Optimizing	Lempel	Ziv	Welch	for	DNA	Compres	sion

A Thesis

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I want to thank a few people.

Preface

This is an example of a thesis setup to use the reed thesis document class (for LaTeX) and the R bookdown package, in general.

List of Abbreviations

LZW Lempel Ziv Welch

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Abstract

The Lempel Ziv Welch compression algorithm is a lossless data compression algorithm used for numerous applications, including the Unix file compression utility compress and the GIF image format. Storing, reading, and transferring enormous amounts of data is often an issue in the biological field, especially when concerning DNA. This thesis explores the application of Lempel Ziv Welch to the compression of DNA. A variety of different optimization of the original LZW algorithm are explore included palatalizing, multiple dictionaries, and some other cool thing here broh.

Dedication

You can have a dedication here if you wish.

Introduction

When dealing with DNA, it

Chapter 1

Backround and Motivations

This thesis deals with some high level topics and uses language specific to compression research. This chapter tries to give brief summaries and examples of the relevant topics to be discussed so readers of all experience levels can put our results into context.

1.1 What is information?

Suppose you had an idea that you wanted to share with another person. Humans have many ways to communicate information; you could send a text message, you could tell them with words, you could tell them with sign language. But regardless of the medium, you have some idea that you want to get across. Does it matter if the other person gets your message exactly? Or can it be part of the message? If someone asks you "Where library", despite the lack of prepositions you still understand what they mean. So did that person convey any less information than a person who asks "Where is the library?" Clearly, information is fundamental to how humans interact and how they understand the world, but defining it proves difficult. For our purposes, let us assume that information is something that can be interpreted to glean information that you didn't know before.

Information on computers can take many forms, such as text, audio, and video. This information can travel through many channels including the internet, wires, screens, etc. To maximize the amount of information that can be transmitted through a channel, we need to encode the information in a way which minimizes its size, while also preserving its essential features. This process is called compression.

1.2 Compression: A history

The idea of compressing data to make transmission easier has been around for a long time. One of the earliest uses of compression was the use of Morse code, developed by Samuel Morse in the 1830s, used to transmit messages over telegraph. Morse uses a binary code, where dots and dashes represent different letters and numbers of the message to be transmitted.

In the 1970s, researchers began to develop more sophisticated compression algoritms, such as Huffman encoding and Lempel-Ziv compression. Both LZW and Huffman use codes to represent different characters in a message.

TODO: Elaborate ## Compression Metrics

1.2.1 Compression Ratio

Compression Ratio is the measure of size reduction achieved by a compression algorithm. It is typically expressed as a ratio of the size of the uncompressed data (OS) to the size of the compressed data (CS).

$$CR = \frac{OS}{CS}$$

So a higher compression ratio means a more effective compression algorithm, and means that we were able to store more data in less space, allowing for easier storage and transfer.

1.2.2 Runtime

The runtime is also an important part of evaluating the effectiveness of a compression algorithm. If you have the option of two compression algorithms, one with a compression ratio of 2.0, and another with a compression ratio of 2.15 but takes twice as long as the other, you may opt for a lower compression ratio to save time.

1.2.3 Memory Usage

Memory usage is closely tied with runtime when it comes to compression algorithms. Memory generally refers to information that programs track as they are running on a computer. So do reduce our runtime and make a more effective compression algorithm, we want to be saving only the most important data that our algorithm needs in order to reduce our memory usage.

1.3 Lossless vs. Lossy Compression

1.3.1 Lossy

Lossy compression is based on the idea that not all information is vital. For instance, when saving a picture on your computer, your computer may save it in the .jpeg format to save space. Jpegs lose some of the information in the original picture and produce an overall lower quality picture, but the general information in the picture is preserved. Another example

TODO: Continue examples

1.3.2 Lossless

Lossless compression is the compression of data with the goal of preserving all the information in the data. As a result, lossless compression algorithms usually don't compress as well as their lossy counterparts. Examples of lossless compression algoritms are Huffman Encoding and Lempel Ziv Welch, which is the focus of this thesis.

TODO: Elaborate more here

1.4 Examples of Compression Algorithms

1.4.1 Run Length Encoding

Run Length Encoding (RLE) is on of the simplest and most intuitive forms of compression. We can take advantage of redundant runs of characters in a sequence by just giving the number of times each character appears. Suppose you want to send the following message

AAGCTTTTTTTTGGGGGCCCT

Even if this message did mean something, we can get the information across without repeating ourselves. When writing a grocery list, you don't write "egg egg egg", you say "4 eggs". RLE uses this same strategdy.

2A1G1C8T5G3C1T

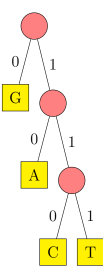
We could compress this even further if we omit the 1 on characters that only appear once. Although not as sophisticated as other methods, RLE is effective when used on texts that have a lot of repeating characters.

1.4.2 Huffman

Huffman Encoding is a strategy that assigns variable length code to certain symbols in the data. The goal is to assign short codes to frequently appearing symbols and longer codes to less frequent symbols.

Suppose we have a message "ACAGGATGGC". We can calculate the frequency of each letter by counting the number of times each letter shows up and dividing by the total number of letters

Then, we can use the frequencies to build a tree, which will assign short codes for frequent letters and longer code for less frequent letters.



So G = 0, A = 10, T = 111 and C = 110. Notice that none of the encodings are prefixes of one another, which makes it unambiguous in decoding.

So our message would be encoded to 1011010001011100110.

1.4.3 Arithmetic

Arithmetic encoding is another lossless compression algorithm that uses probability to assign codes to symbols in the message. Unlike Huffman, arithmetic encoding assigns a single code to the whole message, rather than separate codes for each symbol.

Here is a simple example. Say we want to encode a string of characters "ACGT". Arithmetic Encoding also requires the encoder and decoder know the probabilities of each of the characters that could possibly be in the message. Let's say the probability of each symbol in the message are

- P(A) = 1/10
- P(C) = 2/10

- P(G) = 4/10
- P(T) = 3/10

We want to represent the message as a fractional number between 0 and 1. We will divide the interval [0,1] into sub intervals using the probabilities of each character in the message. That way, each symbol is represented by the sub-interval that corresponds to its probability.

So since 'A' comes first, we divide [0,1] into [0.0,0.1). Since 'C' is next, we go from [0.0,0.1] to [0.01, 0.03). Then since 'G is next, we go from [0.01, 0.03) to [0.016, 0.02). Finally, since 'T' is last, we go from [0.016, 0.02) to [0.0188, 0.02).

So any number in the interval can be used to represent our message.

Arithmetic encoding can have a better compression ratio that Huffman in some cases, but the computation time is often not worth the payoff.

TODO: Add graphic

1.4.4 Lempel Ziv Welch

Lempel Ziv Welch is another lossless compression algorithm. When compressing, LZW builds a dictionary of codewords, where codewords represent strings previously seen in the message. As it compresses the message, the dictionary grows. The compression algorithm leaves behind the codewords and some of the original characters, allowing the decompression algorithm to build up the same dictionary as it decompresses the message.

