

# A Different Kind Of Stability

## Energy of Protein Formation

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# Motivation

## Background

Protein geochemistry is concerned with the occurrence and variation of proteins in all environments on Earth.

## Example

Amino acid differences between mitochondrial and nuclear proteins

# Motivation

## Background

Protein geochemistry is concerned with the occurrence and variation of proteins in all environments on Earth.

## Hypothesis

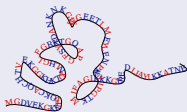
Molecular evolution is a type of chemical reaction.

## Outline

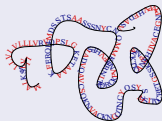
- 1 Protein Formation Reactions
- 2 Calculating Relative Stabilities
- 3 Natural Experiment

# Folding Reactions

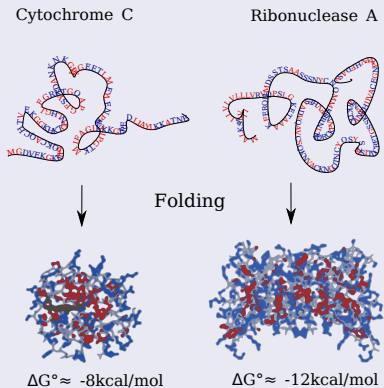
Cytochrome C



Ribonuclease A



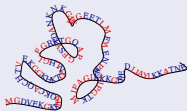
# Folding Reactions



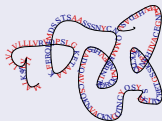
- Folding as a conformational process
- Stability referenced to unfolded protein
- Cellular/laboratory timescales

# Formation Reactions

Cytochrome C

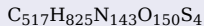


Ribonuclease A



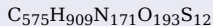
# Formation Reactions

Cytochrome C



$$\Delta G_f^\circ \approx -3650 \text{ kcal/mol}$$

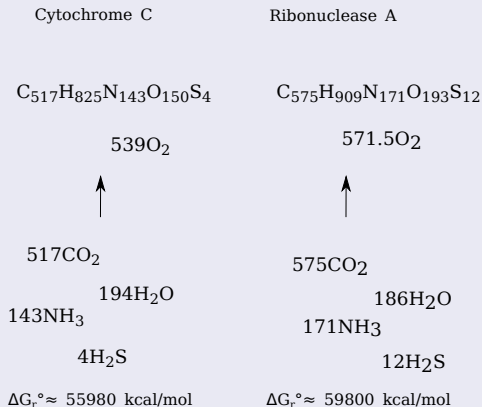
Ribonuclease A



$$\Delta G_f^\circ \approx -4960 \text{ kcal/mol}$$

- Formation as a chemical process
- $\Delta G_f^\circ$ : Standard Gibbs energy of formation from the elements

# Formation Reactions

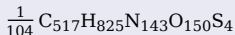


- Formation as a chemical process
- Stability referenced to inorganic species
- $\Delta G_r^\circ$ : Standard Gibbs energy of reaction
- Overall energy change is independent of mechanism



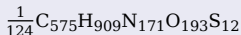
# Residue Equivalents

## Cytochrome C



$$\Delta G_r^\circ \approx 538 \text{ kcal/mol}$$

## Ribonuclease A



$$\Delta G_r^\circ \approx 482 \text{ kcal/mol}$$

- Reactions normalized by protein length
- Energetic meaning of reaction coefficients

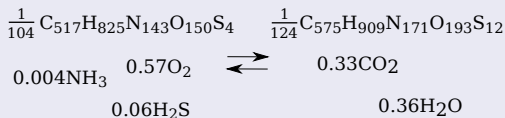
## Environment & Energy

Shift to lower  $\text{O}_2$  potential more strongly decreases Gibbs energy of formation of CYC than RNAS1.

