PART I: Algorithms (mostly strings)

Mini-forward

These texts were written for "Introduction to Computational Biology and Biological Modeling" course I taught for 16 years at Penn. The text is somewhat wordy. I've put things that are technical or side discussions in boxes.

I wrote this mostly to collate the topics I wanted to cover and I apologize that I am missing explicit references to other papers or textbooks from which I sourced my material. It is inexcusable sloppiness but I haven't had the time to rewrite everything for more public distribution except for the students who take my course (who are also given a list of important books for reference). Some of the books and references I draw heavily from are:

Algorithms on strings, trees, and sequences: computer science and computational biology, 1997. Dan Gusfield, Cambridge Univ Press, New York, NY.

Computational Molecular Biology: An Algorithmic Approach, 2000. Pavel A Pevzner, MIT Press, Cambridge MA.

Biological sequence analysis: Probabilistic models of proteins and nucleic acids, 1998. Richard Durbin, Sean R. Eddy, Anders Korgh, Graeme Mitchison, Cambridge Univ Press, New York, NY.

Statistical Learning Theory, 1998. Vladmir N. Vapnik, Wiley, New York, NY.

If you happen to see this document in the wild, please accept my apologies and also if you see any errors, I would appreciate any corrections.

Junhyong Kim © 2018.

Unit 1: String search problems

Genomic data mostly consists of sequences of letters representing the nucleotides, A, C, G, and T for DNA (with U instead of T for RNA) and 20 letters of the amino-acids. Here are the codes for amino acids:

Amino Acid	Three letter code	Single letter code
Alanine	Ala	A
Arginine	Arg	R
Asparagine	Asn	N
Aspartic Acid	Asp	D
Cysteine	Cys	\mathbf{C}
Glutamic Acid	Glu	E
Glutamine	Gln	Q
Glycine	Gly	G
Histidine	His	H
Isoluecine	Ile	I
Leucine	Leu	L
Lysine	Lys	K
Methionine	Met	M
Phenylalanine	Phe	F
Proline	Pro	P
Serine	Ser	S
Theronine	Thr	T
Tryptophan	Tryp	W
Tyrosine	Tyr	Y
Valine	Val	V
Asn or Asp	Asx	В
Glu or Gln	Glx	Z
Leu or Ile	Xle	J
Unknown	Xaa	X

There are a few codes at the bottom of the above table for ambiguous amino-acids (only those that are commonly difficult to distinguish by chemical methods). There are ambiguity codes for nucleotides as well:

Code
R
Y
\mathbf{M}
K
S
W
Η
В
V
D
N

Typical genomic data is some string of amino-acid or nucleotides that is corresponds to biological units such as a "gene." This data is most commonly stored in a text file. The most common format for text files of nucleotides or amino-acids is called the FASTA format (invented by David Lipman and William Pearson for one of the first string analysis program packages). The format is simple. It has one line called the header that starts with the symbol ">". The header line has freeform information about the sequence that follows. The next set of lines contains the string of interest, which terminates either at the end of file or with the appearance of the next header line.

Here is the FASTA record (i.e., one string in FASTA format) for the Drosophila melanogaster olfactory receptor gene Or22a.

>gi|24580887|ref|NM_078729.2| Drosophila melanogaster odorant receptor 22a (Or22a), mRNA ${\tt TCATTTACTTGGATCGGGTGATGTGGTCCTTTGGCTGGACAGAGCCTGAAAACAAAAGGTGGATCCTTCC}$ TTATAAACTGTGGTTAGCGTTCGTGAACATAGTAATGCTCATCCTTCTGCCGATCTCGATAAGCATCGAG TACCTCCACCGATTTAAAACCTTCTCGGCGGGGGGGTTCCTTAGTTCCCTCGAGATTGGAGTCAACATGT ACGGAAGCTCTTTTTAAGTGCGCCTTCACCTTGATTGGATTCAAGAAAAGACAGGAAGCTAAGGTTTTTACT GGATCAGCTGGACAAGAGATGCCTTAGCGATAAGGAGAGGTCCACTGTTCATCGCTATGTCGCCATGGGA ${\tt AACTTTTTCGATATTTTGTATCACATTTTTTACTCCACCTTCGTGGTAATGAACTTCCCGTATTTTCTGC}$ TTGAGAGACGCCATGCTTGGCGCATGTACTTTCCATATATCGATTCCGACGAACAGTTTTACATCTCCAG ${\tt CATCGCCGAGTGTTTTCTGATGACGGAGGCCATCTACATGGATCTCTGTACGGACGTGTGTCCCTTGATC}$ TCCATGCTTATGGCTCGATGCCACATTAGCCTCCTGAAACAGCGACTGAGAAATCTCCGATCGAAGCCAG GAAGGACCGAAGATGAGTACTTGGAGGAGCTCACCGAGTGCATTCGGGATCATCGATTGCTATTGGACTA TGTTGACGCATTGCGACCCGTCTTTTCGGGAACCATTTTTGTGCAGTTCCTCCTGATCGGTACTGTACTG GGTCTCTCAATGATAAATCTAATGTTCTTCTCGACATTTTGGACTGGTGTCGCCACTTGCCTTTTTATGT TCGACGTGTCCATGGAGACGTTCCCCTTTTGCTATTTGTGCAACATGATTATCGATGACTGCCAGGAAAT GTCCAATTGCCTCTTTCAATCGGACTGGACCTCTGCCGATCGTCGCTACAAATCCACGTTGGTATACTTT ${\tt TGGCTATGGTGAAGCTGGCATTTTCTGTGGTTACGGTAATTAAGCAATTTAACTTGGCCGAAAGGTTTCA}$ ATATTATATTGCTGTACCCTAATAAATATTTA

One of the most basic things to do in handling genomic sequences is to look for particular strings of interest from a large collection, say the Genbank (http://www.ncbi.nlm.nih.gov/genbank/), which holds most of the genomic sequence information submitted by researchers all over the world. Or, perhaps look for a particular substring in a much larger string. We will call this the "Search Problem" or the "String Match Problem." There are many kinds of searches one might want to do, each with varying degree of difficulty. For each, we have to consider not only how to do it, but how to do it fast. A fundamental consideration in all genomics-type of problems is that the size of the problem is absolutely huge. Being able to solve a small problem can be very very different from having it work on, say the entire collection of webpages in the world. A mammalian genome has approximately 3x109 bases of DNA information or ~4 x 108 bytes. A typical search will have to work over many thousands of such genomes.

We can approximately classify the kinds of searches we want to do as:

- 1. Exact match
- 2. Inexact match
- 3. Inexact match allowing for rearrangements
- 4. Abstract patterns
- 5. Database dependent searches

An "Exact Match", is like what it sounds like. It is a search for an exact occurrence of a particular string. You can open up a MS Word document, for instance the class syllabus, and do a search for "Computational Biology". In this case, we may either want the first instance of the search string or perhaps all such instances. If you happened to type in "Cmputational Biology", you won't find anything. However, if you do a Google search for "Cmputational Biology", you will get a list of potential search terms that are close to "Cmputational Biology". Here, the Google algorithm carries out an "Inexact Match" search amongst the set of common search queries that it has stored. Inexact matching can have many different flavors. The simplest is where the difference between the target in the database and the query pattern is small and does not change the linear order of the symbols. A tool developed at NIH called BLAST does many types of inexact searches for databases maintained at NCBI (National Center for Biotechnology Information). However, once the difference becomes greater, the computational difficulty becomes much greater—for example, a google search for "Cmpttnl Blgy" will turn up some very different things.

More complex inexact match problems happen when we want matches to rearrangements of the query pattern such as permutations. For example, "Biology Computational" is a permutation of the original query. DNA strings can have such rearrangements due to experimental errors or through natural mutations. "Biology Computational" is an example of a circular permutation. (To obtain a circular permutation of a string "ABCDE" imagine we have a string with "ABCDE" written on it and then connect the string in a circle, then

break it randomly in some place.) Possible circular permutation of "ABCDE" includes strings like "EABCD", "CDEAB", etc. Circular permutations can be an important type of biological rearrangement because a fragment of DNA or RNA can circularize (e.g., plasmid) and then become linearized by a break at a different place. More complex rearrangements can be things like "ComputaBioltional logy" (transposition) or "lanoitatupmoC Biology" (inversion) or "ComppmoCutational Biology" (inverted duplication), etc. More commonly, we might want to search for **abstract patterns**; that is, patterns that we define by certain rules. For example, a simple kind of abstract pattern is one that allows multiple possible variations of a letter: "[C|K]omputational [B|V]iology" where by [C|K] we mean either "C" or "K". There are commonly used syntaxes to describe such abstract patterns. In the programming language Perl and the operating system LINUX/UNIX, one particular type of syntax constructions is called "Regular Expressions." We will learn more about them later, but you can get more information from the web. Some abstract pattern search can be quite complicated such as "palindromes of size 10." Many kinds of searches depend on the database; that is, the **search type is a function of the database** that you are searching through. Examples include: "Best Matching String", "Longest run of A's", "strings that appear more than 3 times", "the most frequent 5 letter word", etc. All of these search problems get progressively harder to solve and harder to carryout efficiently.

We now introduce some definitions and terminologies that will help make the discussion more precise:

Alphabet: A finite collection of symbols; e.g., {A, C, T, G}, {A-Z, 0-9}. A blank space, "", is also considered a symbol as well are other punctuation marks. Sometimes we will make a distinction between types of symbols, using terms like "terminal symbols" and "non-terminal symbols" (more later). Unless there is a potential for confusion, in this text, I will use the term "symbol" and "letter" interchangeably.

Words, strings: A consecutive set of symbols from the alphabet set. In general, I will put quotes around a string, but sometimes the quotes may be dropped.

k-tuple, k-mer: Typically the term "k-tuple" is used to denote an ordered set of k things; e.g., a vector of numbers (1, 2,1, 3) would be called 4-tuple. A k-mer is typically used to denote an ordered set of symbols; i.e., it more explicitly refers to composition of symbols. In much of computational biology, these two terms are used interchangeably.

Most of the time, we might use a variable to denote a letter (symbol), a word, or a string. For example:

We write, ST to denote string concatenations such that ST = "ACCCTA" and TS = "AACCCT". We also use a power notation, T^n , to denote multiple application of the concatenation operation so T^3 = "AAA". More uncommonly, the power notation might be applied to an alphabet. So, if $D = \{A, C, G, T\}$ is the alphabet set for DNA, then D^4 might be used to denote the set of all 4-mers from the DNA alphabet.

Substring: A substring of a string, say "ACCCT", is some string that is a contiguous subset of the string. For example, "CCC" is a substring of "ACCCT". However, "ACCT" is not a substring of "ACCCT" since it is not contiguous. As in many standard mathematical definitions, the string is a substring of itself. That is "ACCCT" is a substring of "ACCCT".

Subsequence: A subsequence of a string, say "ACCCT", is any consecutive subset of the original string. For example, "ACCT" is a subsequence, but "CACCT" or "ACCTT" is not a subsequence. If a string is a subsequence of another string, we are able to align the two strings such that there is a correspondence between the symbols along with "gap symbols" in the substring for any missing letters. For example,

[&]quot;ACCCT"

[&]quot;ACC-T"

```
or
"ACCCT"
or
"ACCCT"
"---CT"
```

Given a string S = ``ACCCT'' and so on, we will use S[1], S[2], etc. to denote the 1^{st} , 2^{nd} , symbol, etc. We will also use the notation S[1...3] or S[2..4] to denote the substring that goes from 1^{st} to 3^{rd} symbol or the 2^{nd} to 4^{th} symbol and so on.

Prefix and suffix: Given some string S of length n, a prefix is any substring that has the form S[1...k] for and $k \le n$, and a suffix is any substring that has the form S[k..n] for any $k \le n$.

Kleene star: There is a special notation called the Kleene star that represents the set of all strings that can be generated from some finite set of alphabet or strings by (countably) consecutive application of all possible concatenations. For example, if $S = \{A, T\}$ then $S^* = \{A, AA, AT, TA, AAT, ATAT,\}$.

There is a special term called "language", the consequence of which we will see later. Here is the definition.

Language: If A is some alphabet and A* is the Kleene star of A, then a language is some subset of A*. Different languages of the same alphabet are different subsets of A*.

This definition of a "language" is not exactly what we usually think of when we think of natural languages. The term comes from the study of formal languages (as opposed to natural languages) that we will discuss later. However, the term is motivated by natural languages. Consider the standard alphabet set used in English along with punctuation marks and maybe some non-standard symbols like \hat{e} . Let $\mathbf{A} = \{A, B, C ...\}$ be this alphabet set, then A* represents all the possible (including nonsense) words and sentences that can be composed from this alphabet. English is a particular subset of A* that corresponds to the usual kinds of English words and sentences while French is a different subset of **A***. While this seems a bit overly abstract, precise notions like this can help G, T} be the alphabet for DNA then **D*** is the set of all possible strings comprised of the four nucleotides. We know that things like "AAA...A" (10,000 A's) are not genes as far as we know, while say 10 A's in a row might be part of a gene. So a gene is a subset of **D***--we could even call it a *Language of The Gene*. The question is whether we might know which subset of **D*** belongs to the *Language of The Gene* and which do not belong. How do we know whether a particular string of letters do or do not belong in English? While not always exact, we think of a string of letters as being part of the English language if they conform to certain rules of grammar. One might think it would be nice if there were a grammar for the Language of The Gene. We will consider many of these issues later.

Unit 2: Exact string search

Now, back to the search problem. We will first tackle the exact search problem. Here is the problem:

Exact Search Problem: Let S be some string and P another string. If P is a substring of S, return the offset of the first match position. That is, if S is a string of length n and P is a string of length m and S[k+1..k+m] = P[1..m] (and $k+m \le n$) then return k, where k is the smallest such number.

In real applications S might be a very large collection, sometimes an entire database such as the Genbank. (Note that we are returning k and the pattern we want starts from the k+1 position; therefore, k is called the offset. We should also note that whether we return the offset or the starting position doesn't really matter. There is just a convention to return the offset.)

It doesn't take too much thought to see that there is a simple algorithm to solve this problem. Here is a pseudo-code. Pseudo code contains statements that outline the steps of an algorithm but typically do not correspond to an actual programming language.

```
i = 0
while (i <= n-m) {
    k = 1
    result = 1
    while (k <= m) {
        if(S[i+k] <> P[k]) then result = 0
        increment k
    }
    if result = 1 then return i
    increment i
}
return "not found"
Fig 1: Naïve Search
```

If you have never programmed before, this pseudo code will need a bit of explaining. We first set an index variable called \mathbf{i} to zero. This index will go from zero to near the end of the S string and refer to positions within the string S. In fact, it will go up to the offset point of the last possible place that the string P can be found in S. For example, if S = ``ACCT'' and P = ``CT'', then n = 4, m = 2, and \mathbf{i} will run from 0 to 2. The part of the code that starts with "while (X) { }" says we should repeat everything within the brackets "{ }" as long as the condition \mathbf{X} is true. These kinds of constructions are called a "loop"; that is, keep looping through statements inside the brackets as long as the condition \mathbf{X} is true.

We next set another index **k** to 1 and use it to refer to the positions within the string P. We also set another variable that we will call **result** to the value 1. This variable is what we call a "flag". The flag variable will tell us after some calculations whether we have found a match to the string P. The second **while** loop now runs from the first position of string P to the end of the string P and tests if the symbols of P are different from the symbols of the string S, starting from the **i+k** position of S. If anywhere along the string P, we find a mismatch with S, we set the flag, **result = 0**. Therefore, after we do this loop, **result** variable will stay at the value 1 if there is a substring in S that matches P, but if there is any symbol that doesn't match it gets set to a value of 0. If **result = 1** after we go through the inner **while** { } loop, we will have found the first match and we return the current value of **i**, which is the offset. Stating "return the value of **i**" also implies that we stop all calculations as soon as this happens—this is called "breaking out" of the various loops. Otherwise, we increase the index **i** by 1 and start again. If we never breakout until we reach **i > n-m**, then this means that the string P does not exist and we might return something like "not found"—in practice, we might return a negative value to represent the case where the string is not found.

All of this is a rather longwinded way of saying, "start from the beginning of string S and see if we can match to string P, if not, move one position forward and then try again." Trying to be precise can seem tedious until we get used to the programming language (or pseudo code). Hang on.

Like real world problems, computational problems have multiple possible solutions. For example, we could loop backward instead of forward. Sometimes, seemingly trivial changes in how we approach a problem can make a large difference, as we will see. Sometimes, seemingly trivial problems such as multiplying two numbers can require complex strategies to do it well (look up this problem in computer algorithm books). What is "doing it well" in computation? If we consider the code above, one simple trivial change is to consider stopping the inner

loop as soon as we find a mismatch (i.e., breakout of the inner loop). It seems that would allow faster solution to the problem and therefore it would be solving it better. The foremost concern in solving a computing problem well is to minimize the use of resources. Two kinds of resources stand out in computing: **memory** and **time**. For example, in the algorithm above it is clear that we need memory to store S and P strings and to be able to access it, essentially, randomly. By the last statement, I mean that when we ask for the value of S[k] some mechanism exists to retrieve that information more or less instantly. We call such mechanism of storage, random access memory, or RAM. One can imagine that if S is very large, say the size of Genbank with some 400 billion bytes, such facile storage and retrieval might not be possible.

What about time? As you already know, time to compute something depends on a lot of different things such as the machine, how the machine does things, etc. Running the same computer game on a Mac, iPad, iPhone, all result in play at different speed. To analyze the amount of time required for a particular program can be a very tricky affair—and actually important. For example, recently I wanted to know how much time it would take to analyze some next-generation sequencing data (more on this later). This is a very large dataset, typically on the order of several hundred GB or so of input/output data. The program also runs on multiple processors. Because it reads and writes many files simultaneously from multiple processors it is fiendishly hard to know how much time the program will take. Actual time depends on the particular dataset, the memory state of various CPUs, the network interconnects (because the storage system is connected by a network), what disks were being accessed, and so on. I especially wanted to benchmark the time on cloud computing facilities. The virtual machines for cloud computing can have very variable actual hardware environments depending on when you access the machines, so it was doubly hard to figure out the computing time. In cases like this, the only feasible approach is an empirical approach; that is, try running the program a large number of times over a suitable number of test datasets and come up with a statistical characterization.

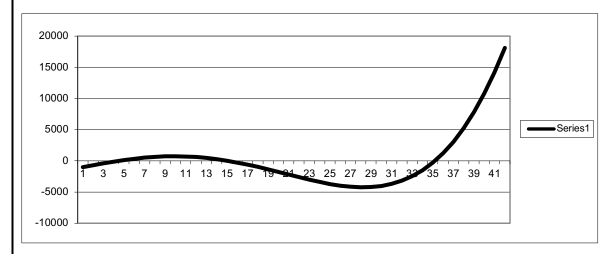
Any real-world program should be benchmarked by an empirical approach no matter what a theoretical analysis (to be learned next) tells you. However, there are many pitfalls to watch out for as well. For example, running the program may be costly—processing a single next generation sequencing dataset can take ~1,000 CPU hours. This means we cannot do a whole lot of empirical testing. A typical way around this is to process smaller problems (say smaller number of sequences) and extrapolate up to a realistic size. However, such extrapolation may breakdown in many ways. A machine may process 200 GB datasets with no problems and then suddenly take 10 times as long for a 500 GB dataset because of various storage bottlenecks. Processing a single data set might take 100 sec while processing 10 data sets may take an hour because of shared memory collisions, network delays, and so on. Careful empirical performance testing can be a whole enterprise and a research project. Regardless, all evaluations of an algorithmic approach should balance both empirical and theoretical analysis. This brings up the point for me the remind you that in any modeling or analysis we do, there is always a hierarchy of models/phenomena from low level to high level. For example, in a computing problem we have the low level to high level layers of micro-code driving the operation of a CPU, to assembly language, the higher programming language, code written in the programming language, to abstract description of the algorithm. In each case one might think that the lower level constrains what is possible to the higher level while higher level imposes a higher-order logic to the lower level. For example, the available micro-code and machine implementation constrains the efficiency of the code written in higher level language like Java while the code you write in Java imposes a logical flow to the execution of the micro-code.

Box: Asymptotic rate of function growth

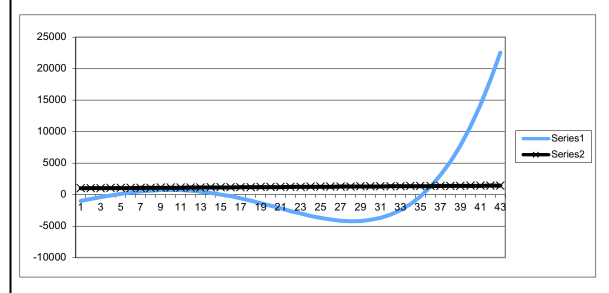
In discussing things like how much computing time is needed as the size of the input problem grows, we are typically not so concerned about details but kind of a rough estimate. To be more exact about "rough estimates", we introduce the notion of asymptotic bounds on some function. Suppose we have a complicated function say like this:

$$f(n) = -1000 + 300n + 2n^2 - 2n^3 + 0.05n^4$$

The curve for this graph up to n = 42 looks like this:

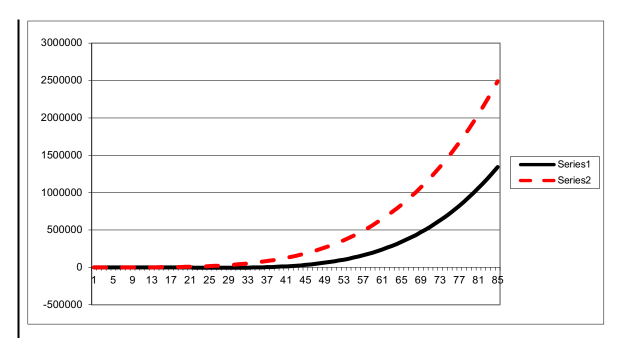


The main feature is that this graph goes up for a while and then goes down and then goes up again. Now let's compare it against a graph of g(n) = 1000 + 10n:



The g(n) graph is shown above marked by "x" symbols. What we notice is that g(n) is bigger than f(n) up to some point around 35 and then f(n) is substantially bigger. In fact, with a little bit of math we can show that after n = 35, f(n) is <u>always bigger</u>. Now suppose that g(n) and f(n) describes the time complexity of two algorithms for a n-sized problem. At small problem sizes, the algorithm with g(n) time complexity takes longer but once the problem size become bigger than a critical amount, the algorithm with f(n) time complexity takes much longer. Since we usually care about the general behavior of the system, it is useful to characterize the behavior of computation ignoring the irregular and potentially complicated behavior at the small sizes. So, this motivates us to only consider the behavior of the system when n > crit, where crit denotes some number.

Now consider the function $h(n) = 0.05n^4$. This function is similar to f(n) except for the lower order terms. The following shows the graph of h(n) (red) and f(n) (solid):



Now that we are plotting out to n = 85 the numbers are so big that we don't notice the dips and valleys in f(n) anymore. The two graphs look very similar except for a little displacement. In fact, with a little bit more math we can show that the values of the two functions never differ by more than a constant factor, c. And, the graph of $h(n) = 0.05n^4$ never differs from the graph of $p(n) = n^4$ by more than a constant factor. Since $p(n) = n^4$ looks very simple, it would be nice to say that h(n) is pretty much like p(n) and since f(n) is pretty much like h(n), we would also like to say that f(n) is characterized by p(n).

All of the above discussion motivates the following definition:

We say $f(n) = O(n^4)$ if once n is greater than some number crit, $f(n) \le cn^4$ (where c is some constant factor). That is, we can always say the f(n) grows at a rate that is less than a constant factor of n^4 . Or, more informally, f(n) is $\underline{bounded\ above}$ by n^4 . We can, of course, bound f(n) from below. For example, f(n) is bounded from below by n^3 , by which we mean f(n) will grow $\underline{at\ least\ as\ fast}$ as a constant factor of n^3 . The function f(n) is also bounded from below by n^4 as well as by n^3 . (And, if you think about it, bounded from below by n^2 and n.) If f(n) is bounded above and bounded below by the same function, n^4 in this case, then we say f(n) has the same growth rate as n^4 . Typically, we want to know the worst-case scenario so we are interested in the growth that is bounded from above. That is, we want to know that $f(n) = O(n^4)$; that way we know that the performance of something described by f(n) will never be worse than n^4 . But, often times, these bounds are also exact—that is, indicates the same growth rate.

In all of our discussion above, I used a concrete example, but as you should be aware of now, the $O(\cdot)$ notation is used for any general function. Thus,

$$f(n) = O(q(n))$$

means the function f(n) is bounded above by the function q(n). We use the notation $\Omega(f(n))$ for bounded below and $\Theta(f(n))$ for bounded above and bounded below by f(n).

Going back to the pseudo-code in Fig 1, we can now carry out a theoretical analysis of the computing time required to run this program. There are two **while** { } loops in the code, one nested inside the other. Therefore, it is clear that the statements inside the second **while** { } loop will be repeated the most number of times. We might reasonably think that the total time for the program will be generally determined by how many times those inner loop statements are carried out. It is easy to see that the two statements, $if(S[i+k] \iff P[k])$ then **result = 0; increment k,** will be repeated (n-m+1)m times. Or, if n is much bigger than m, then $n-m+1 \sim n$ so the inner loop statements will be repeated about nm times. The whole reason to write a program is to solve the same problem multiple times on different inputs. The main important characteristic of the input for the pseudocode in Fig 1, is the length of the two strings, n and m. We usually want to know how much time a given algorithm will take as a function of the characteristics of the input. Here, our analysis tells us that we expect the computing time to be (most likely) a function of n times m. In actual program execution, there are other statements besides the ones in the inner loop that need to be processed. For example, locating the data (the two strings), reading the strings into random access memory, etc. However, as n and m become larger and larger, these other steps will contribute less and less to computing time (with some possible exceptions as we discuss below). How an algorithm's resource use scales with the size of the input problem is called the *computational complexity* of the algorithm. (Or, the computational complexity of the <u>problem</u>; we will discuss the difference between a "problem" and an "algorithm" later.) Typically, we are interested in the asymptotic complexity—that is, what happens to time use or memory use when the input size becomes pretty large. The pseudo-code in Fig 1 has time complexity O(nm) and memory complexity O(n+m), where the notation O(.) means "asymptotic upper bound" (see boxed text above).

