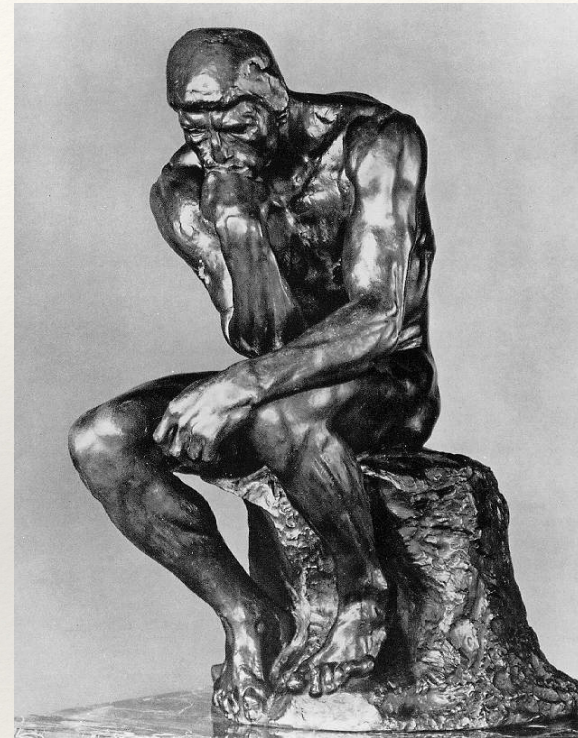


# Computational Structural Bioinformatics

Patrice Koehl

# Science, **then**, and now...

- For a long time, people thought that it would be enough to reason about the existing knowledge to explore everything there is to know.
- One single person could possess all knowledge in her cultural context.  
(encyclopedia of Diderot and D'Alembert)
- Reasoning, and mostly passive observation were the main techniques in scientific research





Science, **then**, and now...

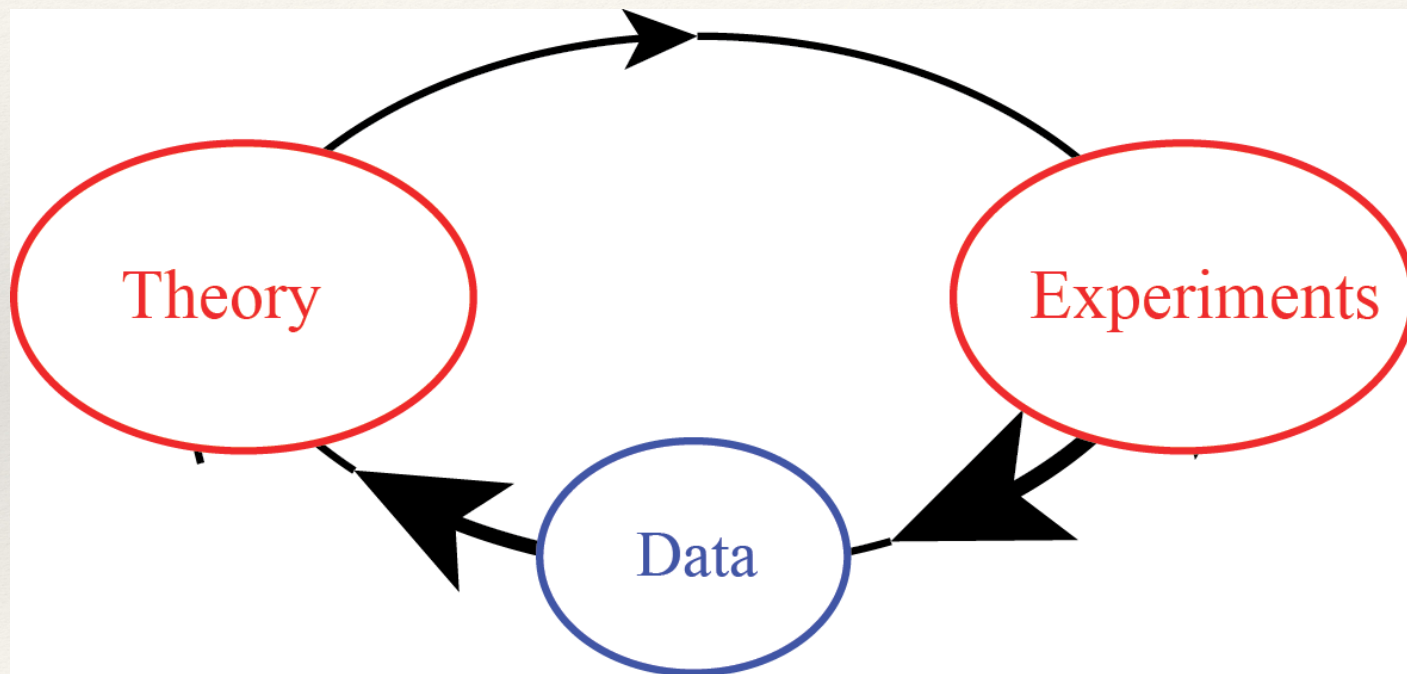
*“All science is either physics, or stamp collecting”*

Rutherford, chemist and physicist, 1876-1937

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# Science, then, and now

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# Science, then, and now

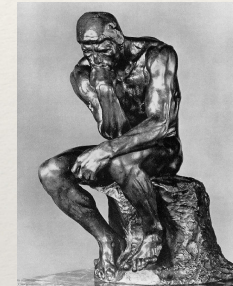
## 1. Thousand years ago – **Experimental Sciences**

Description of natural phenomena



## 2. Last few hundred years – **Theoretical Sciences**

Newton's law, Maxwell's equations...



