>
<th>A2</th>
<th>E1</th>
<th>E2</th>
<th>C1</th>
<th>C2</th>
<th>E3</th>
<th>E4</th>
</tr>
</thead>
</table>

You can see what happens: E gets spread across the disk, and as a result, when accessing E, you don’t get peak (sequential) performance from the disk. Rather, you first read E1 and E2, then seek, then read E3 and E4. This fragmentation problem happened all the time in the old UNIX file system, and it hurt performance. (A side note: this problem is exactly what disk defragmentation tools help with; they will reorganize on-disk data to place files contiguously and make free space one or a few contiguous regions, moving data around and then rewriting inodes and such to reflect the changes)

One other problem: the original block size was too small (512 bytes). Thus, transferring data from the disk was inherently inefficient. Smaller blocks were good because they minimized internal fragmentation (waste within the block), but bad for transfer as each block might require a positioning overhead to reach it. We can summarize the problem as follows:

**The Crux:**

**How To Organize On-disk Data To Improve Performance**

How can we organize file system data structures so as to improve performance? What types of allocation policies do we need on top of those data structures? How do we make the file system “disk aware”?

---

**Operating Systems**

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41.2 FFS: Disk Awareness Is The Solution

A group at Berkeley decided to build a better, faster file system, which they cleverly called the Fast File System (FFS). The idea was to design the file system structures and allocation policies to be “disk aware” and thus improve performance, which is exactly what they did. FFS thus ushered in a new era of file system research; by keeping the same interface to the file system (the same APIs, including open(), read(), write(), close(), and other file system calls) but changing the internal implementation, the authors paved the path for new file system construction, work that continues today. Virtually all modern file systems adhere to the existing interface (and thus preserve compatibility with applications) while changing their internals for performance, reliability, or other reasons.

41.3 Organizing Structure: The Cylinder Group

The first step was to change the on-disk structures. FFS divides the disk into a bunch of groups known as cylinder groups (some modern file systems like Linux ext2 and ext3 just call them block groups). We can thus imagine a disk with ten cylinder groups:

```
G0  G1  G2  G3  G4  G5  G6  G7  G8  G9
```

These groups are the central mechanism that FFS uses to improve performance; by placing two files within the same group, FFS can ensure that accessing one after the other will not result in long seeks across the disk.

Thus, FFS needs to have the ability to allocate files and directories within each of these groups. Each group looks like this:

```
S  ib  db  Inodes  Data
```

We now describe the components of a cylinder group. A copy of the super block (S) is found in each group for reliability reasons (e.g., if one gets corrupted or scratched, you can still mount and access the file system by using one of the others).

The inode bitmap (ib) and data bitmap (db) track whether each inode or data block is free, respectively. Bitmaps are an excellent way to manage free space in a file system because it is easy to find a large chunk of free space and allocate it to a file, perhaps avoiding some of the fragmentation problems of the free list in the old file system.

Finally, the inode and data block regions are just like in the previous very simple file system. Most of each cylinder group, as usual, is comprised of data blocks.
A side: FFS File Creation

As an example, think about what data structures must be updated when a file is created; assume, for this example, that the user creates a new file /foo/bar.txt and that the file is one block long (4KB). The file is new, and thus needs a new inode; thus, both the inode bitmap and the newly-allocated inode will be written to disk. The file also has data in it and thus it too must be allocated; the data bitmap and a data block will thus (eventually) be written to disk. Hence, at least four writes to the current cylinder group will take place (recall that these writes may be buffered in memory for a while before the write takes place). But this is not all! In particular, when creating a new file, we must also place the file in the file-system hierarchy; thus, the directory must be updated. Specifically, the parent directory foo must be updated to add the entry for bar.txt; this update may fit in an existing data block of foo or require a new block to be allocated (with associated data bitmap). The inode of foo must also be updated, both to reflect the new length of the directory as well as to update time fields (such as last-modified-time). Overall, it is a lot of work just to create a new file! Perhaps next time you do so, you should be more thankful, or at least surprised that it all works so well.

41.4 Policies: How To Allocate Files and Directories

With this group structure in place, FFS now has to decide how to place files and directories and associated metadata on disk to improve performance. The basic mantra is simple: keep related stuff together (and its corollary, keep unrelated stuff far apart).

Thus, to obey the mantra, FFS has to decide what is “related” and place it within the same block group; conversely, unrelated items should be placed into different block groups. To achieve this end, FFS makes use of a few simple placement heuristics.

The first is the placement of directories. FFS employs a simple approach: find the cylinder group with a low number of allocated directories (because we want to balance directories across groups) and a high number of free inodes (because we want to subsequently be able to allocate a bunch of files), and put the directory data and inode in that group. Of course, other heuristics could be used here (e.g., taking into account the number of free data blocks).

For files, FFS does two things. First, it makes sure (in the general case) to allocate the data blocks of a file in the same group as its inode, thus preventing long seeks between inode and data (as in the old file system). Second, it places all files that are in the same directory in the cylinder group of the directory they are in. Thus, if a user creates four files, /dir1/1.txt, /dir1/2.txt, /dir1/3.txt, and /dir99/4.txt, FFS would try to place the first three near one another (same group) and the fourth far away (in some other group).
It should be noted that these heuristics are not based on extensive studies of file-system traffic or anything particularly nuanced; rather, they are based on good old-fashioned common sense (isn’t that what CS stands for after all?). Files in a directory are often accessed together (imagine compiling a bunch of files and then linking them into a single executable). Because they are, FFS will often improve performance, making sure that seeks between related files are short.

41.5 Measuring File Locality

To understand better whether these heuristics make sense, we decided to analyze some traces of file system access and see if indeed there is namespace locality; for some reason, there doesn’t seem to be a good study of this topic in the literature.

Specifically, we took the SEER traces [K94] and analyzed how “far away” file accesses were from one another in the directory tree. For example, if file $f$ is opened, and then re-opened next in the trace (before any other files are opened), the distance between these two opens in the directory tree is zero (as they are the same file). If a file $f$ in directory dir (i.e., dir/$f$) is opened, and followed by an open of file $g$ in the same directory (i.e., dir/$g$), the distance between the two file accesses is one, as they share the same directory but are not the same file. Our distance metric, in other words, measures how far up the directory tree you have to travel to find the common ancestor of two files; the closer they are in the tree, the lower the metric.
Figure 41.1 shows the locality observed in the SEER traces over all workstations in the SEER cluster over the entirety of all traces. The graph plots the difference metric along the x-axis, and shows the cumulative percentage of file opens that were of that difference along the y-axis. Specifically, for the SEER traces (marked “Trace” in the graph), you can see that about 7% of file accesses were to the file that was opened previously, and that nearly 40% of file accesses were to either the same file or to one in the same directory (i.e., a difference of zero or one). Thus, the FFS locality assumption seems to make sense (at least for these traces).

Interestingly, another 25% or so of file accesses were to files that had a distance of two. This type of locality occurs when the user has structured a set of related directories in a multi-level fashion and consistently jumps between them. For example, if a user has a src directory and builds object files (.o files) into a obj directory, and both of these directories are sub-directories of a main proj directory, a common access pattern will be proj/src/foo.c followed by proj/obj/foo.o. The distance between these two accesses is two, as proj is the common ancestor. FFS does not capture this type of locality in its policies, and thus more seeking will occur between such accesses.

We also show what locality would be for a “Random” trace for the sake of comparison. We generated the random trace by selecting files from within an existing SEER trace in random order, and calculating the distance metric between these randomly-ordered accesses. As you can see, there is less namespace locality in the random traces, as expected. However, because eventually every file shares a common ancestor (e.g., the root), there is some locality eventually, and thus random trace is useful as a comparison point.

41.6 The Large-File Exception

In FFS, there is one important exception to the general policy of file placement, and it arises for large files. Without a different rule, a large file would entirely fill the block group it is first placed within (and maybe others). Filling a block group in this manner is undesirable, as it prevents subsequent “related” files from being placed within this block group, and thus may hurt file-access locality.

Thus, for large files, FFS does the following. After some number of blocks are allocated into the first block group (e.g., 12 blocks, or the number of direct pointers available within an inode), FFS places the next “large” chunk of the file (e.g., those pointed to by the first indirect block) in another block group (perhaps chosen for its low utilization). Then, the next chunk of the file is placed in yet another different block group, and so on.

Let’s look at some pictures to understand this policy better. Without the large-file exception, a single large file would place all of its blocks into one part of the disk. We use a small example of a file with 10 blocks to illustrate the behavior visually.
Here is the depiction of FFS without the large-file exception:

\[
\begin{array}{cccccccccc}
G0 & G1 & G2 & G3 & G4 & G5 & G6 & G7 & G8 & G9 \\
0 & 1 & 2 & 3 & 4 & 5 & 6 & 7 & 8 & 9
\end{array}
\]

With the large-file exception, we might see something more like this, with the file spread across the disk in chunks:

\[
\begin{array}{cccccccccc}
G0 & G1 & G2 & G3 & G4 & G5 & G6 & G7 & G8 & G9 \\
8 & 9 & 0 & 1 & 2 & 3 & 4 & 5 & 6 & 7
\end{array}
\]

The astute reader will note that spreading blocks of a file across the disk will hurt performance, particularly in the relatively common case of sequential file access (e.g., when a user or application reads chunks 0 through 9 in order). And you are right! It will. We can help this a little, by choosing our chunk size carefully.

Specifically, if the chunk size is large enough, we will still spend most of our time transferring data from disk and just a relatively little time seeking between chunks of the block. This process of reducing an overhead by doing more work per overhead paid is called \textit{amortization} and is a common technique in computer systems.

Let’s do an example: assume that the average positioning time (i.e., seek and rotation) for a disk is 10 ms. Assume further that the disk transfers data at 40 MB/s. If our goal was to spend half our time seeking between chunks and half our time transferring data (and thus achieve 50% of peak disk performance), we would thus need to spend 10 ms transferring data for every 10 ms positioning. So the question becomes: how big does a chunk have to be in order to spend 10 ms in transfer? Easy, just use our old friend, math, in particular the dimensional analysis we spoke of in the chapter on disks:

\[
\frac{40 \text{ MB}}{\text{sec}} \cdot \frac{1024 \text{ KB}}{\text{MB}} \cdot \frac{1 \text{ sec}}{1000 \text{ ms}} \cdot 10 \text{ ms} = 409.6 \text{ KB}
\]  

(41.1)

Basically, what this equation says is this: if you transfer data at 40 MB/s, you need to transfer only 409.6 KB every time you seek in order to spend half your time seeking and half your time transferring. Similarly, you can compute the size of the chunk you would need to achieve 90% of peak bandwidth (turns out it is about 3.69 MB), or even 99% of peak bandwidth (40.6 MB!). As you can see, the closer you want to get to peak, the bigger these chunks get (see Figure 41.2 for a plot of these values).

FFS did not use this type of calculation in order to spread large files across groups, however. Instead, it took a simple approach, based on the structure of the inode itself. The first twelve direct blocks were placed in the same group as the inode; each subsequent indirect block, and all the blocks it pointed to, was placed in a different group. With a block size of 4-KB, and 32-bit disk addresses, this strategy implies that every 1024 blocks of the file (4 MB) were placed in separate groups, the lone exception being the first 48-KB of the file as pointed to by direct pointers.
The Challenges of Amortization

We should note that the trend in disk drives is that transfer rate improves fairly rapidly, as disk manufacturers are good at cramming more bits into the same surface, but the mechanical aspects of drives related to seeks (disk arm speed and the rate of rotation) improve rather slowly [P98]. The implication is that over time, mechanical costs become relatively more expensive, and thus, to amortize said costs, you have to transfer more data between seeks.

41.7 A Few Other Things About FFS

FFS introduced a few other innovations too. In particular, the designers were extremely worried about accommodating small files; as it turned out, many files were 2 KB or so in size back then, and using 4-KB blocks, while good for transferring data, was not so good for space efficiency. This internal fragmentation could thus lead to roughly half the disk being wasted for a typical file system.

The solution the FFS designers hit upon was simple and solved the problem. They decided to introduce sub-blocks, which were 512-byte little blocks that the file system could allocate to files. Thus, if you created a small file (say 1 KB in size), it would occupy two sub-blocks and thus not waste an entire 4-KB block. As the file grew, the file system will continue allocating 512-byte blocks to it until it acquires a full 4-KB of data. At that point, FFS will find a 4-KB block, copy the sub-blocks into it, and free the sub-blocks for future use.
You might observe that this process is inefficient, requiring a lot of extra work for the file system (in particular, a lot of extra I/O to perform the copy). And you’d be right again! Thus, FFS generally avoided this pes-simal behavior by modifying the libc library; the library would buffer writes and then issue them in 4-KB chunks to the file system, thus avoiding the sub-block specialization entirely in most cases.

A second neat thing that FFS introduced was a disk layout that was optimized for performance. In those times (before SCSI and other more modern device interfaces), disks were much less sophisticated and required the host CPU to control their operation in a more hands-on way. A problem arose in FFS when a file was placed on consecutive sectors of the disk, as on the left in Figure 41.3.

In particular, the problem arose during sequential reads. FFS would first issue a read to block 0; by the time the read was complete, and FFS issued a read to block 1, it was too late: block 1 had rotated under the head and now the read to block 1 would incur a full rotation.

FFS solved this problem with a different layout, as you can see on the right in Figure 41.3. By skipping over every other block (in the example), FFS has enough time to request the next block before it went past the disk head. In fact, FFS was smart enough to figure out for a particular disk how many blocks it should skip in doing layout in order to avoid the extra rotations; this technique was called parameterization, as FFS would figure out the specific performance parameters of the disk and use those to decide on the exact staggered layout scheme.

You might be thinking: this scheme isn’t so great after all. In fact, you will only get 50% of peak bandwidth with this type of layout, because you have to go around each track twice just to read each block once. Fortunately, modern disks are much smarter: they internally read the entire track in and buffer it in an internal disk cache (often called a track buffer for this very reason). Then, on subsequent reads to the track, the disk will just return the desired data from its cache. File systems thus no longer have to worry about these incredibly low-level details. Abstraction and higher-level interfaces can be a good thing, when designed properly.

Some other usability improvements were added as well. FFS was one of the first file systems to allow for long file names, thus enabling more expressive names in the file system instead of a the traditional fixed-size approach (e.g., 8 characters). Further, a new concept was introduced...
Called a **symbolic link**. As discussed in a previous chapter, hard links are limited in that they both could not point to directories (for fear of introducing loops in the file system hierarchy) and that they can only point to files within the same volume (i.e., the inode number must still be meaningful). Symbolic links allow the user to create an “alias” to any other file or directory on a system and thus are much more flexible. FFS also introduced an atomic `rename()` operation for renaming files. Usability improvements, beyond the basic technology, also likely gained FFS a stronger user base.

### 41.8 Summary

The introduction of FFS was a watershed moment in file system history, as it made clear that the problem of file management was one of the most interesting issues within an operating system, and showed how one might begin to deal with that most important of devices, the hard disk. Since that time, hundreds of new file systems have developed, but still today many file systems take cues from FFS (e.g., Linux ext2 and ext3 are obvious intellectual descendants). Certainly all modern systems account for the main lesson of FFS: treat the disk like it’s a disk.
References

[MJLF84] “A Fast File System for UNIX”
Marshall K. McKusick, William N. Joy, Sam J. Leffler, Robert S. Fabry
ACM Transactions on Computing Systems.
pages 181-197.
McKusick was recently honored with the IEEE Reynold B. Johnson award for his contributions to file systems, much of which was based on his work building FFS. In his acceptance speech, he discussed the original FFS software: only 1200 lines of code! Modern versions are a little more complex, e.g., the BSD FFS descendant now is in the 50-thousand lines-of-code range.

[P98] “Hardware Technology Trends and Database Opportunities”
David A. Patterson
Keynote Lecture at the ACM SIGMOD Conference (SIGMOD ’98)
June, 1998
A great and simple overview of disk technology trends and how they change over time.

[K94] “The Design of the SEER Predictive Caching System”
G. H. Kuenning
MOBICOMM ’94, Santa Cruz, California, December 1994
According to Kuenning, this is the best overview of the SEER project, which led to (among other things) the collection of these traces.
Crash Consistency: FSCK and Journaling

As we’ve seen thus far, the file system manages a set of data structures to implement the expected abstractions: files, directories, and all of the other metadata needed to support the basic abstraction that we expect from a file system. Unlike most data structures (for example, those found in memory of a running program), file system data structures must persist, i.e., they must survive over the long haul, stored on devices that retain data despite power loss (such as hard disks or flash-based SSDs).

One major challenge faced by a file system is how to update persistent data structures despite the presence of a power loss or system crash. Specifically, what happens if, right in the middle of updating on-disk structures, someone trips over the power cord and the machine loses power? Or the operating system encounters a bug and crashes? Because of power losses and crashes, updating a persistent data structure can be quite tricky, and leads to a new and interesting problem in file system implementation, known as the crash-consistency problem.

This problem is quite simple to understand. Imagine you have to update two on-disk structures, \( A \) and \( B \), in order to complete a particular operation. Because the disk only services a single request at a time, one of these requests will reach the disk first (either \( A \) or \( B \)). If the system crashes or loses power after one write completes, the on-disk structure will be left in an inconsistent state. And thus, we have a problem that all file systems need to solve:

**The Crux: How To Update The Disk Despite Crashes**

The system may crash or lose power between any two writes, and thus the on-disk state may only partially get updated. After the crash, the system boots and wishes to mount the file system again (in order to access files and such). Given that crashes can occur at arbitrary points in time, how do we ensure the file system keeps the on-disk image in a reasonable state?
In this chapter, we’ll describe this problem in more detail, and look at some methods file systems have used to overcome it. We’ll begin by examining the approach taken by older file systems, known as fsck or the file system checker. We’ll then turn our attention to another approach, known as journaling (also known as write-ahead logging), a technique which adds a little bit of overhead to each write but recovers more quickly from crashes or power losses. We will discuss the basic machinery of journaling, including a few different flavors of journaling that Linux ext3 [T98,PAA05] (a relatively modern journaling file system) implements.

42.1 A Detailed Example

To kick off our investigation of journaling, let’s look at an example. We’ll need to use a workload that updates on-disk structures in some way. Assume here that the workload is simple: the append of a single data block to an existing file. The append is accomplished by opening the file, calling lseek() to move the file offset to the end of the file, and then issuing a single 4KB write to the file before closing it.

Let’s also assume we are using standard simple file system structures on the disk, similar to file systems we have seen before. This tiny example includes an inode bitmap (with just 8 bits, one per inode), a data bitmap (also 8 bits, one per data block), inodes (8 total, numbered 0 to 7, and spread across four blocks), and data blocks (8 total, numbered 0 to 7). Here is a diagram of this file system:

```
Inode Bmap | Data Bmap | Inodes | Data Blocks
---|---|---|---
I[v1] | Da |
```

If you look at the structures in the picture, you can see that a single inode is allocated (inode number 2), which is marked in the inode bitmap, and a single allocated data block (data block 4), also marked in the data bitmap. The inode is denoted I[v1], as it is the first version of this inode; it will soon be updated (due to the workload described above).

Let’s peek inside this simplified inode too. Inside of I[v1], we see:

```
owner : remzi
permissions : read-only
size : 1
pointer : 4
pointer : null
pointer : null
pointer : null
```

In this simplified inode, the size of the file is 1 (it has one block allocated), the first direct pointer points to block 4 (the first data block of the file, Da), and all three other direct pointers are set to null (indicating
that they are not used). Of course, real inodes have many more fields; see previous chapters for more information.

When we append to the file, we are adding a new data block to it, and thus must update three on-disk structures: the inode (which must point to the new block as well as have a bigger size due to the append), the new data block Db, and a new version of the data bitmap (call it B[v2]) to indicate that the new data block has been allocated.

Thus, in the memory of the system, we have three blocks which we must write to disk. The updated inode (inode version 2, or I[v2] for short) now looks like this:

<table>
<thead>
<tr>
<th>owner</th>
<th>remzi</th>
</tr>
</thead>
<tbody>
<tr>
<td>permissions</td>
<td>read-only</td>
</tr>
<tr>
<td>size</td>
<td>2</td>
</tr>
<tr>
<td>pointer</td>
<td>4</td>
</tr>
<tr>
<td>pointer</td>
<td>5</td>
</tr>
<tr>
<td>pointer</td>
<td>null</td>
</tr>
<tr>
<td>pointer</td>
<td>null</td>
</tr>
</tbody>
</table>

The updated data bitmap (B[v2]) now looks like this: 00001100. Finally, there is the data block (Db), which is just filled with whatever it is users put into files. Stolen music perhaps?

What we would like is for the final on-disk image of the file system to look like this:

<table>
<thead>
<tr>
<th>Inode Bmap</th>
<th>Data Bmap</th>
<th>Inodes</th>
<th>Data Blocks</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>I[v2]</td>
<td>Da</td>
</tr>
</tbody>
</table>

To achieve this transition, the file system must perform three separate writes to the disk, one each for the inode (I[v2]), bitmap (B[v2]), and data block (Db). Note that these writes usually don’t happen immediately when the user issues a write() system call; rather, the dirty inode, bitmap, and new data will sit in main memory (in the page cache or buffer cache) for some time first; then, when the file system finally decides to write them to disk (after say 5 seconds or 30 seconds), the file system will issue the requisite write requests to the disk. Unfortunately, a crash may occur and thus interfere with these updates to the disk. In particular, if a crash happens after one or two of these writes have taken place, but not all three, the file system could be left in a funny state.

**Crash Scenarios**

To understand the problem better, let’s look at some example crash scenarios. Imagine only a single write succeeds; there are thus three possible outcomes, which we list here:
Just the data block (Db) is written to disk. In this case, the data is on disk, but there is no inode that points to it and no bitmap that even says the block is allocated. Thus, it is as if the write never occurred. This case is not a problem at all, from the perspective of file-system crash consistency.

Just the updated inode (I[v2]) is written to disk. In this case, the inode points to the disk address (5) where Db was about to be written, but Db has not yet been written there. Thus, if we trust that pointer, we will read garbage data from the disk (the old contents of disk address 5).

Further, we have a new problem, which we call a file-system inconsistency. The on-disk bitmap is telling us that data block 5 has not been allocated, but the inode is saying that it has. This disagreement in the file system data structures is an inconsistency in the data structures of the file system; to use the file system, we must somehow resolve this problem (more on that below).

Just the updated bitmap (B[v2]) is written to disk. In this case, the bitmap indicates that block 5 is allocated, but there is no inode that points to it. Thus the file system is inconsistent again; if left unresolved, this write would result in a space leak, as block 5 would never be used by the file system.