We are still not quite done thinking about Naïve Search pseudo-code. We assumed that when we ask for "ith symbol of string S" by S[i], we will have instantaneous access to that information (RAM). What if such memory cannot be used? Suppose all the information is recorded on a physical tape and each time we need the ith information, we need to physically move the tape to the ith position from the beginning? Then the time complexity bound of O(nm) doesn't tell us the right picture. What about the index variable i? Will our computing machines behave the same if $\mathbf{i} = \mathbf{0}$ versus $\mathbf{i} = \mathbf{300}$ billion? Will they properly handle index values like 2 trillion, a number that might occur in genomics data? Most modern computers provide complicated architecture to allow access to memory in an arbitrary (i.e., random) manner and also to provide variables that can count large numbers. But, there is a limit, sometimes a fairly important limit. Machine implementation of computing involve finite modules that carryout finite operations. Things are fine while the problem size is within these finite limits but if the problem size gets bigger, much more machine operations have to be carried out—sometimes very much more. Available memory is often structured such that a certain amount of memory can be accessed very fast, others not so fast (e.g., placed on hard disks, whose recall/access can be very slow). In addition, how a given programming language handles certain kinds of data structure may be very different. Previously, we postulated that the main determinant of the Naïve Search algorithm speed might be how many times the inner loop statements are executed. Suppose S is a very long string, say n = 500 billion. If we naively try to read this into a single large string variable, simply reading the data may take much longer than computing the statements in the inner loop. Scripting programming languages, such as Perl, R, and Python, tend to have difficulty with very large memory use. One thing we can see from the Naïve Search pseudo-code is that for at least one of the strings, we don't really need to access the memory in a random manner. That is, string S is examined from left to right (index value θ to n-m) without ever going back; and, at the time of use, we only need to access m number of positions (the length of P). Therefore, we should be able to implement this algorithm with memory complexity O(min(n,m)); that is, we only need RAM in proportion to the shorter of the two strings. Small memory usage may allow some algorithms to be much faster than other algorithms even if theoretically they have identical asymptotic time complexity.

With all that we are now ready to see if we can improve upon Naïve Search algorithm. The main thing we want to do is to see if we can reduce the number of comparisons. The naïve part of Naïve Search algorithm is in

comparing against string P and then moving one step along string S and doing the same thing all over again. It seems we should be able to avoid all this. We already considered breaking out of the loop as soon as we find a mismatch. We now use a seemingly trivial but a very important change, which turns out to have a very big effect. Inside the inner loop of the Naïve Search algorithm, we scan the string P left to right against left to right positions of string S, starting from the offset \mathbf{i} . What if we start the scan from right to left? Suppose at the very first right most symbol of P, we found it did not match the corresponding position in S, call this position S[k]. We can immediately move the string P rightward (i.e., increase the offset) all the way to a position within P that we do know matches to the symbol in S[k]. Here is an example:

```
123456789
S: "AAACTTGTT"
P: "TAC"
```

String P is currently at offset 3. Scanning from the right we see the symbol "C" in string P does not match the symbol "T" at S[6]. This tells us we can immediately advance string P like this:

```
123456789
S: "AAACTTGTT"
P: "TAC"
```

Now, string P has moved to offset 5 from offset 3, jumping over evaluating offset 4. (Why are we able to do this?) Thus, we've saved a certain amount of redundant comparisons. Why not jump to offset 6; i.e., a full frame of the string P? Since the "T" in P[1] does match S[6], the position we evaluated, we have to make sure no extension from that position results in the full match of the string P.

This seemingly simple but great idea is small part of the core idea in the famous Boyer-Moore algorithm, developed by Bob Boyer and J. Strother Moore in 1977. As we noted before, a mismatch in the string P tells us to stop processing any more of the string. The key idea in Boyer-Moore algorithm is that this also tells us something about the string S and how to skip some of the processing of string S. Going from right to left, especially gives us the information about the maximum amount of skipping we can do in string S. We will do the full Boyer-Moore algorithm later but for now, here is the pseudo-code for a simpler version:

Fig 2

Many things have happened in the pseudo-code for Fig 2 compared to Fig 1. First, as can be seen the inner loop starts from the end of string P(k = m) and moves left (decreases the index k). Second, inside the inner loop is the statement

increment i by bad_character_skip(S[i+k], k)

Here, there is a function called **bad_character_skip()** that has not been defined yet. It takes in two values, the symbol at S[i+k]—this is the position that has been found to not match the current P[k] symbol, and the value of k—the current position within the string P. This function will tell us how much to change the offset so that the string P will be shifted to the right, either by the full string, or until a position within P that matches the symbol S[i+k]. If this seems confusing look back at the example above. We assume this function will be pre-computed. The pre-computation will require a table of size $p \times m$, where p is the number of symbols in the alphabet. I note here that if the number of symbols is large (like the English alphabet and all the symbols one might find in a MS Word document) and the search string is relatively long, then this table can be pretty unwieldy. (The original version of Boyer-Moore algorithm uses a much smaller table, discussed below.) Since the number of symbols in DNA or amino-acids is manageable, we can easily consider a $p \times m$ table. Here is an example for the search string "AAACGT", where the rows index the mismatched character at S[i+k] and the columns index k, the position within P where the right most mismatch occurred:

$S[i+k] \setminus k$	1	2	3	4	5	6	
A	0	0	0	1	2	3	
\mathbf{C}	1	2	3	0	1	2	
G	1	2	3	4	0	1	
T	1	2	3	4	5	0	

You should convince yourself that this table can be computed only from the string P and does not require information from string S. Pre-computing this table means that we will assume when we need the value of **bad_character_skip()**, we will get it instantly without additional computing. Of course, we need to store these pre-computed values, so we will need to use $p \times m$ RAM memory to achieve this instant access.

In design of algorithms there are many ways to tradeoff memory use for computational time. In learning to multiply multi-digit numbers in grade school, we were made to memorize the single digit multiplication table. Then we were taught an algorithm to apply the memorized multiplication table (what we call a "lookup" table) position by position. If we had memorized a larger table, say all two digit multiplications, we would be able to do longer multiplications much faster. If we didn't memorize our multiplication tables, we would have to do a lot of sums for even single digit multiplications and it would take forever. In fact, the architecture of a computer's CPU employs a lot of such lookup tables to judiciously speed up certain operations while trying not to blow up the complexity of the CPU.

The third new thing in the pseudo-code in Fig 2 is the statement "go to line A:". It means to move the execution of the program to the line that is labeled "line A: $\mathbf{k} = \mathbf{m}$ ". That is, break out of the inner loop and start over again at "line A" with the new value of the index i. (Yes, sometimes when we are working with a limited programming language, a "go to" statement can be useful.)

Okay, what do we expect from this new algorithm? A moment of thought will show that it depends on the contents of the string S and P. Suppose S = "AAAAAA....A"; say, a 100 A's and P = "TTTT". As soon as we compare the right most T, we will shift the entire string P by 4 positions. So, we will end up doing $\sim 100/4$ symbol comparisons. If P = "TTTTTTTT", we will shift the string P by 8 positions and end up doing $\sim 100/8$ comparisons, etc. This is great! In this best case scenario the time complexity of this algorithm is O(n/m). This is a key feature of Boyer-Moore algorithm: the search can be faster the longer the search string! Now suppose P = "TAAA". Then we will not know that P does not match the first 4 symbols of S, until we get to the very last (left

12

most) symbol of P. Then we shift by one. So, in this worst case scenario we are right back to doing O(nm) comparisons. What do we expect if both strings S and P are uniformly random strings of A, C, G, and T? I leave that for you to solve (hint, try with the simplest case first where we assume the right most symbol in P immediately mismatches; how many positions will we skip on average?)

Thinking about this problem further, it seems like we should be able to do better than O(nm), even in the worst case. If P = ``TAAA'' here is what we would have compared at offset = 0:

123456 S: AAAAA..A P: TAAA

Starting from P[4], we have to count down to P[1] to know that we have a mismatch. The algorithm above, then shifts S by +1 and starts at P[4] aligned to S[5] and counts down again. However, when we are doing this we are throwing away information we already know. That is, we already know that S[2-4] matched P[2-4], and none of these were the symbol "T". So, our intuition is that we should already have the information to skip by +4 positions. In fact, this is true.

The full Boyer-Moore algorithm tries to match the search pattern, P, from some offset position in the database S, moving from right to left. When a mismatch has been found, we know we can increase the offset, effectively moving the pattern P in the right hand direction against S (or, equivalently moving S left against P). Suppose the length of P is m and the mismatch happens k positions left of P[m]; i.e., at P[m-k]. Since we know all the symbols of P, we know the corresponding letters of S from P[m-k+1] to P[m]. Thus, we know how to best shift P. The idea of a **suffix match table** is to use this information.

To make things concrete, let's start with an example:

123456789 S:TATTCGGTT P:GCGACG

The mismatch has happened in the P[4]-S[4] position. But, we already know that P[5,6] = "CG" has matched and therefore the corresponding substring in S is "^ACG" where I used the symbol "^A" to denote "not A". (Actually, when we are actively scanning we know that the precise corresponding substring in S is "TCG"; we will see below how that information can be used to skip even further.) Looking at the internal structure of P by eye, we know there is another "CG" left of the first "CG"; therefore, we should be able to increase the offset until that "CG" aligns with the "CG" we already know is part of S:

123456789 S:TATTCGGTT P: GCGACG

At this point, we start at S[9] and P[6] again to try matching. The key is that we can compute these offsets by an examination of the internal structure of P. Basically, we are looking for the repeat structure of P. We want to enumerate all the possible repeat structures. I will first show a suffix match table and then we can discuss how this is constructed. We enumerate the mismatch cases from right to left of the string P, and then what should happen when we find a mismatch:

Position of first mismatch of	Implied pattern in S	Action to be taken
P, scanning from right to left		
P[6]	^G	increase offset by 1 (i.e., move P right by 1)
P[5]	^CG	increase offset by 5
P[4]	^ACG	increase offset by 3
P[3]	^GACG	increase offset by 5
P[2]	^CGACG	increase offset by 5
P[1]	^GCGACG	increase offset by 6

How do we know what action to take? The basic idea is to find out if there is a pattern starting left of the current mismatch position that matches the pattern in S (the first column of above table). For example, we might have:

```
S:...TCG
P:GCGACG
```

To the left of "ACG" in P, we have "GCG", which matches "^ACG" and is at 3 positions left of the current position (thus, increase offset to 3).

Wait, why are we not looking for "TCG", which we know is S[5-6]? Because, we want to compute these offsets by only looking at P without consideration of the mismatch symbol in S. We will see later, that we can use the information that the mismatch symbol was T and not {C, G, T} in a different manner.

If there is no match left of the current evaluation point, there still could be some prefix left of the current point matches the suffix right of the current point. That is, we can slide the string P right up to the point until it overlaps with known parts of S (represented by suffixes right of mismatch point).

A better way to summarize the two cases we just discussed is to consider the pattern at the point of mismatch. We put down the original P string (not S) and then slide the mismatch pattern to the left, until we get a match. When we are doing that, we assume that the original string is padded to the left by "wild card" symbols (i.e., symbols that match any symbol). Like this:

```
P:....GCGACG
0: ^ACG
1: ^ACG
2: ^ACG
3: ^ACG
```

This tells us that there is an internal repeat of A CG pattern +3 offset from the current mismatch. We try this with the case that we find a mismatch at P[2]; i.e., C CGACG

```
P:.....GCGACG

0: ^CGACG

1: ^CGACG

2: ^CGACG

3: ^CGACG

4: ^CGACG

5: ^CGACG
```

So the calculated offset is +5.

To fix ideas, we do one more example with P = "TAAAA".

Position of first mismatch of P,	Implied pattern in S	Action to be taken
scanning from right to left		
P[5]	^A	increase offset by 4
P[4]	^AA	increase offset by 3
P[3]	^AAA	increase offset by 2
P[2]	^AAAA	increase offset by 1
P[1]	^TAAAA	increase offset by 5

The suffix match table has an entry for each of the possible cases where the mismatch was found in m, m-1, m-2,...,I positions of the string P, where m is the length of P. The way I've described the construction of the suffix match table, you would have to do a search through the string P for each of those mismatch cases and the algorithm might take something like $O(m^2)$ time to construct this table. Can you think of how to do this in less time?

Okay, now one more part. In the modified Boyer-Moore that I gave in Fig 2, we computed a fairly large bad character table. Here, we compute a simplified table, only with respect to the last (right most) symbol in P. Basically, we want to compute for each possible symbol the right most place it can be found in P. For example, with P = ``GCGACG'', then we have: A = 4, C = 5, G = 6, D = 0. Suppose D = 0 Suppose D = 0 at some point in the algorithm, then we are supposed to look up D = 0 in the bad character table. Then we compute

```
max(1, j - bad\_char\_tab(S/i)),
```

where bad_char_tab(S[i]) means look up the value in the table. Note that depending on where you are j bad_char_shift(S[j]) can be a negative number. If it is, we set the offset shift to 1.

We will call the two strategies the "bad character rule" and the "good suffix rule". Finally, the Boyer-Moore algorithm in words:

Match P to S, right to left, from some offset position in S. If a mismatch is found, compute *shift* = max(bad character rule, good suffix rule). Increase offset by *shift* and start matching P to S at the new offset position, right to left.

The Boyer-Moore algorithm is the gold standard in exact string searches. There are other cases, which we will discuss in a moment, where we might want to use a different strategy but for straightforward single string search, this would be the algorithm of choice. If we go back to the examples we discussed before, say looking for "GTTTT" in a string of 100 A's, you can see that the bad character rule will be evoked immediately and we will evaluate at 20 positions and we will find that the string does not exist. If we were looking for "TAAAA" in a string of 100 A's, the suffix match rule will be evoked and again and this time we will evaluate no more than 100 positions and conclude the string does not exist. However, it turns out there can be some pathological cases. Suppose we were looking for "AAAAA" in a string of 100 A's and want to find all of them. We find a match at the first position, but none of the rules allow us to move more than 1 position. So we end up doing O(mn) comparisons again. There are additional fixes to this problem. It turns out that with all the trimmings, the Boyer-Moore type of strategy will have the worst case time complexity of O(n).

The pathological example just discussed points to a very important idea when coming up with solutions to problems—whether the problem is algorithmic, statistical, or mathematical in nature. We have to always try hard to consider the extremes of the input cases: the best case, the worst case, etc. What we want is a solution that behaves nicely in some sense over all possible cases. For example, we might want the solution to behave nicely even in the worst possible case, or, behave nicely on the average. Or, to behave in a smooth predictable way from the best to the worst so that we know when it is starting to behave badly.

15

A final comment on the Boyer-Moore family of algorithms. As mentioned, the algorithm has a very nice property that it tends to become more efficient (on average) when the length of the string pattern sought is longer—this is a very desirable thing indeed. However, the gain in efficiency tends to be lower when the size of the alphabet is small. As you can surmise from the algorithm, when we have a mismatch, it is best when there is no pattern or symbol in rest of the string P. Larger the alphabet size, more likely this will be. Small alphabets like DNA tend to have a lot of possible matches left of the mismatch position, then the size of the offset shift tends to be small, losing efficiency. Thus, our work is not done.

Unit 3: String search by structured data

We now consider the case where we have some common fixed database of strings like Genbank and we would like to search the database over and over again. In these cases, we might try to organize our database in some way such that multiple queries can be efficient. Organizing the database might take a lot of effort, but if we are going to carry out millions of searches on it, it might be worth the effort. The canonical example of such organization is your standard dictionary or encyclopedia. Effort has been put into the dictionary/encyclopedia to organize the entries in an alphabetical order so future searches can be efficient. In fact, the usual dictionary/encyclopedia also has "pointers" to sections so you can cross-reference from one section to another. Organizing information into particular structures that facilitate access is called (making) "Data Structures". There is a large volume of information on data structures in computer science and the data structures are intimately tied to algorithms. I should also note that many data structures can be theoretically equivalent but can have a large impact on performance in actual programming code, especially as a function of the programming language.

Let's start with some simple examples of data structures. You can study the subject in detail in many computer science courses. We start with unstructured data: {0, 2, 4, 3}. This is a set of four numbers and you will remember from math that a set only "contains" the objects and there is no meaning to the order. If I wanted to know whether the number 4 is a member of the set, I would have to search through the set—somehow. An "array" is the most important kind of data structure. It is equivalent to taking the unstructured set and giving it an order (making it a k-tuple): (0, 2, 4, 3). (Recall in mathematical notation a set is denoted with {} brackets while an ordered set is denoted with () brackets.) Now, the 3rd item in the 4-tuple is the information "4." The tuple can be given a name (i.e., a variable) so we might write N = (0, 2, 4, 3). Then N[1] = 0, N[2] = 2, etc. Programming languages implement this in various notations (syntax). In Perl we would write @N = (0, 2, 4, 3)and then retrieve the information using the syntax N[1] = 2. (Unfortunately, Perl starts array indices from 0 so this example is shifted from my preferred convention). The key idea of an array is that the locations of the objects are indexed by natural numbers so we can use arithmetic operations to retrieve information. For example, we can use a variable for natural numbers, e.g., \$i (continuing with Perl syntax, where the variable names are preceded with the symbol "\$"). If we set \$i = 0, and then retrieve \$N[\$i], we get the value 0 from the array @N. (In Perl, array variables are preceded by the symbol "@" when referring to the whole array and preceded by the symbol "\$" when picking out individual members of the array.) Now we can carry out arithmetic operations on \$i, say, add one such that \$i = 1 then \$N[\$i] = 2. If you have never programmed before, this is VERY IMPORTANT. The same symbolic notation "\$N[\$i]" retrieves different pieces of information dependent on value of \$i. If we weren't able to use such higher-order notation (and data structures), programs would be extremely unwieldy.

An array assumes that there is a machine-implemented method to access the ith information. Imagine that your information is written on a stack of index card and I wanted the 98th card out of 500. You can see this might not be so easy without some special provision. A simpler kind of data structure is where the information has order but it can only be accessed from the "top" or the "bottom" or both. A "**stack**" is a data structure that is just like a stack of index cards. You can store information by putting a new card on the top and retrieve information by taking off and reading the top card. Thus, a stack has the characteristic of LAST-IN-FIRST-OUT (LIFO); that

16

is, the last piece of information in the stack is the first piece of information that can be retrieved. When you retrieve the information, it is also lost from the stack. Putting information into the stack is called "push" and retrieving information is called "pop" (e.g., popping from stack X). A "queue" has a little bit more so that you can access the last item in the order. Information is again pushed in (or sometimes called "enqueue") and when the information is popped (also called "dequeue"), the first piece of information that was stored is retrieved. That is, we get FIRST-IN-FIRST-OUT (FIFO). It may seem like these are very primitive structures (and they are), but both the stack and queue are used often in programming (and in the CPU machine operations), typically for storing a list of things to do.

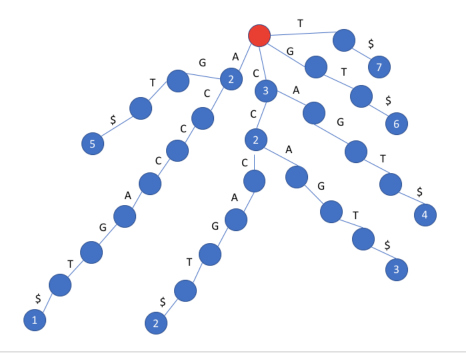
As mentioned, there are many different kinds of data structures used in algorithms and we will continue with two additional examples that are commonly used in computational biology. While an array allows us to retrieve information using natural numbers and arithmetic operations, many kinds of information are not naturally organized into numerical order. However, we often have information that come in pairs, say FIRST NAME + LAST NAME; or, say GENE NAME + DNA SEQUENCE. We can represent this as a set of 2-tuples: for example, {(A, 0), (C, 2), (G, 4), (T, 3)}. An associative array gives us a way to store such paired information. We assume that each 2-tuple is the pair (Key, Value) so that if we denote %N = (A, 0, C, 2, G, 4, T, 3) (here, Perl denotes associative array with variable names that is preceded by the symbol "%" and each pair of objects in the list are assumed to be Key, Value pairs), then \$N{'C'} retrieves the information value "2". (Perl uses the syntax \$N{} } to access elements of an associative array.) Associative arrays are often used to store information that is commonly referenced by strings of letters rather than natural numbers. Examples include (Gene Name, DNA string), (DNA string, Gene expression), (DNA string, occurrence frequency), etc. At first glance, this seems like the way to go. Just put all of our information in (Key, Value) format—but, we will see later that this requires a lot of additional computing and difficulties.

Arrays, stacks, and queues store information in a one-dimensional linear order but efficient retrieval of information is often aided by multi-dimensional structure or a more general structure. In particular, storing information in a tree graph structure is very useful in a wide variety of algorithms. We first briefly discuss graphs and trees as a discrete object. A graph consists of two sets, a set of **vertices**, V, and a set of **edges**, E, such that we write $G = \{V, E\}$. Edges connect the vertices to each other. So, if the set of vertices are $V = \{v_1, v_2, \dots v_n\}$ then we can depict an edge by the pair $e = \{v_b \ v_i\}$; that is, the edge e connects the two vertices v_i and v_i . An edge may be directed in the sense that the edge $e = (v_i, v_i)$ now depicts connections from v_i to v_i but not the reverse connection $(v_i \text{ to } v_i)$. If a graph consists of directed edges than we call it a **di-graph**. Each edge connects only two vertices but each vertex may have many edges. The number of edges of a vertex is called its **degree**. If it is a digraph, then a vertex will have in-degree and out-degree. A special type of graph is a tree graph. In a tree graph the structure of the connections are such that there are no circuits between vertices. That is, you cannot use the edge connections to go from one vertex to another vertex and then return to the original vertex using a different path. Therefore, in a tree graph, every pair of vertices has a unique path connecting them. A tree graph may have a root, which is a special vertex and a path in the tree can be considered to be moving away from or towards the root. A tree graph will have one or more leaf-vertices, which are the vertices at the terminal end of a path leading away from the root vertex. The vertices and edges of a graph may have associated information and/or labels. A formal way of talking about such information is consider a map from the set of vertices (or edges) to another set that contains the information. So if we have the vertex set $V = \{v_1, v_2, ..., v_n\}$ and the set of real numbers, \mathbb{R} , then a map $\rho: V \to \mathbb{R}$ might be a real-valued annotation of the vertices. For example, the vertices might represent genes, then $\rho(v_1) = 1.67$ would represent the idea that vertex v_I has the expression value 1.67. The range of the map for the vertices (or the edges) might be DNA strings, names (e.g., species names as we will see later when we discuss phylogenetic trees), numbers, pictures, etc.

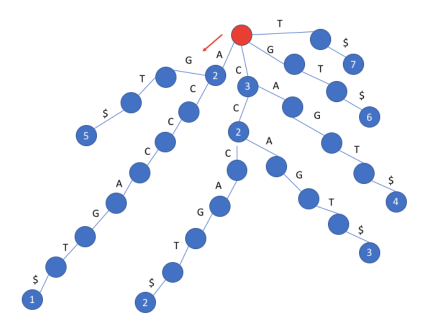
An important data structure for accessing strings is called a suffix tree, or a prefix tree depending on how we arrange things. A prefix tree is just the string inverse of the suffix tree and we can use them equivalently; thus, for this text I will just deal with suffix trees because it relates better to a topic we will discuss later (suffix arrays). (More recently, these structures have also been called a "trie" instead of a tree, from "re**trie**val" to give it a technical flavor.) Suppose we have the string **S** = "ACCCAGT". We start with enumerating all of its suffixes.

ACCCAGT\$
CCCAGT\$
CCAGT\$
CAGT\$
AGT\$
T\$

I put the symbol "\$" at the end of each suffix to denote the end position. The following is the suffix tree:



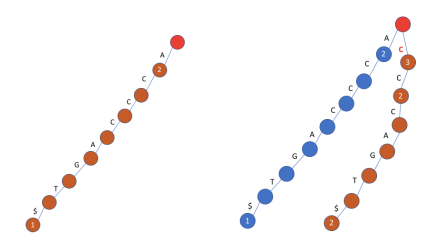
The picture shows a graph with vertices and edges. Each edge is labeled with a symbol. The red circle is the root of the tree. The vertices (also sometimes called nodes) are decorated with other information, which we will discuss in a minute.



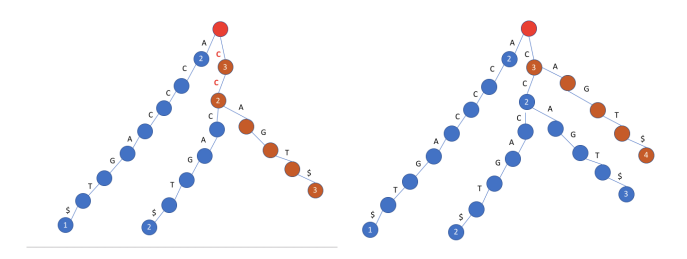
Consider what happens if you walk from the root of the tree toward the leaf of the tree (the last vertex in the path from the root). We walk from the root on the path marked by the red arrow in the figure above.

