# Protein sequence comparison and Protein evolution Tutorial - ISMB98 

William R. Pearson*<br>Department of Biochemistry,<br>Jordan Hall,\#440<br>University of Virginia, Charlottesville, VA 22908, USA

June, 1998

## Contents

1 Introduction ..... 2
1.1 Evolutionary time scales ..... 4
1.2 Similarity, Ancestry and Structure ..... 7
1.3 Modes of Evolution ..... 8
1.3.1 Conventional divergence from a common ancestor ..... 9
1.3.2 Sequence similarity and homology, the $\mathrm{H}^{+}$ATPase ..... 10
1.3.3 Protein families diverge at different rates ..... 12
1.3.4 Mosaic proteins ..... 18
1.4 Introns Early/Late ..... 18
1.5 DNA vs Protein comparison ..... 20
2 Alignment methods ..... 22
2.1 Algorithms ..... 22
2.2 Dynamic Programming Algorithms ..... 25
2.3 Scoring methods ..... 27

[^0]2.4 Heuristic Algorithms ..... 28
2.4.1 BLAST ..... 28
2.4.2 FASTA ..... 30
3 The statistics of sequence similarity scores ..... 31
3.1 Sequence alignments without gaps ..... 32
3.2 Similarity scores increase with sequence length ..... 32
3.3 Empirical statistics for alignments with gaps ..... 32
3.4 Statistical significance by random shuffling ..... 33
4 Identifying distantly related protein sequences ..... 35
4.1 Serine proteases ..... 35
4.2 Glutathione S-transferases ..... 41
4.3 G-protein-coupled receptors ..... 42
5 Repeated structures in proteins ..... 45
6 Summary ..... 47
References ..... 49
7 Suggested Reading ..... 51
7.1 General Protein evolution ..... 51
7.1.1 Introns Early/Late ..... 51
7.2 Alignment methods ..... 51
7.2.1 Algorithms ..... 51
7.2.2 Scoring methods ..... 52
7.3 Evaluating matches - statistics of similarity scores ..... 52

## 1 Introduction

The concurrent development of molecular cloning techniques, DNA sequencing methods, rapid sequence comparison algorithms, and computer workstations has revolutionized the role of biological sequence comparison in molecular biology. As a result, the role of protein sequence data in molec-
ular biology and biochemistry has dramatically changed. Twenty-five years ago, protein sequence determination was usually one of the last steps in the characterization of a protein. Now the process is reversed, so that it is common to clone and sequence a gene of biological interest-e.g., one that is induced by serum stimulation, or a developmental change, or a chromosomal rearrangement associated with a disease. This is the fundamental premise of the human genome project-that one can first sequence all the genes in an organism and then infer their function by sequence analysis.

Today, the most powerful method for inferring the biological function of a gene (or the protein that it encodes) is by sequence similarity searching on protein and DNA sequence databases. With the development of rapid methods for sequence comparison, both with heuristic algorithms and powerful parallel computers, discoveries based solely on sequence homology have become routine. One of the more dramatic discoveries was the identification of a new tumor suppressor gene in humans that is related to yeast and E. coli DNA repair enzymes. This discovery, the result of a similarity search, both told the investigators that they had identified the appropriate gene and demonstrated clearly the nature of the oncogenic mutation. As entire genomes from bacteria, yeast, and simple eukaryotes become available, protein sequence comparison will become an even more powerful tool for understanding biological function.

Protein sequence comparison is our most powerful tool for characterizing protein sequences because of the enormous amount of information that is preserved throughout the evolutionary process. For many protein sequences, an evolutionary history can be traced back 1-2 billion years. Proteins that share a common ancestor are called homologous. Sequence comparison is most informative when it detects homologous proteins. Homologous proteins always share a common three-dimensional folding structure and they often share common active sites or binding domains. Frequently homologous proteins share common functions, but sometimes they do not. Our ability to characterize the biological properties of a protein based on sequence data alone stems almost exclusively from properties conserved through evolutionary time. Predictions of common properties for non-homologous proteinssimilarities that have arisen by convergence- are much less reliable.

This tutorial examines how the information conserved during the evolution of a protein molecule can be used to infer reliably homology, and thus a shared protein fold and possibly a shared active site or function. We will start by reviewing a geological/evolutionary time scale. Many protein sequences can be used to infer reliably events that happened more than a billion years ago. Remarkably, some protein sequences change so slowly that they could be used to "date" events that took place more than 5 billion years ago, had the proteins existed. Next we will look at the evolution of several protein families. During the tutorial, these families will be used to demonstrate that homologous protein ancestry can be inferred with confidence. We will also examine different modes of protein evolution and consider some hypotheses that have been presented to explain the very earliest events in protein evolution.

The next part of the tutorial will examine the technical aspects of protein sequence comparison. Both optimal and heuristic algorithms and their associated parameters that are used to characterize protein sequence similarities are discussed. Perhaps more importantly, we will survey the statistics of local similarity scores, and how these statistics can both be used to improved the selectivity of a search and to evaluate the significance of a match.

We will then examine distantly related members of three protein families, the serine proteases, the glutathione transferases, and the G-protein-coupled receptors (GCRs). The serine proteases are used
to emphasize that even when a highly conserved motif is found throughout a family, similarity extends over a much longer region. The glutathione transferases and GCRs are very diverse families whose members frequently do not share significant pair-wise similarity. The relative strengths of strategies to characterize such relationships will be examined.

Finally, we will discuss how sequence similarity can be used to examine internal repeated or mosaic structures in proteins. Such repeated structures can arise from either divergence-calmodulin EF-hand repeats and EGF-domains-or convergence-tropomyosin and transcription factor coiledcoil.

This tutorial is directed towards examining protein evolution. Most of the algorithms and methods that are applied to protein evolution can be used with DNA sequences as well. However, in general, DNA sequence comparisons are far far less informative than protein sequence comparisons (see Fig. 8). DNA sequences that do not encode proteins or structural RNAs (e.g. ribosomal RNAs) diverge very rapidly, so that it is usually difficult to detect reliably non-coding DNA sequence homologies for sequences that diverged more than 200 million years ago. In contrast, even the most rapidly changing protein sequences can detect sequences that are 200 million years old; typically protein sequence comparisons detect sequences that diverged 1 billion years ago. Thus, the most important lesson from this tutorial is, when searching sequence databases for homologous sequences, to use protein sequences whenever possible.

### 1.1 Evolutionary time scales

When we search for homologous proteins, we are trying to identify proteins that shared a common ancestor in the past. Fig. 1 shows a general evolutionary tree that reaches back to the beginning of the earth's history. The goal of protein sequence comparison is to take a protein sequence, for example from a human chromosome, and search a protein database to find homologous sequences, often from very divergent organisms. Thus, if the similarity search produces significant matches with a protein found in yeast, then an ancestral protein must have existed in an organism at least 1 billion years ago and that the descendants of that organism preserved the sequence in modern day humans and yeast. Likewise, if a yeast protein is homologous to one found in E. coli, that sequence must have existed in 2 billion years ago in the primordial organism that gave rise to bacteria and fungi.

When we examine protein or DNA sequences, we are almost always studying modern (present day) sequences. Thus, it does not make any sense to say that a yeast or bacterial sequence is more primitive than a mammalian sequence; all sequences are contemporary. As we will see later, however, there are examples of sequences that are found only in vertebrates, or only in animals or plants but not both. Such sequences are less ancient than those found both in mammals and bacteria.

For organisms that diverged within the past 600 My (million years), inferences about divergence times for modern organisms are taken from geological data; more ancient divergence times are inferred from extrapolations of evolutionary "clocks." Evolutionary clocks are based both on slowly changing protein sequences and on ribosomal RNA sequences; such divergence time estimates require a rate of change that is constant on average. The oldest fossils are of prokaryotes in rocks about 2.5 billion years old; this geological age is consistent with that inferred from evolutionary divergence rates.

Table 1 summarizes some important milestones in evolutionary time, and, when considered with

Figure 1: The tree of life


Adapted from Dayhoff et al., 1978.

Table 1: Some Important dates in history

| Origin of the universe | $-12^{a}$ | $\pm 2$ |
| :--- | :--- | :--- |
| Formation of the solar system | -4.6 | $\pm 0.4$ |
| First self-replicating system | -3.5 | $\pm 0.5$ |
| Prokaryotic-eukaryotic divergence | -1.8 | $\pm 0.3$ |
| Plant-animal divergence | -1.0 |  |
| Invertebrate-vertebrate divergence | -0.5 |  |
| Mammalian radiation beginning | -0.1 |  |

${ }^{a}$ Billions of years. From Doolittle et al., 1986.

Table 2, gives a better perspective on the evolutionary horizons provided by different protein families. The theoretical lookback times in Table 2 are based on the assumption that one can identify proteins that share about $20 \%$ sequence identity throughout their entire length. It will be clear from later examples that if two protein sequences share $25 \%$ identity across their lengths, they are homologous, and that in some cases, convincing evidence of common ancestry can be deduced from similarities as low as $20 \%$. These look-back times can be confirmed in practice; for example, with sensitive sequence comparison algorithms, significant similarity between plant and animal globins can be found.

Table 2: Evolutionary Horizons

| Protein | PAMs $^{a} / 100$ residues <br> $/ 10^{8}$ <br> years | Theoretical <br> Lookback time | Horizon |
| :--- | :---: | :---: | :--- |
| Pseudogenes | 400 | $45^{c}$ | Primates, Rodents |
| Fibrinopeptides | 90 | 200 | Mammalian Radiation |
| Lactalbumins | 27 | 670 | Vertebrates |
| Ribonucleases | 21 | 850 | Animals |
| Hemoglobins | 12 | $1.5^{d}$ | Plants/Animals |
| Acid Proteases | 8 | 2.3 | Prokayrotic/Eukarotic |
| Triosphosphate isomerase | 3 | 6 | Archaen |
| Glutamate dehydrogenase | 1 | 18 |  |
|  |  |  |  |

[^1]Figure 2: Structural similarity in related proteins - serine proteases


Expectation values $(\mathrm{E}())$, percent identity, the length of the alignment are shown with respect to bovine trypsin. The last two numbers report the length of the alignment and the length of the library sequence whose structure is shown.

### 1.2 Similarity, Ancestry and Structure

The inference of homology - common ancestry - is the most powerful conclusion that one can draw from a similarity search because homologous proteins share similar three-dimensional structures. This can be seen in Fig. 2, where the structures of three members of the serine protease superfamily are
shown. Two of these proteins, bovine chymotrypsin and S. griseus trypsin, share strong sequence similarity while the third related sequence, S. griseus protease A, does not share significant similarity $(\mathrm{E}()<66)$ yet the protein has a very similar structure. Thus, as will be seen throughout this chapter, homologous proteins need not share statistically significant, or even detectable, sequence similarity.

Endochitinase is an example of a very high-scoring, but unrelated protein whose structure is known. This high scoring unrelated sequence does not share any structural similarity with trypsin or other serine proteases. If two proteins are not homologous, one cannot draw any conclusion about their structural similarity, even though they may have high similarity scores.

### 1.3 Modes of Evolution

Figure 3: Orthologous sequences - The cytochrome 'c' family


Cytochrome 'c's comprise a family of orthologous proteins that are found in all organisms. The sequences on this tree are orthologous - two cytochrome 'c's are different because they are in different species.

### 1.3.1 Conventional divergence from a common ancestor

Homologous sequences can be divided into two groups: (1) orthologous sequences - sequences that differ because they are found in different species; and (2) paralogous sequences - sequences that differ because of a gene duplication event. Fig. 3 shows an evolutionary tree for orthologous cytochrome ' $c$ ' sequences. The branching pattern, which reflects the differences between cytochrome ' $c$ ' sequences, matches the evolutionary relationships of the species that express the proteins.

Figure 4: Orthology and paralogy - The globin family


Members of the globin oxygen binding protein family have evolved through a series of gene duplications and speciation events. The human $\alpha$ and $\delta$ genes duplicated less than 50 Mya ( $\delta$ chains are found in primates, but not in other mammals).

In general, the organismal tree and the sequence tree will not match if the sequences are paralogous. Members of the globin oxygen binding protein family are both orthologous - they differ because of speciation - and paralogous p - they differ because of gene duplications. Thus, human $\alpha$-globin, mouse $\alpha$-globin, and chicken $\alpha$-globin are all orthologs, they differ because of the speciation events that gave rise to humans, rodents, and birds. Mouse $\beta$ globin and human $\alpha$ globin are paralogous; they differ because of a gene duplication that created the $\alpha$ and $\beta$ subunits some 600 Mya
(million years ago). An evolutionary tree based on human $\alpha$, chicken $\alpha$, and mouse $\beta$ would imply that humans are more closely related to chickens than to mice. While such a mistake is unlikely in a well-studied family like the globins, it can be quite common in large, diverse, and poorly characterized families like the G-protein-coupled receptors (Fig. 19).

### 1.3.2 Sequence similarity and homology, the $\mathbf{H}^{+}$ATPase

Our first example of the significant sequence similarity shared by homologous proteins will use one of the chains of the $\mathrm{H}^{+}$-ATPase, or proton-pump, used to convert energy to ATP in the mitochrondria and chloroplasts of aerobic organisms.