Here is a simple example. We may be sending messages with the characters {'A', 'C', 'T', 'G'}, so I will start with those in my dictionary. Say we want to send the message

"AAGGAATCC"

When we compress, we start at the beginning of the message and scan through.

TODO: Finish example, maybe add some pseudo code as well

Chapter 2

Optimizing LZW: Approach

To restate the goal of this thesis, we seek to optimize LZW for use in compression of DNA. I chose to write in C++.

TODO: Elaborate here

2.1 Supporting Research

There has been several attempts to optimize LZW by computer science researchers. One paper made use of multiple indexed dictionaries in order to speed up the compression process (Keerthy, 2019). The concept is simple, rather than a single large dictionary, have multiple dictionaries, one for each possible string size. That way, the dictionaries grow more slowly and accesses are faster. This paper also used Genomic data to gather their metrics and compared their algorithm to other popular DNA compression techniques, which makes it particularly relevant for this paper.

Another paper used simple parallelization techniques to improve compression speed (Pani, Mishra, & Mishra, 2012). Rather than compressing the whole file linearly, the researches broke the file into portions and compressed them with LZW in parallel, which greatly increased the compression speed at the cost of a reduced compression ratio.

Yet another paper made use of Chinese Remainder Theorem to augment Lempel Ziv Welch (Ibrahim & Gbolagade, 2020). They saw great reduction in compression time without comprimising compression ratio, although these results could not be verified. The details of their implementation were not clear from the paper. We tried multiple different methods of utilizing CRT given the pseudocode in their paper, but we could not get anything that looked like it may improve compression time. We reached out to the authors of the paper, but we were not able to further our progress

on this method and thus the method is not used in this thesis.

2.2 Corpora

Most compression papers make use of a Corpus, which is a collection of files to run a compression algorithm on in order to evaluate performance and to compare different algorithms to one another.

In the world of DNA compression, there are several academic papers on the subject. One of the first and most popular of the papers was published in 1994, and the selection of DNA sequences used in the paper have become an informal corpus for the subject of DNA compression, cited by more than thirty publications (Grumbach & Tahi, 1994).

Name	Size.bytes.		
chmpxx	121024		
chntxx	155844		
hehcmv	229354		
humdyst	38770		
humghcs	66495		
humhbb	73308		
humhdab	58864		
humprtb	56737		
mpomtcg	186609		
mtpacga	100314		
vaccg	191737		

Another, newer paper aimed to create a corpus specifically for compressing DNA (Pratas & Pinho, 2018). They put together a corpus of DNA sequences for this purpose, as summarized below. Since the papers publishing, it has been cited by several DNA compression papers.

Name	Size.bytes.
AeCa	1591049
AgPh	43970
BuEb	18940

Name	Size.bytes.
DaRe	62565020
DrMe	32181429
EnIn	26403087
EsCo	4641652
GaGa	148532294
НаНі	3890005
HePy	1667825
HoSa	189752667
OrSa	43262523
PlFa	8986712
ScPo	10652155
YeMi	73689

This particular dataset is publicly available at this link.

2.3 Evaluating Performance

Evaluating performance of a program is difficult. There is a notion of theoretical run time, but on an actual computer there are many processes running in the background, so it can be hard to get a consistent reading on performance.

To attempt to counteract this, we ran the function on the same file multiple times, and took the median of the compression and decompression times for all the runs.

2.4 A Starting Point

As a starting point, we thought it was best to get a working implementation of LZW in C++ on regular text files, optimize it as much as we could, and then try variations from there, optimizing it for DNA. We want to try various techniques tried by researches in the field, but it is important to have a fast baseline from which we can compare and improve upon.

2.4.1 Growing Codewords and Bit Output

When reading files on the computer, most characters are stored as bytes, which is made up of 8 bits. For instance 01000001 stands for the letter 'A' in ASCII encoding.

Numbers are more simple to display, so 00000001 is 1, 00000010 is 2, and so on.

But if we are translating numbers to binary, we don't need all of the bits in a byte. In binary, 1 is the same as 01 is the same as 0000000000001. So when we are outputting codewords for LZW, we don't necessarily need to output a whole byte. We can have growing codewords.

As the number of codewords grows, the number of bits needed to represent it also grows. So if we are on codeword 8, we need 4 bits since 8 is 1000. As our dictionary grows, we can grow the number of bits needed to display a codeword and save a lot of space in our compressed document.

So we needed a method of outputting bits one by one, and reading in bits one by one. This is not something that is supported in C++ on its own. We were able to create this functionality by defining a class.

```
// BitInput: Read a single bit at a time from an input stream.
// Before reading any bits, ensure your input stream still has valid inputs.
class BitInput {
public:
 // Construct with an input stream
 BitInput(const char* input);
 BitInput(const BitInput&) = default;
 BitInput(BitInput&&) = default;
 // bool eof();
 // Read a single bit (or trailing zero)
 // Allowed to crash or throw an exception if called past end-of-file.
 bool input_bit();
 int read_n_bits(int n);
}
// BitOutput: Write a single bit at a time to an output stream
// Make sure all bits are written out by the time the destructor is done.
class BitOutput {
public:
 // Construct with an input stream
```

So when we are encoding and need to output a codeword, we can output_n_bits, where n is the number of bits needed to display our greatest codeword. When decoding, we can just read_n_bits.

2.4.2 Getting EOF to work

One of the very early issues with the implementation was how to denote the end of a file. The early implementation would work for some files, but for others the very last part of the file would be lost after encoding and then decoding.

In theoretical implementations of LZW, computer scientists tend to denote the end of a message with a special character, one that isn't seen anywhere else in the file. In this initial implementation, that wasn't possible because we wanted to be able to compress any file with any characters.

The solution was to reserve a codeword to mark the end of the file. So we start with a starting dictionary containing all ASCII characters.

```
std::unordered_map<std::string, int> dictionary;
for (int i = 0; i < 256; ++i){
    std::string str1(1, char(i));
    dictionary[str1] = i;
}</pre>
```

Then use the code 256 to denote the end of file. So the algorithm goes along reading a file. It builds up a current string character by character, adding the character

to the string and checking if it has seen that sequence before. Once it find the end of file, we stop and output the EOF codeword.

The problem was, what about what is left over? Suppose we are reading a file, and the file ends with "ACCT". If "A" is in the dictionary, we see if "AC" is in the dictionary, and so on. This leaves us with three possible cases when we reached the end of the file

- 1. "ACC" was in the dictionary but "ACCT" was not. This means we can output the codeword for "ACC", follow it by the character "T", and we are done. This is the ideal scenario, because nothing is left over when we output the EOF codeword
- 2. "ACCT" was in the dictionary: This means we have one more codeword to output, but since we reached the end of the file, we never got to output it.
- 3. "AC" was in the dictionary, but "ACC" was not: in this case, we would output the codeword for "AC" output the character "C", and then start looping again starting at "T". But we reach the end of the file, so we output EOF before outputting T.