We first encounter an "A" symbol along the edge. If we now go down the left child branch of the vertex below this edge, we encounter "G", then "T", then "\$". This translates into the suffix "AGT\$". If we go down the right child vertex, we will see a sequence of symbols "CCCAGT\$". Concatenate this with the first "A" and we get "ACCCAGT\$", which is the whole string and also a suffix of **S**. If you continue this kind of path walking from the root to all of the leaves of the tree you will see that we will recover all of the suffixes of **S**. I have annotated the leaves of the tree with the index number of the suffix. So the "AGT\$" path ends at a leaf marked "5", "ACCCAGT\$" path ends at a leaf marked 1, and so on.

Before we discuss what we can do with the tree, I will show you how to construct one by hand: Start with the longest suffix and then construct a set of vertices from the root, as many vertices as the length of the suffix. Put the symbols of the suffix on the edges with the last edge marked with the symbol "\$". We then take the second longest suffix. We examine the symbols on the existing tree to see if this suffix can be "hung" off an existing edge on the tree we have made so far. If this is not possible we start a new branch from the root, like this:

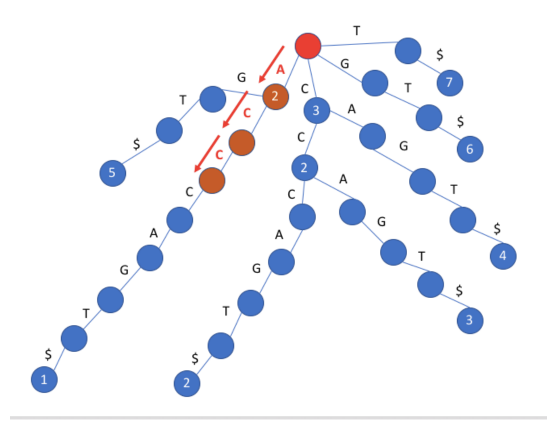


We continue adding branches of the tree that correspond to the next suffix of the original string. At some point in the tree construction, some suffix will naturally extend from an existing branch:

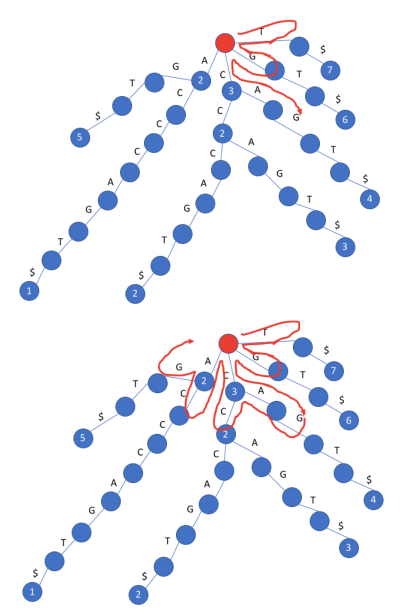


In the picture above the suffix "CCAGT\$" can be added to the tree using the existing "CCCCAGT\$" path as shown above. The same for "CAGT\$". We continue until we finished inserting all of the suffixes.

What can we now do with this tree? Suppose we were looking for the string "ACC" within the string "ACCCAGT". We start at the root of the tree representing "ACCCAGT" and find the edge with "A", then keep going down the tree looking for "C" to make "AC". Keep going until we find "ACC". It took only the same number of operation as the length of the string that we were searching for. Notice that the leaves of this tree has numbering of the suffixes that one gets moving from the root to the leaf. So, from that information we learn that the string "ACC" starts at the 1st position because the branch where we found "ACC" was the branch leading to the 1st suffix. Suppose we were looking for just "C". Again, we start at the root and look for an edge with "C". The edge with "C" has three children that end at leaves marked, 2, 3, and 4. This tells us that the symbol "C" can be found in starting positions 2, 3, and 4. The tree immediately reveals all the repeat structures of the original string. For example, looking at this tree tells us that there is one "T", one "G", three "C"s, and two "A"s. It also tells us there are two "CC"s, and so on.



What else can we do with the tree? We can use the tree to implement inexact searches. Suppose we want to search for "TAG" allowing for up to one mismatch? Then we go down the tree from the root looking for "TAG". When we find one mismatch we keep going down the tree until we either complete the search or find a second mismatch. If we do find a second mismatch we backtrack up the tree and go down another branch, until we find the string that satisfies our one-mismatch condition or we fail. The red line in the figure below shows the search path for "TAG" where we continue the search even if we find a mismatch until we find "CAG". We note that we can't just stop after we find "CAG". We have to continue the search since there could be "GAG" and "AAG". The next figure shows completing the search until we are sure there are no other substrings that match our criterion. For the naïve search algorithm we discussed at the beginning we can also implement a k-mismatch algorithm by simply counting the number of mismatches as we do the exhaustive search. But the tree approach here will be much more efficient (how much?).



As we have just seen, the tree is an extremely useful structure. What are the downsides? Well, the first downside is that the way I've described the construction, it takes $O(n^2)$ time to construct the tree, where n is the length of the database string—this is the amount of time required to enumerate each suffix (or preffix). Fortunately, there is an algorithm to construct this tree in O(n) time, which is too complicated for this course. There is a more serious problem in that storing such a tree requires more memory than the original length of the string. The tree shown above is much bigger than the original string (what is the space complexity?). This can be a problem when the database is very large. It turns out that the structure of the tree can be encoded by using various pointers such that the space usage is O(n), although it can be a fairly large multiple of n. There is also a more compact data structure that can store the tree information called a suffix array (or a prefix array). We will see the suffix array structure later when we treat the Burrows-Wheeler Transform for next-generation sequence data. Unfortunately, suffix arrays are not as simple to access as walking down a tree graph. Regardless, then given something as nifty as the tree data structure, why would we even consider something like Boyer-Moore? First, it takes O(n) time to construct the tree, which isn't an improvement on the Boyer-Moore. And, the Boyer-Moore algorithm uses less extra space (aside from the database string itself), and it is also very simple algorithm. It is only when you use the tree multiple times that it proves itself useful since each search takes time proportional to the search string, not the database. Second, remember that in the best case Boyer-Moore has the time

complexity O(n/m), which can be very good—faster than the size of the database (we call this a sublinear algorithm).

Unit 4: Finite State Automata and computing

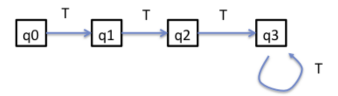
We now consider a very different approach to the string search problem. The motivation here is that a particular approach that we call Finite State Automata (FSA) turns out to be a component of a fundamental model of computing, leading to among other things, the very famous Turing Machine (TM). Furthermore, the FSA approach gives us a standard way of searching for (and defining) a family of abstract string patterns called the "regular grammar" family.

Automata are "machines" (in a conceptual sense) that have "internal states" and read inputs. Upon reading an input symbol, they do something based on the combination of the input symbol and the current internal state and the result is, voila, computation. A FSA, one of the simplest type of automata, is defined by: (1) a set of finite internal states, (2) a set of finite input symbols (or letters), and (3) a set of transition rules that reads "if machine is in state p and reads symbol v change to state q." (A Turing Machine has just a few more rules.) Useful computation is obtained by declaring an "accepting state", which is supposed to represent the completion of some specified task. (We may have multiple accepting states in some FSA.) The main characteristic of an FSA is that it has no memory except the current state. Future behavior depends only on the current state and no past states or past events. You will see later that there are other memory-less models of processes in other disciplines (e.g., Markov models in probability), which share structural similarities to FSA.

The FSA is often specified by giving a table of transitions where the columns represent possible input symbols, the rows represent the possible machine states, and each cell is the next transition state given the combination of row machine state and column input symbol. For example, here is a FSA over {A, C, G, T} that processes a DNA string and reaches the accepting state if it sees "TTT". (Note that this machine continues to process the string after the first occurrence of "TTT".) This FSA has four states labeled **q0-q3**. The **q0** is the initial starting state of the machine and **q3** is the accepting state.

	A	С	G	T
$\mathbf{q}0$	$\mathbf{q}0$	$\mathbf{q}0$	$\mathbf{q}0$	q1
q1	q0	$\mathbf{q}0$	q0	q 2
q 2	q0	$\mathbf{q}0$	q0	q 3
q3 (accepting)	q 0	q 0	q 0	q 3

This transition table can be depicted (see next figure) with a graph where the vertices represent the machine states and the directed edges represent the transition rule conditioned the input symbol (indicated by labels on the edges). Note that since there are four possible input symbols for the DNA alphabet, every vertex should have four edges coming out of the vertex. However, since depicting this can be messy, by convention, we agree that if a particular possible input symbol does not have a corresponding edge drawn, it means to return to the initial state. That is, in the figure below, **q1** has only one edge labeled with the input symbol "T". If the input symbol is "A", "C", or "G", then the transition is back to **q0**.



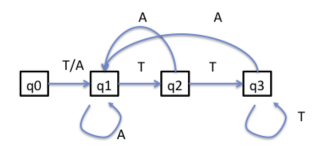
Let's try running the FSA above when the input string is "ATTCTTTG". When following the rows of the table below, read it as "the FSA starts in [State of FSA] and reads the [Symbol processed] which induces a transition to [State of FSA] in the next line."

Step	State of FSA	Symbol processed
0	${f q}{f 0}$	A
1	$\mathbf{q}0$	T
2	$\mathbf{q}1$	T
3	q 2	С
4	$\mathbf{q}0$	T
5	$\mathbf{q}1$	T
6	q 2	T
7	$\mathbf{q}3$	G
8	$\mathbf{q}0$	END

You see that at step 7, the machine reaches the accepting state **q3**. Imagine that a bell rings or something at this point; then you know that a substring "TTT" has been found within the input string. After that point, the machine reads a new symbol "G" and goes back to the initial state **q0**. If you look at the figure, you see that from **q3**, if the input symbol is another "T", then the machine moves to **q3** again. This is for cases where we might have "...TTTT...", in which case we have two instances of "TTT": "...TTTT..." and "...TTTT..."

How do we make the transition table? How did I know how to fill in the transition table (and therefore, the transition graph) to build a FSA that finds "TTT"? One way to think about this is that the class of all possible FSA is a programmable computer, albeit a simple one. A particular instance of FSA is a program in the programming language of FSA. Making a transition table can be considered the same as "programming." For simple kind of FSAs that search for fixed strings, we can describe a step-by-step formula for constructing FSAs that processes the strings. That is, we can construct a program that generates FSA transition tables—you can think of such a program as a **compiler** that translates string patterns to FSA transition tables. A compiler that generates FSA transition tables for a fixed string is relatively easy to write. Compilers for much broader class of string patterns, like the examples shown below, can also be written. Here, we will not study how to write such compilers but concentrate on understanding how to come up with the transition graph for any string searches. It is much easier to draw such figures (graphical programming, so to speak), than to write compilers for FSA.

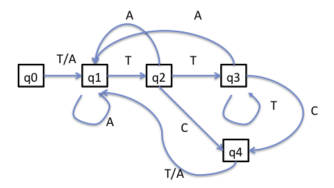
The great strength of FSA is that we can fairly easily generate programs that detect various abstract patterns. For example, suppose we wanted to search of the pattern "[T|A]TT" where by [T|A] I mean T or A. Here is the FSA graph:



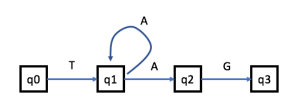
From **q0**, if the FSA sees the input "T" or "A" it makes the transition to state **q1**. We can interpret the state **q1** as the state corresponding to "I have seen one A or one T". Note the other arrows labeled with "A" that feed into state **q1**. For example, at state **q2**, if the input is "T", then the FSA progresses **q3**. If the input is "C" or "G", then it goes back to **q0**. But, if the input is "A",

then this corresponds to "I have seen one A or one T" so it goes to **q1**. Other transition arrows can be figured out with similar reasoning.

Things get more tricky when we are trying to process patterns like "[T|A]T[T|C]".



Now we need to have two accepting states, **q3** and **q4**. I leave it up to you to trace out the logic of the above graph. You can try tracing the computation for some example strings to see that it works.



Finally, we might want to search for patterns like "TA*G", where by A* we mean one or more A's.

This is where we consider an important generalization of FSA. Consider the figure on the left. When we are in **q1** (the state corresponding to "I have seen one T") and we see the input "**A**", we move on to state **q2**. But, we also have the arrow back from **q2** that is also labeled with "**A**". This arrow tells us that when we are in **q1** and receive the input "A", the state of the machine changes to both **q2** and **q1**. That is, now the machine has two simultaneous states.

When we follow the logic of this machine, we see that the machine will continue to keep track of inputs that look like "TA*" until the next input is not "A". When the transition tables of automata contain entries that tell the machines to make transitions to more than one state, we consider such events to be **non-deterministic** in the sense that more than one computational paths are generated. The above diagram shows a non-deterministic FSA (nFSA). There are several different ways to think about what it means to have a non-deterministic computation machine. One way is to imagine that when the machine reaches a state and input that results in transition to two different states, we now have two machines: one that follows the one path (say, transition to **q2** for the nFSA shown above) and another that goes to the other path (say, **q1**). Each time multiple state transitions happen, we give birth to new machines like a Sci-Fi multi-universe scenario. Each of our machines continues computing, and when any one of them reach the accepting state then we know we have found what we were looking for. So, a non-deterministic automaton can be seen a set of (possibly) infinite computing machines. Later, when we talk about the computational complexity of non-deterministic automata, we have to assume that possibly an infinite number of machines are doing the computing.

It should be obvious to you that we could have also created a FSA without two edges for the input "**A**". It will turn out that this is always the case for FSA; there is a deterministic equivalent version of a nFSA—but, a lot of

times giving ourselves the option to use these empty edges gives a cleaner "program". If the automaton is a FSA, we don't necessarily have infinite number of computing machines even in the non-deterministic version. An FSA has no memory and has only its states and input symbols. Therefore, if there are finite states and finite symbols, then for the nFSA there are only a finitely number of different combinations of states and input symbols. Any two machines in the same state receiving the same input symbol do exactly the same thing, thus, there is no need to consider the two machines as something different. For the nFSA processing "TA*G" shown above, at any step in the program execution, there are four possible states: $\mathbf{q0}$ — $\mathbf{q3}$. If an input symbol and the state cause the transition to two different states, then we just keep track of the fact that we now have two machines. In general, for nFSA if we have **q** number of states, we only need to keep track of at most **q** states at any given time. What about computational time complexity? The transition table has the size |S|x|A|, where S is the set of states and A is the set of symbols (the alphabet). In general, to program a FSA to recognize a string of length m, we need $|S| \sim O(m)$; that is, the size of the FSA program is upper bounded by the length of the string. So, constructing the transition table will require O(mp) time where m is the length of the string and p is the number of symbols in the alphabet set. Once we have such a transition table, we have to process each symbol of the input database string. So, if the length of the database string is n, then the computational time complexity is O(n) for the deterministic FSA. For the nFSA, we may have to keep track of up to q states at any input step, where q = O(m)as we already discussed. Therefore, the computational time complexity will be O(nm).

Automatons more complex than a FSA have the capacity to store information as well as change states based on input. For example, a full Turing Machine can be seen as having two sets of memory in a stack data structure. The state of the machine involves both the state of the automaton and the state of the memory. For a non-deterministic automaton, when we transition into multiple machines we have to keep track of all of their states. For FSA there were only a finite number of different states. But, for automatons with memory the state of the memory may have infinite possible configurations. Therefore, something more complex like a non-deterministic Turing Machine is equivalent to (possibly) an infinite number of deterministic Turing Machines.

Finally, we stated that FSA has a connection to formal grammar constructs called regular grammar. Formal grammars are rules for generating strings of symbols as we discussed previously when we discussed "languages". Formal grammars, as opposed to grammars for natural languages, have fixed rules by which "grammatically correct" strings of a language can be generated. There is a kind of a duality between automata that process strings of symbols and formal grammars that generate strings of symbols. An automaton will tell you when an (input) string contains strings or substrings that belong to some rule—that is, belong to a language. If a string is found to belong to a language by an automaton we say the string has been **parsed** by the automaton. So, an automaton for a given language can scan through A* (remember the Kleene star of the alphabet set A) and then tell you which ones are part of the language. A formal grammar gives finite rules by which all such strings can be generated by repeated application of the rules. For example, we may have the rule $R \rightarrow aR$ to mean that we take some string represented by the variable R and replace it with another string that concatenates the symbol "a' in front of R. There is a one-to-one correspondence between a set of formal grammar rules called **regular grammar** production rules and a FSA instance. We won't dig too deep here, but these kinds of ideas have been adopted in computational biology to form the basis for modeling families of genomic strings. For example, we might want to have a set of grammatical rules or (conversely) a FSA that will generate/parse all amino-acid strings that belong to transcription factor family. [This is a hypothetical example, so far, we know of no such grammar.]

Box: Complexity of Algorithms and Problems

So, far we have considered algorithms for solving the problem of searching for particular substrings within a larger database string. We have also discussed performance issues of these algorithms, especially with respect to their space and time complexity. We analyzed the algorithms by considering things like the number of steps for the key operations, etc. We also discussed that we don't really need to know all the details but in general we would like to know the scaling behavior of the algorithms with respect to size of the input. We discussed

asymptotic upper and lower bounds of functions as concepts that help succinctly characterize the scaling behavior. Analyzing an algorithm in this manner is relatively straightforward—most of the time. As we saw from the Boyer-Moore algorithm, some algorithms change their complexity dependent on the input. In which case, we have to strive to find the best and worst cases, as well as consider how we might compute the average. Computing the average can be especially difficult since then we have to define what it means to have random input cases. We will touch upon this subject more later.

What is more interesting is to consider this question: while we can analyze the complexity of an algorithm, what about the problem itself? Can we say something about the computational complexity of a problem? In the exact string search problem, the first naïve algorithm gave us O(nm) time complexity; but, we immediately found a better algorithm with a worst-case time complexity of O(n) ($n \ge m$). If we only knew the first algorithm, we might have said that the time complexity of the exact string search problem is O(nm). However, the Boyer-Moore algorithm tells us that it is O(n). That is, asymptotically bounded from above by f(n) = n. Can we do better? We already know that in the best case Boyer-Moore algorithm has time complexity O(n/m)—is there a clever algorithm that makes this true for all input? Is there an absolute lower bound for the problem so that we will know when we should stop trying to do better? In fact, if we are trying to discuss something as abstract as the notion of absolute computational complexity of a problem, as opposed to that of an algorithm, don't we have to define what it means to compute? For example, we do not know what kind of "computation" a chess grand master carries out, but it is probably not the same kind as what a chess computer program carries out. So, we cannot discuss some inherent notion of computational complexity without fixing some ideas about what we mean by computation. The fundamental nature of computation is a topic we will pursue later. For now, simply extrapolating from the examples that we have seen, we will just say computation is a process that takes in finite set of inputs, carry out iterations of a finite set of operations, for a finite period of time using the usual kinds of computers that we have.

The distinction between computational complexity of an algorithm versus the complexity of a problem is a fundamental distinction in theory of computation. Known algorithms for a given problem will establish upper bounds on the complexity of a problem—after all, a problem can't be more complex than a known solution. Sometimes things like the basic need to read the input can set lower bounds on the problem. For any problem, when we have upper and lower bounds, we would like those bounds to be tight, in the sense that the actual behavior is close to the known bounds. Stating that the exact string search problem is $\Omega(1)$ (lower bound by constant time) and $O(2^{nm})$ (exponential in nm) doesn't really give much information. Proving a good set of asymptotic bounds for a problem (as opposed to an algorithm) can be very difficult. Consider the common grade school algorithm for multiplying two n-digit numbers in decimal notation. We all learned the algorithm of stacking the two numbers together, starting with the right hand most digit of the second number and then sequentially carrying out position-by-position single-digit multiplication. (We know how to do single-digit multiplication because we've memorized it; i.e., we have a look up table for it.) When all is said and done, if you look at the operations, you will see that the standard positional algorithm takes $O(n^2)$ operations, where n is the number of digits for both numbers. This is why we didn't like doing multi-digit multiplications! So, the problem of n-digit multiplication has a known $O(n^2)$ complexity bound. It turns out we can do better by dividing the problem with the strategy of splitting each number into two pieces. For example, if the two numbers are 372643 and 182737, we restate them as the numbers (372*1000+643) and (182*1000+737). You can think about this strategy and how it might help with the multiplication problem. I won't give the details here but I should note that this kind of approach, where we divide the problem into smaller pieces that can be solved more efficiently and then putting it back together is called (naturally) "divide-and-conquer." With this divide-and-conquer approach, it turns out that we can asymptotically approach O(n) complexity for n-digit multiplications. Since reading the two digits takes at least n operations, we might say the problem has $\Omega(n)$ lower bound; and putting the upper and lower bounds together suggests that the problem of *n*-digit decimal multiplication has $\theta(n)$ time complexity.

Analysis for determining the complexity of a problem can be very involved. However, once we start considering various problems of different computational flavor, it turns out that there are fairly large classes of problems that

all seem to share certain characterization of complexity. We may not be able to say a problem definitely has O(n) complexity versus, say $O(n^{1/2})$ complexity; but, we can usually say they belong to one of these larger classes of complexity. To illustrate these larger classes, we discuss three class kinds of problems that are very commonly discussed in computer science textbooks. The first is a classic problem, which was supposed to have been motivated by a parlor puzzle (look it up).

Eulerian path or Eulerian cycle: Suppose we have a graph $G = \{V, E\}$ where V is the set of its vertices and E is the set of its edges. We would like to find a path that starts at some vertex and visits other vertices such that every edge of the graph is traversed exactly once. If the path returns to the starting vertex, we call it an Eulerian cycle; otherwise, we call it an Eulerian path.

At this point, we introduce another key concept in computer science: the <u>decision version of a problem</u>. The decision version of some problem is usually something of a simplification of the original problem, where the problem is restated as a Yes/No, or TRUE/FALSE, binary choice problem. The decision version of the Eulerian path/cycle problem is:

Given a graph $G = \{V, E\}$, is there a path that traverses every edge exactly once?

Note in this case, we didn't ask to find the path but to merely state whether such a path/cycle exists. The decision version of finding a string P as a substring of S is "Given P and S, is P a substring of S?" In most cases, if we know how to solve the decision version of a problem, we can use it to solve the search problem itself.

The great mathematician L. Euler (who among other things invented Topology) gave a concise answer to the decision version of (what we now call) the Eulerian path/cycle problem. The main thing he noted was that if we are to traverse all edges, when we enter a vertex using some edge, then we have to exit that vertex using another edge. So, each vertex that we enter and exit must have even number of edges; i.e., it must have an even degree. The only exception to this is the starting vertex and the ending vertex. We can have exactly two vertices that have odd degrees. For a cycle, since the starting and ending vertices have to coincide, all vertices have to have even degrees. It is easy to see that the condition of even degrees is a necessary condition for the existence of an Eulerian path/cycle. It also turns out that we can show that it is a sufficient condition. We have the following condition that can be checked to give a solution to the decision version of the Eulerian path/cycle problem.

Given a graph G, there is an Eulerian cycle, if and only if the degrees of all vertices are even. There is an Eulerian path, if and only if exactly two vertices have odd degrees and all other vertices have even degrees.

Although the details depend a bit on how we represent the graph G in a program (i.e., the data structure for G), it is reasonable to think that we should have a O(n) time algorithm for checking the degree of each of the n vertices of a graph. So, the Eulerian path/cycle problem can be decided with O(n) time complexity. The function f(n) = n is a polynomial function of n, with the power of n being 1. We call this class, the **P class** where P stands for Polynomial. The Eulerian path/cycle problem belongs to the P class. We are very interested in problems that can be solved in polynomial time, because for such problems we have some hope that we will be able to solve them even if the input size is very large. After all, by Moore's Law, computing power has been growing exponentially. So, a problem that is polynomial in complexity, even if the input size is large, should be solved someday. We call such problems, *tractable problems*. Problems that can be solved in polynomial time form the baseline case of computational problems. Now we turn to what seems like a completely related problem called the Hamiltonian path/cycle problem:

Hamiltonian path/cycle problem: Given a graph $G = \{V, E\}$, find a path/cycle through the graph such that every vertex is visited exactly once. (Of course, the decision version of the problem will be "Given a graph G, is there a path/cycle…")

While seemingly an innocent sister to the Eulerian path/cycle problem, it turns out that there is no known polynomial time algorithm to decide whether a graph has or does not have a Hamiltonian path/cycle. To take a very naïve approach to the problem, if the Hamiltonian path/cycle exists, it must be a permutation of the list of the vertices. Suppose |V| = n, i.e., there are n vertices. Then, examining every permutation of the vertices will involve n! possible cases. If we look up Stirling's approximation for the factorial function, we get:

$$n! \sim \sqrt{2\pi n} \cdot e^{-n} n^n$$

Or, $O(n^n) \sim O(2^{nlogn})$. This is a lot. Actually, we don't really have to look at every n! permutation of the list of vertices since most of these permutations will contain a sequence of vertices that are not connected in G. We can refine our analysis by considering the possible paths starting at one of the n vertices. Then, from the starting vertex, we will have p_0 possible next vertices, where p_0 is the degree of the starting vertex. Suppose we go to one of the secondary vertices, then we would have p_1 next possible vertices, where p_1 is now the degree of the second vertex, etc. As we expand out into possible paths, some of the path will be closed off because we cannot go any more or, we are forced to visit a vertex we have already visited. It is easy to see that we would have $O(nk^{n-1}) \sim O(2^n)$ upper bound on the possible paths where k is the maximum degree of a vertex in the graph. Whether we have $O(2^{nlogn})$ complexity or $O(2^n)$ complexity, the time complexity here is generally of the form $O(2^{poly(n)})$, where poly(n) denotes "polynomial function in n". We note for computational complexity statements, we will include functions of log[poly(n)] to be included in the polynomial function class, since such functions are upper bounded by polynomial functions of n. We call this the **EXP class** (of time complexity).

