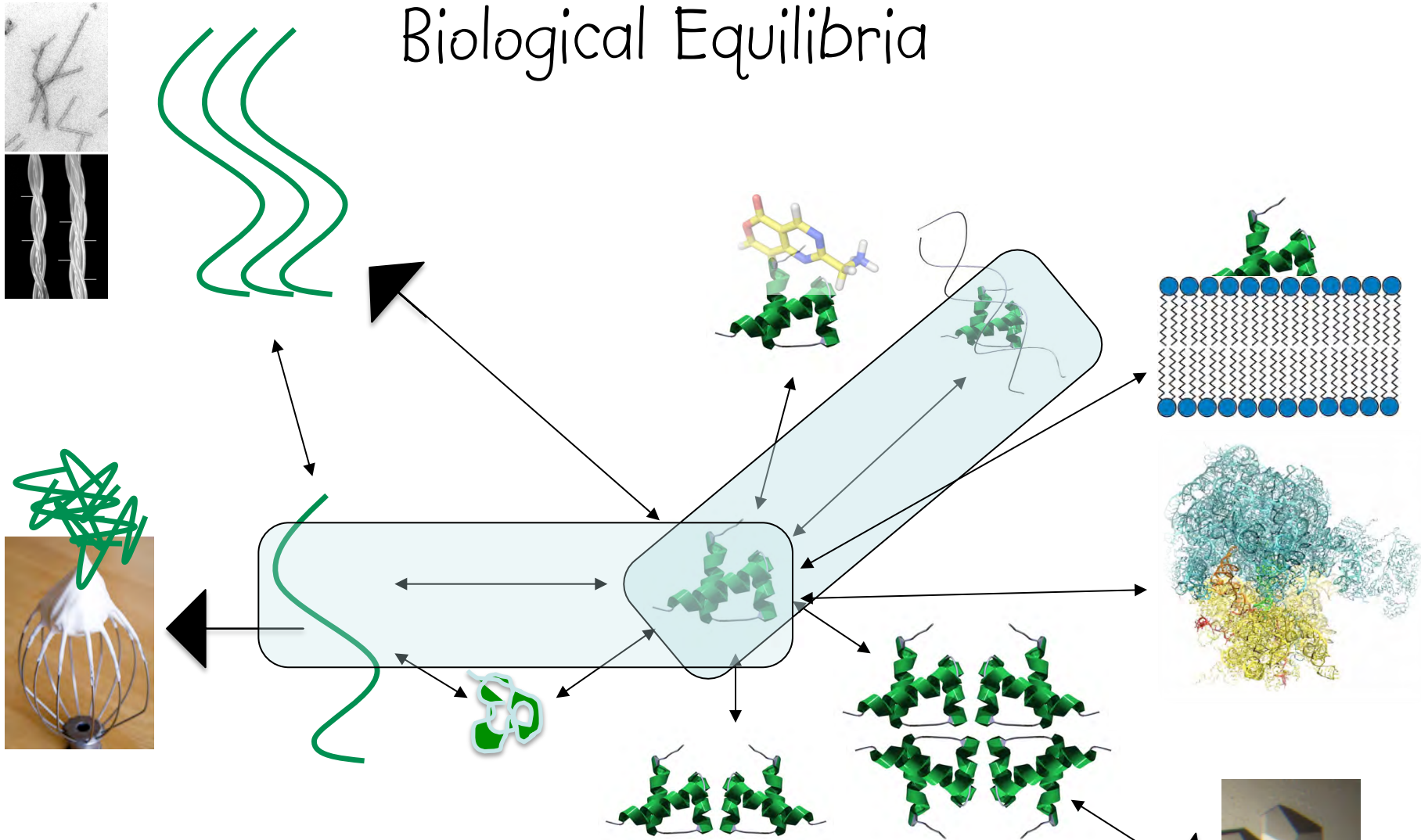


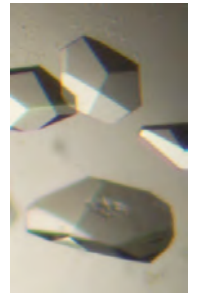
- Equilibrium Thermodynamics
- Biological Calorimetry

Chris Johnson

Biological Equilibria



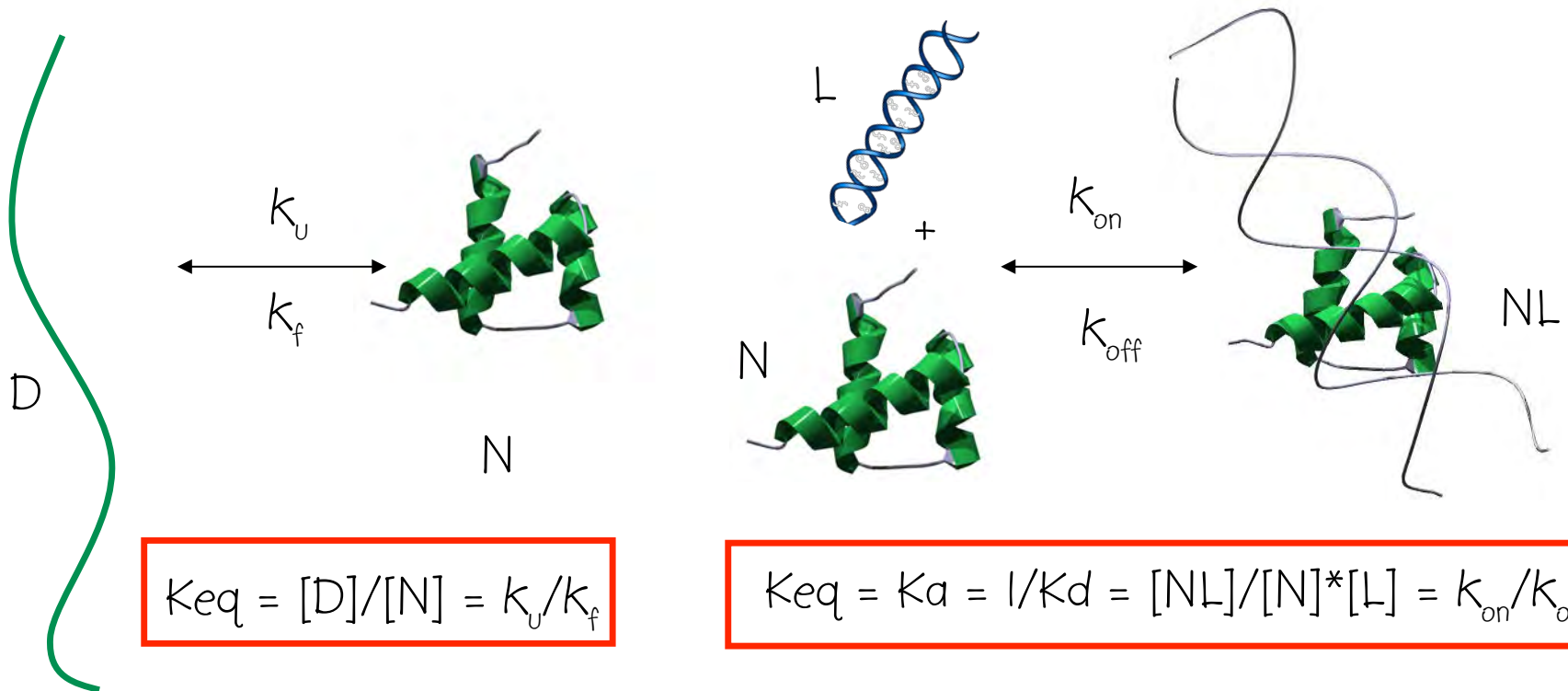
Much of biology is based around 'reversible' equilibrium processes that are mediated by consolidated non-covalent interactions which are individually 'weak' in strength



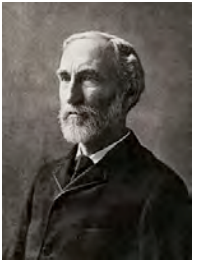
Biological Reversible Equilibria

For example, protein folding or protein-DNA binding

- At equilibrium the 'reaction' position (i.e., the biology) is described by the equilibrium constant, K_{eq} , which is the ratio of the concentrations (products / reactants) which are determined by the ratio of the forward and reverse rate constants (for kinetic methods come to lecture 10)



Biological Equilibrium Thermodynamics



Josiah Willard Gibbs
1839 – 1903

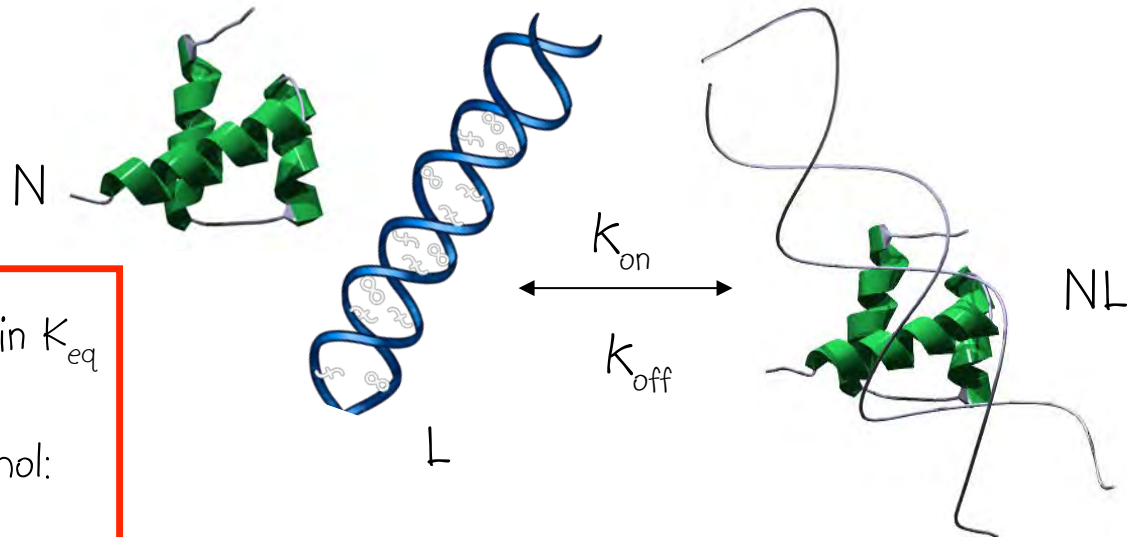
- The Gibbs Free Energy (ΔG) is a logarithmic representation of the equilibrium position.