# Residue Equivalents

Cytochrome C

Ribonuclease A

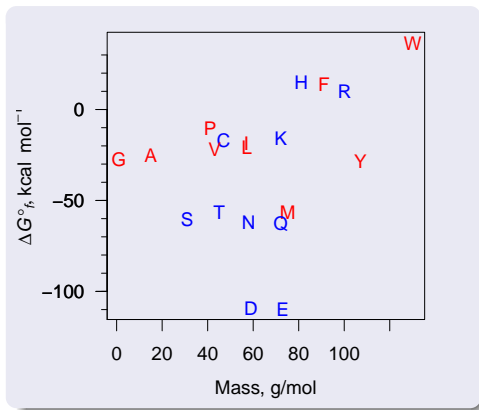


- Reactions normalized by protein length
- Energetic meaning of reaction coefficients
- Transformation reaction; cellular to evolutionary timescales

## Environment & Energy

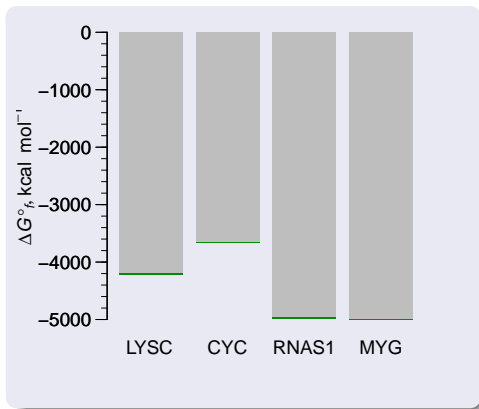
Shift to lower  $\text{O}_2$  potential more strongly decreases Gibbs energy of formation of CYC than RNAS1.

# Standard Gibbs Energies



- Group additivity of aqueous species properties for amino acid residues [Dick et al., 2006]

# Standard Gibbs Energies



- Group additivity of aqueous species properties for amino acid residues [Dick et al., 2006]
- Protein size dependence of standard Gibbs energies

LYSC	Lysozyme	129
CYC	Cytochrome C	104
RNAS1	Ribonuclease A	124
MYG	Myoglobin	153

- Gibbs energies of folding [Privalov and Khechinashvili, 1974] are 1% or less of energies of formation.

# Computational Plan

- Calculate standard Gibbs energies of proteins from amino acid group additivity and equations of state for aqueous species.
- Proceed without accounting for energy of protein folding.
  - Per mole of protein, energy of folding is small compared to Gibbs energy of chemical formation reaction.
  - If all proteins are folded, energy of folding tends to cancel in relative stability calculations.

# Metastable Equilibrium

- Chemical affinity is opposite of Gibbs energy change due to reaction progress,

$$dG = -SdT + VdP - Ad\xi$$

$G$       Gibbs energy

$S$       Entropy

$T$       Temperature

$V$       Volume

$P$       Pressure

$A$       Chemical Affinity

$\xi$       Reaction Progress

# Metastable Equilibrium

- Chemical affinity is opposite of Gibbs energy change due to reaction progress,
- computed as a function of standard Gibbs energy, reaction stoichiometry and chemical activities.

$$dG = -SdT + VdP - Ad\xi$$

$$A = 2.303RT \log(K / Q)$$

$A$	Chemical Affinity
$K$	Equilibrium Constant
$Q$	Activity Product
$R$	Gas Constant

# Metastable Equilibrium

- Chemical affinity is opposite of Gibbs energy change due to reaction progress,
- computed as a function of standard Gibbs energy, reaction stoichiometry and chemical activities.

$$dG = -SdT + VdP - Ad\xi$$

$$A = 2.303RT \log(K / Q)$$

$$A = -\Delta G_r$$

$$K = 10^{(-\Delta G_r^\circ / 2.303RT)}$$

$$Q = 10^{\sum \nu \log a} = \prod a^\nu$$

$$\nu \quad \text{Reaction Coefficient}$$

$$a \quad \text{Chemical Activity}$$



# Metastable Equilibrium

- Chemical affinity is opposite of Gibbs energy change due to reaction progress,
- computed as a function of standard Gibbs energy, reaction stoichiometry and chemical activities.
- Maxwell-Boltzmann distribution allows for a transformation between reference states.