Figure 5: The PAM250 matrix

| Cys | 12 |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Ser | 0 | 2 |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| Thr | -2 | 1 | 3 |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| Pro | -1 | 1 | 0 | 6 |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| Ala | -2 | 1 | 1 | 1 | 2 |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| Gly | -3 | 1 | 0 | -1 | 1 | 5 |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| Asn | -4 | 1 | 0 | -1 | 0 | 0 | 2 |  |  |  |  |  |  |  |  |  |  |  |  |  |
| Asp | -5 | 0 | 0 | -1 | 0 | 1 | 2 | 4 |  |  |  |  |  |  |  |  |  |  |  |  |
| Glu | -5 | 0 | 0 | -1 | 0 | 0 | 1 | 3 | 4 |  |  |  |  |  |  |  |  |  |  |  |
| Gln | -5 | -1 | -1 | 0 | 0 | -1 | 1 | 2 | 2 | 4 |  |  |  |  |  |  |  |  |  |  |
| His | -3 | -1 | -1 | 0 | -1 | -2 | 2 | 1 | 1 | 3 | 6 |  |  |  |  |  |  |  |  |  |
| Arg | -4 | 0 | -1 | 0 | -2 | -3 | 0 | -1 | -1 | 1 | 2 | 6 |  |  |  |  |  |  |  |  |
| Lys | -5 | 0 | 0 | -1 | -1 | -2 | 1 | 0 | 0 | 1 | 0 | 3 | 5 |  |  |  |  |  |  |  |
| Met | -5 | -2 | -1 | -2 | -1 | -3 | -2 | -3 | -2 | -1 | -2 | 0 | 0 | 6 |  |  |  |  |  |  |
| Ile | -2 | -1 | 0 | -2 | -1 | -3 | -2 | -2 | -2 | -2 | -2 | -2 | -2 | 2 | 5 |  |  |  |  |  |
| Leu | -6 | -3 | -2 | -3 | -2 | -4 | -3 | -4 | -3 | -2 | -2 | -3 | -3 | 4 | 2 | 6 |  |  |  |  |
| Val | -2 | -1 | 0 | -1 | 0 | -1 | -2 | -2 | -2 | -2 | -2 | -2 | -2 | 2 | 4 | 2 | 4 |  |  |  |
| Phe | -4 | -3 | -3 | -5 | -4 | -5 | -4 | -6 | -5 | -5 | -2 | -4 | -5 | 0 | 1 | 2 | -1 | 9 |  |  |
| Tyr | 0 | -3 | -3 | -5 | -3 | -5 | -2 | -4 | -4 | -4 | 0 | -4 | -4 | -2 | -1 | -1 | -2 | 7 | 10 |  |
| Trp | -8 | -2 | -5 | -6 | -6 | -7 | -4 | -7 | -7 | -5 | -3 | 2 | -3 | -4 | -5 | -2 | -6 | 0 | 0 | 17 |
|  | C | S | T | P | A | G | N | D | E | Q | H | R | K | M | I | L | V | F | Y | W |

The similarity scores in Figs. 6-8 were calculated using the Smith-Waterman algorithm (Smith \& Waterman, 1981,Sec. 2.2), a method that guarantees to calculate the best (optimal) score between any two protein or DNA sequences, given a scoring matrix and gap penalties. Fig. 5 shows the PAM250 matrix, which was developed almost 20 years ago by Dayhoff and her colleagues (Dayhoff et al., 1978). The PAM250 matrix, or modern versions such as the BLOSUM50 matrix used here, incorporates information about the likelihood that one amino-acid will be mutated into another over evolutionary time. Thus, changes that are very unlikely to occur in evolution, for example the substitution of the very small glycine residue for the very large tryptophan residue, are given large negative scores ( -7 in Fig. 5), while conservative changes, such as the substitution of lysine by arginine (both have basic side chains), are given positive scores $(+3)$. The scores for identical matches also vary in the PAM250 matrix, depending on whether the amino-acids are common (e.g. serine and methionine), and thus likely to be aligned by chance, or rare (e.g. cysteine and tryptophan). There is a well-developed statistical theory for substitution matrices (Altschul, 1991), which will be discussed in section 2.3.

Table 3 reports similarity scores and their statistical significance from a search of the PIR annotated

Figure 6: Searching with human ATP-ase, similarity scores

|  | opt | E () |  |
| :---: | :---: | :---: | :---: |
| < 20 | 17 | $0:=$ | one $=$ represents 22 library sequences |
| 22 | 0 | 0 : |  |
| 24 | 0 | 0 : |  |
| 26 | 2 | 0 : $=$ |  |
| 28 | 7 | 3:* |  |
| 30 | 7 | 18:* |  |
| 32 | 45 | $68:===$ * |  |
| 34 | 166 | $184:========*$ |  |
| 36 | 337 | 379: ======== | $====$ * |
| 38 | 581 | $626:========$ | $==============$ * |
| 40 | 869 | 873: ========= | $========================$ * |
| 42 | 1009 | 1067:========= | $=============================*$ |
| 44 | 1276 | 1177: ========= | $===========$ |
| 46 | 1253 | 1198: ======== | = |
| 48 | 1199 | 1147: ========= | $===================================$ * |
| 50 | 1032 | 1047: ========= | $==============================$ * |
| 52 | 949 | 920: ========= | $=========================$ * $==$ |
| 54 | 838 | $786:=========$ | $====================*===$ |
| 56 | 578 | 657:======== | $=============$ * |
| 58 | 467 | 539: ========= | $===========$ * |
| 60 | 393 | 437: ========= | $=======$ * |
| 62 | 339 | 350 : ========= | $====$ * |
| 64 | 276 | 278: ========= | = * |
| 66 | 214 | $220:=========$ * |  |
| 68 | 188 | 173: =======*= |  |
| 70 | 140 | 136: ======* |  |
| 72 | 131 | 106: ====* $=$ |  |
| 74 | 88 | 83: ===* |  |
| 76 | 71 | $64:==*=$ |  |
| 78 | 48 | $50:=$ * |  |
| 80 | 43 | 39: =* |  |
| 82 | 38 | $30:=*$ |  |
| 84 | 27 | 24 : =* |  |
| 86 | 21 | 18:* |  |
| 88 | 15 | 14:* |  |
| 90 | 17 | 11:* |  |
| 92 | 7 | 8: * | :=======* = represents 1 library sequence |
| 94 | 22 | 7:* | $===$ * $==============$ |
| 96 | 3 | 5 : * | : === * |
| 98 | 8 | 4:* | : ===*==== |
| 100 | 6 | 3:* | : ==*=== |
| 102 | 5 | 2:* | : =*=== |
| 104 | 9 | 2:* | : =*======= |
| 106 | 4 | 1:* | : *=== |
| 108 | 5 | 1:* | : * $====$ |
| 110 | 4 | 1:* | : *=== |
| 112 | 4 | 1:* | : *=== |
| 114 | 4 | 1:* | : *=== |
| 116 | 6 | $0:=$ | * $======$ |
| 118 | 1 | $0:=$ | * $=$ |
| >120 | 32 | $0:=$ | $=$ |

protein sequence database (PIR1, release 44, March, 1995) using the human $\mathrm{H}^{+}$-ATPase as a query sequence. There is excellent agreement between the expected and actual distributions of similarity scores. In this search, all of the library sequences related (homologous) to the query sequence obtained scores higher than any of the unrelated sequences. However, a number of unrelated sequences obtained very high scores; 10 of the 32 sequences with $z$-scores $>120$ ( 7 standard deviations above the mean ${ }^{1}$ ) are not members of the $\mathrm{H}+$-ATPase family.

Fig. 6 shows the distribution of similarity scores between human $\mathrm{H}^{+}$-ATPase (PIR entry PWHU6) and each protein sequence in the PIR1 (rel. 44) database. The ' $=$ ' symbols in the histogram show the distribution of normalized similarity scores calculated during the search, thus, 393 sequences in the PIR1 library had scores of 60 or 61 . The ' $*$ ' symbols report the expected number of sequences with the indicated range of scores for a search of a database of this size, based on random chance. The basis for the statistical estimates will be discussed in section 3 .

While Table 3 shows that all of the members of this family have siginificant similarity with the human enzyme, Fig. 7 gives a more realistic perspective of the family's evolutionary history by considering every possible pairwise alignment. When the $E$. coli enzyme is used to search the database for related $\mathrm{H}^{+}$-ATPases, the ranking of the different sequences changes, but sequences distant from the $E$. coli sequence have more significant similarities than those distant from the human sequence.

### 1.3.3 Protein families diverge at different rates

For many protein families with a variety of divergence rates, the rate of change over evolutionary time is relatively constant. These rates can be used to date the divergence events (e.g. plants and animals) that occurred more than 600 Mya and thus do not have a fossil record. However, different protein families diverge at different rates, so that, in general, the number of differences between a pair of sequences cannot be used to estimate the time the two sequences diverged. This is particularly true for paralogous sequences; once a sequence has duplicated, it may change very rapidly before selective pressure on its new function slows its rate of change. Thus, in Table 4 there are several members of growth hormone superfamily-growth hormone, sommatotropin, and prolactin-with different divergence rates.

[^2]Table 3: Searching with human ATP-ase, high-scoring sequences

| The best scores are: | s -w | z-score | $\mathrm{E}(12805)$ | $\%$ | len |  |
| :--- | :--- | ---: | :--- | :--- | :--- | ---: |
| PWHU6 | H+-trans. ATP synth.-human mito. | 1400 | 1767.8 | $10^{-92}$ | 100.0 | 226 |
| PWBO6 | H+trans. ATP synth.-bovine mito. | 1157 | 1460.9 | $10^{-75}$ | 77.9 | 226 |
| PWMS6 | H+trans. ATP synth.-mouse mito. | 1118 | 1411.6 | $10^{-72}$ | 75.7 | 226 |
| PWXL6 | H+-trans. ATP synth.-frog mito. | 745 | 940.6 | $10^{-46}$ | 53.3 | 226 |
| PWFF6Y | H+-trans. ATP synth.-fruit fly mito. | 473 | 597.1 | $10^{-27}$ | 37.8 | 222 |
| PWFF6 | H+-trans. ATP synth.-fruit fly mito. | 471 | 594.6 | $10^{-26}$ | 37.5 | 224 |
| PWBY3 | H+-trans. ATP synth.-yeast mito. | 438 | 551.7 | $10^{-25}$ | 36.2 | 232 |
| PWAS6N | H+-trans. ATP synth.-aspergillus mito. | 365 | 459.6 | $10^{-19}$ | 30.4 | 230 |
| PWKQ6 | H+-trans. ATP synth.-Cochliobolus mito. | 353 | 444.4 | $10^{-18}$ | 31.3 | 214 |
| PWWT6 | H+-trans. ATP synth.-wheat mito. | 309 | 385.4 | $10^{-15}$ | 28.9 | 235 |
| PWNT6M | H+-trans. ATP synth.-tobacco mito. | 309 | 385.2 | $10^{-15}$ | 28.3 | 233 |
| PWZM6M | H+trans. ATP synth.-corn mito. | 283 | 355.0 | $10^{-15}$ | 31.1 | 291 |
| LWEC6 | H+trans. ATP synth.-E. coli | 178 | 223.0 | $10^{-6}$ | 23.3 | 236 |
| LWRZ6 | H+-trans. ATP synth.-rice chloro. | 144 | 180.8 | 0.00037 | 24.2 | 231 |
| PWPMA6 | H+-trans. ATP synth.-pea chloro. | 143 | 179.5 | 0.00044 | 25.0 | 232 |
| PWYBAA | H+-trans. ATP synth.-Synechocystis | 142 | 177.3 | 0.00058 | 26.5 | 170 |
| PWSPA6 | H+-trans. ATP synth.-spinach chloro. | 138 | 173.2 | 0.00098 | 24.2 | 231 |
| PWYCA6 | H+-trans. ATP synth.-cyanobacteria | 127 | 158.9 | 0.0062 | 26.3 | 167 |
| LWNT6 | H+-trans. ATP synth.-tobacco chloro. | 126 | 158.1 | 0.0069 | 22.1 | 231 |
| LWLV6 | H+-trans. ATP synth.-Marchiantia chloro. | 126 | 158.0 | 0.0069 | 24.0 | 167 |
| PWEGAC | H+-trans. ATP synth.-Euglena chloro. | 123 | 154.1 | 0.011 | 25.7 | 214 |
| S17420 | ubiquinol-cytochrome-c reductase | 113 | 138.0 | 0.09 | 23.4 | 158 |
| S17418 | ubiquinol-cytochrome-c reductase | 108 | 131.7 | 0.20 | 24.5 | 208 |
| QXBO2MM | NADH dehydrogenase (ubiquinone) | 107 | 131.2 | 0.22 | 26.1 | 211 |
| S17415 | ubiquinol-cytochrome-c ceductase | 105 | 127.9 | 0.33 | 27.7 | 137 |
| DNHUN2 | NADH dehydrogenase (ubiquinone) | 103 | 126.1 | 0.41 | 20.1 | 149 |
| QRECAA | amino acid trans. protein-E. Coli | 104 | 125.1 | 0.47 | 23.4 | 111 |
| CBHU | ubiquinol-cytochrome-c reductase | 102 | 124.1 | 0.53 | 26.8 | 205 |
| S17419 | ubiquinol-cytochrome-c reductase | 101 | 122.9 | 0.63 | 23.4 | 158 |
| S17407 | ubiquinol-cytochrome-c reductase | 99 | 120.3 | 0.87 | 23.6 | 140 |
| QQBEN5 | integral membrane protein-saimiriine herp | 98 | 119.4 | 0.99 | 20.8 | 202 |

The horizontal line indicates the separation been the lowest scoring related sequences and the highest scoring unrelated sequence.

