

## Sequence Alignment

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## This Lecture

- Approximate String Matching
- Edit distance and alignment
- Computing global alignments
- Local alignment


## Gene Function

- A fundamental principle of bioinformatics
- The function of a protein depends on its physical structure
- The physical structure depends on the protein sequence
- The protein sequence depends on the gene sequence
- If the sequence of two genes is only slightly different, so will be the protein sequence
- If the sequence of two proteins is only slightly different, so will be their structure
- If the structure of two proteins is only moderately different, they likely have the same (or at least share some) function
- Studying the sequence of genes allows the generation of hypotheses about the function of the proteins they encode


## How Genes Evolve

- Evolution, sequences, and function
- Any two species $X_{1}, X_{2}$ have a common ancestor $A$
- Any gene $G$ from $A$ will undergo independent evolution in $X_{1}$ and $X_{2}$, leading to genes $G_{1}$ and $G_{2}$
- The more similar $G_{1}$ and $G_{2}$ are, the more likely do they still have the same function (that of G)
- For any two genes of non-trivial length, the chance that they have a very similar sequence by chance is extremely small
- Corollary: If genes $G_{1}$ and $G_{2}$ from species $X_{1}$ and $X_{2}$ today are very similar, they most likely derive from the same ancestor $A$ and most likely have the same function
- How can we quantify this?

AGGTTGATAGCCGA G

AGGTTTATAGCTCGA
AGCTTGAT_GCCGA

## Basic Evolutionary Events

- The simplest model: Single bases can be replaced (R), inserted (I), or deleted (D) (or kept (M))
- Any changes must be explained by sequences of $I, D, R$
- I.e., by singular evolutionary events accumulating over time
- We call this an edit script
- Very simple yet quite powerful model
- One more simplification



## Example: Eyeless (ey)

- Family of genes identified first in Drosophila
- When activated in arbitrary cells, non-functional eyes start to grow at various places of the body
- ey is a "master gene" - controls a cascade of activations of other genes eventually leading to eye development
- Also inflicted with several other neural developments
- Zitat [NCBI Gene]
- Enables DNA-binding transcription factor activity, ... Involved in several processes, including adult walking behavior; nervous system development; and regulation of insulin-like growth factor receptor signaling pathway... expressed in several structures, including central nervous system; embryonic head; eyeantennal disc; neuroblasts; and photoreceptor. Human ortholog(s) of this gene implicated in bilateral optic nerve hypoplasia; eye disease (multiple); glucose intolerance; and paranoid schizophrenia...


## Eyes



D


Red: Only shadow Blue: Lenses etc. Green: Mirrors

Oval: Compound eyes Rectangle: Single chamber

Source: Treisman (2004).

- Eyes probably are an example of convergent evolution
- However, genes controlling eye development are highly conserved across a wide range of species


## Homologues of "eyeless isoform D" (DM)

```
(3) job:201105063F73IVJYOG in UniProtKB by taxonomy - Mozilla Firefox
Datei Bearbeiten Ansicht Chronik Lesezeichen Extras Hilfe
8 eyeless uniprot drosophila - Google-Suche x .. job:201105063F73IVJYOG in UniProtKB b... x
& \
% Meistbesuchte Seiten \square Nachsehen \square Frequent 啕 wBi Lehre s% Google \square News
    Protein Knowledgebase (UniProtKB)\vee iob:201105063F73|JJY0G
```

229 results for job:201105063F73IVJY0G in UniProtKB browsing bytaxonomy
国 View result list

+ Ceractinomorpha (4)
- Eumetazoa (225) Bilateria (224)
- Coelomata (213)
- Deuterostomia (135)

Chordata (129)

+ Branchiostoma
${ }^{+}$Branchiostoma (10)
+ Urochordata (8)
- vertebrata (111)
- Euteleostomi (109)
+ Clupeocephala (30)
- Tetrapoda (79)

Amniota (56)

+ Neognathae (8)
- Theria (48)
- Eutheria
\& Euarchontoglires (41)

Saccoglossus kowalevskii (Acorn worm)
Protostomia (78)