$$\Delta G = -RT \ln K_{eq}$$

- It has 'components' of 'Enthalpy' (ΔH) and 'Entropy' (ΔS)

$$\Delta G = \Delta H - T\Delta S$$

- When ΔG is -ve, $\ln K_{eq}$ is +ve, so $K_{eq} > 1$ and so the products are in excess (e.g., for K_a this is NL)



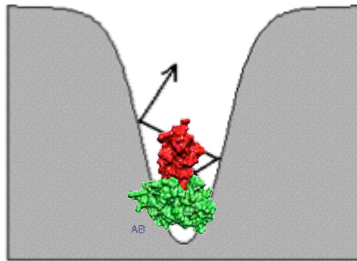
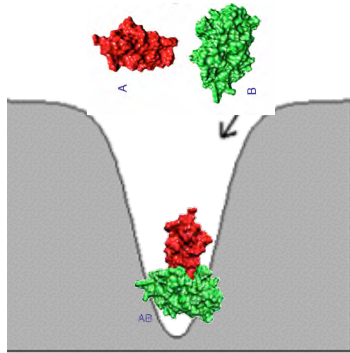
Each order of magnitude in K_{eq}
gives $\Delta \ln 2.303$

At 298K RT is 0.59 kcal/mol:

$$\Delta\Delta G = 1.36 \text{ kcal/mol}$$

What are Enthalpy and Entropy?

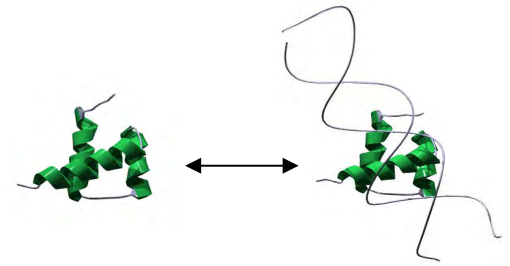
A Simplified View



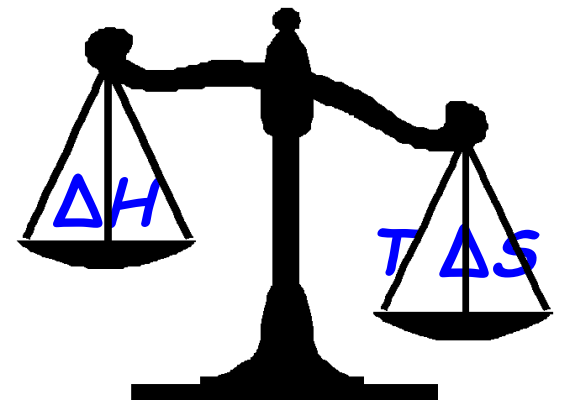
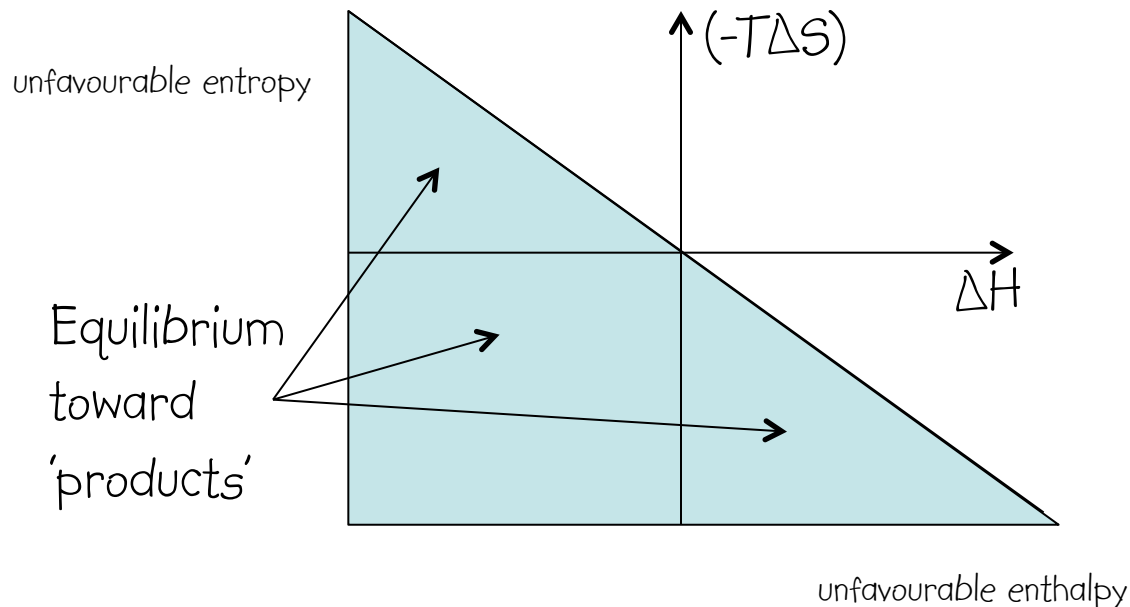
- Systems naturally progress to a lower internal energy level; lower enthalpy,
- The change, ΔH , is -ve and heat given off, "exothermic", equilibrium product favoured.
- Systems want more ways of configuring with the same energy (more 'disorder'). Statistically the more ways of achieving an outcome there are the more probable that outcome. Temperature modulates this probability effect through thermal motion ($-T\Delta S$)
- Entropy is a measure of the number of ways of arranging system energy; ΔS , is +ve (more ways) equilibrium product favoured ($-T\Delta S$ is -ve)

$$\Delta G = -RT \ln K_{eq} = \Delta H - T\Delta S$$

Where does the equilibrium end up?

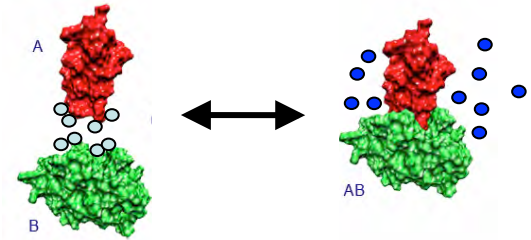


- Simply the balance of enthalpy and temperature-entropy gives ΔG , and thus defines the equilibrium position.
- Temperature is a key factor through $T\Delta S$ term (think thermal melting)
- A -ve ΔG is 'favourable' and can be obtained even when one component is 'unfavourable' if the other is 'favourable' and larger



Are ΔH and ΔS Fixed values?

Changes in heat capacity



- In biological equilibria there are typically large changes in solvation which affects the system heat capacity (C_p), that is the energy needed to increase temperature by 1 °K, units, $\text{cal mol}^{-1} \text{K}^{-1}$. These reactions have a **change** in heat capacity, ΔC_p
- The integral of C_p or C_p/T from zero K to temperature T represents the enthalpy and entropy levels for a system
- Because of the significant ΔC_p for biological equilibria the corresponding ΔH , ΔS (and thus ΔG) are all temperature dependent with ΔS in a non linear manner

$$\Delta G_{T_2} = \left[\Delta H_{T_1} + \Delta C_p \cdot (T_2 - T_1) \right] - T_2 \left[\Delta S_{T_1} + \Delta C_p \cdot \ln \left(\frac{T_2}{T_1} \right) \right]$$

Other than temperature and ΔC_p can anything else affect ΔG ?



Henry Louis Le Chatelier
1850-1936

... many things because

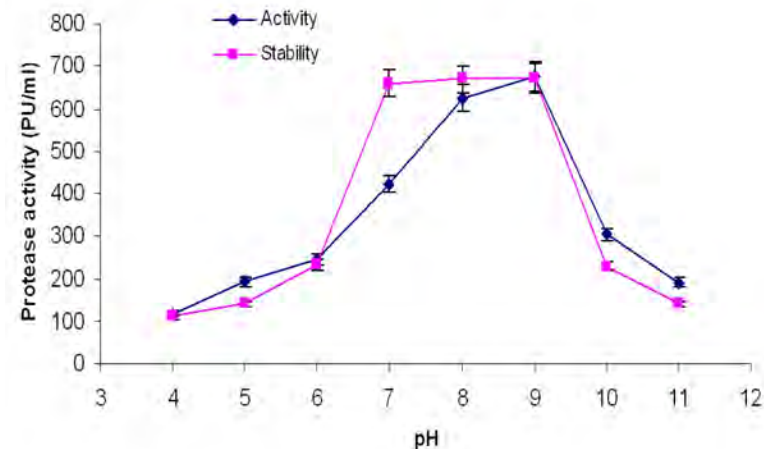
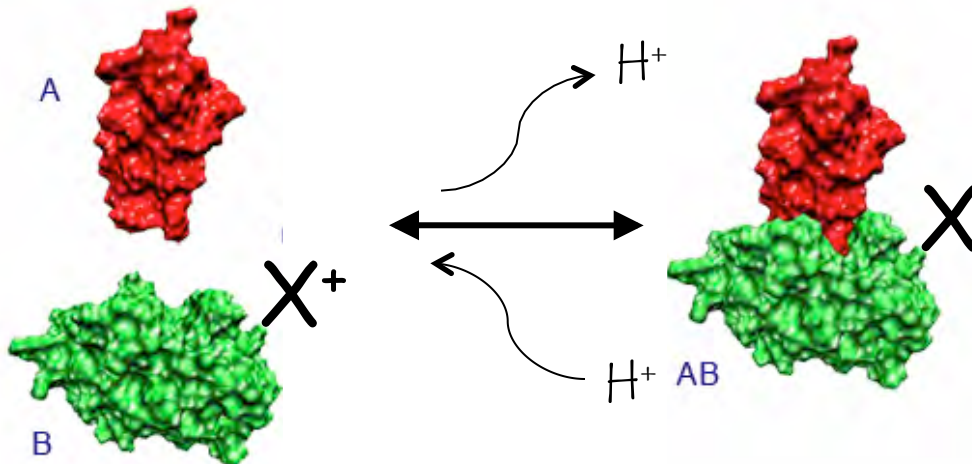
Law of Mass Action; Le Chatelier's principle

"When a system at dynamic equilibrium is disturbed, the equilibrium position will shift in the direction which tends to minimise, or counteract, the effect of the disturbance"

For example....