- $dG = -SdT + VdP - Ad\xi$

- $A = 2.303RT \log(K / Q)$

- $$\frac{a}{\sum a} = \frac{e^{A/RT}}{\sum e^{A/RT}}$$

Equal-activity reference state

More stable: higher affinity ( $A$ )

# Metastable Equilibrium

- Chemical affinity is opposite of Gibbs energy change due to reaction progress,
- computed as a function of standard Gibbs energy, reaction stoichiometry and chemical activities.
- Maxwell-Boltzmann distribution allows for a transformation between reference states.
- When the chemical affinities of the formation reactions are all equal, the proteins are in metastable equilibrium.

- $dG = -SdT + VdP - Ad\xi$

- $A = 2.303RT \log(K / Q)$

- $$\frac{a}{\sum a} = \frac{e^{A/RT}}{\sum e^{A/RT}}$$

Equal-activity reference state

More stable: higher affinity ( $A$ )

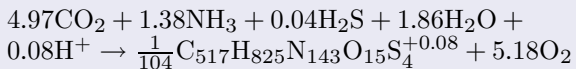
Equal-affinity reference state

More stable: higher activity ( $a$ )

# Computational Plan

- Calculate standard Gibbs energies of proteins from amino acid group additivity and equations of state for aqueous species.
- Proceed without accounting for energy of protein folding.
- Calculate equilibrium activities of proteins.
  - Start with chemical affinities in equal-activity reference state.
  - Use reference state transformation to calculate equilibrium activities.

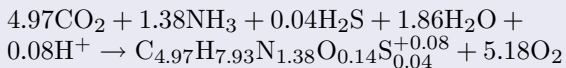
# Formation of Ionized Unfolded CYC\_BOVIN



- Ionization of proteins using additivity

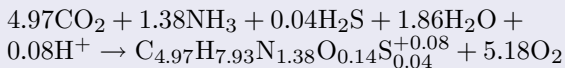
▶ [more info?](#)

# Formation of Ionized Unfolded CYC\_BOVIN



- Ionization of proteins using additivity  
[▶ more info?](#)
- Write per-residue formulas.

# Formation of Ionized Unfolded CYC\_BOVIN



$$\log K = -\Delta G_r^\circ / 2.303RT$$

$$= -393.0$$

- Ionization of proteins using additivity

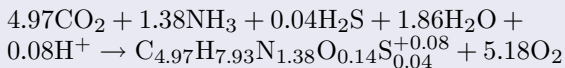
[▶ more info?](#)

- Write per-residue formulas.

$T$  25 °C

$P$  1 bar

# Formation of Ionized Unfolded CYC\_BOVIN



$$\begin{aligned} \log Q &= \log a_{\text{C}_{4.97}\text{H}_{7.93}\text{N}_{1.38}\text{O}_{0.14}\text{S}_{0.04}^{+0.08}} + \\ &5.18 \log f_{\text{O}_2} - 4.97 \log a_{\text{CO}_2} - 1.38 \log a_{\text{NH}_3} - \\ &0.04 \log a_{\text{H}_2\text{S}} - 1.86 \log a_{\text{H}_2\text{O}} - \log a_{\text{H}^+} \\ &= -393.4 \end{aligned}$$

- Ionization of proteins using additivity

[▶ more info?](#)

- Write per-residue formulas.

$\log a_{\text{residue}}$	0
$\log a_{\text{CO}_2}$	-3
$\log a_{\text{H}_2\text{O}}$	0
$\log a_{\text{NH}_3}$	-4
$\log f_{\text{O}_2}$	-80
$\log a_{\text{H}_2\text{S}}$	-7
pH	7

# Formation Properties

## Values per Residue

Protein	$\log K$	$\log Q$	$\log a$	$A/2.303RT$
LYSC	-361.6	-357.4	0	-4.17
CYC	-393.0	-393.4	0	0.34
RNAS1	-352.6	-348.4	0	-4.27
MYG	-407.6	-408.6	0	0.96

- Equal activity reference state: MYG is more stable.