Figure 7: Phylogeny of $\mathrm{H}^{+}$-ATPases


An evolutionary tree of $\mathrm{H}^{+}$-ATPases (subunit 6). Sequences were aligned using the GCG PILEUP program, distances calculated using the GCG DISTANCES program, and the tree constructed using the Neighbor-Joining algorithm (GCG GROWTREE). Expectation values from a search with the human $\mathrm{H}^{+}$-ATPase (PWHU6, Table 3) and a search with the E. coli sequence are shown.

Figure 8: Searching with human ATPase, high-scoring alignments


Alignments of human $\mathrm{H}^{+}$-ATPase with the E. coli homologue and a plant chloroplast homologue. Despite the considerable evolutionary distance (both sequences diverged at least 2 Bya ), the pairs of sequence share more than $20 \%$ identity across almost their entire lengths. ':' symbols denote identities; ' .' denote conservative substitutions. Searches were performed with the BLOSUM50 matrix and gap penalties of $-12 /-2$.

Table 4: Rates of change in protein families

| Protein | Rate $^{a}$ | Protein | Rate |
| :--- | ---: | :--- | ---: |
| Fibrinopeptides | 90 | Thryrotropin beta chain | 7.4 |
| Growth hormone | 37 | Parathyrin | 7.3 |
| Ig kappa chain C region | 37 | Parvalbumin | 7.0 |
| Kappa casein | 33 | BPTI Protease inhibitors | 6.2 |
| Ig gamma chain C region | 31 | Trypsin | 5.9 |
| Lutropin beta chain | 30 | Melanotropin beta | 5.6 |
| Ig lambda chain C region | 27 | Alpha crystallin A chain | 5.0 |
| Complement C3a | 27 | Endorphin | 4.8 |
| Lactalbumin | 27 | Cytochrome b | 4.5 |
| Epidermal growth factor | 26 | Insulin | 4.4 |
| Somatotropin | 25 | Calcitonin | 4.3 |
| Pancreatic ribonuclease | 21 | Neurophysin 2 | 3.6 |
| Lipotropin beta | 21 | Plastocyanin | 3.5 |
| Haptoglobin alpha chain | 20 | Lactate dehydrogenase | 3.4 |
| Serum albumin | 19 | Adenylate cyclase | 3.2 |
| Phospholipase A |  | 2.8 |  |
| Protease inhibitor PST1 type | 19 | Triosephosphate isomerase | 2.6 |
| Prolactin | 17 | Corticotropin | 2.5 |
| Pancreatic hormone | 17 | Glyceraldehyde 3-P DH | 2.2 |
| Carbonic anhydrase C | 16 | Cytochrome C | 2.2 |
| Lutropin alpha chain | 16 | Plant ferredoxin | 1.9 |
| Hemoglobin alpha chain | 12 | Collagen | 1.7 |
| Hemoglobin beta chain | 12 | Troponin C, skeletal muscle | 1.5 |
| Lipid-binding protein A-II | 10 | Alpha crystallin B-chain | 1.5 |
| Gastrin | 9.8 | Glucagon | 1.2 |
| Animal lysozyme | 9.8 | Glutamate DH | 0.9 |
| Myoglobin | 8.9 | Histone H2B | 0.9 |
| Amyloid A | 8.7 | Histone H2A | 0.5 |
| Nerve growth factor | 8.5 | Histone H3 | 0.14 |
| Acid proteases | 8.4 | Ubiquitin | 0.1 |
| Myelin basic protein | 7.4 | Histone H4 |  |
|  |  |  | 2.5 |

${ }^{a}$ percent/100 My
From (Nei, 1987; Dayhoff et al., 1978)

Figure 9: The limits of sequence similarity


### 1.3.4 Mosaic proteins

"Conventional" protein families, e.g. the globins, cytochrome 'c's, $\mathrm{H}^{+}$-ATPases, in which protein sequences have diverged from a common ancestor in a direct fashion, typically with only modest changes in the length of the sequence, have been known for more than 30 years. In the past 10 years, a more complex type of protein evolution has been observed-proteins that contain multiple domains from other proteins. These proteins have been called "mosaic" proteins; the domains are frequently inserted through a process called "exon shuffling." Table 7 lists a number of human proteins that are comprised of mosaic domains, but such proteins are not limited to mammals. Similar mosaic structures are common in DNA binding proteins, both in bacteria and eukaryotes.

Table 5: Classification of Protein Families
I. Ancient Proteins
A. First editions. Direct-line descendacy to human and contemporary prokaryotes. Mostly mainstream metabolism enzymes. Example: triosphosphate isomerase ( $44.8 \%$ identical over 250 aa, $\left.\mathrm{E}(59000)<10^{-36}\right)$.
B. Second edition. Homologous sequences in human and prokaryotic proteins, but apparently different functions. Example: human glutathione reductase and pseudomonas mercury reductase ( $31 \%$ identical over 438 aa, $\left.\mathrm{E}(59000)<10^{-32}\right)$.
II. Middle-age proteins. Proteins found in most eukaryotes but prokaryotic counterparts are unknown. Example: actin (human and yeast share $88 \%$ identical over 375 aa, E()$<10^{-145}$, other yeast actin homologs share as little as little as $26.4 \%$ over 489 aa, E()$<10^{-14}$.
III. Modern proteins
A. Recent vintage. Proteins found in animals or plants but not both. Not found in prokaryotes. Example: collagen.
B. Very recent inventions. Proteins found in vertebrates but not elsewhere. Example: plasma albumin.
C. Recent mosaics. Modern proteins clearly the result of exon shuffling. Example: LDL receptor.

From Doolittle et al., 1986.

### 1.4 Introns Early/Late

The occurrence of mosaic proteins and the discovery of the "exon/intron" structure of genes in the late 1970's led several investigators to suggest that the exon structure of genes reflected the construction of proteins from modular domains (Gilbert \& Glynias, 1993). While acceptance of this proposal is quite widespread, it is based on very little data. There is no question that many modern mosaic proteins are constructed by a process of "exon-shuffling" whereby exons from other genes have been combined to build new structures. In addition, for some proteins exons are associated with well defined structural elements. The association of exons with structural elements may reflect and ancient construction of

Table 6: Ancient human proteins

| A. First edition type |  |  |  |
| :---: | :---: | :---: | :---: |
| Human protein | Prokaryotic homologue | \% identity | $\mathrm{E}(59,000)$ |
| Triosephosphate isomerase | E. coli | 46 | $<10^{-36}$ |
| Phosphoglyceraldehylde dehydrogenase | B. stearothermophilus | 52 | $<10^{-78}$ |
| Alkaline phosphatase | E. coli | 28 | $<10^{-20}$ |
| Dihydrofolate reductase | E. coli | 28 | $<10^{-6}$ |
| Superoxide dismutase ( $\mathrm{Cu}-\mathrm{Zn}$ ) | E. coli | 32 | $<10^{-7}$ |
| Hypoxanthine-guanine phosphoribosyl transferase | E. coli | 34 | $<10^{-17}$ |
| B. Second edition type |  |  |  |
| Glutathione reductase | Mercuric reductase, Pseudomonas | 31 | $<10^{-32}$ |
| Glutamate dehydrogenase (NAD) | Glutamate dehydrogenase, E. coli | 29 | $<10^{-24}$ |
| Ornithine transcarbamylase | Aspartate transcarbamylase, E. coli | 26 | $<10^{-11}$ |

Adapted from Doolittle et al., 1986

Table 7: Mosaic proteins

| A. EGF-type | B. C9-type |
| :--- | :--- |
| Epidermal growth factor precursor | Complement C9 |
| Tumor growth factors | LDL receptor |
| LDL receptor | Notch (Drosophila) |
| Factor IX | lin-12 (C. elegans) |
| Protein C |  |
| Tissue plasminogen activator | C. Fibronectin finger |
| Urokinase | Fibronectin |
| Complement C9 | Tissue plasminogen activator |
| Notch protein (Drosophila) | D. Protease "Kringle" |
| lin-12 (C. elegans) | Plasminogen |
|  | Tissue plasminogen activator |
|  | Urokinase |
|  | Prothrombin |
|  |  |

From Doolittle et al., 1986.
proteins from primordial exons. Alternatively, introns are also capable of invading genes; thus, the association of exons with structures may reflect modern invasions that are less disruptive when they occur between structural elements.

A recent test of the "introns" early hypothesis suggests there is little evidence to support the asso-
ciation of introns with structural boundaries (Stoltzfus et al., 1994.

### 1.5 DNA vs Protein comparison

While all of the comparison methods described below work on either protein or DNA sequences, one's ability to identify distantly related sequences is reduced dramatically when DNA sequences are used. Fig. 8 compares the statistical significance of the best similarity scores obtained in a search of the GenBank DNA sequence database using a mouse glutathione transferase cDNA clone with the significance of the same alignment in a search of the GenPept protein sequence database (GenPept is derived from GenBank by translating DNA sequences into the encoded protein sequences). Many DNA sequences encoding clearly related proteins, e.g. RABGSTB have similarity scores that are expected to occur several times by chance in a DNA database search. DNA sequences are far less informative, both because they lack the inherent biochemical information that is retained in the PAM250 substitution matrix, and because many changes in DNA sequences (third-base changes) do not change the encoded protein.

Differences in the performance of sequence comparison algorithms are insignificant compared to the loss of information that occurs when one compares DNA sequences. If the biological sequence of interest encodes a protein, protein sequence comparison is always the method of choice.

Table 8: DNA vs Protein Sequence Comparison

|  |  | score | $\mathrm{E}(\mathrm{DNA})$ | $\mathrm{E}(\mathrm{prot})$ | $\mathrm{E}(\mathrm{tx})$ |
| :--- | :--- | :--- | :--- | :--- | :--- |
| MUSGST | Mouse glutathione S-transferase class mu | 5090 | $10^{-233}$ | $10^{-90}$ | $10^{-120}$ |
| MUSGSTA | Mouse, glutathione transferase GT9.3 mu | 3693 | $10^{-167}$ | $10^{-73}$ | $10^{-120}$ |
| HUMGSTAB | Homo sapiens glutathione transferase | 1930 | $10^{-84}$ | $10^{-60}$ | $10^{-80}$ |
| MAMGLUTRA | M.auratus mu class GST | 399 | $10^{-11}$ | $10^{-73}$ | $10^{-11}$ |
| RATGSTYD | Rat glutathione S-transferase Yb subunit | 399 | $10^{-11}$ | $10^{-74}$ | $10^{-10}$ |
| HSGSTM4 | H.sapiens GSTM4 gene for GST | 390 | $10^{-11}$ | $10^{-69}$ | $10^{-10}$ |
| RATGSTY | Rattus norvegicus GST | 372 | $10^{-10}$ | $10^{-71}$ | $10^{-10}$ |
| HSGSTM1B | H.sapiens GSTM1b gene for GST | 358 | $10^{-9}$ | $10^{-63}$ | $10^{-10}$ |
| HSGSTMU3 | Human GSTmu3 gene for a GST | 322 | $10^{-7}$ | $10^{-25}$ | $10^{-6}$ |
| BTGST | Bovine GST mRNA for GST | 249 | 0.00013 | $10^{-16}$ | $10^{-22}$ |
| HSGSTPI1 | Human mRNA for anionic GST | 237 | 0.00049 | $10^{-17}$ | $10^{-21}$ |
| MUSGTF | Mus musculus GST mu | 196 | 0.041 | $10^{-4}$ | $10^{-6}$ |
| CRUGSTP | Chinese hamster GST | 196 | 0.043 | $10^{-16}$ | $10^{-21}$ |
| CRUGSTPIE | Cricetulus griseus GST pi | 196 | 0.04 | $10^{-16}$ | $10^{-21}$ |
| HAMGSTPIE | Mesocricetus auratus GST pi | 191 | 0.13 | $10^{-16}$ | $10^{-21}$ |
| BTRNAXOR | B.taurus xanthine oxidoreductase | 184 | 0.11 | $>10$ | $>5$ |
| HUMKAL2 | Human glandular kallikrein gene | 170 | 0.59 | $>10$ | $>5$ |
| RNGSTYC2F | R.norvegicus GST Yc1 | 170 | 0.67 | $10^{-6}$ | $>5$ |
| MMGLUT | M.musculus mRNA for GST | 168 | 1.0 | $10^{-7}$ | $10^{-8}$ |
| MUSTHYGP | Mouse Thy-l.2 glycoprotein | 163 | 1.3 | $>10$ | $>5.0$ |
| HUMTROPIO1 | Human troponin I, slow-twitch isoform | 161 | 1.7 | $>10$ | $>5$ |
|  |  |  |  |  |  |

Expectation values for searches against DNA (score, $\mathrm{E}(\mathrm{DNA})$ ), protein ( $\mathrm{E}($ prot $)$ ), and translated DNA ( $\mathrm{E}(\mathrm{tx}$ ) databases. A mouse glutathione transferase cDNA sequence (MUSGST) was used to search either the primate (GBPRI), rodent (GBROD), and mammalian (GBMAM) divisions of the GenBank DNA sequence database for the DNA sequence comparisons. Protein expectations ( E (prot)) were calculated from a search the translated cDNA sequence against the GenPept sequence database, which includes all of translated GenBank. Unrelated sequences are italicized; $\mathrm{E}(\mathrm{prot})$ for unrelated sequences are $\gg 100$.