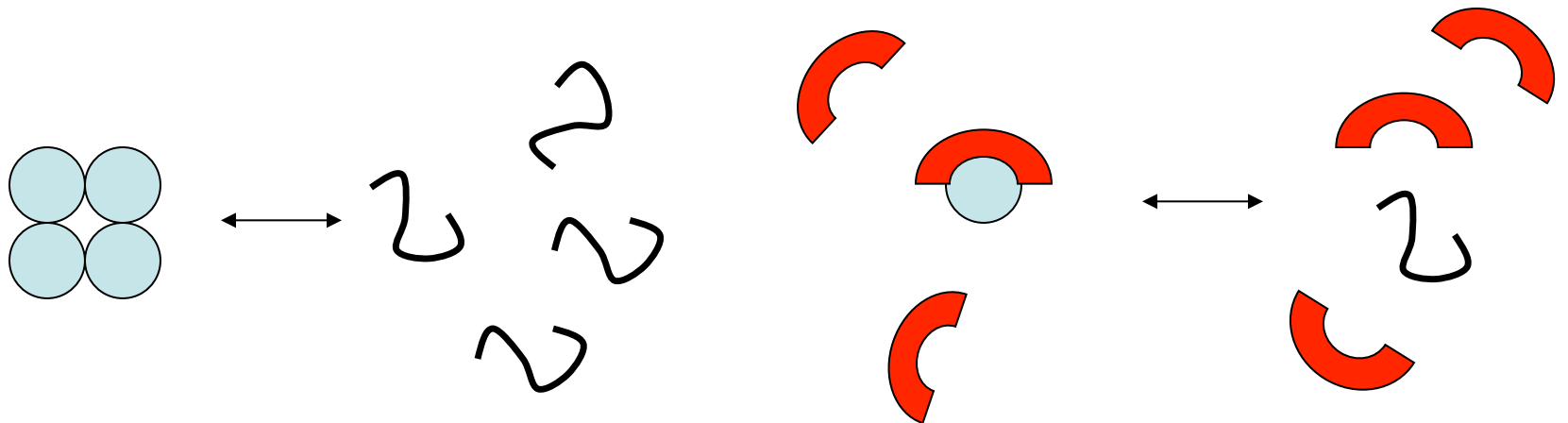
Protein-protein Interaction with a change in protonation

- The pK_a 's of groups change on complex formation
- Equilibrium position changes the proton concentration (although pH change is prevented by the buffer)
- Therefore proton concentration (pH) will affect the equilibrium position (ΔG)
- If there are no pK_a shifts in an equilibrium, there is no pH dependence in the equilibrium position.



Other Examples of Mass Action....

- Equilibrium changes concentration of molecules
- Therefore, equilibrium (stability, melting temperature) depends on the concentration
- Equilibrium involves changes in binding of ligand
- Therefore, the equilibrium (chemical stability, melting temperature) depends on concentration of the ligand

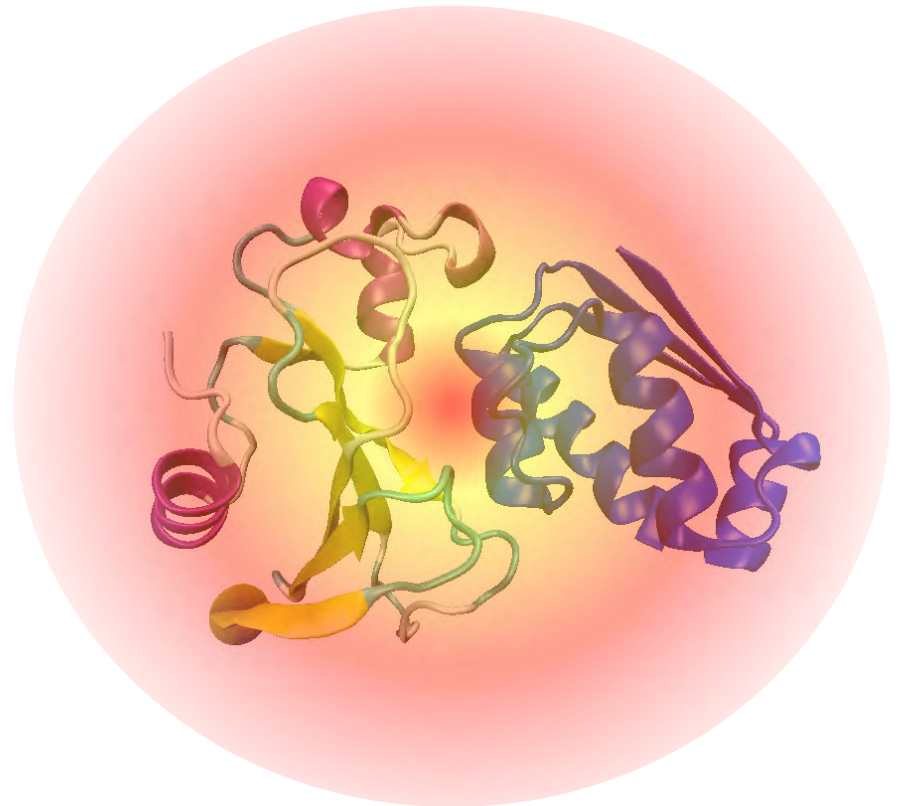
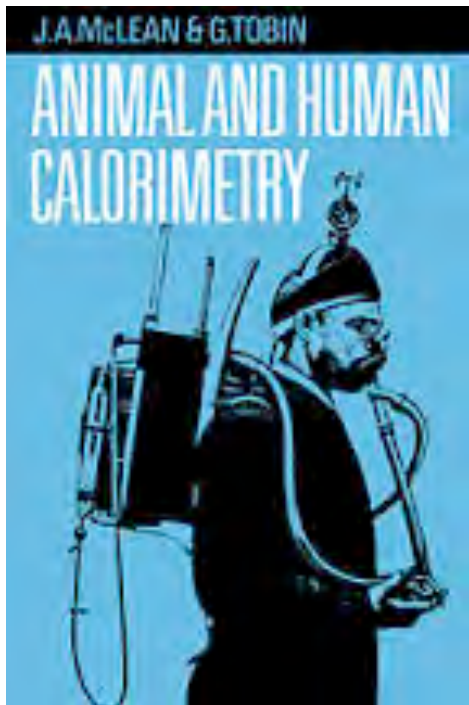


Biological Calorimetry

How to measure Enthalpy?

Under constant pressure the **heat** transferred during a process is the enthalpy (ΔH).

Calorimetry (Calor; heat Metrum; measure) is thus a technique that measures directly enthalpies of processes.



Calorimetry might be useful then?

$$\Delta G = \Delta H - T\Delta S$$

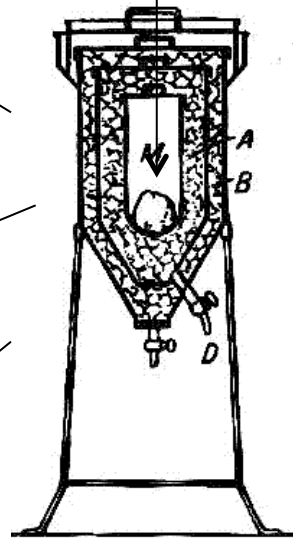
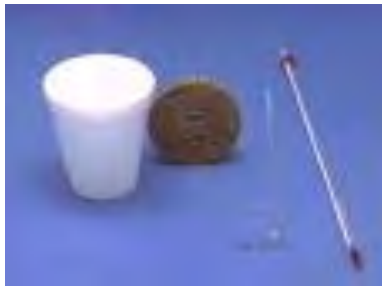
$$\Delta G = -RT \ln K_{eq}$$

- The signal in a calorimeter is the 'heat' of a process, the change in enthalpy (ΔH), which is a direct measurement of one of the driving forces of biological equilibria

Also calorimetry is

- Able to determine the value of $T\Delta S$ (if we measure ΔG (K_{eq}) and ΔH)
- A very general method. All biological equilibria have an associated enthalpy (conformational transitions, melting, binding/interaction, turnover/catalysis, etc.). No method or assay development is required.
- A non-optical, label free method. No specific group(s) or label(s) required, can use turbid suspensions or crude extracts, unusual solvents, high backgrounds of other molecules, etc. Thus applicable to many systems from the molecular to cellular level (so long as material can be put in the cell!)