# Formation Properties

## Values per Residue

Protein	$\log K$	$\log Q$	$\log a$	$A/2.303RT$
LYSC	-361.6	-362.0	-4.62	0.45
CYC	-393.0	-393.5	-0.10	0.45
RNAS1	-352.6	-353.1	-4.72	0.45
MYG	-407.6	-408.0	0.51	0.45

- Equal activity reference state: MYG is more stable.
- Equal affinity reference state: MYG is still more stable!

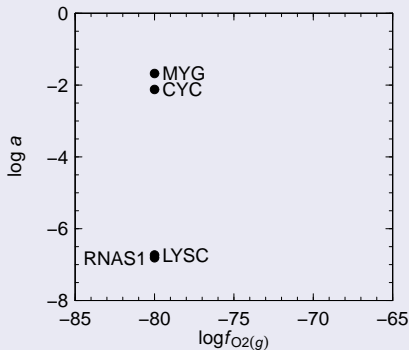
# Formation Properties

## Values per Residue

Protein	$\log K$	$\log Q$	$\log a$	$A/2.303RT$	$\log a_{\text{protein}}$
LYSC	-361.6	-362.0	-4.62	0.45	-6.73
CYC	-393.0	-393.5	-0.10	0.45	-2.12
RNAS1	-352.6	-353.1	-4.72	0.45	-6.81
MYG	-407.6	-408.0	0.51	0.45	-1.68

- Equal activity reference state: MYG is more stable.
- Equal affinity reference state: MYG is still more stable!
- Molality of residue = molality of protein \* protein length
- Activity of residue = activity of protein \* protein length (assuming ideality)

# Equilibrium Activity Diagrams

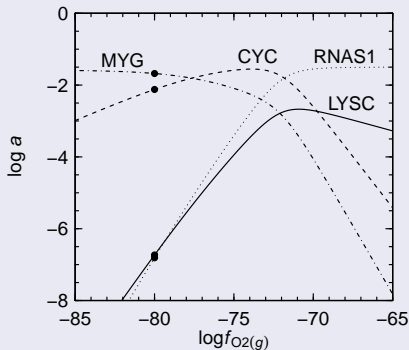


- Metastable equilibrium activities with total activity of residues = 4
- We are using  $T = 25\text{ }^{\circ}\text{C}$  and  $\text{pH} = 7$

It Shows ...

MYG is relatively most stable at  $\log f_{O_2} = -80$ .

# Equilibrium Activity Diagrams



- Metastable equilibrium activities as a function of  $\log f_{O_2}$  with total activity of residues = 4
- $\log f_{O_2}$  can be converted to other measurements of oxidation-reduction potential. [▶ how?](#)

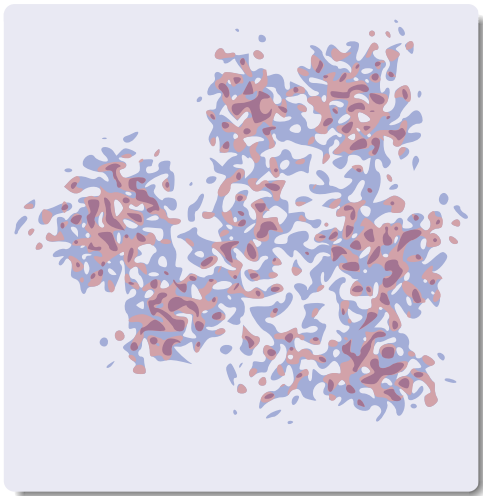
## It Shows ...

Relative stability is sensitive to oxidation potential.

# Computational Plan

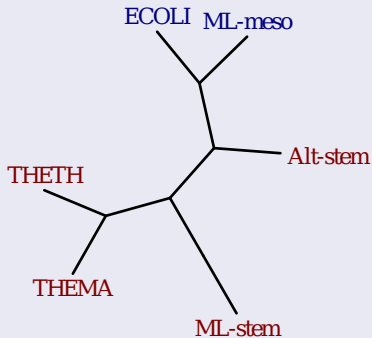
- Calculate standard Gibbs energies of proteins from amino acid group additivity and equations of state for aqueous species.
- Proceed without accounting for energy of protein folding.
- Calculate equilibrium activities of proteins.
  - Relative stabilities depend on the species (chemical compositions, standard Gibbs energies).
  - Relative stabilities depend on the environment (temperature, pressure, activities/fugacities of basis species).

