

Formation of Biological Structures

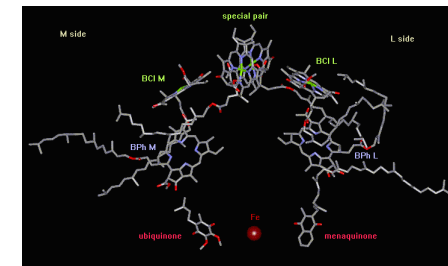
Szabolcs Osváth
Semmelweis University

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Structure – Function Relationship

From the molecular level to ecosystems, there is a strong relationship between structure and function of biological systems.

Hartmut Michel, Johann Deisenhofer, Robert Huber
1982 – 3D structure of the bacterial photosynthetic reaction center
1988 – Nobel prize



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How Big Are They?

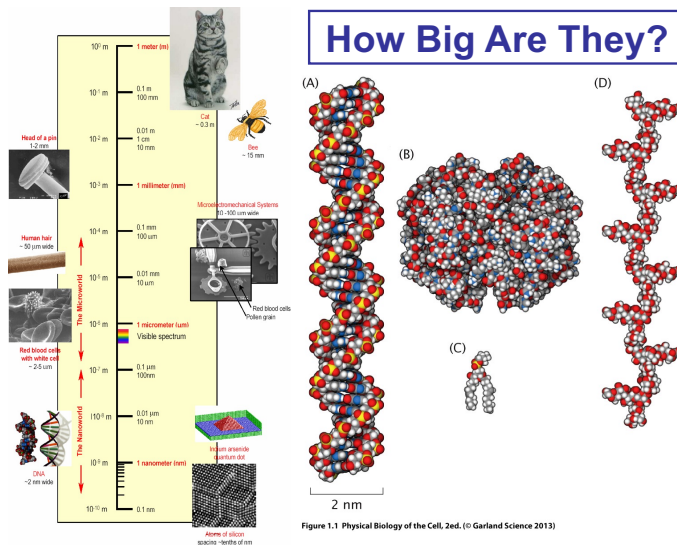


Figure 1.1 Physical Biology of the Cell, 2ed. (© Garland Science 2013)

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"Plenty of Room at the Bottom"

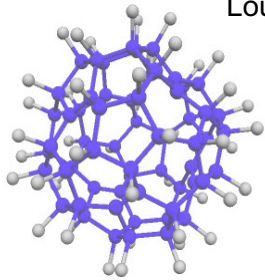
" The principles of physics, as far as I can see, do not speak against the possibility of maneuvering things atom by atom. It is not an attempt to violate any laws; it is something, in principle, that can be done; but in practice, it has not been done because we are too big."

Richard Feynman, 1959

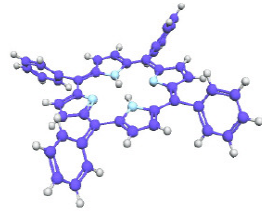
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Wave – Particle Duality

Louis De Broglie: $\lambda = h/p$



fluorofullerene
 $C_{60}F_{48}$
1632 Da

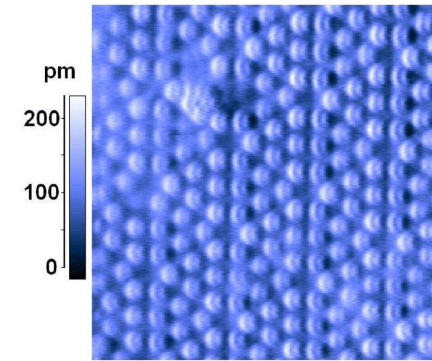


tetraphenylporphyrin $C_{44}H_{30}N_4$

L Hackermuller, S Uttenthaler, K Hornberger, E Reiger, B Brezger,
A Zeilinger, M Arndt; Phys. Rev. Lett. 91 (2003) 90408

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Wave – Particle Duality

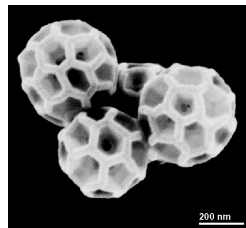
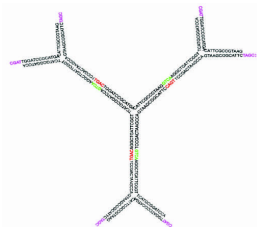


Scanning Tunneling Microscope (STM) image

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Molecular Recognition and Self Assembly

Molecular recognition
(e.g. self assembly of DNA molecules into „balls”).