Following from the above accounting, the naïve <u>algorithm</u> for deciding the Hamiltonian path/cycle problem has EXP time complexity. We are not sure if the Hamiltonian path/cycle <u>problem</u> itself belongs in EXP time complexity class. There might be an efficient algorithm that avoids all the exhaustive enumeration of possible paths. But, as of now, we don't know any algorithm for the Hamiltonian path/cycle that does not have exponential complexity. Yet, here is an interesting twist to this problem. Suppose there is a computational wizard (sometimes called the *Prover*) who shows you (the <u>Verifier</u>) a putative Hamiltonian path/cycle. It is a simple matter (i.e., a polynomial process) to check the validity of the asserted path/cycle solution. A putative Hamiltonian path/cycle for a graph is something we call a <u>witness</u> to the fact that the graph has a Hamiltonian path. And, the validity of the witness can be checked with a polynomial time algorithm. In fact, we can check the witness with O(n) time complexity, where n is the number of vertices. Here is an analogy. Suppose somebody asks you to solve an algebraic equation for real-valued solutions. As you might remember from high school, finding solutions for equations can be hard and tedious. But, if somebody gives you some numbers and claims those numbers are solutions to the algebraic equation, you can easily plug in the answers and check to see that the putative solutions work. Those numbers are witness to the fact that the equation has solutions.

To reiterate, what we just said is:

Given some problem, if somebody gives us a possible solution to the problem, we can check in polynomial time if the solution is correct,

...and therefore verify that the problem can be solved. What is the big deal about this? It turns out that there are problems where this is not the case. Consider the standard chess game. Suppose there is a game in progress and we want to know "Is there a sequence of moves to guarantee that White will win?" Now, a computational wizard tells us that White will checkmate in 23 moves. If this statement is true, it is a witness to the original question and we would answer "Yes, there is a sequence of moves to guarantee that White will win." However, as you might imagine, it is no trivial matter to check the statement that there is a strategy for White to achieve checkmate in 23 moves. To see this, suppose that were to explain to you how there is a checkmate in 23 moves. I might start with, "I move my knight to position X", "then you might move your queen to X*", "or move your pawn to X**", "or", etc. That is, I have to describe all the possible events before I get to the 23^{rd} move that puts the black in a checkmate. In fact, just describing this sequence of solutions will require exponential amount of time. Thus, there are decision problems, where somebody can give a plausible witness statement, yet, there is no polynomial time procedure to check the witness itself.

The discussion on the Eulerian path/cycle problem, Hamiltonian path/cycle problem, and the checkmate problem, suggests that regardless of the particular algorithm, these problems have very different complexity flavors and should be treated differently. We already know that the Eulerian path/cycle problem is in the complexity class P, since we know an algorithm that can solve the problem in polynomial time. A possible witness for the Hamiltonian Path/Cycle problem for a graph of n vertices is an ordered list of vertices, 1>3>2>4>...>n, that is a candidate Hamiltonian path/cycle. We can check this by ensuring that each vertex is represented once in the list and for every adjacent pair of vertices in the list that there is a connecting edge. Therefore, this witness can be checked in O(n) time. For the chess problem, to give a witness to the fact that there is a guaranteed checkmate for white in, say 23 moves, we will have to first show a possible white move, then list all the possible black moves, then show all the white moves corresponding to each of the black moves, etc. That is, we will need at least an exponential sized witness, which will take at least exponential time to check.

It turns out that the Hamiltonian path/cycle problem is in a class called **NP**, which stand for non-deterministic polynomial. The NP class is characterized by the fact that a witness for the problem can be checked in polynomial time. The key notion for problems in the NP class is that a non-deterministic computational machine can solve NP class problems in polynomial time. Recall the discussion from non-deterministic FSA. A view on such machines was a computing machine whose states can multifurcate to generate virtual copies and simultaneously carry out multiple computing paths. If any one of the machines finishes, the problem is solved. If we consider the Hamiltonian path problem, we might consider a process that starts at some vertex and then considers all possible next vertices simultaneously, then again consider all the possible next-next vertices, etc. Each of these paths creates a new "virtual machine" and eventually one of them can verify a witness in polynomial time—as long as the path itself is polynomial in length. Thus, if we have a non-deterministic computing machine, we would be able to solve the problem in polynomial time. Of course, considered this way $P \subseteq NP$; every polynomial time algorithm can also be solved in polynomial time by a non-deterministic machine. The curious thing is that we can't seem to prove that this inclusion is proper. That is, we do not know whether there exist problems that are in NP but not in P. One of the most famous problems in computer science and mathematics is whether P = NP. There are absolutely strong reasons to think that $P \neq NP$. We believe strongly that there are problems that are known to be NP and have no polynomial time solutions. But, no formal proof has been given despite many extremely gifted people working on it. In practice, we know a lot of problems for which no polynomial time algorithm is known, such as the phylogeny problem or the multiple alignment problem that we will discuss later; but we don't have a proof that polynomial time algorithms do not exist for these problems. Unfortunately, it turns out that many, if not most problems, in computational biology are in NP or even harder.

Computational complexity does not stop at NP. There are additional hierarchies in computational complexity such that we might talk about NTIME(f(n)), where f(n) is the time complexity to check the witness of the problem. For NP, f(n) = O(poly(n)). On the other hand, if it takes exponential time to check the witness of a problem, we would say it belongs in the NTIME($2^{poly(n)}$) class. This can go on, of course. There are two important complexity classes that are relevant to computational biology because many problems belong to those classes. The first is a class known as NP-complete and the second is a class called NP-hard. Both of these require us to explore a notion called "**reduction**". Reduction in computation involves taking one problem, say the Hamiltonian problem, and reducing it to (typically as special cases of) another problem. More precisely, suppose we have an input, x, to a particular decision problem (e.g., x is a graph for the Hamiltonian problem). We want to have a procedure—i.e, an algorithm, to turn this input x into a new input f(x) that is suitable for another problem such that if x is a YES case for the first problem, f(x) is a YES case for the second problem. I will give a contrived and somewhat silly example just so you can get an idea of what is a reduction. Consider the problem of searching through a DNA string S = ``ACTG....' for the occurrence of a dinucleotide, say "TG". The decision version of this problem is:

Q1: Given a DNA string S = ``ACTGGGT'', is "TG", a substring of S?

You can, of course, solve this problem using any of the string search algorithms we learned previously.

Now here is a conversion of the string search problem. The inputs to the problem are the string S and the query dinucleotide. There are 16 possible dinucleotides and I will map the dinucleotides to natural numbers like this:

AA -> 2	AC -> 3	AG -> 5	AT -> 7
CA -> 11	CC -> 13	CG -> 17	CT-> 19
GA -> 23	GC -> 29	GG-> 31	GT -> 37
TA -> 41	TC -> 43	TG -> 47	TT -> 53

Note, that the above table is a list of the first 16 prime numbers. Given a string S = ``ACTGGGT'', we also encode each possible dinucleotide in the string including overlaps. So, S => (AC, CT, TG, GG, GG, GT) => (3, 19, 47, 31, 31, 37). Then we take the product of all the numbers, S = 3*19*47*31*31*37 = 95257203. Now to ask whether the dinucleotide "TG" is a substring to S, we use an algorithm for solving a simple arithmetic problem:

Q2: Given x and y integers, is x mod y = 0? That is, does y divide x?

To solve the string problem Q1, we convert the inputs S -> 95257203 and TG -> 47 and ask

Is $95257203 \mod 47 = 0$?

Since S has a unique prime factorization, solving the division problem with the transformed inputs will answer yes whenever Q1 is yes. You can also see that the conversion of the input from one problem to the other can be done systematically with an algorithm. In fact, for the above example, there is a polynomial time algorithm. There is also a polynomial time algorithm to answer Q2. If there is a polynomial time algorithm to convert the inputs, x, of a problem A into inputs, f(x), of problem B, such that B(f(x)) is YES only when A(x) is YES, then we say:

A has a polynomial time reduction to B.

Reduction can be a very important technique in solving many problems. If we are lucky, we might find a reduction of a novel problem to a known problem, making the job of finding solutions easier. Of a more theoretical interest is the following astonishing fact:

There is a subclasses of NP problems, called NP-complete, where all NP problems can be reduced in polynomial time to problems in NP-complete.

This is astonishing because it says that the solutions to some problems can be used to solve all problems in NP. It turns out that the Hamiltonian decision problem is an example of NP-complete problem. This means that it is possible to devise polynomial time algorithms to convert the input of any problem in NP (which includes P problems) to an input of the Hamiltonian problem such that the answer of the Hamiltonian problem answers all these other problems! What? What kind of magic is this? Well, as miraculous as this statement can seem, in practice the situation is not so simple. It is a non-trivial task to actually construct a polynomial time algorithm to reduce a given problem to one of the NP-complete problems. One of the key activities in computational theory is to prove that some problem can be reduced to a NP-complete problem by creating such input converting algorithms—which takes a lot of brain power. A somewhat vague way to think about this is that certain problems have sufficient degree of complexity that most other problems can be represented as an instance of the

complex problem. We just have to be clever enough to find that representation. One last thing, there are problems that are known to be not in NP but all NP-complete problems can be reduced to this new class of problems. These problems are called *NP-hard*. Many combinatorial optimization problems are NP-hard. The phylogeny estimation problem we will encounter later is a NP-hard problem. On a practical level, if we know a problem is NP-complete or NP-hard, we know that we will almost certainly never find a polynomial time algorithm and we will have to figure out strategies to obtain approximate solutions. This motivates a whole area of algorithms called "approximation algorithms" and "heuristics". We will discuss these later. Stepping back a bit, when we think about computational problems, we see that most of the problems can be described as a search of some kind. The string search problem was an explicit example of such problems. But, the Eulerian path, Hamiltonian path, and Chess checkmate problem can all be seen as a search problem too, where the search is through the set of all possible solutions.

Extra Topic: How do we know when a pattern (or anything) is interesting or significant?

Given some clustering program output, say *k*-means, how do we know that the clusters will be meaningful? And, how do we choose between different kinds of output, say *k*-means output at different values of k? This question is very generic in that we could have applied it to any modeling problem. (How do we know some gene regulatory model is significant? How do we know some regression model is significant?) There is no easy way to answer questions of this kind but a heuristic procedure goes like this:

- 1. Find some diagnostic of the pattern—for example, diameter of a cluster, size of a cluster, within and between distances of clusters, etc.
- 2. Choose some probability model
- 3. Compute the probability of the pattern diagnostic

The above is the classical probability model point of view of answering questions about interesting/significance of data, which is the most commonly used view. There are alternative frameworks such as thermodynamic-theoretic framework where we essentially try to consider the theoretical configuration of possible patterns and from that define a notion of entropy. I will not pursue such a framework here but it can be an interesting approach.

For clustering patterns, step 1 can be already rather problematic. If my goal is to divide my input data in its feature space into k different groups, then something like a statistic of average within cluster distance to between cluster distance might be a reasonable diagnostic of how distinct the pattern is. However, many patterns have multiple scales. For example, a hierarchical cluster tree has many levels of clusters and we may seek to characterize patterns at various different scales or maybe the entire clustering tree. This is a difficult problem and there are several ideas in the literature that you can look up. For the moment, let's assume something simple like "cluster density" where we take the number of points and divide it by the diameter of the cluster (defined as the distance between the furthest points). In a hierarchical tree, we might have such cluster density for each hierarchical level of the cluster tree. We will call measures like "cluster density" the "test statistic" for the pattern. I note that the test statistic may be a vector of numbers or something even more complicated (e.g., a tree graph).

In choosing a probability model there are two different directions we may go. The traditional direction is to choose a "null model" of "no pattern". For example, we may choose a model where each point is uniformly chosen within a bounded region of the feature space such that the probability of observing a point in any region is proportional to the area/volume of the region (in relation to the global boundaries). Once we choose such a null model then we can compute the probability of the test statistic under this model and then make statements like "if the null hypothesis was true, we would see the observed result with probability p". If p is tiny then Neyman-Pearson hypothesis testing framework suggests that we reject the null hypothesis—and therefore claim that the pattern is interesting (see also my notes on hypothesis testing in the Probability Handout). There is much discussion in the literature about whether the hypothesis testing framework is useful for delineating significant results and we won't repeat such discussion here. However, I want to remark on a few things:

- 1. Please remember the often-repeated statement that computing the probability of the statistic as, say 0.01, under the null model does not mean that the alternative model is true with probability 0.99. Under the classic hypothesis testing framework, we cannot say anything about the probability of any model or the probability of the data under any other model—just the probability of the test statistic assuming that the null hypothesis is true.
- 2. The choice of the null model is important for determining whether the rejection of the null model will suggest some interesting alternative model. For example, I could have a completely idiotic null model; say, a model that says expression of genes in a cell can only take the values 0 or a million. Showing that the observed data is highly unlikely under this null model is not interesting. Or, for a subtler example,

suppose that we count the number of hydrophobic amino-acids in a protein and then show that it is highly unlikely under the model that any amino-acid can be used with 1/20 probability. The result is not very interesting and it will be hard to convince anybody that the particular protein is significant. In a loose sense, the better matched the null hypothesis is to the data, the more interesting it will be when we reject the null hypothesis at some p-value. Also, ideally the null model should be something one "intuitively expects" for the data—that way, when we reject it, it will be surprising and interesting. For example, if my null model is that amino-acids are used according to the background frequency of amino-acid usage for all proteins in the particular genome, showing that my protein has very unlikely high content of hydrophobic residues would be more interesting. Even better would be to compare against a model of proteins with the same length, same functional annotation, etc. Of course, we also want to be able to give a good biological interpretation to rejecting the null hypothesis such as, "therefore, this protein may be a membrane protein."

There is a technical problem when we have a complex test statistic. A canonical example is when we compare two transcriptomes and the test statistic is the t-test statistic for each gene. We now have a vector of **n** such t-test statistics. This will raise the question of what kind of probability model to use. The simplest probability model is the standard model for t-tests: I have two samples drawn from identical mean and identical variance. The problem is that this is a model for a single marginal value and I have a **n**-dimensional vector of t-test statistics. I need a model for the entire vector. I could try to construct a model for the entire vector by trying to derive the probability of seeing a value of at least X in any of the **n** coordinates. We can also decide on trying to test certain marginal aspects of the **n**-dimensional vector, such as the probability that k number of the coordinates will be greater than X. Various ways of doing this kind of modeling falls under the domain of "family-wise error rate" with procedures like Bonferroni, Sidak, Benjamini-Hochberg, etc. The family-wise error rate problem generally deals with cases when we have a test statistic that is an aggregate of multiple application of the same kind of procedure (e.g., t-test). But, the probability model might involve more complicated scenarios such as the set of cluster densities for each cluster in a hierarchical cluster tree. Therefore, for general situations we need a model of the joint distribution of the configuration of the test statistic—and also a choice of null model rejection regions (see the Probability Handout for this notion) that will give us appropriate interpretations.

The second direction for probability model is to explicitly consider a larger family of probability models. For example, suppose we have gene expression data from multiple cells and we want to carryout k-means clustering. We might have a family of models where we assume that there are k different groups for k = 1 to say 50. Each group is assumed to have some centroid position and a multivariate normal distribution for observed points. We might also assume that the centroid position and the variance-covariance matrix of the multivariate normal distributions are sampled from some probability model. Then, we can compute the likelihood of one of these models like this:

$$L(M_\theta|D) = P(D|M_\theta) = P(x_1, \dots, x_n|f(\vec{x}) = \vec{K}, N(\mu_1, \Sigma_1), \dots, N(\mu_k, \Sigma_k))$$

where $\vec{x} = (x_1, ..., x_n)$ is the vector of observed points (cell transcriptomes), $f(\vec{x}) = \vec{K}$ is a vector of assignment of each of the n points to one of k groups, and $N(\mu_1, \Sigma_1), ..., N(\mu_k, \Sigma_k)$ are the k different multivariate normal distributions. Next, we need a probability model for how we might draw from such a family of models. That is, a prior probability model of the set of probability models. Such a model would give probabilities for various values of k, the probability of multivariate normal distributions, etc. Then we can use the Bayes' formula and obtain the posterior probability of the model given the data:

$$P(M_{\theta}|D) = \frac{P(D|M_{\theta})P(M_{\theta})}{\int_{\theta} P(D|M_{\theta})P(M_{\theta})}$$
(Eq 1)

We can then explicitly give probability values to various clustering results; e.g., what is the probability that k = 3. This is the Bayesian framework for finding the significant/interesting models. Since we get a probability value

for each of the possible models of the data, we might claim that the most probable model or certain set of most probable models is the most interesting set. Not unexpectedly, there are some problems here as well:

- 1. It is somewhat more straightforward to choose a null model than to choose a whole family of possible models. For the multivariate normal gene expression model above, is this a good set of possible models? Should we include other possible distributions besides normal? Why not expand to include a lot more possible models?
- 2. We usually do not have a good guideline for the prior distribution. This prior distribution specifies the probability of any of the particular models from the set of possible models in the absence of data. How do we even begin to contemplate such a distribution? Perhaps by generating a hypothesis about the number of distinct transcriptomes? Should we just decide all models are equally probable? Are there technical problems with such "equally probable" model?
- 3. Computing the posterior distribution can be technically challenging. You can see in Eq 1 above, that the denominator contains an integration of the joint distribution of the data and the models over the family of models. In many cases, we may not know how to do this analytically. There are some algorithmic techniques such as Markov Chain Monte-Carlo (MCMC), but the performance of these algorithms for complicated integration is unknown. In more complicated models, computing the P(D | M), i.e., the likelihood, part of the numerator can be also difficult.
- 4. It may be that if the set of possible models is too large then the probabilities associated with any one of them might be insignificant and we may not have a good distinction between different models—this is not a critique of the approach per se. It is just that, there is a temptation to expand the set of possible models to make sure we do not miss the right model—but there will be a tradeoff in terms of data information.

Whether we choose the hypothesis testing framework or the Bayesian framework, we need to compute the probability distributions implied by the chosen probability model. In the older days, much of the effort was concentrated on solving the technical problem of finding a closed form solution or an analytical approximation for this calculation. These days such computations are commonly carried out using computers and simulation techniques. I should note that if it is possible, it is still desirable to have some kind of analytical approach because it makes it much easier to explore the model variations. Computational techniques for probability distribution has been covered in several places in these notes. I repeat some here so you don't have to go looking for them.

For the hypothesis testing framework, our main goal is to obtain a probability distribution of the pattern text statistic. The easiest approach to this is to use a computer to obtain pseudo replicates of the original data. That is, we can pretend to repeat the observation many different times and compute the pattern statistic for each of these and generate an empirical distribution over the pseudo replicates. There are several different ways we can obtain pseudo replicates. Here are two common ones.

Simulation--If we have a fairly concrete model and a reasonable set of parameter values for the model, we could simulate pseudo replicates of the original observation. For example, if we choose a multivariate normal distribution model of transcriptome that we discussed above, then we can use computational sampling from such distribution to obtain pseudo transcriptomes. We can cluster the resulting datasets, compute the test statistic and repeat many thousand times to obtain a distribution of the test statistic. Then we can examine the original test statistic value to see where it falls in the distribution of the pseudo replicates.

Resampling—We can resample the original datasets with some kind of randomization scheme to obtain pseudo replicates. A common resampling scheme is called "bootstrap" resampling. In bootstrap resampling, the observed data is drawn with equal probability and with replacement. For example, with transcriptome data over say 100 cells, we might generate a pseudo replicate by choosing one of the 100 cells at random and repeating

35

until we have 100 cells. Allowing for replacement means the same cell might be represented multiple times in the pseudo replicate. The assumption here is that the relevant null distribution is the observed distribution of data. There are various possible schemes to bootstrap resample. For example, for each gene, we could have bootstrap resampled the 100 different cell values. The resampling scheme constitutes the null hypothesis and therefore the scheme should be chosen mindful of the various caveats that I raised above.

For Bayesian computations, we have to try to compute the posterior probability. A common computational technique is the Markov Chain Monte-Carlo (MCMC). MCMC is a technique for sampling from a probability distribution in a way to obtain an approximation of the distribution. We first consider the simple idea of sampling from a probability distribution. There are many cases in which we can sample from a probability distribution with reasonable efficiency (and accuracy) but we may have harder time exactly computing the distribution itself. As somewhat of an artificial example, the normal distribution with mean 0 and variance 1, N(0, 1), is given by the probability distribution function $p(x) = \frac{1}{\sqrt{2\pi}}e^{-x^2}$. If we wanted to use it to know Prob{X < c} we would have to integrate p(x), which turns out to be not so easy. But, we might have a method to sample from the N(0,1) distribution. We could sample, say a million times and the resulting frequencies (typically summarized as a histogram) gives us an approximation of the exact distribution. More samples we have more accurate will be our estimate.

We give an algorithm for drawing an approximate N(0, 1) random sample:

- 1. Let $U \sim [0, 1]$ be a uniform random number between 0 and 1. Most computer programming languages have a function for generating a uniform pseudo-random number in the [0, 1] interval.
- 2. Draw from $U \sim [0, 1]$ twelve times and sum up the result. Let X be the sum.
- 3. Return Y = X 6.0

This algorithm relies on the central limit theorem where the sum of random variables converges to a normal distribution. The mean of 12 uniform random numbers in [0, 1] is 6.0. The variance of $U \sim [0, 1]$ is $\int_0^1 (x - \frac{1}{2})^2 dx = \frac{1}{12}$, so the variance of X, the sum of 12 such random numbers is 1; therefore the standard deviation of X is 1. Therefore, Y = X - 6.0 is an approximate sample from N(0, 1). We can now sample Y some suitable number of times, say a 1000 times. We get an empirical histogram from this sampling to approximate p(x) and we can easily approximate $P(x) = \frac{1}{12}$.

Returning to the problem of Bayesian posterior distributions, we want to generate samplings from the posterior distribution such that with enough samples we can get a good idea of the posterior distribution. The idea here is to generate such samples using an iterative process. Recall the definition of an equilibrium distribution of a Markov chain. The goal of MCMC algorithm is to construct a Markov chain such that its equilibrium distribution is, in fact, the probability distribution that we want. At this point, I have to emphasize that the Markov chain part of this algorithm has nothing to do with the Markov chain that we used to model sequence evolution over a tree. It just happens that we are going to use a Markov chain to sample from a desired distribution, but the technique itself is a general technique not related to sequence evolution models.

Let's start with something incredibly simple. Suppose we have a two state A/B Markov process. At each transition event, we toss a coin and move to the other state with probability 1/2. Suppose we start in state A. You can think of this as a marginal distribution with probability 1 in state A and zero in state B, denote this by $\pi'(0) = (1,0)$. After the first step, we have $\pi'(1) = (0.5, 0.5)$. After the second step we have $\pi'(2) = (0.5, 0.5)$ again, so we've quickly reached the equilibrium distribution. Let's see what happens with the actual events. We start with A. At the next step, we might be in B (with probability 0.5). The next step we might be again in B, and so on. So, we might come up with a sequence ABBABBAABAA... and so on. Immediately after the first step there should be an equal probability of occurrence of A and B, so counting A's and B's should give us a good approximation of the equilibrium distribution of (0.5, 0.5). Let's change the model slightly to drive this point home. Suppose at each step we make a change to the other state with probability 1/4 and stay in the same state with probability 3/4.

We again start in state A and the marginal distribution starts $\pi'(0) = (1, 0)$. After one step we have $\pi'(1) = (0.75, 0.25)$. Step 2, $\pi'(2) = (0.625, 0.375)$. Step 3, $\pi'(3) = (0.563, 0.437)$. Step 4, $\pi'(4) = (0.531, 0.469)$. Step 5, $\pi'(5) = (0.516, 0.484)$. We can see this will soon go to $\pi' = (0.5, 0.5)$. Now let's do an actual trial:

While it may not seem that obvious, what our calculation tells us that when we look at this sequence of letters, after some initial steps, the proportions of A's and B's should be equal—that is, the marginal distribution should be (0.5, 0.5). Of course, for any short segment, we tend to have runs of letters because the process tends to stay in one or the other state. However, the long run proportion of the letters will still be 50:50. The result of our construction is a Markov chain whose equilibrium distribution is the target distribution, $Prob\{H = 1/2\}$. If we ever had a problem computing the probability distribution of a flip of a fair coin, we could have used the above process and counted the long term frequency of A's and B's.

As we just mentioned, this long run frequency can be only used after the system has reached (or nearly reached) equilibrium. The period prior to equilibrium is called the "burn-in" period. In a simple model like the above, the burn-in period is short. In a complex model, it is not clear and this is one of the most difficult issues in using MCMC. We know that we can use frequency counts after some initial period, but what is the duration of that initial period? There is no easy way to answer this question—but people use various rules-of-thumb.

Now suppose we want an estimate of some *pdf*, how do we get a Markov chain to converge to this distribution? There are many different ways to do this. One straightforward way is called the Metropolis-Hastings algorithm. The algorithm involves two components. The first component is the state space for the Markov chain (MC). For phylogenetic trees, the state space is the tree topologies and the sequence evolution model parameters. The second component is a way to generate transitions within the state space with a probability model that guarantees convergence to the desired equilibrium distribution. Recall back to our discussion of Markov models. The essence of the Markov model is a set of conditional probabilities for transition events between any two points in the state space. Such conditional probabilities can be explicitly stated as they were for models like the Jukes-Cantor model. Or, we may have a way of generating a sample transition from the Markov model without having an explicit formula for the transition (i.e., conditional) probabilities. The Metropolis-Hastings algorithm gives us a way of generating such sample transitions, such that the distribution of the samples will converge to the desired probability distribution in the long run (i.e., the equilibrium distribution).