## 2 Alignment methods

A variety of comparison algorithms and scoring parameters can be used to evaluate protein or DNA sequence similarity. In general, the choice the of "best" algorithm depends on the problem to be solved. Thus, algorithms that calculate a local comparison score-i.e., they find the strongest similarity between the two sequences, ignoring differences outside of the most similar region-are usually most appropriate for searching protein and DNA databases, ${ }^{2}$ while global comparison algorithms are more appropriate when homology has been established, as when building evolutionary trees. Pattern-based, rather than similarity-based, comparison methods may be preferred when searching for functionally conserved non-homologous domains.

In searching protein sequence databases to identify distantly related homologous proteins, it is important to remember that avoiding high similarity scores with unrelated sequences can be more important as calculating high scores for related sequences. There are more than 50,000 protein sequences in comprehensive protein databases, while the typical family of proteins has fewer than 100 members. Thus, comparison algorithms, scoring matrices and gap penalties that produce the best alignments may not be the most effective for searching protein sequence databases (Pearson, 1995; ?).

### 2.1 Algorithms

Two general classes of comparison algorithms are used to calculate similarity scores to infer sequence homology: rigorous algorithms that are guaranteed to calculate an optimal similarity score, e.g. the NeedlemanWunsch (Needleman \& Wunsch, 1970) and SmithWaterman (Smith \& Waterman, 1981) algorithms, and rapid heuristic algorithms that do not guarantee to calculate an optimal score for every sequence in a library, e.g. FASTA (Pearson \& Lipman, 1988) and BLAST(Altschul et al., 1990). Table 2.1 summarizes widely used algorithms for biological sequence comparison.

Two optimal algorithms for calculating similarity scores have been described, the NeedlemanWunsch algorithm (Needleman \& Wunsch, 1970), which calculates a "global" similarity score between two sequences, and the Smith-Waterman algorithm (Smith \& Waterman, 1981), which calculates a "local" similarity score. Global scores require the alignment to begin at the beginning of each sequence and extend to the end of each sequence. Global alignments cannot be used to detect the relationship between DNA binding domains in homeobox proteins or the calcium binding domains shared between calmodulin and calpain. Likewise, global alignment algorithms often do not detect the relationships between mosaic proteins. Global similarity scores can be calculated with or without penalties for gaps at the ends of the sequences.

Local alignment algorithms identify the most similar region shared between two sequences. Thus, homologous calcium binding domains embedded in non-homologous proteins can be detected with local alignment methods. In addition, a local alignment algorithm can be used to find the exons in a genomic DNA sequence by aligning it with its encoded mRNA. Local alignment algorithms are required to identify homologies in mosaic proteins, and they can be used to detect internal domain duplications as well. Table 10 compares the scores of global, global without end-gap-penalties, and

[^3]Table 9: Algorithms for comparing protein and DNA sequences

| algorithm | value <br> calculated | scoring <br> matrix | gap <br> penalty | time <br> required |  |
| :---: | :---: | :---: | :---: | :---: | :--- |
| Needleman- <br> Wunsch | global similarity | arbitrary | penalty/gap <br> $q$ | $O\left(n^{2}\right)$ | Needleman and <br> Wunsch, 1970 |
| Sellers | (global) distance | unity | penalty/residue <br> $r k$ | $O\left(n^{2}\right)$ | Sellers, 1974 |
| Smith- | local similarity | $\hat{S}_{i j}<0.0$ | affine <br> $q+r k$ | $O\left(n^{2}\right)$ | Smith and Waterman, 1981 <br> Waterman |
| FASTA | approx. local <br> similarity | $\hat{S}_{i j}<0.0$ |  | limited gap size <br> $q+r k$ | $O\left(n^{2}\right) / K$ |

local similarity scores for a variety of related and unrelated proteins.
Rigorous sequence comparison algorithms, like the Smith-Waterman algorithm, require time proportional to $O(m N)$, where $m$ is the length of the query sequence and $N$ is the number of amino acids in the protein sequence library. Modern high-performance unix workstations can compare a 300 residue protein sequence (human opsin) to the 40,000 entry, 15,000,000 amino acid Swiss-Prot 31 database in less than 10 minutes.

Although very rapid ${ }^{3}$ algorithms are available for calculating optimal global similarity scores between two sequences, particularly with unit cost scores, such algorithms are rarely appropriate for biological sequence comparison. Unit cost algorithms must discard the substantial biochemical information encoded in the PAM250 matrix. Most rapid optimal algorithms calculate only global similarities; such comparisons are not useful for DNA sequence comparison because the "ends" required for a global sequence comparison are usually arbitrary.

[^4]Table 10: Global and local sequence similarity scores

| PIR Entry |  |  |  | Similarity Score |  |  | Distance |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | Global |  | Local |  |
|  |  |  |  | $\begin{gathered} \text { End } \\ \text { Penalty } \end{gathered}$ | No End Penalty |  |  |
| HBHU | vs | HBHU | Hemoglobin beta-chain-human | 725 | 725 | 725 | 0 |
|  |  | HAHU | Hemoglobin alpha-chain-human | 314 | 320 | 322 | 152 |
|  |  | MYHU | Myoglobin-Human | 121 | 164 | 166 | 212 |
|  |  | GPYL | Leghemoglobin-Yellow lupin | 8 | 28 | 43 | 239 |
|  |  | LZCH | Lysozyme precursor-Chicken | -107 | 16 | 32 | 220 |
|  |  | NRBO | Pancreatic ribonuclease-Bovine | -124 | 16 | 31 | 280 |
|  |  | CCHU | Cytochrome c-Human | -160 | 10 | 26 | 321 |
| MCHU | vs | MCHU | Calmodulin-Human | 671 | 671 | 671 | 0 |
|  |  | TPHUCS | Troponin C, skeletal muscle | 395 | 430 | 438 | 161 |
|  |  | PVPK2 | Parvalbumin beta-Pike | -57 | 103 | 115 | 313 |
|  |  | CIHUH | Calpain heavy chain-Human | -2085 | 89 | 100 | 2463 |
|  |  | AQJFNV | Aequorin precursor-Jelly fish | -65 | 48 | 76 | 391 |
|  |  | KLSWM | Calcium binding protein-Scallop | -89 | 45 | 52 | 323 |
| QRHULD | vs | EGMSMG | Epidermal growth factor precursor | -591 | 475 | 655 | 2549 |

Figure 10: Global and local alignment paths
A. Global

|  | A | B | D | D | E | F | G | H | I |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| A |  | $\begin{aligned} & \text { \} } \\ {-1} \end{aligned}$ |  |  |  |  |  |  | \} |
| B | $\backslash$ | $\backslash_{2}$ |  | $\begin{aligned} & \backslash \\ & -2 \end{aligned}$ | ${ }_{-2}$ | ${ }_{-2}$ |  |  | $-2$ |
| D | \} |  | $3$ | $1$ | $-1$ | $\backslash_{-3}$ | $\backslash_{-3}$ | $\_{-3}$ | $\backslash_{-3}$ |
| E | \} | $\begin{array}{r} 1 \text { ! } \\ -2 \end{array}$ |  | $2$ | $I_{2}$ |  | _-2 | $\begin{aligned} & 1 \\ & --4 \end{aligned}$ | $-4$ |
| G | \} | ${ }_{-2}$ |  | $\begin{array}{r} \backslash! \\ 0 \end{array}$ | $\backslash_{1}$ | $\backslash_{1}$ | $\backslash_{1}$ |  | _-3 |
| K | $\backslash$ | $\backslash_{-2}$ | $\begin{array}{r} 1 \text { ! } \\ -3 \end{array}$ | $\begin{array}{r} \backslash! \\ -2 \end{array}$ | $\begin{array}{r} \backslash! \\ -1 \end{array}$ | $\backslash_{0}$ | $\_{0}$ | $\bar{彳}_{0}$ | $-2$ |
| H | $\backslash$ | $\backslash_{-2}$ | $\backslash_{-3}$ | $\begin{array}{r} \backslash! \\ -4 \end{array}$ | $\begin{array}{r} 1! \\ -3 \end{array}$ | $\begin{array}{r} \backslash! \\ -2 \end{array}$ | $\backslash_{-1}$ | $1$ | $\overline{\}_{-1}$ |
| I | $\backslash$ | $\backslash_{-2}$ | $\_{-3}$ | $\backslash_{-4}$ | $\begin{array}{r} \backslash! \\ -5 \end{array}$ | $\begin{array}{r} 1! \\ -4 \end{array}$ | $\begin{array}{r} 1! \\ -3 \end{array}$ |  | $2$ |

Optimal global alignments (score 2):

$$
\begin{aligned}
& \text { A B D D E F G H I (top) } \\
& \text { A B D - E G K H I (side) } \\
& \text { or } A B-D E G K H I
\end{aligned}
$$

B. Local

|  | A | B |  | D |  | D |  | E |  | F |  |  | H |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| A | $\backslash$ |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
|  |  | 1 | 0 |  | 0 |  | 0 |  | 0 |  | 0 |  |  | 0 |  | 0 |
| B |  | ) |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
|  |  | 0 | 2 | - | 0 |  | 0 |  | 0 |  | 0 |  |  | 0 |  | 0 |
| D |  |  | ! | \} |  | \} |  |  |  |  |  |  |  |  |  |  |
|  |  | 0 | 0 |  | 3 | - | 1 |  | 0 |  | 0 |  |  | 0 |  | 0 |
| E |  |  |  |  | . | \} |  | $\backslash$ |  |  |  |  |  |  |  |  |
|  |  | 0 | 0 |  | 1 |  | 2 |  | 2 | - | 0 |  |  | 0 |  | 0 |
| G |  |  |  |  |  | $\backslash$ | ! | $\backslash$ |  | $\backslash$ |  |  |  |  |  |  |
|  |  | 0 | 0 |  | 0 |  | 0 |  | 1 |  | 1 |  |  | 0 |  | 0 |
| K |  |  |  |  |  |  |  |  |  | $\backslash$ |  |  | 1 |  |  |  |
|  |  | 0 | 0 |  | 0 |  | 0 |  | 0 |  | 0 | 0 |  | 0 |  | 0 |
| H |  |  |  |  |  |  |  |  |  |  |  |  | 1 |  |  |  |
|  |  | 0 | 0 |  | 0 |  | 0 |  | 0 |  | 0 |  |  | 1 |  | 0 |
| I |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
|  | 0 | 0 | 0 |  | 0 |  | 0 |  | 0 |  | 0 |  |  | 0 |  | 2 |

Optimal local alignment (score 3):

$$
\begin{array}{llll}
\text { A } & \text { B } & \text { D } & \text { (top) } \\
\text { A } & \text { B } & \text { D } & \text { (side) }
\end{array}
$$

### 2.2 Dynamic Programming Algorithms

The algorithms used to calculate the maximum similarity scores between two sequences are most easily visualized with an alignment matrix or path graph. Figs. 10-11 demonstrate the correspondence between an alignment path graph and an actual alignment. The goal along the path is to maximize the similarity score for the alignment that ends at each potential vertex. For the figures, similarity scores are increased by +1 for diagonal edges if the two residues along the path are identical; if they are different, the diagonal edge cost is -1 . The cost along either a horizontal or vertical edge, which corresponds to an insertion in the top sequence (horizonal edge) or an insertion in the left-side sequence (vertical edge) is -2 . To produce a global alignment from a path graph, simply begin at the bottomright corner of the graph and follow the "active" paths, noted by $\backslash, \quad$ or ! to the upper-left corner, aligning the two residues along the diagonal path, or aligning a residue with a gap if a horizontal or vertical path is taken.

For the global alignment in Fig. 10A, there are two alignments that produce the optimal score. Optimal comparison algorithms guarantee to produce the best score, given the match, mismatch, and gap costs, but frequently there are several optimal alignments for a single score. For the local alignment in Fig. 10B, there are several sub-optimal alignments with scores of 2 . Note that the local alignment in Fig. 10B would extend from one end of each sequence to the other if the gap cost were reduced to -1 .

Figure 11: An alignment path matrix


Fig. 11 provides an exercise for the reader.
While there are an exponential number of potential alignments with gaps between two protein or DNA sequences, dynamic programming algorithms are available that can calculate the optimal score in $O(M N)$ steps. This efficiency is achieved by determining the optimal score for each prefix of each string, and then extending each prefix by considering the three paths that can be used to extend an alignment: (1) by extending the alignment by one residue in each sequence; (2) by extending the alignment by one residue in the first sequence and aligning it with a gap in the second; or (3) extending the alignment by one residue in the second sequence and aligning it with a gap in the first. This decision must be made for each of the $M N$ prefixes of sequences of length $M$ and $N$.

The first algorithm for comparing protein sequences (Needleman \& Wunsch, 1970) calculates a "global" similarity score. A simplified global algorithm is shown in Fig. 12. Since a global algorithm requires that the alignment extend from the beginning to the end of the alignment, it is sufficient to report the score in the lower right $(S(M, N))$ of the scoring matrix.