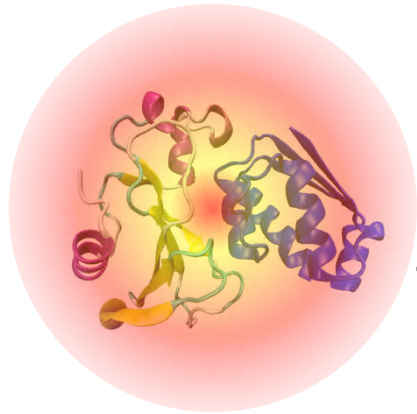
Calorimetry was one of the earliest techniques reported in the 'literature' and is now used in many areas



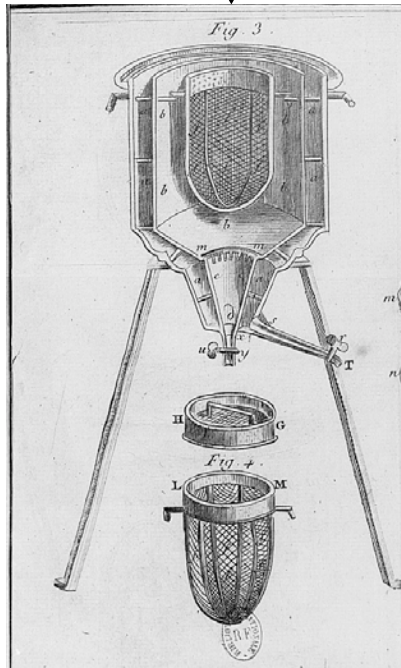
- Lavoisier's ice calorimeter 1780's
- Respiration is 'combustion'



'Biocalorimetry'



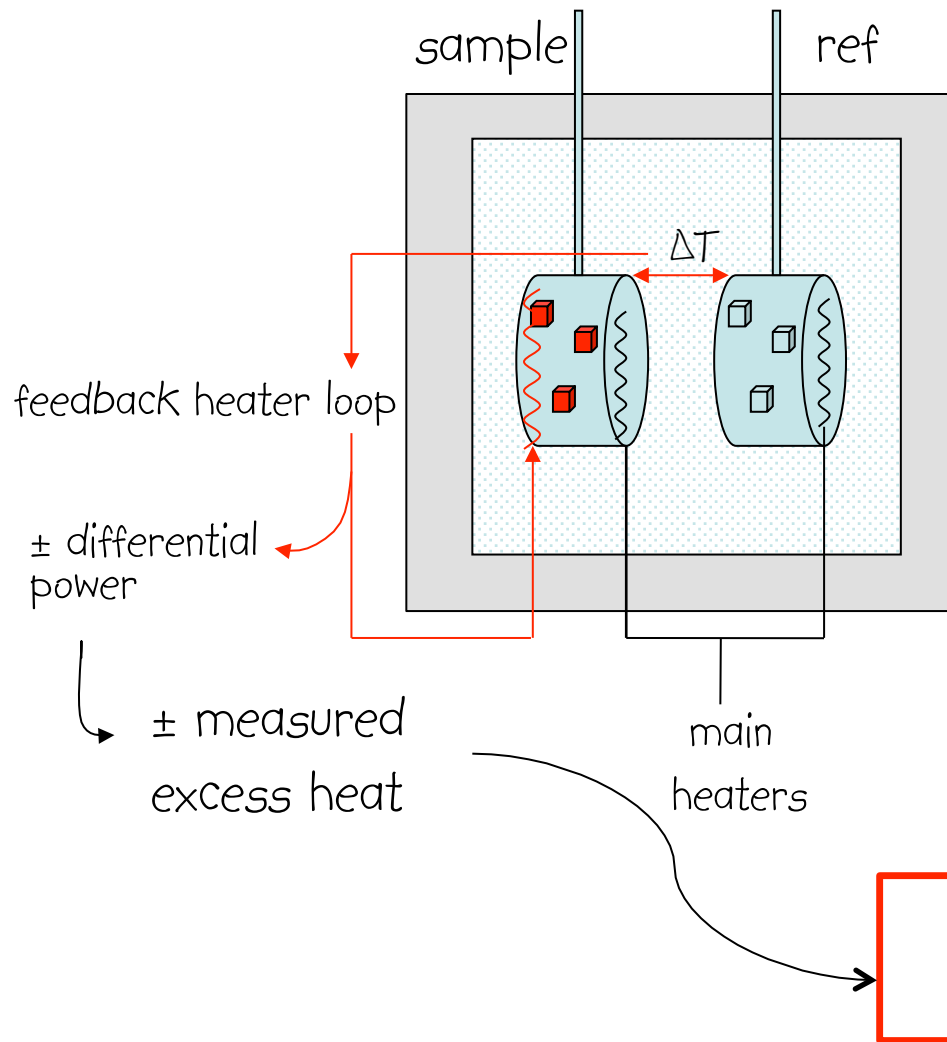
- Heat (enthalpy, ΔH) from a typical protein-protein interaction might be $-10 \text{ kcal mol}^{-1}$ *.
- Thus, 50 nmol (1ml of 50 μM solution) will give off -5×10^{-4} cal heat upon binding thus melting 6 μg ice or heating 1 ml water by $5 \times 10^{-4} \text{ }^\circ\text{C}$.
- Conventional direct heat transfer calorimetry is not possible.



Antoine-Laurent de Lavoisier
1743-1794

* K_d 50nM @ 298K $\Delta S = 0$

Power Compensation Biocalorimeters



- Difference in temperature relative to an 'identical' reference cell, measured by very precise thermopile, is kept constant by the calorimeter in a feedback loop controlling electrical heating to the sample cell.
- Increases or decreases in differential power in this circuit are directly proportional to the excess heat taken up or given off during 'reactions'

Types of Biocalorimetry

- Isothermal Titration Calorimeters (**ITC**); study interactions through the titration and mixing of two molecules at constant temperature (4-70°C : Kd; mM to nM)
- Differential Scanning Calorimeters (**DSC**); study temperature-induced transitions, melting, by increasing or decreasing temperature (T_m; 0-130°C)

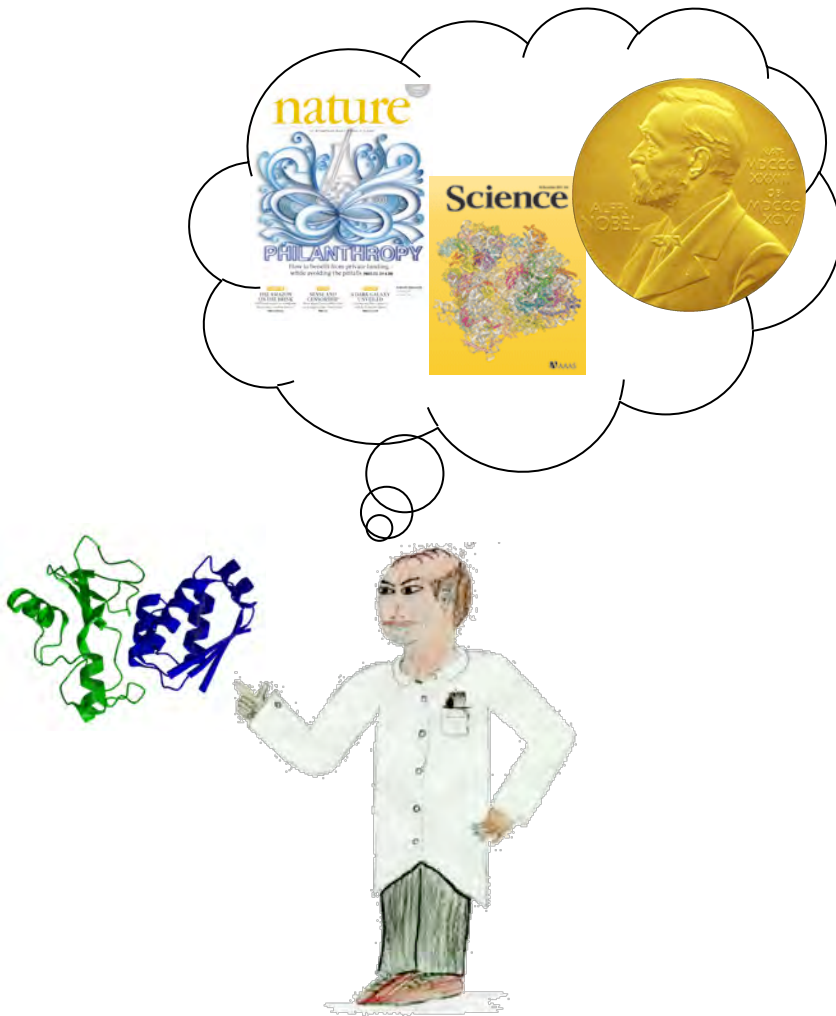


Int J Pept Protein Res. 1973;5(4):229-37.
Calorimetric investigation of ribonuclease thermal denaturation.

Anal Biochem. 1989 May 15;179(1):131-7.
Rapid measurement of binding constants and heats of binding using a new titration calorimeter.



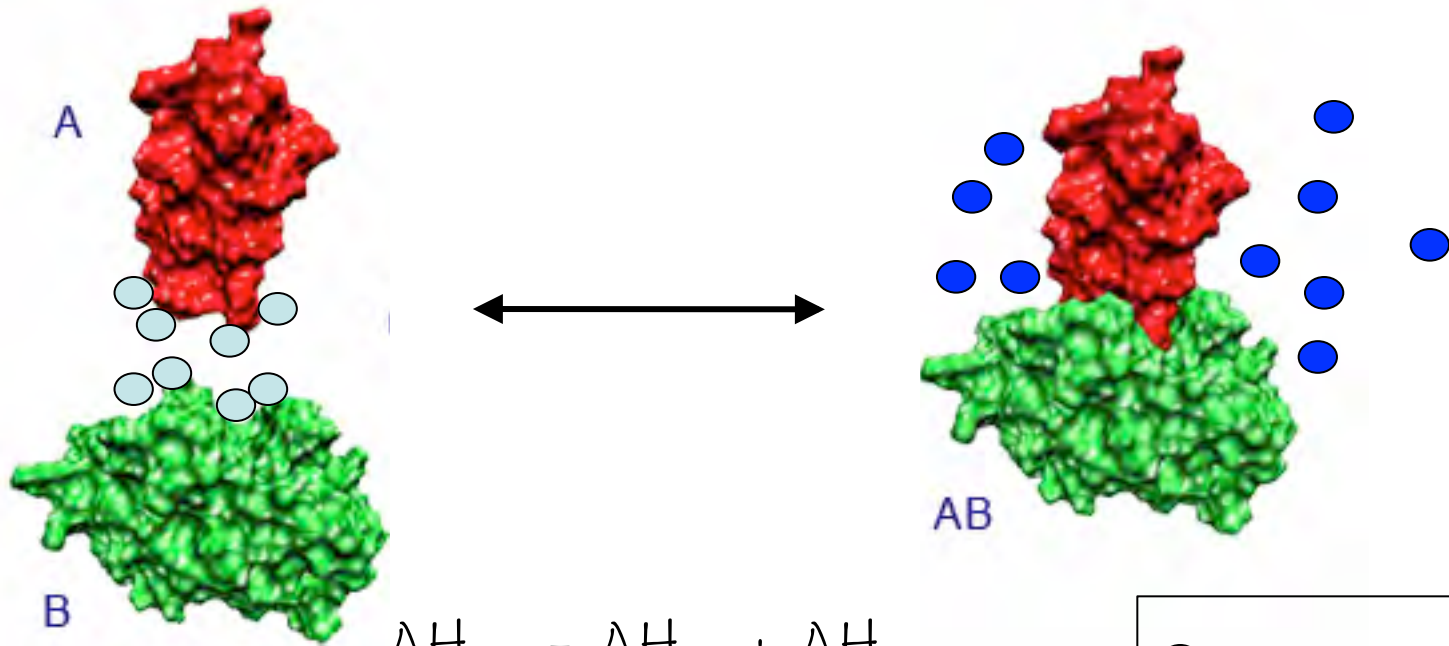
I can see interactions in my structure and mutate them so I can measure and manipulate their energies directly?!!