Metropolis-Hastings algorithm starts the MC in some state; say, a particular tree topology and a combination of edge parameters. Then it samples from the transition probability using two steps: (1) a step to propose a new state; (2) a step to probabilistically accept or reject the proposed step. We will denote by q(x,y) the probability of proposing a new state y, given that the MC was in state x. We will call this the **proposal function**, meaning that this probability function governs the proposal of a new state from the current state x. One possible proposal function is q(x,y) = q(y)—that is the proposed new state does not depend on the current state. For example, in a phylogenetic tree case, one could propose any possible binary tree with some probability q(y). This is called an independence chain. More commonly, the proposed new state depends on the current state. For example, we could propose one of the Nearest Neighbor Interchange trees with equal probability.

In the second step, we accept the new state with the following probability:

$$Prob\{Accept \ state \ y\} = min\{\frac{p(y \mid data)q(x,y)}{p(x \mid data)q(y,x)},1\}$$
 (Eq 11.6)

First note that the reason we have min(A, 1) on the right hand side is because the ratio A may be bigger than 1 and we want this to be a probability and bounded by 1. The numerator of the ratio is comprised of the product of the posterior probability of the state y (i.e., p(y | data)) and the proposal probability of changing to y from x. The denominator is the complement of this: it is the product of the posterior probability of the state x and the

proposal probability of changing to x from y. If q(x,y) = q(y,x), that is the probability of proposing y from x is the same as proposing x from y, then the ratio is simply the ratio of the two posterior probabilities. Remember that the posterior probabilities can be obtained from the Bayes' formula. Wait, but doesn't Eq 11.6 involve the posterior distribution? Didn't we say this is what we are trying to compute with the MCMC algorithm? Fortunately, what we need is the ratio of the two posterior probabilities, not the individual posterior probabilities. Thus from Bayes' formula the integrals in the denominators cancel and we only need to compute:

$$\frac{p(y \mid data)}{p(x \mid data)} = \frac{p(data \mid y)p(y)}{p(data \mid x)p(x)}$$
(Eq 11.7)

where $p(data \mid x)$ is the likelihood and p(x) is the prior distribution. We especially note that for the formula in Eq 11.7, we only need the likelihood and the prior probability for a particular state value x (or y). Thus we are not trying to compute the full range of the function but just one value. This is especially important when the state space is extremely large such as for binary trees. It is easy to compute the likelihood for a particular tree, hard for all possible trees.

To apply the Metropolis-Hastings MCMC method for trees, we start at some random initial binary tree with particular transition matrices on the edges, denote this (T, θ) , where T represents the tree topology and θ represents the **R** matrix and edge parameters. We first compute the likelihood of the data $p(data \mid T, \theta)$ using the dynamic programming algorithm. We also assume we have a prior probability distribution from which we can compute $p(T, \theta)$. Then we propose a new tree and set of edge transition matrices (T, θ) from some proposal function. We compute the likelihood $p(data \mid T, \theta)$ and prior probability $p(T, \theta)$. Now, we compute

$$\frac{p(data \mid T', \theta')p(T', \theta')q(\{T, \theta\}, \{T', \theta'\})}{p(data \mid T, \theta)p(T, \theta)q(\{T', \theta'\}, \{T, \theta\})}$$
(Eq 11.8)

That is, the product of the ratio of the posterior probability and the ratio of the proposal probabilities. We draw a random number from 0 to 1 and then accept the new state if the random number is lower than Eq 11.8. We repeat this over and over again. (Millions and millions and millions of times.)

After a sufficient burn-in time interval, we collect the states (trees) visited by the MC—say hundreds of thousands of them. We may collect with some spacing between the samples so that we avoid the problem of correlated states within any finite sampling interval. This is an approximate sample from the posterior distribution (if the burn-in time was sufficient). We may find the most probable tree among this collection or represent the whole collection by taking a majority-rule consensus tree we previously discussed.

Several remarks are due:

1. For possibly complicated MCMCs, it is hard to know how long to wait before we finish the "burn in" period. To see how difficult this might be, think of the problem of a multinomial distribution with a very large state space—say a trillion possible outcomes, each with probability p_i , i = 1 to trillion. We have some way of taking samples from this model and we will use the sample to estimate p_i . On one hand, it would seem like we should take at least a few trillion samples to get a good idea of the various p_i values. On the other hand, maybe only ten outcomes have significantly large p_i and everything else is near zero. In which case, if the MCMC tended to "hover" around those ten cases, then we would get a very good estimate of the posterior distribution. For the phylogeny tree problem, the state space who probability distribution we want to estimate is the binary trees and the scaled time parameter for each edge (assuming that the rate matrix \mathbf{R} identical for every edge by assumption). As we discussed previously we have O(t) number of binary t-leaf trees, so the size of the state space can be very large. Whether the situation is like ten major probable states out of trillions or 100's of billions out of trillions depends very much on the data.

- 2. If a tiny proportion of the trees have most of the probability mass, the posterior distribution would be most efficiently estimated if we could have the Markov chain hover around those states. The actual occupancy of the states should be proportional to the relative posterior probabilities as given in Eq 11.6, so the main determinant of whether we would be sampling those states with high frequency depends on the proposal function. In fact, the "black art" of MCMC algorithms is designing the proposal function. As mentioned before, a possible proposal function for trees is something like considering neighboring edit trees with equal probability. This is generally a good thing since neighboring trees are likely to be similarly probable. So, if we have a tree with high probability then we would probably want to sample pretty well around it. But, we do not want our Markov chain to be stuck in some local region of the state space without getting to some other region that also has high probability. We would get stuck if the transition probability was low from one set of states to another set of states—that is, if there were some very low probability trees surrounding high probability trees. A good proposal function should allow us to jump through such barriers. In sum, as you can see, getting a good sample of the posterior distribution has all of the same issues as getting a good optimization of an objective function.
- 3. One other, not so minor, issue is that even if the Markov chain is somehow guaranteed to be in an equilibrium distribution, it may be still difficult to get a good sample of the posterior distribution. First, there might be very many high probability states, say millions. We generally need a sample size that is some large multiple of those high probability states so we might need a very large number of samples. Furthermore, the Markov chain may spend a large percent of its time in one subset of the probable states and then switch very rarely to another subset of the probable states and so on. This will create strong correlation in the successive state samples, which will require even more samples to alleviate.
- 4. If the Markov chain is not in equilibrium or if we don't have enough samples, then we will get a very biased estimate of the posterior distribution. On the other hand, if we were solving the maximum likelihood optimization problem and if we were stuck at some local optimum for the maximum likelihood methods, we would also get a very biased estimate. So, such problems are not unique to the Bayesian-MCMC methods. Unfortunately, these kinds of situations are rather hard to diagnose.

Unit 5: Large-scale searches and Re-sequencing Alignment

We now turn to the problem of doing very large-scale searches including aligning sequences that we know are not too different from each other. The most common application occurs in resequencing. In the resequencing problem, we already know the reference genome sequence, we have resequenced minor variants of the reference genome, and we would like to align the resequenced data to the reference genome. For the most part, in this kind of situation, we do not have to consider somewhat complex problem of positional homology problem that we discuss in the next unit (briefly, the problem of identifying positions in two or more strings that are related to each other by genetic descent). In the resequencing problem, we expect the two strings to be aligned to be nearly identical to each other with obvious positional homology. In this case, the alignment problem is really a string search problem for strings with some bounded difference—say different up to k positions. We could use any of the previous string search methods that allows us to search with up to k number of different letters. The key challenge is for the cases when we have a very large input size. We might have a large input size because the database we want to consider is very large (e.g., Genbank) or because the strings we want to align to the database is large (e.g., next generation sequencing). For these cases, we need to consider strategies that are better than O(n), where n might be the size of the database string. After all, if we have 500 million 100-bp short reads of the human genome, which is 3 billion bases, we don't want to do 500 million searches at O(3 billion) time complexity. As we discussed in the section on prefix/suffix trees, when carrying out multiple searches (e.g., the short reads) over the same database string (e.g., the reference genome) it is usually best to structure the database string. The main problem is how to structure the database string when the number of searches or the database size is very large. Here we discuss several strategies such as indexing and filtration (also called exclusion method) and then look in detail at a space-efficient version of prefix/suffix tree.

We first start with the term "index." An index is kind of a handle into another object; or, more abstractly a function from the set of index values to the set of data objects. For example, an index of a book is a list of words and the pages of their occurrence. RAM memory of a computer is indexed by "addresses" which are some finite sized (binary) numbers and the memory locations contain various data objects. Programing languages allow us to map some variable we might write to the memory by these address indexes. For example, if I use an array and refer to X[1], X[2], etc., the compiler turns these into references for actual memory locations. In general, indexes are well-organized, say non-negative integers, and the object that they reference are more scattered, say all the different things we might put into memory. Thus, an indexing scheme gives us an ordered way to "handle" heterogeneous collection of objects by associating ordered objects to unordered objects. Finally, an indexing scheme might involve an algorithmic procedure that creates the association such as the FM-index we discuss below for Burrows-Wheeler Transforms.

Programming languages usually allow us to use various non-negative integer values as indexes of array variables and the compiler knows how to associate the memory locations. Which memory locations to associate for arrays is relatively straightforward because when the index is non-negative integers, association of the variables to the memory addresses is predictable. For example, a real-valued array running from index 0 to 1,000 can be associated to 1,001 consecutive 32-bit locations. We can think of this as taking one set of well-ordered numbers, the array indexes, and mapping it into another set of well-ordered numbers, the memory addresses. Now, let's consider a more general situation where we might want to store a genomic string in memory and then retrieve positions in the genome by using substrings as indexes. To do this, we need to take the general collection of things that we want to use as indexes, say substrings, and map them to numbers, which then can be mapped to memory locations.

Hashing is the term used to describe various systems to generate a relationship between a general set (e.g., substrings) and the set of non-negative integers with computational efficiency close to O(1) time; i.e., (almost) like RAM memory. We call such a map, a **hash function**. Let **S** be a set of objects of interest; for example, **S** might be the collection of short sequence reads from next generation sequencing. A (perfect) hash function h(t) is a function from **S** to the natural numbers such that if $a, b \in S$ then h(a) = h(b), if and only if a = b. The hash function can be used to use elements of **S** to reference locations in the memory. For example, if we have RAM

40

that is addressed by natural numbers, we might do something like $mem(h(a)) \leftarrow a$; that is compute h(a) and then put the object a into the memory location h(a). Or, in general reserve mem(h(a)) as a place to put information associated with the object a; this is called a hash table. An associative array that we discussed in data structures is an instance of this kind of procedure. For example, we might assign binary values to mem(h(a)) to indicate whether sequences from a particular person contains the substring a. Then when we have a second set of sequences from another person, we can just examine mem(h(a)) to see if a found in this person was also in the genome of the first person. Such a test could be done in O(1) (i.e., constant) time.

Suppose we have the reference human genome and a bunch of 100bp short reads from next generation sequencing. In the ideal case, we might break up the reference genome into $\mathbf{S} = \{\text{set of all possible 100bp} \}$ substrings of the reference genome, construct a hash table with the location of the 100bp substrings, then use the table to align the next generation sequence reads to the reference genome in constant time for each read. This means that we have a method of taking a 100bp string of $\{A, C, G, T\}$ symbols and converting it into a number. One possible scheme is to encode each symbol as a number: $\{A \rightarrow 0, C \rightarrow 1, G \rightarrow 2, T \rightarrow 3\}$ and then the 100bp string can be represented as a concatenation of these numbers; i.e., each position in the string is encoded by a decimal shift. Or, a bit more compactly we could use binary numbers to encode the symbols, needing 2-bits per base position and then shift in binary positions. Such a system would take up 200-bit memory space, which is $2^{200} \sim 10^{20}$ —a totally impractical amount of memory. In practice, we do not have infinite amount of memory, so the range of the hash function must be limited, say 0 to v, where v is the size of the total RAM. If $|\mathbf{S}| > v$, then we will no longer have a one-to-one relationship. In general, $|\mathbf{S}|$ may not be ungodly large. For example, we might have the human genome string of length 3 billion and we want to create an index to all k-mer substring locations. This will be at max 3 billion such k-mers and probably considerably smaller if k is not too large.

Constructing a good hash function involves trying to find a way to map S to a bounded range of number [0 to m], such that m is as small as possible, we can compute the relationship efficiently, and we avoid as much as possible having more than one element of S mapping to the same number. When two or more different elements of S map to the same number, we say that the hash function has a "collision". There are many different ways to resolve such a collision, usually involving examining the details of the object after initial hash function computation that results in using more computing time. A good hash function is difficult to construct. Many programming languages come with a generic hash function (e.g., Perl's associative array) but such functions are not necessarily very efficient (in memory and time). A more efficient hash function can be constructed for specialized applications such as genomic substrings, but this requires work beyond the scope of this text. For this course, we will just assume that if the size of S is not too large, we have efficient hash functions available. One last thought on hash functions is that there is a close relationship between the idea of hashing and the idea of encryption. When we encrypt some text, one way to think about this is that we are taking finite predictable numbers (think of letters as having numerical representations) and trying to map them to random (nearly) infinite range numbers. A hash function takes a collection that might be random and very large, say all possible substrings of a genome, and map it to smaller finite predictable numbers.

In general, if we were to take a human genome and break it into all possible 100 bp substrings, the size of the collection could be very large, up to 3 billion (imagine just taking overlapping 100 bp pieces). But, 3 billion is something that would fit into the RAM memory of modern computers so an appropriate hash function can be used to store the human genome using an indexing system of 100 bp nucleotide strings. However, since we would like to allow for some variations, we would have to hash a much larger collection than just the genome. If we were to allow up to two mismatches in each 100bp string, we now need 16x3 billion indices, suddenly not so easy to fit into standard computing hardware. Once we allow for more reasonable variations like indels, the size of the possible index space becomes impossible. Therefore, direct hashing of short 100 bp reads even with efficient ways to resolve collisions is not a practical solution.

One possible approach is to consider a smaller subset of information about the genomic string, such as *p*-mers of certain size. For example, the genome may be "indexed" by a set of all possible 10-mers, where each possible 10-

41

mer would be associated with a list of locations of those 10-mers. There are approximately 1 million all possible 10-mers of ACGT. Using 2-bits for each letter, an index of all possible p-mers can be stored in 4p/4 bytes (= 2^{2p-2} bytes). Current computers easily have access to 8GB or so of memory so we can store an index of 17-mer or so in RAM. Longer the index length, smaller the number of possible positions in the genome, but more memory we need.

Of course, any particular genome like the human genome may not contain all possible p-mers, if p is of a reasonable size. So an index of all possible 17-mer is not likely to be space efficient—in those cases, we might use a hash function to reduce memory use by only mapping the 17-mers that appear in the reference genome. Suppose we now have a 100bp read and would like to find an exact match in the human genome. We can examine the first 17 bases of the 100bp read and then look up the index of 17-mers, which has the information on the positions where we can find the exact 17-mer. Then, we can extend the strings from the position of each of the 17-mer locations to see if we can find a match to the 100 bp read string. This is called filtration or exclusion. We used the index to filter out or exclude all the parts of the reference genome that cannot contain the 100 bp read. Filtration efficiency can be defined as the expected number of potential matches divided by the expected number of index match positions. For example, suppose for a random 100 bp read string, the expected number of match positions in the reference genome is 5 and the expected number of 17-mer indexed positions is 50, we would have a filtration efficiency of 1/10. Obviously, if we had a 100-mer index, filtration efficiency would be 1. Various different schemes have been considered to increase the filtration efficiency without increasing the size of the index. In principle, a 17-mer occurs once every 16 billion bases for a random sequence with equi-probable bases. Thus, every 17-mer should be expected to be unique within a human genome and this kind of indexing should give us a very efficient alignment. Unfortunately, the human genome is highly redundant. More than 40% of the genome is comprised of nearly identical repeats of short strings. Thus, the filtration efficiency of any reasonable sized k-mer will not be very high.

As mentioned above, indexing exact substrings is not enough because we would like to find matches with k-mismatches rather than an exact match. Suppose our query string is size m, then an immediate fact is that the query string and the database string will share an exact match of a substring of size at least $l = \lfloor m/(k+1) \rfloor$ where $\lfloor x \rfloor$ means the integer less or equal to x. Thus, if we have 100 bp sequence reads and would like to allow up to 2 mismatches, we could filter by looking for exact matches of 33-mers. An index of 33-mers is not feasible, but when the read sizes were smaller (30 bp in the first next generation sequencers), this kind of filtration using a smaller k-mer was a feasible and efficient approach.

Is there a way to use reasonable sized p-mers for filtration but somehow avoid the problem with redundant substrings of the human genome? One idea is called gap filtration. In this strategy we create a p-mer that spans a length that is much larger than p by considering p positions, each separated by a gap of size d. See the figure below and note that p is the total number of positions that we are using while p+(p-1)d is the total span of the index:



Gapped p-mer with p = 4, d = 3

The utility of a gapped index is two-fold. First, as mentioned, many biological sequences have positional redundancies in their sequence. An index is most useful if the occurrence of any index pattern is approximately random in the database sequence—otherwise, we might have some indices that map to tens of thousands of positions, greatly reducing efficiency. By placing gaps in the index, we can help avoid the redundancies that are often spatially adjacent. For example, a mammalian genome has hundreds of thousands (or even millions) of short repeated sequences. A gapped *p*-mer whose total span is greater than the typical size of the repeated

sequences has a better chance of covering unique sequences that yields unique positions. Second, by using multiple gapped p-mer with different gap sizes, we can implement a multiple filtration procedure. In this strategy we would generate a series of gapped p-mers with different gap size d and then in our searches we can retrieve all positions that satisfy multiple gapped indices.

Suppose we had enough memory space to construct a p-mer table of a reference genome of size \mathcal{N} and we wish to use this table to find a string of length m and let's also assume that m/p = q. Under the optimal scenario the probability of occurrence of any particular p-mer is 4^{p} . Given some read, we might break it up into q pieces of pmers and for any one of them we would expect $\sim 4-!N$ number of "hits" in the human genome. One possible strategy is to use the p-mer with the smallest number of hits and then go to each one and see if we can extend it to the full m-length read string. This would take $O(4-\ell Nm)$ time. As another possible approach, we could try to find the conjunction of the positions of the q set of p-mers. Suppose the genomic coordinates returned by the q different p-mers are denoted g_{ij} for the ith position of the jth p-mer. If the read string is in the genome, there must be a set of q different coordinates in the hit lists that are within m distance of each other (because the read string is length m). The naïve way to do this would be to consider the first two sets $G_1 = \{g_{ij} \mid j = 1\}$ and $G_2 = \{g_{ij} \mid j = 1\}$ 2}. Then for each coordinate in G_1 and G_2 , we would try to assess which pairs are within ρ distance from each other (because adjacent p-mers in the read string should be p positions apart). Since $|G_1| \sim |G_2| = O(4-pN)$, such pairwise comparison would take $O(4^{-2p}N^2)$ time. The list of joint hits is expected to be of size $O(4^{-2p}N)$ so the comparison with the 3^{rd} list, G3 will take $O(4^{-3}t/N^2)$ and the next step will be even smaller. Therefore, the time complexity of the naïve method will be determined by the first step and we can say the time complexity is O(4- $^{2}PN^{2}$). If 4PN < m then this is better than just "hit and extend" method. The human genome is 3×10^{9} , so if $p > 10^{9}$ 13 or so, then 4 + N < m. In fact, we can do much better than this. For example, if the coordinate lists are provided to us sorted from smallest to largest, we can do the conjunction search through G_1 and G_2 in $O(|G_1| + |G_2|) = O(4pN)$ time. All of this suggests that filtration can be a very effective strategy even if our available memory is limited. Well, obviously there has to be a caveat. The caveat is that because of redundancies of a typical genome (we also say the genome has low complexity) the expected occurrence of any given p-mer is not $4 \pm N$. Some are much more prevalent than others. Sometimes we can potentially custom design various gapped p-mer indices whose occurrence is nearly uniformly random for a particular genome. This is one area of current algorithmic development.

Space efficient encoding of suffix/prefix strings

In general, if the number of mismatches we want to allow is small, the most efficient way to align the reads would be the prefix/suffix tree we discussed previously. Recall that once such a data structure is constructed, we can efficiently search in time proportional to the length of the query string. The problem was the potential space complexity of the tree, especially for small alphabet size like DNA strings. Recently, an approach using something called the Burrows-Wheeler Transform (BWT) was developed to help solve the space efficiency problem. The BWT is a representation of prefixes or suffixes of a string, which stores information corresponding to a suffix (prefix) tree in a data structure that only needs space in proportion to the size of the database string. It is also related a data structure called suffix (prefix) arrays. Accessing the information in the BWT requires a bit more work than moving down a tree graph but generally it can be accessed very efficiently. Here, I first describe BWT as a representation of suffixes of a string, then how we can use it to search for particular substrings.

BWT is a reversible transformation of the original (database) string into a new string (the transformed string). [I should note that in mathematical terminology a transformation is an one-to-one map over a set of things, such that we have an invertible relationship between pairs of objects in the set. For BWT the domain is the set of strings of length n.] It turns out that if we take a generic string and compute its BWT, the resulting string tends to group recurrent symbols together and therefore have better compression properties. Therefore, BWT has been used as part of compression algorithms. But, for substring searches, it is not too useful to think of it this way. Instead we can think of BWT as producing a string that encodes all the suffixes of a string in such a way that we can search through the suffixes from that string in an efficient manner. Suppose we are given the string

43

"ACAACGT", then we can list all of its circular permutations (meaning shift each symbol ahead by one position, rotate the first symbol to the last position):

0:ACAACGT\$
1:CAACGT\$A
2:AACGT\$AC
3:ACGT\$ACA
4:CGT\$ACAA
5:GT\$ACAAC
6:T\$ACAACG
7:\$ACAACGT

Where the symbol "\$" was used to denote the end of the sequence. I have also labeled the circular permutations from 0 to 7. (Yes, I started the index from zero, because this is how the algorithm is described in other literature and I didn't want you to get confused. Also, this makes it convenient to take into account the fact that we added a "\$".) Another way to think about the circular permutations is that it is the list of all the suffixes of the string (i.e., think of the strings as ending at \$) followed by the prefixes of the string. I will call the above list CP-table for Circular Permutation-table. Now we lexicographically (alphabetically) sort the strings (assume \$ precedes other letters):

7:\$ACAACGT 2:AACGT\$AC 0:ACAACGT\$ 3:ACGT\$ACA 1:CAACGT\$A 4:CGT\$ACAA 5:GT\$ACAAC 6:T\$ACAACG

This will be called **sorted** CP-table. The last column of the sorted CP-table is defined as the BWT of the input string. That is, "TC\$AAACG" is the BWT of "ACAACGT\$". The first column of numbers of the sorted CP-tables is the row indices of the original CP-table. It also happens to be the indices of the suffixes of the original string (see the suffixes in the original CP-table). The sequence of these suffix indices, (7, 2, 0, 3, 1, 4, 5, 6), is called the **suffix array** and keeps track of the original positions. What can we do with this BWT? First, very interestingly, this string "TC\$AAACG" contains information to regenerate the entire set of sorted circular permutations. We can do this by the following steps:

1: Start with the BWT, which is the last column of the sorted CP-table:

T C \$ A A C G

CGT

GT\$

GT\$

T\$A

2: Sort the BW has all the symb			st column, by defin	ition (note that eac	h column of the CP-	table
\$ A A C C C T						
3: Add back the	e BWT					
T\$ CA \$A AC AC CG GT						
4: Sort the augr	mented column	s, this generates the	e first two columns			
\$A AC AC CA CG GT T\$						
5: Add back the	e BWT and sor	t and continue:				
T\$A CAA \$AC AAC ACA ACG	\$AC AAC ACA ACG CAA CGT	T\$AC CAAC \$ACA AACG ACAA ACGT	\$ACA AACG ACAA ACGT CAAC CGT\$	T\$ACA CAACG \$ACAA AACGT ACAAC ACGT\$	\$ACAA AACGT ACAAC ACGT\$ CAACG CGT\$A	

As can be seen in the above sequence, we continue to recreate the columns of the sorted CP-table by this sequence. Once we have the full CP-table, any one of the rows can be used to recover the original string. Thus, BWT is a reversible transformation. Why does the method work? (Hint, for any string S in the CP-table, S[1] is the symbol that immediately follows the S[n] symbol in the original string; or vice-versa, S[n] is the symbol that immediately precedes the S[1].)