There are also three more crash scenarios in this attempt to write three blocks to disk. In these cases, two writes succeed and the last one fails:

The inode (I[v2]) and bitmap (B[v2]) are written to disk, but not data (Db). In this case, the file system metadata is completely consistent: the inode has a pointer to block 5, the bitmap indicates that 5 is in use, and thus everything looks OK from the perspective of the file system’s metadata. But there is one problem: 5 has garbage in it again.

The inode (I[v2]) and the data block (Db) are written, but not the bitmap (B[v2]). In this case, we have the inode pointing to the correct data on disk, but again have an inconsistency between the inode and the old version of the bitmap (B1). Thus, we once again need to resolve the problem before using the file system.

The bitmap (B[v2]) and data block (Db) are written, but not the inode (I[v2]). In this case, we again have an inconsistency between the inode and the data bitmap. However, even though the block was written and the bitmap indicates its usage, we have no idea which file it belongs to, as no inode points to the file.

1However, it might be a problem for the user, who just lost some data!
The Crash Consistency Problem

Hopefully, from these crash scenarios, you can see the many problems that can occur to our on-disk file system image because of crashes: we can have inconsistency in file system data structures; we can have space leaks; we can return garbage data to a user; and so forth. What we’d like to do ideally is move the file system from one consistent state (e.g., before the file got appended to) to another atomically (e.g., after the inode, bitmap, and new data block have been written to disk). Unfortunately, we can’t do this easily because the disk only commits one write at a time, and crashes or power loss may occur between these updates. We call this general problem the crash-consistency problem (we could also call it the consistent-update problem).

42.2 Solution #1: The File System Checker

Early file systems took a simple approach to crash consistency. Basically, they decided to let inconsistencies happen and then fix them later (when rebooting). A classic example of this lazy approach is found in a tool that does this: fsck. fsck is a UNIX tool for finding such inconsistencies and repairing them [M86]; similar tools to check and repair a disk partition exist on different systems. Note that such an approach can’t fix all problems; consider, for example, the case above where the file system looks consistent but the inode points to garbage data. The only real goal is to make sure the file system metadata is internally consistent.

The tool fsck operates in a number of phases, as summarized in McKusick and Kowalski’s paper [MK96]. It is run before the file system is mounted and made available (fsck assumes that no other file-system activity is on-going while it runs); once finished, the on-disk file system should be consistent and thus can be made accessible to users.

Here is a basic summary of what fsck does:

- **Superblock:** fsck first checks if the superblock looks reasonable, mostly doing sanity checks such as making sure the file system size is greater than the number of blocks allocated. Usually the goal of these sanity checks is to find a suspect (corrupt) superblock; in this case, the system (or administrator) may decide to use an alternate copy of the superblock.

- **Free blocks:** Next, fsck scans the inodes, indirect blocks, double indirect blocks, etc., to build an understanding of which blocks are currently allocated within the file system. It uses this knowledge to produce a correct version of the allocation bitmaps; thus, if there is any inconsistency between bitmaps and inodes, it is resolved by trusting the information within the inodes. The same type of check is performed for all the inodes, making sure that all inodes that look like they are in use are marked as such in the inode bitmaps.

---

2Pronounced either “eff-ess-see-kay”, “eff-ess-check”, or, if you don’t like the tool, “eff-suck”. Yes, serious professional people use this term.
• **Inode state:** Each inode is checked for corruption or other problems. For example, `fsck` makes sure that each allocated inode has a valid type field (e.g., regular file, directory, symbolic link, etc.). If there are problems with the inode fields that are not easily fixed, the inode is considered suspect and cleared by `fsck`; the inode bitmap is correspondingly updated.

• **Inode links:** `fsck` also verifies the link count of each allocated inode. As you may recall, the link count indicates the number of different directories that contain a reference (i.e., a link) to this particular file. To verify the link count, `fsck` scans through the entire directory tree, starting at the root directory, and builds its own link counts for every file and directory in the file system. If there is a mismatch between the newly-calculated count and that found within an inode, corrective action must be taken, usually by fixing the count within the inode. If an allocated inode is discovered but no directory refers to it, it is moved to the lost+found directory.

• **Duplicates:** `fsck` also checks for duplicate pointers, i.e., cases where two different inodes refer to the same block. If one inode is obviously bad, it may be cleared. Alternately, the pointed-to block could be copied, thus giving each inode its own copy as desired.

• **Bad blocks:** A check for bad block pointers is also performed while scanning through the list of all pointers. A pointer is considered “bad” if it obviously points to something outside its valid range, e.g., it has an address that refers to a block greater than the partition size. In this case, `fsck` can’t do anything too intelligent; it just removes (clears) the pointer from the inode or indirect block.

• **Directory checks:** `fsck` does not understand the contents of user files; however, directories hold specifically formatted information created by the file system itself. Thus, `fsck` performs additional integrity checks on the contents of each directory, making sure that “.” and “..” are the first entries, that each inode referred to in a directory entry is allocated, and ensuring that no directory is linked to more than once in the entire hierarchy.

As you can see, building a working `fsck` requires intricate knowledge of the file system; making sure such a piece of code works correctly in all cases can be challenging [G+08]. However, `fsck` (and similar approaches) have a bigger and perhaps more fundamental problem: they are too slow. With a very large disk volume, scanning the entire disk to find all the allocated blocks and read the entire directory tree may take many minutes or hours. Performance of `fsck`, as disks grew in capacity and RAIDs grew in popularity, became prohibitive (despite recent advances [M+13]).

At a higher level, the basic premise of `fsck` seems just a tad irrational. Consider our example above, where just three blocks are written to the disk; it is incredibly expensive to scan the entire disk to fix problems that occurred during an update of just three blocks. This situation is akin to dropping your keys on the floor in your bedroom, and then com-
mencing a search-the-entire-house-for-keys recovery algorithm, starting in the basement and working your way through every room. It works but is wasteful. Thus, as disks (and RAIDs) grew, researchers and practitioners started to look for other solutions.

42.3 Solution #2: Journaling (or Write-Ahead Logging)

Probably the most popular solution to the consistent update problem is to steal an idea from the world of database management systems. That idea, known as write-ahead logging, was invented to address exactly this type of problem. In file systems, we usually call write-ahead logging journaling for historical reasons. The first file system to do this was Cedar [H87], though many modern file systems use the idea, including Linux ext3 and ext4, reiserfs, IBM’s JFS, SGI’s XFS, and Windows NTFS.

The basic idea is as follows. When updating the disk, before overwriting the structures in place, first write down a little note (somewhere else on the disk, in a well-known location) describing what you are about to do. Writing this note is the “write ahead” part, and we write it to a structure that we organize as a “log”; hence, write-ahead logging.

By writing the note to disk, you are guaranteeing that if a crash takes place during the update (overwrite) of the structures you are updating, you can go back and look at the note you made and try again; thus, you will know exactly what to fix (and how to fix it) after a crash, instead of having to scan the entire disk. By design, journaling thus adds a bit of work during updates to greatly reduce the amount of work required during recovery.

We’ll now describe how Linux ext3, a popular journaling file system, incorporates journaling into the file system. Most of the on-disk structures are identical to Linux ext2, e.g., the disk is divided into block groups, and each block group has an inode and data bitmap as well as inodes and data blocks. The new key structure is the journal itself, which occupies some small amount of space within the partition or on another device. Thus, an ext2 file system (without journaling) looks like this:

```
Super  Group 0  Group 1  ...  Group N
```

Assuming the journal is placed within the same file system image (though sometimes it is placed on a separate device, or as a file within the file system), an ext3 file system with a journal looks like this:

```
Super  Journal  Group 0  Group 1  ...  Group N
```

The real difference is just the presence of the journal, and of course, how it is used.
Data Journaling

Let’s look at a simple example to understand how data journaling works. Data journaling is available as a mode with the Linux ext3 file system, from which much of this discussion is based.

Say we have our canonical update again, where we wish to write the ‘inode (I[v2]), bitmap (B[v2]), and data block (Db) to disk again. Before writing them to their final disk locations, we are now first going to write them to the log (a.k.a. journal). This is what this will look like in the log:

| Journal | TxB | I[v2] | B[v2] | Db | TxE |

You can see we have written five blocks here. The transaction begin (TxB) tells us about this update, including information about the pending update to the file system (e.g., the final addresses of the blocks I[v2], B[v2], and Db), as well as some kind of transaction identifier (TID). The middle three blocks just contain the exact contents of the blocks themselves; this is known as physical logging as we are putting the exact physical contents of the update in the journal (an alternate idea, logical logging, puts a more compact logical representation of the update in the journal, e.g., “this update wishes to append data block Db to file X”, which is a little more complex but can save space in the log and perhaps improve performance). The final block (TxE) is a marker of the end of this transaction, and will also contain the TID.

Once this transaction is safely on disk, we are ready to overwrite the old structures in the file system; this process is called checkpointing. Thus, to checkpoint the file system (i.e., bring it up to date with the pending update in the journal), we issue the writes I[v2], B[v2], and Db to their disk locations as seen above; if these writes complete successfully, we have successfully checkpointed the file system and are basically done. Thus, our initial sequence of operations:

1. **Journal write**: Write the transaction, including a transaction-begin block, all pending data and metadata updates, and a transaction-end block, to the log; wait for these writes to complete.
2. **Checkpoint**: Write the pending metadata and data updates to their final locations in the file system.

In our example, we would write TxB, I[v2], B[v2], Db, and TxE to the journal first. When these writes complete, we would complete the update by checkpointing I[v2], B[v2], and Db, to their final locations on disk.

Things get a little trickier when a crash occurs during the writes to the journal. Here, we are trying to write the set of blocks in the transaction (e.g., TxB, I[v2], B[v2], Db, TxE) to disk. One simple way to do this would be to issue each one at a time, waiting for each to complete, and then issuing the next. However, this is slow. Ideally, we’d like to issue
To enforce ordering between two disk writes, modern file systems have to take a few extra precautions. In olden times, forcing ordering between two writes, A and B, was easy: just issue the write of A to the disk, wait for the disk to interrupt the OS when the write is complete, and then issue the write of B.

Things got slightly more complex due to the increased use of write caches within disks. With write buffering enabled (sometimes called immediate reporting), a disk will inform the OS the write is complete when it simply has been placed in the disk’s memory cache, and has not yet reached disk. If the OS then issues a subsequent write, it is not guaranteed to reach the disk after previous writes; thus ordering between writes is not preserved. One solution is to disable write buffering. However, more modern systems take extra precautions and issue explicit write barriers; such a barrier, when it completes, guarantees that all writes issued before the barrier will reach disk before any writes issued after the barrier.

All of this machinery requires a great deal of trust in the correct operation of the disk. Unfortunately, recent research shows that some disk manufacturers, in an effort to deliver “higher performing” disks, explicitly ignore write-barrier requests, thus making the disks seemingly run faster but at the risk of incorrect operation [C+13, R+11]. As Kahan said, the fast almost always beats out the slow, even if the fast is wrong.

All five block writes at once, as this would turn five writes into a single sequential write and thus be faster. However, this is unsafe, for the following reason: given such a big write, the disk internally may perform scheduling and complete small pieces of the big write in any order. Thus, the disk internally may (1) write TxB, I[v2], B[v2], and TxE and only later (2) write Db. Unfortunately, if the disk loses power between (1) and (2), this is what ends up on disk:

<table>
<thead>
<tr>
<th>Journal</th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>TxB</td>
<td>I[v2]</td>
<td>B[v2]</td>
<td>??</td>
<td></td>
</tr>
<tr>
<td>id=1</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Why is this a problem? Well, the transaction looks like a valid transaction (it has a begin and an end with matching sequence numbers). Further, the file system can’t look at that fourth block and know it is wrong; after all, it is arbitrary user data. Thus, if the system now reboots and runs recovery, it will replay this transaction, and ignorantly copy the contents of the garbage block ‘??’ to the location where Db is supposed to live. This is bad for arbitrary user data in a file; it is much worse if it happens to a critical piece of file system, such as the superblock, which could render the file system unmountable.
You may have noticed a particular inefficiency of writing to the log. Namely, the file system first has to write out the transaction-begin block and contents of the transaction; only after these writes complete can the file system send the transaction-end block to disk. The performance impact is clear, if you think about how a disk works: usually an extra rotation is incurred (think about why).

One of our former graduate students, Vijayan Prabhakaran, had a simple idea to fix this problem [P+05]. When writing a transaction to the journal, include a checksum of the contents of the journal in the begin and end blocks. Doing so enables the file system to write the entire transaction at once, without incurring a wait; if, during recovery, the file system sees a mismatch in the computed checksum versus the stored checksum in the transaction, it can conclude that a crash occurred during the write of the transaction and thus discard the file-system update. Thus, with a small tweak in the write protocol and recovery system, a file system can achieve faster common-case performance; on top of that, the system is slightly more reliable, as any reads from the journal are now protected by a checksum.

This simple fix was attractive enough to gain the notice of Linux file system developers, who then incorporated it into the next generation Linux file system, called (you guessed it!) Linux ext4. It now ships on millions of machines worldwide, including the Android handheld platform. Thus, every time you write to disk on many Linux-based systems, a little code developed at Wisconsin makes your system a little faster and more reliable.

To avoid this problem, the file system issues the transactional write in two steps. First, it writes all blocks except the TxE block to the journal, issuing these writes all at once. When these writes complete, the journal will look something like this (assuming our append workload again):

<table>
<thead>
<tr>
<th>Journal</th>
<th>TxB</th>
<th>B[v2]</th>
<th>Db</th>
</tr>
</thead>
<tbody>
<tr>
<td>id=1</td>
<td>I[v2]</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

When those writes complete, the file system issues the write of the TxE block, thus leaving the journal in this final, safe state:

<table>
<thead>
<tr>
<th>Journal</th>
<th>TxB</th>
<th>B[v2]</th>
<th>Db</th>
<th>TxE</th>
</tr>
</thead>
<tbody>
<tr>
<td>id=1</td>
<td>I[v2]</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>id=1</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

An important aspect of this process is the atomicity guarantee provided by the disk. It turns out that the disk guarantees that any 512-byte
write will either happen or not (and never be half-written); thus, to make sure the write of TxE is atomic, one should make it a single 512-byte block. Thus, our current protocol to update the file system, with each of its three phases labeled:

1. **Journal write**: Write the contents of the transaction (including TxB, metadata, and data) to the log; wait for these writes to complete.
2. **Journal commit**: Write the transaction commit block (containing TxE) to the log; wait for write to complete; transaction is said to be **committed**.
3. **Checkpoint**: Write the contents of the update (metadata and data) to their final on-disk locations.

Recovery

Let’s now understand how a file system can use the contents of the journal to **recover** from a crash. A crash may happen at any time during this sequence of updates. If the crash happens before the transaction is written safely to the log (i.e., before Step 2 above completes), then our job is easy: the pending update is simply skipped. If the crash happens after the transaction has committed to the log, but before the checkpoint is complete, the file system can **recover** the update as follows. When the system boots, the file system recovery process will scan the log and look for transactions that have committed to the disk; these transactions are thus **replayed** (in order), with the file system again attempting to write out the blocks in the transaction to their final on-disk locations. This form of logging is one of the simplest forms there is, and is called **redo logging**. By recovering the committed transactions in the journal, the file system ensures that the on-disk structures are consistent, and thus can proceed by mounting the file system and readying itself for new requests.

Note that it is fine for a crash to happen at any point during checkpointing, even after some of the updates to the final locations of the blocks have completed. In the worst case, some of these updates are simply performed again during recovery. Because recovery is a rare operation (only taking place after an unexpected system crash), a few redundant writes are nothing to worry about.

**Batching Log Updates**

You might have noticed that the basic protocol could add a lot of extra disk traffic. For example, imagine we create two files in a row, called file1 and file2, in the same directory. To create one file, one has to update a number of on-disk structures, minimally including: the inode bitmap (to allocated a new inode), the newly-created inode of the file, the

---

3Unless you worry about everything, in which case we can’t help you. Stop worrying so much, it is unhealthy! But now you’re probably worried about over-worrying.
data block of the parent directory containing the new directory entry, as well as the parent directory inode (which now has a new modification time). With journaling, we logically commit all of this information to the journal for each of our two file creations; because the files are in the same directory, and let’s assume even have inodes within the same inode block, this means that if we’re not careful, we’ll end up writing these same blocks over and over.

To remedy this problem, some file systems do not commit each update to disk one at a time (e.g., Linux ext3); rather, one can buffer all updates into a global transaction. In our example above, when the two files are created, the file system just marks the in-memory inode bitmap, inodes of the files, directory data, and directory inode as dirty, and adds them to the list of blocks that form the current transaction. When it is finally time to write these blocks to disk (say, after a timeout of 5 seconds), this single global transaction is committed containing all of the updates described above. Thus, by buffering updates, a file system can avoid excessive write traffic to disk in many cases.

**Making The Log Finite**

We thus have arrived at a basic protocol for updating file-system on-disk structures. The file system buffers updates in memory for some time; when it is finally time to write to disk, the file system first carefully writes out the details of the transaction to the journal (a.k.a. write-ahead log); after the transaction is complete, the file system checkpoints those blocks to their final locations on disk.

However, the log is of a finite size. If we keep adding transactions to it (as in this figure), it will soon fill. What do you think happens then?

```
  Journal
Tx1  Tx2  Tx3  Tx4  Tx5 ...
```

Two problems arise when the log becomes full. The first is simpler, but less critical: the larger the log, the longer recovery will take, as the recovery process must replay all the transactions within the log (in order) to recover. The second is more of an issue: when the log is full (or nearly full), no further transactions can be committed to the disk, thus making the file system “less than useful” (i.e., useless).

To address these problems, journaling file systems treat the log as a circular data structure, re-using it over and over; this is why the journal is sometimes referred to as a **circular log**. To do so, the file system must take action some time after a checkpoint. Specifically, once a transaction has been checkpointed, the file system should free the space it was occupying within the journal, allowing the log space to be reused. There are many ways to achieve this end; for example, you could simply mark the oldest
and newest transactions in the log in a journal superblock; all other space is free. Here is a graphical depiction of such a mechanism:

![Journal Superblock Diagram](image)

In the journal superblock (not to be confused with the main file system superblock), the journaling system records enough information to know which transactions have not yet been checkpointed, and thus reduces recovery time as well as enables re-use of the log in a circular fashion. And thus we add another step to our basic protocol:

1. **Journal write**: Write the contents of the transaction (containing TxB and the contents of the update) to the log; wait for these writes to complete.
2. **Journal commit**: Write the transaction commit block (containing TxE) to the log; wait for the write to complete; the transaction is now committed.
3. **Checkpoint**: Write the contents of the update to their final locations within the file system.
4. **Free**: Some time later, mark the transaction free in the journal by updating the journal superblock.

Thus we have our final data journaling protocol. But there is still a problem: we are writing each data block to the disk twice, which is a heavy cost to pay, especially for something as rare as a system crash. Can you figure out a way to retain consistency without writing data twice?

### Metadata Journaling

Although recovery is now fast (scanning the journal and replaying a few transactions as opposed to scanning the entire disk), normal operation of the file system is slower than we might desire. In particular, for each write to disk, we are now also writing to the journal first, thus doubling write traffic; this doubling is especially painful during sequential write workloads, which now will proceed at half the peak write bandwidth of the drive. Further, between writes to the journal and writes to the main file system, there is a costly seek, which adds noticeable overhead for some workloads.

Because of the high cost of writing every data block to disk twice, people have tried a few different things in order to speed up performance. For example, the mode of journaling we described above is often called data journaling (as in Linux ext3), as it journals all user data (in addition to the metadata of the file system). A simpler (and more common) form of journaling is sometimes called ordered journaling (or just metadata
journaling), and it is nearly the same, except that user data is not written to the journal. Thus, when performing the same update as above, the following information would be written to the journal:

|---------|-----|-------|-------|-----|

The data block Db, previously written to the log, would instead be written to the file system proper, avoiding the extra write; given that most I/O traffic to the disk is data, not writing data twice substantially reduces the I/O load of journaling. The modification does raise an interesting question, though: when should we write data blocks to disk?

Let’s again consider our example append of a file to understand the problem better. The update consists of three blocks: I[v2], B[v2], and Db. The first two are both metadata and will be logged and then checkpointed; the latter will only be written once to the file system. When should we write Db to disk? Does it matter?

As it turns out, the ordering of the data write does matter for metadata-only journaling. For example, what if we write Db to disk after the transaction (containing I[v2] and B[v2]) completes? Unfortunately, this approach has a problem: the file system is consistent but I[v2] may end up pointing to garbage data. Specifically, consider the case where I[v2] and B[v2] are written but Db did not make it to disk. The file system will then try to recover. Because Db is not in the log, the file system will replay writes to I[v2] and B[v2], and produce a consistent file system (from the perspective of file-system metadata). However, I[v2] will be pointing to garbage data, i.e., at whatever was in the the slot where Db was headed.

To ensure this situation does not arise, some file systems (e.g., Linux ext3) write data blocks (of regular files) to the disk first, before related metadata is written to disk. Specifically, the protocol is as follows:

1. **Data write**: Write data to final location; wait for completion (the wait is optional; see below for details).
2. **Journal metadata write**: Write the begin block and metadata to the log; wait for writes to complete.
3. **Journal commit**: Write the transaction commit block (containing TxE) to the log; wait for the write to complete; the transaction (including data) is now committed.
4. **Checkpoint metadata**: Write the contents of the metadata update to their final locations within the file system.
5. **Free**: Later, mark the transaction free in journal superblock.

By forcing the data write first, a file system can guarantee that a pointer will never point to garbage. Indeed, this rule of “write the pointed to object before the object with the pointer to it” is at the core of crash consistency, and is exploited even further by other crash consistency schemes [GP94] (see below for details).
In most systems, metadata journaling (akin to ordered journaling of ext3) is more popular than full data journaling. For example, Windows NTFS and SGI’s XFS both use non-ordered metadata journaling. Linux ext3 gives you the option of choosing either data, ordered, or unordered modes (in unordered mode, data can be written at any time). All of these modes keep metadata consistent; they vary in their semantics for data.

Finally, note that forcing the data write to complete (Step 1) before issuing writes to the journal (Step 2) is not required for correctness, as indicated in the protocol above. Specifically, it would be fine to issue data writes as well as the transaction-begin block and metadata to the journal; the only real requirement is that Steps 1 and 2 complete before the issuing of the journal commit block (Step 3).