Local alignment algorithms must consider alignments that begin and end at each of the $M N$ positions in the alignment matrix. Despite this added complexity, they only add two additional steps to

Figure 12: Algorithms for Global and Local similarity scores

```
\(S(0,0) \leftarrow 0\)
for \(j \leftarrow 1\) to \(N\) do
    \(S(0, j) \leftarrow S(0, j-1)+\sigma\binom{-}{b_{j}}\)
for \(i \leftarrow 1\) to \(M\) do
[ \(\quad S(i, 0) \leftarrow S(i-1,0)+\sigma\binom{a_{i}}{-}\)
    for \(j \leftarrow 1\) to \(N\) do
        \(S(i, j) \leftarrow \max \left[S(i-1, j-1)+\sigma\binom{a_{i}}{b_{j}}, S(i-1, j)+\sigma\binom{a_{i}}{-}, S(i, j-1)+\sigma\binom{-}{b_{j}}\right]\)
]
write "Global similarity score is" \(S(M, N)\)
best \(\leftarrow 0\)
for \(j \leftarrow 1\) to \(N\) do
    \(S^{\prime}(0, j) \leftarrow S^{\prime}(0, j-1)+\sigma\binom{-}{b_{j}}\)
for \(i \leftarrow 1\) to \(M\) do
[ \(\quad S^{\prime}(i, 0) \leftarrow S^{\prime}(i-1,0)+\sigma\binom{a_{i}}{-}\)
    for \(j \leftarrow 1\) to \(N\) do
    \(\left[\quad S^{\prime}(i, j) \leftarrow \max \left[0, S^{\prime}(i-1, j-1)+\sigma\binom{a_{i}}{b_{j}}, S^{\prime}(i-1, j)+\sigma\binom{a_{i}}{-}, S^{\prime}(i, j-1)+\sigma\binom{-}{b_{j}}\right]\right.\)
        best \(\leftarrow \max \left(S^{\prime}(i, j), b e s t\right)\)
]
write "Local similarity score is" best
```

the global alignment algorithm. Since every possible starting position must be considered, similarity scores cannot fall below zero and a 0 term is added to the max comparison in Fig. 12. Since they can end at any position in the matrix, the best score must be saved at each step. In practice, global and local comparison algorithms require the same amount of computation.

### 2.3 Scoring methods

The scoring matrices used for protein sequence comparison are much more sophisticated than +1 for a match and -1 for a mismatch. The most effective matrices are based on the actual frequency of substitutions that occur between related proteins. Two different approaches have been used to produce these matrices. The original PAM250 matrix (Fig. 5) was produced by examining several hundred alignments between very closely related proteins, and then calculating the frequency with which each
amino-acid residue changed into each of the others at a very short evolutionary distance-one where only $1 \%$ of the residues had kchanged (Dayhoff et al., 1978). This replacement frequency, when corrected for the amino-acid abundance, can be used to calculate the PAM1 scoring matrix (PAM is "Point Accepted Mutation"). If the matrix is multiplied against itself 250 times, a PAM250 matrix, which reflects the frequency of change for proteins that have diverged $250 \%$. If a two protein sequences have diverged by $250 \%$, it is expected that they will share about $20 \%$ sequence identity (Fig. 9). Since $20 \%$ identity is at the edge of where significant similarity can be detected, the PAM250 matrix has been widely used. The PAM250 matrix is based on small number of amino acid substitutions; modern extrapolated matrices based both on sequence alignments (Jones et al., 1992) and structural alignments (Johnson \& Overington, 1993) are available.

Substitution matrices have also been calculated directly by examining "blocks" of aligned sequences that differ by no more than $X \%$ (Henikoff \& Henikoff, 1992). Thus, the BLOSUM62 matrix, which is used by the BLASTP rapid comparison program, is derived from substitution data for blocks of aligned sequences that are no more than $62 \%$ identical. BLOSUM62 performs substantially better than extrapolated matrices with BLASTP and FASTA (Henikoff \& Henikoff, 1993), but both BLOSUM and extrapolated matrices can perform well when used with optimal gap penalties (Pearson, 1995).

Altschul (1991) has provided a information-theory based perspective for evaluating scoring matrices in general for alignments without gaps. Using a statistical theory for such alignments (Karlin \& Altschul, 1990), it is possible to convert any similarity score to a value in "bits" that can be used to compare scores produced by different alignments. Unfortunately, the analytical formulas that are used for this conversion cannot easily be applied to alignments that contain gaps. Collins et al., 1988 and Altschul, 1993 have also pointed out that different scoring matrices are optimal at different evolutionary distances. Thus, short proteins sequences that are $50 \%$ identical can be more easily identified with a "shallower" PAM matrix, e.g. PAM60.

### 2.4 Heuristic Algorithms

Two rapid heuristic algorithms are frequently used for searching protein and DNA sequence databases, FASTA (Pearson \& Lipman, 1988) and BLASTP (Altschul et al., 1990). These methods are 5-50 times faster than the rigorous Smith-Waterman algorithm, and can produce results of similar quality in many cases.

Fig. 13 summarizes the difference between the FASTA, BLASTP, and Smith-Waterman algorithms. BLASTP and FASTA are faster than Smith-Waterman because they examine only a portion of the potential alignments between two sequences. FASTA focuses on regions where there are either pairs ( $k t u p=2$ ) or single aligned $k t u p=1$ identities; BLASTP examines regions that include triples of conserved amino acids.

### 2.4.1 BLAST

Advances in the statistical theory of sequence alignments without gaps (Karlin \& Altschul, 1990) provided the theoretical basis for the BLASTP program (Altschul et al., 1990). BLASTP is now the most

Figure 13: Heuristic strategies for sequence comparison


Table 11: Sequence similarity with BLASTP

Step 1 For each three amino acids in the query sequence, identify all of the substitutions of each word that have a similarity score greater than a threshold score $T=11$. In practice, word-matches with scores $\geq T$ are seen on average 150 times per library sequence.
Step 2 Build a discrete finite automaton (DFA) to recognize the list of identical and substituted three letter words.
Step 3 Use the DFA to identify all of the matching words in sequences in the database. If a match is found, attempt to extend the match both forwards and backwards using the BLOSUM62 matrix to produce a score that is higher than a threshold score. Save all of the high scoring regions shared by the query sequence and each library sequence. The best of these scores is reported as the best single MSP (maximal segment pair) score. These high scoring regions do not contain gaps.

Step 4 Attempt to combine multiple MSP regions. For each "consistent" combination, calculate the probability of obtaining that may consistent matches using either "poisson" or "sum" statistics.(Karlin \& Altschul, 1993) Report the lowest probability score based on statistics used.

Step 5 Report all of the significant alignments. Frequently, a query and library sequence will contain several MSPs because of the requirement that they do not contain gaps.
widely used program for rapid sequence comparison, in large part because of its accurate estimates for the statistical significance of similarity scores (see 3. BLASTP, like FASTA, uses a word-based scanning procedure to identify regions of local similarity (Table 11) with out gaps. BLASTP is effective because it combines high sensitivity with excellent selectivity. BLASTP combines good sensitivity with exceptional selectivity. Except when the query sequence contains a low complexity region, BLASTP rarely calculates scores for unrelated sequences.

### 2.4.2 FASTA

The current version of FASTA provides several significant improvements over earlier versions. FASTA now calculates optimized scores (step 4 in Table 12)) for most of the sequences in the database and provides accurate estimates for statistical significance (3). Calculation of optimized scores improves substantially the performance of FASTA. Without the calculation, FASTA performs significantly worse than BLASTP; however, with the calculation of optimized scores (and normalization of the scores based on library sequence length), FASTA performs significantly better than BLASTP and almost as well as the Smith-Waterman algorithm (Pearson, 1995). In addition, FASTA now uses the SmithWaterman algorithm to produce final alignments; previous versions limited the size of gaps, which sometimes led to incomplete alignments.

Every database search for members of a diverse protein family involves a tradeoff between sensitivitythe ability to identify distantly related members of the family-and selectivity-the ability to avoid high similarity scores for unrelated sequences. Table 3.3 compares how effectively the three algorithms maintain this balance for a large protein family—the G-protein-coupled receptors. Thus, BLASTP calculates a very highly significant score for the closely related opsin and dopamine D 2 receptors, and a significant score for the more distantly related thromboxane $\mathrm{A}_{2}$ receptor, but it does not detect the

Table 12: Sequence similarity with FASTAv20

Step 1 Identify regions shared by the two sequences with the highest density of identities ( $k t u p=1$ ) or pairs of identities ( $k t u p=2$ ).

Step 2 Rescan the ten regions with the highest density of identities using the BLOSUM50 matrix. Trim the ends of the region to include only those residues contributing to the highest score. Each region is a partial alignment without gaps.
Step 3 If there are several initial regions with scores greater than the CUTOFF value, check to see whether the trimmed initial regions can be joined to form an approximate alignment with gaps. Calculate a similarity score that is the sum of the joined initial regions minus a penalty (usually 20) for each gap (initn). The score of the single best initial region found in Step 2 is also reported (initl).

Step 4 For sequences with scores greater than a threshold, construct an optimal local alignment of the query sequence and the library sequence, considering only those residues that lie in a band centered on the best initial region found in Step 2. For protein searches with $k t u p=2$ a 16 residue band is used by default. A 32 residue band is used with $k t u p=1$. This is the optimized (opt) score.

Step 5 After all (or the first $10-20,000$ ) scores have been calculated, normalize the raw similarity scores by regressing the similarity score against $\ln$ (library-sequence length) and calculating the average variance in similarity scores. Z-values (normalized scores with mean 0 and variance 1 ) are calculated, and the calculation is repeated with library sequences with $z$-values greater than 5.0 and less than -5.0 removed. These z -values are used to rank the library sequences.
Step 6 The Smith-Waterman algorithm (without limitation on gap size) is used to display alignments.
similarity between opsin and the very distantly related Dictyostelium cAMP (CAR1) receptor. In addition, BLASTP would never suggest a relationship between opsin and cytochrome oxidase. FASTA ( $k t u p=2$ does a better job at recognizing the relationship between opsin and thromboxane A2, fails to detect the cAMP-1 receptor, and is more ambiguous about a possible relationship with cytochrome oxidase. FASTA with $k t u p=1$ and Smith-Waterman calculate statistically significant relationships between opsin and cAMP-1, but also good (but not significant) scores for opsin and cytochrome oxidase.

## 3 The statistics of sequence similarity scores

The development of accurate statistical estimates for local sequence similarity scores (Karlin \& Altschul, 1990; Mott, 1992) has allowed dramatic improvement in our ability to reliably recognize distantly related proteins. The statistical estimates calculated by BLASTP are used widely in large scale sequence comparison, e.g. to characterize all of the genes on a yeast chromosome or all of the genes in a bacterial genome. The incorporation of statistical estimates into FASTA and SSEARCH (a Smith-Waterman implementation) have significantly improved the performance of these programs as well.

### 3.1 Sequence alignments without gaps

The statistics of local similarity scores for alignments without gaps but with an arbitrary substitution matrix have been described by Karlin \& Altschul, 1990. Local similarity scores are described by the extreme value distribution. Using the parameters $\lambda$ and $K$, which can be derived from the scoring matrix and the amino acid composition of the query sequence, the probability that a normalized similarity score:

$$
\begin{equation*}
S^{\prime}=\lambda S-\ln K m n \tag{1}
\end{equation*}
$$

(Karlin \& Altschul, 1990; Altschul et al., 1994) where $m$ is the length of the query sequence and $n$ is the length of the library sequence can be calculated as:

$$
\begin{equation*}
P\left(S^{\prime} \geq x\right)=1-\exp \left(-e^{-x}\right) \tag{2}
\end{equation*}
$$

Since a typical database search typically involves thousands of pairwise comparisons, the expectation of finding a score $S^{\prime} \geq X$ for a search of $D$ sequences is: $E\left(S^{\prime} \geq X\right)=P D$. (Thus, searches of highly redundant databases may be less informative, because $D$ is larger but the number of different sequences is not.)

### 3.2 Similarity scores increase with sequence length

The normalization in equation 1 shows that scores for alignments without gaps between random sequences increase as $\ln K m n$, or since $K$ and $m$ are fixed for a given search, $\ln n$, the length of the library sequence. This is seen empirically with scores for alignments that contain gaps (Collins et al., 1988; Mott, 1992) and is shown in Fig. 14. For local similarities, the variance of the score should be independent of library sequence length. Thus, normalization of similarity scores by fitting a line to the relationship of similarity score to $\ln n$ will reduce the scores of long, unrelated sequences, and make it possible to detect more distant relationships (Pearson, 1995).

### 3.3 Empirical statistics for alignments with gaps

Accurate statistical estimates for alignments with gaps can can be calculated by normalizing similarity scores to remove the $\ln n$ dependence for similarity scores. This can be seen in Fig. 6, where the ' $\star$ 's show the fit of an extreme value distribution to the observed data (' $==$ '). FASTA and SSEARCH estimate statistical significance by fitting a line to $S$ vs $\ln n$ and calculating the average variance for the scores. The regression line and variance are used to calculate

$$
\begin{equation*}
Z-\text { score }=(S-(a+b \ln n)) / \sqrt{v a r} \tag{3}
\end{equation*}
$$

The distribution of $Z-$ score's should follow the extreme value distribution, so that:

$$
\begin{equation*}
P(Z>x)=1-\exp \left(-e^{-1.282 Z-0.5772}\right) \tag{4}
\end{equation*}
$$

and, as before, $E(Z>x)=P D$.

Figure 14: Similarity scores and library sequence length


The distribution of Smith-Waterman similarity scores is plotted as a function of $\log (n), n$ is the length of the library sequence. Filled symbols indicate individual related sequences (only the most distant related sequences are shown); open symbols show the average and std. error of similarity scores for unrelated sequences.

### 3.4 Statistical significance by random shuffling

Statistical estimates derived from database searches measure the difference between an observed similarity score and that expected for a sequence with the amino acid composition of the database. Such tests may overestimate significance in cases where the query sequence's amino acid composition differs from that of the database. Thus, membrane proteins with their hydrophobic transmembrane domains may have statistically significant scores with non-homologous membrane proteins. A more challenging test compares the similarity score between a query and library sequence with the distribution of scores obtained by comparing the query sequence to random sequences with the same length and amino acid composition as the library sequence. Such sequences are easily generated by randomly shuffling the library sequence, either globally, by exchanging randomly each amino acid with any other position in the sequence, or locally, by performing the exchanges within a window of 10-20 residues.