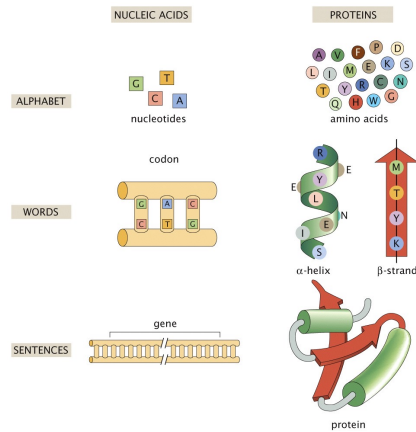
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Biopolymers

reaction	$t_{1/2}$ @ 25 °C	$t_{1/2}$ @ 100 °C	typical number of monomeric units in a polymer molecule	number of different monomers
DNA hydrolysis	140 000 years	22 years	$3 \cdot 10^9$ (human DNA)	4
RNA hydrolysis	4 years	9 days	few dozen (tRNA)	4
protein hydrolysis	400 years	5.5 weeks	few hundred	20

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Biopolymers



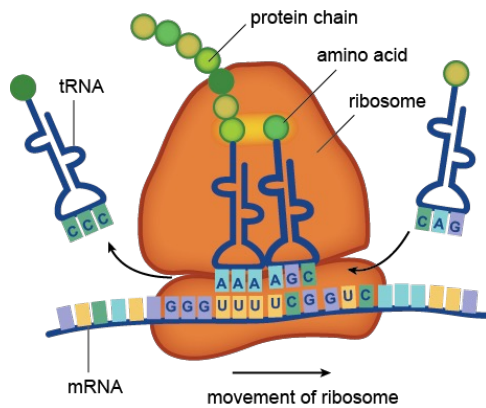
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Role of RNA in Living Systems

- messenger (mRNS)
- ribosomal (rRNS)
- transfer (tRNS)
- regulator
- enzyme (ribozyme)
- switch (riboswitch)
- virus gene RNS

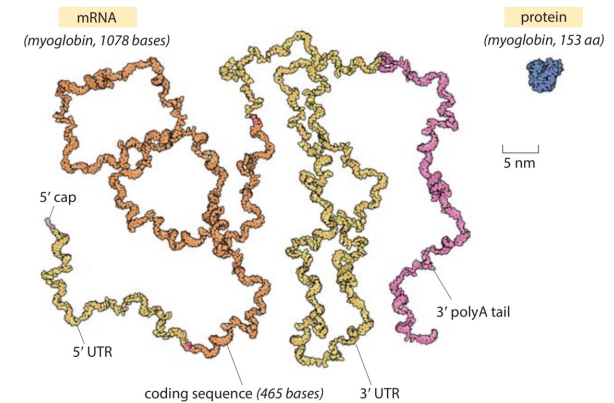
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Ribosome Function



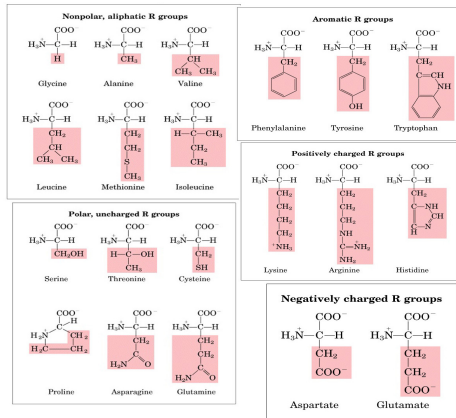
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Relationship Between RNA Code and Protein Sequence



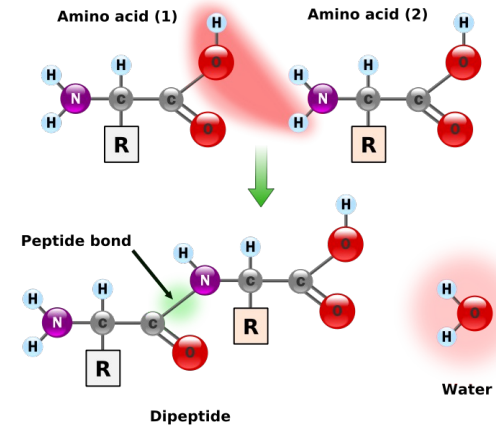
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The Twenty Standard Amino Acids