## 3. Last few decades – **Computational Sciences**

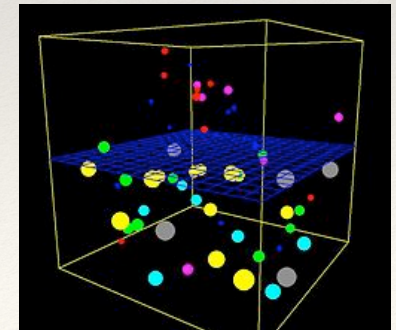
Simulation of complex phenomena

## 4. Today – **Data-Intensive Sciences**

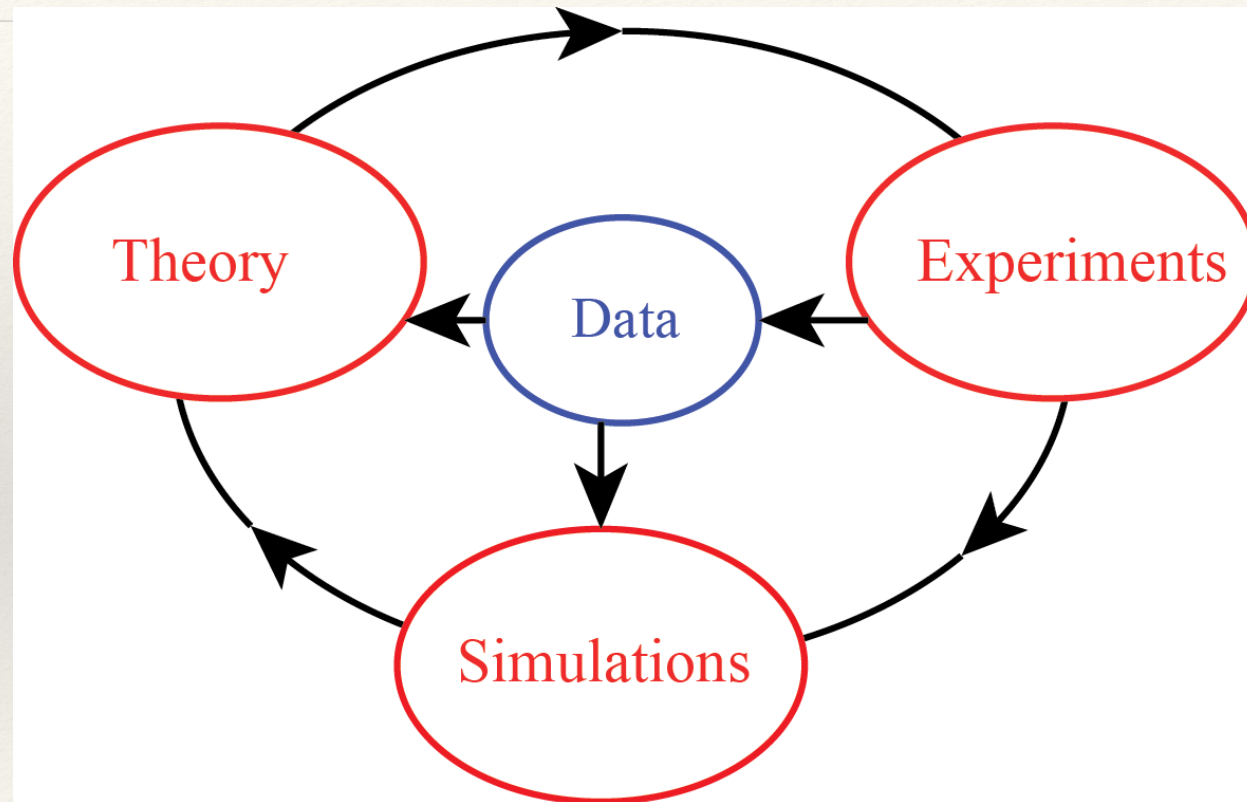
Scientist overwhelmed with data sets from many different sources

*Data captured by instruments*

*Data generated by simulations*

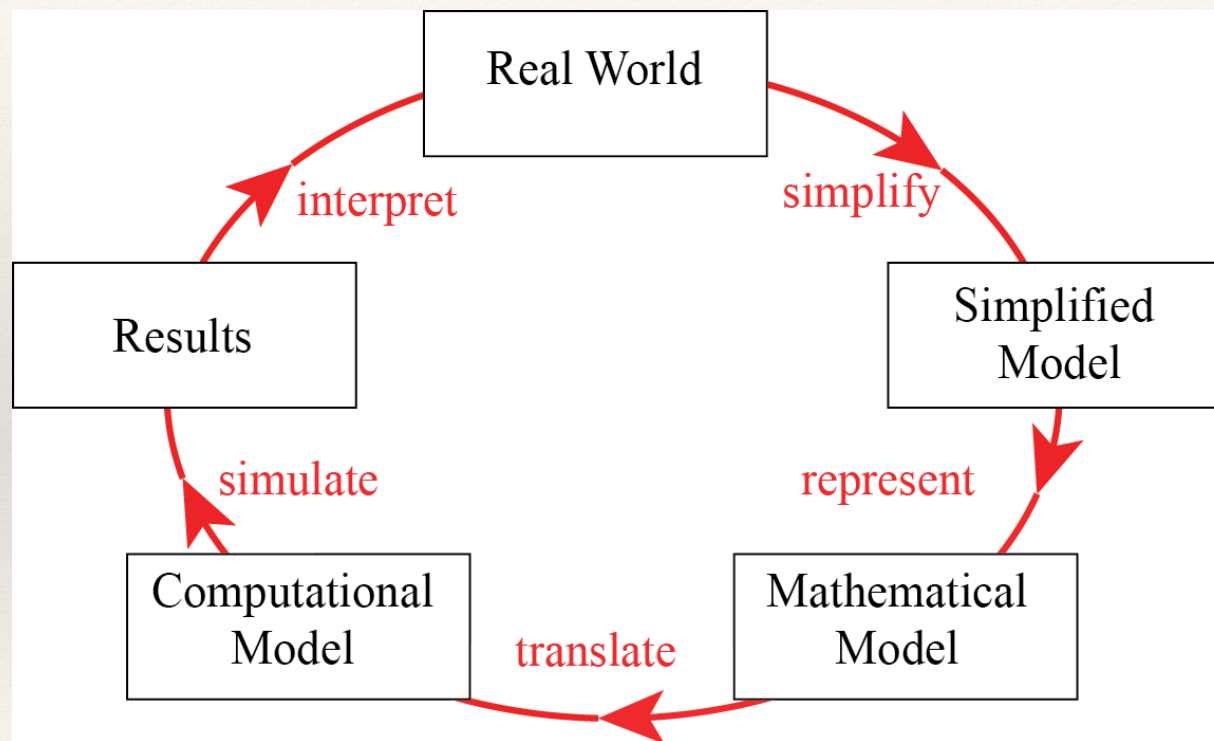


# Science, then, and now





# Mathematical Modeling



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## Context: Biology

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- ❖ “Life sciences” have their origins in ancient Greece

*Aristotle wrote influential treatises on zoology, anatomy and botany, that remained influential till the Renaissance*

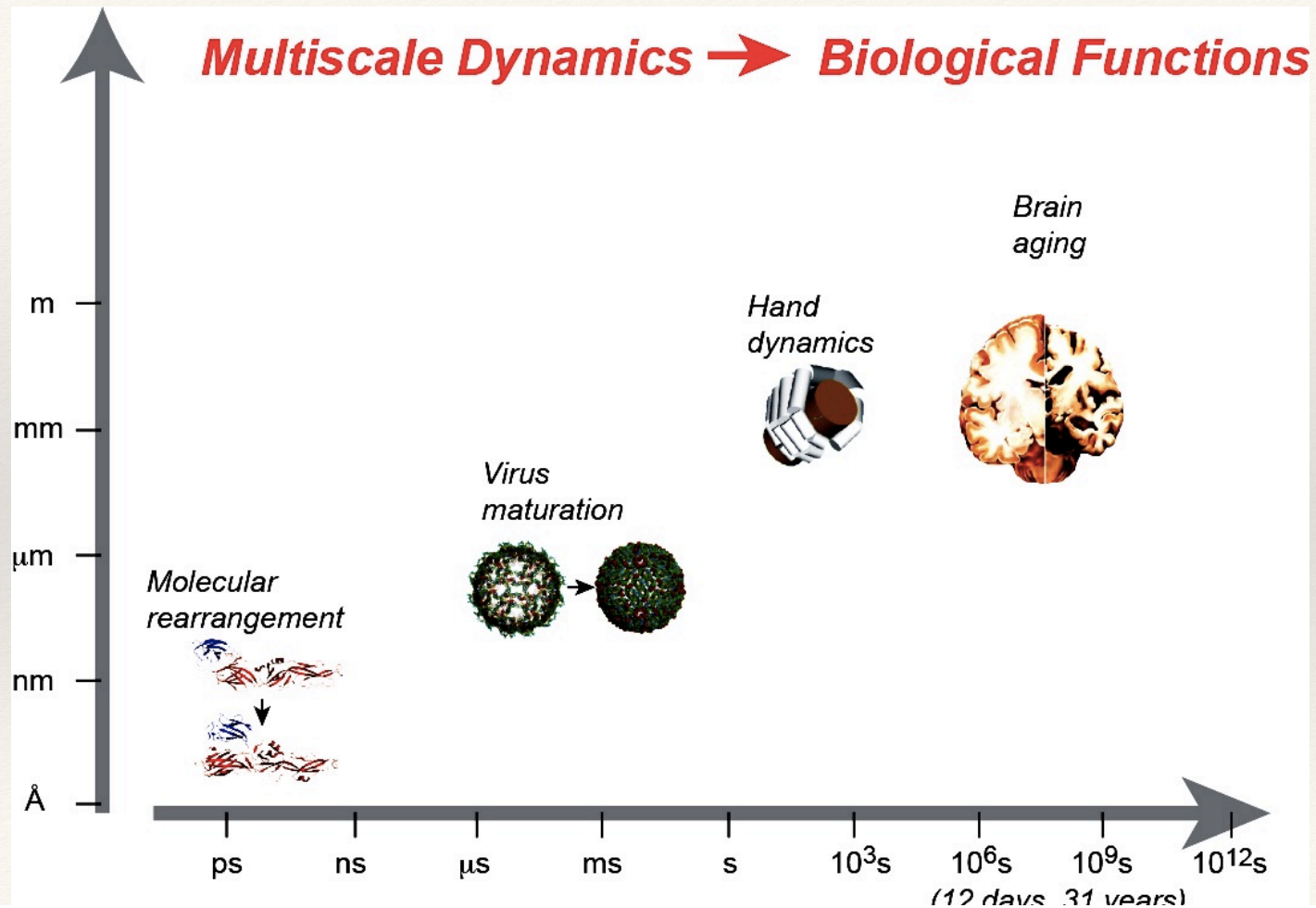
- ❖ “Life sciences” have always relied both on observation and discovery

