Typically, to be “biologically related” means to share a common ancestor. In biology, we call this **homologous**. Two proteins sharing a common ancestor are said to be **homologs**. Homology often implies structural similarity & sometimes (not always) sequence similarity. A **statistically significant sequence or structural similarity** can be used to infer homology (common ancestry).

![Example of homologs: Myoglobin & Hemoglobin](http://en.wikipedia.org/wiki/File:Myoglobin.png & File:1GZX_Haemoglobin.png)

In practice, searching for sequence or structural similarity is one of the most powerful computational approaches for discovering functions for genes, since we can often glean many new insights about a protein based on what is known about its homologs.

Here's an example from my own lab, where we discovered that myelinating the neurons in your brain employs the same biochemical mechanism used by bacteriophages to make capsids. The critical breakthrough was recognizing that the human and phage proteins contained homologous domains.

![Example of homologous domains](Li, Park, Marcotte PLoS Biology (2013))
Sequence alignment algorithms such as BLAST, PSI-BLAST, FASTA, and the Needleman–Wunsch & Smith-Waterman algorithms arguably comprise some of the most important driver technologies of modern biology and underlie the sequencing revolution.

So, let’s start learning bioinformatics algorithms by learning how to align two protein sequences.

Live demo:

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MVLSPADKTNVKAAWGKVGAHAGEYGAELERMFLSFPTTTKTYFPHFDLSHGSAQVKGHG
KKVADALTNAHAVVDDMPNALSALSDLHAHKLRVDPVNFKLLSHCLLVTLAAHLPAEFTP
AVHASLDKFLASVSTVLT5KYR

Title: All non-redundant GenBank CDS translations+PDB+SwissProt+PIR+PRF excluding environmental samples from WGS projects
Molecule Type: Protein
Update date: 2012/02/28
Number of sequences: 17,362,047
Protein sequence alignment

To align two sequences, we need to perform 3 steps:

1. We need some way to decide which alignments are better than others.
   For this, we'll invent a way to give the alignments a “score” indicating their quality.

2. Align the two proteins so that they get the best possible score.

3. Decide if the score is “good enough” for us to believe the alignment is biologically significant.
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We’ll treat mutations as independent events.

This allows us to create an **additive scoring scheme**. The score for a sequence alignment will be the sum of the scores for aligning each of the individual positions in two sequences.

What kind of mutations should we expect?
**Substitutions, insertions** and **deletions**.

Insertions and deletions can be treated as equivalent events by considering one or the other sequence as the reference, and are usually called **gaps**.

Let’s consider two models:

First, a **random** model, where amino acids in the sequences occur independently at some given frequencies.

The probability of observing an alignment between \( x \) and \( y \) is just the product of the frequencies (\( q \)) with which we find each amino acid.

We can write this as:

\[
P(x, y \mid R) = \prod_i q_{x_i} \prod_j q_{y_j}
\]
Second, a **match** model, where amino acids at a given position in the alignment arise from some common ancestor with a probability given by the joint probability $p_{ab}$.

So, under this model, the probability of the alignment is the product of the probabilities of seeing the individual amino acids aligned.

We can write that as:

$$P(x, y \mid M) = \prod_i p_{x_i y_i}$$

What’s this mean? What does the capital pi mean again?

To decide which model better describes an alignment, we’ll take the ratio:

$$\frac{P(x, y \mid M)}{P(x, y \mid R)} = \frac{\prod_i p_{x_i y_i}}{\prod_i q_{x_i} \prod_j q_{y_j}} = \prod_i \frac{p_{x_i y_i}}{q_{x_i} q_{y_j}}$$

What did these mean again?

Such a ratio of probabilities under 2 different models is called an **odds ratio**.

Where else have you heard odds ratios used?

Basically: if the ratio > 1, model $M$ is more probable
if < 1, model $R$ is more probable.
Now, to convert this to an additive score $S$, we can simply take the logarithm of the odds ratio (called the log odds ratio): 

$$ S = \sum_i s(x_i, y_i) $$

This is just the score for aligning one amino acid with another amino acid:

$$ s(a, b) = \log \left( \frac{P_{ab}}{P_a P_b} \right) $$

Here written $a$ and $b$ rather than $x_i$ and $y_i$ to emphasize that this score reflects the inherent preference of the two amino acids ($a$ and $b$) to be aligned.

Almost done with step 1...

The last trick:
Take a big set of pre-aligned protein sequence alignments (that are correct!) and measure all of the pairwise amino acid substitution scores (the $s(a,b)$'s). Put them in a 20x20 amino acid substitution matrix:
This is the BLOSUM50 matrix.
(The numbers are scaled & rounded off to the nearest integer):

What’s the score for aspartate (D) aligning with itself?
How about aspartate with phenylalanine (F)? Why?

Using this matrix, we can score any alignment as the sum of scores of individual pairs of amino acids.

For example, the top alignment in our earlier example:

F1gA1  EAGNVLKPGRLDTLPPFTVLGDAISLRDLSPQDFQIQLGRTQQRVAKGQVRQVYIASGD
++K++GRLDTEPF++L+ N A+SLR ++QP+ R+W+KAGQ V V+A G+
F1gA2  TLQDIKHKQGRLDTPPLAEGPFQAGSVLAQHQPQRQV1NQPLTRNLRLNLA+KAGQDQVVGQLGSE

gets the score:

\[ S(\text{F1gA1}, \text{F1gA2}) = -1 - 2 - 2 + 2 + 4 + 6 + ... = 186 \]

We also need to penalize gaps. For now, let’s just use a constant penalty \( d \) for each amino acid gap in an alignment, \( i.e. \):

the penalty for a gap of length \( g \) = \(-g \times d\)
To align two sequences, we need to perform 3 steps:

1. We need some way to decide which alignments are better than others. For this, we’ll invent a way to give the alignments a “score” indicating their quality.

