

# Computational biology

## Co-evolution to predict protein structures

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Statistiques pour les sciences du Vivant et de l'Homme

December 11, 2019

- Get an overview of computational biology topics
  - Topics (genomics, metagenomics, proteomics, etc.)
  - Know some important databases
  - Know standard tools (Blast, PyMol) and libraries (BioPython)
- Have a basic culture of order of magnitude in computational biology
  - Quantity of data
  - Size of genomes
  - Size of organisms
- Toward autonomy for design and implementation of methods
  - Case study of SNP detection
  - Case study of protein structure prediction

# Today's outline: from gene sequence to protein structure

- Sequence-structure-function paradigm
  - Genomes, genes, proteins
  - Databases
- Evolution
  - Selective pressure
  - Multiple sequence alignment
  - Co-evolution

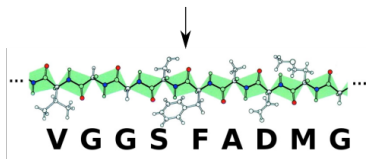
# From genome to function, the very big picture

ACGATGTATTCAGCGATTACGATAAAGCTACGTAGTGGCA

On a genome ( $\sim 5$ Mbp), specific motifs define beginning and end of a gene

# From genome to function, the very big picture

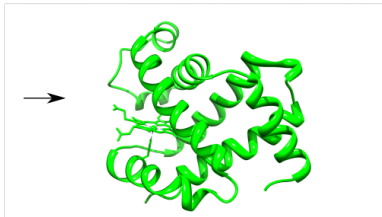
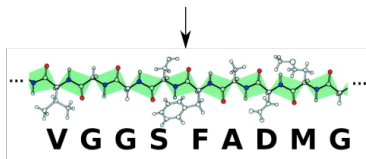
ACG**ATGTATTCAGCGATTACGATAAAGCTACGTAGT**GGCA



*Transcription + translation*, to form a chain of amino acids ( $\sim 300-3000\text{AA}$ )

# From genome to function, the very big picture

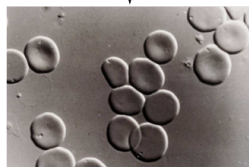
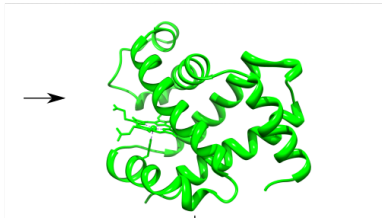
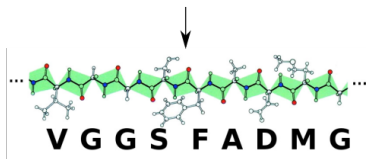
ACG**ATGTATTCAGCGATTACGATAAAGCTACGTAGT**GGCA



*Protein folding* under physico-chemical interactions, diameter  $\sim$  few nanometers

# From genome to function, the very big picture

ACG**ATGTATTCAGCGATTACGATAAAGCTACGTAGT**GGCA



O<sub>2</sub> transport

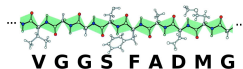
Protein endowed with a function (biochemical reactions, transport, etc.)

# Data at every steps

Nucleic seq.

..ATTGTCGATGAC..

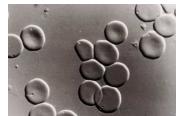
Amino acid seq.



Protein



Function





# Data at every steps

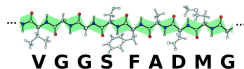
Nucleic seq.

..ATTGTCGATGAC..



[ncbi.nlm.nih.gov](http://ncbi.nlm.nih.gov)

Amino acid seq.



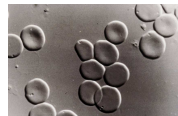
[uniprot.org](http://uniprot.org)

Protein



[rcsb.org](http://rcsb.org)