*taxonomy, classifications, theory of evolution, ...*

- ❖ Biology is changing with the arrival of massive amount of data from the different genomics experiments



## Shapes and Dynamics of Biological Systems: Connecting the dots...



# What is ‘bioinformatics’?

- The term was originally proposed in 1988 by Dr. Hwa Lim
- The original definition was :

*“a collective term for data compilation, organisation, analysis and dissemination”*

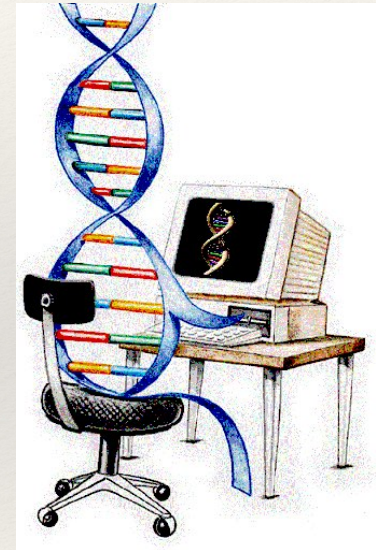


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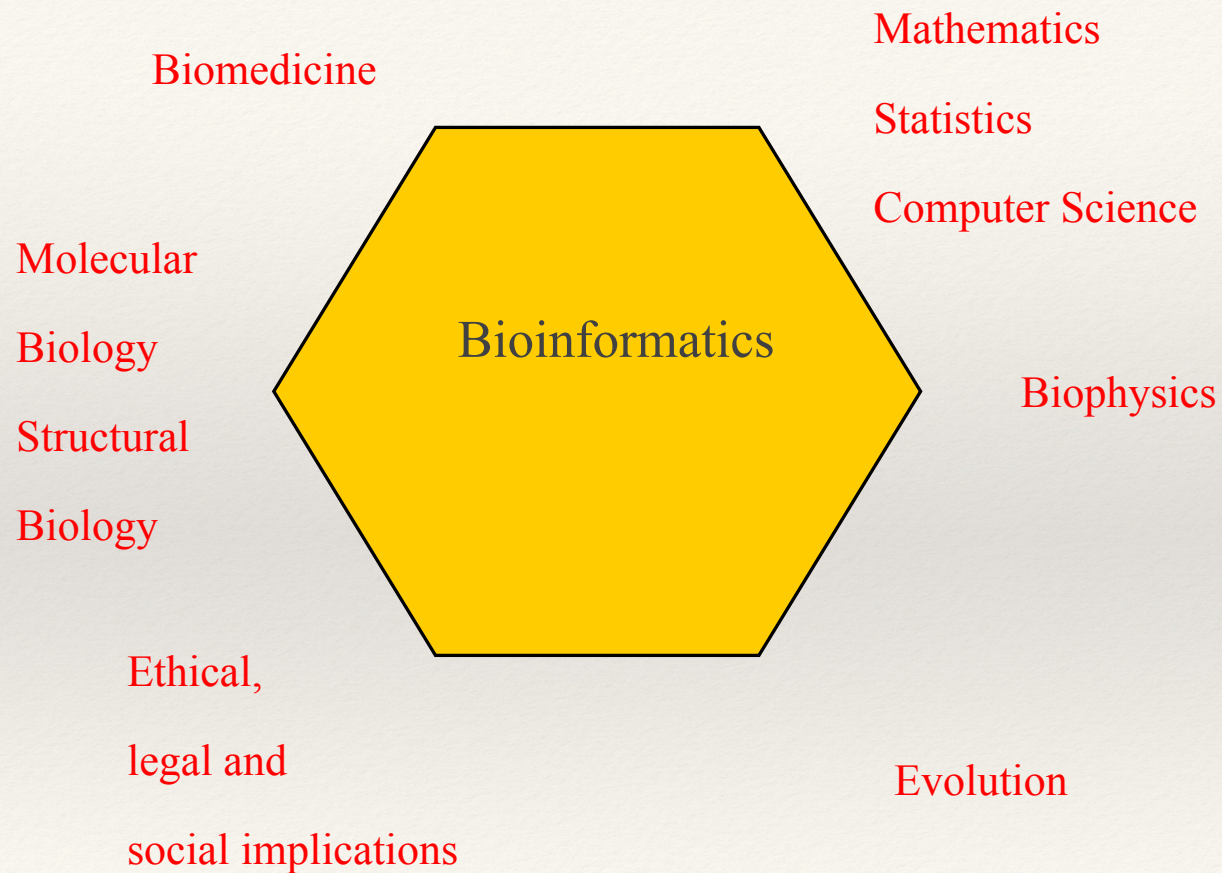
## That means....

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- Using information technology to help solve biological problems by designing novel algorithms and methods of analyses (*computational biology*)
- It also serves to establish innovative software and create new or maintain existing databases of information, allowing open access to the records held within them (*bioinformatics*)



# Bioinformatics is interdisciplinary





# What data?

*Biologists have been  
classifying data on  
plants and animals  
since the Greeks*

**HOLT BIOSOURCES**  
TEACHING TRANSPARENCIES

**Classification of Modern Humans** **57**

Homo sapiens   Homo erectus   Australopithecus   Gorilla   Elephant   Snake   Earthworm  
Fish   Sea star   Snail

**Kingdom**  
Animalia

Includes chordates, sea stars, earthworms, snails, jellyfish, sponges, clams, and insects

**Phylum**  
Chordata

Includes mammals, fishes, reptiles, birds, and amphibians

**Class**  
Mammalia

Includes primates and elephants, along with cats, dogs, horses, kangaroos, whales, bats, seals, dolphins, and many others

**Order**  
Primates

Includes members of the family Hominidae, along with prosimians, monkeys, and apes such as the gorilla