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The Peptide Bond



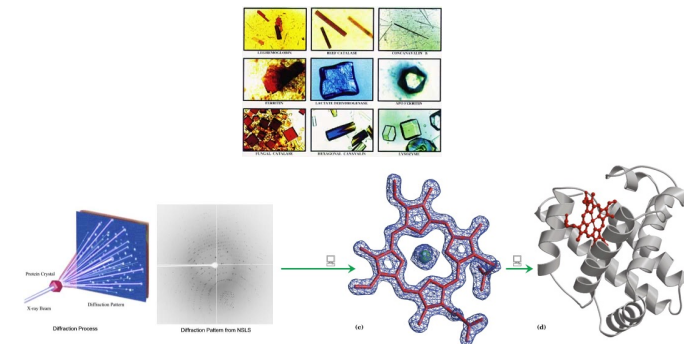
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Role of Proteins in Living Systems

- chemical catalysis
- transport
- energy conversion and storage
- coordinated movement
- mechanical skeleton
- immune response
- molecular recognition
- passing information
- gene regulation
- growth and differentiation

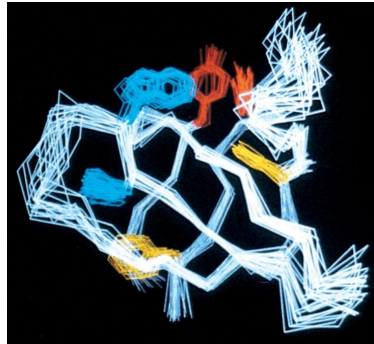
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X-ray Crystallography



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NMR Structure Determination



NMR structure of the 64 amino acid SH3 domain of the Src protein

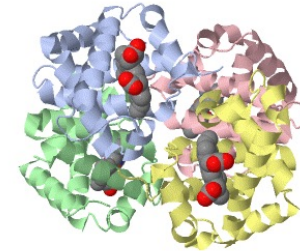
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wwPDB.org

As of March 17, 2025, the Protein Data Bank (PDB) contains **232,829 released structures**.

Visualize:

[Jmol](#)
[Pymol](#)
[VMD](#)
[Rasmol](#)



deoxyhemoglobin
(PDB entry 1A3N)

Jmol

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Interactions Stabilizing the Native State

- short range repulsion
- Van der Waals interaction
- electrostatic interaction
- hydrogen bonding
- hydrophobic interaction
- disulfide bridge

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Short Range Repulsion

Due to the exchange (Pauli) interaction, at short distances there is a strong repulsion between electrons.

The potential energy of the repulsion increases quickly with decreasing distance ($\sim 1/r^{12}$).

Atoms can be considered hard spheres with a given radius (Van der Waals radius).

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Van der Waals Interaction

Occurs between any two atoms due to the interaction of induced dipole moments.

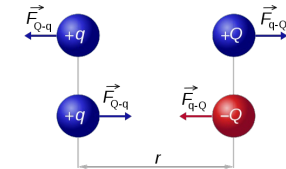
Dependence on the distance of the interaction energy: $\sim 1/r^6$

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Electrostatic Interaction

Distance dependence of the interaction energy of the Coulomb force:

$$E = \frac{q \cdot Q}{4\pi\epsilon_0\epsilon_r \cdot r}$$



The relative dielectric constant inside the protein is approx. 4, and 80 in water.

Salt bridges between ion pairs (Lys, Arg and Glu, Asp).

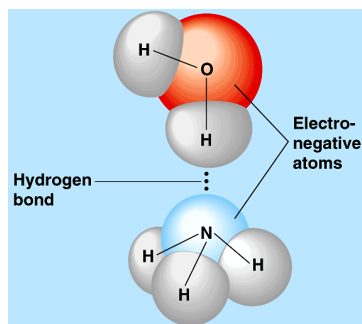
There is a large hydrate shell around charges in water.

Mobile ions can strongly shield charges.

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Hydrogen Bonding

Attraction force between a H atom of a more electronegative atom or group (hydrogen bond donor) and another atom bearing a lone pair of electrons (hydrogen bond acceptor).

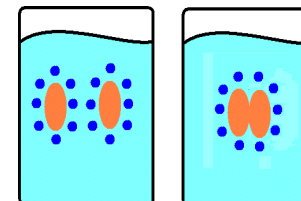


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Hydrophobic Interaction

observed tendency of nonpolar surfaces to adhere in an aqueous solution and exclude water molecule

entropic effect originating from the disruption of hydrogen bonds of liquid water by the nonpolar solute



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Disulfide Bridge

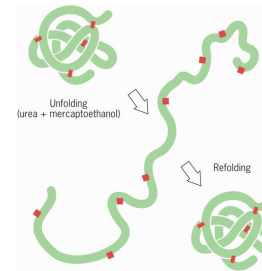
stabilizes the native structure
decreases the conformational entropy of the unfolded protein:

$$\Delta S = -2.1 \text{ J/K} - 1.5 \cdot R \cdot \ln n$$

n is the number of AAs between the two bonded AAs.