+ Annelida
+ Arthropoda
+ Decapodiformes
Lineus sanguineus (Ribbon worm) (1)
- Platyhelminthes (5)

Dugesiidae (3)
Schistosoma manson
Brachionus plicatilis (Marine rotifer) (Brachionus muelleri)
(1)

Nematoda (roundworms)
Caenorhabditis (3)
MFTLQPTPTAIGTVVPPWSAGTLIERLPSLEDMAHKDNVIAMRNLPCLGTAGGSGLG GIAGKPSPTMEAVEASTASHPHSTSSYFATTYYHLTDDECHSGVNQLGGVFVGGRPL PDSTRQKIVELAHSGARPCDISRILQVSNGCVSKILGRYYETGSIRPRAIGGSKPRVAT AEVVSKISQYKRECPSIFAWEIRDRLLQENVCTNDNIPSVSSINRVLRNLAAQKEQQST GSGSSSTSAGNSISAKVSVSIGGNVSNVASGSRGTLSSSTDLMQTATPLNSSESGGAS NSGEGSEQEAIYEKLRLLNTQHAAGPGPLEPARAAPLVGQSPNHLGTRSSHPQLVHG NHQALQQHQQQSWPPRHYSGSWYPTSLSEIPISSAPNIASVTAYASGPSLAHSLSPP NDIESLASIGHQRNCPVATEDIHLKKELDGHQSDETGSGEGENSNGGASNIGNTEDD QARLILKRKLQRNRTSFTNDQIDSLEKEFERTHYPDVFARERLAGKIGLPEARIQVWFS NRRAKWRREEKLRNQRRTPNSTGASATSSSTSATASLTDSPNSLSACSSLLSGSAGG PSVSTINGLSSPSTLSTNVNAPTLGAGIDSSESPTPIPHIRPSCTSDNDNGRQSEDCRR VCSPCPLGVGGHQNTHHIQSNGHAQGHALVPAISPRLNFNSGSFGAMYSNMHHTAL SMSDSYGAVTPIPSFNHSAVGPLAPPSPIPQQGDLTPSSLYPCHMTLRPPPMAPAHHH IVPGDGGRPAGVGLGSGQSANLGASCSGSGYEVLSAYALPPPPMASSSAADSSFSAAS SASANVTPHHTIAQESCPSPCSSASHFGVAHSSGFSSDPISPAVS..

- 250 most similar protein sequences in UniProt
- Sequence identities all >50\%,
- All p-Values < 1E-50


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## Edit Scripts and Edit Distances

- Definition
- Let A, B $\in \Sigma^{*}=\Sigma \cup^{\prime \prime}$ "
- An edit script e is a sequence of operations $I, D, R, M$
- $e$ is an edit script for $A$ and $B$ iff $e(A)=B$
- Slightly underdetermined - which replacement? Which base to insert?
- The length of an edit script is the number of $I, D, R$ it contains
- The edit distance between $A$ and $B$ is the length of the shortest edit script for $A$ and $B$
- Remarks
- If we know $e(A)=B$, determining $e^{\prime}$ with $e^{\prime}(B)=A$ is trivial
- The shortest edit script is not unique, but its length is
- MIMMMR

A_TGTA
AGTGTC

IRMMMDI
ATGTA
AGTGT_ $\overline{\mathbf{C}}$

## Alignment

- Edit scripts are intuitive from an evolutionary point-of-view, but somewhat clumsy from a computational point-of-view
- Definition
- $A$ (global) alignment of strings $A, B$ is an arrangement of $A$ and $B$, enriched with „_" at arbitrary positions, under each other such that no column contains two "_"
- The score of an alignment is the number of "_" plus the number of mismatching columns it contains
- The alignment distance between $A$ and $B$ is the minimal score of any alignment of $A$ and $B$
- Edit distance and alignment distance are essentially identical
- Examples
$-{ }_{-}^{\text {AGTGTC }}$
Score:


5


2


2

## A Visual Approach: Dotplots

- $A$ dotplot of two strings $A, B$ is a matrix $M$ with
- The i'th character in $A$ is represented by the ith column
- The j'th character in B is represented by the j'th row
- M[i,j]=1 (blue) iff $A[i]=B[j]$

|  | A | T | G | C | G | G | T | G | C | A | A | T | G |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| A |  |  |  |  |  |  |  |  |  |  |  |  |  |
| T |  |  |  |  |  |  |  |  |  |  |  |  |  |
| G |  |  |  |  |  |  |  |  |  |  |  |  |  |
| G |  |  |  |  |  |  |  |  |  |  |  |  |  |
| T |  |  |  |  |  |  |  |  |  |  |  |  |  |
| G |  |  |  |  |  |  |  |  |  |  |  |  |  |
| C |  |  |  |  |  |  |  |  |  |  |  |  |  |
| A |  |  |  |  |  |  |  |  |  |  |  |  |  |
| T |  |  |  |  |  |  |  |  |  |  |  |  |  |

## Dotplot and Identical Substrings

- How do identical substrings look like in a dotplot?

|  | A | T | G | C | G | G | T | G | C | A | A | T | G |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| A |  |  |  |  |  |  |  |  |  |  |  |  |  |
| T |  |  |  |  |  |  |  |  |  |  |  |  |  |
| G |  |  |  |  |  |  |  |  |  |  |  |  |  |
| G |  |  |  |  |  |  |  |  |  |  |  |  |  |
| T |  |  |  |  |  |  |  |  |  |  |  |  |  |
| G |  |  |  |  |  |  |  |  |  |  |  |  |  |
| C |  |  |  |  |  |  |  |  |  |  |  |  |  |
| A |  |  |  |  |  |  |  |  |  |  |  |  |  |
| T |  |  |  |  |  |  |  |  |  |  |  |  |  |


|  | A | T | G | C | G | G | T | G | C | A | A | T | G |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| A |  |  |  |  |  |  |  |  |  |  |  |  |  |
| T |  |  |  |  |  |  |  |  |  |  |  |  |  |
| G |  |  |  |  |  |  |  |  |  |  |  |  |  |
| G |  |  |  |  |  |  |  |  |  |  |  |  |  |
| T |  |  |  |  |  |  |  |  |  |  |  |  |  |
| G |  |  |  |  |  |  |  |  |  |  |  |  |  |
| C |  |  |  |  |  |  |  |  |  |  |  |  |  |
| A |  |  |  |  |  |  |  |  |  |  |  |  |  |
| T |  |  |  |  |  |  |  |  |  |  |  |  |  |

- Diagonals from up-left to down-right
- Longest diagonal is the longest common substring


## Alignments and Dotplots

- Every alignment of $A, B$ can be uniquely mapped into a path through $M$
- The path starts in the upper-left corner (coord: 0,0)
- Go through the alignment column by column
- Next column is " X, _" - move to the right
- Next column is " $\_, X^{\prime \prime}$ - move down
- Next column is " $X, Y$ " - move right-down

ATG_CGGTG_CAATG
ATGG_TGCA_T


ATGCGGTGCAATG
ATGGTGCCAT


## Examples



> ATGCGGTGCAATG ATG_GTGCA__T


- Clearly, the number $c(P)$ of 1's (blue cells) crossed in a diagonal step by a path $P$ is the same as $|\mathrm{P}|-\mathrm{e}(\mathrm{A}, \mathrm{B})$
- Finding the path that minimizes $|\mathrm{P}|-\mathrm{c}(\mathrm{P})$ (or maximizes $c(P)$ ) solves the problem of computing the edit distance


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## Algorithm

- How do we compute the edit distance of two strings?
- Naïve: Enumerate all paths, compute $c(P)$ for each

- Bad news: There exist $>3^{\min (m, n)}$ paths
- Good news: We can compute e(A,B) with ~3*m*n operations