**Family**  
Hominidae

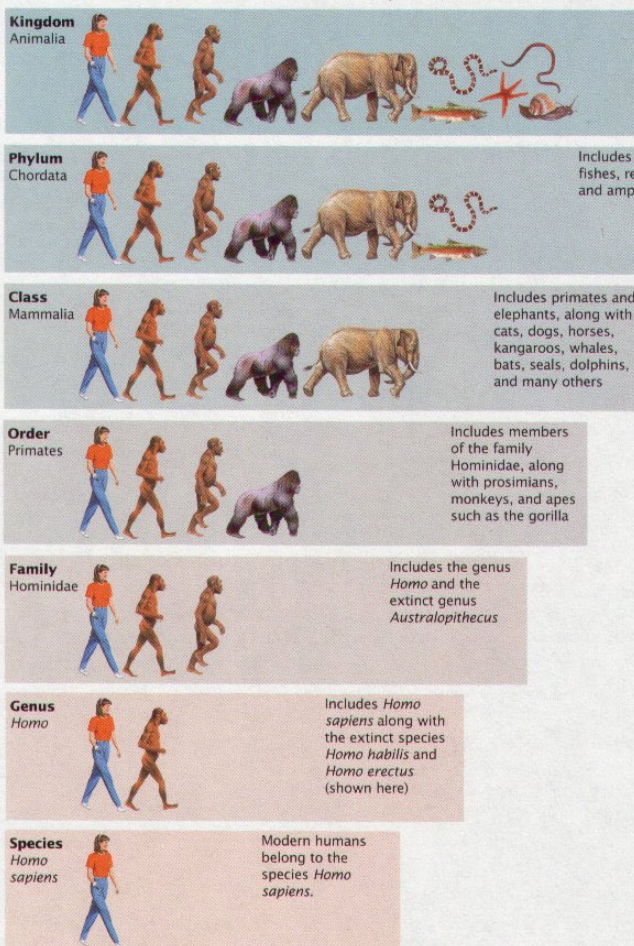
Includes the genus *Homo* and the extinct genus *Australopithecus*

**Genus**  
*Homo*

Includes *Homo sapiens* along with the extinct species *Homo habilis* and *Homo erectus* (shown here)

**Species**  
*Homo sapiens*

Modern humans belong to the species *Homo sapiens*.





# The Tree of Life

*"The affinities of all beings of the same class have sometimes been represented by a great tree... As buds give rise by growth to fresh buds, and these if vigorous, branch out and overtop on all sides many a feebler branch, so by generation I believe it has been with the great Tree of Life, which fills with its dead and broken branches the crust of the earth, and covers the surface with its ever branching and beautiful ramifications."*

