# 6 Evaluation of alignment scores

## 6.1 Statistical analysis

Statistical tests are performed to give an explanation to observed alignment scores.

### Hypothesis testing

- Alternative hypothesis
- Null hypothesis

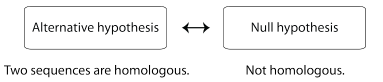


Figure 6.1: The null hypothesis and the alternative hypothesis

#### P-value

"The p-value is defined as the probability of obtaining a result equal to or more extreme than what was actually observed, assuming that the null hypothesis is true"

- the p-value page on Wikipedia (https://en.wikipedia.org/wiki/P-value)

#### Significance level $(\alpha)$

The significance level should be chosen to indicate strong/weak evidence against the null hypothesis.

Significance levels 0.05 and 0.01 are often used in life sciences.

- Statistically significant:  $\alpha = 0.05$
- Statistically highly significant:  $\alpha = 0.01$

#### Common misunderstandings of p-value

"The p-value is not the probability that the null hypothesis is true or the probability that the alternative hypothesis is false."

- the p-value page on Wikipedia (https://en.wikipedia.org/wiki/P-value)

## Underlying (background) score distributions

Table 6.1: Alignment methods and distributions

Method	Underlying distribution
Global alignment	Unknown
Local alignment (ungapped)	Gumbel

# 6.2 Evaluation of global alignment

The underlying distribution of global alignment scores is unknown.

### Random generation of sequences

One needs to consider using the appropriate length and compositions of amino acids or nucleotides needs when creating randomised sequences.

### Example

Input sequences

q: ACGT d: AGTACC

Frequencies:  $f_A = 0.2, f_C = 0.4, f_G = 0.1, f_T = 0.3$ 

Length: 6

d1: CCAGTC
d2: TCACCG
d3: CTTGAA

. . .

### Frequency distributions

- Universal (e.g. the whole protein database)
- Global (e.g. protein super families)
- Local (e.g. query and database sequences)

#### Additional constrains

Constrains on sequences generation are often considered.

- Di-amino acid frequencies
- Sub-region specific frequencies

### Non-parametric test and p-value

The simplest non-parametric test is calculating the rank of the score for the original alignment as the p-value.

$$p = (b+1)/(n+1)$$

where b is the number of randomly generated scores above the score of the original alignment, and n is the sample size.

**N.B.** n should be sufficiently large (e.g. >1000) to estimate an accurate p-value.

### Example

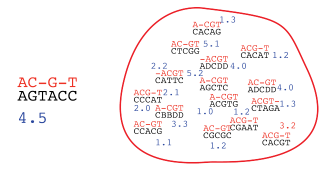


Figure 6.2: Randomly generated sequences and alignment scores

1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20
1	1.1	1.3	1.4	1.7	2.1	2.2	2.2	2.3	2.5	2.8	3	3.2	3.3	3.4	3.6	4.2	4.4	4.7	5.2
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p-value: (2+1)/(20+1) = 0.1429

- Significance level  $\alpha = 0.2$ : reject the null hypothesis
- Significance level  $\alpha = 0.05$ : the null hypothesis is not rejected

#### Exercise 6.1

1. Calculate the frequencies of nucleotides from the four sequences below.

d1: CCAGCd2: TCACGd3: CTTAAd4: AACAA

Frequencies:  $\{f_A = , f_C = , f_G = , f_T = \}$ 

2. Calculate the p-value of the alignment below.

q: AACG d: A-CG Score: 40

Assume that the scores are pre-calculated for the alignments of the query sequence and nine randomly generated sequences as follows. Use them for the p-value calculation.

No.	1	2	3	4	5	6	7	8	9
Score	4	14	33	45	74	76	82	83	94

### Using the normal distribution

The underlying distribution of global alignment scores is unknown, but the z-score is sometimes calculated.

The z-score is:

$$z = \frac{x - \mu}{\sigma}$$

where:

 $\mu$  is the mean of the population.

 $\sigma$  is the standard deviation of the population.

#### Mean and variance

The sample mean  $(\bar{x})$  and the sample variance  $(s^2)$  are calculated as follows.

$$\bar{x} = \frac{1}{n} \sum_{i=1}^{n} x_i$$

$$s = \sqrt{\frac{\sum_{i=1}^{n} (x_i - \bar{x})^2}{n-1}}$$

### Example of z-score

- $\bar{x}$ : 2.78
- s: 1.4964

$$z = \frac{4.5 - 2.78}{1.4964} = 1.1494$$

The p-value is 0.125196.

# 6.3 Evaluation of local alignment

The underlying distribution of local alignment scores is an extreme value distribution.

#### Gumbel distribution

The Gumbel distribution is a member of the extreme value distribution family.

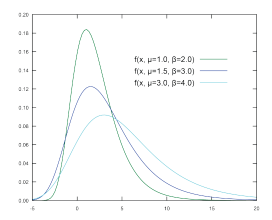


Figure 6.3: Gumbel distribution (source: Herr blaschke, Wikimedia Commons)

The cumulative distribution function (CDF) of the Gumbel distribution:

$$F_Y(y) = \exp[-e^{-\lambda(y-\mu)}]$$

Parameters

- $\mu$ : the modal value of the distribution, characteristic value
- $\lambda$ : a measure of the variance, decay constant

#### Extreme value distribution

An extreme value distribution is a limiting distribution for the minimum or the maximum of a sufficiently large sample. Ungapped alignments with large sequence lengths are known to have this type of distribution.