GT\$A

T\$AC

CGT\$A

GT\$AC

GT\$AC

T\$ACA

CGT\$

GT\$A

The ability to reconstruct the input string from the BWT string is not so much important for our purposes, but it demonstrates the idea that the BWT string encodes information about the suffixes. Going back to the sorted CP-table:

7:\$ACAACGT 2:AACGT\$AC 0:ACAACGT\$ 3:ACGT\$ACA 1:CAACGT\$A 4:CGT\$ACAA 5:GT\$ACAAC

Suppose we want to search for some particular string, say "CAAC". Then the fact that the table shows all the suffixes of the original string means that if CAAC is a substring, there will be some string in some row that starts with CAAC (indicated by red letters in the above sorted CP-table). Furthermore, since the table is sorted, if there are multiple instances of CAAC, rows starting with that string will follow consecutively. Last, lexicographic order of the strings also means that we can find a particular substring by a search procedure that is sensitive to alphabetical order. Thus, the sorted table is like a suffix/prefix tree in the sense that the information in the substrings has been organized in an ordered manner. For example, in the sorted CP-table above, we see that AC is in the rows 2 and 3 (using 0 as the first row). If we read off the corresponding values in the suffix array (the first column of the above table) for rows 2 and 3, we get 0 and 3 (see blue numbers in the table). These numbers correspond to the offset position in the original string where we can find AC; that is, in the original string "ACAACGT" there is a substring AC in the $1+0=1^{st}$ position and $1+3=4^{th}$ position. We formalize this idea by defining two functions, min(W) and max(W) for some substring W, as the minimum and maximum row position in the sorted CP-table of the strings that starts with the substring W. Given the sorted CP-table, we would like to compute min(W) and max(W) for any query string. The suffix indices of the rows from min(W) to max(W) give us the information on the positions of the substrings we are looking for.

There are two problems here. First, the entire sorted circular permutation table is $O(n^2)$ in size, where n is the length of the database string. Therefore, as given, the CP-table uses a lot of space—as much space as the worst case in the suffix tree. Second, even if we were given the entire table and we use the fact that it is lexicographically ordered, it can take O(n) time to find $\min(\mathbf{W})$ and $\max(\mathbf{W})$ by a naïve search. If the table consisted of the human genome, we will have about a billion rows starting with the letter "A"—a lot of possibilities to wade through to find, say, "AC". Fortunately, there is an iterative process where we can compute $\min(\mathbf{W})$ and $\max(\mathbf{W})$ efficiently using only the BWT, not the full table.

As might be expected, the algorithm proceeds by building on solutions to shorter strings. Suppose we already know $\min(\mathbf{W})$ and $\max(\mathbf{W})$ for some substring \mathbf{W} . We wish to compute $\min()$ and $\max()$ functions for the substring $\mathbf{x}\mathbf{W}$, where \mathbf{x} is a symbol; that is, an extension of \mathbf{W} by one symbol. If we know how to do this efficiently, we should be able sequentially find any substring. Let's start with an example: we know the position of $\mathbf{W} = \text{``AC''}$ in the sorted CP-table above and we want to look for $\mathbf{A}\mathbf{W}$; that is, "AAC". Given that the table is lexicographically sorted, wherever "AAC" is we know that its row position is greater than any rows that contain symbols that are lexicographically less than "A" (i.e., "\$"). If we were looking for "GAC", then its position is greater than any rows that start with A or C and less than the rows that start with T. The first step in our algorithm is to use this kind of information to establish a baseline starting position for any string $\mathbf{x}\mathbf{W}$. Suppose we were looking for "CGT". Then since it starts with "C", we know its row position has to be between [number of A's]+1 and [number of A's] + [number of Cs]. In the original string "ACAACGT", there are 3 A's and 2 C's so "CGT" is in between row indices 4 and 5. We create a function $\mathbf{N}(\mathbf{x})$ as the number of symbols in the database string that is lexicographically less than the letter \mathbf{x} (not counting \$). So, in the string for the CP-table above, we will have $\mathbf{N}(\mathbf{A}) = 0$, $\mathbf{N}(\mathbf{C}) = 3$, $\mathbf{N}(\mathbf{G}) = 5$, and $\mathbf{N}(\mathbf{T}) = 6$. This function allows us to know that any string "Axxxx"

46

will be within the row range 1 to 3, "Cxxx" will be within the range 4 to 5, etc. The function N(x) basically tells us the starting row of the extension x**W**, since the substring x**W** has to start after all the rows of the symbols that are less than x (in lexicographic order).

Assume that we know $\min(AC) = 2$ and $\max(AC) = 3$. We would like to use this to find the offset of xAC amongst the rows that start with the symbol x. We can first use N(x) to tells us the starting position. The pattern xAC is somewhere after N(x). We want to compute this offset. This is where the properties of the CP-table become critical. Recall that for any row of the circular permutation, S[n] is the letter that immediately precedes S[1]. So, if xAC exists as a substring in the original string, there will be rows in the CP-table whose strings have the form "AC...x"; that is, strings that start with the substring AC and end with the symbol x. In general, suppose we have xW then there will be strings of the form "W....x" (i.e., starts with W and end with x) somewhere in the CP-table. Suppose we count the strings in the CP-table of the form "W....x" with W < AC. Then, this also counts all the strings of the form xW where W < AC; call this number g. Recall that N(x)+1 is the starting row index of all strings of the form "xAC...". If we also count the strings of the form "AC...x", call this number h, then N(x) + g + h is the last row index of all strings of the form "xAC..."

How do we figure out g and h? If we look at the row position of min(AC) (i.e., 2 in the table above) and count all the symbols in BWT (the last column of the sorted CP-table) that are x, but is less than the position 2, that corresponds exactly to counting all strings of the form " \mathbf{W}x" where $\mathbf{W} <$ AC. Thus, we can compute the minimum offset (i.e., g) of any string that starts with xAC. Then we add +1 for the actual position because xAC will start one position after this. For the maximum offset, we count strings of the form " \mathbf{W}x" where $\mathbf{W} \le$ AC; that is, we include AC. To do this, we look for incidences of x in the BWT that before the position of max(AC). In our example for AAC, the minimum offset can be obtained by counting the number of A's in the BWT before position 2. There are zero such A's. So, the offset of minimum position of AAC is +1 (= 0 + 1). For the maximum offset, we search for A's in positions equal to or less than 3. There is one of those. So the maximum offset is +1. The base position is 0, since there are no symbols less than A (except for the x symbol). So we come to the conclusion that AAC can be found in the row range 1 to 1 of the sorted CP-table. We look up the corresponding suffix array value, which is 2, and declare that there exists AAC in position 2 of the original string. The key here is that we didn't actually need the full CP-table. We just did all of our computations from the last column, i.e., the BWT string. Thus, there is no need to keep the full CP-table with $O(n^2)$ memory usage. The BWT string only uses O(n) memory.

To put this all together in a more precise form, let B[x, i] be the number of symbol x's found at or before the position i in the BWT string. Then we have the following recursive formula:

$$\begin{aligned} \min(\mathbf{x}\mathbf{W}) &= N(\mathbf{x}) + B[\mathbf{x}, \min(\mathbf{W})\text{-}1] + 1 \\ \max(\mathbf{x}\mathbf{W}) &= N(\mathbf{x}) + B[\mathbf{x}, \max(\mathbf{W})] \end{aligned} \tag{Eq 1}$$

To fix ideas, let's compute the function values for AAC, recursively starting with "C", then "AC", then "AAC".

$$\min(C) = N(C) + 1$$
$$\max(C) = N(G)$$

For any single symbol X,

$$min(X) = N(X) + 1$$
 and $max(X) = N(Y)$ (Eq 2)

where Y is the symbol that lexicographically follows X. We can consider Eq 2 as the boundary conditions for the recursive formula of Eq 1.

For our string "ACAACGT" the BWT is "TC\$AAACG", so we have:

$$min(C) = N(C) + 1 = 3 + 1 = 4$$

 $max(C) = N(G) = 5$

Fig 2 shows that strings starting with "C" are indeed found between 4^{th} and 5^{th} row index. Before we go on, we can just pre-compute the function N(x) and B[x,i] (go back and recall the definitions of these functions):

7:\$ACAACGT
2:AACGT\$AC
0:ACAACGT\$
3:ACGT\$ACA
1:CAACGT\$A
4:CGT\$ACAA
5:GT\$ACAAC
6:T\$ACAACG
Fig 2

N(A)		N(C)]	N(G)	N(T)		
0		3		5	6			
	i=0	i=1	i=9	i=3	i=4	i=5	i=6	i=7
A	0	0	0	1–3	2	3	3	3
С	0	1	1	1	1	1	2	2
G	0	0	0	0	0	0	0	1
T	1	1	1	1	1	1	1	1

Now,

$$min(AC) = N(A) + B[A, min(C)-1] + 1 = 0 + 1 + 1 = 2$$

 $max(AC) = N(A) + B[A, max(C)] = 0 + 3 = 3$

We check Fig 2 again. Next,

$$\begin{aligned} &\min(\texttt{AAC}) = N(\texttt{A}) + B[\texttt{A}, \min(\texttt{AC})\text{-}1] + 1 = 0 + 0 + 1 = 1 \\ &\max(\texttt{AAC}) = N(\texttt{A}) + B[\texttt{A}, \max(\texttt{AC})] = 0 + 1 = 1 \end{aligned}$$

We are right again!

To find any substring using BWT, we need the suffix array (because it keeps track of the offset positions of the suffixes), we need the function N(x) and the function B[x,i]. N(x) is a simple function and can be computed in O(n) time and takes up space only in proportion to the size of the alphabet. B[x,i] is a function that takes up O(kn) space where k is the size of the alphabet and n is the length of the database. One complication is that if n is large, it can take a large amount of memory to represent the number. So, in fact, the B[x,i] table takes O(knm) space where m is the amount of space needed to represent the positions of the string. If the length of the string is n, then our positional notation for numbers takes log(n) space. That is, "1000000" denotes a million numbers but our notation for the number only takes 6 positions. So, the total size needed by B[] is O(k nlog(n)). We can also compute the table for this function in O(n) time once we have the BWT (we just need to count). The suffix array takes up O(n log(n)) space as well, where again log(n) is due to representation of the numbers. As one might imagine from the case of prefix/suffix tree, there exists an O(n) time algorithm to compute the BWT. The O(n) algorithm uses O(n) space to compute BWT. The human genome is on the order of 1 GB of bit-encoded storage (each of the four bases take 2-bits and each byte stores 4 letters) and most computers do not have more than 8 GB of memory, so the exact constant in front of O(n) matters. It turns out there are more space efficient algorithms that have time complexity O(n log(n)).

In sum, the BWT allows us to obtain a suffix array and a representation of the prefix/suffix tree that is space efficient. Searching through the BWT representation is a bit more involved than a simple search down a tree, but as you can see as long as N(x), B(x,i) functions are precomputed, it can be done in time proportional to the length of the query string. The BWT has been effectively used in many programs for next-generation sequence alignment. On the other hand, as the length of the query string goes up we may have problems related to handling k-mismatches. Under a model of uniform mismatch rates, the value of k will have to increase with the increasing length of the sequence reads. A naïve extension of the k-mismatch search on BWT will grow exponentially in the number of mismatches. Next generation sequencing is being used for other applications that may involve fairly large differences in the reference genome and the query string. For example, cancer cells may have a large number of insertions, deletions, duplications, and other genomic rearrangements. In fact, a key characteristic of cancer genomes is the presence of unusually large number of genomic rearrangements. Exact matching or even k-mismatch searches will fail to align sequences around genomic rearrangements. Next generation sequencing is also being used for RNA sequencing. RNA sequences have structural variations like splicing and RNA edits. In fact, RNA sequencing data is revealing that there might be extensive variations in the RNA transcripts that are considerably more different than previously thought. Thus, the RNA copy of the DNA information may be quite different in sequence from the genomic DNA sequence. In all of these cases with structural variations, indels, or large number of mismatches, using a filtration strategy with an appropriate index set is likely to provide better sensitivity. A common procedure now is to use BWT type of efficient search to align reads and then take the left over reads that cannot be aligned in that manner and use a more compute intensive approach that couple filtration techniques with other approaches.

BLAST: Approximate Optimal Alignment

In principle, for any of the next generation sequence read alignments, the algorithm of choice is probably the Smith-Waterman local alignment algorithm. Unfortunately, this algorithm has time complexity O(nm), where n is the database size and m is the query string size. If $n = 3 \times 10^9$ and m = 100 bp and we have 10^8 reads, we have to do $O(3 \times 10^{19})$ comparisons. Thus, we carryout bounded difference searches using techniques like BWT hoping that most reads are not that different from the reference genome. A key part of this is that, in general, algorithms to search for solutions that fall within some bound, e.g., less than 3 mismatches, are easier than algorithms that search for optimal solutions. If an optimization problem is NP-hard, bounded search version of the problem can often be NP or P. This kind of consideration becomes especially important for finding optimally matching sequences in large general database like the Genbank. Very commonly, we might have some read or sequence from a new experiment and we want to compare it against all the known sequences to see whether the sequence is homologous or similar to known sequences. In this case, our databases are already large and growing at an exponential pace, so unless m is pretty small we again cannot run Smith-Waterman type of alignment.

There are various programs that attempt to carry out approximate optimal local alignment against a large database. The most commonly used program is the BLAST family of programs. The actual BLAST program involves different versions for proteins, DNA, etc. and has many tweaks to greatly fine tune its performance (The BLAST servers at NCBI carryout millions and millions of searches a year.) To understand the details of BLAST you will have to consult the latest manuals available online. But, here is an outline of BLAST. The first step is to convert the optimization problem into a bounded search problem. Instead of searching for N optimally aligning subsequences, BLAST tries to generate a list of all subsequences that align with some score better than E. If E is computed in a reasonable manner, the number of subsequences satisfying [score > E] criterion is about N. BLAST computes the criterion using a probability model that approximately computes the probability of a match of quality Q or better if a *random* string of size *m* were aligned against a random string database of size *n*. While there are several ways to use this information, one common way is to compute a statistic of this probability distribution called **Expect** (**E**) value, which is interpreted as the expected number of hits of quality Q or better given a random string of size *m* and random string database of size *n*. The user supplies the desired value for E, then the program calculates hit quality threshold according to its probability model. It then uses a form of filtration (exclusion) as the basic strategy find subsequences that satisfy the quality criterion.

49

For filtration, BLAST uses a series of concepts that they call "segment pairs", "locally maximal segment pairs", and "maximal segment pairs" (MSP). A segment pair of two strings **S** and **P** is an alignment between substrings of **S** and **P** with no gaps. Imagine breaking up both **S** and **P** into all possible words of size k. Alignment of words of the same size from **S** and **P** are segment pairs. The score of a segment pair is computed from a cost matrix. Locally maximal segment pairs are segment pairs whose score is locally maximal in the sense that it will decrease if we extend or shrink the size of the segment pair. Maximal segment pair is the segment pair that has the best score amongst all locally maximal segment pairs. The basic idea of BLAST is to find maximal segment pairs between two strings that together exceed some statistical threshold. The algorithm first constructs a table of words of a fixed size k (a parameter of the algorithm) for both the database and the query string. Then, it finds all instances of the segment pairs of the word size that exceeds a score criterion. These matches become seeds that can be extended to locally maximal segment pairs. The extension rules can vary to allow for gaps. The score of the locally maximal segment pairs is computed to display all those that exceed the preset statistical threshold. However, performance tweaks can sometimes cause some local matches to be missed. BLAST is incredibly fast and it is the most commonly used tool in sequence analysis. However, because of its heuristic nature, best matching is not guaranteed and the results may depend considerably on the parameters used for any given search and the characteristics of the database itself.

Unit 6: Sequence Alignments, Combinatorial Optimization

In this unit we consider the classic sequence alignment problem. In an alignment problem we are given two (or more) strings and we wish to match them to each other, the best we can. Therefore, you can think of this problem as a database-dependent search problem where the result and the algorithm involve optimizing the search over the database (i.e., the second string). This immediately tells us that the computational problem will be harder because we are forced to examine the whole database. There are two kinds of alignment problems. The first involves matching the strings with some notion of best, which must be modeled, while the second involves matching strings that are nearly identical and therefore the notion of best is very simple and problems are closer to the string search problems we discussed in previous units. For the first class of problems we have to work on both the modeling problem and the computational problem. For the second, which typically arises in next-generation sequencing applications, the main problem is the computational problem, which we already discussed in Unit 5—but, we note that the kinds of large-scale searches described in Unit 5 is also trying to solve a version of the alignment problem, therefore, the general discussion on the alignment model also applies to that class of problems.

We first define sequence alignments (we will generalize the definition later). Let S and P be two strings. We consider augmenting the strings by adding a "-" (gap) symbol at various positions. For example, if S = ``AACCG'', augmented S' might be "--AAC-CG". A sequence alignment of S and P are two augmented strings S' and P', such that |S'| = |P'|; that is, the augmented strings have the same length. In this manner, there is an invertible map between the positions of S' and P' where S'[i] $\sim P'[i]$ are <u>aligned</u>. Here is an example of possible alignments for S = ``ATTTGGAA'' and P = ``TTGCA''.

```
S' = ATTTGGAA
P' = -TT-GCA-
S' = ATTTGGAA
P' = --TTGC-A
S' = ---ATTTGGAA-
P' = TTGC----A--
S' = ----ATTTGGAA
P' = TTGCA-----
```

The third alignment above has a "-" in the last position of both S' and P'. In general, we consider such aligned positions of two gap symbols to be superfluous, so in most alignments both the 1st position and the last position have at least one symbol from the original S and P. The fourth alignment above shows that a valid possible alignment is one that offsets the original strings completely. The main utility of alignments is that there is supposed to be biological meaning to the positional correspondence $S[i] \sim P[i]$. The simplest such meaning arises when S might be a known genomic sequence and P is a string obtained from an instrument that sequenced the genome. Then the meaning of the correspondence $S[i] \sim P[i]$ is clear: P[i] is the copy of S[i]. We might see cases of $S[i] \neq P[i]$ within some alignment, in which case, we might attribute this to sequencing error. However, in more general situations, S and P are related strings, but not supposed to be exact copies of each other. In most cases S and P are genetically related to each other by ancestor-descendent relationships. That is, S and P are genetic copies of some common ancestor in the past. Recall that all parent-child relationships involve copying the genome from one generation to another. In the process of this copying, alterations can happen in the child. The simplest kind of alteration would change the base identity, say a mutation from A to C. If we were to look at two different children of the same parent, one child might inherit the string S = "AATTGGAA" and another might inherit P = ``CATTGGAA''. While $S[1] \neq P[1]$ there is still a notion of correspondence between S[1] and P[1]. Namely, they are both (imperfect) copies of the mother's string, call it M, so that S[1] and P[1] are identical in their relative position in the genome due to common descent from M[1]. We call this "positional **homology**" and say that S[1] and P[1]'s are positionally homologous.

Homology is a very important concept. Given two objects, if some of their features are identical (e.g., relative position in a genomic string), and this identity is due to a shared common ancestor, we call the feature (or by implication the object with the feature) homologous. Homology is a logical concept denoting a relational map of biological measurements and/or objects. The shared features due to the shared common ancestor distinguishes the origin of common feature from that arising due to other factors such as chance or physical forces. It is important to note that two or more things are either homologous or not homologous. There is no concept of "percent homology". (One might have two objects that have certain percent similarity.) The biological goal of sequence alignment is to identify positional homologies between nucleotides or amino-acids. We need a model to make this happen.

We go back to the consideration of mother-child genome and think about the kind of things that can happen during the biological replication from the mother's string, M, to the child's string S. We already discussed the case of a single nucleotide mutating (a point mutation). Another common possibility is that some nucleotides are inserted into the sequence, say transforming M = ``AATTGGAA'' to S = ``AATTTGGAA'' or S = ``AATTTGGAA''"AATTTGTGTGAA", etc. This insertion may be random, which can happen through the random mismatch repair mechanism. More commonly, the inserted sequence may come through duplication of nearby sequence, say transforming M = "AATTGGAA" to S = "AATTAATTGGAA" (called tandem duplications; we also call such nearby sequences cis- sequence). Or, we may get an insertion of a copy of a string from far away (we call this trans- sequence, in biology), say through retrotransposition. When a random fill-in happens, the inserted nucleotides have no "ancestor"—we can think of them as spontaneously generated. However, when local duplication occurs or some remote sequence is copied in, the ancestor-descendent relationship of the sequences become complicated; one part of the genome may be the ancestor to another part of the genome. Other kinds of changes can happen, albeit not commonly. We might have local inversion, say M = "ATTGGAA" to S = "ATTAAGG", typically due to the generation of loops in the DNA. Other possible changes are duplication and inversion; chromosomal translocations, where large pieces of the DNA become cut and pasted elsewhere; and other things. Of course, much more commonly, we can have deletions of pieces of the string. When we compare two strings, it is hard to tell whether one of the strings has an insertion or the other had a deletion, so we commonly call them **indels**.

The possible biological processes that change the child's sequence from the mother's sequence can be thought of as a kind of an **edit operation on strings**. Given two homologous strings, they are related to each other by the initial ancestral common sequence and then diverge from each other through a sequence of biological edit operations carried out on the strings throughout their respective genealogical history. (Of course, two strings

may not have any common ancestor, in which case they are not homologous.) So, the first step in inferring positional homology of sequences is to model the edit operations. One possible model is some kind of a stochastic process model that specifies the probability of each kind of possible edits per generation. It turns out such probability models for the kinds of mutational edits we discussed are extremely complicated and hard to convert into an inference method. Instead, the standard approach is to specify an approximate notion of goodness-of-alignment with a numerical score and try to optimize this score. Modeling a problem as an optimization problem is an extremely common form of modeling and we will have more to say about this later. In this particular case, we will be modeling the alignment problem as a combinatorial optimization problem, where the term combinatorial refers to the fact that the objects that are being explored are sequence alignments, a kind of a "combinatorial object", as opposed to numerical optimization, which involve real-valued numbers. In optimization, we have some real-valued function, f(X), where X is the object being evaluated (the **argument** to the function). For each X, the function f(X) measures some kind of notion of goodness of X with respect to our goals; therefore, we call this f(X) function, the **objective function** of the optimization. Choosing a particular objective function corresponds to choosing a particular model for the problem at hand (e.g., the problem of inferring positional homology).

There are many models for sequence alignments in the literature. Most models only allow two types of edits, point mutations and indels. Other types of edits (i.e., translocation) are very difficult to model or compute, although some algorithms do exist. Here, we will only consider the point mutation/indel type of edits under three different variations—the first two are simple models mostly to help us understand the problem and the basic algorithm. The third model, called the affine gap penalty model, is the bread-and-butter of sequence alignments. We will discuss the algorithm for the affine gap penalty model and then consider some additional variations.

The objective function for sequence alignments takes as an input a potential alignment and computes a numerical score. For example, we might have these two possible alignments:

```
S' = ATTTGGAA
P' = -TT-GCA-
S' = ATTTGGA-A
P' = --TTG-CA-
```

One possible objective function (and therefore a model) is to consider the edits implied by the positional homologies of the alignment. I've colored such edits in red. The two alignments above. If we now give each edit a unit score, the two different alignments above would have the scores 4 and 6, respectively. This kind of a score (i.e., objective function) is called an **edit distance**. In this particular case, we would propose to minimize the edit distance score function, thereby minimizing the number of edits between the two sequences. This corresponds to the somewhat vague idea that, all things being equal, mutations/indel are rare and therefore any two related sequences should have a minimal number of edits between them.

Another possible objective function is to compute a notion of similarity between the aligned sequences. For the two alignments above, we might compute the number of positions that have the same nucleotides. The scores then will be 4 and 3, respectively. In this case, we would want to maximize the objective function because (again somewhat vaguely) we think if two sequences are related they should be as similar as possible. One advantage of a similarity score is we can start modeling the notion of similarity itself. For example, we might consider the following "cost" matrix C:

	A	С	G	Т	-
A	1			-1	-2
С	-1	1	-1	-1	-2

G	-1	-1	1	-1	-2
T	-1	-1	-1	1	-2
-	-2	-2	-2	-2	undefined

Each element, C(ij) gives a score for each position of an alignment as a function of the paired symbols in the position. Note that the symbols in the table also include the "-" symbol that denote an indel. In general, we would have a k by k cost matrix where k is the size of the alphabet set in our strings. The alignment

```
S' = ATTTGGAA
P' = -TT-GCA-
```

would have the score -2 + 1 + 1 + (-2) + 1 + (-1) + 1 + (-2) = -3. Using this cost matrix, we can model the score of an indel differently from a point mutation and we can also model the score of different kinds of point mutations differently. Thus, this class of objective functions gives us more flexibility in designing models for biological scenarios (e.g., certain kinds of mutations might be more common than others).

As mentioned the most common kind of alignment model involves an objective function called an affine gap penalty function. Here, the main idea is that biological insertions or deletions don't happen individually but can happen involving multiple base pairs, albeit longer indels might be less likely. We score the indel by a "gap penalty function" of the form:

 $-(Wo + We \cdot h)$ [note the minus sign in front of the parenthesis]

where W_0 is a number called the "gap opening penalty" and W_e is another number called the "gap extension penalty" and h is the size of the gap. For example, let $W_0 = 3$ and $W_e = 1$. Now, consider the following alignment:

```
S' = ATTTGGA-A
P' = --TTG-CA-
```

There are two kinds of gaps, a single two-position gap (marked in blue) and three one-position gaps (marked in red). The two position gap would incur the penalty -(3+1*2) = -5, while the one position gap would incur the penalty -(3+1*1) = -4. The function is called affine gap penalty because it is an affine function of the length of the indels (i.e, the gap). In this model, we will also have to score the positions that have non-gap symbols on both sides such as the 3^{rd} position (T:T) and the 7^{th} position (A:C). (I will use the notation X:Y to refer to an aligned position with symbol X on the top and Y on the bottom.) For the, so-called "**match positions**" (the positions of the alignment which do not have gap symbols on either string), we can again specify a cost matrix similar to the one above (with one less row and column since we are treating the gaps separately).