## Enumerating all Paths Recursively



## The naïve (recursive) Way

- Observation
- Let $|A|=n,|B|=m$
- Let $d(i, j)=e(A[\ldots i], B[\ldots j])$ for $0 \leq i \leq n$ and $0 \leq j \leq m$ with $d(i, 0)=i$ and $d(0, j)=j$
- We can compute $e(A, B)=d(n, m)$ recursively as follows

$$
d(i, j)=\min \left\{\begin{array}{c}
d(i, j-1)+1 \\
d(i-1, j)+1 \\
d(i-1, j-1)+t(i, j)
\end{array}\right.
$$

$$
t(i, j)=\left\{\begin{array}{l}
1: \text { if } \quad A[i] \neq B[j] \\
0: \text { else }
\end{array}\right.
$$

## Algorithm

```
function d(i,j) {
    if (i = 0) return j;
    else if (j = 0) return i;
    else
\[
\begin{aligned}
\text { return min }( & d(i, j-1)+1, \\
& d(i-1, j)+1, \\
& d(i-1, j-1)+t(A[i], B[j])) ;
\end{aligned}
\]
}
function t(c)
    if (c
    else
}
```


## What is Happening?



## Much Redundant Computation



There are only $\sim n * m$ different parameter combinations

## Dynamic Programming - Using a Table

- Instead of computing top-down (from $n, m$ ), we compute all different values for $\mathrm{d}(\mathrm{i}, \mathrm{j})$ bottom-up
- We store all values in a table
- We can immediately "compute" $\mathrm{d}(\mathrm{i}, 0)$ and $\mathrm{d}(0, \mathrm{j})$
- Which values can we compute next?



## Example

$$
d(i, j)=\min \left\{\begin{array}{c}
d(i, j-1)+1 \\
d(i-1, j)+1 \\
d(i-1, j-1)+t(i, j)
\end{array}\right\}
$$

|  |  | A | T | G | C | G | G | T |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 |
| A | 1 |  |  |  |  |  |  |  |
| T | 2 |  |  |  |  |  |  |  |
| G | 3 |  |  |  |  |  |  |  |
| G | 4 |  |  |  |  |  |  |  |


|  |  | A | T | G | C | G | G | T |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 |
| A | 1 | 0 |  |  |  |  |  |  |
| T | 2 |  |  |  |  |  |  |  |
| G | 3 |  |  |  |  |  |  |  |
| G | 4 |  |  |  |  |  |  |  |


|  |  | A | T | G | C | G | G | T |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 |
| A | 1 | 0 | 1 | 2 | 3 | 4 | 5 | 6 |
| T | 2 |  |  |  |  |  |  |  |
| G | 3 |  |  |  |  |  |  |  |
| G | 4 |  |  |  |  |  |  |  |


|  |  | A | T | G | C | G | G | T |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 |
| A | 1 | 0 | 1 | 2 | 3 | 4 | 5 | 6 |
| T | 2 | 1 | 0 | 1 | 2 | 3 | 4 | 5 |
| G | 3 |  |  |  |  |  |  |  |
| G | 4 |  |  |  |  |  |  |  |


|  |  | A | T | G | C | G | G | T |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 |
| A | 1 | 0 | 1 | 2 | 3 | 4 | 5 | 6 |
| T | 2 | 1 | 0 | 1 | 2 | 3 | 4 | 5 |
| G | 3 | 2 | 1 | 0 | 1 | 2 | 3 | 4 |
| G | 4 |  |  |  |  |  |  |  |


|  |  | A | T | G | C | G | G | T |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 |
| A | 1 | 0 | 1 | 2 | 3 | 4 | 5 | 6 |
| T | 2 | 1 | 0 | 1 | 2 | 3 | 4 | 5 |
| G | 3 | 2 | 1 | 0 | 1 | 2 | 3 | 4 |
| G | 4 | 3 | 2 | 1 | 1 | 1 | 2 | 3 |