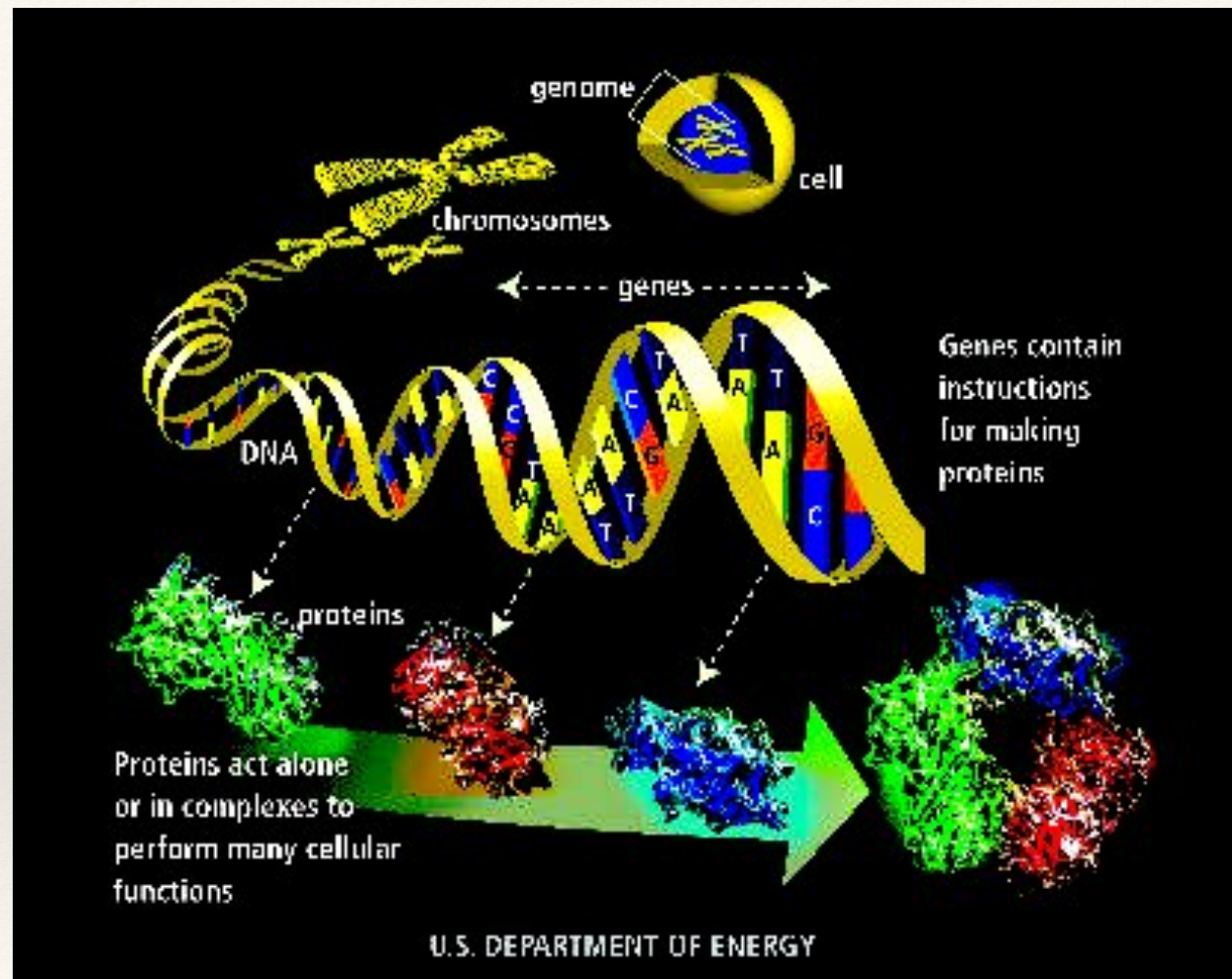


*Charles Darwin, 1859*



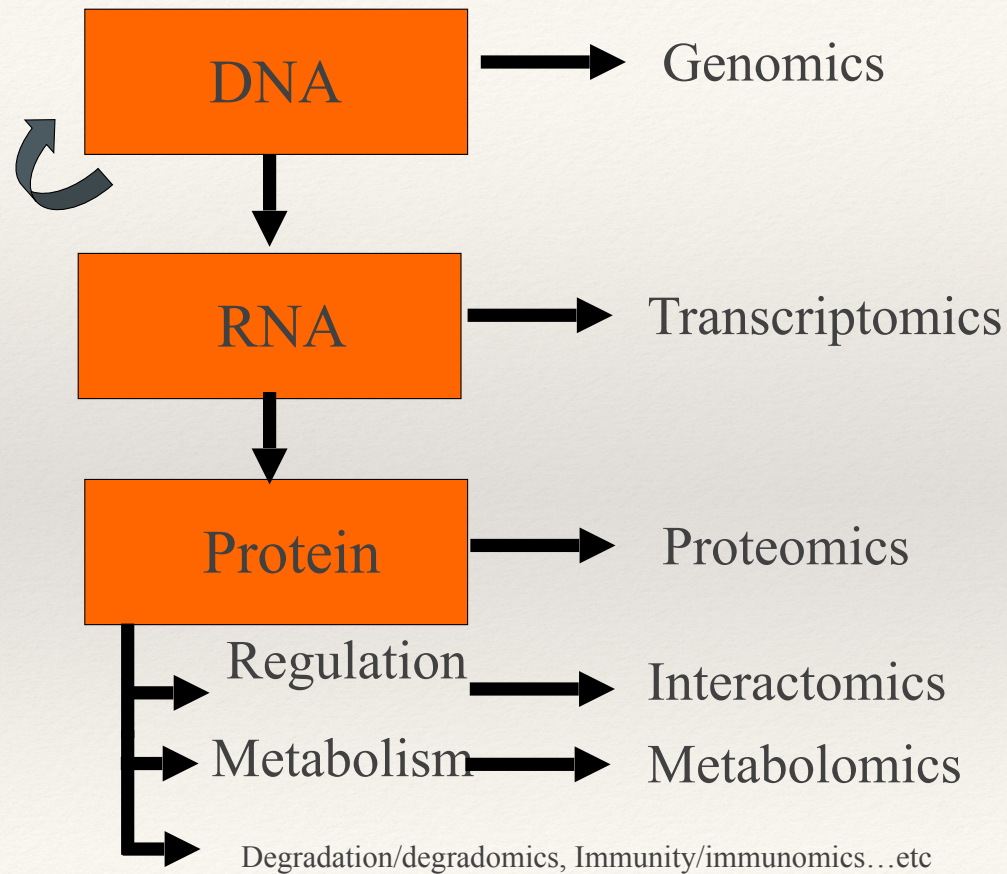
<http://tolweb.org>





# Central Dogma and the “omics”

## Integrative Systems Biology





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## Genes (1)

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- ❖ Genes are the basic units of heredity
- ❖ A gene is a sequence of bases that carries the information required for constructing a particular protein (gene “encode” the protein)
- ❖ The human genome comprises ~ 20,000 genes

Organism	Estimated size	Estimated gene #	Number of chromosome
<i>Homo sapiens</i> (human)	2900 million bases	~20,000	46
<i>Rattus norvegicus</i> (rat)	2,750 million bases	~30,000	42
<i>Mus musculus</i> (mouse)	2500 million bases	~30,000	40
<i>Oryza sativa</i> L. (rice)	450 million bases	~40,000	12
<i>Drosophila melanogaster</i> (fruit fly)	180 million bases	13,600	8
<i>Arabidopsis thaliana</i> (plant)	125 million bases	25,500	5
<i>Caenorhabditis Elegans</i> (roundworm)	97 million bases	19,100	6
<i>Saccharomyces cerevisiae</i> (yeast)	12 million bases	6300	16
<i>Escherichia coli</i> (bacteria)	4.7 million bases	3200	1
<i>H. Influenzae</i> (bacteria)	1.8 million bases	1700	1



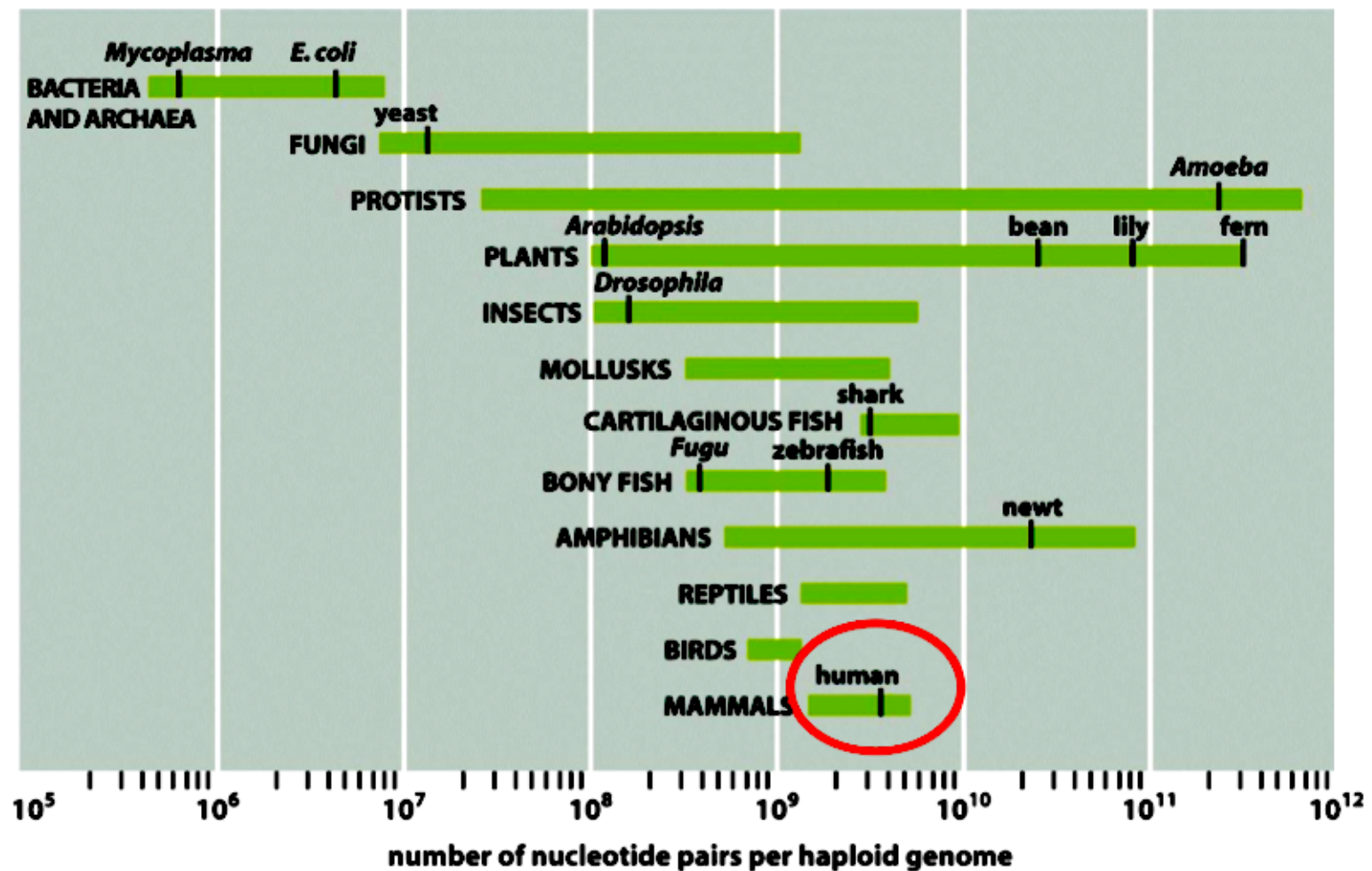
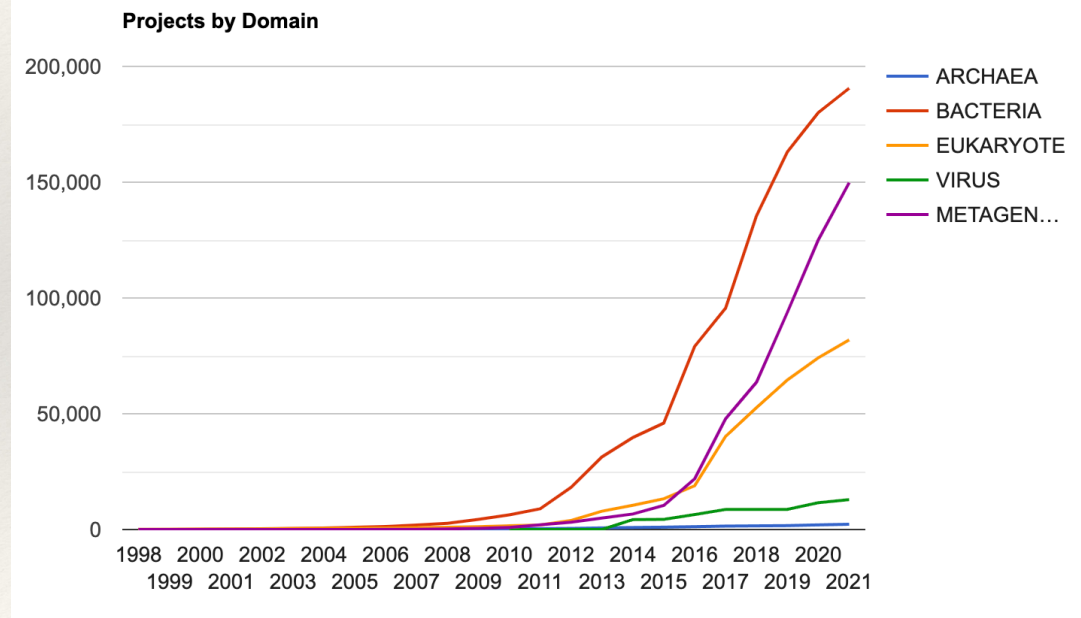