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Anfinsen's Dogma



Refolding of Ribonuclease A

Christian B. Anfinsen

The information of the 3D protein structure is encoded in the 1 D AA sequence.

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Importance of the Protein Folding Problem

One of the most important questions of molecular biophysics.

We sequence genomes, we build databases, but we can't predict protein structure and function based on the genetic information.

There are roughly two dozen conformational diseases:

Misfolded proteins and deposition of amyloid plaques was observed in various diseases (pl. Creutzfeld-Jakob disease, Alzheimer disease, Parkinson disease).

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Levinthal's Paradox - Calculation

Cyrus Levinthal

Consider a protein of 151 AAs. Assume all the 150 bonds connecting them have only two possible conformations. Assume that a reorientation of the bonds happens in 10^{-13} s.

A random search through the phase space would last:
 $2^{150} \cdot 10^{-13} \text{ s} = 4.6 \cdot 10^{24} \text{ years}$.

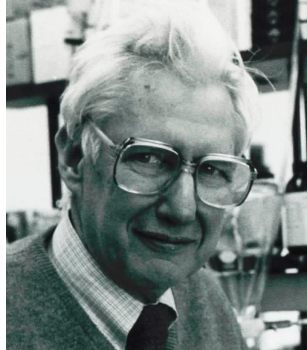
Age of Earth: $4.6 \cdot 10^9$ years
Age of the Universe: $13.7 \cdot 10^9$ years
Proteins typically fold on the ms to s timescale.

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Levinthal's Paradox - Conclusion

The phase space of a protein is way too big to find the native structure by random search.

Cyrus Levinthal
1922 - 1990



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Kinetic Pathways and Intermediate States

All proteins have a most stable conformation.

The protein can find this conformation by following a kinetic pathway and adopting specific intermediate states.

In vivo, trapping of the protein in intermediate states is prevented by protein disulfide isomerases, peptidyl prolyl isomerases and chaperones.

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Energy Landscape Models

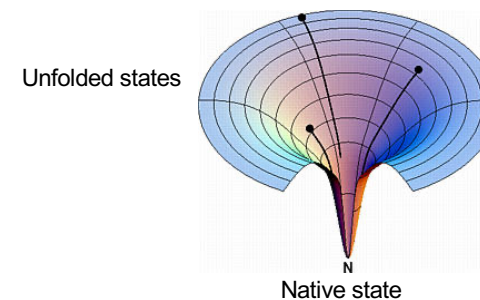
At constant pressure and temperature every thermodynamic system tends to minimize Free enthalpy (Gibbs free energy).

A free enthalpy (Gibbs free energy) value is associated to every conformation of the protein.

The protein does not search through the entire phasespace, but starts to "flow" towards lower free enthalpies.

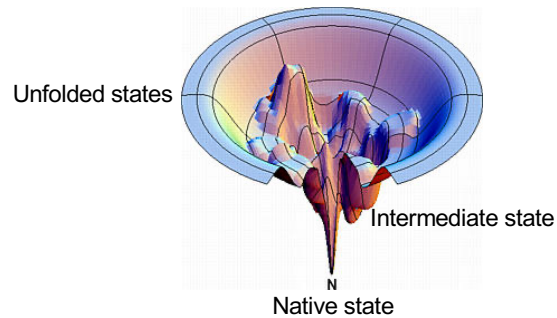
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Smooth Funnel



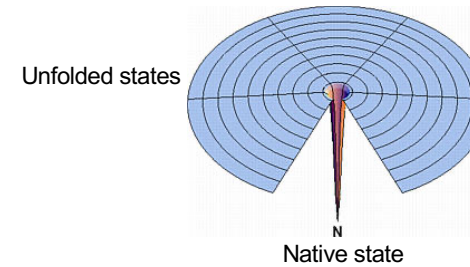
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Rugged Funnel



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Energy Landscape View of Levinthal's Paradox



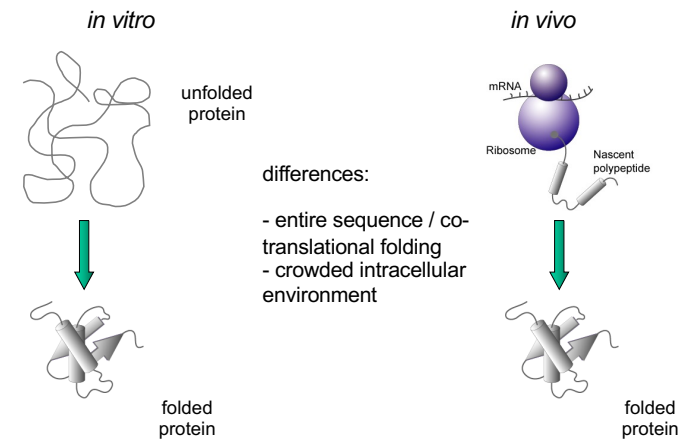
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Comparison of the Two Folding Models