- Measured heat (Enthalpy) is a global non-specific probe and calorimetry measures the totality of heat effects from **all changes** in a system, i.e., specific structural interactions (observable by X-ray or NMR), but also changes in solvation, shifts in pKa's (i.e., changes in protonation), changes in dynamics, etc.
- This emphasises the combination of molecular forces driving the interaction process in solution (because these are not resolved in structures they are often forgotten or ignored).

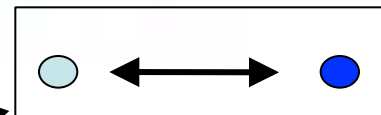
Interactions in Solution

- Interactions in solution involve changes in solvation and the observed thermodynamics are the sum of all contributions
- Proteins only interact *in vacuo* in the figures of journals!



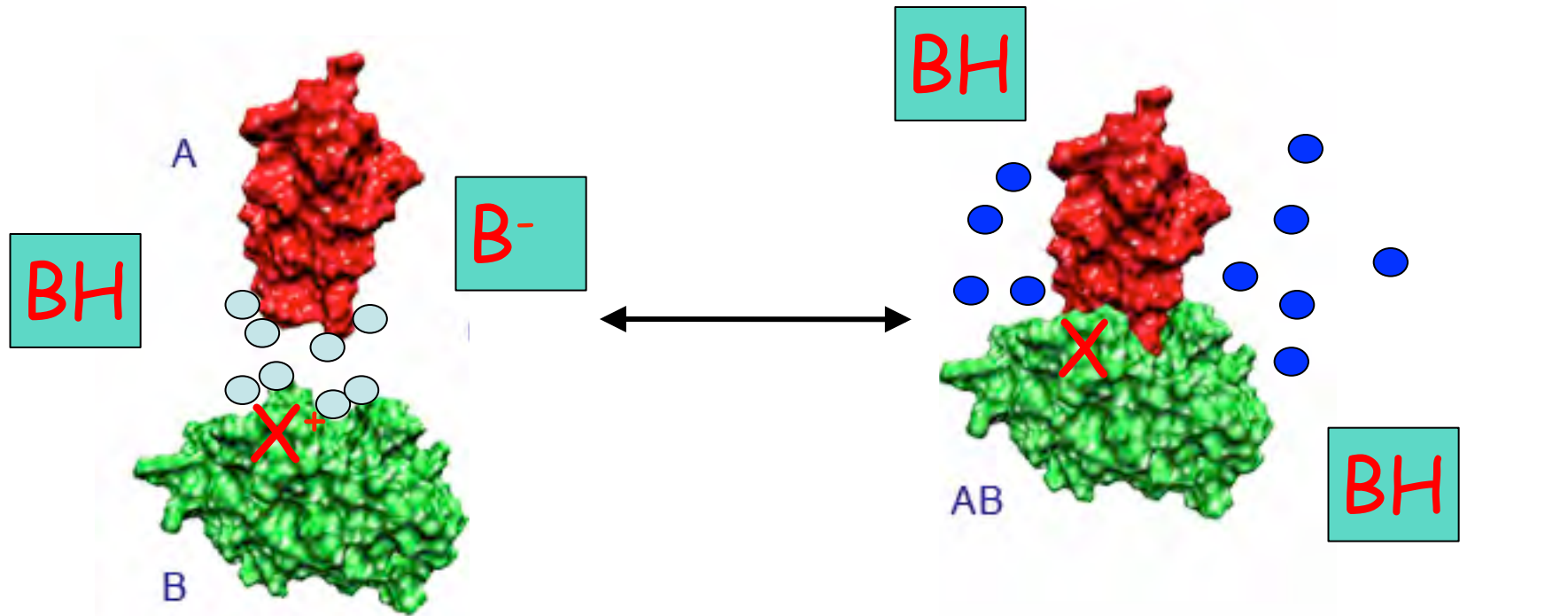
$$\Delta H_{obs} = \Delta H_{AB} + \Delta H_{solvation}$$

$$\Delta S_{obs} = \Delta S_{AB} + \Delta S_{solvation}$$



Interactions in Buffers

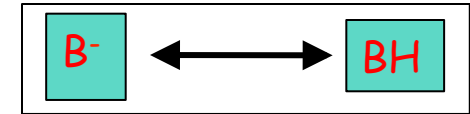
- Buffers take up or donate protons during the interaction and this also has an associated enthalpy



$$\Delta H_{obs} = \Delta H_{AB} + \Delta H_{solvation} + \Delta H_{ionisation}$$

$$\Delta S_{obs} = \Delta S_{AB} + \Delta S_{solvation} + \Delta S_{ionisation}$$

Calorimetry can measure Protonation Changes (Δv)



- Δv is the sum of all protonation changes involved in the reaction

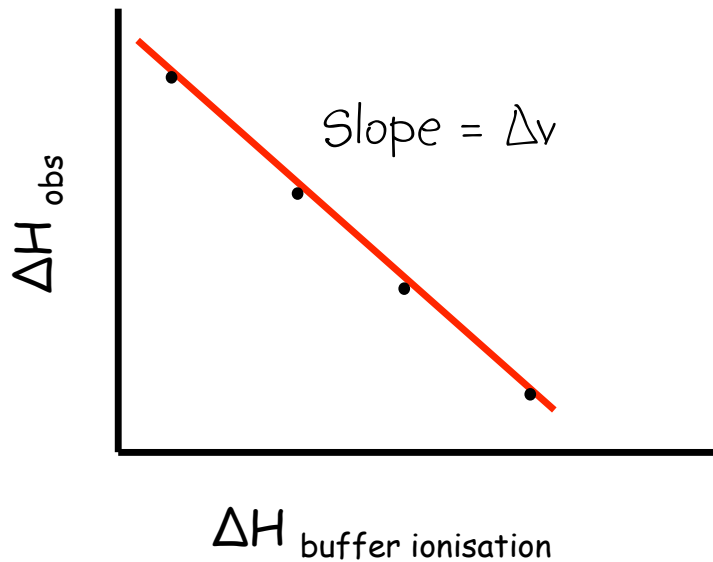


TABLE I. Enthalpy and Heat Capacity Changes for the Dissociation of Protonated Buffer Substances in 0.1 M KCl at 25° C

| Buffer substances | pK* | ΔH (kJ mol ⁻¹) | ΔC_p (J K ⁻¹ mol ⁻¹) | $\partial \Delta C_p / \partial T$ (10 ⁻³ J K ⁻² mol ⁻¹) |
|----------------------|-------|---------------------------------------|--|---|
| Acetate | 4.62 | 0.49 ± 0.02 | -128 ± 2 | — |
| MES | 6.07 | 15.53 ± 0.03 | 16 ± 2 | — |
| Cacodylate | 6.14 | -1.96 ± 0.02 | -78 ± 2 | — |
| Glycerol 2-phosphate | 6.26 | -0.72 ± 0.02 | -179 ± 2 | 0.79 ± 0.39 |
| PIPES | 6.71 | 11.45 ± 0.04 | 19 ± 4 | — |
| ACES | 6.75 | 31.41 ± 0.05 | -27 ± 4 | — |
| Phosphate | 6.81 | 5.12 ± 0.03 | -187 ± 3 | 2.01 ± 0.22 |
| BES | 7.06 | 25.17 ± 0.07 | 2 ± 5 | — |
| MOPS | 7.09 | 21.82 ± 0.03 | 39 ± 3 | — |
| Imidazole | 7.09 | 36.59 ± 0.06 | -16 ± 5 | — |
| TES | 7.42 | 32.74 ± 0.03 | -33 ± 3 | — |
| HEPES | 7.45 | 21.01 ± 0.07 | 49 ± 5 | — |
| EPPS | 7.87 | 21.55 ± 0.05 | 56 ± 4 | — |
| Triethanolamine | 7.88 | 33.59 ± 0.04 | 48 ± 3 | — |
| Tricine | 8.00 | 31.97 ± 0.05 | -45 ± 4 | — |
| Bicine | 8.22 | 27.05 ± 0.05 | 2 ± 4 | — |
| TAPS | 8.38 | 41.49 ± 0.06 | 23 ± 5 | — |
| CAPS | 10.39 | 48.54 ± 0.07 | 29 ± 6 | — |

*Standard error is within ±0.01.