We solved this issue by adding 2 extra bits after the EOF codeword. These bits denote the case that occurred

```
// after we've encoded, we either have
// no current block (case 0)
// we have a current block that is a single character (case 1)
// otherwise we have a current block > 1 byte (default)
switch (currentBlock.length()){
case 0:
   bit output.output bit(false);
   bit output.output bit(false);
   break;
case 1:
   bit output.output bit(false);
   bit output.output bit(true);
   bit output.output n bits((int) currentBlock[0], CHAR BIT);
   break;
default:
   bit output.output bit(true);
```

```
bit_output.output_bit(true);

int code = dictionary[currentBlock];
bit_output.output_n_bits(code, codeword_size);
break;
}
```

So when the decoded is reading and encounters the EOF codeword, it can look at the next two bits to see if anything is left over.

At this point, there was a working implementation that was able to compress and decompress files. Here is the performance of this version on the two copora.

File.Name	Original.File.Size	Compressed.Size	Compression.Ratio	Compression.Time	Decompression.Time
DNACorpus1/humprtb	56737	21902	2.590	7	2
${\rm DNACorpus1/humdyst}$	38770	15300	2.534	5	1
DNACorpus1/vaccg	191737	70067	2.736	18	7
DNACorpus1/hehcmv	229354	85526	2.682	23	9
${\rm DNACorpus1/mpomtcg}$	186609	70254	2.656	18	7
DNACorpus1/humhdab	58864	22699	2.593	8	2
${\rm DNACorpus1/chmpxx}$	121024	43516	2.781	13	4
${\rm DNACorpus1/mtpacga}$	100314	36862	2.721	11	4
DNACorpus1/chntxx	155844	58336	2.671	16	6
${\rm DNACorpus 1/humghcs}$	66495	25552	2.602	7	2
DNACorpus1/humhbb	73308	28134	2.606	8	3

File.Name	Original.File.Size	${\bf Compressed. Size}$	Compression.Ratio	Compression.Time	Decompression.Time
DNACorpus2/YeMi	73689	27235	2.706	9	3
DNACorpus2/DaRe	62565020	19586457	3.194	15034	1898
DNACorpus2/EnIn	26403087	8609993	3.067	5427	781
DNACorpus2/HePy	1667825	566972	2.942	145	40
${\rm DNACorpus2/OrSa}$	43262523	14148071	3.058	9790	1350
DNACorpus2/EsCo	4641652	1593404	2.913	569	121
DNACorpus2/GaGa	148532294	46851765	3.170	40810	4773
DNACorpus2/ScPo	10652155	3590856	2.966	1761	295
DNACorpus2/HaHi	3890005	1306708	2.977	445	97
${\rm DNACorpus2/HoSa}$	189752667	57200209	3.317	53818	5913
DNACorpus2/AeCa	1591049	556535	2.859	141	40
DNACorpus2/DrMe	32181429	10619042	3.031	7190	1015
DNACorpus2/BuEb	18940	7893	2.400	2	0
DNACorpus2/PlFa	8986712	2895744	3.103	1367	238
DNACorpus2/AgPh	43970	17442	2.521	6	2

2.4.3 Using Constants

The early version of the code was not clean. There were hard coded variables, unspecified integer types, and generally messy naming conventions that made the code difficult to read and debug.

The next major step in the code was to start using constants for everything, including

- STARTING CODEWORD: What codeword we should start at
- EOF CODEWORD: What we should output when we reach end of file
- STARTING_DICT_SIZE: At this stage, we had a starting dict size of 256 to hold all possible bytes, but later we will specialize for DNA

It also made sense to start using a specific type for codewords. At this stage, we opted for a 32 bit integer.

There are several tools at a developers disposal when looking to debug and optimize code. One tool used for this thesis was callgrind which is a tool of valgrind a profiling tool. Profiling tools are used to look at how your code works, where the bottlenecks are, and what can be changed/improved for the performance of your code.

Callgrind in particular is a profiling tool which associates assembly instructions to lines of code, indicating to the programmer which lines take a lot of instructions and which take less. For those unfamiliar, assembly instructions are what code is turned into so that it can be ran on your computer's processor. In general, more instructions means that code takes longer to run.

The callgrind output drew attention to one particular part of the code. A C++ unordered_map uses iterators, basically pointers into the dictionary. If an entry is not present in the dictionary, the find() function will return a iterator to the end of the dictionary.

The check for this in our algorithm looked like this.

```
// if we've already seen the sequence, keep going
// TODO: use cend() and save this iterator
if (dictionary.find(currentBlock + next_character) != dictionary.end()){
    currentBlock = currentBlock + next_character;
}
```

Here is the callgrind output for that line.

```
105,030,135 ( 0.18%) if (dictionary.find(currentBlock + next_character) != dictionary.find(currentBlock + next_character) != dictionary.find(curre
```

```
11,653,779,430 (19.65%) => /usr/include/c++/9/bits/unordered_map.h:std::unordered_map.h:std::unordered_map.h:std::unordered_map.h:std::_cxx11::basi 956,941,242 ( 1.61%) => /usr/include/c++/9/bits/unordered_map.h:std::unordered_map
241,180,314 ( 0.41%) => /usr/include/c++/9/bits/hashtable_policy.h:bool std::__deta
```

As shown, this line is taking a significant amount of instructions, and it needs to pull the end() of the dictionary each time it is ran. If we use cend() instead and save that iterator in a variable called end, we can save a significant amount of instructions.

```
89,470,115 ( 0.61%) if (dictionary.find(currentBlock + next_character) != e 3,353,009,053 (22.78%) => /usr/include/c++/9/bits/basic_string.h:std::_cxx11::basi 2,833,786,025 (19.26%) => /usr/include/c++/9/bits/unordered_map.h:std::unordered_ma 420,120,704 ( 2.85%) => /usr/include/c++/9/bits/basic_string.h:std::_cxx11::basic_50,570,065 ( 0.34%) => /usr/include/c++/9/bits/hashtable_policy.h:bool std::__detai
```

Of course, there are several issues with this method. It is difficult to associate instructions with a single line of code. Some lines are interdependent, and assembly often behaves differently than the code that produces it. Another thing is that compliers are very advanced, and sometimes small optimizations like this are done by the compiler automatically.

Despite these issues, this change was still worth making, if not to save time then for sake of clarity and readability of the code. Also, despite the inaccuracy of callgrind, like many profiling tools, its job is not necessarily to provide exact measurements of code performance, but to give indications to trouble spots which can be improved.

Here are the runs after this optimization.