## Finding the (an) optimal Alignment(s)

- Traceback
- We find the path from back to front
- Start at cell (n,m)
- See which cells were used to compute d(n,m)
- Walk any of these - finds one optimal path
- Walking all means finding all optimal paths
- Alternative: Store pointers while filling the table

|  |  | $A$ | $T$ | $G$ | $C$ | $G$ | $G$ | $T$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Q | 1 | 2 | 3 | 4 | 5 | $G$ | 7 |
| A | 1 | 0 | 1 | 2 | 3 | 4 | 5 | 6 |
| T | 2 | 1 | 0 | 1 | 2 | 3 | 4 | 5 |
| G | 3 | 2 | 1 | 0 | 1 | 2 | 3 | 4 |
| G | 4 | 3 | 2 | 1 | 1 | 1 | 2 | 3 |


|  |  | A | T | G | C | G | G | T |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Q | 1 | 2 | 3 | 4 | 5 | 6 | 7 |
| A | 1 | Q | 1 | 2 | 3 | 4 | 5 | 6 |
| T | 2 | 1 | 0 | 1 | 2 | 3 | 4 | 5 |
| G | 3 | 2 | 1 | 0 | 1 | 2 | 3 | 4 |
| G | 4 | 3 | 2 | 1 | 1 | 1 | 2 | 3 |


|  |  | A | T | G | C | G | G | T |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Q | 1 | 2 | 3 | 4 | 5 | 6 | 7 |
| A | 1 | Q | 1 | 2 | 3 | 4 | 5 | 6 |
| T | 2 | 1 | 0 | 1 | 2 | 3 | 4 | 5 |
| G | 3 | 2 | 1 | 0 | 1 | 2 | 3 | 4 |
| G | 4 | 3 | 2 | 1 | 1 | 1 | 2 | 3 |

## Complexity

- Building the table
- For every $\mathrm{d}(\mathrm{i}, \mathrm{j})$, we need to access three other cells and make some (constantly many) additions and comparisons
- There are $(m+1) *(n+1)$ cells
- Thus: ~3*m*n=0(m*n) operations
- Finding one optimal alignment
- We must walk from $(n, m)$ to $(1,1)$
- Such a path can have at most length m+n
- We cannot go wrong!
- Together: approximately m+n operations
- Together: $\mathrm{O}\left(\mathrm{m}^{*} \mathrm{n}\right)$ (for $\mathrm{m}{ }^{*} \mathrm{n}>\mathrm{m}+\mathrm{n}$ )


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## Eyeless Again - a Closer Look

Filter - Overview Results • Job information Customize order
Graphical overview


- The similar regions in the different homologues are not distributed randomly
- Actually, a single stretch of 128 AA, the PAX domain, is virtually unchanged in all homologues
- Controls binding to DNA and hence regulatory effects
- Typical: Only some parts of a sequence are conserved, and these carry function


## Example




## Coincidence?



No coincidence!


> IIIIIIIII


## Distance or Similarity

- Given two sequences A, B
- Until now, we computed a global distance
- The higher $e(A, B)$, the less similar are $A$ and $B$
- The longer $A$ and $B$, the higher their distance in general
- Different lengths are punished: $e(A, B) \geq||A|-|B||$
- Often, we want a local similarity instead
- Illustration: If we don't compare two defined genes (exons), but two strings presumably containing each one gene (exon)
- Local: We need to search for substrings $A^{\prime} \in A, B^{\prime} \in B$ which are very similar to each other
- $A^{\prime}$ and $B^{\prime}$ also should have a certain length to be interesting
- e( $\left.A^{\prime}, B^{\prime}\right)$ does not help - optimal distance is 0 for $A^{\prime}=B^{\prime}=" "$


## Aligned Sequences

- Assume we have an alignment $L$ of two sequences $A, B$
- Let $A^{L}\left(B^{L}\right)$ be the aligned version of $A(B)$
- $A^{L}$ and $B^{L}$ are strings over the alphabet $\Sigma^{*}$
- Example
- A=ATTAG, B=TTCAA
- L=

ATT_AG
_TTCAA

- Then $A^{L}=A T T \_A G, B L=\_T T C A A$
- Note that $\left|A^{L}\right|=\left|B^{L}\right|$


## Aligned Sequence Similarity

- A scoring function is a function s: $\Sigma^{*} x \Sigma^{*} \rightarrow$ Integer
- We also call s a substitution matrix
- (High) positive scores: "good" pairs; (low) negative sc.: "bad" pairs
- The similarity sim' of two aligned sequences $A^{L}, B^{\downarrow}$ wrt. $s$ with $\left|A^{L}\right|=\left|B^{\perp}\right|=n$ is defined as

$$
\operatorname{sim}^{\prime}\left(A^{L}, B^{L}\right)=\sum_{i=1}^{n} s\left(A^{L}[i], B^{L}[i]\right)
$$