Table 13: Search Algorithms and Statistical Significance

| algorithm | closely <br> related | related | distantly <br> related | unrelated |
| :--- | :---: | :---: | :---: | :---: |
|  | dopamine D2 $^{a}$ | thromboxane A2 $^{b}$ | cAMP-1 $^{c}$ | cytochrome oxidase $^{d}$ |
|  |  |  |  |  |
| Smith-Waterman $^{\text {chSS }^{e}}$ | $3 \times 10^{-9}$ | $2 \times 10^{-4}$ | 0.01 | 0.57 |
| PRSS(window=20) | $8 \times 10^{-10}$ | $10^{-4}$ | 0.007 | 0.45 |
|  | $8 \times 10^{-8}$ | 0.001 | 0.23 | 3.0 |
| fasta, ktup=1, opt | $3 \times 10^{-9}$ | $7 \times 10^{-5}$ | 0.02 | 0.39 |
| fasta, ktup=2, opt | $2 \times 10^{-6}$ | $10^{-4}$ | 2.2 | 0.36 |
| BLASTP | $2 \times 10^{-22}$ | 0.07 | $>1.0$ | $>1.0$ |

${ }^{a}$ D2DR_HUMAN, ${ }^{b}$ TA2R_MOUSE, ${ }^{c}$ CAR1_DICDI, ${ }^{d}$ APPC_ECOLI

Expected number of times that a similarity score as high or higher than that obtained by the indicated library sequence would be obtained by chance in a search of Swiss-Prot ( $\approx 58,000$ entries) with the OPSD_HUMAN (human opsin) query sequence. ${ }^{e}$ Expected times this score would be obtained after 1,000 shuffles of the indicated library sequence with either global (prss) or local (window=20) amino acid exchanges.

Because this Monte Carlo test measures the significance of the order of the two amino acid sequences, rather than the difference between the highest scoring sequences and the rest of the database, it tends to be more demanding.

As before, similarity scores for random sequences should follow the extreme value distribution, and a fit of the distribution of scores can be used to estimate the significance of an unshuffled score. However, to extrapolate an expectation value from shuffled sequences to that for a library search, the " E() -value" must be multiplied by the ratio of the number of sequences in the library to the number of shuffled sequences. Thus, in the example below, an E() -value from 500 shuffles must be multiplied by 80 to be comparable to an E() -value from the 40,000 entry Swiss-Prot. As expected, the E() -value from the actual search- $2 \times 10^{-4}$ —is slightly more significant than that from the shuffled distribution- $3 \times 10^{-3}$.

```
Comparison of OOHU (human opsin) with TA2R_MOUSE (thromboxane A2 receptor)
BLOSUM50 matrix, gap penalties: -12,-2
unshuffled s-w score: 160; shuffled score range: 38 - 92
Lambda: 0.15076 K: 0.017357; P(160)= 7.4282e-08
For 500 sequences, a score >=160 is expected 3.71e-05 times
```

Although accurate statistical estimates can be very valuable in interpreting the results of similarity searches, they must be evaluated with caution. Distantly related homologous sequences often do not

Figure 15: Patterns for serine proteases

```
ID TRYPSIN_HIS; PATTERN.
AC PSO0134;
DE Serine proteases, trypsin family, histidine active site.
PA [LIVM]-[ST]-A-[STAG]-H-C.
NR /TOTAL=158(158); /POSITIVE=154(154); /UNKNOWN=2 (2); /FALSE_POS=2 (2);
NR /FALSE_NEG=11(11);
CC /TAXO-RANGE=??EP?; /MAX-REPEAT=1;
CC /SITE=5,active_site;
ID TRYPSIN_SER; PATTERN.
AC PSO0135;
DE Serine proteases, trypsin family, serine active site.
PA G-D-S-G-G.
NR /TOTAL=160(160); /POSITIVE=151(151); /UNKNOWN=1(1); /FALSE_POS=8(8);
NR /FALSE_NEG=16(16);
CC /TAXO-RANGE=??EP?; /MAX-REPEAT=1;
CC /SITE=3,active_site;
```

Patterns from PROSITE that identify 152/163 (TRYPSIN_HIS or 143/159 TRYPS IN_SER members of the serine protease protein family.
share statistically significant similarity. Thus, overreliance on statistical estimates, particularly after a single search, can miss genuine homologies. Conversely, sequences with low-complexity regions often share significant similarity but are not homologous. Finally, some structures, such as the coiledcoil structure in tropomyosin, share statistical significance because of a common repeated structure, because of convergence (analogy), rather than homology.

## 4 Identifying distantly related protein sequences

In this section, we will examine similarity searches in three diverse families of protein sequences, serine proteases, glutathione S-transferases, and the G-protein-coupled receptors. The serine proteases are considered because they provide a classic example of a family of proteins with a highly conserved active site; the glutathione transferases are a very diverse family where many members do not share significant similarity with all other members, while the G-protein-coupled receptors are a very large and diverse family of membrane proteins.

### 4.1 Serine proteases

Serine proteases cleave peptide bonds using a "catalytic triad" of histidine, serine, and aspartic acid; these residues are underlined in Fig. 17. Because these residues are so highly conserved, patterns that focus on two of the regions (Fig. 15) can be used to identify every member of the serine protease family. Fig. 16 shows the highest scoring unnormalized similarity scores. As is often the case for divergent protein families, several members of the family do not share statistically significant similarity
with bovine trypsin. These sequences are italicized in Fig. 16; their membership in the serine protease family is based on common three-dimensional structures. As expected from the discussion in section 3.2, several of the highest scoring unrelated sequences are substantially longer than genuine serine proteases. These scores have much higher (less significant) expectation values when the $\ln n$ correction is used.

The absolute conservation of residues in the "catalytic triad" might suggest that similarities between members of this family are limited to those regions. This is not the case, as can be seen in Fig. 17. Similarity in the serine proteases typically extends from one end of the protein to the other, with strong conservation throughout the sequence. Indeed, the region around one of the residues in the catalytic triad-the apartic acid-is not well conserved. While the residues in the catalytic triad is an essential feature of serine proteases, the serine protease fold (two domains containing anti-parallel $\beta$-barrels) are required to bring these residues together.

The requirement for a common folded structure in homologous proteins usually causes similarities to extend from one end of the protein to the other, or for mosaic proteins, from one end of a domain to the other. Fig. 18 displays the locally similar regions for the related and unrelated in Table 16; the highest scoring unrelated sequences tend to have relatively short ( $<100$ residue) regions of higher similarity ( $\approx 30 \%$ identical) while related sequences have longer ( $140-400$ ), though sometimes lower ( $25 \%$ ) similarity. In general, shorter, higher similarities are less significant than longer, lower similarities.

Figure 16: Serine protease search - high scoring sequences

| LOCUS | Description | len | score | $\mathrm{E}(12,000)$ |
| :---: | :---: | :---: | :---: | :---: |
| TRBOTR | trypsin precursor - bovine | 229 | 1559 | $10^{-97}$ |
| TRRT2 | trypsin II precursor - rat | 246 | 1240 | $10^{-76}$ |
| KQHU | tissue kallikrein precursor - | 262 | 669 | $10^{-37}$ |
| NGMSG | 7S NGF gamma chain I | 237 | 645 | $10^{-36}$ |
| KQRTTN | tonin - rat | 235 | 623 | $10^{-34}$ |
| KYBOA | chymotrypsin A precursor - bovine | 245 | 609 | $10^{-34}$ |
| PLHU | plasmin precursor - human | 790 | 580 | $10^{-31}$ |
| TRFF | trypsin-like proteinase | 256 | 579 | $10^{-31}$ |
| KFHU | coagulation factor IXa | 461 | 578 | $10^{-31}$ |
| ELRT2 | pancreatic elastase II | 271 | 559 | $10^{-30}$ |
| KYBOB | chymotrypsin B precursor - bovine | 245 | 556 | $10^{-30}$ |
| KFHU1 | coagulation factor XIa | 625 | 547 | $10^{-29}$ |
| WMMS28 | complement factor D homolog | 259 | 541 | $10^{-29}$ |
| EXBO | coagulation factor Xa | 492 | 518 | $10^{-27}$ |
| DBHU | complement factor D | 246 | 517 | $10^{-27}$ |
| KXBO | protein C (activated) | 456 | 515 | $10^{-27}$ |
| UKHU | u-plasminogen activator precu | 431 | 507 | $10^{-26}$ |
| TBHU | thrombin precursor - human (fr | 615 | 472 | $10^{-24}$ |
| TRSMG | trypsin - Streptomyces griseus | 221 | 409 | $10^{-20}$ |
| C1HURB | complement subcomponent C1r p | 705 | 356 | $10^{-16}$ |
| HPHU1 | haptoglobin-1 precursor - human | 347 | 335 | $10^{-15}$ |
| TRPGAZ | azurocidin - pig | 219 | 316 | $10^{-14}$ |
| HPRT | haptoglobin - rat (fragments) | 297 | 289 | $10^{-12}$ |
| C2HU | complement C 2 - human | 752 | 198 | $10^{-6}$ |
| BBHU | complement factor B - human | 739 | 169 | 0.00014 |
| KXBOZ | protein Z - bovine | 396 | 142 | 0.0041 |
| TRYXB4 | alpha-lytic proteinase | 396 | 107 | 0.83 |
| OKBY8W | probable protein kinase YCR008W | 603 | 107 | 1.3 |
| RRIHM2 | RNA-directed RNA polymerase | 4488 | 99 | 37 |
| IJFFTM | cadherin-related tumor suppressor | 5147 | 99 | 42 |
| GNNYE7 | genome polyprot. - enterovirus 70 | 2194 | 98 | 20 |
| VGIHHC | E2 glycoprotein - coronavirus | 1173 | 96 | 14 |
| QRRBVD | VLDL receptor - rabbit | 873 | 96 | 10 |
| PRSMBG* | proteinase B-S. griseus | 185 | 96 | 1.9 |
| MMMSB2 | laminin chain B2 precursor - mouse | 1607 | 95 | 23 |
| RERTK | renin precursor - rat | 402 | 94 | 6.0 |
| MMMSA | laminin chain A - mouse | 3084 | 93 | 61 |
| LNRZ | lectin precursor - rice | 227 | 90 | 6.0 |
| PRSMAG* | proteinase A - S . griseus | 182 | 89 | 5.5 |

Figure 17: Alignment of serine proteases


Figure 18: Serine protease alignments

| TRBOTR | 1559 | 100.0 |  |
| :---: | :---: | :---: | :---: |
| TRRT2 | 1240 | 74.7 |  |
| TRDFS | 1070 | 66.5 |  |
| KQHU | 669 | 41.5 | ---- |
| NGMSG | 665 | 39.7 |  |
| KQRTTN | 623 | 40.9 |  |
| KYBOA | 609 | 42.1 |  |
| PLHU | 580 | 39.7 |  |
| TRFF | 579 | 42.1 |  |
| KFHU | 578 | 40.9 |  |
| KYRTB | 564 | 39.5 |  |
| ELRT2 | 559 | 38.1 |  |
| KYBOB | 556 | 37.8 |  |
| KFHU1 | 547 | 37.6 |  |
| WMMS28 | 541 | 35.7 |  |
| EXBO | 518 | 39.4 |  |
| DBHU | 517 | 34.1 |  |
| KXBO | 515 | 37.3 |  |
| UKHU | 507 | 37.0 |  |
| TBHU | 472 | 35.8 |  |
| TRSMG | 409 | 35.3 |  |
| C1HURB | 356 | 30.4 |  |
| HPHU1 | 335 | 28.1 |  |
| TRPGAZ | 316 | 30.0 |  |
| HPRT | 289 | 26.0 |  |
| C2HU | 198 | 25.7 | -- |
| BBHU | 169 | 25.1 |  |
| KXBOZ | 142 | 25.2 |  |
| TRYXB4 | 107 | 21.5 |  |
| OKBY8W | 107 | 33.3 | - |
| RRIHM2 | 99 | 25.9 | ---- |
| IJFFTM | 99 | 27.0 |  |
| GNNYE7 | 98 | 29.9 | ----------------- |
| VGIHHC | 96 | 29.8 |  |
| QRRBVD | 96 | 25.2 |  |
| PRSMBG* | 96 | 24.9 | --- |
| MMMSB2 | 95 | 25.3 | ------ |
| RERTK | 94 | 23.8 | ----- |
| MMMSA | 93 | 25.6 | --------- |
| LNRZ | 90 | 26.1 | ---------- |
| PRSMAG* | 89 | 25.3 |  |