Figure 1-37 Molecular Biology of the Cell 5/e (© Garland Science 2008)

# The genomics projects

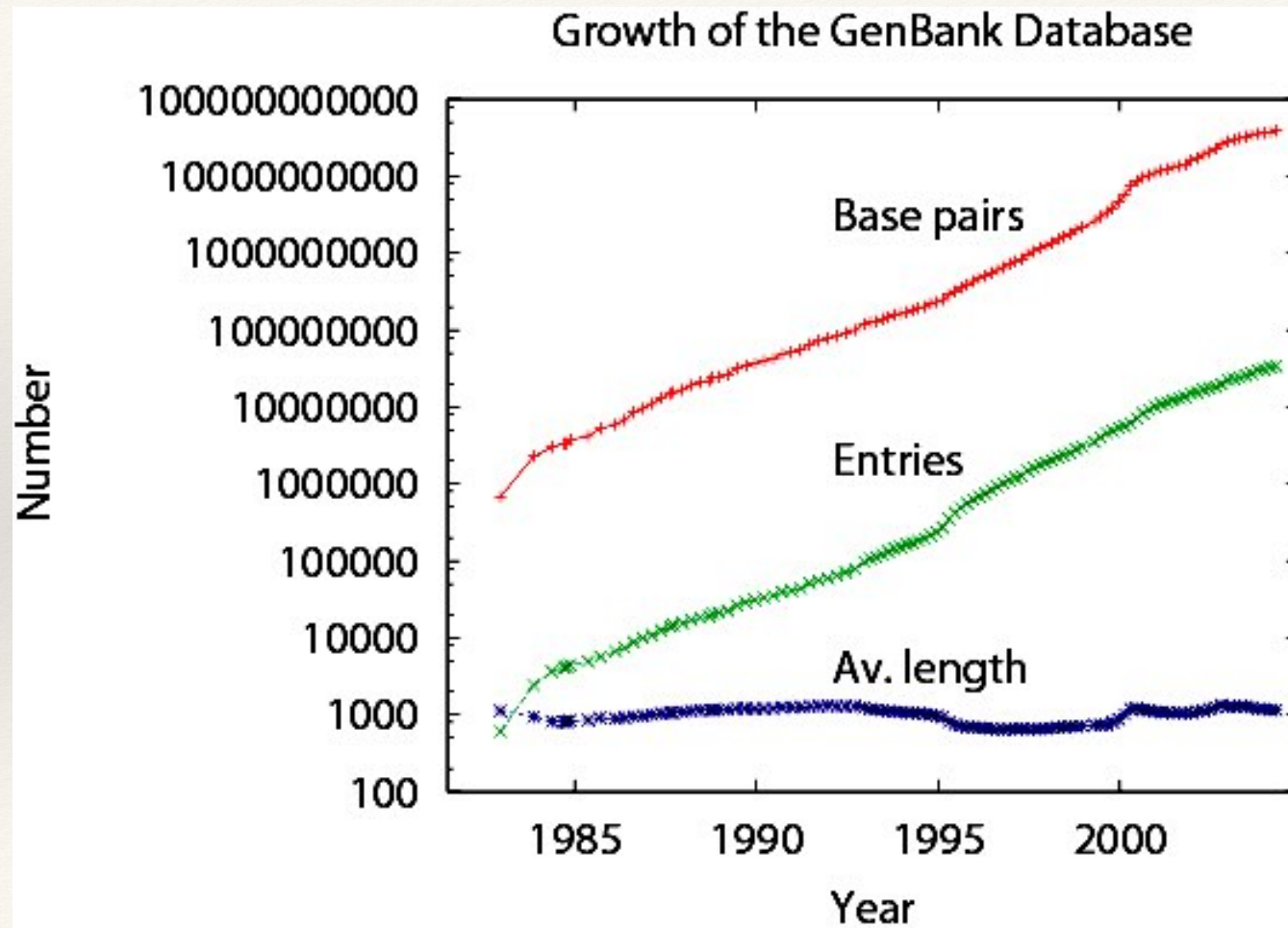
Studies ⓘ	<u>49,867</u>
Biosamples ⓘ	<u>147,318</u>
Sequencing Projects ⓘ	<u>436,918</u>
Analysis Projects ⓘ	<u>328,692</u>
Organisms	<u>418,931</u>



(GOLD)