Having chosen an inference model through the choice of an objective function, we now need to optimize it. This means considering all possible alignments of two strings, computing the objective function, and settling upon the alignment function that maximizes or minimizes the objective function. Unfortunately for us, the number of possible alignments of two strings S and P of length n and m, is rather large. We can write an exact formula for the number of possible alignments (how?) but suffice it to say that it is exponential in the lengths of n and m. Fortunately, the situation is not as bad as one of those NP problems. It turns out that sequence alignment problems have the property that global solutions can be built up from smaller solutions. The main idea is to reduce the problem to smaller pieces and extend the solutions, while guaranteeing that the extension leads to global optimum. [A quick word on local and global optima. Objective functions can have local optima. Local optima are like hills in a mountain range. The form peaks relative to nearby points, but not necessarily the tallest peak (which would be the global optimum).]

Suppose that we have two strings S and P, each of length n and m, respectively. We will consider the ith and jth prefixes of S and P. Since we will talk about this a lot, I will use the short-hand notation [i,j] to refer to the ith and jth prefixes of S and P. For example, suppose we want to align the string S = ``ACTTG'' with the string S = ``ACTTG''. Then the following are examples of the notation:

$$[1,1] \rightarrow S = \text{``A''}, P = \text{``A''}$$

 $[1,3] \rightarrow S = \text{``A''}, P = \text{``AAC''}$
 $[3,1] \rightarrow S = \text{``ACT''}, P = \text{``A''}$

We break up the problem of optimal alignment of S and P to the problem of optimal alignment of the ith and jth prefixes of the two strings; i.e., the best alignment for [i,j]. Note that the original problem is solved if we have the optimal alignment for [n, m]. To find optimal alignment for [i,j], we start with the assumption that we know all optimal alignment for any combination of shorter prefixes. For example, [i-1,j], [i-2,j], [i-1,j-1], and so on. Now in considering [i,j], we can ask whether we need the information from all the shorter alignments. The answer depends on the objective function model. In the case of the edit distance objective function and similarity objective function, it turns out that to find the optimal score for [i,j], we only need to consider the optimal scores for [i-1,j], [i,j-1], and [i-1,j-1] and how to extend the scores to [i,j] in an optimal manner. This is because the score assigned to each potential alignment position is independent of the other positions (e.g., we do not assign the score taking into account previous positions) and it is additive. Thus, we can take the strategy that we will optimally align [i,j] prefix pair by using one-position shorter alignments and then adding one more position to get to [i,j]. Since, there are two strings, we can have three different possibilities: String 1 is one position shorter, String 2 is one position shorter, or both strings are one position shorter. That is, [i-1,j], [i,j-1], [i-1,j-1]. From each of these shorter alignments we build to the alignment of [i,j]. There are thee possibilities:

- (1) From [i-1,j], we insert the *i*th letter from S and an opposing gap symbol to get the alignment of [i,j]
- (2) From [i, j-1], we insert the jth letter from P and an opposing gap symbol to get the alignment of [i, j]
- (3) From [i-1, j-1] we insert the *i*th letter from S against the *j*th letter from P to get the alignment of [i, j]

The cost of any of these three incremental steps can be read from the cost matrix (or the edit operation considerations). So, the possible score at [i, j] given the score of the shorter alignments are:

- (1) Optimal score up to [i-1,j] + cost of *i*th letter from S aligned against a gap symbol
- (2) Optimal score up to [i, j-1] + cost of jth letter from P aligned against a gap symbol
- (3) Optimal score up to [i-1, j-1] + cost of *i*th letter from S aligned against the *j*th letter from P.

The optimal score at [i, j] is the optimal value amongst one of the (1), (2), or (3) possibilities. (Note, the "optimal" may be the maximum or the minimum depending on your objective function.)

This is the basis of **Recursion Formulas** for computing the alignment objective function. I'll just pick one for the similarity alignment for illustration (here C(x,y) is the cost matrix):

$$V(0,j) = \sum_{1 \le k \le j} C(-,P(k))$$

$$V(i,0) = \sum_{1 \le k \le i} C(S(k),-)$$

$$V(i,j) = max \begin{cases} V(i-1,j-1) + C(S(i),P(j)) \\ V(i-1,j) + C(S(i),-) \\ V(i,j-1) + C(-,P(j)) \end{cases}$$
Eq 1

The third formula for V(i, j) in Eq 1 expresses exactly what we discussed above. To solve V(i, j) we need to know some starting values; what we call **boundary conditions**. This is given by the formula for V(0, j) and V(i, 0). The values for the boundary conditions follow common sense. Each corresponds to aligning prefixes of S or P against the empty string. The recursion formula succinctly describes the idea of finding optima for larger objects ([i, j] prefixes) from optimal solutions of smaller objects ([i-1, j], [i-1, j-1] prefixes).

To carry out the computations, we can use a **recursion function** in a programming language to compute V(i, j) until we get V(n, m)—that is the optimal value for the full lengths of S and P. It turns out that using a recursion function is not the most efficient way to compute V(i, j), but for principles sake let's discuss recursion a little bit since it is a common programming device.

BOX: Recursive Functions

One of the technical things you may have trouble digesting, if you've never seen this before, is recursive functions and recursive algorithms. The basic idea is simple. We can often define an operation as "similar" to a previous operation but with some additions. Another way to think about is as building a solution utilizing previous solutions. Eq 1 above is a kind of such iterative relationship. In fact, a mathematical series is the most basic example of defining new quantities or functions by previous quantities or functions. For example, let's consider the following definition:

$$F(n) = F(n-1) + 1, \quad n = 1,...,\infty$$

 $F(0) = 1$ (Eq 2)

The first part of Eq 2 is a function whose argument is a non-zero integer. The second part F(0) = 1 gives the boundary conditions.

Eq 2 is easy to understand going forward in the index n. Here are some values:

n	F(n-1)	F(n)
0	Undefined	1
1	1	1+1 = 2
2	2	2+1 = 3
3	3	3+1=4
4	4	4+1 = 5

We see that in fact F(n) = n+1. It is a simple function that returns the argument plus one. In practice, we might like to know the value of F(10). Then by the definition given in Eq 2, we need to know F(9), which in turn requires the knowledge of F(8), etc. until we get to F(0). The value of F(0) is given by the boundary conditions so we can move back up to F(1), F(2), etc, until we obtain the value of F(10). The fact that the function recursively defines the values by previous values is the reason we call this a recursive function.

For Eq 2, we can solve the recursive formula to derive an algebraic formula that directly gives the value in a closed form. In other cases, it is not so simple. For example, consider the Fibonacci series defined as

$$F(n) = F(n-1) + F(n-2), n = 2,...,\infty$$

$$F(0) = 0, F(1) = 1$$
 (Eq 3)

Can you write a closed form formula for Eq 3?

We now consider a program for computing F(n) in Eq 2. First, an algorithmic function is more or less just like a mathematical function. It takes input values and returns an output value. So we might write something like this:

```
Function F(A,B,C) {

Do some stuff (Prog 1)

Return Z
```

where A, B, and C are the arguments (= input) to the function and I wrote three of them just to emphasize that there might be multiple arguments. The syntax of Prog 1 should be pretty intuitive. It says "get as input A, B, C from external conditions", "Do things inside curly brackets", and "Return the value Z". The value Z might be a number or something more complex. In computer programming we construct functions like Prog 1 and then "call" them by writing their names down with a particular argument value. In other words, when we evaluate a program function F(X), we also say we "call F(X)".

We now write Eq 2 as a program:

```
Function F(n) {

If (n < 0) "Say I give up"

If (n = 0) Return 1

If (n > 0) Return F(n-1)+1
}
```

Now consider what happens if we call the function in Prog 2 with F(3). That is, I want the function value when I pass the argument "3".

```
Step 1: 3 is not negative so I don't give up
Step 2: 3 is not 0 so I don't return 1
```

Step 3: I decrease 3 to 2 and then call the same function with the argument 2, i.e., call F(2). I wait for the return value of F(2) and return that value + 1.

The recursive part of the function is in Step 3, where we call the same function but with a different argument. If we think about it, when F(2) is evaluated the same thing will happen and F(1) will be called. Only when F(0) is called will a concrete value (=1) be returned. So the value of F(3) is not known until we reach back to F(0). After that F(1) then takes the F(0) return value, adds 1 and returns 2 to F(2), which then adds 1 and returns that value to F(3), etc. So, there is a hierarchical nesting of function calls and return of computed values. Concentrating on the values returned, we can rewrite what is happening like this:

```
Return_3(Return_2(Return_1(Return_0 1) + 1) + 1) + 1
```

Where the subscripts indicate the argument value for which each return value is being generated.

Recursion seems complicated when we think of this kind of nested hierarchical function evaluations. But, if we concentrate on the forward definition such as in Eq 2 and Eq 3, you can see that the actual construction is easy. Here is another example that computes n!

```
Function F(n)
```

```
If (n<0) "Say I give up"

If (n=0) Return 1

If (n>0) Return F(n-1)*n

}

(Prog 3)
```

A recursive function need not have integers as arguments. Here is a program that takes in some positive real number and splits the value in half and computes some value. It keeps splitting until the intervals are smaller than a predetermined size. Therefore, the program returns how many halvings are required for the original value to become smaller than the predetermined size. I don't think there is any practical use for this function but please try to follow what it is doing as an exercise.

```
Function Split(X) 

{
if(X<0) \text{ "Say I give up"}
if(X<0.0001) \text{ return 1}
if(X>=0.0001) \text{ return } F(X/2)+1
} 

(Prog 4)
```

Lastly, a recursive function or a recursive program is very nice because it often leads to a clean simple way to describe a complicated problem. For example, the closed form function for the Fibonacci series in Eq 2 is difficult to derive but the statement as a recursive function is simple. However, in terms of computation, there is a great cost to defining functions this way. As noted above, each function evaluation has to wait until all the "lesser" functions are evaluated before finishing. In a computer program all this waiting is stacked into a special memory called the "stack" (ta da! We already saw this before). If you called Prog 3 with a large argument value, say F(1000000), that could cause a lot of memory to be used up in waiting. All recursive functions have equivalent definition in a non-recursive form, although that might not be easy to find. Efficient programmers try to avoid recursion unless the number of function calls is likely to be small.

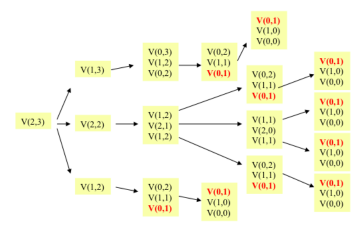
Dynamic Programming to compute the alignment function

If we were to use standard recursion function to solve the recursive formulas in Eq 1 above, we might construct a function that looks like this (assume this function has access to the cost matrix C(i, j) and the two strings S and P):

```
Function V(i, j):

If( i = 0 and j = 0) return 0
Else if (i = 0) return the sum of C(-, C(P[k]) from k = 1 to i
Else if (j = 0) return the sum of C(C(S[k], -) from k = 1 to j
Else {
     Return maximum of the following three quantities:
     V(i-1, j-1) + C(S[i], P[j])
     V(i-1,j) + C(S[i], -)
     V(i, j-1) + C(-, P[j])
}
Fig 1
```

Now consider the sequence of recursive function calls when we call V(2, 3):



The figure above shows the branching sequence of function calls, each function calling additional function, until it reaches the boundary conditions. As can be seen from the above figure, a straightforward recursive computation generates a lot of redundant calculations. I've highlighted the number of times V(0, 1) would have been computed in red. Roughly speaking, if I call the function with V(n, m), then this will generate three new function calls, each of which creates another 3 new function call, etc. until we reach i = 0 or j = 0. If $n \sim m$, we will be carrying on $O(3^n)$ function calls and the computational complexity will be exponential in the length of the string to be aligned.

Tabular Computation

When we think about the problem of computing the recursive function, to compute any V(i, j), we only need to know V(i-1,j-1), V(i-1,j), and V(i,j-1). So, if we compute the function in a systematic manner, we only need to compute each V(i,j) once. For example, for V(2,3), we could first compute all V(0,0), V(0,1), V(0,2), V(0,3), V(1,0), V(2,0) since these functions don't depend on other functions. This is shown in the table below, where X marks the computations:

V(i,j)	0	1	2	3
0	X	X	X	X
1	X			
2	X			

Now to compute V(1,1), we know all the necessary previous function values: V(0,0), V(1,0), and V(0,1). So we can fill in the table:

V(i,j)	0	1	2	3
0	X	X	X	X
1	X	X		
2	X			

If we proceed row by row, left to right, each time we will have all the necessary previous values so we can fill in the table in O(nm) time where n and m are the lengths of the two strings. The technique of finding a global optimum by building up from partial solutions (solutions to the shorter strings in the alignment case) in a

systematic manner to avoid computing redundant values is called **Dynamic Programming**. Here, "programming" refers not to coding but a technique for optimization. This is somewhat due to historical leftover terminology. Unfortunately, the term is in wide use so you will see terms like "**linear programming**", "**integer programming**", etc., all of which refer to optimization techniques. Next, we look at the above "tabular computation" in detail and also how to recover the optimal alignment itself from computing the optimal value of the objective function.

Tabular Computing

The key to systematically computing the objective function is to understand what is implied when we are filling in the [i, j] cell. From the above example, let's consider this stage of the table:

V(i,j)	0	1	2	3
0	X	X	X	X
1	X	X	X	X
2	X	X	X	

When we are filling in the [2,2] cell (marked in red), what we are doing is considering how to extend the partial solutions represented by the neighboring cells (diagonally left, top, and left). Suppose this is for the strings S and P and we will assume that the column indices are referring to positions of S and the row indices are referring to the positions of P. (Of course, we could reverse this and it would not make a difference as long as we keep it consistent.) The solution for cell [2, 2] can be extended from one of three possible directions. The simplest case is extending from the partial solution to [1,1]. The cell [1,1] represents the score of the optimal alignment for the prefix S[1] and P[1]. By definition of an alignment, the alignment of S[1] and P[1] is "filled" in the sense that both the left and right end of the alignments should have nothing dangling. In fact, the optimal alignment of S[1] and P[1] will be one of the following three cases:

$$S[1]$$
 - $S[1]$ - S

We are going to extend one of these alignments to the alignment of the prefixes S[1...2] and P[1...2]. We don't know which of the three cases is the optimal alignment for S[1] and P[1], but we don't care because the arrangement of the base pairs in the S[1] and P[1] alignment will not affect the score of the extension to S[1...2] and P[1...2]. To make the extension we have to add the symbols S[2] and P[2] to the existing optimal alignment. There are again three possible ways:

XXXS[2] -	XXXS[2]	XXX - S[2]
XXX - P[2]	XXXP[2]	XXXP[2] -

I use the notation:

XXX

XXX

to denote the previous optimal alignment, since we don't know its exact form and all we care about is that it is a block of two strings. In general, the score models of alignments typically have the constraint that the cost of aligning two base pairs against each other is less than the cost of two positions that are aligned with a gap against a base pair. That is, we don't allow the 1st or the 3rd column cases—if this weren't the case, most alignments

would simply interleave base pairs against gap characters, leading to nonsensical alignment. So, in effect, if we include this constraint, then the only possible extension of the previous alignment with S[2] and P[2] addition is:

XXXS[2] XXXP[2]

There are two other previous alignments that can be extended to that of [2,2]. We can have an optimal alignment for [2,1] and extend to [2,2] or an optimal alignment for [1,2] and extend to [2,2]. These two alignment extension will have one of the following two forms:

YYYYS[2] YYYY -ZZZZ -

ZZZZP[2]

YYYY and ZZZZ represent the optimal alignment of the smaller strings.

Of these three possible ways of extending smaller optimal alignments to [2,2], one or more lead to the optimal score for [2,2]. We say "one or more" because we can have identical scores and, therefore, multiple optimal alignments. To compute which ones are optimal extensions we need to know the scores for the previous alignments and then add the cost/benefit of the extension to [2,2]. We choose the best score of the three possible scores as the score for [2,2].

Here is a computed example for the strings S = ``AAA'' P = ``AT'' with a cost matrix where each match is +1, each mismatch (substitution) is -1, and each gap is -2. It is useful to add the strings to the scoring table so we can keep track of what we are doing.

	-	A	А	A
-	0	-2	-4	-6
A	-2			
Т	-4			

At the step, we fill in the first row and column. Note that each row and column of the table starts with a gap symbol. The boundary cells represent an alignment of the strings against this gap symbol—i.e., alignment to a null string. For example, going across the top row, the cells are [0,0], [0,1], [0,2], [0,3]. The [0,0] cell is just the starting condition. Then [0,1] cell represents the optimal alignment of S[1] against the null string. The [0,2] cell represents the optimal alignment of S[1...2] against the null string, and so on. So if we were to show the optimal alignments for the top row, it would look like:

[0,0]	[0,1]	[0,2]	[0,3]
NULL	A	AA	AAA
	_		

Given the cost matrix, the values to fill in for the marginal case should be obvious. Now we consider how to fill in the value for the cell [1,1].

	-	А	А	А
-	0	-2	-4	-6
А	-2	(-4, -4, +1)		
T	-4			

In the above table, I listed three numbers (-4, -4, +1) corresponding to the cost of extending from top, left, diagonal, respectively. Let's go over these one by one. The alignment to the left has the score -2. Extending this to the right, implies adding the first "A" of the top string (the S string) to the existing alignment. So, the extended alignment is:

XA X-

where again I used the X's to denote the block of previous alignment. The previous alignment score was -2 and the pair (A: -) has the score -2 according to the cost matrix, yielding a total of -4. By the same argument, extending the alignment from top, corresponds to

X-XA

which also yields a total score of -4. Finally, extending from the diagonal implies:

XA XA

yielding 0 (score of [0,0]) + 1 = +1 score. Obviously, the optimal value is +1, so we would have put the score of +1 for the [1,1] cell.

Here is the finished table where I list all three scores for each cell and mark the optimal value by red:

	-	A	A	А
-	0	-2	-4	-6
A	-2	(-4, -4, +1)	(-6, -1,-1)	(-8, -3,-3)
Т	-4	(-1 ,-6,-3)	(-3,-3, 0)	(-5, -2,-2)

The optimal score on the bottom right most cell, -2, is the optimal score value for the final alignment. As can be seen from the steps we've taken if we are aligning strings of length n and m, with the similarity objective function given in Eq 1, the computational time complexity to obtain the optimal score is O(nm).

All of the above shows us how the recursive scheme and its computation works. But, isn't there a different way to obtain an optimal alignment? Let's consider aligning S = ``ATA'' and P = ``AA'', using the scoring scheme, +1 for match, -1 for mismatch, and -2 for a gap. First convince yourself that the optimal alignment is:

ATA A-A

This is a small enough problem that we can exhaustively try out the alignments and find out that the above alignment with the score 0 is the optimal alignment. We now consider trying to optimally align one letter at a time from each string, say one letter from S then one letter from P, finding the best choice going forward rather that considering all the paths to current alignment as we did above. We first have:

A A

Then, we would do:

AT

A-

Then,

AT

AA

Because that is a better score than

AT-

A-A

Which would have the score -3. What if we did one letter from P then one letter from S? We would have the following:

A-

AA

AT

AA

What if we were allowed to examine one letter from each of the string simultaneously? Well, you can immediately see that we would still end up at

ΑT

AA

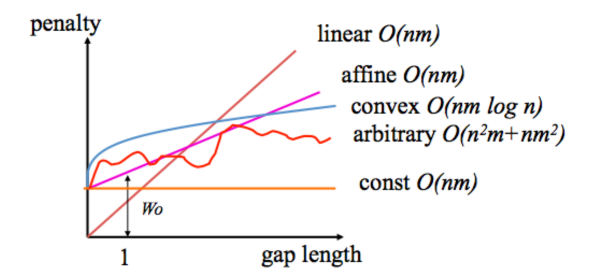
After these steps, we would add the final symbol to get the alignment:

ATA

AA-

with the score of -2.

The approach we have just taken is called a "Greedy" approach. We break the problem into smaller ones but at each step we greedly choose the best next step. There are certain types of problems where this approach is a good thing to do, but in the case of sequence alignment we see that the locally optimal step is not globally optimal; that is, the greedy approach gets us trapped in a local optimum. At each step, working off guaranteed best immediate solution and extending it forward does not work. Instead, we have to work backwards and ask, to get to X, what are the possible optimal paths to X? In the case of the similarity alignment in Eq.1, we were able to trace these optimal paths by considering all the possible one-step back--three of them. Why do we only need to look one step back? This has to do with the alignment model. The basic question for the alignment algorithm is whether it should introduce one or more gaps to align against one of the strings or whether it should introduce match/mismatches. When the cost of indels has a constant cost for each (thus, a linear function of length), this decision can be made one step at a time. Even in the case of the affine penalty function, we can still make the decision one step at a time because once we take into account whether a gap is a new one or an extension, the cost after that is constant per each indel. On the other hand, suppose we have a gap penalty function that is a variable function of the length, then considering one step is not enough since putting in enough gaps may increase the score rather than decreasing it. In which case, we have to keep track of all the possible optimal paths to X. The figure below shows the kinds of gap models we might consider and the computational time complexity of optimizing each kind of penalty function.



The main thing to remember is that each of these corresponds to a biological model. One way to think about the gap penalty models is as a heuristic probability model. Suppose f(k) is the unknown probability mass function for gaps of size k for evolutionary mutations. Then, the penalty function might be seen as a heuristic representation of this probability mass function, typically creating a cost value that is inversely proportional to the probability of the event. As mentioned above, an exact stochastic model incorporating a continuous time probability model has not be developed. The various gap penalty functions will correspond with a stochastic model if the penalty function reflects the **marginal distribution** of the stochastic model; that is, if it reflects the probability of seeing gaps of size k given that the stochastic process has run for some time period t. Unfortunately, for many biological reasonable models we do not have derivation of such marginal distributions.

Obtaining the optimal alignment

The recursive equations (e.g., Eq 1) and the tabular computation only return the score of the optimal alignment, not the optimal alignment itself. If we know the optimal score, is there a way to know the optimal alignment? Suppose we know that the optimal score for aligning S = ``ATA'' and P = ``AA'' is 0. Recall that this optimal score for S[1...3] and P[1...2] is an extension from one of the three possible partial alignments: S[1...3]-P[1], S[1...2]-P[1...2], and S[1...2]-P[1...2]. That is we know that the score is from one of these three cases:

$$V(3,1) + (-2) \rightarrow (-:A)$$

 $V(2,2) + (-2) \rightarrow (A:-)$
 $V(2,1) + 1 \rightarrow (A:A)$

where I have indicated the implied added alignment on the right side of the arrow and the function V(i,j) returns the optimal score of aligning the strings S[1...i] and P[1...j]. We can compute the optimal score of any of the three partial alignments using the same procedure, so we assume we will always also know the value of V(i-1,j), V(i,j-1), and V(i-1,j-1). Suppose the optimal score of V(2,1) = -1 while V(2,2) = V(3,1) = 0. Then V(2,1) + 1 yields the KNOWN optimal score 0. The other two cases would have led to the suboptimal score of -2. So, by this argument we know the alignment of the last position has to be (A:A), and that amongst the three possible ways of getting the final optimal alignment, we built it from the optimal alignment of [2,1]. We can now repeat the procedure to find out the optimal alignment corresponding to V(2,1). Successively applying this procedure we will eventually get to V(0,0) and the whole optimal alignment. Thus, we can reconstruct the optimal alignment using only an algorithm to compute the score and not the alignment. For many optimization problems, if we know the optimal score, we can usually use a similar process to find the optimal configuration

that led to the optimal score. However, as you can see this post-hoc reconstruction is an inefficient way to obtain the optimal alignment. We can easily retrieve the optimal alignment if we keep track of the choices we make during the computation of the optimal score. The following algorithm is called the "**traceback**" algorithm. Recall that when we are computing the optimal score during tabular computation:

V(i,j)	0	1	2	3
0	X	X	X	X
1	X	X	X	X
2	X	X	X	

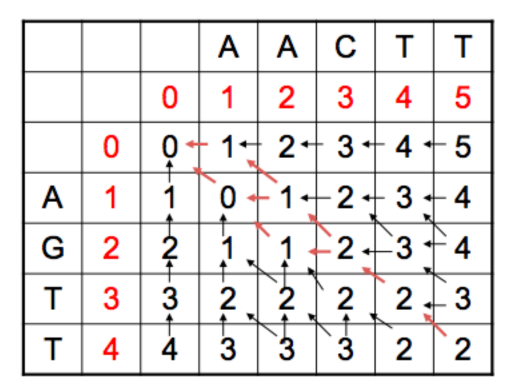
We consider the score of the three possible extensions from the partial alignments:

V(i,j)	0	1	2	3
0	X	X	X	X
1	X	X	X	X
2	X	X	X	

Suppose the optimal score for the (2, 2) cell (marked in red) was obtained from extending the score of the (1, 2) cell. Then, at the point of computation, we can keep a pointer to the partial alignment that yielded the particular score:

V(i,j)	0	1	2	3
0	X	X	X	X
1	X	X	X	X
2	X	X	X	

We can keep such pointers for each cell that we compute. Here is an example of an actual table:



Every cell has pointers (the arrows) but the main cell that is of interest to us is the bottom right most cell, which represents the score of the full alignment. We can start from the bottom-right most cell and trace the pointers back to the top-left most cell to figure out the optimal alignment. In this example, the pointer in the cell (4, 5) traces back to the cell (3, 4). This tells us the last position of the optimal alignment is (T:T). The cell (3, 4) traces back to cell (2, 3), resulting in the 2nd to last position optimal alignment of (T:T). From cell (2, 3), there are two trace back pointers. This can happen if there were two equally optimal choices. This tells us that there are at least two equally optimal alignments. The trace back to the cell (1, 2) implies the 3^{rd} from last alignment of (C:T). That is, we have the following optimal partial alignments up to this point:

...CTT

...GTT

and

...CTT

...-TT

The traceback from the cell (2, 2) points to (1, 1), which in turn points to (0, 0). This sequence results in the alignment:

AACTT

AG-TT

The traceback from the cell (1,2) splits into two possibilities again. There is a pointer to (1,1) and another to (0,1), both of which in turns points to (0,0). The alignments for these two paths are:

AACTT

A-GTT

AACTT -AGTT

So, the table above implies three total optimal alignments, all of which have the score 2.