Tricky Case: Block Reuse

There are some interesting corner cases that make journaling more tricky, and thus are worth discussing. A number of them revolve around block reuse; as Stephen Tweedie (one of the main forces behind ext3) said:

“What’s the hideous part of the entire system? ... It’s deleting files. Everything to do with delete is hairy. Everything to do with delete... you have nightmares around what happens if blocks get deleted and then reallocated.” [T00]

The particular example Tweedie gives is as follows. Suppose you are using some form of metadata journaling (and thus data blocks for files are not journaled). Let’s say you have a directory called foo. The user adds an entry to foo (say by creating a file), and thus the contents of foo (because directories are considered metadata) are written to the log; assume the location of the foo directory data is block 1000. The log thus contains something like this:

<table>
<thead>
<tr>
<th>Journal</th>
<th>TxB id=1</th>
<th>I[foo] ptr:1000</th>
<th>TxE id=1</th>
</tr>
</thead>
<tbody>
<tr>
<td>D[foo]</td>
<td>final addr:1000</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

At this point, the user deletes everything in the directory as well as the directory itself, freeing up block 1000 for reuse. Finally, the user creates a new file (say foobar), which ends up reusing the same block (1000) that used to belong to foo. The inode of foobar is committed to disk, as is its data; note, however, because metadata journaling is in use, only the inode of foobar is committed to the journal; the newly-written data in block 1000 in the file foobar is not journaled.

<table>
<thead>
<tr>
<th>Journal</th>
<th>TxB id=1</th>
<th>I[foo] ptr:1000</th>
<th>TxE id=1</th>
</tr>
</thead>
<tbody>
<tr>
<td>D[foo]</td>
<td>final addr:1000</td>
<td></td>
<td></td>
</tr>
<tr>
<td>TxB id=2</td>
<td>I[foobar] ptr:1000</td>
<td>TxE id=2</td>
<td></td>
</tr>
</tbody>
</table>
Now assume a crash occurs and all of this information is still in the log. During replay, the recovery process simply replays everything in the log, including the write of directory data in block 1000; the replay thus overwrites the user data of current file `foobar` with old directory contents! Clearly this is not a correct recovery action, and certainly it will be a surprise to the user when reading the file `foobar`.

There are a number of solutions to this problem. One could, for example, never reuse blocks until the delete of said blocks is checkpointed out of the journal. What Linux ext3 does instead is to add a new type of record to the journal, known as a revoke record. In the case above, deleting the directory would cause a revoke record to be written to the journal. When replaying the journal, the system first scans for such revoke records; any such revoked data is never replayed, thus avoiding the problem mentioned above.

**Wrapping Up Journaling: A Timeline**

Before ending our discussion of journaling, we summarize the protocols we have discussed with timelines depicting each of them. Table 42.1 shows the protocol when journaling data as well as metadata, whereas Table 42.2 shows the protocol when journaling only metadata.

In each table, time increases in the downward direction, and each row in the table shows the logical time that a write can be issued or might complete. For example, in the data journaling protocol (42.1), the writes of the transaction begin block (TxB) and the contents of the transaction can logically be issued at the same time, and thus can be completed in any order; however, the write to the transaction end block (TxE) must not be issued until said previous writes complete. Similarly, the checkpointing writes to data and metadata blocks cannot begin until the transaction end block has committed. Horizontal dashed lines show where write-ordering requirements must be obeyed.

A similar timeline is shown for the metadata journaling protocol. Note that the data write can logically be issued at the same time as the writes
42.4 Solution #3: Other Approaches

We’ve thus far described two options in keeping file system metadata consistent: a lazy approach based on \textit{fsck}, and a more active approach known as journaling. However, these are not the only two approaches. One such approach, known as Soft Updates [GP94], was introduced by Ganger and Patt. This approach carefully orders all writes to the file system to ensure that the on-disk structures are never left in an inconsistent state. For example, by writing a pointed-to data block to disk before the inode that points to it, we can ensure that the inode never points to garbage; similar rules can be derived for all the structures of the file system. Implementing Soft Updates can be a challenge, however; whereas the journaling layer described above can be implemented with relatively little knowledge of the exact file system structures, Soft Updates requires intricate knowledge of each file system data structure and thus adds a fair amount of complexity to the system.

Another approach is known as \textit{copy-on-write} (yes, COW), and is used in a number of popular file systems, including Sun’s ZFS [B07]. This technique never overwrites files or directories in place; rather, it places new updates to previously unused locations on disk. After a number of updates are completed, COW file systems flip the root structure of the file system to include pointers to the newly updated structures. Doing so makes keeping the file system consistent straightforward. We’ll be learning more about this technique when we discuss the log-structured file system (LFS) in a future chapter; LFS is an early example of a COW.

\begin{table}[h]
\centering
\begin{tabular}{|c|c|c|}
\hline
Tx\(B\) & Journal Contents & Tx\(E\) \\
\hline
issue & issue & complete \\
complete & complete & complete \\
\hline
\end{tabular}
\caption{Metadata Journaling Timeline}
\end{table}

to the transaction begin and the contents of the journal; however, it must be issued and complete before the transaction end has been issued.

Finally, note that the time of completion marked for each write in the timelines is arbitrary. In a real system, completion time is determined by the I/O subsystem, which may reorder writes to improve performance. The only guarantees about ordering that we have are those that must be enforced for protocol correctness (and are shown via the horizontal dashed lines in the tables).
Another approach is one we just developed here at Wisconsin. In this technique, entitled backpointer-based consistency (or BBC), no ordering is enforced between writes. To achieve consistency, an additional backpointer is added to every block in the system; for example, each data block has a reference to the inode to which it belongs. When accessing a file, the file system can determine if the file is consistent by checking if the forward pointer (e.g., the address in the inode or direct block) points to a block that refers back to it. If so, everything must have safely reached disk and thus the file is consistent; if not, the file is inconsistent, and an error is returned. By adding back pointers to the file system, a new form of lazy crash consistency can be attained [C+12].

Finally, we also have explored techniques to reduce the number of times a journal protocol has to wait for disk writes to complete. Entitled optimistic crash consistency [C+13], this new approach issues as many writes to disk as possible and uses a generalized form of the transaction checksum [P+05], as well as a few other techniques, to detect inconsistencies should they arise. For some workloads, these optimistic techniques can improve performance by an order of magnitude. However, to truly function well, a slightly different disk interface is required [C+13].

42.5 Summary

We have introduced the problem of crash consistency, and discussed various approaches to attacking this problem. The older approach of building a file system checker works but is likely too slow to recover on modern systems. Thus, many file systems now use journaling. Journaling reduces recovery time from $O(\text{size-of-the-disk-volume})$ to $O(\text{size-of-the-log})$, thus speeding recovery substantially after a crash and restart. For this reason, many modern file systems use journaling. We have also seen that journaling can come in many different forms; the most commonly used is ordered metadata journaling, which reduces the amount of traffic to the journal while still preserving reasonable consistency guarantees for both file system metadata as well as user data.
References

[B07] “ZFS: The Last Word in File Systems”
Jeff Bonwick and Bill Moore
ZFS uses copy-on-write and journaling, actually, as in some cases, logging writes to disk will perform better.

[C+12] “Consistency Without Ordering”
Vijay Chidambaram, Tushar Sharma, Andrea C. Arpaci-Dusseau, Remzi H. Arpaci-Dusseau
FAST ’12, San Jose, California
A recent paper of ours about a new form of crash consistency based on back pointers. Read it for the exciting details!

[C+13] “Optimistic Crash Consistency”
Vijay Chidambaram, Thanu S. Pillai, Andrea C. Arpaci-Dusseau, Remzi H. Arpaci-Dusseau
SOSP ’13, Nemacolin Woodlands Resort, PA, November 2013
Our work on a more optimistic and higher performance journaling protocol. For workloads that call \texttt{fsync()} a lot, performance can be greatly improved.

[GP94] “Metadata Update Performance in File Systems”
Gregory R. Ganger and Yale N. Patt
OSDI ’94
A clever paper about using careful ordering of writes as the main way to achieve consistency. Implemented later in BSD-based systems.

[G+08] “SQCK: A Declarative File System Checker”
Haryadi S. Gunawi, Abhishek Rajimwale, Andrea C. Arpaci-Dusseau, Remzi H. Arpaci-Dusseau
OSDI ’08, San Diego, California
Our own paper on a new and better way to build a file system checker using SQL queries. We also show some problems with the existing checker, finding numerous bugs and odd behaviors, a direct result of the complexity of \texttt{fsck}.

[H87] “Reimplementing the Cedar File System Using Logging and Group Commit”
Robert Hagmann
SOSP ’87, Austin, Texas, November 1987
The first work (that we know of) that applied write-ahead logging (a.k.a. journaling) to a file system.

Ao Ma, Chris Dragga, Andrea C. Arpaci-Dusseau, Remzi H. Arpaci-Dusseau
FAST ’13, San Jose, California, February 2013
A recent paper of ours detailing how to make \texttt{fsck} an order of magnitude faster. Some of the ideas have already been incorporated into the BSD file system checker [MK96] and are deployed today.

[MK96] “Fscck - The UNIX File System Check Program”
Marshall Kirk McKusick and T. J. Kowalski
Revised in 1996
Describes the first comprehensive file-system checking tool, the eponymous \texttt{fsck}. Written by some of the same people who brought you FFS.

[MJLF84] “A Fast File System for UNIX”
Marshall K. McKusick, William N. Joy, Sam J. Leffler, Robert S. Fabry
ACM Transactions on Computing Systems.
August 1984, Volume 2:3
You already know enough about FFS, right? But yeah, it is OK to reference papers like this more than once in a book.
Crash Consistency: FSCK and Journaling

[P+05] “IRON File Systems”
Vijayan Prabhakaran, Lakshmi N. Bairavasundaram, Nitin Agrawal, Haryadi S. Gunawi, Andrea C. Arpaci-Dusseau, Remzi H. Arpaci-Dusseau
SOSP ’05, Brighton, England, October 2005
A paper mostly focused on studying how file systems react to disk failures. Towards the end, we introduce a transaction checksum to speed up logging, which was eventually adopted into Linux ext4.

[PA05] “Analysis and Evolution of Journaling File Systems”
Vijayan Prabhakaran, Andrea C. Arpaci-Dusseau, Remzi H. Arpaci-Dusseau
USENIX ’05, Anaheim, California, April 2005
An early paper we wrote analyzing how journaling file systems work.

Abhishek Rajimwale, Vijay Chidambaram, Deepak Ramamurthi, Andrea C. Arpaci-Dusseau, Remzi H. Arpaci-Dusseau
DSN ’11, Hong Kong, China, June 2011
Our own paper on the problem of disks that buffer writes in a memory cache instead of forcing them to disk, even when explicitly told not to do that! Our solution to overcome this problem: if you want A to be written to disk before B, first write A, then send a lot of “dummy” writes to disk, hopefully causing A to be forced to disk to make room for them in the cache. A neat if impractical solution.

[T98] “Journaling the Linux ext2fs File System”
Stephen C. Tweedie
The Fourth Annual Linux Expo, May 1998
Tweedie did much of the heavy lifting in adding journaling to the Linux ext2 file system; the result, not surprisingly, is called ext3. Some nice design decisions include the strong focus on backwards compatibility, e.g., you can just add a journaling file to an existing ext2 file system and then mount it as an ext3 file system.

[T00] “EXT3, Journaling Filesystem”
Stephen Tweedie
Talk at the Ottawa Linux Symposium, July 2000
A transcript of a talk given by Tweedie on ext3.

[T01] “The Linux ext2 File System”
Available: http://e2fsprogs.sourceforge.net/ext2.html
A simple Linux file system based on the ideas found in FFS. For a while it was quite heavily used; now it is really just in the kernel as an example of a simple file system.
In the early 90’s, a group at Berkeley led by Professor John Ousterhout and graduate student Mendel Rosenblum developed a new file system known as the log-structured file system [RO91]. Their motivation to do so was based on the following observations:

- **Memory sizes were growing**: As memory got bigger, more data could be cached in memory. As more data is cached, disk traffic would increasingly consist of writes, as reads would be serviced in the cache. Thus, file system performance would largely be determined by its performance for writes.

- **There was a large and growing gap between random I/O performance and sequential I/O performance**: Transfer bandwidth increases roughly 50%-100% every year; seek and rotational delay costs decrease much more slowly, maybe at 5%-10% per year [P98]. Thus, if one is able to use disks in a sequential manner, one gets a huge performance advantage, which grows over time.

- **Existing file systems perform poorly on many common workloads**: For example, FFS [MJLF84] would perform a large number of writes to create a new file of size one block: one for a new inode, one to update the inode bitmap, one to the directory data block that the file is in, one to the directory inode to update it, one to the new data block that is apart of the new file, and one to the data bitmap to mark the data block as allocated. Thus, although FFS would place all of these blocks within the same block group, FFS would incur many short seeks and subsequent rotational delays and thus performance would fall far short of peak sequential bandwidth.

- **File systems were not RAID-aware**: For example, RAID-4 and RAID-5 have the small-write problem where a logical write to a single block causes 4 physical I/Os to take place. Existing file systems do not try to avoid this worst-case RAID writing behavior.

An ideal file system would thus focus on write performance, and try to make use of the sequential bandwidth of the disk. Further, it would perform well on common workloads that not only write out data but also
update on-disk metadata structures frequently. Finally, it would work well on RAIDs as well as single disks.

The new type of file system Rosenblum and Ousterhout introduced was called LFS, short for the Log-structured File System. When writing to disk, LFS first buffers all updates (including metadata!) in an in-memory segment; when the segment is full, it is written to disk in one long, sequential transfer to an unused part of the disk, i.e., LFS never overwrites existing data, but rather always writes segments to free locations. Because segments are large, the disk is used efficiently, and performance of the file system approaches its zenith.

**THE CRUX:**
**HOW TO MAKE ALL WRITES SEQUENTIAL WRITES?**

How can a file system turns all writes into sequential writes? For reads, this task is impossible, as the desired block to be read may be anywhere on disk. For writes, however, the file system always has a choice, and it is exactly this choice we hope to exploit.

### 43.1 Writing To Disk Sequentially

We thus have our first challenge: how do we transform all updates to file-system state into a series of sequential writes to disk? To understand this better, let’s use a simple example. Imagine we are writing a data block \( D \) to a file. Writing the data block to disk might result in the following on-disk layout, with \( D \) written at disk address \( A0 \):

\[
\begin{array}{c}
\text{D} \\
A0
\end{array}
\]

However, when a user writes a data block, it is not only data that gets written to disk; there is also other metadata that needs to be updated. In this case, let’s also write the inode \((I)\) of the file to disk, and have it point to the data block \( D \). When written to disk, the data block and inode would look something like this (note that the inode looks as big as the data block, which generally isn’t the case; in most systems, data blocks are 4 KB in size, whereas an inode is much smaller, around 128 bytes):

\[
\begin{array}{c}
\text{D} \\
\text{blk[0]:A0} \\
I \\
A0
\end{array}
\]
All interesting systems are comprised of a few general ideas and a number of details. Sometimes, when you are learning about these systems, you think to yourself “Oh, I get the general idea; the rest is just details,” and you use this to only half-learn how things really work. Don’t do this! Many times, the details are critical. As we’ll see with LFS, the general idea is easy to understand, but to really build a working system, you have to think through all of the tricky cases.

This basic idea, of simply writing all updates (such as data blocks, inodes, etc.) to the disk sequentially, sits at the heart of LFS. If you understand this, you get the basic idea. But as with all complicated systems, the devil is in the details.

### 43.2 Writing Sequentially And Effectively

Unfortunately, writing to disk sequentially is not (alone) enough to guarantee efficient writes. For example, imagine if we wrote a single block to address \( A \), at time \( T \). We then wait a little while, and write to the disk at address \( A + 1 \) (the next block address in sequential order), but at time \( T + \delta \). In-between the first and second writes, unfortunately, the disk has rotated; when you issue the second write, it will thus wait for most of a rotation before being committed (specifically, if the rotation takes time \( T_{\text{rotation}} \), the disk will wait \( T_{\text{rotation}} - \delta \) before it can commit the second write to the disk surface). And thus you can hopefully see that simply writing to disk in sequential order is not enough to achieve peak performance; rather, you must issue a large number of contiguous writes (or one large write) to the drive in order to achieve good write performance.

To achieve this end, LFS uses an ancient technique known as write buffering\(^1\). Before writing to the disk, LFS keeps track of updates in memory; when it has received a sufficient number of updates, it writes them to disk all at once, thus ensuring efficient use of the disk.

The large chunk of updates LFS writes at one time is referred to by the name of a segment. Although this term is over-used in computer systems, here it just means a large-ish chunk which LFS uses to group writes. Thus, when writing to disk, LFS buffers updates in an in-memory segment, and then writes the segment all at once to the disk. As long as the segment is large enough, these writes will be efficient.

Here is an example, in which LFS buffers two sets updates into a small segment; actual segments are larger (a few MB). The first update is of

\(^1\)Indeed, it is hard to find a good citation for this idea, since it was likely invented by many and very early on in the history of computing. For a study of the benefits of write buffering, see Solworth and Orji [SO90]; to learn about its potential harms, see Mogul [M94].
four block writes to file \( j \); the second is one block being added to file \( k \). LFS then commits the entire segment of seven blocks to disk at once. The resulting on-disk layout of these blocks is as follows:

![Diagram showing on-disk layout of blocks]

### 43.3 How Much To Buffer?

This raises the following question: how many updates LFS should buffer before writing to disk? The answer, of course, depends on the disk itself, specifically how high the positioning overhead is in comparison to the transfer rate; see the FFS chapter for a similar analysis.

For example, assume that positioning (i.e., rotation and seek overheads) before each write takes roughly \( T_{position} \) seconds. Assume further that the disk transfer rate is \( R_{peak} \) MB/s. How much should LFS buffer before writing when running on such a disk?

The way to think about this is that every time you write, you pay a fixed overhead of the positioning cost. Thus, how much do you have to write in order to amortize that cost? The more you write, the better (obviously), and the closer you get to achieving peak bandwidth.

To obtain a concrete answer, let’s assume we are writing out \( D \) MB. The time to write out this chunk of data (\( T_{write} \)) is the positioning time \( T_{position} \) plus the time to transfer \( D \) (\( DR_{peak} \)), or:

\[
T_{write} = T_{position} + \frac{D}{R_{peak}}
\]

And thus the effective rate of writing (\( R_{effective} \)), which is just the amount of data written divided by the total time to write it, is:

\[
R_{effective} = \frac{D}{T_{write}} = \frac{D}{T_{position} + \frac{D}{R_{peak}}}
\]

What we’re interested in is getting the effective rate (\( R_{effective} \)) close to the peak rate. Specifically, we want the effective rate to be some fraction \( F \) of the peak rate, where \( 0 < F < 1 \) (a typical \( F \) might be 0.9, or 90% of the peak rate). In mathematical form, this means we want \( R_{effective} = F \times R_{peak} \).

At this point, we can solve for \( D \):

\[
R_{effective} = \frac{D}{T_{position} + \frac{D}{R_{peak}}} = F \times R_{peak}
\]
\[ D = F \times R_{\text{peak}} \times (T_{\text{position}} + \frac{D}{R_{\text{peak}}}) \]  

(43.4)

\[ D = (F \times R_{\text{peak}} \times T_{\text{position}}) + (F \times R_{\text{peak}} \times \frac{D}{R_{\text{peak}}}) \]  

(43.5)

\[ D = \frac{F}{1 - F} \times R_{\text{peak}} \times T_{\text{position}} \]  

(43.6)

Let’s do an example, with a disk with a positioning time of 10 milliseconds and peak transfer rate of 100 MB/s; assume we want an effective bandwidth of 90% of peak \((F = 0.9)\). In this case, \(D = \frac{0.9}{0.1} \times 100 \text{ MB/s} \times 0.01 \text{ seconds} = 9 \text{ MB}\). Try some different values to see how much we need to buffer in order to approach peak bandwidth. How much is needed to reach 95% of peak? 99%?

43.4 Problem: Finding Inodes

To understand how we find an inode in LFS, let us briefly review how to find an inode in a typical UNIX file system. In a typical file system such as FFS, or even the old UNIX file system, finding inodes is easy, because they are organized in an array and placed on disk at fixed locations.

For example, the old UNIX file system keeps all inodes at a fixed portion of the disk. Thus, given an inode number and the start address, to find a particular inode, you can calculate its exact disk address simply by multiplying the inode number by the size of an inode, and adding that to the start address of the on-disk array; array-based indexing, given an inode number, is fast and straightforward.

Finding an inode given an inode number in FFS is only slightly more complicated, because FFS splits up the inode table into chunks and places a group of inodes within each cylinder group. Thus, one must know how big each chunk of inodes is and the start addresses of each. After that, the calculations are similar and also easy.

In LFS, life is more difficult. Why? Well, we’ve managed to scatter the inodes all throughout the disk! Worse, we never overwrite in place, and thus the latest version of an inode (i.e., the one we want) keeps moving.

43.5 Solution Through Indirection: The Inode Map

To remedy this, the designers of LFS introduced a level of indirection between inode numbers and the inodes through a data structure called the \textbf{inode map (imap)}. The imap is a structure that takes an inode number as input and produces the disk address of the most recent version of the inode. Thus, you can imagine it would often be implemented as a simple array, with 4 bytes (a disk pointer) per entry. Any time an inode is written to disk, the imap is updated with its new location.
People often say that the solution to all problems in Computer Science is simply a level of indirection. This is clearly not true; it is just the solution to most problems. You certainly can think of every virtualization we have studied, e.g., virtual memory, as simply a level of indirection. And certainly the inode map in LFS is a virtualization of inode numbers. Hopefully you can see the great power of indirection in these examples, allowing us to freely move structures around (such as pages in the VM example, or inodes in LFS) without having to change every reference to them. Of course, indirection can have a downside too: extra overhead. So next time you have a problem, try solving it with indirection. But make sure to think about the overheads of doing so first.

The imap, unfortunately, needs to be kept persistent (i.e., written to disk); doing so allows LFS to keep track of the locations of inodes across crashes, and thus operate as desired. Thus, a question: where should the imap reside on disk?

It could live on a fixed part of the disk, of course. Unfortunately, as it gets updated frequently, this would then require updates to file structures to be followed by writes to the imap, and hence performance would suffer (i.e., there would be more disk seeks, between each update and the fixed location of the imap).

Instead, LFS places chunks of the inode map right next to where it is writing all of the other new information. Thus, when appending a data block to a file \( k \), LFS actually writes the new data block, its inode, and a piece of the inode map all together onto the disk, as follows:

![Diagram](image)

In this picture, the piece of the imap array stored in the block marked \( imap \) tells LFS that the inode \( k \) is at disk address \( A1 \); this inode, in turn, tells LFS that its data block \( D \) is at address \( A0 \).

43.6 The Checkpoint Region

The clever reader (that’s you, right?) might have noticed a problem here. How do we find the inode map, now that pieces of it are also now spread across the disk? In the end, there is no magic: the file system must have some fixed and known location on disk to begin a file lookup.
LFS has just such a fixed place on disk for this, known as the **checkpoint region (CR)**. The checkpoint region contains pointers to (i.e., addresses of) the latest pieces of the inode map, and thus the inode map pieces can be found by reading the CR first. Note the checkpoint region is only updated periodically (say every 30 seconds or so), and thus performance is not ill-affected. Thus, the overall structure of the on-disk layout contains a checkpoint region (which points to the latest pieces of the inode map); the inode map pieces each contain addresses of the inodes; the inodes point to files (and directories) just like typical UNIX file systems.