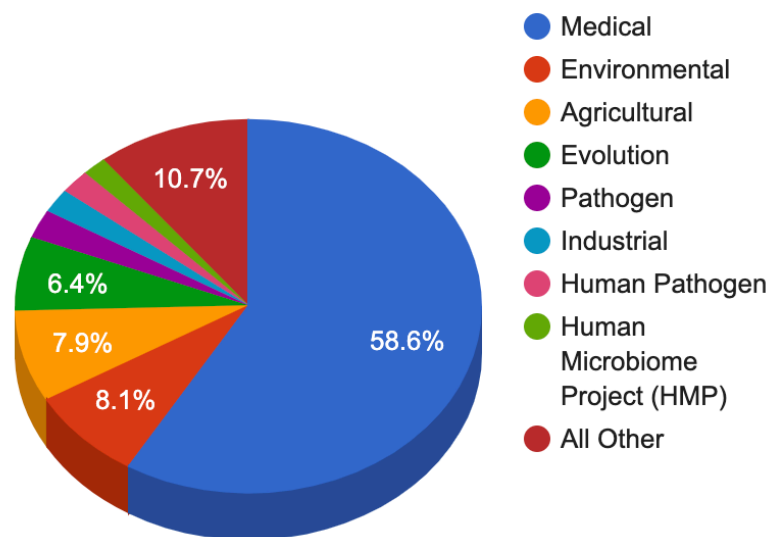


# Gene Databases



## Project Relevance of Bacterial Genome Projects

Project Relevance of Bacterial Projects





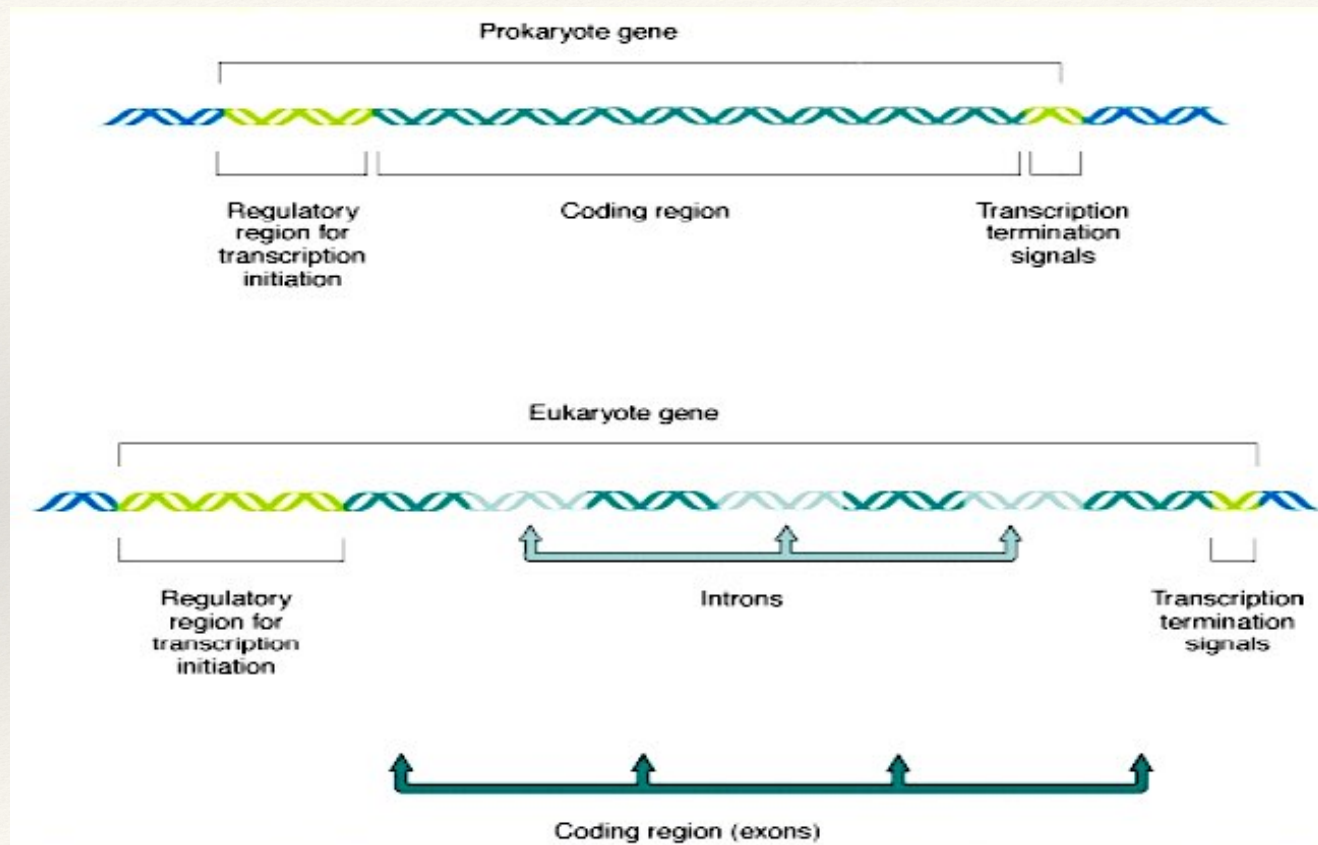
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## Genes (2)

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- ❖ The ~20,000 genes of the human genome encode > 100,000 polypeptides
- ❖ Not all of the DNA in a genome encodes protein
  - microbes: 90% coding gene
  - human: 3% coding gene
- ❖ About  $\frac{1}{2}$  of the non-coding DNA in humans is highly conserved (functionally important)

# Gene Processing



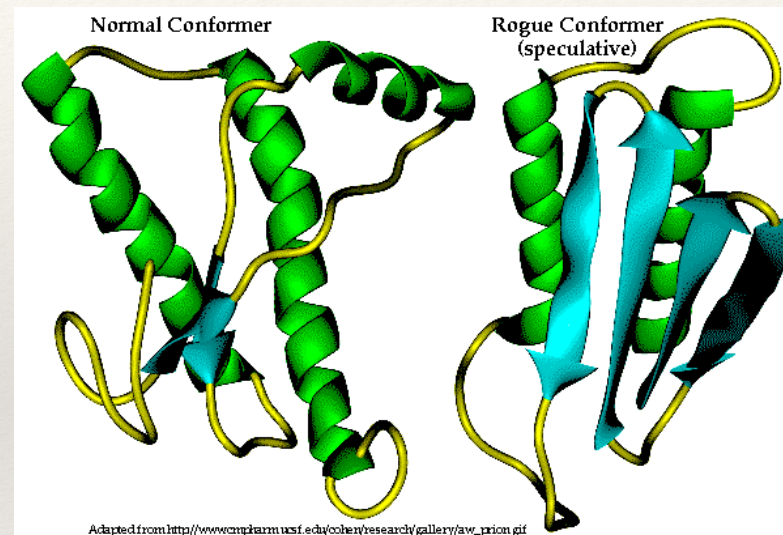


# Proteins: The Molecules of Life

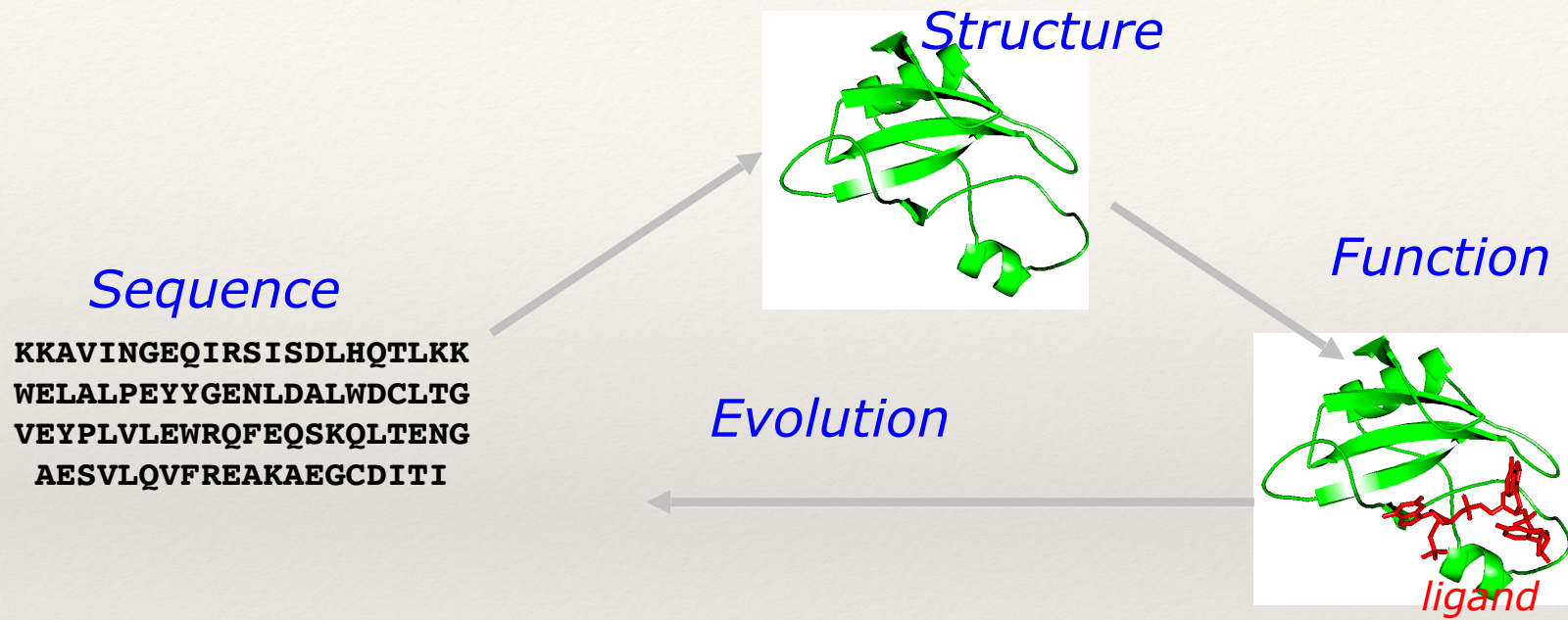
- Ubiquitous molecules that are involved in all cellular functions

- Communication agents between cells

- Failure of a protein (missing, inactive, ...) can lead to serious health problems (prions, ...)