## Example

$$
\Sigma^{\prime}=\left\{A, C, G, T,{ }^{\prime}\right\}
$$

|  | $\mathbf{A}$ | $\mathbf{C}$ | $\mathbf{G}$ | $\mathbf{T}$ | - |
| :---: | :---: | :---: | :---: | :---: | :---: |
| $\mathbf{A}$ | 4 | -2 | -2 | -1 | -3 |
| $\mathbf{C}$ |  | 4 | -1 | -2 | -3 |
| $\mathbf{G}$ |  |  | 4 | -2 | -3 |
| $\mathbf{T}$ |  |  |  | 4 | -3 |

$$
\begin{array}{ll}
\begin{array}{l}
\text { AC_GTC } \\
\text { AGGT_C }
\end{array} & =\mathbf{- 1} \\
\begin{array}{ll}
\text { ACGTC } \\
\text { AGGTC }
\end{array} & =\mathbf{1 5} \\
\text { A_CGTC } & =\mathbf{1 0}
\end{array}
$$

## Sequence Similarity

- The similarity sim of two sequences $A, B$ (wrt. s) is the highest similarity score sim' over all alignments of $A$ and $B$

$$
\operatorname{sim}(A, B)=\max _{L=\operatorname{align}(A, B)} \operatorname{sim}^{\prime}\left(A^{L}, B^{L}\right)
$$

- We are not yet there: This still is a global similarity score


## Computing $\operatorname{sim}(\mathrm{A}, \mathrm{B})$

- Same ideas as for edit distance
- But: We want a high similarity, not a low distance
- But: We have individual scores per pair, not only $1 / 0$
- We can compute $\operatorname{sim}(|\mathrm{A}|,|\mathrm{B}|)$ with

$$
\begin{array}{r}
\operatorname{sim}(i, 0)=\sum_{k=1 . . i} s\left(A[k],{ }_{2}\right) \quad \operatorname{sim}(0, j)=\sum_{k=1 . . j} s\left(\_, B[k]\right) \\
\operatorname{sim}(i, j)=\left\{\begin{array}{c}
\operatorname{sim}(i, j-1)+\begin{array}{l}
\left(\_, B[j]\right) \\
\operatorname{sim}(i-1, j)+\left(A[i], \_\right) \\
\operatorname{sim}(i-1, j-1)++(A[i], B[j])
\end{array}
\end{array}\right)
\end{array}
$$

## Example

|  | A | G | T | C |
| :---: | :---: | :---: | :---: | :---: |
| A | 4 | -1 | -1 | -1 |
| G |  | 4 | -1 | -1 |
| T |  |  | 4 | -1 |
| C |  |  |  | 4 |
| - | -3 | -3 | -3 | -3 |

Edit Distance

|  |  | A | G | G | T | C |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | 0 | 1 | 2 | 3 | 4 | 5 |
| A | 1 | 0 | 1 | 2 | 3 | 4 |
| G | 2 | 1 | 0 | 1 | 2 | 3 |
| T | 3 | 2 | 1 | 1 | 1 | 2 |
| C | 4 | 3 | 2 | 2 | 2 | 1 |
| C | 5 | 4 | 3 | 3 | 3 | 2 |


|  |  | A | G | G | T | C |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | 0 | -3 | -6 | -9 | -12 | -15 |
| A | -3 | 4 | 1 | -2 | -5 | -8 |
| G | -6 | 1 | 8 | 5 |  |  |
| T | -9 |  |  |  |  |  |
| C | -12 |  |  |  |  |  |
| C | -15 |  |  |  |  |  |

## Lokal Similarity = Local Alignment

- Definition
- The local similarity score sim* of A, B is defined as

$$
\operatorname{sim}^{*}(A, B)=\max _{\forall A^{\prime} \text { substringOf } A, B^{\prime} \text { substringOf } B}\left(\operatorname{sim}\left(A^{\prime}, B^{\prime}\right)\right)
$$