Table 14: Glutathione S-transferases

| The best scores are: | s-w | Z-score | E(58,753) |  |
| :--- | :--- | ---: | ---: | :--- |
| GTM1_MOUSE | Glutathione S-transferase GT8.7 | 1490 | 1827.8 | $10^{-101}$ |
| GTM1_RAT | Glutathione S-transferase YB1 | 1406 | 1818.9 | $10^{-95}$ |
| GTM1_HUMAN | Glutathione S-transferase | 1235 | 1591.1 | $10^{-82}$ |
| GTM2_CHICK | Glutathione S-transferase 2 | 954 | 1232.1 | $10^{-61}$ |
| GTP1_MOUSE | Glutathione S-transferase P | 361 | 463.7 | $10^{-19}$ |
| GTA2_MOUSE | Glutathione S-transferase Ya | 229 | 291.9 | $10^{-9}$ |
| SC2_OCTDO | S-crystallin 2 (OL2). | 224 | 290.9 | $10^{-9}$ |
| GTA1_MOUSE | Glutathione S-transferase GT41A | 218 | 277.7 | $10^{-8}$ |
| GTC_MOUSE | Glutathione S-transferase Yc | 215 | 273.9 | $10^{-8}$ |
| GTA1_HUMAN | Glutathione S-transferase A1-1 | 206 | 262.1 | $10^{-7}$ |
| GT28_SCHHA | Glutathione S-transferase 28 kd | 203 | 258.7 | $10^{-7}$ |
| GTA3_MOUSE | Glutathione S-transferase GST 5.7 | 183 | 232.3 | $10^{-6}$ |
| GT28_SCHJA | Glutathione S-transferase 28 kd | 169 | 214.8 | $10^{-5}$ |
| GTS2_DROME | Glutathione S-transferase 2 | 164 | 213.4 | $10^{-4}$ |
| SC1_OCTVU | S-crystallin 1. | 159 | 204.1 | $10^{-4}$ |
| GTA2_CHICK | Glutathione S-transferase, CL-3. | 144 | 183.0 | 0.00051 |
| SC18_OMMSL | S-crystallin SL18. | 131 | 166.9 | 0.010 |
| GTT1_MUSDO | Glutathione S-transferase 1 | 122 | 153.8 | 0.055 |
| GTH1_MAIZE | Glutathione S-transferase I | 120 | 150.9 | 0.056 |
| GTXA_TOBAC | Auxin-regulated protein | 117 | 146.7 | 0.130 |
| GT32_MAIZE | Glutathione S-transferase III | 115 | 144.1 | 0.19 |
| GTT1_DROME | Glutathione S-transferase 1-1 | 100 | 125.2 | 2.1 |
| GTH1_WHEAT | Glutathione S-transferase 1 | 98 | 121.7 | 3.3 |
| GT_PROMI | Glutathione S-transferase GST-6.0 | 97 | 121.6 | 3.4 |
| DCMA_METSP | Dichloromethane dehalogenase | 98 | 119.5 | 4.4 |
| MOD5_YEAST | tRNA isopentenyltransferase | 100 | 118.4 | 5.1 |
| GTY2_ISSOR | Glutathione S-transferase Y-2 | 94 | 118.3 | 5.2 |
| GTX2_TOBAC | Auxin-induced PGNT35/PCNT111. | 93 | 115.5 | 7.4 |
| GTT1_RAT | Glutathione S-transferase 5 | 93 | 114.8 | 8.1 |
| SPCB_HUMAN | Spectrin beta chain, erythrocyt (2137) | 108 | 113.5 | 9.6 |
| DAPF_YERPE | Diaminopimelate epimerase | 90 | 112.9 | 10.0 |
| LIGE_PSEPA | $\beta$-etherase | 91 | 110.9 | 13.0 |
| EF1G_HUMAN | Elongation factor 1 $\gamma$ | 94 | 110.5 | 14.0 |
|  |  |  |  |  |

All of the unitalicized sequences are known to be members of the glutathione transferase family.

### 4.2 Glutathione S-transferases

The glutathione transferase family of enzymes is a very diverse family of proteins found, in various forms, in animals, plants, and prokaryotes. Fortunately, many of the members of this family have a common enzyme activity so that they can be recognized by name. Table 14 shows that for this family, there are many homologues that do not show significant similarity when the database is searched with a single query sequence.

Frequently, clear identification of a distant homology will require several database searches, with either different algorithms or additional query sequences. For example, in Table 14, one might wish to test the possibility that glutathione S-transferases share homology with elongation factors, which are among the high scoring sequences. The result of a search using EF1G_HUMAN is shown in Table 15. Here, there is a clear relationship between this elongation factor and the class-theta glutathione transferases. An additional search with a class-theta sequence reveals the most distant relationships in this family more clearly.

Table 15: Distant glutathione transferase homologs

| Re-search with LIGE_PSEPA |  |  |  |  |
| :--- | :--- | ---: | ---: | :--- |
| The best scores are: | s-etherase | Z-score | E(58,762) |  |
| LIGE_PSEPA | Glutathione S-transferase 1 | 1993 | 2540.6 | $10^{-135}$ |
| GTT1_DIACA | 170 | 210.9 | $10^{-4}$ |  |
| GTX6_SOYBN | Probable glutathione S-transferase | 168 | 208.2 | $10^{-4}$ |
| GTX3_TOBAC | Probable glutathione S-transferase | 165 | 204.5 | $10^{-5}$ |
| GTXA_ARATH | Glutathione S-transferase | 161 | 199.3 | 0.00016 |
| GTX2_TOBAC | Probable glutathione S-transferase | 157 | 194.2 | 0.00031 |
| GTX1_SOLTU | Probable glutathione S-transferase | 149 | 184.2 | 0.0011 |
| GTX1_TOBAC | Probable glutathione S-transferase | 147 | 181.4 | 0.0016 |
|  |  |  |  |  |
|  | $\quad$ Re-search with EF1G_HUMAN |  |  |  |
| The best scores are: |  | s-w | Z-score | E(58,709) |
| EF1G_HUMAN | Elongation factor 1 $\gamma($ EF-1 $\gamma)$ | 2977 | 3423.6 | $10^{-183}$ |
| EF1G_XENLA | Elongation factor 1 $\gamma$ (EF-1 $\gamma$ ) | 2370 | 2723.3 | $10^{-144}$ |
| EF1H_YEAST | Elongation factor $1 \gamma$ 2 (EF-1 $\gamma$ ) | 769 | 876.9 | $10^{-42}$ |
| EF1G_TRYCR | Elongation factor $1 \gamma$ (EF-1 $\gamma$ ) | 715 | 814.6 | $10^{-38}$ |
| SYV_HUMAN | valyl-tRNA synthetase | 440 | 483.4 | $10^{-20}$ |
| GTH1_MAIZE | Glutathione S-transferase I | 222 | 252.0 | $10^{-7}$ |
| GTH3_MAIZE | Glutathione S-transferase III | 193 | 218.2 | $10^{-5}$ |
| GTH1_WHEAT | Glutathione S-transferase 1 | 186 | 209.8 | $10^{-5}$ |
| GTH1_TOBAC | Glutathione S-transferase | 184 | 208.1 | $10^{-4}$ |
| GTY2_ISSOR | Glutathione S-transferase Y-2 | 175 | 198.8 | 0.00017 |
| GTH2_WHEAT | Glutathione S-transferase 2 | 175 | 192.6 | 0.00028 |
| GTX6_SOYBN | Probable glutathione S-transferase | 171 | 192.6 | 0.00037 |
| GTX2_TOBAC | Probable glutathione S-transferase | 169 | 190.8 | 0.0005 |
| GTT1_DROME | Glutathione S-transferase 1-1 | 162 | 182.9 | 0.0013 |

Figure 19: G-protein-coupled receptors


### 4.3 G-protein-coupled receptors

The G-protein-coupled receptors (GCRs) are one of the largest known gene families; members of the family transduce signals from light, peptides, cationic amines, lipid mediators, odors, and many more small molecules. An evolutionary tree that summarizes the diversity of this family is shown in Fig. 19. Based on hydrophobicity plots and the structure of bacteriorhodopsin (a protein that does not share significant similarity with members of this family), the GCRs are thought to contain seven transmembrane domains, so that the N-terminus of the proteins is extracellular, while the C-terminus is intra-
cellular.

Table 16: GCRs distant from human opsin

| The best scores are: |  | s-w | Z-score | E(58,649) |
| :--- | :--- | :--- | :--- | :--- |
| MC3R_RAT | melanocortin-3 receptor | 140 | 164.4 | 0.014 |
| OLF6_CHICK | olfactory receptor-like protein | 139 | 163.4 | 0.016 |
| MC3R_MOUSE | melanocortin-3 receptor | 139 | 163.2 | 0.016 |
| ML1A_XENLA | melatonin receptor type 1A | 133 | 161.4 | 0.02 |
| GU27_RAT | gustatory receptor GUST27 | 137 | 161.0 | 0.021 |
| AG2T_RAT | type-1C angiotensin II receptor | 132 | 159.2 | 0.027 |
| OLF2_RAT | olfactory receptor-like protein F12 | 135 | 158.5 | 0.03 |
| MAS_MOUSE | MAS proto-oncogene | 133 | 155.9 | 0.041 |
| PAFR_MACMU | platelet activating factor receptor | 130 | 155.6 | 0.043 |
| MAS_RAT | MAS proto-oncogene. | 131 | 153.5 | 0.056 |
| OLF2_CHICK | olfactory receptor-like protein C | 129 | 151.4 | 0.074 |
| CAR1_DICDI | cyclic AMP receptor 1 | 130 | 150.9 | 0.079 |
| YS96_CAEEL | hypothetical 110.4 KD protein | 133 | 147.7 | 0.12 |
| 5H2A_CAVPO | 5-hydroxytryptamine 2A receptor ( | 121 | 143.5 | 0.2 |
| PER4_RAT | prostaglandin E2 receptor EP4 | 124 | 142.1 | 0.24 |
| CAR3_DICDI | cyclic AMP receptor 3 | 124 | 142.0 | 0.25 |
| OLF4_CHICK | olfactory receptor-like protein c | 121 | 141.7 | 0.25 |
| ML1B_RAT | melatonin receptor type 1B | 115 | 141.6 | 0.26 |
| UL33_HSV7J | G-protein coupled receptor homolog U12 | 121 | 141.0 | 0.28 |
| OLF5_CHICK | olfactory receptor-like protein C | 120 | 140.5 | 0.3 |
| MAS_HUMAN | MAS proto-oncogene. | 120 | 140.2 | 0.31 |
| NU2M_CHOCR | NADH-ubiquinone oxidoreductase chain 2 | 122 | 139.5 | 0.34 |
| PER4_HUMAN | prostaglandin E2 receptor EP4 | 120 | 137.2 | 0.45 |
| OLF1_CHICK | olfactory receptor-like protein C | 117 | 136.8 | 0.48 |

Because GCRs have transmembrane domains, the highest scoring unrelated sequences are frequently other membrane proteins. Table 16 lists sequences from Swiss-Prot that have marginally significant matches with a human opsin sequence (there are more than 500 related sequences with expectations ranging from $0-0.01$ that are not shown). As with most divergent families, the question becomes, "how do I know that XXX is/is not a GCR?" This is more difficult with the GCRs, because they have long variable length loops in both their extracellular and intracellular domains.

As before, two strategies can be used to evaluate the hypothesis of homology: re-searching the library and statistical significance from shuffling. A search of the Swiss-Prot database reveals that MAS_HUMAN shares significant similarity $(E(58,500)<0.01)$ with $205 \mathrm{GCRs} ; 100$ additional scores with less statistical significance also belong to the GCR family before the first non-GCR is encountered. In contrast, the highest ranking scores from the NU2M_CHOCR are (more than 100 NADH oxidoreductase sequences are not shown):

```
The best scores are: s-w Z-score E(58649)
NU2M_CHOCR NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 2 ( 497) 3181 2999.6 1.7e-160
NUON_RHOCA NADH DEHYDROGENASE I CHAIN N (EC 1.6.5. ( 478) 928 877.8 2.6e-42
NU2C_MARPO NADH-PLASTOQUINONE OXIDOREDUCTASE CHAIN ( 501) 827 782.4 5.3e-37
NU2M_PODAN NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 2 ( 556) 788 745.0 6.4e-35
NU2M_ANOGA NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 2 ( 341) 460 439.2 6.9e-18
NU2M_RAT NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 2 (E ( 345) 393 376.1 2.3e-14
NU2M_CROLA NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 2 ( 348) 312 299.7 4.1e-10
NU5M_XENLA NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 5 ( 604) 230 218.9 1.3e-05
NDHF_BACSU NADH DEHYDROGENASE SUBUNIT 5 (EC 1.6.5. ( 505) 190 182.4 0.0014
COX1_LEITA CYTOCHROME C OXIDASE POLYPEPTIDE I ( 549) 154 147.9 0.12
Y825_HAEIN HYPOTHETICAL PROTEIN HIO825. ( 244) 145 144.7 0.17
CCMF_RHIME CYTOCHROME C-TYPE BIOGENESIS PROTEIN CY ( 676) 152 144.7 0.18
NU5M_ANOAR NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 5 ( 266) 145 144.2 0.19
RFBX_SALTY RFBX PROTEIN. ( 430) 148 143.9 0.19
ATP6_OENBE ATP SYNTHASE A CHAIN (EC 3.6.1.34) ( 281) 141 140.0 0.32
YMO4_PARTE HYPOTHETICAL 18.8 KD PROTEIN (ORF4). ( 156) 135 138.3 0.4
YC43_ODOSI HYPOTHETICAL 30.1 KD PROTEIN YCF43 (ORF ( 263) 138 137.7 0.43
YJFS_ECOLI HYPOTHETICAL 53.6 KD PROTEIN IN AIDB-RP ( 488) 142 137.4 0.45
NU4M_ANOAR NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 4 ( 221) 135 136.0 0.54
COP_CLOPE COPY NUMBER PROTEIN (ORF4). ( 198) 134 135.7 0.55
YJG2_YEAST HYPOTHETICAL 94.9 KD PROTEIN IN MRPL8-N ( 830) 143 134.8 0.62
CAPE_STAAU CAPE PROTEIN. ( 440) 138 134.3 0.66
OPSD_MOUSE RHODOPSIN. ( 348) 134 132.1 0.89
```

The results from the MAS_HUMAN and NU2M_CHOCR, which show that MAS_HUMAN is clearly a member of the GCR family, contrast with the statistical significance calculated with the PRSS program. Comparing the ОOHU with RTA_RAT score with the distribution of scores calculated after shuffling RTA_RAT 1000 times with a local window of 20 suggests that the unshuffled score (109) is expected 6 times in 1000 shuffles. In contrast, the NU2M_CHOCR score is expected only 1.7 times in 1000 shuffles. From this perspective, the NU2M_CHOCR score is somewhat more significant, but, in fact, neither similarity score is statistically significant. It is not until MAS_HUMAN is compared with other members of the family, e.g. the angiotensin, fMet-Leu-Phe, thrombin, or substance-P receptors with E-values from $10^{-12}-10^{-6}$, that the relationship is apparent.