Here is an example of the checkpoint region (note it is all the way at the beginning of the disk, at address 0), and a single imap chunk, inode, and data block. A real file system would of course have a much bigger CR (indeed, it would have two, as we’ll come to understand later), many imap chunks, and of course many more inodes, data blocks, etc.

---

**43.7 Reading A File From Disk: A Recap**

To make sure you understand how LFS works, let us now walk through what must happen to read a file from disk. Assume we have nothing in memory to begin. The first on-disk data structure we must read is the checkpoint region. The checkpoint region contains pointers (i.e., disk addresses) to the entire inode map, and thus LFS then reads in the entire inode map and caches it in memory. After this point, when given an inode number of a file, LFS simply looks up the inode-number to inode-disk-address mapping in the imap, and reads in the most recent version of the inode. To read a block from the file, at this point, LFS proceeds exactly as a typical UNIX file system, by using direct pointers or indirect pointers or doubly-indirect pointers as need be. In the common case, LFS should perform the same number of I/Os as a typical file system when reading a file from disk; the entire imap is cached and thus the extra work LFS does during a read is to look up the inode’s address in the imap.

---

**43.8 What About Directories?**

Thus far, we’ve simplified our discussion a bit by only considering inodes and data blocks. However, to access a file in a file system (such as /home/remzi/foo, one of our favorite fake file names), some directories must be accessed too. So how does LFS store directory data?
Fortunately, directory structure is basically identical to classic UNIX file systems, in that a directory is just a collection of (name, inode number) mappings. For example, when creating a file on disk, LFS must both write a new inode, some data, as well as the directory data and its inode that refer to this file. Remember that LFS will do so sequentially on the disk (after buffering the updates for some time). Thus, creating a file `foo` in a directory would lead to the following new structures on disk:

![Diagram showing directory and file structures]

The piece of the inode map contains the information for the location of both the directory file `dir` as well as the newly-created file `f`. Thus, when accessing file `foo` (with inode number `f`), you would first look in the inode map (usually cached in memory) to find the location of the inode of directory `dir` (A3); you then read the directory inode, which gives you the location of the directory data (A2); reading this data block gives you the name-to-inode-number mapping of `(foo, k)`. You then consult the inode map again to find the location of inode number `k` (A1), and finally read the desired data block at address A0.

There is one other serious problem in LFS that the inode map solves, known as the recursive update problem [Z+12]. The problem arises in any file system that never updates in place (such as LFS), but rather moves updates to new locations on the disk.

Specifically, whenever an inode is updated, its location on disk changes. If we hadn’t been careful, this would have also entailed an update to the directory that points to this file, which then would have mandated a change to the parent of that directory, and so on, all the way up the file system tree.

LFS cleverly avoids this problem with the inode map. Even though the location of an inode may change, the change is never reflected in the directory itself; rather, the imap structure is updated while the directory holds the same name-to-inumber mapping. Thus, through indirection, LFS avoids the recursive update problem.

43.9 A New Problem: Garbage Collection

You may have noticed another problem with LFS; it keeps writing newer version of a file, its inode, and in fact all data to new parts of the disk. This process, while keeping writes efficient, implies that LFS leaves older versions of file structures all over the disk, scattered throughout the disk. We call such old stuff garbage.
For example, let’s imagine the case where we have an existing file referred to by inode number $k$, which points to a single data block $D_0$. We now overwrite that block, generating both a new inode and a new data block. The resulting on-disk layout of LFS would look something like this (note we omit the imap and other structures for simplicity; a new chunk of imap would also have to be written to disk to point to the new inode):

In the diagram, you can see that both the inode and data block have two versions on disk, one old (the one on the left) and one current and thus live (the one on the right). By the simple act of overwriting a data block, a number of new structures must be persisted by LFS, thus leaving old versions of said blocks on the disk.

As another example, imagine we instead append a block to that original file $k$. In this case, a new version of the inode is generated, but the old data block is still pointed to by the inode. Thus, it is still live and very much apart of the current file system:

So what should we do with these older versions of inodes, data blocks, and so forth? One could keep those older versions around and allow users to restore old file versions (for example, when they accidentally overwrite or delete a file, it could be quite handy to do so); such a file system is known as a versioning file system because it keeps track of the different versions of a file.

However, LFS instead keeps only the latest live version of a file; thus (in the background), LFS must periodically find these old dead versions of file data, inodes, and other structures, and clean them; cleaning should thus make blocks on disk free again for use in a subsequent writes. Note that the process of cleaning is a form of garbage collection, a technique that arises in programming languages that automatically free unused memory for programs.

Earlier we discussed segments as important as they are the mechanism that enables large writes to disk in LFS. As it turns out, they are also quite integral to effective cleaning. Imagine what would happen if the LFS
cleaner simply went through and freed single data blocks, inodes, etc., during cleaning. The result: a file system with some number of free holes mixed between allocated space on disk. Write performance would drop considerably, as LFS would not be able to find a large contiguous region to write to disk sequentially and with high performance.

Instead, the LFS cleaner works on a segment-by-segment basis, thus clearing up large chunks of space for subsequent writing. The basic cleaning process works as follows. Periodically, the LFS cleaner reads in a number of old (partially-used) segments, determines which blocks are live within these segments, and then write out a new set of segments with just the live blocks within them, freeing up the old ones for writing. Specifically, we expect the cleaner to read in \( M \) existing segments, compact their contents into \( N \) new segments (where \( N < M \)), and then write the \( N \) segments to disk in new locations. The old \( M \) segments are then freed and can be used by the file system for subsequent writes.

We are now left with two problems, however. The first is mechanism: how can LFS tell which blocks within a segment are live, and which are dead? The second is policy: how often should the cleaner run, and which segments should it pick to clean?

43.10 Determining Block Liveness

We address the mechanism first. Given a data block \( D \) within an on-disk segment \( S \), LFS must be able to determine whether \( D \) is live. To do so, LFS adds a little extra information to each segment that describes each block. Specifically, LFS includes, for each data block \( D \), its inode number (which file it belongs to) and its offset (which block of the file this is). This information is recorded in a structure at the head of the segment known as the segment summary block.

Given this information, it is straightforward to determine whether a block is live or dead. For a block \( D \) located on disk at address \( A \), look in the segment summary block and find its inode number \( N \) and offset \( T \). Next, look in the imap to find where \( N \) lives and read \( N \) from disk (perhaps it is already in memory, which is even better). Finally, using the offset \( T \), look in the inode (or some indirect block) to see where the inode thinks the \( T \)th block of this file is on disk. If it points exactly to disk address \( A \), LFS can conclude that the block \( D \) is live. If it points anywhere else, LFS can conclude that \( D \) is not in use (i.e., it is dead) and thus know that this version is no longer needed. A pseudocode summary of this process is shown here:

\[
\begin{align*}
(N, T) &= \text{SegmentSummary}[A]; \\
\text{inode} &= \text{Read(imap}[N]); \\
\text{if } (\text{inode}[T] == A) \\
\quad &\text{// block } D \text{ is alive} \\
\text{else} \\
\quad &\text{// block } D \text{ is garbage}
\end{align*}
\]
Here is a diagram depicting the mechanism, in which the segment summary block (marked \textit{SS}) records that the data block at address $A0$ is actually a part of file $k$ at offset 0. By checking the imap for $k$, you can find the inode, and see that it does indeed point to that location.

There are some shortcuts LFS takes to make the process of determining liveness more efficient. For example, when a file is truncated or deleted, LFS increases its \textit{version number} and records the new version number in the imap. By also recording the version number in the on-disk segment, LFS can short circuit the longer check described above simply by comparing the on-disk version number with a version number in the imap, thus avoiding extra reads.

43.11 A Policy Question: Which Blocks To Clean, And When?

On top of the mechanism described above, LFS must include a set of policies to determine both when to clean and which blocks are worth cleaning. Determining when to clean is easier; either periodically, during idle time, or when you have to because the disk is full.

Determining which blocks to clean is more challenging, and has been the subject of many research papers. In the original LFS paper [RO91], the authors describe an approach which tries to segregate \textit{hot} and \textit{cold} segment. A hot segment is one in which the contents are being frequently over-written; thus, for such a segment, the best policy is to wait a long time before cleaning it, as more and more blocks are getting over-written (in new segments) and thus being freed for use. A cold segment, in contrast, may have a few dead blocks but the rest of its contents are relatively stable. Thus, the authors conclude that one should clean cold segments sooner and hot segments later, and develop a heuristic that does exactly that. However, as with most policies, this is just one approach, and by definition is not “the best” approach; later approaches show how to do better [MR+97].

43.12 Crash Recovery And The Log

One final problem: what happens if the system crashes while LFS is writing to disk? As you may recall in the previous chapter about journaling, crashes during updates are tricky for file systems, and thus some-
thing LFS must consider as well.

During normal operation, LFS buffers writes in a segment, and then (when the segment is full, or when some amount of time has elapsed), writes the segment to disk. LFS organizes these writes in a log, i.e., the checkpoint region points to a head and tail segment, and each segment points to the next segment to be written. LFS also periodically updates the checkpoint region. Crashes could clearly happen during either of these operations (write to a segment, write to the CR). So how does LFS handle crashes during writes to these structures?

Let’s cover the second case first. To ensure that the CR update happens atomically, LFS actually keeps two CRs, one at either end of the disk, and writes to them alternately. LFS also implements a careful protocol when updating the CR with the latest pointers to the inode map and other information; specifically, it first writes out a header (with timestamp), then the body of the CR, and then finally one last block (also with a timestamp). If the system crashes during a CR update, LFS can detect this by seeing an inconsistent pair of timestamps. LFS will always choose to use the most recent CR that has consistent timestamps, and thus consistent update of the CR is achieved.

Let’s now address the first case. Because LFS writes the CR every 30 seconds or so, the last consistent snapshot of the file system may be quite old. Thus, upon reboot, LFS can easily recover by simply reading in the checkpoint region, the imap pieces it points to, and subsequent files and directories; however, the last many seconds of updates would be lost.

To improve upon this, LFS tries to rebuild many of those segments through a technique known as roll forward in the database community. The basic idea is to start with the last checkpoint region, find the end of the log (which is included in the CR), and then use that to read through the next segments and see if there are any valid updates within it. If there are, LFS updates the file system accordingly and thus recovers much of the data and metadata written since the last checkpoint. See Rosenblum’s award-winning dissertation for details [R92].

43.13 Summary

LFS introduces a new approach to updating the disk. Instead of overwriting files in places, LFS always writes to an unused portion of the disk, and then later reclaims that old space through cleaning. This approach, which in database systems is called shadow paging [L77] and in file-system-speak is sometimes called copy-on-write, enables highly efficient writing, as LFS can gather all updates into an in-memory segment and then write them out together sequentially.

The downside to this approach is that it generates garbage; old copies of the data are scattered throughout the disk, and if one wants to reclaim such space for subsequent usage, one must clean old segments periodically. Cleaning became the focus of much controversy in LFS, and con-
Whenever your system has a fundamental flaw, see if you can turn it around into a feature or something useful. NetApp’s WAFL does this with old file contents; by making old versions available, WAFL no longer has to worry about cleaning, and thus provides a cool feature and removes the LFS cleaning problem all in one wonderful twist. Are there other examples of this in systems? Undoubtedly, but you’ll have to think of them yourself, because this chapter is over with a capital “O”. Over. Done. Kaput. We’re out. Peace!

cerns over cleaning costs [SS+95] perhaps limited LFS’s initial impact on the field. However, some modern commercial file systems, including NetApp’s WAFL [HLM94], Sun’s ZFS [B07], and Linux btrfs [M07] adopt a similar copy-on-write approach to writing to disk, and thus the intellectual legacy of LFS lives on in these modern file systems. In particular, WAFL got around cleaning problems by turning them into a feature; by providing old versions of the file system via snapshots, users could access old files whenever they deleted current ones accidentally.
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Data Integrity and Protection

Beyond the basic advances found in the file systems we have studied thus far, a number of features are worth studying. In this chapter, we focus on reliability once again (having previously studied storage system reliability in the RAID chapter). Specifically, how should a file system or storage system ensure that data is safe, given the unreliable nature of modern storage devices?

This general area is referred to as **data integrity** or **data protection**. Thus, we will now investigate techniques used to ensure that the data you put into your storage system is the same when the storage system returns it to you.

**Crux: How To Ensure Data Integrity**

How should systems ensure that the data written to storage is protected? What techniques are required? How can such techniques be made efficient, with both low space and time overheads?

44.1 Disk Failure Modes

As you learned in the chapter about RAID, disks are not perfect, and can fail (on occasion). In early RAID systems, the model of failure was quite simple: either the entire disk is working, or it fails completely, and the detection of such a failure is straightforward. This **fail-stop** model of disk failure makes building RAID relatively simple [S90].

What you didn’t learn is about all of the other types of failure modes modern disks exhibit. Specifically, as Bairavasundaram et al. studied in great detail [B+07, B+08], modern disks will occasionally seem to be mostly working but have trouble successfully accessing one or more blocks. Specifically, two types of single-block failures are common and worthy of consideration: **latent-sector errors (LSEs)** and **block corruption**. We’ll now discuss each in more detail.
Cheap | Costly
---- | ----
LSEs  9.40% | 1.40%
Corruption  0.50% | 0.05%

Table 44.1: Frequency of LSEs and Block Corruption

LSEs arise when a disk sector (or group of sectors) has been damaged in some way. For example, if the disk head touches the surface for some reason (a head crash, something which shouldn’t happen during normal operation), it may damage the surface, making the bits unreadable. Cosmic rays can also flip bits, leading to incorrect contents. Fortunately, in-disk error correcting codes (ECC) are used by the drive to determine whether the on-disk bits in a block are good, and in some cases, to fix them; if they are not good, and the drive does not have enough information to fix the error, the disk will return an error when a request is issued to read them.

There are also cases where a disk block becomes corrupt in a way not detectable by the disk itself. For example, buggy disk firmware may write a block to the wrong location; in such a case, the disk ECC indicates the block contents are fine, but from the client’s perspective the wrong block is returned when subsequently accessed. Similarly, a block may get corrupted when it is transferred from the host to the disk across a faulty bus; the resulting corrupt data is stored by the disk, but it is not what the client desires. These types of faults are particularly insidious because the are silent faults; the disk gives no indication of the problem when returning the faulty data.

Prabhakaran et al. describes this more modern view of disk failure as the fail-partial disk failure model [P+05]. In this view, disks can still fail in their entirety (as was the case in the traditional fail-stop model); however, disks can also seemingly be working and have one or more blocks become inaccessible (i.e., LSEs) or hold the wrong contents (i.e., corruption). Thus, when accessing a seemingly-working disk, once in a while it may either return an error when trying to read or write a given block (a non-silent partial fault), and once in a while it may simply return the wrong data (a silent partial fault).

Both of these types of faults are somewhat rare, but just how rare? Table 44.1 summarizes some of the findings from the two Bairavasundaram studies [B+07,B+08].

The table shows the percent of drives that exhibited at least one LSE or block corruption over the course of the study (about 3 years, over 1.5 million disk drives). The table further sub-divides the results into “cheap” drives (usually SATA drives) and “costly” drives (usually SCSI or FibreChannel). As you can see from the table, while buying better drives reduces the frequency of both types of problem (by about an order of magnitude), they still happen often enough that you need to think carefully about them.
Some additional findings about LSEs are:

- Costly drives with more than one LSE are as likely to develop additional errors as cheaper drives
- For most drives, annual error rate increases in year two
- LSEs increase with disk size
- Most disks with LSEs have less than 50
- Disks with LSEs are more likely to develop additional LSEs
- There exists a significant amount of spatial and temporal locality
- Disk scrubbing is useful (most LSEs were found this way)

Some findings about corruption:

- Chance of corruption varies greatly across different drive models within the same drive class
- Age affects are different across models
- Workload and disk size have little impact on corruption
- Most disks with corruption only have a few corruptions
- Corruption is not independent with a disk or across disks in RAID
- There exists spatial locality, and some temporal locality
- There is a weak correlation with LSEs

To learn more about these failures, you should likely read the original papers [B+07,B+08]. But hopefully the main point should be clear: if you really wish to build a reliable storage system, you must include machinery to detect and recovery from both LSEs and block corruption.

44.2 Handling Latent Sector Errors

Given these two new modes of partial disk failure, we should now try to see what we can do about them. Let’s first tackle the easier of the two, namely latent sector errors.

CRUX: HOW TO HANDLE LATENT SECTOR ERRORS

How should a storage system handle latent sector errors? How much extra machinery is needed to handle this form of partial failure?

As it turns out, latent sector errors are rather straightforward to handle, as they are (by definition) easily detected. When a storage system tries to access a block, and the disk returns an error, the storage system should simply use whatever redundancy mechanism it has to return the correct data. In a mirrored RAID, for example, the system should access the alternate copy; in a RAID-4 or RAID-5 system based on parity, the system should reconstruct the block from the other blocks in the parity group. Thus, easily detected problems such as LSEs are readily recovered through standard redundancy mechanisms.
The growing prevalence of LSEs has influenced RAID designs over the years. One particularly interesting problem arises in RAID-4/5 systems when both full-disk faults and LSEs occur in tandem. Specifically, when an entire disk fails, the RAID tries to reconstruct the disk (say, onto a hot spare) by reading through all of the other disks in the parity group and recomputing the missing values. If, during reconstruction, an LSE is encountered on any one of the other disks, we have a problem: the reconstruction cannot successfully complete.

To combat this issue, some systems add an extra degree of redundancy. For example, NetApp’s RAID-DP has the equivalent of two parity disks instead of one [C+04]. When an LSE is discovered during reconstruction, the extra parity helps to reconstruct the missing block. As always, there is a cost, in that maintaining two parity blocks for each stripe is more costly; however, the log-structured nature of the NetApp WAFL file system mitigates that cost in many cases [HLM94]. The remaining cost is space, in the form of an extra disk for the second parity block.

44.3 Detecting Corruption: The Checksum

Let’s now tackle the more challenging problem, that of silent failures via data corruption. How can we prevent users from getting bad data when corruption arises, and thus leads to disks returning bad data?

**Crux: How To Preserve Data Integrity Despite Corruption**

Given the silent nature of such failures, what can a storage system do to detect when corruption arises? What techniques are needed? How can one implement them efficiently?

Unlike latent sector errors, detection of corruption is a key problem. How can a client tell that a block has gone bad? Once it is known that a particular block is bad, recovery is the same as before: you need to have some other copy of the block around (and hopefully, one that is not corrupt!). Thus, we focus here on detection techniques.

The primary mechanism used by modern storage systems to preserve data integrity is called the checksum. A checksum is simply the result of a function that takes a chunk of data (say a 4KB block) as input and computes a function over said data, producing a small summary of the contents of the data (say 4 or 8 bytes). This summary is referred to as the checksum. The goal of such a computation is to enable a system to detect if data has somehow been corrupted or altered by storing the checksum with the data and then confirming upon later access that the data’s current checksum matches the original storage value.
There’s No Such Thing As A Free Lunch, or TNSTAAFL for short, is an old American idiom that implies that when you are seemingly getting something for free, in actuality you are likely paying some cost for it. It comes from the old days when diners would advertise a free lunch for customers, hoping to draw them in; only when you went in, did you realize that to acquire the “free” lunch, you had to purchase one or more alcoholic beverages. Of course, this may not actually be a problem, particularly if you are an aspiring alcoholic (or typical undergraduate student).

Common Checksum Functions

A number of different functions are used to compute checksums, and vary in strength (i.e., how good they are at protecting data integrity) and speed (i.e., how quickly can they be computed). A trade-off that is common in systems arises here: usually, the more protection you get, the costlier it is. There is no such thing as a free lunch.

One simple checksum function that some use is based on exclusive or (XOR). With XOR-based checksums, the checksum is computed simply by XOR’ing each chunk of the data block being checksummed, thus producing a single value that represents the XOR of the entire block.

To make this more concrete, imagine we are computing a 4-byte checksum over a block of 16 bytes (this block is of course too small to really be a disk sector or block, but it will serve for the example). The 16 data bytes, in hex, look like this:

365e c4cd ba14 8a92 ecef 2c3a 40be f666

If we view them in binary, we get the following:

0011 0110 0101 1110 1100 0100 1100 1101
1011 1010 0001 0100 1000 1010 1001 0010
1110 1100 1110 1111 0010 1100 0011 1010
0100 0000 1011 1110 1111 0110 0110 0110

Because we’ve lined up the data in groups of 4 bytes per row, it is easy to see what the resulting checksum will be: simply perform an XOR over each column to get the final checksum value:

0010 0000 0001 1011 1001 0100 0000 0011

The result, in hex, is 0x201b9403.

XOR is a reasonable checksum but has its limitations. If, for example, two bits in the same position within each checksummed unit change, the checksum will not detect the corruption. For this reason, people have investigated other checksum functions.
Another simple checksum function is addition. This approach has the advantage of being fast; computing it just requires performing 2’s-complement addition over each chunk of the data, ignoring overflow. It can detect many changes in data, but is not good if the data, for example, is shifted.

A slightly more complex algorithm is known as the Fletcher checksum, named (as you might guess) for the inventor, John G. Fletcher [F82]. It is quite simple and involves the computation of two check bytes, $s_1$ and $s_2$. Specifically, assume a block $D$ consists of bytes $d_1 \ldots d_n$; $s_1$ is simply defined as follows: $s_1 = s_1 + d_i \mod 255$ (computed over all $d_i$); $s_2$ in turn is: $s_2 = s_2 + s_1 \mod 255$ (again over all $d_i$) [F04]. The fletcher checksum is known to be almost as strong as the CRC (described next), detecting all single-bit errors, all double-bit errors, and a large percentage of burst errors [F04].

One final commonly-used checksum is known as a cyclic redundancy check (CRC). While this sounds fancy, the basic idea is quite simple. Assume you wish to compute the checksum over a data block $D$. All you do is treat $D$ as if it is a large binary number (it is just a string of bits after all) and divide it by an agreed upon value ($k$). The remainder of this division is the value of the CRC. As it turns out, one can implement this binary modulo operation rather efficiently, and hence the popularity of the CRC in networking as well. See elsewhere for more details [M13].

Whatever the method used, it should be obvious that there is no perfect checksum: it is possible two data blocks with non-identical contents will have identical checksums, something referred to as a collision. This fact should be intuitive: after all, computing a checksum is taking something large (e.g., 4KB) and producing a summary that is much smaller (e.g., 4 or 8 bytes). In choosing a good checksum function, we are thus trying to find one that minimizes the chance of collisions while remaining easy to compute.