# Elongation Factor Tu



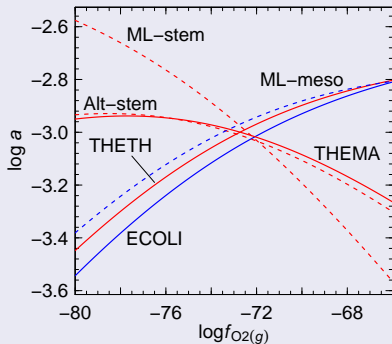
- EF-Tu from *Escherichia coli*

# Elongation Factor Tu



- EF-Tu's from *Escherichia coli*, *Thermotoga maritima*, *Thermus thermophilus*
- and reconstructed by maximum likelihood (**ML**) **stem** of bacterial tree, stem of **mesophilic** bacteria, and **Alternative** tree topology [Gaucher et al., 2003]
- This tree built using parsimony (PHYLP software), 394 aligned amino acids.

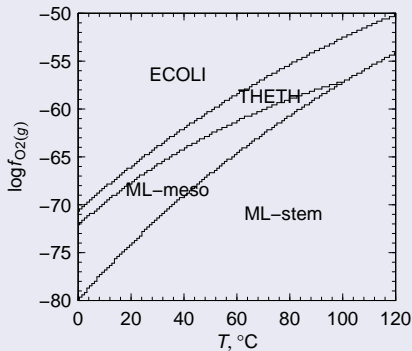
# Equilibrium Activity Diagrams



- Equilibrium activity diagram
- In most reactions, proteins from higher temperature favored by lower  $\log f_{O2(g)}$ .

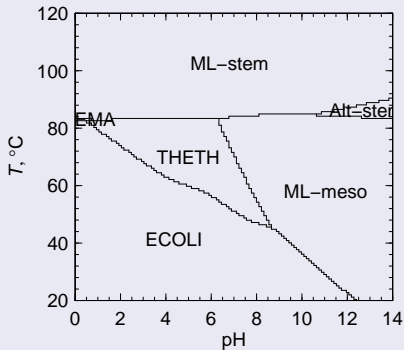


# Equilibrium Activity Diagrams



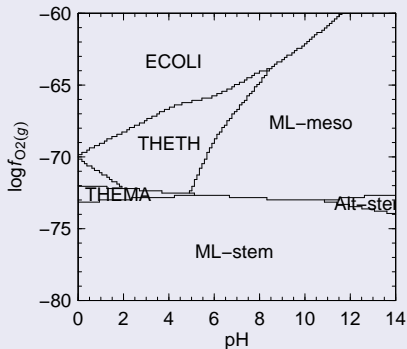
- Equilibrium predominance diagram
- At constant  $f_{O_2(g)}$ , increasing  $T$  tends to favor “ML-stem”.

# Equilibrium Activity Diagrams



- Equilibrium predominance diagram
- $\log f_{O_2} = -60$

# Equilibrium Activity Diagrams



- Equilibrium predominance diagram
- $T = 25\text{ }^{\circ}\text{C}$

## Multidimensionality

What about  $\log a_{H_2O}$ ,  $\log a_{CO_2}$ , etc.?

# Computational Plan

- Calculate standard Gibbs energies of proteins from amino acid group additivity and equations of state for aqueous species.
- Proceed without accounting for energy of protein folding.
- Calculate equilibrium activities of proteins.
- Investigate temperature, oxidation potential, pH, other chemical potentials.
  - Are reducing conditions associated with hotter environments?
  - The system is multidimensional; could also vary the chemical potentials of carbon, nitrogen, sulfur.