Example (m and n are not large in this example)

		A	$\mathbf{C}$	G	$\mathbf{C}$	Α	$\mathbf{C}$	G
	0	0	0	0	0	0	0	0
С	0	0	0.5	0	0.5	0	0.5	0
G	0	0	0	1	0.5	0.2	0	1
A	0	0.5	0	0.5	0.7	0.2	0	0.5

#### Parameter estimation

The p-value of the Gumbel distribution can be calculated as:

$$P[Y > y] = 1 - F_Y(y) = 1 - \exp[-e^{-\lambda(y-\mu)}]$$

The parameters  $\mu$  and  $\lambda$  can be estimated from the arithmetic mean  $m_Y$  and the variance  $\sigma_Y^2$  of the observed sample.

$$\lambda \approx 1.282/\sigma_Y$$

$$\mu \approx m_Y - 0.577/\lambda$$

### Example of parameter estimation

Below is the optimal local alignment with the score between q:ACAGACTACTA and d:TCAGACTGGGAACCE.

CAGACT CAGACT Score: 6

The mean and the variance of the alignment scores are estimated as follows from randomly generated sequences.

 $m_Y$ : 1.7221  $\sigma_Y$ : 1.6025

Then,  $\lambda$  and  $\mu$  are estimated from  $m_Y$  and  $\sigma_Y$ .

 $\lambda \approx 1.282/1.6025 = 0.8$   $\mu \approx 1.7221 - 0.577/0.8 \approx 1$ 

The p-value is approximately 0.0181 when  $\lambda = 0.8$  and  $\mu = 1$ . The test result is statistically significant ( $\alpha = 0.05$ ), and therefore, the null hypothesis is rejected.

Conclusion: The query and the database sequences are homologous (p-value: 0.0181).

#### 6.4 Evaluation of database search

BLAST reports bit scores and e-values as search result. Bit score are calculated from raw scores, and e-values represent the expected numbers of database hits.

### Example of BLAST output

- q: HSBGPG Human gene for bone gla protein (BGP)
- d: osteocalcin [Felis catus]
- Sequence ID: XP\_003999760.1

	Score		Expect	Identities	Positives	$\mathbf{Gaps}$
_	38.5 bits	(88)	3.5	19/25 (76%)	20/25 (80%)	0/25 (0%)
Que	ry	677	TAFVS	SKQEGSEVVKRPF	RRYLYQWLG	751
			AFVS	SKQEGSEVV+R F	RRYL LG	
Sbj	ct	36	AAFVS	SKQEGSEVVRRLF	RRYLAPGLG	60

#### Karlin-Altschul statistics

- $\lambda$  is a scalar parameter for score matrix
- K is a scalar parameter for search space size

BLAST pre-calculates both parameters in a search space independent manner.

### Example of Karlin-Altschul statistics

• Matrix: BLOSUM62

• Lambda: 0.267

• K: 0.041

### Sequence databases

The NCBI site provides several databases for BLAST search.

- Nucleotide collection (nr/nt)
- Non-redundant protein sequences (nr)

#### Example of database statistics

• Database: nr

• Number of letters: 41,667,927,126

• Number of sequences: 113,671,629

#### 6.5 Bit score and e-value

BLAST reports bit-scores and e-values that can be used for evaluation on search results.

#### Bit score

Bit scores are normalized scores that have the same unit (bit). The scores can be comparable even when different scoring schemes are used.

$$S' = \frac{(\lambda S - \ln K)}{\ln 2}$$

 $2^{S'}$  indicates the expected search space size that one would find one alignment with score at least S by chance alone.

# Example of bit score calculation

• Lambda ( $\lambda$ ): 0.267

• K: 0.041

• Score: 88

$$S' = \frac{(\lambda S - \ln K)}{\ln 2} = \frac{(0.267 \times 88 - \ln 0.041)}{\ln 2} = 38.506$$

$$2^{S'} = 2^{38.506} = 390, 300, 663, 957$$

### E-value

"The Expect value (E) is a parameter that describes the number of hits one can expect to see by chance when searching a database of a particular size"

- BLAST Frequently Asked questions (http://blast.ncbi.nlm.nih.gov)

$$E(S) = Kmne^{-S} = \frac{mn}{2^{S'}}$$

## Example of E-value calculation

• n: 25

• m: 41,667,927,126

• Lambda ( $\lambda$ ): 0.267

• K: 0.041

• Score: 88

$$E(88) = \frac{41,667,927,126 \times 25}{2^{38.506}} = 2.669$$

#### Exercise 6.2

• λ: 1.28

• K: 0.5

• m: 1000

• n: 100

Calculate  $\exp(-1.28)$  as 0.28.

1. What is the e-value of the score 1?

2. Is the alignment with score 1 likely homologous?