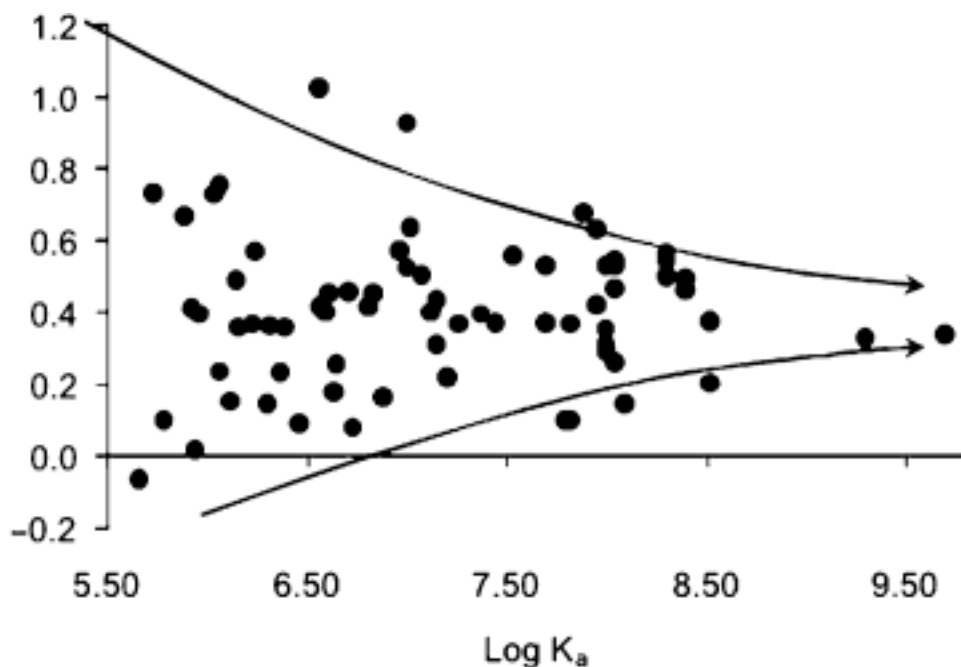
Fukada and Takahashi (1998)

PROTEINS: Structure, Function, and Genetics

33,159

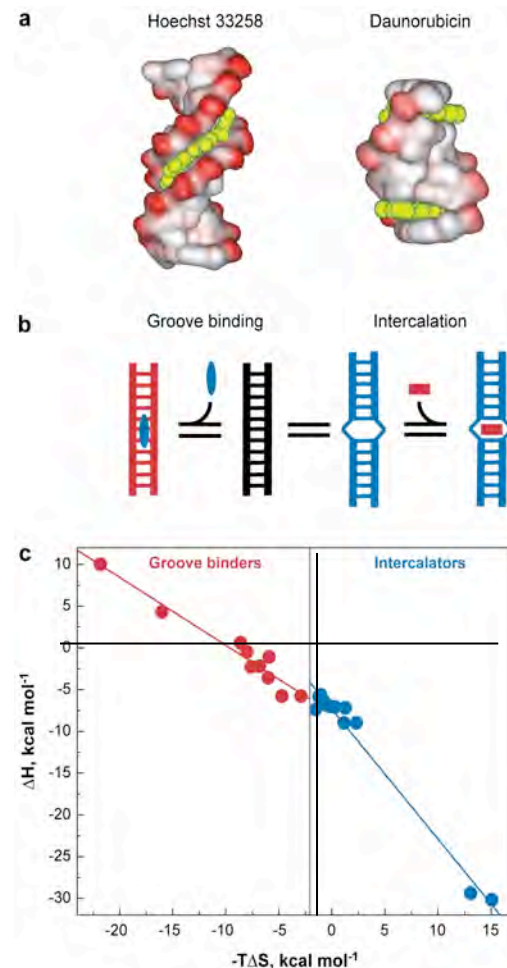
Interpretation of thermodynamics may be (currently) impossible, but there may be qualitative trends.....

Small molecule drug development



Allophenylnorstatine inhibitors of plasmepsin II
 Ruben et al. *Chemical Biology & Drug Design* 2006, 67, 2-4

DNA Binding mode



Practical Biocalorimetry

Types of Biocalorimeter

- Isothermal Titration Calorimeters (**ITC**); study interactions through the titration and mixing of two molecules at constant temperature ($4-70^{\circ}\text{C}$: K_d ; mM to nM)
- Differential Scanning Calorimeters (**DSC**); study temperature-induced transitions, “melting”, by increasing or decreasing temperature (T_m ; $0-130^{\circ}\text{C}$)

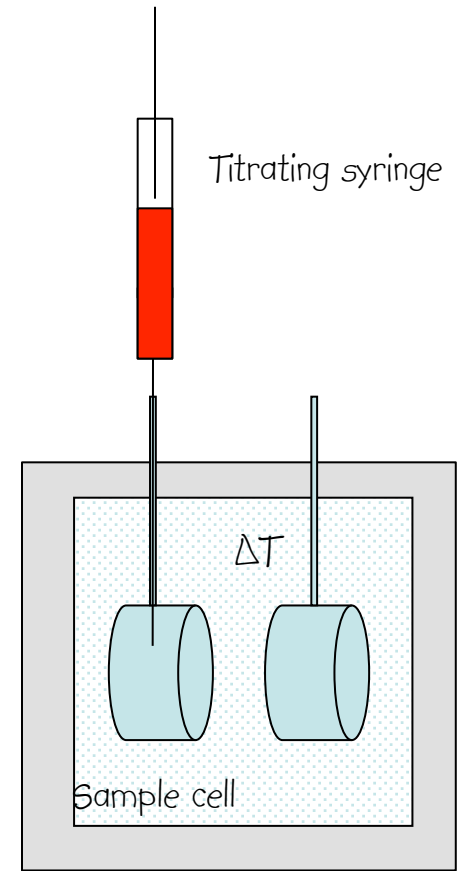


ITC

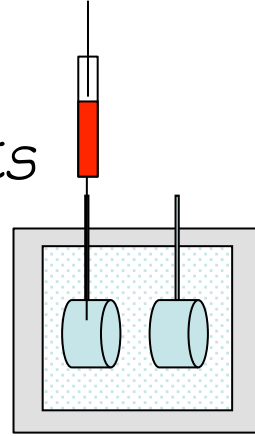
Typical ITC Experiment

- iTC 200 instrument (active cell volume 200 μL)
- 350 μL of 'target' 10 - 50 μM * loading cell (275 μL consumed)
- 70 μL of 'ligand' 50 - 500 μM * loading syringe (55 μL consumed)
- ITC is a 'sample hungry' technique so evidence of binding from other techniques and/or the literature is useful.
- Heat is a non-specific probe and calorimetry measures the totality of heat effects from all events Controls needed!

* Typical starting concentrations. Actual concentrations depend on ΔH (signal amplitude), K_d of binding and the type of experiment



Sources of signal and controls for ITC experiments



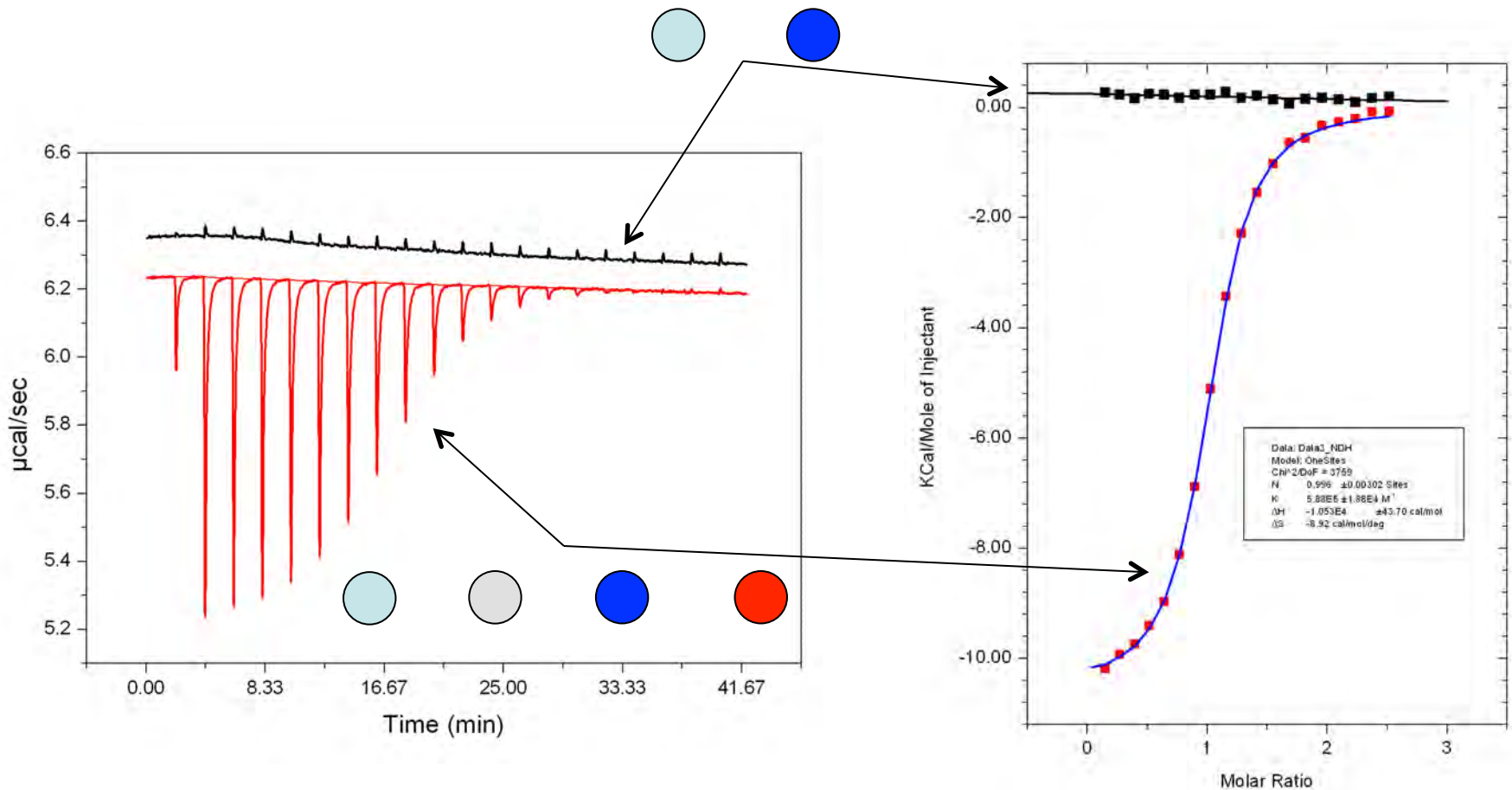
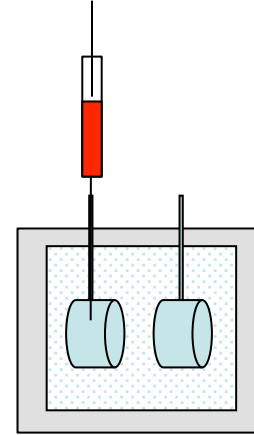
- ① Buffer injected into buffer. The background heat of 'injection'. Mechanical disturbances and temperature gradients. ●
- ② Buffer injected into protein. Background heat plus the heat of dilution of the protein in the cell. Usually = background as dilution factor in the cell is small. ● + ●
- ③ Ligand injected into buffer. Background heat plus heat of dilution of the ligand. Significant! Dilution of ligand is large ● + ●
- ④ Ligand injected into protein. Background plus heat of dilution of protein and ligand with heat of binding. ● + ● + ● + ●