Checksum Layout

Now that you understand a bit about how to compute a checksum, let’s next analyze how to use checksums in a storage system. The first question we must address is the layout of the checksum, i.e., how should checksums be stored on disk?

The most basic approach simply stores a checksum with each disk sector (or block). Given a data block $D$, let us call the checksum over that data $C(D)$. Thus, without checksums, the disk layout looks like this:

<table>
<thead>
<tr>
<th>D0</th>
<th>D1</th>
<th>D2</th>
<th>D3</th>
<th>D4</th>
<th>D5</th>
<th>D6</th>
</tr>
</thead>
</table>
With checksums, the layout adds a single checksum for every block:


Because checksums are usually small (e.g., 8 bytes), and disks only can write in sector-sized chunks (512 bytes) or multiples thereof, one problem that arises is how to achieve the above layout. One solution employed by drive manufacturers is to format the drive with 520-byte sectors; an extra 8 bytes per sector can be used to store the checksum.

In disks that don’t have such functionality, the file system must figure out a way to store the checksums packed into 512-byte blocks. One such possibility is as follows:

| C[D0] | C[D1] | C[D2] | C[D3] | C[D4] | D0 | D1 | D2 | D3 | D4 |

In this scheme, the n checksums are stored together in a sector, followed by n data blocks, followed by another checksum sector for the next n blocks, and so forth. This scheme has the benefit of working on all disks, but can be less efficient; if the file system, for example, wants to overwrite block D1, it has to read in the checksum sector containing C(D1), update C(D1) in it, and then write out the checksum sector as well as the new data block D1 (thus, one read and two writes). The earlier approach (of one checksum per sector) just performs a single write.

### 44.4 Using Checksums

With a checksum layout decided upon, we can now proceed to actually understand how to use the checksums. When reading a block D, the client (i.e., file system or storage controller) also reads its checksum from disk C_s(D), which we call the stored checksum (hence the subscript C_s). The client then computes the checksum over the retrieved block D, which we call the computed checksum C_c(D). At this point, the client compares the stored and computed checksums; if they are equal (i.e., C_s(D) == C_c(D), the data has likely not been corrupted, and thus can be safely returned to the user. If they do not match (i.e., C_s(D) != C_c(D)), this implies the data has changed since the time it was stored (since the stored checksum reflects the value of the data at that time). In this case, we have a corruption, which our checksum has helped us to detect.

Given a corruption, the natural question is what should we do about it? If the storage system has a redundant copy, the answer is easy: try to use it instead. If the storage system has no such copy, the likely answer is to return an error. In either case, realize that corruption detection is not a magic bullet; if there is no other way to get the non-corrupted data, you are simply out of luck.

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44.5 A New Problem: Misdirected Writes

The basic scheme described above works well in the general case of corrupted blocks. However, modern disks have a couple of unusual failure modes that require different solutions.

The first failure mode of interest is called a misdirected write. This arises in disk and RAID controllers which write the data to disk correctly, except in the wrong location. In a single-disk system, this means that the disk wrote block $D_x$ not to address $x$ (as desired) but rather to address $y$ (thus “corrupting” $D_y$); in addition, within a multi-disk system, the controller may also write $D_{i,x}$ not to address $x$ of disk $i$ but rather to some other disk $j$. Thus our question:

**Crux: How To Handle Misdirected Writes**

How should a storage system or disk controller detect misdirected writes? What additional features are required from the checksum?

The answer, not surprisingly, is simple: add a little more information to each checksum. In this case, adding a physical identifier (physical ID) is quite helpful. For example, if the stored information now contains the checksum $C(D)$ as well as the disk and sector number of the block, it is easy for the client to determine whether the correct information resides within the block. Specifically, if the client is reading block 4 on disk 10 ($D_{10,4}$), the stored information should include that disk number and sector offset, as shown below. If the information does not match, a misdirected write has taken place, and a corruption is now detected. Here is an example of what this added information would look like on a two-disk system. Note that this figure, like the others before it, is not to scale, as the checksums are usually small (e.g., 8 bytes) whereas the blocks are much larger (e.g., 4 KB or bigger):

You can see from the on-disk format that there is now a fair amount of redundancy on disk: for each block, the disk number is repeated within each block, and the offset of the block in question is also kept next to the block itself. The presence of redundant information should be no surprise, though; redundancy is the key to error detection (in this case) and recovery (in others). A little extra information, while not strictly needed with perfect disks, can go a long ways in helping detect problematic situations should they arise.
44.6 One Last Problem: Lost Writes

Unfortunately, misdirected writes are not the last problem we will address. Specifically, some modern storage devices also have an issue known as a lost write, which occurs when the device informs the upper layer that a write has completed but in fact it never is persisted; thus, what remains is left is the old contents of the block rather than the updated new contents.

The obvious question here is: do any of our checksumming strategies from above (e.g., basic checksums, or physical identity) help to detect lost writes? Unfortunately, the answer is no: the old block likely has a matching checksum, and the physical ID used above (disk number and block offset) will also be correct. Thus our final problem:

**CRUX: HOW TO HANDLE LOST WRITES**

How should a storage system or disk controller detect lost writes? What additional features are required from the checksum?

There are a number of possible solutions that can help [K+08]. One classic approach [BS04] is to perform a write verify or read-after-write; by immediately reading back the data after a write, a system can ensure that the data indeed reached the disk surface. This approach, however, is quite slow, doubling the number of I/Os needed to complete a write.

Some systems add a checksum elsewhere in the system to detect lost writes. For example, Sun’s Zettabyte File System (ZFS) includes a checksum in each file system inode and indirect block for every block included within a file. Thus, even if the write to a data block itself is lost, the checksum within the inode will not match the old data. Only if the writes to both the inode and the data are lost simultaneously will such a scheme fail, an unlikely (but unfortunately, possible!) situation.

44.7 Scrubbing

Given all of this discussion, you might be wondering: when do these checksums actually get checked? Of course, some amount of checking occurs when data is accessed by applications, but most data is rarely accessed, and thus would remain unchecked. Unchecked data is problematic for a reliable storage system, as bit rot could eventually affect all copies of a particular piece of data.

To remedy this problem, many systems utilize disk scrubbing of various forms [K+08]. By periodically reading through every block of the system, and checking whether checksums are still valid, the disk system can reduce the chances that all copies of a certain data item become corrupted. Typical systems schedule scans on a nightly or weekly basis.
44.8 Overheads Of Checksumming

Before closing, we now discuss some of the overheads of using checksums for data protection. There are two distinct kinds of overheads, as is common in computer systems: space and time.

Space overheads come in two forms. The first is on the disk (or other storage medium) itself; each stored checksum takes up room on the disk, which can no longer be used for user data. A typical ratio might be an 8-byte checksum per 4 KB data block, for a 0.19% on-disk space overhead.

The second type of space overhead comes in the memory of the system. When accessing data, there must now be room in memory for the checksums as well as the data itself. However, if the system simply checks the checksum and then discards it once done, this overhead is short-lived and not much of a concern. Only if checksums are kept in memory (for an added level of protection against memory corruption [Z+13]) will this small overhead be observable.

While space overheads are small, the time overheads induced by checksumming can be quite noticeable. Minimally, the CPU must compute the checksum over each block, both when the data is stored (to determine the value of the stored checksum) as well as when it is accessed (to compute the checksum again and compare it against the stored checksum). One approach to reducing CPU overheads, employed by many systems that use checksums (including network stacks), is to combine data copying and checksumming into one streamlined activity; because the copy is needed anyhow (e.g., to copy the data from the kernel page cache into a user buffer), combined copying/checksumming can be quite effective.

Beyond CPU overheads, some checksumming schemes can induce extra I/O overheads, particularly when checksums are stored distinctly from the data (thus requiring extra I/Os to access them), and for any extra I/O needed for background scrubbing. The former can be reduced by design; the latter can be tuned and thus its impact limited, perhaps by controlling when such scrubbing activity takes place. The middle of the night, when most (not all!) productive workers have gone to bed, may be a good time to perform such scrubbing activity and increase the robustness of the storage system.

44.9 Summary

We have discussed data protection in modern storage systems, focusing on checksum implementation and usage. Different checksums protect against different types of faults; as storage devices evolve, new failure modes will undoubtedly arise. Perhaps such change will force the research community and industry to revisit some of these basic approaches, or invent entirely new approaches altogether. Time will tell. Or it won’t. Time is funny that way.
References

[B+08] “An Analysis of Data Corruption in the Storage Stack”
Lakshmi N. Bairavasundaram, Garth R. Goodson, Bianca Schroeder,
Andrea C. Arpaci-Dusseau, Remzi H. Arpaci-Dusseau
FAST ’08, San Jose, CA, February 2008
The first paper to truly study disk corruption in great detail, focusing on how often such corruption
occurs over three years for over 1.5 million drives. Lakshmi did this work while a graduate student at
Wisconsin under our supervision, but also in collaboration with his colleagues at NetApp where he was
an intern for multiple summers. A great example of how working with industry can make for much
more interesting and relevant research.

Wendy Bartlett, Lisa Spainhower
IEEE Transactions on Dependable and Secure Computing, Vol. 1, No. 1, January 2004
This classic in building fault tolerant systems is an excellent overview of the state of the art from both
IBM and Tandem. Another must read for those interested in the area.

[C+04] “Row-Diagonal Parity for Double Disk Failure Correction”
P. Corbett, B. English, A. Goel, T. Grcanac, S. Kleiman, J. Leong, S. Sankar
FAST ’04, San Jose, CA, February 2004
An early paper on how extra redundancy helps to solve the combined full-disk-failure/partial-disk-failure
problem. Also a nice example of how to mix more theoretical work with practical.

[F04] “Checksums and Error Control”
Peter M. Fenwick
A great simple tutorial on checksums, available to you for the amazing cost of free.

[F82] “An Arithmetic Checksum for Serial Transmissions”
John G. Fletcher
Fletcher’s original work on his eponymous checksum. Of course, he didn’t call it the Fletcher checksum,
rather he just didn’t call it anything, and thus it became natural to name it after the inventor. So don’t
blame old Fletch for this seeming act of braggadocio.

Dave Hitz, James Lau, Michael Malcolm
USENIX Spring ’94
The pioneering paper that describes the ideas and product at the heart of NetApp’s core. Based on this
system, NetApp has grown into a multi-billion dollar storage company. If you’re interested in learning
more about its founding, read Hitz’s autobiography “How to Castrate a Bull: Unexpected Lessons on
Risk, Growth, and Success in Business” (which is the actual title, no joking). And you thought you
could avoid bull castration by going into Computer Science.

[K+08] “Parity Lost and Parity Regained”
Andrew Krioukov, Lakshmi N. Bairavasundaram, Garth R. Goodson, Kiran Srinivasan,
Randy Thelen, Andrea C. Arpaci-Dusseau, Remzi H. Arpaci-Dusseau
FAST ’08, San Jose, CA, February 2008
This work of ours, joint with colleagues at NetApp, explores how different checksum schemes work (or
don’t work) in protecting data. We reveal a number of interesting flaws in current protection strategies,
some of which have led to fixes in commercial products.
“Cyclic Redundancy Checks”
Author Unknown
Not sure who wrote this, but a super clear and concise description of CRCs is available here. The internet is full of information, as it turns out.

“IRON File Systems”
Vijayan Prabhakaran, Lakshmi N. Bairavasundaram, Nitin Agrawal, Haryadi S. Gunawi, Andrea C. Arpaci-Dusseau, Remzi H. Arpaci-Dusseau
SOSP ’05, Brighton, England, October 2005
Our paper on how disks have partial failure modes, which includes a detailed study of how file systems such as Linux ext3 and Windows NTFS react to such failures. As it turns out, rather poorly! We found numerous bugs, design flaws, and other oddities in this work. Some of this has fed back into the Linux community, thus helping to yield a new more robust group of file systems to store your data.

“Design and Implementation of the Log-structured File System”
Mendel Rosenblum and John Ousterhout
SOSP ’91, Pacific Grove, CA, October 1991
Another reference to this ground-breaking paper on how to improve write performance in file systems.

“Implementing Fault-Tolerant Services Using The State Machine Approach: A Tutorial”
Fred B. Schneider
ACM Surveys, Vol. 22, No. 4, December 1990
This classic paper talks generally about how to build fault tolerant services, and includes many basic definitions of terms. A must read for those building distributed systems.

“Zettabyte Reliability with Flexible End-to-end Data Integrity”
Yupu Zhang, Daniel S. Myers, Andrea C. Arpaci-Dusseau, Remzi H. Arpaci-Dusseau
MSST ’13, Long Beach, California, May 2013
Our own work on adding data protection to the page cache of a system, which protects against memory corruption as well as on-disk corruption.
Summary Dialogue on Persistence

**Student:** Wow, file systems seem interesting(!), and yet complicated.

**Professor:** That’s why me and my spouse do our research in this space.

**Student:** Hold on. Are you one of the professors who wrote this book? I thought we were both just fake constructs, used to summarize some main points, and perhaps add a little levity in the study of operating systems.

**Professor:** Uh... er... maybe. And none of your business! And who did you think was writing these things? (sighs) Anyhow, let’s get on with it: what did you learn?

**Student:** Well, I think I got one of the main points, which is that it is much harder to manage data for a long time (persistently) than it is to manage data that isn’t persistent (like the stuff in memory). After all, if your machines crashes, memory contents disappear! But the stuff in the file system needs to live forever.

**Professor:** Well, as my friend Kevin Hultquist used to say, “Forever is a long time”; while he was talking about plastic golf tees, it’s especially true for the garbage that is found in most file systems.

**Student:** Well, you know what I mean! For a long time at least. And even simple things, such as updating a persistent storage device, are complicated, because you have to care what happens if you crash. Recovery, something I had never even thought of when we were virtualizing memory, is now a big deal!

**Professor:** Too true. Updates to persistent storage have always been, and remain, a fun and challenging problem.

**Student:** I also learned about cool things like disk scheduling, and about data protection techniques like RAID and even checksums. That stuff is cool.

**Professor:** I like those topics too. Though, if you really get into it, they can get a little mathematical. Check out some the latest on erasure codes if you want your brain to hurt.

**Student:** I’ll get right on that.
Professor: (frowns) I think you’re being sarcastic. Well, what else did you like?

Student: And I also liked all the thought that has gone into building technology-aware systems, like FFS and LFS. Neat stuff! Being disk aware seems cool. But will it matter anymore, with Flash and all the newest, latest technologies?

Professor: Good question! And a reminder to get working on that Flash chapter... (scribbles note down to self) ... But yes, even with Flash, all of this stuff is still relevant, amazingly. For example, Flash Translation Layers (FTLs) use log-structuring internally, to improve performance and reliability of Flash-based SSDs. And thinking about locality is always useful. So while the technology may be changing, many of the ideas we have studied will continue to be useful, for a while at least.

Student: That’s good. I just spent all this time learning it, and I didn’t want it to all be for no reason!

Professor: Professors wouldn’t do that to you, would they?
A Dialogue on Distribution

Professor: And thus we reach our final little piece in the world of operating systems: distributed systems. Since we can’t cover much here, we’ll sneak in a little intro here in the section on persistence, and focus mostly on distributed file systems. Hope that is OK!

Student: Sounds OK. But what is a distributed system exactly, oh glorious and all-knowing professor?

Professor: Well, I bet you know how this is going to go...

Student: There’s a peach?

Professor: Exactly! But this time, it’s far away from you, and may take some time to get the peach. And there are a lot of them! Even worse, sometimes a peach becomes rotten. But you want to make sure that when anybody bites into a peach, they will get a mouthful of deliciousness.

Student: This peach analogy is working less and less for me.

Professor: Come on! It’s the last one, just go with it.

Student: Fine.

Professor: So anyhow, forget about the peaches. Building distributed systems is hard, because things fail all the time. Messages get lost, machines go down, disks corrupt data. It’s like the whole world is working against you!

Student: But I use distributed systems all the time, right?

Professor: Yes! You do. And... ?

Student: Well, it seems like they mostly work. After all, when I send a search request to google, it usually comes back in a snap, with some great results! Same thing when I use facebook, or Amazon, and so forth.
**Professor:** Yes, it is amazing. And that’s despite all of those failures taking place! Those companies build a huge amount of machinery into their systems so as to ensure that even though some machines have failed, the entire system stays up and running. They use a lot of techniques to do this: replication, retry, and various other tricks people have developed over time to detect and recover from failures.

**Student:** Sounds interesting. Time to learn something for real?

**Professor:** It does seem so. Let’s get to work! But first things first ...

(bites into peach he has been holding, which unfortunately is rotten)
Distributed systems have changed the face of the world. When your web browser connects to a web server somewhere else on the planet, it is participating in what seems to be a simple form of a **client/server** distributed system. When you contact a modern web service such as Google or Facebook, you are not just interacting with a single machine, however; behind the scenes, these complex services are built from a large collection (i.e., thousands) of machines, each of which cooperate to provide the particular service of the site. Thus, it should be clear what makes studying distributed systems interesting. Indeed, it is worthy of an entire class; here, we just introduce a few of the major topics.

A number of new challenges arise when building a distributed system. The major one we focus on is **failure**; machines, disks, networks, and software all fail from time to time, as we do not (and likely, will never) know how to build “perfect” components and systems. However, when we build a modern web service, we’d like it to appear to clients as if it never fails; how can we accomplish this task?

**THE CRUX:**

**HOW TO BUILD SYSTEMS THAT WORK WHEN COMPONENTS FAIL**

How can we build a working system out of parts that don’t work correctly all the time? The basic question should remind you of some of the topics we discussed in RAID storage arrays; however, the problems here tend to be more complex, as are the solutions.

Interestingly, while failure is a central challenge in constructing distributed systems, it also represents an opportunity. Yes, machines fail; but the mere fact that a machine fails does not imply the entire system must fail. By collecting together a set of machines, we can build a system that appears to rarely fail, despite the fact that its components fail regularly. This reality is the central beauty and value of distributed systems, and why they underly virtually every modern web service you use, including Google, Facebook, etc.
In virtually all circumstances, it is good to view communication as a fundamentally unreliable activity. Bit corruption, down or non-working links and machines, and lack of buffer space for incoming packets all lead to the same result: packets sometimes do not reach their destination. To build reliable services atop such unreliable networks, we must consider techniques that can cope with packet loss.

Other important issues exist as well. System performance is often critical; with a network connecting our distributed system together, system designers must often think carefully about how to accomplish their given tasks, trying to reduce the number of messages sent and further make communication as efficient (low latency, high bandwidth) as possible.

Finally, security is also a necessary consideration. When connecting to a remote site, having some assurance that the remote party is who they say they are becomes a central problem. Further, ensuring that third parties cannot monitor or alter an on-going communication between two others is also a challenge.

In this introduction, we’ll cover the most basic new aspect that is new in a distributed system: communication. Namely, how should machines within a distributed system communicate with one another? We’ll start with the most basic primitives available, messages, and build a few higher-level primitives on top of them. As we said above, failure will be a central focus: how should communication layers handle failures?

47.1 Communication Basics

The central tenet of modern networking is that communication is fundamentally unreliable. Whether in the wide-area Internet, or a local-area high-speed network such as Infiniband, packets are regularly lost, corrupted, or otherwise do not reach their destination.

There are a multitude of causes for packet loss or corruption. Sometimes, during transmission, some bits get flipped due to electrical or other similar problems. Sometimes, an element in the system, such as a network link or packet router or even the remote host, are somehow damaged or otherwise not working correctly; network cables do accidentally get severed, at least sometimes.

More fundamental however is packet loss due to lack of buffering within a network switch, router, or endpoint. Specifically, even if we could guarantee that all links worked correctly, and that all the components in the system (switches, routers, end hosts) were up and running as expected, loss is still possible, for the following reason. Imagine a packet arrives at a router; for the packet to be processed, it must be placed in memory somewhere within the router. If many such packets arrive at
// client code
int main(int argc, char *argv[]) {
    int sd = UDP_Open(20000);
    struct sockaddr_in addr, addr2;
    int rc = UDP_FillSockAddr(&addr, "machine.cs.wisc.edu", 10000);
    char message[BUFFER_SIZE];
    sprintf(message, "hello world");
    rc = UDP_Write(sd, &addr, message, BUFFER_SIZE);
    if (rc > 0) {
        int rc = UDP_Read(sd, &addr2, buffer, BUFFER_SIZE);
    }
    return 0;
}

// server code
int main(int argc, char *argv[]) {
    int sd = UDP_Open(10000);
    assert(sd > -1);
    while (1) {
        struct sockaddr_in s;
        char buffer[BUFFER_SIZE];
        int rc = UDP_Read(sd, &s, buffer, BUFFER_SIZE);
        if (rc > 0) {
            char reply[BUFFER_SIZE];
            sprintf(reply, "reply");
            rc = UDP_Write(sd, &s, reply, BUFFER_SIZE);
        }
    }
    return 0;
}

Figure 47.1: Example UDP/IP Client/Server Code

once, it is possible that the memory within the router cannot accommodate all of the packets. The only choice the router has at that point is to drop one or more of the packets. This same behavior occurs at end hosts as well; when you send a large number of messages to a single machine, the machine’s resources can easily become overwhelmed, and thus packet loss again arises.

Thus, packet loss is fundamental in networking. The question thus becomes: how should we deal with it?

47.2 Unreliable Communication Layers

One simple way is this: we don’t deal with it. Because some applications know how to deal with packet loss, it is sometimes useful to let them communicate with a basic unreliable messaging layer, an example of the end-to-end argument one often hears about (see the Aside at end of chapter). One excellent example of such an unreliable layer is found in the UDP/IP networking stack available today on virtually all modern systems. To use UDP, a process uses the sockets API in order to create a communication endpoint; processes on other machines (or on the same machine) send UDP datagrams to the original process (a datagram is a fixed-sized message up to some max size).
int UDP_Open(int port) {
    int sd;
    if ((sd = socket(AF_INET, SOCK_DGRAM, 0)) == -1) { return -1; }
    struct sockaddr_in myaddr;
    bzero(&myaddr, sizeof(myaddr));
    myaddr.sin_family = AF_INET;
    myaddr.sin_port = htons(port);
    myaddr.sin_addr.s_addr = INADDR_ANY;
    if (bind(sd, (struct sockaddr*) &myaddr, sizeof(myaddr)) == -1) {
        close(sd);
        return -1;
    }
    return sd;
}

int UDP_FillSockAddr(struct sockaddr_in*addr, char *hostName, int port) {
    bzero(addr, sizeof(struct sockaddr_in));
    addr->sin_family = AF_INET; // host byte order
    addr->sin_port = htons(port); // short, network byte order
    struct in_addr*inAddr;
    struct hostent*hostEntry;
    if ((hostEntry = gethostbyname(hostName)) == NULL) { return -1; }
    inAddr = (struct in_addr *) hostEntry->h_addr;
    addr->sin_addr = *inAddr;
    return 0;
}

int UDP_Write(int sd, struct sockaddr_in*addr, char *buffer, int n) {
    int addrLen = sizeof(struct sockaddr_in);
    return sendto(sd, buffer, n, 0, (struct sockaddr*) addr, addrLen);
}

int UDP_Read(int sd, struct sockaddr_in*addr, char *buffer, int n) {
    int len = sizeof(struct sockaddr_in);
    return recvfrom(sd, buffer, n, 0, (struct sockaddr*) addr,
                     (socklen_t *) &len);
    return rc;
}

Figure 47.2: A Simple UDP Library

Figures 47.1 and 47.2 show a simple client and server built on top of UDP/IP. The client can send a message to the server, which then responds with a reply. With this small amount of code, you have all you need to begin building distributed systems!