File.Name	Original.File.Size	Compressed.Size	Compression.Ratio	Compression.Time	Decompression.Time
DNACorpus1/humprtb	56737	21902	2.590	7	2
DNACorpus1/humdyst	38770	15300	2.534	5	1
DNACorpus1/vaccg	191737	70067	2.736	19	7
DNACorpus1/hehcmv	229354	85526	2.682	23	9
${\rm DNACorpus1/mpomtcg}$	186609	70254	2.656	19	7
DNACorpus1/humhdab	58864	22699	2.593	7	2
${\rm DNACorpus1/chmpxx}$	121024	43516	2.781	12	4
DNACorpus1/mtpacga	100314	36862	2.721	11	4
DNACorpus1/chntxx	155844	58336	2.671	16	6
${\rm DNACorpus1/humghcs}$	66495	25552	2.602	9	2
DNACorpus1/humhbb	73308	28134	2.606	10	3

File.Name	Original.File.Size	Compressed.Size	Compression.Ratio	Compression.Time	Decompression.Time
DNACorpus2/YeMi	73689	27235	2.706	9	3
DNACorpus2/DaRe	62565020	19586457	3.194	14940	1880
DNACorpus2/EnIn	26403087	8609993	3.067	5401	769
DNACorpus2/HePy	1667825	566972	2.942	144	40
${\rm DNACorpus2/OrSa}$	43262523	14148071	3.058	9782	1344
DNACorpus2/EsCo	4641652	1593404	2.913	569	118
DNACorpus2/GaGa	148532294	46851765	3.170	41060	4787
DNACorpus2/ScPo	10652155	3590856	2.966	1738	292
DNACorpus2/HaHi	3890005	1306708	2.977	444	97
${\rm DNACorpus2/HoSa}$	189752667	57200209	3.317	53774	5854
DNACorpus2/AeCa	1591049	556535	2.859	139	40
DNACorpus2/DrMe	32181429	10619042	3.031	7188	1009
${\rm DNACorpus2/BuEb}$	18940	7893	2.400	2	1
DNACorpus2/PlFa	8986712	2895744	3.103	1352	237
${\rm DNACorpus2/AgPh}$	43970	17442	2.521	5	2

2.4.4 Extraneous String Concatenations

The LZW algorithm is build on iteration: we go through each character, adding it to our current block. If we've seen that current block before, we keep going. If not, we add that block to the dictionary and start over.

Another thing that I noticed from the callgrind output was that a lot of time/instructions are being spent on string concatenation. In general, string concatenation in most language, including C++, have a lot of overhead. A lot of implementations involve creating a new string every time you concatenate two existing string, which can have a significant performance penalty.

In the version of the algorithm at the time, every time we have already seen a sequence, we have to concatenate a character. I noticed that I was doing this concatenation multiple times without needing to.

```
// we concatenate the strings here
if (dictionary.find(currentBlock + next_character) != end ){
    // and here
    currentBlock = currentBlock + next_character;
}
else{
    // other code here ommitted
```

```
// and here!
dictionary[currentBlock + next_character] = codeword;
}
```

If I just concatenate them and save the output into a new string, that will save me from doing the concatenation 2 more times.

```
// save concatenation here

std::string string_seen_plus_new_char = current_string_seen + next_character

if (dictionary.find(string_seen_plus_new_char) != end ){
    current_string_seen = string_seen_plus_new_char;
}

else{

// other code omitted here

dictionary[string_seen_plus_new_char] = codeword;
}
```

Here are the statistics on the version of the algorithm after this change.

File.Name	Original.File.Size	Compressed.Size	Compression.Ratio	Compression.Time	Decompression.Time
DNACorpus1/humprtb	56737	21902	2.590	6	2
DNACorpus1/humdyst	38770	15300	2.534	4	1
DNACorpus1/vaccg	191737	70067	2.736	16	7
DNACorpus1/hehcmv	229354	85526	2.682	20	9
${\rm DNACorpus1/mpomtcg}$	186609	70254	2.656	17	7
DNACorpus1/humhdab	58864	22699	2.593	6	2
${\rm DNACorpus1/chmpxx}$	121024	43516	2.781	12	4
DNACorpus1/mtpacga	100314	36862	2.721	10	4
DNACorpus1/chntxx	155844	58336	2.671	15	6
${\rm DNACorpus1/humghcs}$	66495	25552	2.602	8	2
DNACorpus1/humhbb	73308	28134	2.606	8	3

File.Name	Original.File.Size	Compressed.Size	Compression.Ratio	Compression.Time	Decompression.Time
DNACorpus2/YeMi	73689	27235	2.706	8	3
DNACorpus2/DaRe	62565020	19586457	3.194	14450	1876
DNACorpus2/EnIn	26403087	8609993	3.067	5186	762
DNACorpus2/HePy	1667825	566972	2.942	129	38
${\rm DNACorpus2/OrSa}$	43262523	14148071	3.058	9392	1329
DNACorpus2/EsCo	4641652	1593404	2.913	525	119
DNACorpus2/GaGa	148532294	46851765	3.170	39700	4741
DNACorpus2/ScPo	10652155	3590856	2.966	1640	289
DNACorpus2/HaHi	3890005	1306708	2.977	410	96
${\rm DNACorpus2/HoSa}$	189752667	57200209	3.317	51901	5848
DNACorpus2/AeCa	1591049	556535	2.859	126	40
DNACorpus2/DrMe	32181429	10619042	3.031	6861	1001
${\rm DNACorpus2/BuEb}$	18940	7893	2.400	2	0
DNACorpus2/PlFa	8986712	2895744	3.103	1258	234
${\rm DNACorpus2/AgPh}$	43970	17442	2.521	5	2

2.4.5 Dictionary Accesses

Dictionary accesses can be expensive, especially with the standard library. We learned from callgrind that along with string operations, our program spent a lot of time doing dictionary accesses.

We looked for ways to reduce the number of lookups. At the time, the way the algorithm worked was that it looked up the current string and the next character in the dictionary. If that string is in the dictionary, we keep going. If not, we output the codeword for the current string.

But, the current string on this iteration is the current string from the last iteration, plus one character. So when we were on the previous iteration of the loop, we could save that lookup and prevent a second lookup.

See the code below

```
while(next_character != EOF){

// code omitted

// if we've already seen the sequence, keep going

std::string string_seen_plus_new_char = current_string_seen + next_character;

// save this iterator`

if (dictionary.find(string_seen_plus_new_char) != not_in_dictionary ){

current_string_seen = string_seen_plus_new_char;
```

```
else{

// shouldn't look up again
int code = dictionary[current_string_seen];

// code omitted
}

next_character = input.get();
}
```

We can save that lookup, like so.

```
while(next_character != EOF){
    // code ommitted
    // if we've already seen the sequence, keep going
    std::string string_seen_plus_new_char = current_string_seen + next_charact
    codeword_seen_now = dictionary.find(string_seen_plus_new_char);
    if (codeword_seen_now != not_in_dictionary ){
        current_string_seen = string_seen_plus_new_char;
       codeword_seen_previously = codeword_seen_now; // save codeword here
    }
    else{
        // lookup the current block in the dictionary and output it, along w
       int code = codeword_seen_previously->second; // on the next iteration
       // code omitted
    }
   next_character = input.get();
```

8

DNACorpus1/humhbb

File.Name	Original.File.Size	Compressed.Size	Compression.Ratio	Compression.Time	Decompression.Time
DNACorpus1/humprtb	56737	21902	2.590	5	2
DNACorpus1/humdyst	38770	15300	2.534	4	1
DNACorpus1/vaccg	191737	70067	2.736	17	7
DNACorpus1/hehcmv	229354	85526	2.682	20	9
${\bf DNAC orpus 1/mpomtcg}$	186609	70254	2.656	17	7
DNACorpus1/humhdab	58864	22699	2.593	6	2
DNACorpus1/chmpxx	121024	43516	2.781	11	4
DNACorpus1/mtpacga	100314	36862	2.721	11	3
DNACorpus1/chntxx	155844	58336	2.671	16	6
DNACorpus1/humghcs	66495	25552	2.602	7	2

28134

2.606

Here are the stats of the version of the algorithm after this change.