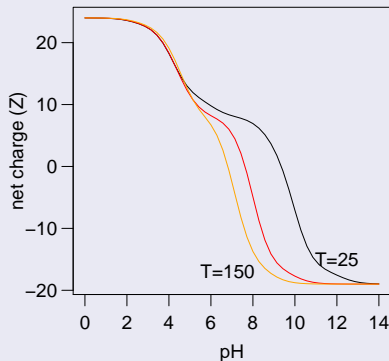
# Computational Plan

- Calculate standard Gibbs energies of proteins from amino acid group additivity and equations of state for aqueous species.
- Proceed without accounting for energy of protein folding.
- Calculate equilibrium activities of proteins.
- Investigate temperature, oxidation potential, pH, other chemical potentials.
- Explore the protein universe using model systems.
- CHNOSZ is the software package used for the preceding calculations.

# References

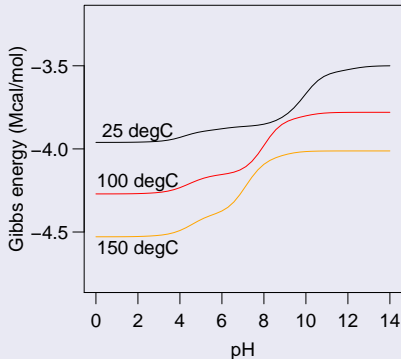
- J. M. Dick, D. E. LaRowe, and H. C. Helgeson. Temperature, pressure, and electrochemical constraints on protein speciation: Group additivity calculation of the standard molal thermodynamic properties of ionized unfolded proteins. *Biogeosciences*, 3(3):311 – 336, 2006. URL <http://www.biogeosciences.net/3/311/2006/>.
- E. A. Gaucher, J. M. Thomson, M. F. Burgan, and S. A. Benner. Inferring the palaeoenvironment of ancient bacteria on the basis of resurrected proteins. *Nature*, 425(6955):285 – 288, SEP 18 2003. ISSN 0028-0836. URL <http://dx.doi.org/10.1038/nature01977>.
- P. L. Privalov and N. N. Khechinashvili. A thermodynamic approach to the problem of stabilization of globular protein structure: A calorimetric study. *J. Mol. Biol.*, 86(3):665 – 684, 1974. ISSN 0022-2836. URL [http://dx.doi.org/10.1016/0022-2836\(74\)90188-0](http://dx.doi.org/10.1016/0022-2836(74)90188-0).

# Protein Ionization (CYC\_BOVIN)



- Net charges computed additively [Dick et al., 2006] using temperature-dependent sidechain  $pK_a$  values

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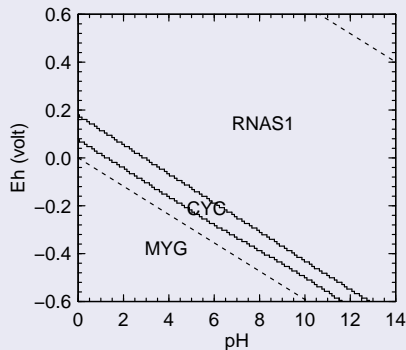


- Net charges computed additively [Dick et al., 2006] using temperature-dependent sidechain  $pK_a$  values
- Also affects standard Gibbs energies of the ionized proteins

[◀ return](#)

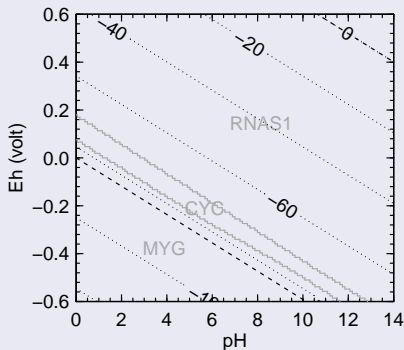


# Oxygen Fugacity



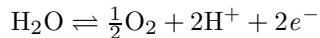
- Eh-pH diagram for proteins
- Dashed lines indicate stability limits of H<sub>2</sub>O:  
log  $f_{O_2} = 0$  (upper),  
log  $f_{O_2} = -83.1$  (lower)

# Oxygen Fugacity



- Eh-pH diagram for proteins

- Convert between Eh and  $\log f_{O_2}$  using law of mass action for



- $Eh = \frac{RT}{F} pe = -\frac{RT}{F} \log a_{e^-}$

◀ return