UDP is a great example of an unreliable communication layer. If you use it, you will encounter situations where packets get lost (dropped) and thus do not reach their destination; the sender is never thus informed of the loss. However, that does not mean that UDP does not guard against any failures at all. For example, UDP includes a checksum to detect some forms of packet corruption.

However, because many applications simply want to send data to a destination and not worry about packet loss, we need more. Specifically, we need reliable communication on top of an unreliable network.
Checksums are a commonly-used method to detect corruption quickly and effectively in modern systems. A simple checksum is addition: just sum up the bytes of a chunk of data; of course, many other more sophisticated checksums have been created, including basic cyclic redundancy codes (CRCs), the Fletcher checksum, and many others [MK09].

In networking, checksums are used as follows. Before sending a message from one machine to another, compute a checksum over the bytes of the message. Then send both the message and the checksum to the destination. At the destination, the receiver computes a checksum over the incoming message as well; if this computed checksum matches the sent checksum, the receiver can feel some assurance that the data likely did not get corrupted during transmission.

Checksums can be evaluated along a number of different axes. Effectiveness is one primary consideration: does a change in the data lead to a change in the checksum? The stronger the checksum, the harder it is for changes in the data to go unnoticed. Performance is the other important criterion: how costly is the checksum to compute? Unfortunately, effectiveness and performance are often at odds, meaning that checksums of high quality are often expensive to compute. Life, again, isn’t perfect.

47.3 Reliable Communication Layers

To build a reliable communication layer, we need some new mechanisms and techniques to handle packet loss. Let us consider a simple example in which a client is sending a message to a server over an unreliable connection. The first question we must answer: how does the sender know that the receiver has actually received the message?

The technique that we will use is known as an acknowledgment, or ack for short. The idea is simple: the sender sends a message to the receiver; the receiver then sends a short message back to acknowledge its receipt. Figure 47.3 depicts the process.

```
Sender
[send message]
[receive ack]

Receiver
[receive message]
[send ack]
```

**Figure 47.3: Message Plus Acknowledgment**

When the sender receives an acknowledgment of the message, it can then rest assured that the message did indeed receive the original message. However, what should the sender do if it does not receive an acknowledgment?
To handle this case, we need an additional mechanism, known as a **timeout**. When the sender sends a message, the sender now sets a timer to go off after some period of time. If, in that time, no acknowledgment has been received, the sender concludes that the message has been lost. The sender then simply performs a **retry** of the send, sending the same message again with hopes that this time, it will get through. For this approach to work, the sender must keep a copy of the message around, in case it needs to send it again. The combination of the timeout and the retry have led some to call the approach **timeout/retry**; pretty clever crowd, those networking types, no? Figure 47.4 shows an example.

Unfortunately, timeout/retry in this form is not quite enough. Figure 47.5 shows an example of packet loss which could lead to trouble. In this example, it is not the original message that gets lost, but the acknowledgment. From the perspective of the sender, the situation seems the same: no ack was received, and thus a timeout and retry are in order. But from the perspective of the receiver, it is quite different: now the same message has been received twice! While there may be cases where this is OK, in general it is not; imagine what would happen when you are downloading a file and extra packets are repeated inside the download. Thus, when we are aiming for a reliable message layer, we also usually want to guarantee that each message is received **exactly once** by the receiver.

To enable the receiver to detect duplicate message transmission, the sender has to identify each message in some unique way, and the receiver needs some way to track whether it has already seen each message before. When the receiver sees a duplicate transmission, it simply acks the message, but (critically) does **not** pass the message to the application that receives the data. Thus, the sender receives the ack but the message is not received twice, preserving the exactly-once semantics mentioned above.

There are myriad ways to detect duplicate messages. For example, the sender could generate a unique ID for each message; the receiver could track every ID it has ever seen. This approach could work, but it is prohibitively costly, requiring unbounded memory to track all IDs.
A simpler approach, requiring little memory, solves this problem, and the mechanism is known as a sequence counter. With a sequence counter, the sender and receiver agree upon a start value (e.g., 1) for a counter that each side will maintain. Whenever a message is sent, the current value of the counter is sent along with the message; this counter value ($N$) serves as an ID for the message. After the message is sent, the sender then increments the value (to $N+1$).

The receiver uses its counter value as the expected value for the ID of the incoming message from that sender. If the ID of a received message ($N$) matches the receiver’s counter (also $N$), it acks the message and passes it up to the application; in this case, the receiver concludes this is the first time this message has been received. The receiver then increments its counter (to $N+1$), and waits for the next message.

If the ack is lost, the sender will timeout and re-send message $N$. This time, the receiver’s counter is higher ($N+1$), and thus the receiver knows it has already received this message. Thus it acks the message but does not pass it up to the application. In this simple manner, sequence counters can be used to avoid duplicates.

The most commonly used reliable communication layer is known as TCP/IP, or just TCP for short. TCP has a great deal more sophistication than we describe above, including machinery to handle congestion in the network [VJ90], multiple outstanding requests, and hundreds of other small tweaks and optimizations. Read more about it if you’re curious; better yet, take a networking course and learn that material well.

47.4 Communication Abstractions

Given a basic messaging layer, we now approach the next question in this chapter: what abstraction of communication should we use when building a distributed system?
As you can probably guess from the discussion, setting the timeout value correctly is an important aspect of using timeouts to retry message sends. If the timeout is too small, the sender will re-send messages needlessly, thus wasting CPU time on the sender and network resources. If the timeout is too large, the sender waits too long to re-send and thus perceived performance at the sender is reduced. The “right” value, from the perspective of a single client and server, is thus to wait just long enough to detect packet loss but no longer.

However, there are often more than just a single client and server in a distributed system, as we will see in future chapters. In a scenario with many clients sending to a single server, packet loss at the server may be an indicator that the server is overloaded. If true, clients might retry in a different adaptive manner; for example, after the first timeout, a client might increase its timeout value to a higher amount, perhaps twice as high as the original value. Such an exponential back-off scheme, pioneered in the early Aloha network and adopted in early Ethernet [A70], avoid situations where resources are being overloaded by an excess of re-sends. Robust systems strive to avoid overload of this nature.

The systems community developed a number of approaches over the years. One body of work took OS abstractions and extended them to operate in a distributed environment. For example, distributed shared memory (DSM) systems enable processes on different machines to share a large, virtual address space [LH89]. This abstraction turns a distributed computation into something that looks like a multi-threaded application; the only difference is that these threads run on different machines instead of different processors within the same machine.

The way most DSM systems work is through the virtual memory system of the OS. When a page is accessed on one machine, two things can happen. In the first (best) case, the page is already local on the machine, and thus the data is fetched quickly. In the second case, the page is currently on some other machine. A page fault occurs, and the page fault handler sends a message to some other machine to fetch the page, install it in the page table of the requesting process, and continue execution.

This approach is not widely in use today for a number of reasons. The largest problem for DSM is how it handles failure. Imagine, for example, if a machine fails; what happens to the pages on that machine? What if the data structures of the distributed computation are spread across the entire address space? In this case, parts of these data structures would suddenly become unavailable. Dealing with failure when parts of your address space go missing is hard; imagine a linked list that where a next pointer points into a portion of the address space that is gone. Yikes!

A further problem is performance. One usually assumes, when writing code, that access to memory is cheap. In DSM systems, some accesses
are inexpensive, but others cause page faults and expensive fetches from remote machines. Thus, programmers of such DSM systems had to be very careful to organize computations such that almost no communication occurred at all, defeating much of the point of such an approach. Though much research was performed in this space, there was little practical impact; nobody builds reliable distributed systems using DSM today.

47.5 Remote Procedure Call (RPC)

While OS abstractions turned out to be a poor choice for building distributed systems, programming language (PL) abstractions make much more sense. The most dominant abstraction is based on the idea of a remote procedure call, or RPC for short [BN84].

Remote procedure call packages all have a simple goal: to make the process of executing code on a remote machine as simple and straightforward as calling a local function. Thus, to a client, a procedure call is made, and some time later, the results are returned. The server simply defines some routines that it wishes to export. The rest of the magic is handled by the RPC system, which in general has two pieces: a stub generator (sometimes called a protocol compiler), and the run-time library. We’ll now take a look at each of these pieces in more detail.

Stub Generator

The stub generator’s job is simple: to remove some of the pain of packing function arguments and results into messages by automating it. Numerous benefits arise: one avoids, by design, the simple mistakes that occur in writing such code by hand; further, a stub compiler can perhaps optimize such code and thus improve performance.

The input to such a compiler is simply the set of calls a server wishes to export to clients. Conceptually, it could be something as simple as this:

```plaintext
interface {
    int func1(int arg1);
    int func2(int arg1, int arg2);
};
```

The stub generator takes an interface like this and generates a few different pieces of code. For the client, a client stub is generated, which contains each of the functions specified in the interface; a client program wishing to use this RPC service would link with this client stub and call into it in order to make RPCs.

Internally, each of these functions in the client stub do all of the work needed to perform the remote procedure call. To the client, the code just

---

1In modern programming languages, we might instead say remote method invocation (RMI), but who likes these languages anyhow, with all of their fancy objects?
appears as a function call (e.g., the client calls \texttt{func1(x)}); internally, the code in the client stub for \texttt{func1()} does this:

- **Create a message buffer.** A message buffer is usually just a contiguous array of bytes of some size.
- **Pack the needed information into the message buffer.** This information includes some kind of identifier for the function to be called, as well as all of the arguments that the function needs (e.g., in our example above, one integer for \texttt{func1}). The process of putting all of this information into a single contiguous buffer is sometimes referred to as the *marshaling* of arguments or the *serialization* of the message.
- **Send the message to the destination RPC server.** The communication with the RPC server, and all of the details required to make it operate correctly, are handled by the RPC run-time library, described further below.
- **Wait for the reply.** Because function calls are usually *synchronous*, the call will wait for its completion.
- **Unpack return code and other arguments.** If the function just returns a single return code, this process is straightforward; however, more complex functions might return more complex results (e.g., a list), and thus the stub might need to unpack those as well. This step is also known as *unmarshaling* or *deserialization*.
- **Return to the caller.** Finally, just return from the client stub back into the client code.

For the server, code is also generated. The steps taken on the server are as follows:

- **Unpack the message.** This step, called *unmarshaling* or *deserialization*, takes the information out of the incoming message. The function identifier and arguments are extracted.
- **Call into the actual function.** Finally! We have reached the point where the remote function is actually executed. The RPC runtime calls into the function specified by the ID and passes in the desired arguments.
- **Package the results.** The return argument(s) are marshaled back into a single reply buffer.
- **Send the reply.** The reply is finally sent to the caller.

There are a few other important issues to consider in a stub compiler. The first is complex arguments, i.e., how does one package and send a complex data structure? For example, when one calls the \texttt{write()} system call, one passes in three arguments: an integer file descriptor, a pointer to a buffer, and a size indicating how many bytes (starting at the pointer) are to be written. If an RPC package is passed a pointer, it needs to be able to figure out how to interpret that pointer, and perform the
correct action. Usually this is accomplished through either well-known types (e.g., a buffer_t that is used to pass chunks of data given a size, which the RPC compiler understands), or by annotating the data structures with more information, enabling the compiler to know which bytes need to be serialized.

Another important issue is the organization of the server with regards to concurrency. A simple server just waits for requests in a simple loop, and handles each request one at a time. However, as you might have guessed, this can be grossly inefficient; if one RPC call blocks (e.g., on I/O), server resources are wasted. Thus, most servers are constructed in some sort of concurrent fashion. A common organization is a thread pool. In this organization, a finite set of threads are created when the server starts; when a message arrives, it is dispatched to one of these worker threads, which then does the work of the RPC call, eventually replying; during this time, a main thread keeps receiving other requests, and perhaps dispatching them to other workers. Such an organization enables concurrent execution within the server, thus increasing its utilization; the standard costs arise as well, mostly in programming complexity, as the RPC calls may now need to use locks and other synchronization primitives in order to ensure their correct operation.

Run-Time Library

The run-time library handles much of the heavy lifting in an RPC system; most performance and reliability issues are handled herein. We'll now discuss some of the major challenges in building such a run-time layer.

One of the first challenges we must overcome is how to locate a remote service. This problem, of naming, is a common one in distributed systems, and in some sense goes beyond the scope of our current discussion. The simplest of approaches build on existing naming systems, e.g., hostnames and port numbers provided by current internet protocols. In such a system, the client must know the hostname or IP address of the machine running the desired RPC service, as well as the port number it is using (a port number is just a way of identifying a particular communication activity taking place on a machine, allowing multiple communication channels at once). The protocol suite must then provide a mechanism to route packets to a particular address from any other machine in the system. For a good discussion of naming, read either the Grapevine paper or about DNS and name resolution on the Internet, or better yet just read the excellent chapter in Saltzer and Kaashoek's book [SK09].

Once a client knows which server it should talk to for a particular remote service, the next question is which transport-level protocol should RPC be built upon. Specifically, should the RPC system use a reliable protocol such as TCP/IP, or be built upon an unreliable communication layer such as UDP/IP?

Naively the choice would seem easy: clearly we would like for a request to be reliably delivered to the remote server, and clearly we would
like to reliably receive a reply. Thus we should choose the reliable transport protocol such as TCP, right?

Unfortunately, building RPC on top of a reliable communication layer can lead to a major inefficiency in performance. Recall from the discussion above how reliable communication layers work: with acknowledgments plus timeout/retry. Thus, when the client sends an RPC request to the server, the server responds with an acknowledgment so that the caller knows the request was received. Similarly, when the server sends the reply to the client, the client acks it so that the server knows it was received. By building a request/response protocol (such as RPC) on top of a reliable communication layer, two “extra” messages are sent.

For this reason, many RPC packages are built on top of unreliable communication layers, such as UDP. Doing so enables a more efficient RPC layer, but does add the responsibility of providing reliability to the RPC system. The RPC layer achieves the desired level of responsibility by using timeout/retry and acknowledgments much like we described above. By using some form of sequence numbering, the communication layer can guarantee that each RPC takes place exactly once (in the case of no failure), or at most once (in the case where failure arises).

Other Issues

There are some other issues an RPC run-time must handle as well. For example, what happens when a remote call takes a long time to complete? Given our timeout machinery, a long-running remote call might appear as a failure to a client, thus triggering a retry, and thus the need for some care here. One solution is to use an explicit acknowledgment (from the receiver to sender) when the reply isn’t immediately generated; this lets the client know the server received the request. Then, after some time has passed, the client can periodically ask whether the server is still working on the request; if the server keeps saying “yes”, the client should be happy and continue to wait (after all, sometimes a procedure call can take a long time to finish executing).

The run-time must also handle procedure calls with large arguments, larger than what can fit into a single packet. Some lower-level network protocols provide such sender-side fragmentation (of larger packets into a set of smaller ones) and receiver-side reassembly (of smaller parts into one larger logical whole); if not, the RPC run-time may have to implement such functionality itself. See Birrell and Nelson’s excellent RPC paper for details [BN84].

One issue that many systems handle is that of byte ordering. As you may know, some machines store values in what is known as big endian ordering, whereas others use little endian ordering. Big endian stores bytes (say, of an integer) from most significant to least significant bits, much like Arabic numerals; little endian does the opposite. Both are equally valid ways of storing numeric information; the question here is how to communicate between machines of different endianness.
Aside: **The End-to-End Argument**

The end-to-end argument makes the case that the highest level in a system, i.e., usually the application at “the end”, is ultimately the only locale within a layered system where certain functionality can truly be implemented. In their landmark paper, Saltzer et al. argue this through an excellent example: reliable file transfer between two machines. If you want to transfer a file from machine $A$ to machine $B$, and make sure that the bytes that end up on $B$ are exactly the same as those that began on $A$, you must have an “end-to-end” check of this; lower-level reliable machinery, e.g., in the network or disk, provides no such guarantee.

The contrast is an approach which tries to solve the reliable-file-transfer problem by adding reliability to lower layers of the system. For example, say we build a reliable communication protocol and use it to build our reliable file transfer. The communication protocol guarantees that every byte sent by a sender will be received in order by the receiver, say using timeout/retry, acknowledgments, and sequence numbers. Unfortunately, using such a protocol does not a reliable file transfer make; imagine the bytes getting corrupted in sender memory before the communication even takes place, or something bad happening when the receiver writes the data to disk. In those cases, even though the bytes were delivered reliably across the network, our file transfer was ultimately not reliable. To build a reliable file transfer, one must include end-to-end checks of reliability, e.g., after the entire transfer is complete, read back the file on the receiver disk, compute a checksum, and compare that checksum to that of the file on the sender.

The corollary to this maxim is that sometimes having lower layers provide extra functionality can indeed improve system performance or otherwise optimize a system. Thus, you should not rule out having such machinery at a lower-level in a system; rather, you should carefully consider the utility of such machinery, given its eventual usage in an overall system or application.

RPC packages often handle this by providing a well-defined endianness within their message formats. In Sun’s RPC package, the XDR (eXternal Data Representation) layer provides this functionality. If the machine sending or receiving a message matches the endianness of XDR, messages are just sent and received as expected. If, however, the machine communicating has a different endianness, each piece of information in the message must be converted. Thus, the difference in endianness can have a small performance cost.

A final issue is whether to expose the asynchronous nature of communication to clients, thus enabling some performance optimizations. Specifically, typical RPCs are made synchronously, i.e., when a client issues the procedure call, it must wait for the procedure call to return before continuing. Because this wait can be long, and because the client
may have other work it could be doing, some RPC packages enable you to invoke an RPC asynchronously. When an asynchronous RPC is issued, the RPC package sends the request and returns immediately; the client is then free to do other work, such as call other RPCs or other useful computation. The client at some point will want to see the results of the asynchronous RPC; it thus calls back into the RPC layer, telling it to wait for outstanding RPCs to complete, at which point return arguments can be accessed.

47.6 Summary

We have seen the introduction of a new topic, distributed systems, and its major issue: how to handle failure which is now a commonplace event. As they say inside of Google, when you have just your desktop machine, failure is rare; when you’re in a data center with thousands of machines, failure is happening all the time. The key to any distributed system is how you deal with that failure.

We have also seen that communication forms the heart of any distributed system. A common abstraction of that communication is found in remote procedure call (RPC), which enables clients to make remote calls on servers; the RPC package handles all of the gory details, including timeout/retry and acknowledgment, in order to deliver a service that closely mirrors a local procedure call.

The best way to really understand an RPC package is of course to use one yourself. Sun’s RPC system, using the stub compiler rpcgen, is a common one, and is widely available on systems today, including Linux. Try it out, and see what all the fuss is about.
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Sun’s Network File System (NFS)

One of the first uses of distributed client/server computing was in the realm of distributed file systems. In such an environment, there are a number of client machines and one server (or a few); the server stores the data on its disks, and clients request data through well-formed protocol messages. Figure 48.1 depicts the basic setup.

As you can see from the picture, the server has the disks, and clients send messages network to access their directories and files on those disks. Why do we bother with this arrangement? (i.e., why don’t we just let clients use their local disks?) Well, primarily this setup allows for easy sharing of data across clients. Thus, if you access a file on one machine (Client 0) and then later use another (Client 2), you will have the same view of the file system. Your data is naturally shared across these different machines. A secondary benefit is centralized administration; for example, backing up files can be done from the few server machines instead of from the multitude of clients. Another advantage could be security; having all servers in a locked machine room prevents certain types of problems from arising.
48.1 A Basic Distributed File System

We now will study the architecture of a simplified distributed file system. A simple client/server distributed file system has more components than the file systems we have studied so far. On the client side, there are client applications which access files and directories through the client-side file system. A client application issues system calls to the client-side file system (such as `open()`, `read()`, `write()`, `close()`, `mkdir()`, etc.) in order to access files which are stored on the server. Thus, to client applications, the file system does not appear to be any different than a local (disk-based) file system, except perhaps for performance; in this way, distributed file systems provide transparent access to files, an obvious goal; after all, who would want to use a file system that required a different set of APIs or otherwise was a pain to use?

The role of the client-side file system is to execute the actions needed to service those system calls. For example, if the client issues a `read()` request, the client-side file system may send a message to the server-side file system (or, more commonly, the file server) to read a particular block; the file server will then read the block from disk (or its own in-memory cache), and send a message back to the client with the requested data. The client-side file system will then copy the data into the user buffer supplied to the `read()` system call and thus the request will complete. Note that a subsequent `read()` of the same block on the client may be cached in client memory or on the client’s disk even; in the best such case, no network traffic need be generated.

![Distributed File System Architecture](image)

**Figure 48.2: Distributed File System Architecture**

From this simple overview, you should get a sense that there are two important pieces of software in a client/server distributed file system: the client-side file system and the file server. Together their behavior determines the behavior of the distributed file system. Now it’s time to study one particular system: Sun’s Network File System (NFS).
48.2 On To NFS

One of the earliest and quite successful distributed systems was developed by Sun Microsystems, and is known as the Sun Network File System (or NFS) [S86]. In defining NFS, Sun took an unusual approach: instead of building a proprietary and closed system, Sun instead developed an open protocol which simply specified the exact message formats that clients and servers would use to communicate. Different groups could develop their own NFS servers and thus compete in an NFS marketplace while preserving interoperability. It worked: today there are many companies that sell NFS servers (including Oracle/Sun, NetApp [HLM94], EMC, IBM, and others), and the widespread success of NFS is likely attributed to this “open market” approach.

48.3 Focus: Simple and Fast Server Crash Recovery

In this chapter, we will discuss the classic NFS protocol (version 2, a.k.a. NFSv2), which was the standard for many years; small changes were made in moving to NFSv3, and larger-scale protocol changes were made in moving to NFSv4. However, NFSv2 is both wonderful and frustrating and thus serves as our focus.