73308

File.Name	Original.File.Size	Compressed.Size	Compression.Ratio	Compression.Time	Decompression.Time
DNACorpus2/YeMi	73689	27235	2.706	7	3
DNACorpus2/DaRe	62565020	19586457	3.194	14146	1869
DNACorpus2/EnIn	26403087	8609993	3.067	5087	758
DNACorpus2/HePy	1667825	566972	2.942	127	39
${\rm DNACorpus2/OrSa}$	43262523	14148071	3.058	9278	1329
DNACorpus2/EsCo	4641652	1593404	2.913	519	115
DNACorpus2/GaGa	148532294	46851765	3.170	39265	4702
DNACorpus2/ScPo	10652155	3590856	2.966	1610	290
DNACorpus2/HaHi	3890005	1306708	2.977	405	97
${\rm DNACorpus2/HoSa}$	189752667	57200209	3.317	50936	5833
DNACorpus2/AeCa	1591049	556535	2.859	125	39
DNACorpus2/DrMe	32181429	10619042	3.031	6773	999
${\rm DNACorpus2/BuEb}$	18940	7893	2.400	2	0
DNACorpus2/PlFa	8986712	2895744	3.103	1254	231
${\rm DNACorpus2/AgPh}$	43970	17442	2.521	5	2

2.4.6 Using Const Char *

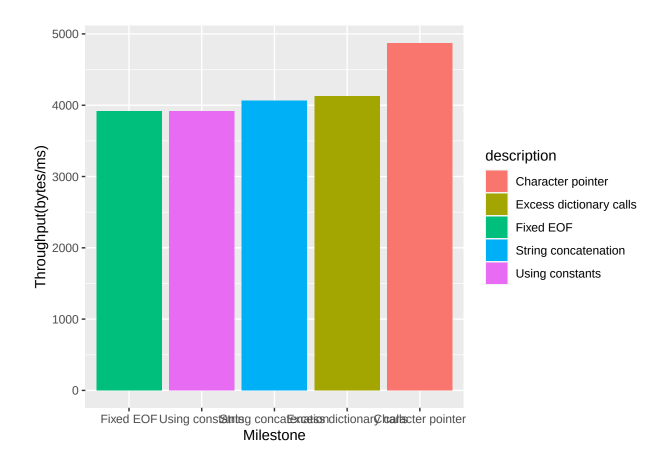
The algorithm works by reading through the entire file, so we know that at some point, we will need to see every byte of the entire file.

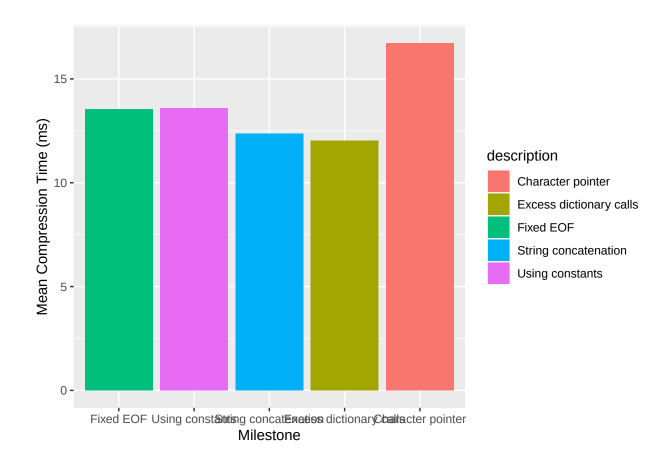
When reading a byte stream of the file, the file may not be in memory the way we want it. The ifstream class in C++ also has many extraneous feature that we don't need. If we map the file directly into memory using mmap and pass around a pointer to that data, it will simplify and speed up the scanning process. Also, using a char* opens the possibility to getting rid of std::string entirely, which means way less overhead and decreased compression time.

TODO: Need to talk about getting rid of string concatenation at some point.

File.Name	Original.File.Size	Compressed.Size	Compression.Ratio	Compression.Time	Decompression.Time
DNACorpus1/humprtb	56737	21902	2.590	7	2
DNACorpus1/humdyst	38770	15300	2.534	7	1
DNACorpus1/vaccg	191737	70067	2.736	15	6
DNACorpus1/hehcmv	229354	85526	2.682	15	8
${\rm DNACorpus1/mpomtcg}$	186609	70254	2.656	16	6
DNACorpus1/humhdab	58864	22699	2.593	9	2
DNACorpus1/chmpxx	121024	43516	2.781	12	4
DNACorpus1/mtpacga	100314	36862	2.721	10	3
DNACorpus1/chntxx	155844	58336	2.671	14	5
${\rm DNACorpus1/humghcs}$	66495	25552	2.602	9	2
DNACorpus1/humhbb	73308	28134	2.606	9	2

File.Name	Original.File.Size	Compressed.Size	Compression.Ratio	Compression.Time	Decompression.Time
DNACorpus2/YeMi	73689	27235	2.706	9	2
DNACorpus2/DaRe	62565020	19586457	3.194	12264	1764
DNACorpus2/EnIn	26403087	8609993	3.067	4070	708
DNACorpus2/HePy	1667825	566972	2.942	96	35
${\rm DNACorpus2/OrSa}$	43262523	14148071	3.058	7627	1252
DNACorpus2/EsCo	4641652	1593404	2.913	348	109
DNACorpus2/GaGa	148532294	46851765	3.170	32906	4456
DNACorpus2/ScPo	10652155	3590856	2.966	1205	265
DNACorpus2/HaHi	3890005	1306708	2.977	271	88
${\rm DNACorpus2/HoSa}$	189752667	57200209	3.317	44733	5517
DNACorpus2/AeCa	1591049	556535	2.859	92	35
DNACorpus2/DrMe	32181429	10619042	3.031	5266	940
DNACorpus2/BuEb	18940	7893	2.400	5	0
DNACorpus2/PlFa	8986712	2895744	3.103	879	220
${\rm DNACorpus2/AgPh}$	43970	17442	2.521	7	1





2.5 Trying Different Dictionaries

A lot of the stress of the LZW algorithm is on the dictionary. We are constantly looking strings up and placing others. Because of the reliance on this data structure, we know that the dictionary accesses and lookups are a bottleneck, so improvements in those areas could greatly increase the efficiency of our program.

So the another step towards an efficient LZW seemed to be to abstract out the std_map and have multiple different dictionary implementations to try and experiment with in our attempt to optimize LZW for DNA compression.

2.5.1 Direct Map

In our analysis of the two corpora, we found some interesting statistics in the redundancy of the data.

Below is a table which shows stats on the runs lengths of the "runs" of data, where a run is a string added to the dictionary during a run of the LZW algorithm.

average_run_length	maximum_run_length	median_run_length	sd_run_length
6.135398	17	6	1.237217

average_run_length	$maximum_run_length$	$median_run_length$	sd_run_length
10.96825	190	11	2.494305

And here is the combined stats of both copora, along with a histogram of all the runs.