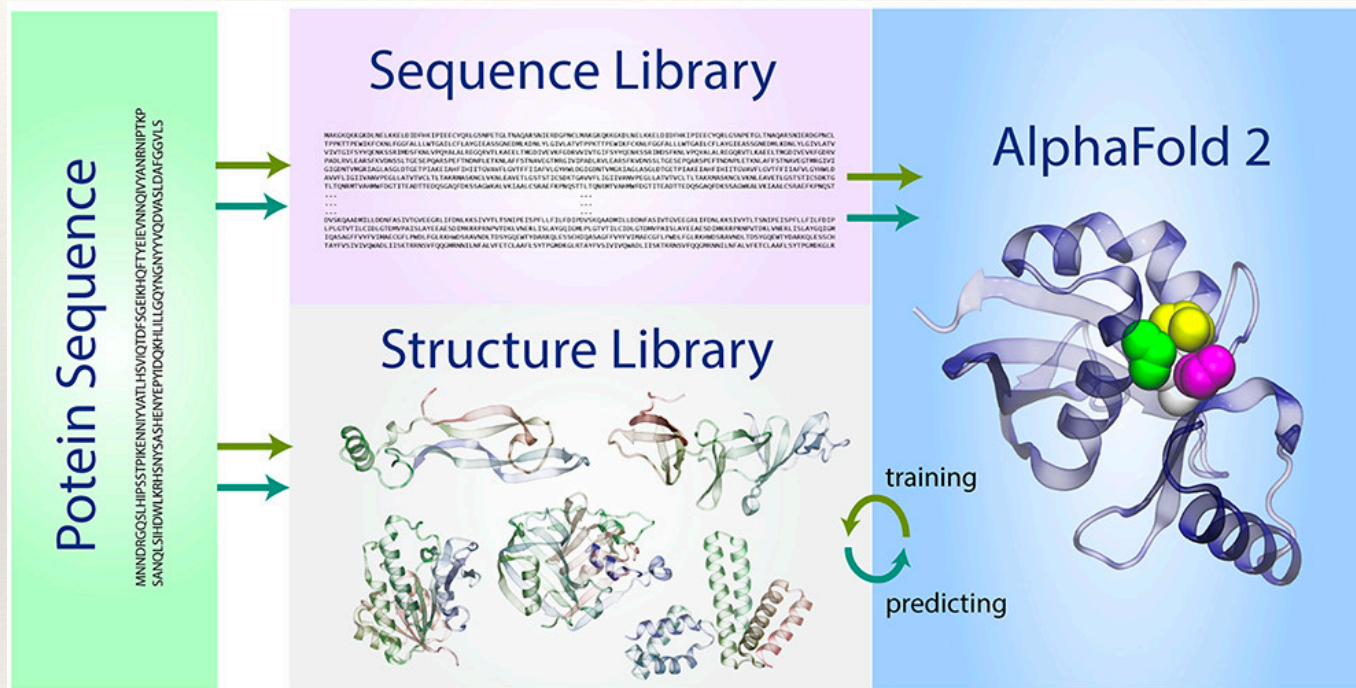


# The Protein Cycle





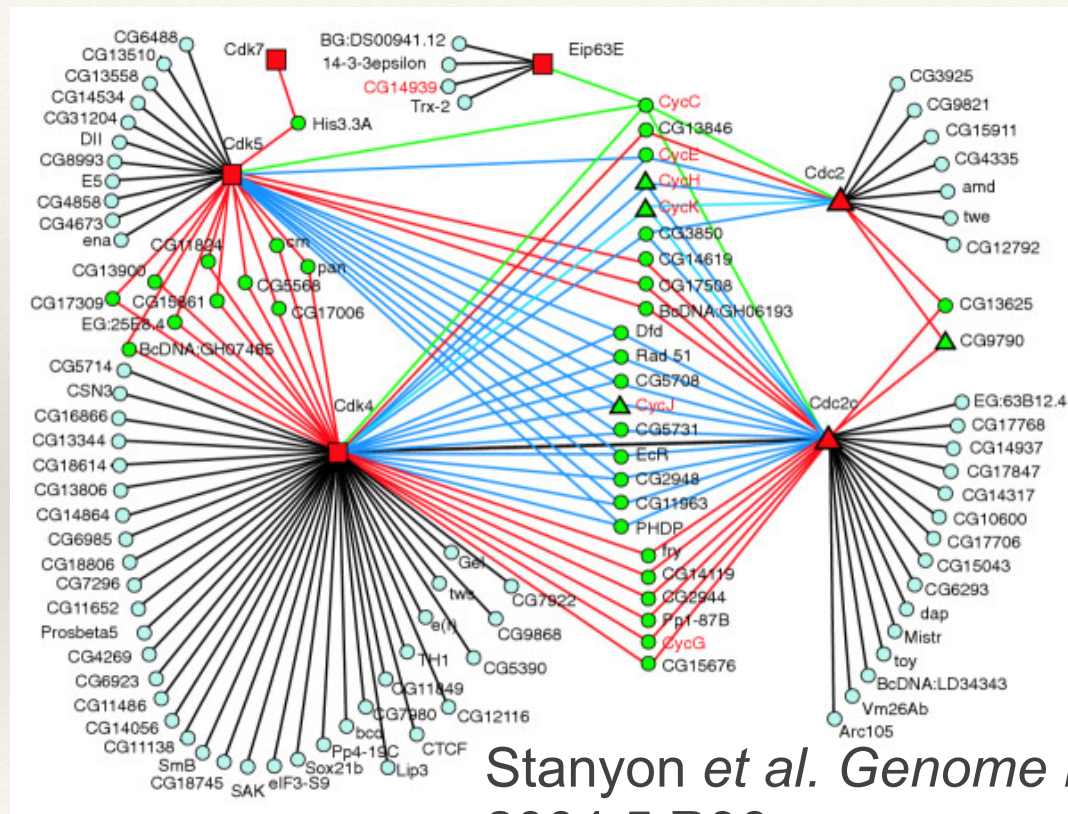
# AlphaFold2: AI for Biology





# Interactomics

Which proteins (biomolecules) interact with which proteins (biomolecules)?





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*Is there a danger, in molecular biology,  
that the accumulation of data will get  
so far ahead of its assimilation into a  
conceptual framework that the data  
will eventually prove an encumbrance ?*

John Maddox, 1988

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## Top ten challenges for bioinformatics

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- 1) Precise models of where and when transcription will occur in a genome (initiation and termination) **ability to predict where and when transcription will occur in genome**
- 2) Precise, predictive models of alternative RNA splicing: **ability to predict the splicing pattern of any primary transcript in any tissue**
- 3) Precise models of signal transduction pathways; **ability to predict cellular responses to external stimuli**
- 4) Determining protein:DNA, protein:RNA, protein:protein recognition codes
- 5) Accurate ab-initio protein structure prediction



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## Top ten challenges for bioinformatics

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- 6) Rational design of small molecule inhibitors of proteins
- 7) Mechanistic understanding of protein evolution: **understanding exactly how new protein functions evolve**
- 8) Mechanistic understanding of speciation: **molecular details of how speciation occurs**
- 9) Development of effective gene ontologies: **systematic ways to describe gene and protein function**
- 10) Education: development of bioinformatics curricula

*Source: Birney (EBI), Burge (MIT), Fickett (Glaxo)*

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# Rough Outline of the Course

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- 1) Overview of DNA, RNA and proteins
- 2) Fluctuations in biology
- 3) Sequence analysis
- 4) Structure prediction
- 5) Simulations
- 6) Drug design