In NFSv2, the main goal in the design of the protocol was simple and fast server crash recovery. In a multiple-client, single-server environment, this goal makes a great deal of sense; any minute that the server is down (or unavailable) makes all the client machines (and their users) unhappy and unproductive. Thus, as the server goes, so does the entire system.
48.4 Key To Fast Crash Recovery: Statelessness

This simple goal is realized in NFSv2 by designing what we refer to as a stateless protocol. The server, by design, does not keep track of anything about what is happening at each client. For example, the server does not know which clients are caching which blocks, or which files are currently open at each client, or the current file pointer position for a file, etc. Simply put, the server does not track anything about what clients are doing; rather, the protocol is designed to deliver in each protocol request all the information that is needed in order to complete the request. If it doesn’t now, this stateless approach will make more sense as we discuss the protocol in more detail below.

For an example of a stateful (not stateless) protocol, consider the open() system call. Given a pathname, open() returns a file descriptor (an integer). This descriptor is used on subsequent read() or write() requests to access various file blocks, as in this application code (note that proper error checking of the system calls is omitted for space reasons):

```
char buffer[MAX];
int fd = open("foo", O_RDONLY); // get descriptor "fd"
read(fd, buffer, MAX); // read MAX bytes from foo (via fd)
read(fd, buffer, MAX); // read MAX bytes from foo
... read(fd, buffer, MAX); // read MAX bytes from foo
close(fd); // close file
```

Figure 48.3: Client Code: Reading From A File

Now imagine that the client-side file system opens the file by sending a protocol message to the server saying “open the file ’foo’ and give me back a descriptor”. The file server then opens the file locally on its side and sends the descriptor back to the client. On subsequent reads, the client application uses that descriptor to call the read() system call; the client-side file system then passes the descriptor in a message to the file server, saying “read some bytes from the file that is referred to by the descriptor I am passing you here”.

In this example, the file descriptor is a piece of shared state between the client and the server (Ousterhout calls this distributed state [O91]). Shared state, as we hinted above, complicates crash recovery. Imagine the server crashes after the first read completes, but before the client has issued the second one. After the server is up and running again, the client then issues the second read. Unfortunately, the server has no idea to which file fd is referring; that information was ephemeral (i.e., in memory) and thus lost when the server crashed. To handle this situation, the client and server would have to engage in some kind of recovery protocol, where the client would make sure to keep enough information around in its memory to be able to tell the server what it needs to know (in this case, that file descriptor fd refers to file foo).
It gets even worse when you consider the fact that a stateful server has to deal with client crashes. Imagine, for example, a client that opens a file and then crashes. The open() uses up a file descriptor on the server; how can the server know it is OK to close a given file? In normal operation, a client would eventually call close() and thus inform the server that the file should be closed. However, when a client crashes, the server never receives a close(), and thus has to notice the client has crashed in order to close the file.

For these reasons, the designers of NFS decided to pursue a stateless approach: each client operation contains all the information needed to complete the request. No fancy crash recovery is needed; the server just starts running again, and a client, at worst, might have to retry a request.

48.5 The NFSv2 Protocol

We thus arrive at the NFSv2 protocol definition. Our problem statement is simple:

**THE CRUX: HOW TO DEFINE A STATELESS FILE PROTOCOL**

How can we define the network protocol to enable stateless operation? Clearly, stateful calls like open() can't be a part of the discussion (as it would require the server to track open files); however, the client application will want to call open(), read(), write(), close() and other standard API calls to access files and directories. Thus, as a refined question, how do we define the protocol to both be stateless and support the POSIX file system API?

One key to understanding the design of the NFS protocol is understanding the file handle. File handles are used to uniquely describe the file or directory a particular operation is going to operate upon; thus, many of the protocol requests include a file handle.

You can think of a file handle as having three important components: a volume identifier, an inode number, and a generation number; together, these three items comprise a unique identifier for a file or directory that a client wishes to access. The volume identifier informs the server which file system the request refers to (an NFS server can export more than one file system); the inode number tells the server which file within that partition the request is accessing. Finally, the generation number is needed when reusing an inode number; by incrementing it whenever an inode number is reused, the server ensures that a client with an old file handle can't accidentally access the newly-allocated file.

Here is a summary of some of the important pieces of the protocol; the full protocol is available elsewhere (see Callaghan's book for an excellent and detailed overview of NFS [C00]).
NFSPROC_GETATTR
  expects: file handle
  returns: attributes

NFSPROC_SETATTR
  expects: file handle, attributes
  returns: nothing

NFSPROC_LOOKUP
  expects: directory file handle, name of file/directory to look up
  returns: file handle

NFSPROC_READ
  expects: file handle, offset, count
  returns: data, attributes

NFSPROC_WRITE
  expects: file handle, offset, count, data
  returns: attributes

NFSPROC_CREATE
  expects: directory file handle, name of file, attributes
  returns: nothing

NFSPROC_REMOVE
  expects: directory file handle, name of file to be removed
  returns: nothing

NFSPROC_MKDIR
  expects: directory file handle, name of directory, attributes
  returns: file handle

NFSPROC_RMDIR
  expects: directory file handle, name of directory to be removed
  returns: nothing

NFSPROC_READDIR
  expects: directory handle, count of bytes to read, cookie
  returns: directory entries, cookie (to get more entries)

Figure 48.4: The NFS Protocol: Examples

We briefly highlight the important components of the protocol. First, the LOOKUP protocol message is used to obtain a file handle, which is then subsequently used to access file data. The client passes a directory file handle and name of a file to look up, and the handle to that file (or directory) plus its attributes are passed back to the client from the server.

For example, assume the client already has a directory file handle for the root directory of a file system (/) (indeed, this would be obtained through the NFS mount protocol, which is how clients and servers first are connected together; we do not discuss the mount protocol here for sake of brevity). If an application running on the client opens the file /foo.txt, the client-side file system sends a lookup request to the server, passing it the root file handle and the name foo.txt; if successful, the file handle (and attributes) for foo.txt will be returned.

In case you are wondering, attributes are just the metadata that the file system tracks about each file, including fields such as file creation time, last modification time, size, ownership and permissions information, and so forth, i.e., the same type of information that you would get back if you called stat() on a file.

Once a file handle is available, the client can issue READ and WRITE protocol messages on a file to read or write the file, respectively. The READ protocol message requires the protocol to pass along the file handle
of the file along with the offset within the file and number of bytes to read. The server then will be able to issue the read (after all, the handle tells the server which volume and which inode to read from, and the offset and count tells it which bytes of the file to read) and return the data to the client (or an error if there was a failure). WRITE is handled similarly, except the data is passed from the client to the server, and just a success code is returned.

One last interesting protocol message is the GETATTR request; given a file handle, it simply fetches the attributes for that file, including the last modified time of the file. We will see why this protocol request is important in NFSv2 below when we discuss caching (can you guess why?).

48.6 From Protocol to Distributed File System

Hopefully you are now getting some sense of how this protocol is turned into a file system across the client-side file system and the file server. The client-side file system tracks open files, and generally translates application requests into the relevant set of protocol messages. The server simply responds to each protocol message, each of which has all the information needed to complete request.

For example, let us consider a simple application which reads a file. In the diagram (Figure 48.1), we show what system calls the application makes, and what the client-side file system and file server do in responding to such calls.

A few comments about the figure. First, notice how the client tracks all relevant state for the file access, including the mapping of the integer file descriptor to an NFS file handle as well as the current file pointer. This enables the client to turn each read request (which you may have noticed do not specify the offset to read from explicitly) into a properly-formatted read protocol message which tells the server exactly which bytes from the file to read. Upon a successful read, the client updates the current file position; subsequent reads are issued with the same file handle but a different offset.

Second, you may notice where server interactions occur. When the file is opened for the first time, the client-side file system sends a LOOKUP request message. Indeed, if a long pathname must be traversed (e.g., /home/remzi/foo.txt), the client would send three LOOKUPs: one to look up home in the directory /, one to look up remzi in home, and finally one to look up foo.txt in remzi.

Third, you may notice how each server request has all the information needed to complete the request in its entirety. This design point is critical to be able to gracefully recover from server failure, as we will now discuss in more detail; it ensures that the server does not need state to be able to respond to the request.
<table>
<thead>
<tr>
<th>Client</th>
<th>Server</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \text{fd} = \text{open}(&quot;/foo&quot;, ...); ) Send \text{LOOKUP} (rootdir FH, “foo”)</td>
<td>Receive \text{LOOKUP} request look for “foo” in root dir return foo’s FH + attributes</td>
</tr>
<tr>
<td>Receive \text{LOOKUP} reply allocate file desc in open file table store foo’s FH in table store current file position (0) return file descriptor to application</td>
<td></td>
</tr>
</tbody>
</table>

**read(fd, buffer, MAX);**
- Index into open file table with \( \text{fd} \)
  - get NFS file handle (FH)
  - use current file position as offset
- Send \text{READ} (FH, offset=0, count=MAX)

Receive \text{READ} request use FH to get volume/inode num read inode from disk (or cache) compute block location (using offset) read data from disk (or cache) return data to client

Receive \text{READ} reply update file position (+bytes read) set current file position = MAX return data/error code to app

**read(fd, buffer, MAX);**
- Same except offset=MAX and set current file position = 2*MAX

**read(fd, buffer, MAX);**
- Same except offset=2*MAX and set current file position = 3*MAX

**close(fd);**
- Just need to clean up local structures
  - Free descriptor ”fd” in open file table
  - (No need to talk to server)

Table 48.1: Reading A File: Client-side And File Server Actions
TIP: IDEMPOTENCY IS POWERFUL

Idempotency is a useful property when building reliable systems. When an operation can be issued more than once, it is much easier to handle failure of the operation; you can just retry it. If an operation is not idempotent, life becomes more difficult.

48.7 Handling Server Failure with Idempotent Operations

When a client sends a message to the server, it sometimes does not receive a reply. There are many possible reasons for this failure to respond. In some cases, the message may be dropped by the network; networks do lose messages, and thus either the request or the reply could be lost and thus the client would never receive a response.

It is also possible that the server has crashed, and thus is not currently responding to messages. After a bit, the server will be rebooted and start running again, but in the meanwhile all requests have been lost. In all of these cases, clients are left with a question: what should they do when the server does not reply in a timely manner?

In NFSv2, a client handles all of these failures in a single, uniform, and elegant way: it simply retries the request. Specifically, after sending the request, the client sets a timer to go off after a specified time period. If a reply is received before the timer goes off, the timer is canceled and all is well. If, however, the timer goes off before any reply is received, the client assumes the request has not been processed and resends it. If the server replies, all is well and the client has neatly handled the problem.

The ability of the client to simply retry the request (regardless of what caused the failure) is due to an important property of most NFS requests: they are idempotent. An operation is called idempotent when the effect of performing the operation multiple times is equivalent to the effect of performing the operation a single time. For example, if you store a value to a memory location three times, it is the same as doing so once; thus “store value to memory” is an idempotent operation. If, however, you increment a counter three times, it results in a different amount than doing so just once; thus, “increment counter” is not idempotent. More generally, any operation that just reads data is obviously idempotent; an operation that updates data must be more carefully considered to determine if it has this property.

The heart of the design of crash recovery in NFS is the idempotency of most common operations. LOOKUP and READ requests are trivially idempotent, as they only read information from the file server and do not update it. More interestingly, WRITE requests are also idempotent. If, for example, a WRITE fails, the client can simply retry it. The WRITE message contains the data, the count, and (importantly) the exact offset to write the data to. Thus, it can be repeated with the knowledge that the outcome of multiple writes is the same as the outcome of a single one.
In this way, the client can handle all timeouts in a unified way. If a WRITE request was simply lost (Case 1 above), the client will retry it, the server will perform the write, and all will be well. The same will happen if the server happened to be down while the request was sent, but back up and running when the second request is sent, and again all works as desired (Case 2). Finally, the server may in fact receive the WRITE request, issue the write to its disk, and send a reply. This reply may get lost (Case 3), again causing the client to re-send the request. When the server receives the request again, it will simply do the exact same thing: write the data to disk and reply that it has done so. If the client this time receives the reply, all is again well, and thus the client has handled both message loss and server failure in a uniform manner. Neat!

A small aside: some operations are hard to make idempotent. For example, when you try to make a directory that already exists, you are informed that the mkdir request has failed. Thus, in NFS, if the file server receives a MKDIR protocol message and executes it successfully but the reply is lost, the client may repeat it and encounter that failure when in fact the operation at first succeeded and then only failed on the retry. Thus, life is not perfect.
Even when you design a beautiful system, sometimes all the corner cases don’t work out exactly as you might like. Take the mkdir example above; one could redesign mkdir to have different semantics, thus making it idempotent (think about how you might do so); however, why bother? The NFS design philosophy covers most of the important cases, and overall makes the system design clean and simple with regards to failure. Thus, accepting that life isn’t perfect and still building the system is a sign of good engineering. Apparently, this wisdom is attributed to Voltaire, for saying ”... a wise Italian says that the best is the enemy of the good” [V72], and thus we call it Voltaire’s Law.

48.8 Improving Performance: Client-side Caching

Distributed file systems are good for a number of reasons, but sending all read and write requests across the network can lead to a big performance problem: the network generally isn’t that fast, especially as compared to local memory or disk. Thus, another problem: how can we improve the performance of a distributed file system?

The answer, as you might guess from reading the big bold words in the sub-heading above, is client-side caching. The NFS client-side file system caches file data (and metadata) that it has read from the server in client memory. Thus, while the first access is expensive (i.e., it requires network communication), subsequent accesses are serviced quite quickly out of client memory.

The cache also serves as a temporary buffer for writes. When a client application first writes to a file, the client buffers the data in client memory (in the same cache as the data it read from the file server) before writing the data out to the server. Such write buffering is useful because it decouples application write() latency from actual write performance, i.e., the application’s call to write() succeeds immediately (and just puts the data in the client-side file system’s cache); only later does the data get written out to the file server.

Thus, NFS clients cache data and performance is usually great and we are done, right? Unfortunately, not quite. Adding caching into any sort of system with multiple client caches introduces a big and interesting challenge which we will refer to as the cache consistency problem.

48.9 The Cache Consistency Problem

The cache consistency problem is best illustrated with two clients and a single server. Imagine client C1 reads a file F, and keeps a copy of the file in its local cache. Now imagine a different client, C2, overwrites the file F, thus changing its contents; let’s call the new version of the file F

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You can probably see the problem that is upcoming (Figure 48.6). In fact, there are two subproblems. The first subproblem is that the client C2 may buffer its writes in its cache for a time before propagating them to the server; in this case, while F[v2] sits in C2’s memory, any access of F from another client (say C3) will fetch the old version of the file (F[v1]). Thus, by buffering writes at the client, other clients may get stale versions of the file, which may be undesirable; indeed, imagine the case where you log into machine C2, update F, and then log into C3 and try to read the file, only to get the old copy! Certainly this could be frustrating. Thus, let us call this aspect of the cache consistency problem update visibility; when do updates from one client become visible at other clients?

The second subproblem of cache consistency is a stale cache; in this case, C2 has finally flushed its writes to the file server, and thus the server has the latest version (F[v2]). However, C1 still has F[v1] in its cache; if a program running on C1 reads file F, it will get a stale version (F[v1]) and not the most recent copy (F[v2]), which is (often) undesirable.

NFSv2 implementations solve these cache consistency problems in two ways. First, to address update visibility, clients implement what is sometimes called flush-on-close (a.k.a., close-to-open) consistency semantics; specifically, when a file is written to and subsequently closed by a client application, the client flushes all updates (i.e., dirty pages in the cache) to the server. With flush-on-close consistency, NFS ensures that a subsequent open from another node will see the latest file version.

Second, to address the stale-cache problem, NFSv2 clients first check to see whether a file has changed before using its cached contents. Specifically, when opening a file, the client-side file system will issue a GETATTR request to the server to fetch the file’s attributes. The attributes, importantly, include information as to when the file was last modified on the server; if the time-of-modification is more recent than the time that the file was fetched into the client cache, the client invalidates the file, thus removing it from the client cache and ensuring that subsequent reads will go to the server and retrieve the latest version of the file. If, on the other
hand, the client sees that it has the latest version of the file, it will go ahead and use the cached contents, thus increasing performance.

When the original team at Sun implemented this solution to the stale-cache problem, they realized a new problem; suddenly, the NFS server was flooded with GETATTR requests. A good engineering principle to follow is to design for the common case, and to make it work well; here, although the common case was that a file was accessed only from a single client (perhaps repeatedly), the client always had to send GETATTR requests to the server to make sure no one else had changed the file. A client thus bombards the server, constantly asking “has anyone changed this file?”, when most of the time no one had.

To remedy this situation (somewhat), an attribute cache was added to each client. A client would still validate a file before accessing it, but most often would just look in the attribute cache to fetch the attributes. The attributes for a particular file were placed in the cache when the file was first accessed, and then would timeout after a certain amount of time (say 3 seconds). Thus, during those three seconds, all file accesses would determine that it was OK to use the cached file and thus do so with no network communication with the server.

48.10 Assessing NFS Cache Consistency

A few final words about NFS cache consistency. The flush-on-close behavior was added to “make sense”, but introduced a certain performance problem. Specifically, if a temporary or short-lived file was created on a client and then soon deleted, it would still be forced to the server. A more ideal implementation might keep such short-lived files in memory until they are deleted and thus remove the server interaction entirely, perhaps increasing performance.

More importantly, the addition of an attribute cache into NFS made it very hard to understand or reason about exactly what version of a file one was getting. Sometimes you would get the latest version; sometimes you would get an old version simply because your attribute cache hadn’t yet timed out and thus the client was happy to give you what was in client memory. Although this was fine most of the time, it would (and still does!) occasionally lead to odd behavior.

And thus we have described the oddity that is NFS client caching. It serves as an interesting example where details of an implementation serve to define user-observable semantics, instead of the other way around.

48.11 Implications on Server-Side Write Buffering

Our focus so far has been on client caching, and that is where most of the interesting issues arise. However, NFS servers tend to be well-equipped machines with a lot of memory too, and thus they have caching concerns as well. When data (and metadata) is read from disk, NFS
servers will keep it in memory, and subsequent reads of said data (and metadata) will not go to disk, a potential (small) boost in performance.

More intriguing is the case of write buffering. NFS servers absolutely may not return success on a WRITE protocol request until the write has been forced to stable storage (e.g., to disk or some other persistent device). While they can place a copy of the data in server memory, returning success to the client on a WRITE protocol request could result in incorrect behavior; can you figure out why?

The answer lies in our assumptions about how clients handle server failure. Imagine the following sequence of writes as issued by a client:

```
write(fd, a_buffer, size); // fill first block with a’s
write(fd, b_buffer, size); // fill second block with b’s
write(fd, c_buffer, size); // fill third block with c’s
```

These writes overwrite the three blocks of a file with a block of a’s, then b’s, and then c’s. Thus, if the file initially looked like this:

```
xxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxx
yyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyy
zzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzzz
```

We might expect the final result after these writes to be like this, with the x’s, y’s, and z’s, would be overwritten with a’s, b’s, and c’s, respectively.

```
aaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaa
yyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyy
cccccccccccccccccccccccccccccccccccccccccccccccccccccccccccc
```

Now let’s assume for the sake of the example that these three client writes were issued to the server as three distinct WRITE protocol messages. Assume the first WRITE message is received by the server and issued to the disk, and the client informed of its success. Now assume the second write is just buffered in memory, and the server also reports it success to the client before forcing it to disk; unfortunately, the server crashes before writing it to disk. The server quickly restarts and receives the third write request, which also succeeds.

Thus, to the client, all the requests succeeded, but we are surprised that the file contents look like this:

```
aaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaa
yyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyyy <--- oops
cccccccccccccccccccccccccccccccccccccccccccccccccccccccccccc
```

Yikes! Because the server told the client that the second write was successful before committing it to disk, an old chunk is left in the file, which, depending on the application, might be catastrophic.

To avoid this problem, NFS servers must commit each write to stable (persistent) storage before informing the client of success; doing so enables the client to detect server failure during a write, and thus retry until
it finally succeeds. Doing so ensures we will never end up with file contents intermingled as in the above example.

The problem that this requirement gives rise to in NFS server implementation is that write performance, without great care, can be the major performance bottleneck. Indeed, some companies (e.g., Network Appliance) came into existence with the simple objective of building an NFS server that can perform writes quickly; one trick they use is to first put writes in a battery-backed memory, thus enabling to quickly reply to WRITE requests without fear of losing the data and without the cost of having to write to disk right away; the second trick is to use a file system design specifically designed to write to disk quickly when one finally needs to do so [HLM94, RO91].

48.12 Summary

We have seen the introduction of the NFS distributed file system. NFS is centered around the idea of simple and fast recovery in the face of server failure, and achieves this end through careful protocol design. Idempotency of operations is essential; because a client can safely replay a failed operation, it is OK to do so whether or not the server has executed the request.

We also have seen how the introduction of caching into a multiple-client, single-server system can complicate things. In particular, the system must resolve the cache consistency problem in order to behave reasonably; however, NFS does so in a slightly ad hoc fashion which can occasionally result in observably weird behavior. Finally, we saw how server caching can be tricky: writes to the server must be forced to stable storage before returning success (otherwise data can be lost).

We haven’t talked about other issues which are certainly relevant, notably security. Security in early NFS implementations was remarkably lax; it was rather easy for any user on a client to masquerade as other users and thus gain access to virtually any file. Subsequent integration with more serious authentication services (e.g., Kerberos [NT94]) have addressed these obvious deficiencies.
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Hitz et al. were greatly influenced by previous work on log-structured file systems.

Mendel Rosenblum, John Ousterhout
Symposium on Operating Systems Principles (SOSP), 1991
LFS again. No, you can never get enough LFS.

[V72] “La Begueule”
Francois-Marie Arouet a.k.a. Voltaire
Published in 1772
Voltaire said a number of clever things, this being but one example. For example, Voltaire also said “If you have two religions in your land, the two will cut each others throats; but if you have thirty religions, they will dwell in peace.” What do you say to that, Democrats and Republicans?
The Andrew File System (AFS)

The Andrew File System was introduced by researchers at Carnegie-Mellon University (CMU) in the 1980’s [H+88]. Led by the well-known Professor M. Satyanarayanan of Carnegie-Mellon University (“Satya” for short), the main goal of this project was simple: scale. Specifically, how can one design a distributed file system such that a server can support as many clients as possible?

Interestingly, there are numerous aspects of design and implementation that affect scalability. Most important is the design of the protocol between clients and servers. In NFS, for example, the protocol forces clients to check with the server periodically to determine if cached contents have changed; because each check uses server resources (including CPU and network bandwidth), frequent checks like this will limit the number of clients a server can respond to and thus limit scalability.

AFS also differs from NFS in that from the beginning, reasonable user-visible behavior was a first-class concern. In NFS, cache consistency is hard to describe because it depends directly on low-level implementation details, including client-side cache timeout intervals. In AFS, cache consistency is simple and readily understood: when the file is opened, a client will generally receive the latest consistent copy from the server.