Since ● is ~ 0 then 4 should be corrected with 3 to give ●

Example: ITC measurement and separate control

600uM peptide into 45 uM protein or into buffer

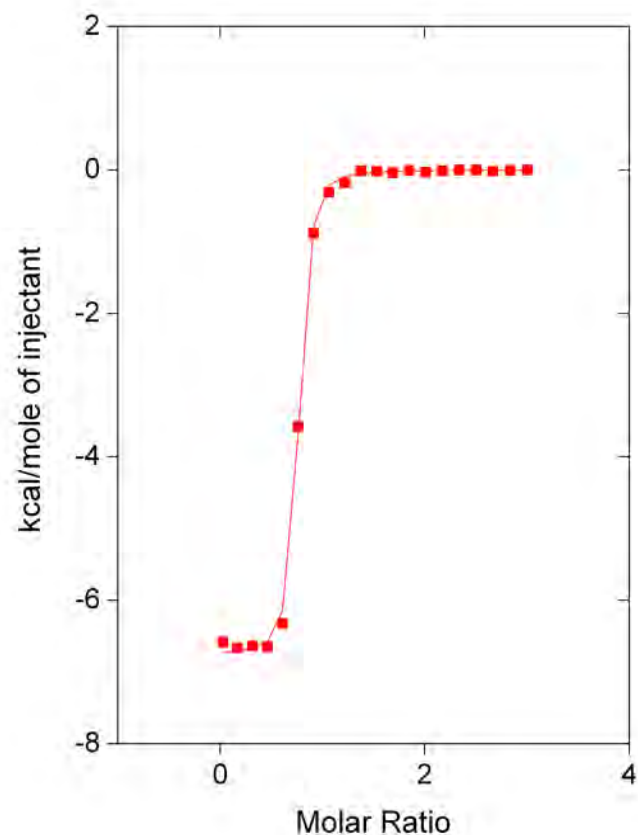
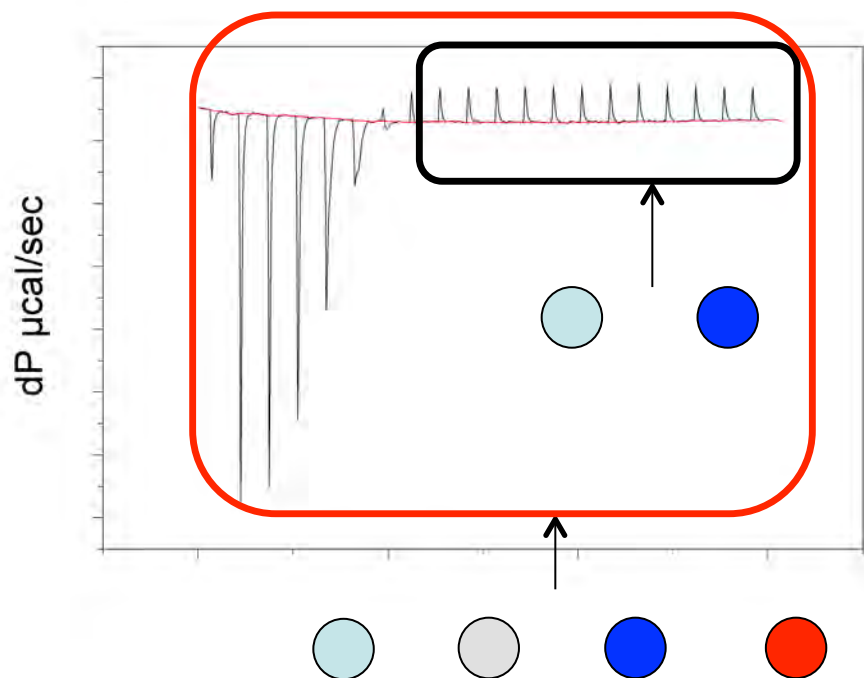
Integrate heat from each injection. Subtract control and fit.



Single experiment measurement

500 μ M ligand into 30 μ M protein

Integrate heat from each injection. Subtract 'control' heats observed at 'saturation' toward the end of the titration



Data Fitting in Origin or PEAQ*

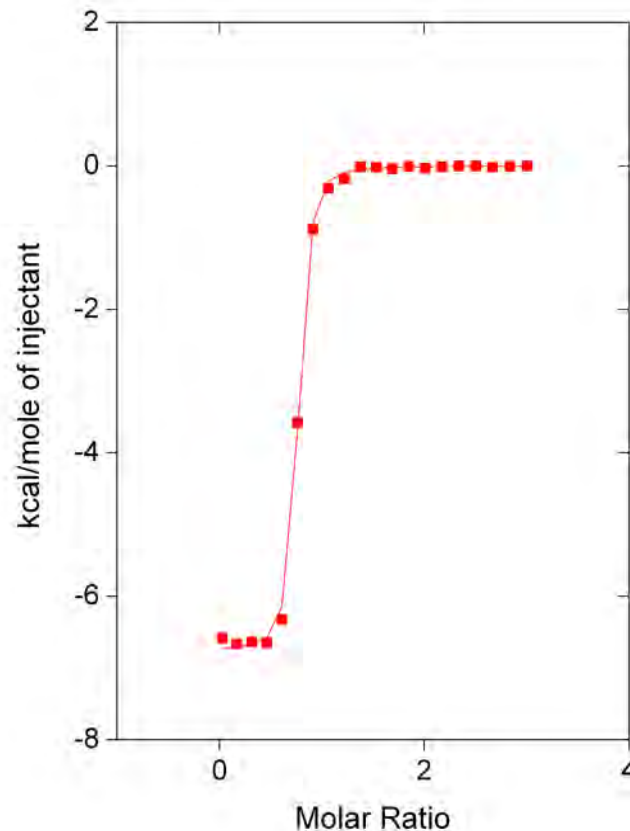
Simple model of N site(s) with identical affinity and enthalpy

| | | |
|------------|--------|------------------------|
| N | 0.703 | 0.00294 |
| Ka | 2.98E6 | 3.61E5 M ⁻¹ |
| Kd | 330 nM | +/- 40 nM |
| ΔH | -6759 | 41.60 cal/mol |

Calculated values

| | |
|------------|---------------|
| ΔG | 9 kcal/mol |
| ΔS | 7 cal/mol/deg |

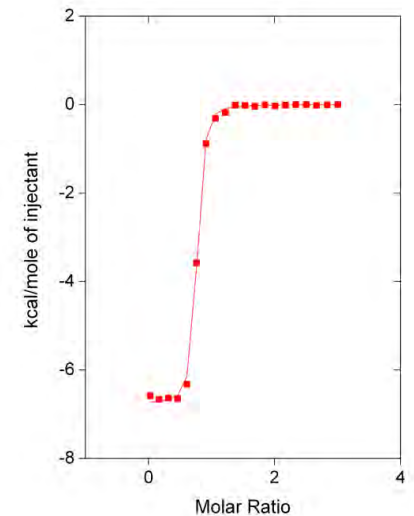
- Disregard the errors and the suggested precision
- Plot in conventional format and refit
- Come to lecture 14



*Software provided with Microcal calorimeters

The 'meaning' of $N = 0.7$

- 'N' is the 'number of sites', or 'stoichiometry' of the interaction and is well determined for this tight binding
- It assumes the concentration of the macromolecule in the cell and the ligand in the syringe are measured correctly and both are 100 % 'pure' and 100% 'native' or 'binding competent'.
- If not, 'N' can be viewed as an indicator of 'active site concentration' or an additional fitting parameter that allows concentration to float during fitting and thereby allow the fit to go through the points.
- In practice $N = 1 \pm 0.2$ is equivalent to $N \sim 1$. $N = 0.5$ and $N = 2$ etc., also have some potential physical meaning (e.g., dimer with one site etc, etc)