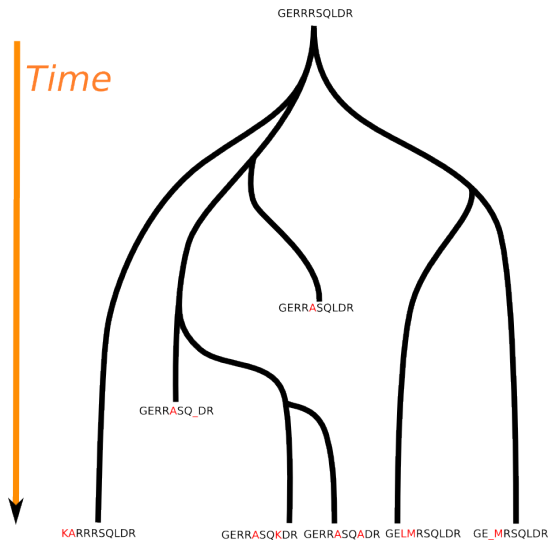
Function



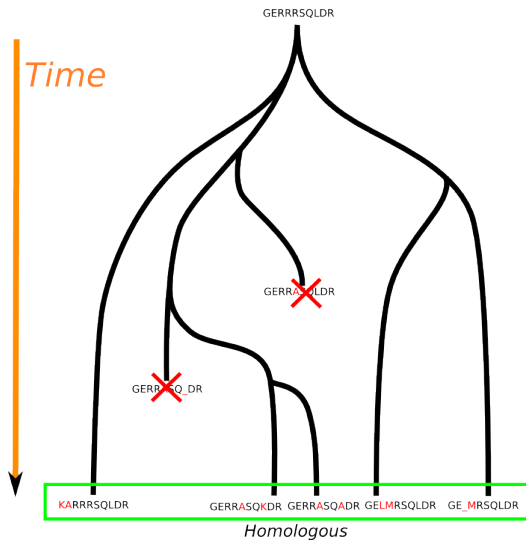
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# Protein evolution through mutations



# Protein evolution through mutations



# Sequence conservation

Aligning the sequences (MSA, multiple sequence alignment):

```
RYDSR TTIFSP..EGRL YQVEYAMEAIGNA.GSAIGILS
RYDSR TTIFSPLR EGRL YQVEYAMEAISHA.GTCLGILS
RYDSR TTIFSP..EGRL YQVEYAQEAISNA.GTAIGILS
RYDSR TTIFSP..EGRL YQVEYAMEAISHA.GTCLGILA
RYDSR TTIFSP..EGRL YQVEYAMEAIGHA.GTCLGILA
RYDSR TTIFSP..EGRL YQVEYAMEAIGNA.GSALGVLA
RYDSR TTTFSP..EGRL YQVEYALEAINNA.SITIGLIT
SYDSR TTIFSP..EGRL YQVEYALEAINHA.GVALGIVA
```

Tools	Database
ClustalW [Larkin et al. 07]	Pfam <a href="http://pfam.xfam.org">pfam.xfam.org</a>

# Sequence conservation

Aligning the sequences (MSA, multiple sequence alignment):

```
RYDSR TTIFSP . . EGRL YQVEYAMEAIGNA . GSAIGILS
RYDSR TTIFSP LR EGRL YQVEYAMEAISHA . GTCLGILS
RYDSR TTIFSP . . EGRL YQVEYAQEAISNA . GTAIGILS
RYDSR TTIFSP . . EGRL YQVEYAMEAISHA . GTCLGILA
RYDSR TTIFSP . . EGRL YQVEYAMEAIGHA . GTCLGILA
RYDSR TTIFSP . . EGRL YQVEYAMEAIGNA . GSALGVLA
RYDSR TTTFSP . . EGRL YQVEYALEAINNA . SITIGLIT
SYDSR TTIFSP . . EGRL YQVEYALEAINHA . GVALGI VA
```

Tools	Database
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Why some positions are conserved, some other aren't?

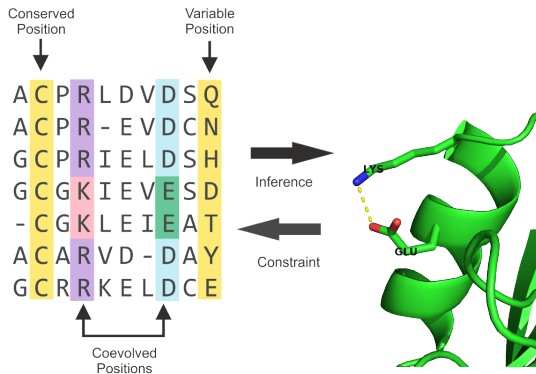
# Structure is determined by amino acid interactions



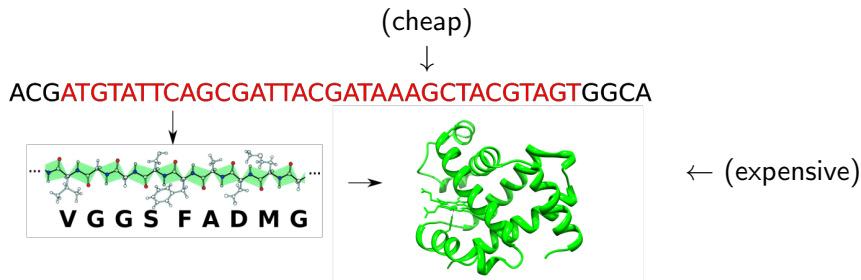
# Preserving the function: coevolution of residues