49.1 AFS Version 1

We will discuss two versions of AFS [H+88, S+85]. The first version (which we will call AFSv1, but actually the original system was called the ITC distributed file system [S+85]) had some of the basic design in place, but didn’t scale as desired, which led to a re-design and the final protocol (which we will call AFSv2, or just AFS) [H+88]. We now discuss the first version.

One of the basic tenets of all versions of AFS is whole-file caching on the local disk of the client machine that is accessing a file. When you open() a file, the entire file (if it exists) is fetched from the server and stored in a file on your local disk. Subsequent application read() and write() operations are redirected to the local file system where the file is
TestAuth  Test whether a file has changed
       (used to validate cached entries)
GetFileStat Get the stat info for a file
Fetch    Fetch the contents of file
Store    Store this file on the server
SetFileStat Set the stat info for a file
ListDir  List the contents of a directory

Figure 49.1: AFSv1 Protocol Highlights

stored; thus, these operations require no network communication and are fast. Finally, upon close(), the file (if it has been modified) is flushed back to the server. Note the obvious contrasts with NFS, which caches blocks (not whole files, although NFS could of course cache every block of an entire file) and does so in client memory (not local disk).

Let’s get into the details a bit more. When a client application first calls open(), the AFS client-side code (which the AFS designers call Venus) would send a Fetch protocol message to the server. The Fetch protocol message would pass the entire pathname of the desired file (for example, /home/remzi/notes.txt) to the file server (the group of which they called Vice), which would then traverse the pathname, find the desired file, and ship the entire file back to the client. The client-side code would then cache the file on the local disk of the client (by writing it to local disk). As we said above, subsequent read() and write() system calls are strictly local in AFS (no communication with the server occurs); they are just redirected to the local copy of the file. Because the read() and write() calls act just like calls to a local file system, once a block is accessed, it also may be cached in client memory. Thus, AFS also uses client memory to cache copies of blocks that it has in its local disk. Finally, when finished, the AFS client checks if the file has been modified (i.e., that it has been opened for writing); if so, it flushes the new version back to the server with a Store protocol message, sending the entire file and pathname to the server for permanent storage.

The next time the file is accessed, AFSv1 does so much more efficiently. Specifically, the client-side code first contacts the server (using the TestAuth protocol message) in order to determine whether the file has changed. If not, the client would use the locally-cached copy, thus improving performance by avoiding a network transfer. The figure above shows some of the protocol messages in AFSv1. Note that this early version of the protocol only cached file contents; directories, for example, were only kept at the server.

49.2 Problems with Version 1

A few key problems with this first version of AFS motivated the designers to rethink their file system. To study the problems in detail, the designers of AFS spent a great deal of time measuring their existing prototype to find what was wrong. Such experimentation is a good thing;
measurement is the key to understanding how systems work and how to improve them. Hard data helps take intuition and make into a concrete science of deconstructing systems. In their study, the authors found two main problems with AFSv1:

- **Path-traversal costs are too high**: When performing a Fetch or Store protocol request, the client passes the entire pathname (e.g., `/home/remzi/notes.txt`) to the server. The server, in order to access the file, must perform a full pathname traversal, first looking in the root directory to find `home`, then in `home` to find `remzi`, and so forth, all the way down the path until finally the desired file is located. With many clients accessing the server at once, the designers of AFS found that the server was spending much of its CPU time simply walking down directory paths.

- **The client issues too many TestAuth protocol messages**: Much like NFS and its overabundance of GETATTR protocol messages, AFSv1 generated a large amount of traffic to check whether a local file (or its stat information) was valid with the TestAuth protocol message. Thus, servers spent much of their time telling clients whether it was OK to used their cached copies of a file. Most of the time, the answer was that the file had not changed.

There were actually two other problems with AFSv1: load was not balanced across servers, and the server used a single distinct process per client thus inducing context switching and other overheads. The load imbalance problem was solved by introducing volumes, which an administrator could move across servers to balance load; the context-switch problem was solved in AFSv2 by building the server with threads instead of processes. However, for the sake of space, we focus here on the main two protocol problems above that limited the scale of the system.
49.3 Improving the Protocol

The two problems above limited the scalability of AFS; the server CPU became the bottleneck of the system, and each server could only service 20 clients without becoming overloaded. Servers were receiving too many TestAuth messages, and when they received Fetch or Store messages, were spending too much time traversing the directory hierarchy. Thus, the AFS designers were faced with a problem:

THE CRUX: HOW TO DESIGN A SCALABLE FILE PROTOCOL

How should one redesign the protocol to minimize the number of server interactions, i.e., how could they reduce the number of TestAuth messages? Further, how could they design the protocol to make these server interactions efficient? By attacking both of these issues, a new protocol would result in a much more scalable version AFS.

49.4 AFS Version 2

AFSv2 introduced the notion of a callback to reduce the number of client/server interactions. A callback is simply a promise from the server to the client that the server will inform the client when a file that the client is caching has been modified. By adding this state to the server, the client no longer needs to contact the server to find out if a cached file is still valid. Rather, it assumes that the file is valid until the server tells it otherwise; insert analogy to polling versus interrupts here.

AFSv2 also introduced the notion of a file identifier (FID) (similar to the NFS file handle) instead of pathnames to specify which file a client was interested in. An FID in AFS consists of a volume identifier, a file identifier, and a “uniquifier” (to enable reuse of the volume and file IDs when a file is deleted). Thus, instead of sending whole pathnames to the server and letting the server walk the pathname to find the desired file, the client would walk the pathname, one piece at a time, caching the results and thus hopefully reducing the load on the server.

For example, if a client accessed the file /home/remzi/notes.txt, and home was the AFS directory mounted onto / (i.e., / was the local root directory, but home and its children were in AFS), the client would first Fetch the directory contents of home, put them in the local-disk cache, and setup a callback on home. Then, the client would Fetch the directory remzi, put it in the local-disk cache, and setup a callback on the server on remzi. Finally, the client would Fetch notes.txt, cache this regular file in the local disk, setup a callback, and finally return a file descriptor to the calling application. See Table 49.1 for a summary.

The key difference, however, from NFS, is that with each fetch of a directory or file, the AFS client would establish a callback with the server, thus ensuring that the server would notify the client of a change in its
Table 49.1: Reading A File: Client-side And File Server Actions

cached state. The benefit is obvious: although the first access to /home/remzi/notes.txt generates many client-server messages (as described above), it also establishes callbacks for all the directories as well as the file notes.txt, and thus subsequent accesses are entirely local and require no server interaction at all. Thus, in the common case where a file is cached at the client, AFS behaves nearly identically to a local disk-based file system. If one accesses a file more than once, the second access should be just as fast as accessing a file locally.
When discussing distributed file systems, much is made of the cache consistency the file systems provide. However, this baseline consistency does not solve all problems with regards to file access from multiple clients. For example, if you are building a code repository, with multiple clients performing check-ins and check-outs of code, you can’t simply rely on the underlying file system to do all of the work for you; rather, you have to use explicit file-level locking in order to ensure that the “right” thing happens when such concurrent accesses take place. Indeed, any application that truly cares about concurrent updates will add extra machinery to handle conflicts. The baseline consistency described in this chapter and the previous one are useful primarily for casual usage, i.e., when a user logs into a different client, they expect some reasonable version of their files to show up there. Expecting more from these protocols is setting yourself up for failure, disappointment, and tear-filled frustration.

49.5 Cache Consistency

When we discussed NFS, there were two aspects of cache consistency we considered: update visibility and cache staleness. With update visibility, the question is: when will the server be updated with a new version of a file? With cache staleness, the question is: once the server has a new version, how long before clients see the new version instead of an older cached copy?

Because of callbacks and whole-file caching, the cache consistency provided by AFS is easy to describe and understand. There are two important cases to consider: consistency between processes on different machines, and consistency between processes on the same machine.

Between different machines, AFS makes updates visible at the server and invalidates cached copies at the exact same time, which is when the updated file is closed. A client opens a file, and then writes to it (perhaps repeatedly). When it is finally closed, the new file is flushed to the server (and thus visible); the server then breaks callbacks for any clients with cached copies, thus ensuring that clients will no longer read stale copies of the file; subsequent opens on those clients will require a re-fetch of the new version of the file from the server.

AFS makes an exception to this simple model between processes on the same machine. In this case, writes to a file are immediately visible to other local processes (i.e., a process does not have to wait until a file is closed to see its latest updates). This makes using a single machine behave exactly as you would expect, as this behavior is based upon typical Unix semantics. Only when switching to a different machine would you be able to detect the more general AFS consistency mechanism.

There is one interesting cross-machine case that is worthy of further discussion. Specifically, in the rare case that processes on different ma-
chines are modifying a file at the same time, AFS naturally employs what is known as a last writer wins approach (which perhaps should be called last closer wins). Specifically, whichever client calls close() last will update the entire file on the server last and thus will be the “winning” file, i.e., the file that remains on the server for others to see. The result is a file that was generated in its entirety either by one client or the other. Note the difference from a block-based protocol like NFS: in NFS, writes of individual blocks may be flushed out to the server as each client is updating the file, and thus the final file on the server could end up as a mix of updates from both clients. In many cases, such a mixed file output would not make much sense, i.e., imagine a JPEG image getting modified by two clients in pieces; the resulting mix of writes would not likely constitute a valid JPEG.

A timeline showing a few of these different scenarios can be seen in Table 49.2. The columns of the table show the behavior of two processes (P₁ and P₂) on Client₁ and its cache state, one process (P₃) on Client₂ and its cache state, and the server (Server), all operating on a single file called, imaginatively, F. For the server, the table just shows the contents of the file after the operation on the left has completed. Read through it and see if you can understand why each read returns the results that it does. A commentary field on the right will help you if you get stuck.

<table>
<thead>
<tr>
<th>Client₁</th>
<th>Client₂</th>
<th>Server</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>P₁</td>
<td>P₂</td>
<td>P₃</td>
<td>Cache</td>
</tr>
<tr>
<td>open(F)</td>
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<td>-</td>
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<tr>
<td>write(A)</td>
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<td></td>
<td>A</td>
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<tr>
<td>close()</td>
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<td>A</td>
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<tr>
<td>open(F)</td>
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<td></td>
<td>A</td>
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<tr>
<td>read() → A</td>
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<td></td>
<td>A</td>
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<td>close()</td>
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<td>A</td>
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<tr>
<td>open(F)</td>
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<td>A</td>
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<td>write(B)</td>
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<td>A</td>
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<tr>
<td>read() → B</td>
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<td>A</td>
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<td>close()</td>
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<td>open(F)</td>
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<td>read() → A</td>
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<td>open(F)</td>
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<td>write(B)</td>
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<tr>
<td>read() → B</td>
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<td>close()</td>
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<td>close()</td>
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<tr>
<td>open(F)</td>
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<tr>
<td>write(D)</td>
<td></td>
<td></td>
<td>A</td>
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<td>close()</td>
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<td>A</td>
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<td>close()</td>
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<tr>
<td>open(F)</td>
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<td>write(C)</td>
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<td>A</td>
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<td>close()</td>
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<td>A</td>
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<td>close()</td>
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<td></td>
<td>A</td>
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</table>

Table 49.2: Cache Consistency Timeline
49.6 Crash Recovery

From the description above, you might sense that crash recovery is more involved than with NFS. You would be right. For example, imagine there is a short period of time where a server (S) is not able to contact a client (C1), for example, while the client C1 is rebooting. While C1 is not available, S may have tried to send it one or more callback recall messages; for example, imagine C1 had file F cached on its local disk, and then C2 (another client) updated F, thus causing S to send messages to all clients caching the file to remove it from their local caches. Because C1 may miss those critical messages when it is rebooting, upon rejoining the system, C1 should treat all of its cache contents as suspect. Thus, upon the next access to file F, C1 should first ask the server (with a TestAuth protocol message) whether its cached copy of file F is still valid; if so, C1 can use it; if not, C1 should fetch the newer version from the server.

Server recovery after a crash is also more complicated. The problem that arises is that callbacks are kept in memory; thus, when a server reboots, it has no idea which client machine has which files. Thus, upon server restart, each client of the server must realize that the server has crashed and treat all of their cache contents as suspect, and (as above) reestablish the validity of a file before using it. Thus, a server crash is a big event, as one must ensure that each client is aware of the crash in a timely manner, or risk a client accessing a stale file. There are many ways to implement such recovery; for example, by having the server send a message (saying “don’t trust your cache contents!”) to each client when it is up and running again, or by having clients check that the server is alive periodically (with a heartbeat message, as it is called). As you can see, there is a cost to building a more scalable and sensible caching model; with NFS, clients hardly noticed a server crash.

49.7 Scale And Performance Of AFSv2

With the new protocol in place, AFSv2 was measured and found to be much more scalable that the original version. Indeed, each server could support about 50 clients (instead of just 20). A further benefit was that client-side performance often came quite close to local performance, because in the common case, all file accesses were local; file reads usually went to the local disk cache (and potentially, local memory). Only when a client created a new file or wrote to an existing one was there need to send a Store message to the server and thus update the file with new contents.

Let us also gain some perspective on AFS performance by comparing common file-system access scenarios with NFS. Table 49.3 shows the results of our qualitative comparison.

In the table, we examine typical read and write patterns analytically, for files of different sizes. Small files have \( N_s \) blocks in them; medium files have \( N_m \) blocks; large files have \( N_L \) blocks. We assume that small
and medium files fit into the memory of a client; large files fit on a local disk but not in client memory.

We also assume, for the sake of analysis, that an access across the network to the remote server for a file block takes $L_{net}$ time units. Access to local memory takes $L_{mem}$, and access to local disk takes $L_{disk}$. The general assumption is that $L_{net} > L_{disk} > L_{mem}$.

Finally, we assume that the first access to a file does not hit in any caches. Subsequent file accesses (i.e., “re-reads”) we assume will hit in caches, if the relevant cache has enough capacity to hold the file.

The columns of the table show the time a particular operation (e.g., a small file sequential read) roughly takes on either NFS or AFS. The rightmost column displays the ratio of AFS to NFS.

We make the following observations. First, in many cases, the performance of each system is roughly equivalent. For example, when first reading a file (e.g., Workloads 1, 3, 5), the time to fetch the file from the remote server dominates, and is similar on both systems. You might think AFS would be slower in this case, as it has to write the file to local disk; however, those writes are buffered by the local (client-side) file system cache and thus said costs are likely hidden. Similarly, you might think that AFS reads from the local cached copy would be slower, again because AFS stores the cached copy on disk. However, AFS again benefits here from local file system caching; reads on AFS would likely hit in the client-side memory cache, and performance would be similar to NFS.

Second, an interesting difference arises during a large-file sequential re-read (Workload 6). Because AFS has a large local disk cache, it will access the file from there when the file is accessed again. NFS, in contrast, only can cache blocks in client memory; as a result, if a large file (i.e., a file bigger than local memory) is re-read, the NFS client will have to re-fetch the entire file from the remote server. Thus, AFS is faster than NFS in this case by a factor of $\frac{L_{net}}{L_{disk}}$, assuming that remote access is indeed slower than local disk. We also note that NFS in this case increases server load, which has an impact on scale as well.
Third, we note that sequential writes (of new files) should perform similarly on both systems (Workloads 8, 9). AFS, in this case, will write the file to the local cached copy; when the file is closed, the AFS client will force the writes to the server, as per the protocol. NFS will buffer writes in client memory, perhaps forcing some blocks to the server due to client-side memory pressure, but definitely writing them to the server when the file is closed, to preserve NFS flush-on-close consistency. You might think AFS would be slower here, because it writes all data to local disk. However, realize that it is writing to a local file system; those writes are first committed to the page cache, and only later (in the background) to disk, and thus AFS reaps the benefits of the client-side OS memory caching infrastructure to improve performance.

Fourth, we note that AFS performs worse on a sequential file overwrite (Workload 10). Thus far, we have assumed that the workloads that write are also creating a new file; in this case, the file exists, and is then over-written. Overwrite can be a particularly bad case for AFS, because the client first fetches the old file in its entirety, only to subsequently overwrite it. NFS, in contrast, will simply overwrite blocks and thus avoid the initial (useless) read.

Finally, workloads that access a small subset of data within large files perform much better on NFS than AFS (Workloads 7, 11). In these cases, the AFS protocol fetches the entire file when the file is opened; unfortunately, only a small read or write is performed. Even worse, if the file is modified, the entire file is written back to the server, doubling the performance impact. NFS, as a block-based protocol, performs I/O that is proportional to the size of the read or write.

Overall, we see that NFS and AFS make different assumptions and not surprisingly realize different performance outcomes as a result. Whether these differences matter is, as always, a question of workload.

49.8 AFS: Other Improvements

Like we saw with the introduction of Berkeley FFS (which added symbolic links and a number of other features), the designers of AFS took the opportunity when building their system to add a number of features that made the system easier to use and manage. For example, AFS provides a true global namespace to clients, thus ensuring that all files were named the same way on all client machines. NFS, in contrast, allows each client to mount NFS servers in any way that they please, and thus only by convention (and great administrative effort) would files be named similarly across clients.

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1We assume here that NFS reads are block-sized and block-aligned; if they were not, the NFS client would also have to read the block first. We also assume the file was not opened with the O_TRUNC flag; if it had been, the initial open in AFS would not fetch the soon to be truncated file’s contents.
One challenge of evaluating any system is the choice of workload. Because computer systems are used in so many different ways, there are a large variety of workloads to choose from. How should the storage system designer decide which workloads are important, in order to make reasonable design decisions?

The designers of AFS, given their experience in measuring how file systems were used, made certain workload assumptions; in particular, they assumed that most files were not frequently shared, and accessed sequentially in their entirety. Given those assumptions, the AFS design makes perfect sense.

However, these assumptions are not always correct. For example, imagine an application that appends information, periodically, to a log. These little log writes, which add small amounts of data to an existing large file, are quite problematic for AFS. Many other difficult workloads exist as well, e.g., random updates in a transaction database.

One place to get some information about what types of workloads are common are through various research studies that have been performed. See any of these studies for good examples of workload analysis [B+91, H+11, R+00, V99], including the AFS retrospective [H+88].

AFS also takes security seriously, and incorporates mechanisms to authenticate users and ensure that a set of files could be kept private if a user so desired. NFS, in contrast, had quite primitive support for security for many years.

AFS also includes facilities for flexible user-managed access control. Thus, when using AFS, a user has a great deal of control over who exactly can access which files. NFS, like most UNIX file systems, has much less support for this type of sharing.

Finally, as mentioned before, AFS adds tools to enable simpler management of servers for the administrators of the system. In thinking about system management, AFS was light years ahead of the field.

49.9 Summary

AFS shows us how distributed file systems can be built quite differently than what we saw with NFS. The protocol design of AFS is particularly important; by minimizing server interactions (through whole-file caching and callbacks), each server can support many clients and thus reduce the number of servers needed to manage a particular site. Many other features, including the single namespace, security, and access-control lists, make AFS quite nice to use. The consistency model provided by AFS is simple to understand and reason about, and does not lead to the occasional weird behavior as one sometimes observes in NFS.
Perhaps unfortunately, AFS is likely on the decline. Because NFS became an open standard, many different vendors supported it, and, along with CIFS (the Windows-based distributed file system protocol), NFS dominates the marketplace. Although one still sees AFS installations from time to time (such as in various educational institutions, including Wisconsin), the only lasting influence will likely be from the ideas of AFS rather than the actual system itself. Indeed, NFSv4 now adds server state (e.g., an “open” protocol message), and thus bears an increasing similarity to the basic AFS protocol.
References

[B+91] “Measurements of a Distributed File System”
Mary Baker, John Hartman, Martin Kupfer, Ken Shirriff, John Ousterhout
SOSP ’91, Pacific Grove, CA, October 1991
An early paper measuring how people use distributed file systems. Matches much of the intuition found in AFS.

Tyler Harter, Chris Dragga, Michael Vaughn, Andrea C. Arpaci-Dusseau, Remzi H. Arpaci-Dusseau
SOSP ’11, New York, NY, October 2011
Our own paper studying the behavior of Apple Desktop workloads; turns out they are a bit different than many of the server-based workloads the systems research community usually focuses upon. Also a good recent reference which points to a lot of related work.

[H+88] “Scale and Performance in a Distributed File System”
John H. Howard, Michael L. Kazar, Sherri G. Menees, David A. Nichols, M. Satyanarayanan, Robert N. Sidebotham, Michael J. West
ACM Transactions on Computing Systems (ACM TOCS), page 51-81, Volume 6, Number 1, February 1988
The long journal version of the famous AFS system, still in use in a number of places throughout the world, and also probably the earliest clear thinking on how to build distributed file systems. A wonderful combination of the science of measurement and principled engineering.

[R+00] “A Comparison of File System Workloads”
Drew Roselli, Jacob R. Lorch, Thomas E. Anderson
USENIX ’00, San Diego, CA, June 2000
A more recent set of traces as compared to the Baker paper [B+91], with some interesting twists.

M. Satyanarayanan, J.H. Howard, D.A. Nichols, R.N. Sidebotham, A. Spector, M.J. West
SOSP ’85, pages 35-50
The older paper about a distributed file system. Much of the basic design of AFS is in place in this older system, but not the improvements for scale.

[V99] “File system usage in Windows NT 4.0”
Werner Vogels
SOSP ’99, Kiawah Island Resort, SC, December 1999
A cool study of Windows workloads, which are inherently different than many of the UNIX-based studies that had previously been done.
Summary Dialogue on Distribution

Student: Well, that was quick. Too quick, in my opinion!
Professor: Yes, distributed systems are complicated and cool and well worth your study; just not in this book (or course).
Student: That’s too bad; I wanted to learn more! But I did learn a few things.
Professor: Like what?
Student: Well, everything can fail.
Professor: Good start.
Student: But by having lots of these things (whether disks, machines, or whatever), you can hide much of the failure that arises.
Professor: Keep going!
Student: Some basic techniques like retrying are really useful.
Professor: That’s true.
Student: And you have to think carefully about protocols: the exact bits that are exchanged between machines. Protocols can affect everything, including how systems respond to failure and how scalable they are.
Professor: You really are getting better at this learning stuff.
Student: Thanks! And you’re not a bad teacher yourself!
Professor: Well thank you very much too.
Student: So is this the end of the book?
Professor: I’m not sure. They don’t tell me anything.
Student: Me neither. Let’s get out of here.
Professor: OK.
Student: Go ahead.
Professor: No, after you.
Student: Please, professors first.
Professor: No, please, after you.
Student: (exasperated) Fine!
Professor: (waiting) ... so why haven’t you left?
Student: I don’t know how. Turns out, the only thing I can do is participate in these dialogues.
Professor: Me too. And now you’ve learned our final lesson...
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