Variations of sequence alignment algorithms

There are many kinds of variations, i.e., alignment models, that have been explored for biological sequence alignments. Here, we discuss three ideas.

First, we discussed affine gap penalty above and discussed how this was the bread-and-butter of alignment models. How might we compute the optimal function for this function? I won't give the full recursive formula but just sketch the idea here. Remember that when considering the score at cell (i, j), we consider extensions from the cells (i-1, j-1), (i, j-1), and (i-1, j). The extension from (i-1, j-1) implies the alignment (S[i]:P[j]), and therefore this is just like the similarity alignment we discussed above. But for (i, j-1) and (i-1, j) implies adding a gap symbol to one of the two strings. That is, the alignment at this step is either (S[i]:-) or (-:P[j]). Since the affine gap penalty function assigns different scores based on whether a gap is a new gap (i.e., opening penalty) or an extension of an existing gap (i.e., extension penalty) we need to know if this is a new gap or an extension. We implement additional book keeping to give ourselves this knowledge for each cell. At each point in the computation, we compute the following three functions:

D(): The cost corresponding to extending from the diagonal

L(): The cost corresponding to extending from the left. That is, adding a gap to string 1

T(): The cost corresponding to extending from the top. That is, adding a gap to string 2

Remember that at each cell, we have one of three possible ways of constructing the optimal alignment. The three functions above keep track of the right most position of these optimal alignments. The function D() represents the optimal alignment with both positions occupied by a base pair. Therefore, if we add a gap after this position, it will be a new gap. The function L() represents the optimal alignment with a gap in string 1. Therefore, if we add a gap after this position, it will either be an extension (if the gap is added to string 1) or a new gap (if the gap is added to string 2). Similar reasoning will apply for the function T().

As a concrete example, when considering an extension from cell (i-1,j) to cell (i,j), we are extending from the left and will be adding a gap to string 1. We need to consider three function values at cell (i-1,j) and whether the extension to the cell (i,j) implies a new gap:

D(i-1, j): New opening because this score represents previous diagonal extension from the cell (i-2, j-1). L(i-1,j): Extension because this score represents previous gap in string 1 as an extension from the cell (i-2, j) T(i-1,j): New opening because this score presents previous extension from the cell (i-1, j-1); that is, a gap in string 2, not string 1, in the previous position.

After we consider all of these three possible kinds of scores, we compute the maximum of all three as the optimal score for the cell (i, j). Thus, for each cell we keep track of four different function values, D, L, T, and the maximum of the three values (as V). In general, to compute the optimal values at any given cell, we may need to know the previous k-steps of computation. Keeping track of each of the k-step results and projecting forward is the main determinant of the computational complexity statements shown in the figure above on gap penalty models.

66

The second variation for alignments involves modifying the scoring scheme to allow certain kinds of "free" variations. For example, suppose we want to allow the strings to have "free" gaps in the beginning or the end of the alignment. That is, allow alignments that look like this without any penalty for such hanging ends:

(Why would we want such alignments?) We can just change the boundary conditions above.

$$V(0,j) = 0$$

 $V(i,0) = 0$ (Var 1)

The boundary conditions above, sets all the cells in the first row and the first column to zero. This is the same as saying that you can start the alignment with arbitrary number of gaps on one of the two strings; no matter how many starting gaps, the score is still zero. Such an alignment might be useful if we are trying to piece together short overlapping substrings from a long common string as might happen in shotgun sequencing.

A famous variation of a similar kind is the Smith-Waterman local alignment. In S-W local alignment, not only do we set the boundary conditions to zero, but we also change the recursion to:

$$V(i,j) = max \begin{cases} V(i-1,j-1) + C(S(i),P(j)) \\ V(i-1,j) + C(S(i),-) \\ V(i,j-1) + C(-,P(j)) \end{cases}$$
 (Smith-Waterman)

Note the zero as an option. This means anywhere in the table, while computing an optimal score we can restart the alignment from the suffixes starting from that position (zero)—this is the very definition of a local alignment. It is the best possible alignment of any substring of the two original strings because the algorithm allows you to restart the alignment if the optimal score drops below zero. Of course, the scoring scheme has to be such that we expect positive values for good local alignments. Under the Smith-Waterman alignment scheme the optimal local alignment score can be found in any of the n by m cells, rather than the bottom-right most cell. This is because we are looking for any sub-alignments of substrings that might be locally best. When we do the traceback we also stop at when we reach a cell with value of zero. The length of the optimal local alignment is determined by the position of the cell(s) at which the top score is found and the cell at which a score of zero is found tracing back from the top scoring cell.

The last variation concerns how we choose the parameters for the alignment objective function. By parameters, I mean a set of numbers that change the behavior of the algorithm independent of the input data. An example is the gap opening/extension penalty for affine penalty function and the values in the cost matrix. Since there are two numbers for the opening extension penalty and 4x4 = 16 possible values for the cost matrix, we would say we have 18 parameters for this model. For the cost matrix, we might use some compilation of closely related sequences (therefore where the alignment is unambiguous) and tabulate the kinds of base pair or residue changes and then come up with an **empirical cost matrix**. For the gap opening or extension costs, it is not so clear. In practice, most biologists try various values and then try to see if the resulting alignment "makes sense". Typically, making sense means that the resulting alignment isn't sprinkled with small indels, because most biological processes don't work like that. Other factors include things like not putting gaps into "functional domains" which are stretches sequences with special function such as catalytic sites. One interesting idea called **parametric alignment**. Here, we consider a suitable range of values for the alignment model parameters. For example, we might consider the gap-opening penalty between 1 and 10; while also considering gap extension penalty of 0 to 5. Then we compute all possible alignments of all possible combination of the parameters within this range. When we do this, many of the combinations of the parameter values will yield the same optimal

alignment (in fact, for the affine gap penalty function, equivalent alignments form a convex set within the parameter space), We might then choose the alignment that is the most "robust" in the sense of being most prevalent amongst the possible parameter value candidates.

The alignment objective function models and the dynamic programming algorithms for computing their optimal values cannot make the judgment of whether the resulting alignment represents the true positional homology of the strings. In general, for any inference problem (which is what we generally do in computational biology), we would like to obtain the inferred object (such as the alignment) and also a measure of the quality of the inferred object. If we use statistical models, many classical techniques try to provide both the inference and the quality of the inference (e.g., an estimation of the mean and the standard error of the mean). Without such probabilistic framework, it is difficult to formulate a good quality estimation procedure. For alignments, several kinds of approaches have been discussed and developed in the literature but there isn't a well-agreed on method for assessing the quality of the positional homology inference. When we have multiple sequence alignments, something we discuss next, we can use the pattern of the alignment across the multiple sequences to try to assess the quality of alignment. But, multiple alignments have computational problems and they also do not have a canonical statistical framework or assessing quality. Thus, measuring the statistical properties of sequence alignments remains an open problem.

Multiple Sequence Alignment

Typically, when aligning moderately disparate sequences, we want to align more than two such strings. We usually have a collection of strings that are either from multiple species/strains/individuals or members of a gene family; and, we want to compare and analyze them simultaneously. Somewhat surprisingly, unlike the pairwise alignment, it turns out that the multiple sequence alignment is a very difficult computational problem. Before we discuss the biological model, we first examine a key component of multiple sequence alignment: how to superposition different alignments. Suppose we are given a pairwise alignment:

```
S1:AC-GGT--T
S2:AGGG-TCCT
```

Now suppose we have a third string S3 = "AAGGGTCT". We can compute an alignment of S3 against S2:

```
S2:AG-GGTCCT
S3:AAGGGT-CT
```

We now put the two sets of alignments together by "gap-augmenting" the alignments where we add gap symbols where necessary to reflect the previous two alignments:

```
S1:AC--GGT--T
S2:AG-GG-TCCT
S3:AAGGG-T-CT
```

In the three-way alignment above, the red gap in S1 is there because the alignment between S2 and S3 had that gap. The blue gap is in S3 because the alignment between S1 and S2 had that gap. This way of superpositioning alignments has a nice property. We can take the three-way alignment and reduce it back to the two pairwise alignments like this:

```
S1:AC--GGT--T S2:AG-GG-TCCT S2:AG-GG-TCCT S3:AAGGG-T-CT
```

Then in each of these, we eliminate the positions with gaps on both strings:

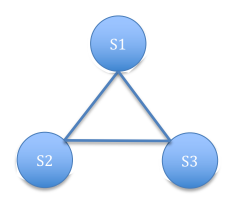
S1:AC-GGT--T S2:AG-GGTCCT S2:AGGGT-CT

We recover the two previous pairwise alignments. This means that if S1:S2 alignment was optimal and S2:S3 alignment was also optimal, the three-way alignment preserves these pairwise optimal alignments. We can schematically draw the way we constructed the three-way alignment like this, where each edge represents optimal pairwise alignment:



The above picture might be called an alignment graph for multiple sequences.

What if we wanted to consider the third possible pairwise alignment, i.e., S1 vs S3? Schematically, we are talking about an alignment graph that looks like this:



It turns out that in this case, finding the three-way alignment that preserves the optimal property of all three pairwise alignments is not so easy. We can't just add a few gaps here and there. We might try to define an optimality criterion as "find the k-way alignment such that the sum of all pairwise alignment scores are optimal". There are C(k, 2) (i.e., k choose 2) number of pairwise scores. This is called the "sum of pairs" optimality criterion. It turns out the optimization can be achieved by using a multi-dimensional version of the dynamic programming algorithm. We write a recursive relationship of $V(i_1, i_2, i_3, ..., i_k)$ for the k strings, where on the right hand side you will have all the 2^k -1 different "one-index-before" possibilities. If the average length of each string is n, then this algorithm has time complexity and space complexity of $O(n^k)$, or exponential in the

number of strings. Obviously, this is not an easy thing to compute for any reasonable sized dataset.

In general, if we have an alignment graph with *k* vertices (therefore *k* strings), we might want to find the *k*-way alignment such that all pairs of strings connected by the edges have optimal scores. The sum of ALL PAIRWISE criterion can be thought of as an alignment graph that is a complete graph—i.e., every vertex is connected to every other vertex. It turns out that if the alignment graph is a tree, no circuits, then we can just compute pairwise optimal alignments and then paste them together like what we did at the beginning. If the graph has circuits like the second picture above, we can't easily paste them together because what is optimal for two edges might conflict with the third edge. Fortunately, as we see below, there isn't a good biological reason to consider a complete graph as the alignment graph. We pretty much always want to consider an alignment tree (sometimes called the guide tree). But, before we go on consider another problem. Suppose we have an alignment graph like this:



But we don't know the string S2; for example, because S2 is an unknown ancestor to S1 and S2. In this case, we might consider the idea of allowing S2 to be any string—but, it should be a string such that the pairwise

alignment to S1 and also to S3 should be optimal. This is again a computationally hard problem. In fact, the general version of this problem falls into the NP-hard class.

Regardless of the computational preliminaries, recall that alignment of two sequences involves trying to estimate positional homologies of the two sequences. For two sequences, we assume they are related to each other by a common ancestor. For three sequences, they might be related to each other by a single common ancestor (Fig 1a) but more likely possibility is that they are related to each other by a successive bifurcating tree (Fig 1b).

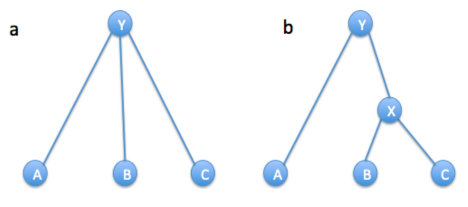


Fig 1

Either way, if we have k different strings, the proper biological model would be to seek to align them in such a way that the alignment follows their (most likely) tree graph genealogy. Unfortunately, there are two problems with this idea.

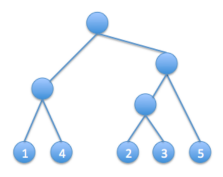
The first problem is that until we align the sequences, we don't actually know if B and C are more closely related and therefore should be aligned first. Maybe we should align A and B together first, or even A and C. In fact, there are three possible bifurcating trees (also called binary trees) for three sequences. We will see in more detail later that if we have k strings, there are $O(2^{klogk})$ number of possible binary trees. So, that is a lot of possible trees to consider. As mentioned, a tree graph that outlines the order of multiple sequence alignment like Fig 1b, is called an alignment guide tree. A guide tree may also be a phylogenetic tree (i.e., the tree depicts the genealogical history of the sequences) or it might be some other tree that might have some other biological rationale.

The second problem involves X. The string X is a hypothetical ancestral string to B and C. Ideally, if we knew the hypothetical ancestors X and Y, we would align B to X, C to X, A to Y, and X to Y in the manner we discussed above. Unfortunately, we don't know X or Y, since those are hypothetical ancestral sequences. I mentioned this can be a hard problem. Here, we would have to consider all possible strings at both X and Y. Let **X** and **Y** depict the collection of all possible strings at the vertices X and Y in Fig 1b. The idea is to choose the alignment and the strings $x \in X$, $y \in Y$ such that the alignment score is optimized. That is, we find the strings x, γ , and the alignment simultaneously by optimizing the objective function. We can call this problem the "phylogenetic alignment for a fixed tree"; i.e., the problem of inferring the optimal alignment and the optimal hypothetical ancestral string together, assuming we know the tree. Yup, this problem is known to be NP-hard. The *full phylogenetic alignment problem* is to infer three things at once: a phylogenetic tree as the guide tree for the alignment; the hypothetical ancestral strings; and the alignment of all the strings. Since the problem is NP-hard for any fixed tree and there are $O(2^{klogk})$ possible alignment trees, it is easy to see that this is also an NP-hard problem. So the simultaneous inference of the phylogenetic alignment tree, the ancestral strings, and the multiple alignment, is a very hard computational problem indeed. Recently, some programs have been proposed to approximately solve this problem and we will discuss these later. Since the exact solution to the problem is hard, there are two heuristic things we can do. First, we can fix the alignment tree without trying to examine

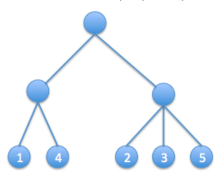
every possible tree. Somehow, we need to fix the tree in a reasonable way without getting information from the full multiple alignment Second, we shouldn't try to find the ancestral sequence that best optimizes the alignment score on the tree, but approximate it with some simple procedure using the information from pairwise alignment of known sequences. For example, in Fig 1b, we might align B and C, then use the pairwise alignment to create a hypothetical ancestral string X, which in turn can be aligned to A.

There are many many different multiple alignment algorithms out there. All of them differ mostly on how they fix the alignment guide tree and how they agglomerate the alignments at the "children" vertices of the guide tree into successive alignments "up" the guide tree. Here we examine just one idea that is approximately the original idea in a popular multiple alignment algorithm called CLUSTAL (which has various version numbers like CLUSTAL V, CLUSTAL W, etc.). I say "approximate" because these algorithms are constantly evolving and any description might differ from the latest version. The first step is using the input sequences to create an alignment guide tree. Ideally, the guide tree should be the phylogenetic tree that correctly depicts the genealogical relationship of the trees. But, it is hard to hope that we can optimally infer a phylogenetic tree without a multiple alignment in the first place. We have a chicken and the egg problem here--without an alignment, we have a hard time inferring the tree and without a tree we have a hard time efficiently solving the alignment problem. We therefore seek a coarse solution by trying to establish a reasonable guide tree that may not be a phylogenetic tree but nevertheless still will be useful to obtaining a multiple alignment. One possibility is to first do all C(k, 2) pairwise alignments. The pairwise alignments can be used to compute a k by k pairwise distance matrix (for k strings). While there are many ways define distances between two aligned sequences, a simple one would be to count the number of mismatch and gaps and divide by the length of the alignment. A pairwise distance matrix gives us a view of the relationship of the k strings, which can be used to infer an alignment guide tree. A common procedure is to construct a UPGMA (Unweighted Pair-Group Method with Arithmatic mean) cluster tree.

Clustering is a procedure to group objects together into sets. For example, if we have five objects {1, 2, 3, 4, 5}, we might group them into two sets {1, 4} and {2, 3, 5}. The first set might have biological meaning like "tRNAs" and the second set might have biological meaning like "snoRNAs". Finding the biologically meaningful groupings of a set of objects is a very common and important activity in computational biology. Methods for clustering fall into a broad category called Machine Learning, and more specifically within "unsupervised learning methods". The "unsupervised" refers to the fact that for some cases (like the UPGMA method) we don't have any other external information to help with the clustering. In other cases, we might have some prior information about the grouping; and, in those cases the goal is not so much the groups themselves but a mathematical method for recapitulating the known groupings. Those kinds of methods are called "supervised learning" methods. Unfortunately, due to lack of time we won't treat machine learning in this part and I encourage you to continue exploring the area by yourself. Going back to clustering, clustering can be carried out hierarchically so that sets can be grouped into larger sets. For example, {{1,4},{{2,3},5}} depicts sets within sets, which is meant to be interpreted as "2 and 3 belong together, which in turn belong together with 5" and so on. Clustering can also be represented by tree graphs. The tree below can be interpreted as the nested set of sets {{1,4},{{2,3},5}}:



In fact, the two sets $\{1,4\}$ and $\{2,3,5\}$ can also be seen as a set of sets, $\{\{1,4\},\{2,3,5\}\}$, and the tree:



UPGMA is a method to create hierarchical clustering using a sequential procedure. We start with a pairwise distance matrix, say:

	1	2	3	4	5
1	0	1	4	5	6
2	1	0	5	6	7
3	4	5	0	2	3
4	5	6	2	0	1
5	6	7	3	1	0

The procedure first finds the closest pair and merges them into a single "cluster". In the matrix above, there are two closest pairs, {1, 2} and {4, 5}. We can choose one at random, say 1 and 2 and merge them:

	1-2	3	4	5
1-2	0			
3		0	2	3
4		2	0	1
5		3	1	0

The main question is how to fill in the new distances between the $\{1, 2\}$ cluster and the other ones. The main idea of the "unweighted arithmetic mean" is that the distance between $\{1, 2\}$ and, say, 5 should be an average of the original distances $d_{1,5}$ and $d_{2,5}$. So we would have:

	1-2	3	4	5
1-2	0	4.5	5.5	6.5
3	4.5	0	2	3
4	5.5	2	0	1
5	6.5	3	1	0

In general, if we have two clusters $\{i,j,k...\}$ and $\{a,b,c....\}$, the distance between these two are computed as the average of all the original pairwise distances, $d_{i,a}, d_{i,b}, ...d_{k,a}, ...$ etc. There are other kinds of such sequential clustering methods. The main difference in most of the methods is how they compute the distance between clusters as they try to sequentially collect them into new clusters. For example, one might compute the distance between $\{i,j,k...\}$ and $\{a,b,c...\}$ as the minimum of all of the original pairwise distances (this is called "single-linkage" clustering). The particular cumulative procedure of clustering here is called agglomerative clustering as

opposed to divisive clustering. Sequential divisive clustering is an alternative method where we first try to identify a division of the objects and then successively find more divisions. I should note that for whatever reasons, researchers generally find the agglomerative clustering seems to give more biologically meaningful clustering than divisive clustering. (Why do you think this is?)

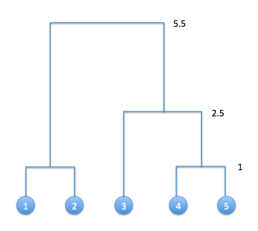
If we continue from the above matrix, we would next find 4 and 5 as the closest pair and we would generate a new matrix:

	1-2	3	4-5
1-2	0	4.5	6
3	4.5	0	2.5
4-5	6	2.5	0

Continuing further:

	1-2	3-4-5
1-2	0	5.5
3-4-5	5.5	0

The sequence of clusters can be now depicted as a tree:



The levels of the tree graph are made proportional to the computed distance between the clusters. The tree on the left is the UPGMA tree of the pairwise distance matrix. One key feature of this tree is that the leaves are all equi-distant from the root of the tree (assume that there is a root along the horizontal line at the very top and also assume that the distance between clusters are used to weight the edges). Such a tree is called an ultrametric tree for reasons we will discuss later. For now, it suffices to say that most trees we see later will have edge weights such that the tree is not ultrametric.

Going back to the multiple alignment problem, we can compute the k by k distance matrix from pairwise alignments and then compute a clustering using the distance matrix. The most common

clustering is UPGMA and we can use the resulting tree as the alignment guide tree (thus the name CLUSTAL). Suppose we have five strings and the strings resulted in the UPGMA tree shown above. We use this guide tree to align string 1 and 2 together and string 4 and 5 together. Now what? Well, the guide tree tells us to align string 3 with the string 4-5 alignment. The second key part of the CLUSTAL is in how such next-level alignment is done. The idea now is to use something called a profile alignment. Suppose we have a set **P** of aligned strings on one side and another set **Q** of aligned strings on the other side. To compute the alignment between the set **P** and **Q**, let **P**[i] and **Q**[j] be the ith and jth aligned positions in the aligned strings **P** and **Q** respectively. Note that **P**[i] refers to a whole column of symbols. For example, if the aligned set **P** is:

ACCTG T-CAG

-CCAG

AC-TG

Then, P[2] = [C-CC]' where I use the notation []' to mean transpose of the column of symbols. Similarly, let Q[3] = [TT-T-T]'. Now when we are computing a dynamic programming table for P and Q, we might be

comparing P[2] against Q[3], then we compute the cost by computing the average cost of all pairwise symbol comparisons. Thus, in the UPGMA alignment tree above, we will first compute the alignment of 1 and 2; 4 and 5. Then, we will compute the alignment of 3 against $\{4, 5\}$ by using this profile alignment procedure. Then we will finally align $\{1,2\}$ against $\{3, \{4, 5\}\}$ by profile alignment again.

There are many possible ways to "improve" on this scheme. We might compute the guide tree in a different manner. One of the popular ways to compute the guide tree is using a distance-based approximate phylogeny algorithm called Neighbor-Joining. We will learn this later. Another way is to compute the distance matrix not with a pairwise alignment but with a different way of comparing two sequences. For example, a by using local alignment or say a k-mer comparison. K-mer comparison is popular for comparing two or more long strings. In this procedure, we consider all possible k-mers of some fixed k. Then we score each string by the frequency of the k-mers. Strings can be compared by comparing the frequency of the k-mers. Other variations include different ways to compute alignments between two sets of aligned sequences.

I put quotations around the word "improve" because, for the most part we don't know whether any of the variations actually improve the alignments in the sense that they help accurately infer the positional homology relationship of the strings. Or, at least, it would be nice if the procedures led to optimizing the phylogenetic tree alignment problem; but we don't know whether CLUSTAL does that very well either. This leads us to a discussion of quality of solution to problems. As mentioned repeatedly, there are two different kinds of solutions: (1) solution to the biological problem; (2) solution to an optimization problem, which is motivated by the biological problem. Unless we know the true biological solution, it is very hard to evaluate how well an algorithm does in solving the biological problem. For sequence alignments there are databases of alignments that are claimed to be "true" alignments. Typically, such alignments are generated by hand curation using human intuition and other accessory information like the 3D structure of proteins that is coded by the sequences. More commonly, algorithms can be tested against simulations of sequence evolution, where the simulation might include all kinds of mutational edits that can happen to a sequence. For the most part, when we have biological problems for which we design a computational solution, we do not know the true answer. Therefore, we have very limited ability to test the algorithms. Often there is some "first-to-market" computational solution like CLUSTAL, which works in a satisfactory manner to the researcher and then a lot of refinements come along that are hard to evaluate. Sometimes the results can be evaluated by empirical tests. Unfortunately, a quark of biology is that what works in one situation usually do not generalize to everything else. Therefore, designing and using any computational algorithm requires a bit of theoretical insight (i.e., is the design principle of the algorithm reasonable) and a lot of biological intuition.

Compared to the biological solution, the solution to the optimization problem is well defined. For example, for the multiple sequence alignment problem, we might consider something like the sum-of-pairs alignment score or say the phylogenetic tree alignment score (the best alignment score over all possible binary trees and all possible ancestral sequences). Such an optimization may be exactly solved, which may be possible for small problems. Or, we might solve it with an approximation algorithm. An approximation algorithm is a procedure that may not exactly solve the optimization problem but approximately solves the problem with a known bound on the degree of approximation. For example, we might talk of a 1.5-approximation algorithm, by which we mean an algorithm that is guaranteed to be no worse than 1.5 times the optimal solution. Approximation algorithms can be very difficult to construct because of the need to guarantee the approximation bound. Finally, we might solve the problem with a heuristic algorithm. A heuristic algorithm tries to solve the optimization problem with reasonable sounding procedures but typically cannot be analyzed to give an exact bound on their performance. For example, the UPGMA+profile alignment procedure of CLUSTAL might be seen as a heuristic algorithm for sum-of-pairs multiple alignment optimization problem. We discussed previously many very difficult problems that are NP-complete or NP-hard. A key interest of theoretical computer science is to derive a polynomial time approximation algorithms for these problems with good bounds. In computational biology practice, most problems are very hard, say NP-hard, and good approximation algorithms are not available so much of the practice involves designing heuristic algorithms. Unfortunately, then it is difficult to understand the performance

of the algorithms and therefore, we seek either "gold-standard" datasets with known (or generally agreed upon) solutions or reasonable simulations with known solutions to test the algorithms.