Table 3.3 compares the statistical significance inferred from database searches with those determined by Monte-Carlo shuffling. As expected, the significance of the scores when compared with locally (window) shuffled sequences is 10 -fold lower than the comparison with globally shuffled scores. It is unclear how to compare the expectation from shuffles with the expectation from a search. In the table, the expectation from a search of a 43,000 entry library is compared to the expectation from 1,000 shuffles. For global shuffles, the expectations are quite comparable while local shuffles are more conservative, yet all but one of the similarity scores judged significant from the database search are still significant when compared with the local-shuffle distribution.

Nevertheless, these examples show both that current statistical models for the similarity scores of unrelated sequences are quite accurate, but also that homologous sequences frequently do not share significant pair-wise similarity scores. Thus, a lack of statistical significance cannot be used to infer non-homology, but strong statistical significance is a good indicator of common ancestry.

Figure 20: Internal duplications in calmodulin


Comparison of human calmodulin with itself. Each diagonal line represents a potential local alignment of calmodulin with itself. Values below the diagonal lines show the number of identities and length of the aligned region (e.g. 33/102) and the expectation value for the similarity score of the alignment.

## 5 Repeated structures in proteins

So far, we have focussed on the identification and statistics of the single most significant similarity score shared by two sequences. As can be seen in Fig. 10B, however, there are frequently several nonoverlapping local alignments with optimal similarity scores. In addition, there can be non-overlapping sub-optimal alignments with significant scores that can be used to infer the duplication events that gave rise to the protein sequence. An algorithm for the best $N$ non-overlapping local alignments was described by (Waterman \& Eggert, 1987).

Figs. 20 and 21 show a graphical plot of the local similarities within the calmodulin calcium binding protein. Calmodulin contains four EF -hand $\mathrm{Ca}^{+}$-binding domains that are well conserved. The highest scoring alignment in Fig. 21 aligns domains $A-B$ with $C-D$; the second highest aligns $A-B-C$ with $B-C-D$; the third aligns $A$ with $D$.

A similar pattern of local similarity can be seen in Fig. 22, which shows the mosaic relationship between the EGF-precursor and the LDL-receptor.

Figure 21: Calmodulin internal alignments
Comparison of:


Figure 22: Mosaic domains shared by the EGF-precursor and LDL-receptor


Some non-homologous structures, particularly proteins containing the coiled-coil structure, have a periodic structure that is easily seen in local similarity plots. Fig. 23 shows local similarities in tropomyosin. All the alignments shown have local similarity scores greater than 120, and each would be significant in a conventional database search.

## 6 Summary

Protein sequence comparison is the most powerful tool available today for inferring structure and function from sequence because of the constraints of protein evolution-a protein fold into a functional structure. Protein sequence similarity can routinely be used to infer relationships between proteins that last shared a common ancestor $1-2.5$ billion years ago. Our ability to identify distantly related proteins has improved over the past five years with the development of accurate statistical estimates, which have provided better normalization methods, and with the use of optimized scoring parameters. In using sequence similarity to infer homology, one should remember:

1. Always compare protein sequences if the genes encode proteins. Protein sequence comparison will typically double the look back time over DNA sequence comparison.
2. While most sequences that share statistically significant similarity are homologous, many distantly related homologous sequences do not share significant homology. (Low complexity re-

Figure 23: Coiled-coil structures share local similarity

gions display significant similarity in the absence of homology). Homologous sequences are usually similar over an entire sequence or domain. Matches that are more than $50 \%$ identical in a $20-40$ amino acid region occur frequently by chance.
3. Homologous sequences share a common ancestor, and thus a common protein fold. Depending on the evolutionary distance and divergence path, two or more homologous sequences may have very few absolutely conserved residues. However, if homology has been inferred between A and $\mathbf{B}$, between $\mathbf{B}$ and $\mathbf{C}$, and between $\mathbf{C}$ and $\mathbf{D}, \mathbf{A}$ and $\mathbf{D}$ must be homologous, even if they share no significant similarity.
4. Similarity searching techniques can be improved either by increasing the ability of a method to recognize distantly related sequences-increased sensitivity-or by lowering scores for unrelated sequences-increased selectivity. Since there are generally 1000 -times more unrelated than related sequences in a sequence database, improvements that reduce the scores of unrelated sequences can have dramatic effects. The most dramatic improvements in comparison methods recently have used this approach.

## References

Altschul, S. F. (1991). Amino acid substitution matrices from an information theoretic perspective. J. Mol. Biol. 219, 555-565.

Altschul, S. F. (1993). A protein alignment scoring system sensitive at all evolutionary distances. J. Mol. Evol. 36, 290-300.

Altschul, S. F., Boguski, M. S., Gish, W. \& Wootton, J. C. (1994). Issues in searching molecular sequence databases. Nature Genet. 6, 119-129.

Altschul, S. F., Gish, W., Miller, W., Myers, E. W. \& Lipman, D. J. (1990). A basic local alignment search tool. J. Mol. Biol. 215, 403-410.

Collins, J. F., Coulson, A. F. W. \& Lyall, A. (1988). The significance of protein sequence similarities. Comp. Appl. Biosci. 4, 67-71.

Dayhoff, M., Schwartz, R. M. \& Orcutt, B. C. (1978). A model of evolutionary change in proteins. In Atlas of Protein Sequence and Structure, (Dayhoff, M., ed.), vol. 5, supplement 3, pp. 345-352. National Biomedical Research Foundation Silver Spring, MD.

Doolittle, R. F., Feng, D. F., Johnson, M. S. \& McClure, M. A. (1986). Relationships of human protein sequences to those of other organisms. Cold Spring Harb. Symp. Quant. Biol. 51, 447-455.

Gilbert, W. \& Glynias, M. (1993). On the ancient nature of introns. Gene, 135, 137-144.
Henikoff, S. \& Henikoff, J. G. (1992). Amino acid substitutions matrices from protein blocks. Proc. Natl. Acad. Sci. USA, 89, 10915-10919.

Henikoff, S. \& Henikoff, J. G. (1993). Performance evalutation of amino-acid substitution matrices. Proteins, 17, 49-61.

Johnson, M. S. \& Overington, J. P. (1993). A structural basis for sequence comparisons. an evaluation of scoring methodologies. J. Mol. Biol. 233, 716-738.

Jones, D. T., Taylor, W. R. \& Thornton, J. M. (1992). The rapid generation of mutation data matrices from protein sequences. Comp. Appl. Biosci. 8, 275-282.

Karlin, S. \& Altschul, S. F. (1990). Methods for assessing the statistical significance of molecular sequence features by using general scoring schemes. Proc. Natl. Acad. Sci. USA, 87, 2264-2268.

Karlin, S. \& Altschul, S. F. (1993). Applications and statistics for multiple high-scoring segments in molecular sequences. Proc. Natl. Acad. Sci USA, 90, 5873-5877.

Mott, R. (1992). Maximum-likelihood estimation of the statistical distribution of smith-waterman local sequence similarity scores. Bull. Math. Biol. 54, 59-75.

Needleman, S. \& Wunsch, C. (1970). A general method applicable to the search for similarities in the amino acid sequences of two proteins. J. Mol. Biol. 48, 444-453.

Nei, M. (1987). Molecular Evolutionary Genetics. Columbia Univ. Press, New York, NY.
Pearson, W. R. (1995). Comparison of methods for searching protein sequence databases. Prot. Sci. 4, 1145-1160.

Pearson, W. R. \& Lipman, D. J. (1988). Improved tools for biological sequence comparison. Proc. Natl. Acad. Sci. USA, 85, 2444-2448.

Smith, T. F. \& Waterman, M. S. (1981). Identification of common molecular subsequences. J. Mol. Biol. 147, 195-197.

Stoltzfus, A., Spencer, D. F., Zuker, M., Logsdon, J. M. \& Doolittle, W. F. (1994). Testing the intron theory of genes: the evidence from protein structure. Science, 265, 202-207.

Waterman, M. S. \& Eggert, M. (1987). A new algorithm for best subsequences alignment with application to tRNA-rRNA comparisons. J. Mol. Biol. 197, 723-728.

## 7 Suggested Reading

### 7.1 General Protein evolution

R. F. Doolittle, D. F. Feng, M. S. Johnson, and M. A. McClure. Relationships of human protein sequences to those of other organisms. Cold Spring Harb. Symp. Quant. Biol., 51:447-455, 1986.
P. Green, D. Lipman, L. Hillier, R. Waterston, D. States, and J. M. Claverie. Ancient conserved regions in new gene sequences and the protein databases. Science, 259:1711-1716, 1993.

### 7.1.1 Introns Early/Late

W. Gilbert and M. Glynias. On the ancient nature of introns. Gene, 135:137-144, 1993.
A. Stoltzfus, D. F. Spencer, M. Zuker, J. M. Logsdon, and W. F. Doolittle. Testing the intron theory of genes: the evidence from protein structure. Science, 265:202-207, 1994.

### 7.2 Alignment methods

### 7.2.1 Algorithms

S. Needleman and C. Wunsch. A general method applicable to the search for similarities in the amino acid sequences of two proteins. J. Mol. Biol., 48:444-453, 1970.
T. F. Smith and M. S. Waterman. Identification of common molecular subsequences. J. Mol. Biol., 147:195-197, 1981.
W. R. Pearson and W. Miller. Dynamic programming algorithms for biological sequence comparison. In L. Brand and M. L. Johnson, editors, Meth. Enz., volume 210, pages 575-601. Academic Press, San Diego, 1992.
D. J. Lipman and W. R. Pearson. Rapid and sensitive protein similarity searches. Science, 227:14351441, 1985.
W. R. Pearson and D. J. Lipman. Improved tools for biological sequence comparison. Proc. Natl. Acad. Sci. USA, 85:2444-2448, 1988.
W. R. Pearson. Rapid and sensitive sequence comparison with FASTP and FASTA. In R. F. Doolittle, editor, Meth. Enz., volume 183, pages 63-98. Academic Press, San Diego, 1990.
S. F. Altschul, W. Gish, W. Miller, E. W. Myers, and D. J. Lipman. A basic local alignment search tool. J. Mol. Biol., 215:403-410, 1990.
W. R. Pearson. Comparison of methods for searching protein sequence databases. Prot. Sci., 4:11451160, 1995.
W. R. Pearson. Empirical statistical estimates for sequence similarity searches J. Mol. Biol. 276:7184, 1998.

### 7.2.2 Scoring methods

M. Dayhoff, R. M. Schwartz, and B. C. Orcutt. A model of evolutionary change in proteins. In M. Dayhoff, editor, Atlas of Protein Sequence and Structure, volume 5, supplement 3, pages 345-352. National Biomedical Research Foundation, Silver Spring, MD, 1978.
S. F. Altschul. Amino acid substitution matrices from an information theoretic perspective. J. Mol. Biol., 219:555-565, 1991.
D. T. Jones, W. R. Taylor, and J. M. Thornton. The rapid generation of mutation data matrices from protein sequences. Comp. Appl. Biosci., 8:275-282, 1992.
S. Henikoff and J. G. Henikoff. Performance evalutation of amino-acid substitution matrices. Proteins, 17:49-61, 1993.

### 7.3 Evaluating matches - statistics of similarity scores

R. F. Doolittle. Similar amino acid sequences: chance or common ancestry? Science, 214:149-159, 1981.
W. R. Pearson. Identifying distantly related protein sequences. Cur. Opinion in Struct. Biol., 1:321326, 1991.
S. Karlin, P. Bucher, V. Brendel, and S. F. Altschul. Statistical methods and insights for protein and DNA sequences. Ann. Rev. of Biophys. Biophys. Chem., 20:175-203, 1991.
S. F. Altschul, M. S. Boguski, W. Gish, and J. C. Wootton. Issues in searching molecular sequence databases. Nature Genet., 6:119-129, 1994.
S. F. Altschul, and W. Gish. Local Alignment Statistics Meth. Enzymol., 266:460-480, 1996.


[^0]:    *FAX: (804) 924-5069; email: wrp@ virginia.EDU

[^1]:    ${ }^{a}$ PAMs, point accepted mutations. ${ }^{b}$ Useful lookback time, 360 PAMs, $15 \%$ identity.
    ${ }^{c}$ Millions of years. ${ }^{d}$ Billions of years. Adapted from Doolittle et al., 1986

[^2]:    ${ }^{1}$ The z-scores plotted have a mean of 50 and a standard deviation of 10 .

[^3]:    ${ }^{2}$ For genomic DNA sequences, there is no logical alternative.

[^4]:    ${ }^{2} O(N d)$, where $N$ is the length of a sequence and $d$ is the number of differences between the two sequences.