2. **Align the two proteins so that they get the best possible score.**

3. Decide if the score is “good enough” for us to believe the alignment is biologically significant.

We’ll use something called **dynamic programming.**

This is **mathematically guaranteed** to find the best scoring alignment, and uses **recursion.** This means problems are broken into sub-problems, which are in turn broken into sub-problems, etc, until the simplest sub-problems can be solved.

We’re going to find the best **local** alignment—the best matching internal alignment—without forcing all of the amino acids to align (i.e. to match **globally**).

| i.e., this | ATGCAT  
| ATGCAT |
| Not this | ACGTTATGCATGACGTA  
| --- | ATGCAT ----T-- |
Here's the main idea:
We’ll make a path matrix, showing the possible alignments and their scores. There are simple rules for how to fill in the matrix. **This will test all possible alignments & give us the top-scoring alignment between the two sequences.**

<table>
<thead>
<tr>
<th>i=0</th>
<th>H</th>
<th>E</th>
<th>A</th>
<th>G</th>
<th>A</th>
<th>W</th>
<th>G</th>
<th>H</th>
<th>E</th>
<th>i=n</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>P</td>
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<td>j=0</td>
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<tr>
<td>y</td>
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<td>j=m</td>
</tr>
</tbody>
</table>

Here are the rules:
For a given square in the matrix $F(i,j)$, we look at the squares to its left $F(i-1,j)$, top $F(i,j-1)$, and top-left $F(i-1,j-1)$. Each should have a score.

We consider 3 possible events & choose the one scoring the highest:

1. $x_i$ is aligned to $y_j$ $\quad F(i-1,j-1) + s(x_i, y_j)$
2. $x_i$ is aligned to a gap $\quad F(i-1,j) - d$
3. $y_j$ is aligned to a gap $\quad F(i,j-1) - d$

For this example, we’ll use $d = 8$. We also set the left-most & top-most entries to zero.
Just two more rules:

If the score is negative, set it equal to zero.

At each step, we also keep track of which event was chosen by drawing an arrow from the cell we just filled back to the cell which contributed its score to this one.

That’s it! Just repeat this to fill the entire matrix.
Here we go! Start with the borders & the first entry.

<table>
<thead>
<tr>
<th></th>
<th>H</th>
<th>E</th>
<th>A</th>
<th>G</th>
<th>A</th>
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<th>H</th>
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<th>E</th>
</tr>
</thead>
<tbody>
<tr>
<td>P</td>
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</tbody>
</table>

Why is this zero? What’s the score from our BLOSSUM matrix for substituting H for P?

Next round!

<table>
<thead>
<tr>
<th></th>
<th>H</th>
<th>E</th>
<th>A</th>
<th>G</th>
<th>A</th>
<th>W</th>
<th>G</th>
<th>H</th>
<th>E</th>
<th>E</th>
</tr>
</thead>
<tbody>
<tr>
<td>P</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
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<td>H</td>
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</tbody>
</table>

Terrible! Again, none of the possible give positive scores. We have to go a bit further in before we find a positive score...
A few more rounds, and a positive score at last!

<table>
<thead>
<tr>
<th>H</th>
<th>E</th>
<th>A</th>
<th>G</th>
<th>A</th>
<th>W</th>
<th>G</th>
<th>H</th>
<th>E</th>
<th>E</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
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<td>0</td>
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<td>0</td>
</tr>
</tbody>
</table>

P 0 0 0 0
A 0 0 0 5
W 0 0 0
H 0
E 0
A 0
E 0

How did we get this one?

& a few more rounds...

<table>
<thead>
<tr>
<th>H</th>
<th>E</th>
<th>A</th>
<th>G</th>
<th>A</th>
<th>W</th>
<th>G</th>
<th>H</th>
<th>E</th>
<th>E</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
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</tr>
</tbody>
</table>

P 0 0 0 0 0
A 0 0 0 5 0
W 0 0 0 0 2
H 0 10 2 0 0
E 0
A 0
E 0

What does this mean?
The whole thing filled in!

Now, find the optimal alignment using a **traceback** process:

- Look for the highest score, then follow the arrows back.
- The alignment “grows” from right to left.
To align two sequences, we need to perform 3 steps:

1. We need some way to decide which alignments are better than others. For this, we’ll invent a way to give the alignments a “score” indicating their quality.

2. Align the two proteins so that they get the best possible score.

3. Decide if the score is “good enough” for us to believe the alignment is biologically significant.
This algorithm always gives the best alignment.

Every pair of sequences can be aligned in some fashion.

So, when is a score “good enough”?

How can we figure this out?

Here’s one approach:

Shuffle one sequence. Calculate the best alignment & its score. Repeat 1000 times.

If we never see a score as high as the real one, we say the real score has <1 in a 1000 chance of happening just by luck.

But if we want something that only occurs < 1 in a million, we’d have to shuffle 1,000,000 times...
Luckily, alignment scores follow a well-behaved distribution, the **extreme value distribution**, so we can do a few trials & fit to this.

\[ p(\text{max score} \leq X) \approx e^{-kNe^{\lambda(X-\mu)}} \]

This p-value gives the significance of your alignment. But, if we search a database and perform many alignments, **we still need something more** (next time).