- Remark
- Inequality in length of $A$ and $B$ does not matter any more
- Sounds terribly complex, but there is a neat trick


```
    IIIIIIII
```



## Example

Match: +1
I/R/D: -1

|  |  | A | T | G | T | G | G |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
|  | 0 | -1 | -2 | -3 | -4 | -5 | -6 |
| G |  |  |  | -1 |  |  |  |
| T |  |  |  |  | 0 |  |  |
| G |  |  |  |  |  | 1 |  |
| A |  |  |  |  |  |  | 0 |



## Smith-Waterman Algorithm

- Smith, Waterman: „Identification of common molecular subsequences", J. Mol. Bio 147, 1981
- Idea
- Note: Local paths need not span the entire strings
- Look at a single path
- A series of matches (positive values for scoring function s) creates a series of increasing similarity values
- Any step with $\mathrm{s}<0$ lowers the score
- Whenever the cumulative score falls below 0 , we drop this prefix
- Instead of carrying on, we conceptually start a new local path
- To this end, we simply set true_score=max(0,score)
- The highest value in the matrix is the end of the best local path


## Computation

- The same ideas as before
- We compute sim* $(A, B)$ using a similar recurrence as for global alignments
- $\operatorname{sim} *[A, B]$ eventually is the maximal value in $S$

$$
S(i, 0)=\sum_{k=1 . . i} s\left(A[k],{ }_{c}\right) \quad S(0, j)=\sum_{k=1 . . j} s(, B[k])
$$

$$
S(i, j)=\max \left\{\begin{array}{c}
S(i, j-1)+s\left(\_, B[j]\right) \\
S(i-1, j+s \nmid[i],-) \\
S(i-1, j-\underbrace{+s(4[i], B[j])}
\end{array}\right.
$$

|  |  | A | T | G | T | C | G |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
|  | 0 | -1 | -2 | -3 | -4 | -5 | -6 |
| A | -1 | 1 | 0 | -1 | -2 | -3 | -4 |
| T | -2 | 0 | 2 | 1 | 0 | -1 | -2 |
| G | -3 | -1 | 1 | 3 | 2 | 1 | 0 |

ATGTCG
ATG $\qquad$
ATGTCG
AT $\qquad$ G

ATGTCG
A_T_G

|  |  | $A$ | T | G | T | C | G |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
|  | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| A | 0 | 1 | 0 | 0 | 0 | 0 | 0 |
| T | 0 | 0 | 2 | 1 | 1 | 0 | 0 |
| G | 0 | 0 | 1 | 3 | 2 | 1 | 1 |

ATGTCG
ATG $\qquad$

## Local versus global Alignment

- Global Alignment
- Comparison of two entire sequences
- Use when you think the entire sequences are related
- Interest: The differences; assumption: Relatedness
- Example: Proteins of the same family
- Local Alignment
- Compare uncharacterized sequences
- Use when comparing "randomly sampled" sequences
- Interest: Similar regions; assumptions: None
- Often a first step before global alignment
- Example: Find similar genes in other species genomes


## Beware: Not all Events are Equal

Wildtype

| CTT | A G T | G A | T A |  | A A A | DNA <br> Protein |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Leu | Ser | Asp | Tyr | Gly | Lys |  |
| C ${ }^{\text {I }}$ |  |  | - |  | A | DNA |
| Leu | Ser | Asp | Stop-Codon |  |  | Protein |
| CTTAGTGAACTACGGTAAA |  |  |  |  |  | DN |
| eu | Ser | His | Asp | Leu | Th | Protein |
| CTTTAGCGACTACGGTA A A |  |  |  |  |  | DNA |
| Leu | Ser | Asp | Tyr | Gly | Lys | Protein |
| CTTAGTGAATACGGTAAA |  |  |  |  |  | DNA |
| Leu | Ser | Glu | Tyr | Gly | Lys | Protein |

## Further Reading

- Everywhere
- Relaxed: Christianini \& Hahn, Chapter 3
- Step by step: Waack, Chapter 9