X	average_run_length	$maximum_run_length$	$median_run_length$	sd_run_length
1	10.95318	190	11	2.505904

Given this data, it is clear that it would be advantageous to speed up the dictionary for smaller run sizes, since most of the runs are below size 20.

To achieve this, we opted for a unique approach. Rather than use a hashmap, we can map the strings directly into memory. Since all of the strings only contain four characters ('A', 'C', 'T', and 'G'), we can represent the characters with two bits. So for a length n string, we can represent it with 2n bits.

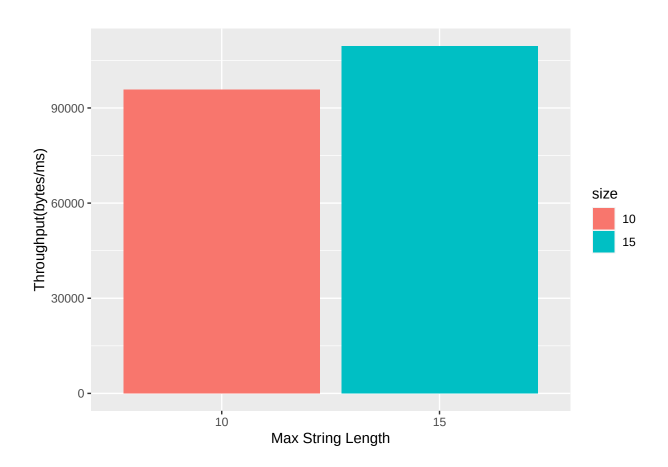
So we can create an indexed dictionary directly in memory for all strings below a certain length. We can use the 2n bit representation of the string to index into an array of codewords.

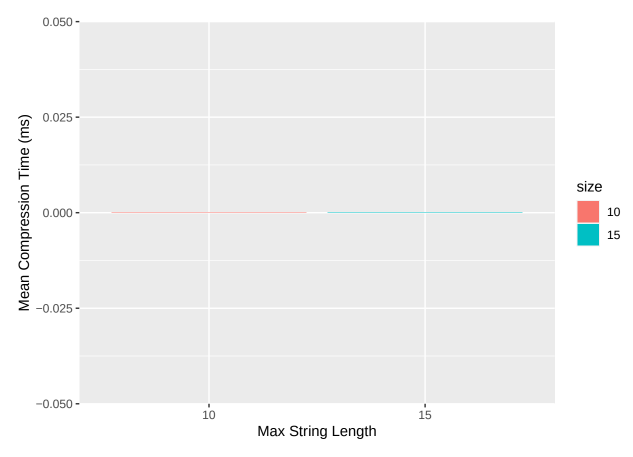
For each strings size 1 to n, we have an array with enough slots for every possible string. For example, for strings of length 3, we have an array of size 4^3 , since there are 4^3 possible strings. In each of those 4^3 slots, we have space for a codeword. All strings of length 3 can be represented by 6 bits, and since 6 bits can represent $2^6 = 4^3$ values, we can use the bit representation to index into the dictionary. If the codeword at that place in the dictionary is 0, we have never seen it before. If it is non-zero, we have found the codeword for that string. For all strings greater than n, we can just use a hashmap on top to handle those.

Here is the data for this version of the dictionary of length 10.

File.Name	Original.File.Size	Compressed.Size	Compression.Ratio	Compression.Time	Decompression.Time
DNACorpus1/humprtb	56737	25883	2.192	1	2
DNACorpus1/humdyst	38770	18437	2.103	1	1
DNACorpus1/vaccg	191737	77120	2.486	4	4
DNACorpus1/hehcmv	229354	93251	2.460	2	5
${\rm DNACorpus1/mpomtcg}$	186609	77315	2.414	4	4
DNACorpus1/humhdab	58864	26752	2.200	1	2
DNACorpus1/chmpxx	121024	49414	2.449	3	3
DNACorpus1/mtpacga	100314	42203	2.377	2	2
DNACorpus1/chntxx	155844	64879	2.402	3	3
${\rm DNACorpus1/humghcs}$	66495	29864	2.227	1	2
DNACorpus1/humhbb	73308	32681	2.243	1	2

File.Name	${\bf Original. File. Size}$	${\bf Compressed. Size}$	${\bf Compression. Ratio}$	${\bf Compression. Time}$	${\bf Decompression. Time}$
DNACorpus2/YeMi	73689	31700	2.325	1	2
DNACorpus2/DaRe	62565020	21182510	2.954	656	442
DNACorpus2/EnIn	26403087	8986109	2.938	271	184
${\rm DNACorpus2/HePy}$	1667825	572507	2.913	26	17
${\rm DNACorpus2/OrSa}$	43262523	15008736	2.882	431	319
DNACorpus2/EsCo	4641652	1621508	2.863	51	42
DNACorpus2/GaGa	148532294	52441876	2.832	1475	1080
${\rm DNACorpus2/ScPo}$	10652155	3691943	2.885	108	83
DNACorpus2/HaHi	3890005	1324571	2.937	51	32
${\rm DNACorpus2/HoSa}$	189752667	63973137	2.966	2035	1319
DNACorpus2/AeCa	1591049	561527	2.833	22	18
DNACorpus2/DrMe	32181429	11150498	2.886	332	237
${\rm DNACorpus2/BuEb}$	18940	9894	1.914	0	1
DNACorpus2/PlFa	8986712	2972072	3.024	106	67
DNACorpus2/AgPh	43970	20885	2.105	1	1





TODO: Have data for multiple different direct map sizes. Also talk about the fixed codeword size and what happens when we run out.

2.5.2 Multiple Indexed Dictionaries

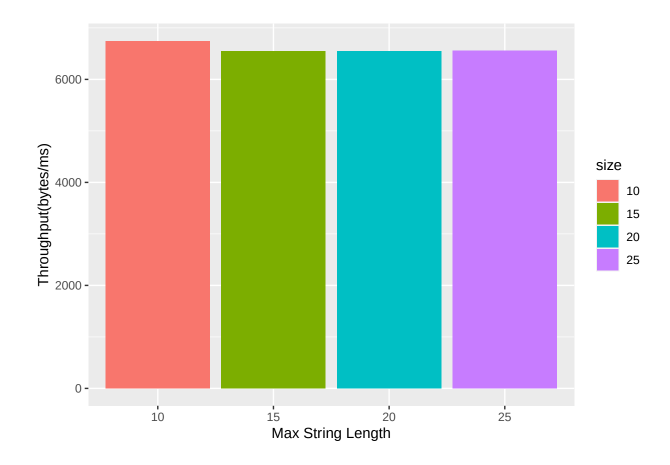
As stated before, there have been research papers about the

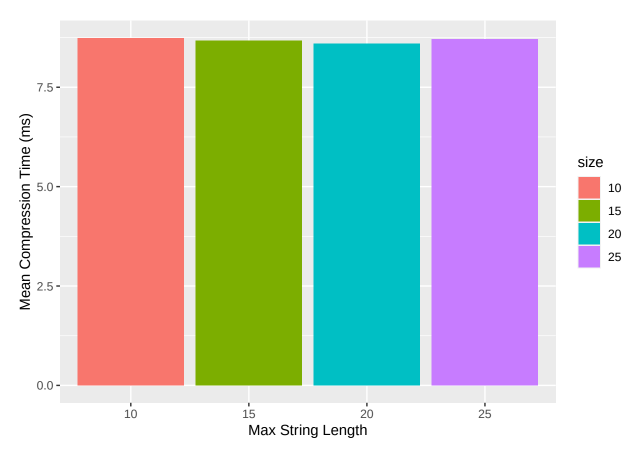
Similar to the Direct Mapped approach, we use dictionaries for each string size up to a certain size n, and for all strings of length greater than n, we use a regular dictionary. As with the direct map dictionary, we need to specify a max string length. We collected metrics for different choices of max string length.