As protein function is vital, **evolution selects mutations preserving structures.**

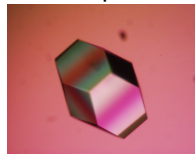
Leading to **compensatory** mutations:



# Computers and protein structure prediction

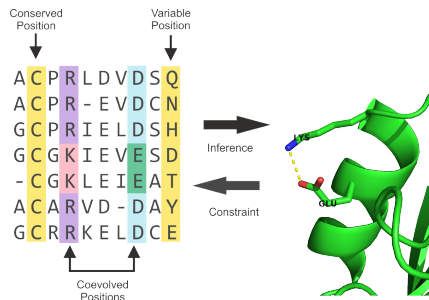


Structure determined by X-Rays  
through a crystal of proteins





# A simple approach for protein structure prediction



- Build or get multiple manio acid sequence alignments (**e.g.** in Pfam database)
- Quantify coevolution between positions in the sequence
- Infer what are the position in contact

What measure for co-evolution? Correlation would work?

## Conservation vs. co-evolution

Conserved position carries no information in terms of co-evolution (entropy is zero).

# Conservation vs. co-evolution

Conserved position carries no information in terms of co-evolution (entropy is zero). A standard approach is to measure it through Mutual Information:

$$MI(i, j) = \sum_{a,b} p(x_i = a, x_j = b) \log \frac{p(x_i=a, x_j=b)}{p(x_i=a) p(x_j=b)}$$

Where

- $x_i$  is the amino acid at position  $i$
- $p(x_i = a)$  is estimated in the MSA by  $\frac{\text{\#sequences having "a" at position } i}{N}$
- $N$  the number of sequences in the MSA
- $p(x_i = a, x_j = b)$  is estimated in the MSA by  $\frac{\text{\#sequence having "a" at } i \text{ and "b" at } j}{N}$

In practice you need  $N > 1,000$  to have reasonable estimation of  $p(x_i = a, x_j = b)$ .

# Over-repdiction at entropic position

When applying the rule

$$MI(i, j) > \tau \Rightarrow \text{contact between } i \text{ and } j$$

some positions predict too many contacts, often position with high entropy. Several corrections can be applied<sup>1</sup>.

## In your project

You can try using the simple correction:

$$MI'(i, j) = MI(i, j) - \frac{1}{N} \sum_k (MI(k, j) + MI(i, k))$$

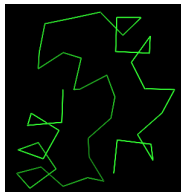
and fix a  $\tau$  to predict a contact as soon as:

$$MI'(i, j) > \tau$$

<sup>1</sup>See <https://doi.org/10.1093/bioinformatics/bti671>

## Hints concerning your Salmonella project

- The mutated gene you identified in the resistant bacteria have a large multiple sequence alignment in Pfam, search for its name in the Pfam browser.
- WP1.T2 consists in creating a tool taking as input an MSA in Fasta format and outputing a **contact matrix**<sup>2</sup>.
- You can model the structure using FT-comar<sup>3</sup> software, and compare to the native structure <sup>4</sup> using RasMol or PyMol software:



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<sup>2</sup>see the *cheatsheet* for details about the file format

<sup>3</sup>[clovisg.github.io/teaching/protein-structure-prediction/ft-comar.tgz](https://clovisg.github.io/teaching/protein-structure-prediction/ft-comar.tgz)

<sup>4</sup>[clovisg.github.io/teaching/protein-structure-prediction/target.pdb](https://clovisg.github.io/teaching/protein-structure-prediction/target.pdb)

# Summary

Check what you've learn:

- What is a genome, a gene, a protein, its structure
- How real sequencing data look like
- What is a SNP, what can be the impact
- Main tools and databases in computational biology
- Potential application of computational biology for public health studies

The project involved basic skills from different area:

- biology
- statistics (Poisson distribution)
- algorithmics (linear time algorithms required)

# Projects

Remember that your project should be like professional answers to the call:

- Clarity
- Fulfillment of the call
- Trustworthiness in the description of the approach

You should send:

- a report, including:
  - description of the strategy
  - approximations and choices
  - application to the project data (what gene is impacted by the SNP)
- your code
- a step-by-step guide to reproduce the results of the report

The TATFAR  
waits for  
interesting answers to its call!