Pathways	Landscape
Given pathways	Energy landscape
Well distinguished intermediates	Multitude of intermediates
Consecutive steps	Parallel folding routes
Classical chemical kinetics applied to protein folding	Statistical physics developed to understand spin glasses

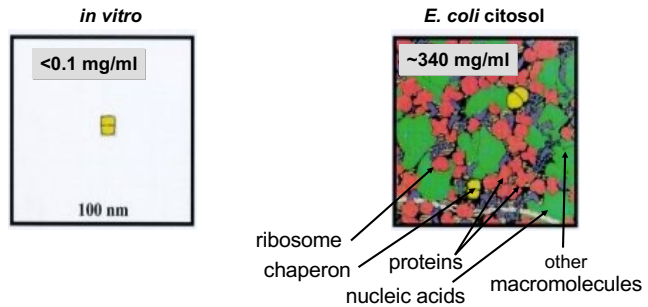
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In vitro and *in vivo* Folding



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Molecular Crowding



In vitro experiments

- lack of binding partner molecules
- lack of posttranslational modifications
- very different physico-chemical environment than in a cell

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Effect of Molecular Crowding

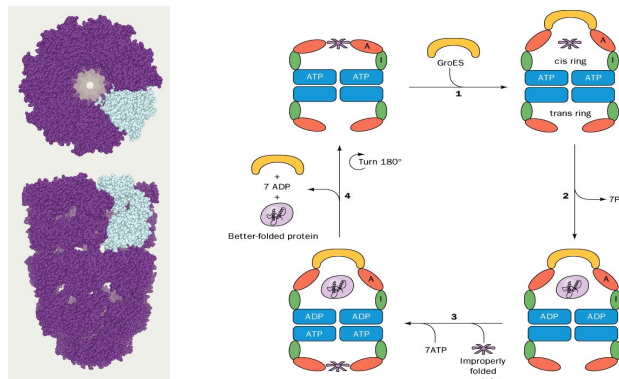
Molecular crowding:

a large fraction of the volume of the cytoplasm is filled by other molecules than water.

- dissociation constants decrease
- speed of protein-protein association increases
- association of fully or partially denatured proteins speeds up

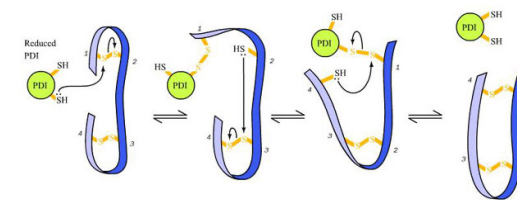
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GroEL/ES Chaperon Cycle



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Protein Disulfide Isomerase Function



structure of the human protein disulfide isomerase



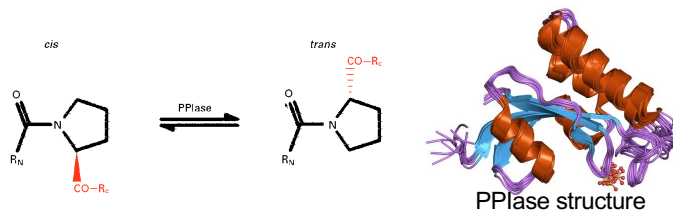
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Proline Cis/Trans Isomerase

Due to the activation barrier between the cis and trans prolines, the presence of cis prolines in the native structure:

- speeds up early folding steps
- slows down the final formation of the native structure.

PPIase (peptidyl-prolyl isomerase)



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Fate of the Protein in Eukaryotic Cells

cytosol	protein synthesis and folding,
extracellular volume	export of folded protein
mitochondrion	limited protein synthesis
chloroplast	limited protein synthesis
endoplasmic reticulum	import of unfolded protein
peroxisome	import of folded protein
nucleus	import of folded protein
lysosome	import of unfolded protein

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AI-Powered Protein Structure Prediction

Google DeepMind
AlphaFold Project

2024 Nobel Prize
in chemistry



Demis Hassabis & John Jumper

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Designing New Proteins

2024 Nobel Prize
in chemistry



David Baker

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