Here are the results for this dictionary.

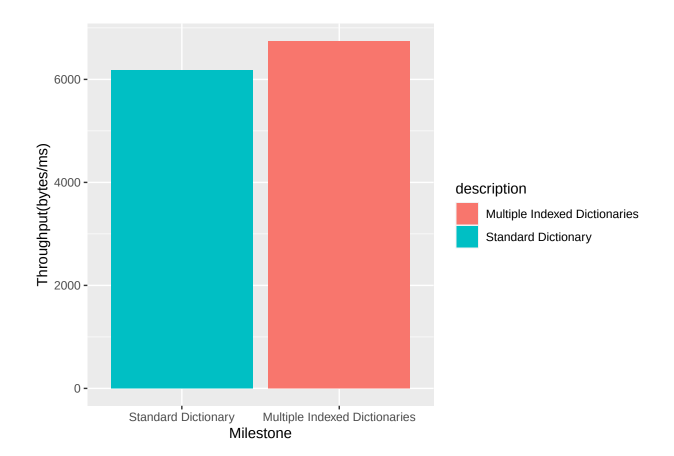
File.Name	Original.File.Size	Compressed.Size	Compression.Ratio	Compression.Time	Decompression.Time
DNACorpus1/humprtb	56737	21687	2.616	5	2
DNACorpus1/humdyst	38770	15116	2.565	3	1
DNACorpus1/vaccg	191737	69820	2.746	14	6
DNACorpus1/hehcmv	229354	85279	2.689	16	7
${\rm DNACorpus1/mpomtcg}$	186609	70007	2.666	15	6
DNACorpus1/humhdab	58864	22484	2.618	5	2
DNACorpus1/chmpxx	121024	43269	2.797	10	4
DNACorpus1/mtpacga	100314	36647	2.737	8	3
DNACorpus1/chntxx	155844	58089	2.683	12	5
DNACorpus1/humghcs	66495	25336	2.625	6	2
DNACorpus1/humhbb	73308	27918	2.626	7	2

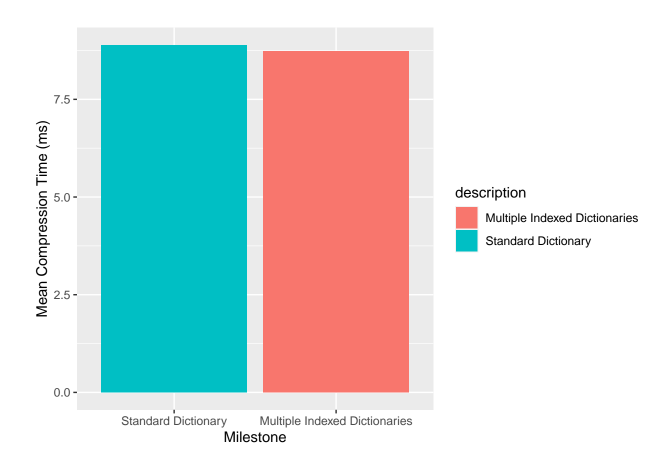
File.Name	Original.File.Size	Compressed.Size	Compression.Ratio	Compression.Time	Decompression.Time
DNACorpus2/YeMi	73689	27019	2.727	7	2
DNACorpus2/DaRe	62565020	19585959	3.194	8878	1721
DNACorpus2/EnIn	26403087	8609527	3.067	3347	696
DNACorpus2/HePy	1667825	566631	2.943	94	34
DNACorpus2/OrSa	43262523	14147605	3.058	5551	1223
DNACorpus2/EsCo	4641652	1593032	2.914	364	105
DNACorpus2/GaGa	148532294	46851235	3.170	22817	4323
DNACorpus2/ScPo	10652155	3590421	2.967	1099	258
DNACorpus2/HaHi	3890005	1306336	2.978	302	86
${\rm DNACorpus2/HoSa}$	189752667	57199679	3.317	31978	5449
DNACorpus2/AeCa	1591049	556194	2.861	91	34
DNACorpus2/DrMe	32181429	10618575	3.031	3865	905
DNACorpus2/BuEb	18940	7740	2.447	1	0
DNACorpus2/PlFa	8986712	2895340	3.104	892	211
DNACorpus2/AgPh	43970	17258	2.548	4	1





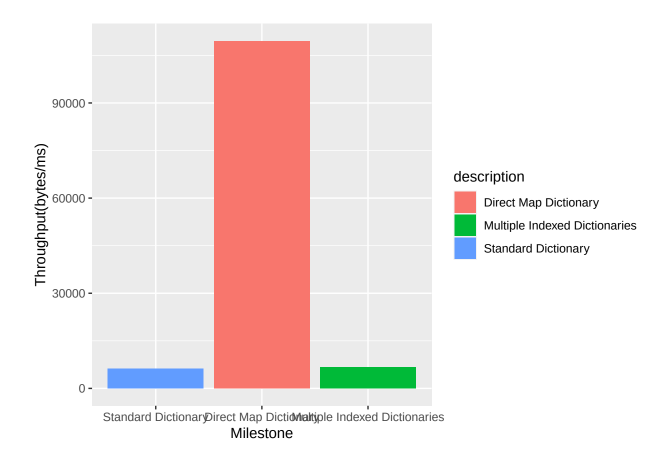
Here is a comparison of the Multiple Dictionaries versus one standard dictionary.

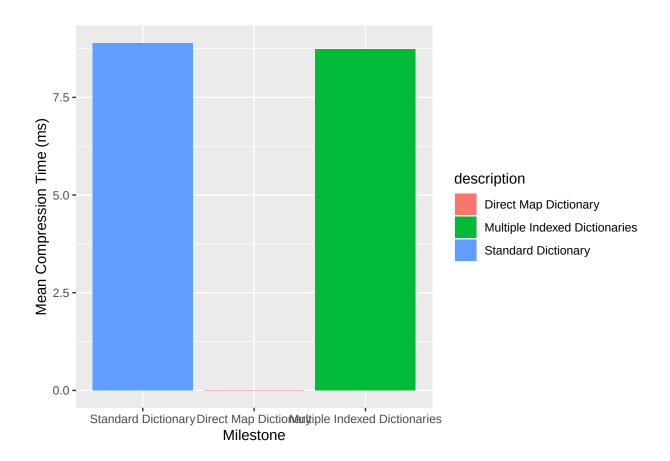




2.5.3 Comparison

Here is a comparison of all three techniques: Standard Dictionary, Multiple Standard Dictionaries, and Direct Mapped Dictionary.





Chapter 3

Graphics, References, and Labels

3.1 Figures

If your thesis has a lot of figures, R Markdown might behave better for you than that other word processor. One perk is that it will automatically number the figures accordingly in each chapter. You'll also be able to create a label for each figure, add a caption, and then reference the figure in a way similar to what we saw with tables earlier. If you label your figures, you can move the figures around and R Markdown will automatically adjust the numbering for you. No need for you to remember! So that you don't have to get too far into LaTeX to do this, a couple R functions have been created for you to assist. You'll see their use below.

In the **R** chunk below, we will load in a picture stored as reed.jpg in our main directory. We then give it the caption of "Reed logo", the label of "reedlogo", and specify that this is a figure. Make note of the different **R** chunk options that are given in the R Markdown file (not shown in the knitted document).

include graphics(path = "figure/reed.jpg")



Figure 3.1: Reed logo

Here is a reference to the Reed logo: Figure 3.1. Note the use of the fig: code

here. By naming the ${\bf R}$ chunk that contains the figure, we can then reference that figure later as done in the first sentence here. We can also specify the caption for the figure via the R chunk option fig.cap.

3.1. Figures 39

Below we will investigate how to save the output of an **R** plot and label it in a way similar to that done above. Recall the flights dataset from Chapter ??. (Note that we've shown a different way to reference a section or chapter here.) We will next explore a bar graph with the mean flight departure delays by airline from Portland for 2014.

Here is a reference to this image: Figure ??.

A table linking these carrier codes to airline names is available at https://github.com/ismayc/pnwflights14/blob/master/data/airlines.csv.

Next, we will explore the use of the out.extra chunk option, which can be used to shrink or expand an image loaded from a file by specifying "scale= ". Here we use the mathematical graph stored in the "subdivision.pdf" file.

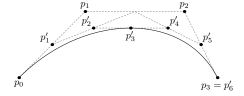


Figure 3.2: Subdiv. graph

Here is a reference to this image: Figure 3.2. Note that echo=FALSE is specified so that the **R** code is hidden in the document.

More Figure Stuff

Lastly, we will explore how to rotate and enlarge figures using the out.extra chunk option. (Currently this only works in the PDF version of the book.)

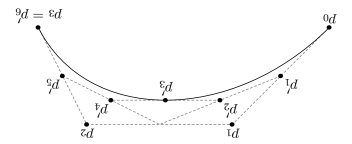


Figure 3.3: A Larger Figure, Flipped Upside Down

As another example, here is a reference: Figure 3.3.

3.2 Footnotes and Endnotes

You might want to footnote something. ¹ The footnote will be in a smaller font and placed appropriately. Endnotes work in much the same way. More information can be found about both on the CUS site or feel free to reach out to data@reed.edu.

3.3 Bibliographies

Of course you will need to cite things, and you will probably accumulate an armful of sources. There are a variety of tools available for creating a bibliography

¹footnote text

database (stored with the .bib extension). In addition to BibTeX suggested below, you may want to consider using the free and easy-to-use tool called Zotero. The Reed librarians have created Zotero documentation at https://libguides.reed.edu/citation/zotero. In addition, a tutorial is available from Middlebury College at https://sites.middlebury.edu/zoteromiddlebury/.

R Markdown uses pandoc (https://pandoc.org/) to build its bibliographies. One nice caveat of this is that you won't have to do a second compile to load in references as standard LaTeX requires. To cite references in your thesis (after creating your bibliography database), place the reference name inside square brackets and precede it by the "at" symbol. For example, here's a reference to a book about worrying: (Molina & Borkovec, 1994). This Molina1994 entry appears in a file called thesis.bib in the bib folder. This bibliography database file was created by a program called BibTeX. You can call this file something else if you like (look at the YAML header in the main .Rmd file) and, by default, is to placed in the bib folder.

For more information about BibTeX and bibliographies, see our CUS site (https://web.reed.edu/cis/help/latex/index.html)². There are three pages on this topic: bibtex (which talks about using BibTeX, at https://web.reed.edu/cis/help/latex/bibtex.html), bibtexstyles (about how to find and use the bibliography style that best suits your needs, at https://web.reed.edu/cis/help/latex/bibtexstyles.html) and bibman (which covers how to make and maintain a bibliography by hand, without BibTeX, at https://web.reed.edu/cis/help/latex/bibman.html). The last page will not be useful unless you have only a few sources.

If you look at the YAML header at the top of the main .Rmd file you can see that we can specify the style of the bibliography by referencing the appropriate csl file. You can download a variety of different style files at https://www.zotero.org/styles. Make sure to download the file into the csl folder.

Tips for Bibliographies

- Like with thesis formatting, the sooner you start compiling your bibliography for something as large as thesis, the better. Typing in source after source is mind-numbing enough; do you really want to do it for hours on end in late April? Think of it as procrastination.
- The cite key (a citation's label) needs to be unique from the other entries.

²Reed College (2007)

- When you have more than one author or editor, you need to separate each author's name by the word "and" e.g. Author = {Noble, Sam and Youngberg, Jessica},.
- Bibliographies made using BibTeX (whether manually or using a manager) accept LaTeX markup, so you can italicize and add symbols as necessary.
- To force capitalization in an article title or where all lowercase is generally used, bracket the capital letter in curly braces.
- You can add a Reed Thesis citation³ option. The best way to do this is to use the phdthesis type of citation, and use the optional "type" field to enter "Reed thesis" or "Undergraduate thesis."

3.4 Anything else?

If you'd like to see examples of other things in this template, please contact the Data @ Reed team (email data@reed.edu) with your suggestions. We love to see people using R Markdown for their theses, and are happy to help.

 $^{^{3}}$ Noble (2002)

Conclusion

If we don't want Conclusion to have a chapter number next to it, we can add the {-} attribute.

More info

And here's some other random info: the first paragraph after a chapter title or section head *shouldn't be* indented, because indents are to tell the reader that you're starting a new paragraph. Since that's obvious after a chapter or section title, proper typesetting doesn't add an indent there.

Appendix A

The First Appendix

This first appendix includes all of the R chunks of code that were hidden throughout the document (using the include = FALSE chunk tag) to help with readibility and/or setup.

In the main Rmd file

```
# This chunk ensures that the thesisdown package is
# installed and loaded. This thesisdown package includes
# the template files for the thesis.
if (!require(remotes)) {
 if (params$`Install needed packages for {thesisdown}`) {
    install.packages("remotes", repos = "https://cran.rstudio.com")
 } else {
   stop(
     paste('You need to run install.packages("remotes")",
            "first in the Console.')
   )
 }
if (!require(thesisdown)) {
 if (params$`Install needed packages for {thesisdown}`) {
   remotes::install_github("ismayc/thesisdown")
 } else {
   stop(
     paste(
        "You need to run",
```

In Chapter 3:

Appendix B

The Second Appendix, for Fun

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