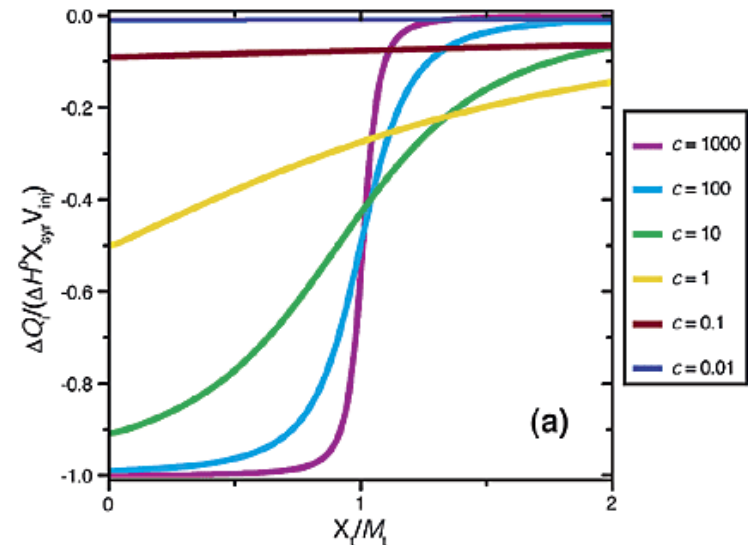
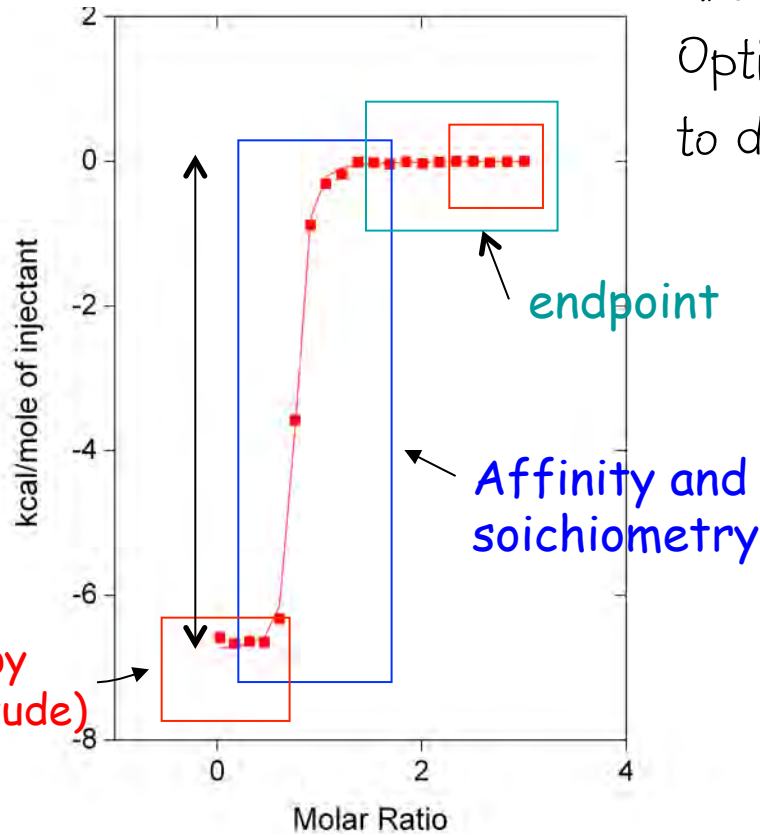


NLLS Fitting values

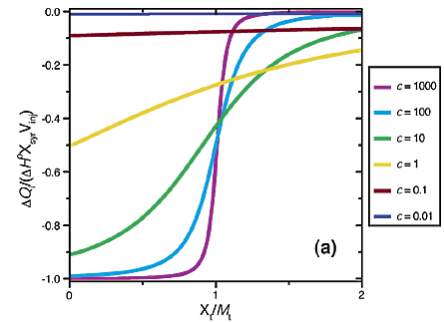
| | |
|------------|----------|
| N | 0.703 |
| K_a | $2.98E6$ |
| K_d | 330 nM |
| ΔH | -6759 |

Information content and optimising ITC experiments

Derivative plots can be described by parameter known as 'C value' = $[\text{protein cell}] / K_d$
Optimal range for ITC $10 < c < 500$ when trying to determine K_d , N and ΔH in one experiment



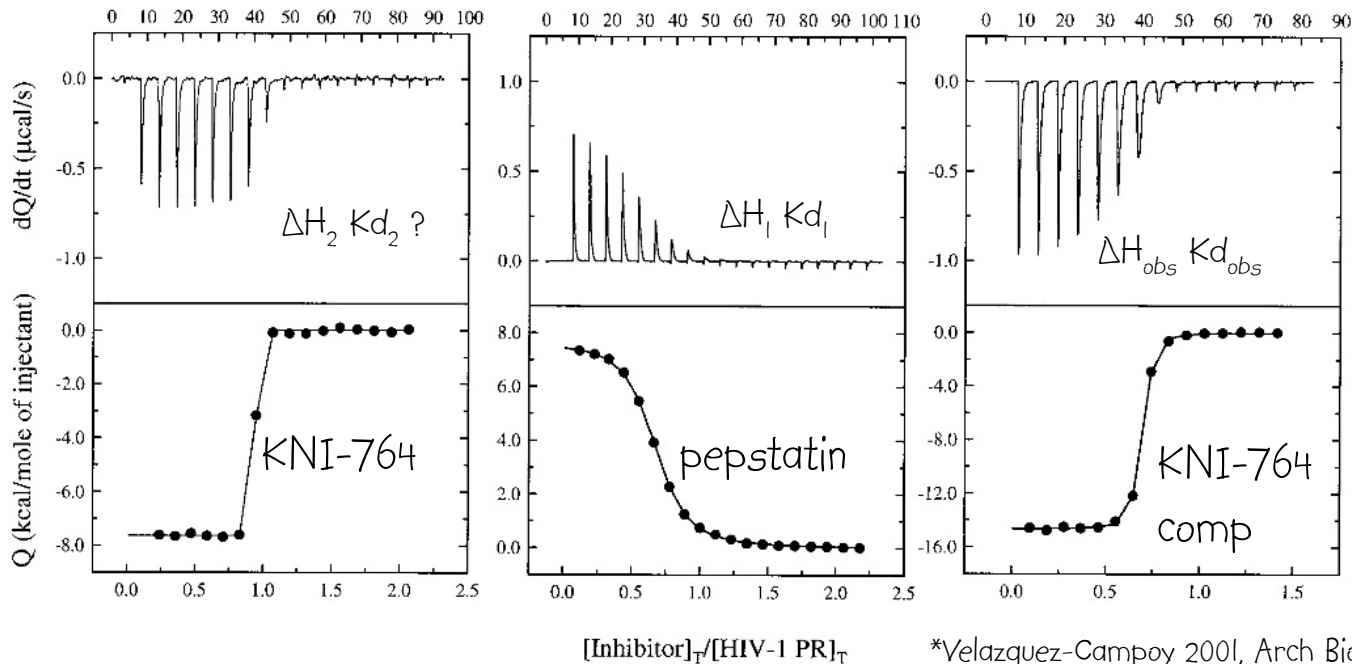
High affinity binding



- As binding becomes tighter c-value ($[cell]/K_d$) becomes too large unless protein concentration is decreased.
- However, (cell concentration $\times \Delta H$) determines the total heat in the experiment and with current detection limits [protein] must be $> 5-10 \mu M$ unless ΔH is very large.
- This limits simple ITC measurements to K_d 's in the 100's nM range
- However, since ITC is a non-optical and probe free method it is perfect for displacement (competition) binding methods which can extend the K_d range down to pM and beyond.

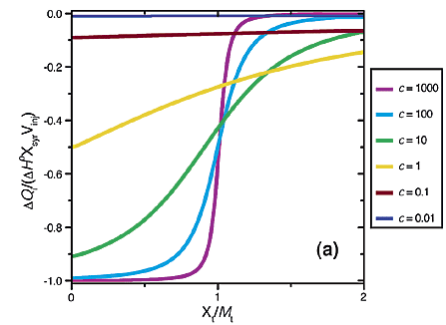
Competition binding experiments

- Measure first a 'weaker' binding ligand (pepstatin). Displace this ligand in a second titration of the endpoint complex with the tight binder (KNI-764)
- ΔH_{obs} competition $\sim = \Delta H_2 + (-\Delta H_1)$
- Kd_{obs} competition $\sim = Kd_1 \times Kd_2$ KNI-764 inhibitor binding HIV-1 protease Kd 32 pM
- Note. Both affinity and enthalpy change making this an ideal tool for screening small molecules, for example, that compete for a target binding site



*Velazquez-Campoy 2001, Arch Biochem Biophys, 390, p169
 Velazquez-Campoy 2006 Nat Protoc, 1(1):186-91

Weak binding



- As binding becomes weaker, the c -value ($[cell]/K_d$) becomes small unless $[protein]$ is increased. K_d , ΔH and n are all less well constrained and it may be difficult to define titration endpoints.
- The background heat of dilution of the ligand (which may be required at 10's mM concentration in the syringe) is critical and so separate titrations of ligand into buffer are required.
- Many small molecule fragments require 5-10% DMSO to dissolve at mM concentrations. This exacerbates the background heat of dilution if DMSO is not carefully matched in cell and syringe.
- If material and solubility are not limiting, very weak binding or non-specific interactions can be quantified

- Turnball et al., J Am Chem Soc. 2003, 125(48):14859-66
- Tellinghuisen. Anal Biochem. 2008; 373(2):395-7

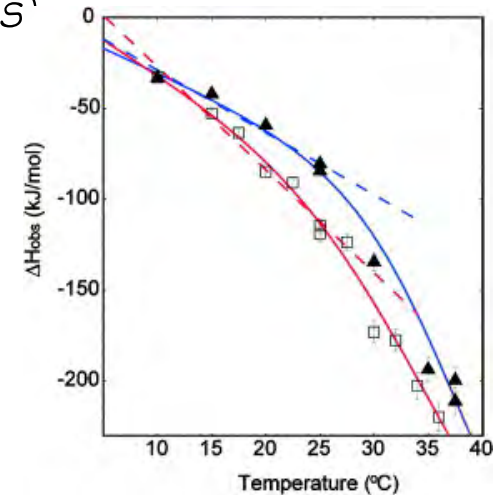
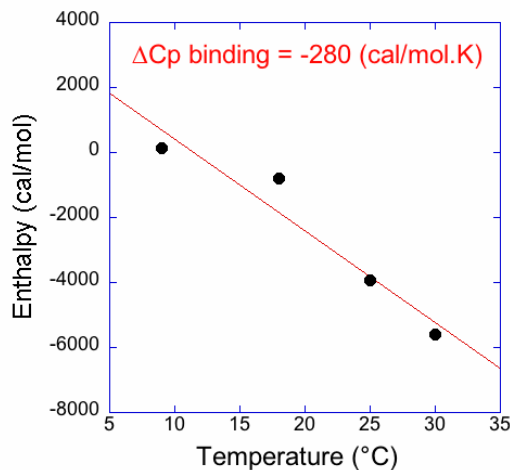
No just K_d 's and ΔH

ΔC_p of binding

$$\Delta H = \int_0^T \Delta C_p \cdot \partial T$$

$$\Delta C_p = \frac{\partial \Delta H}{\partial T}$$

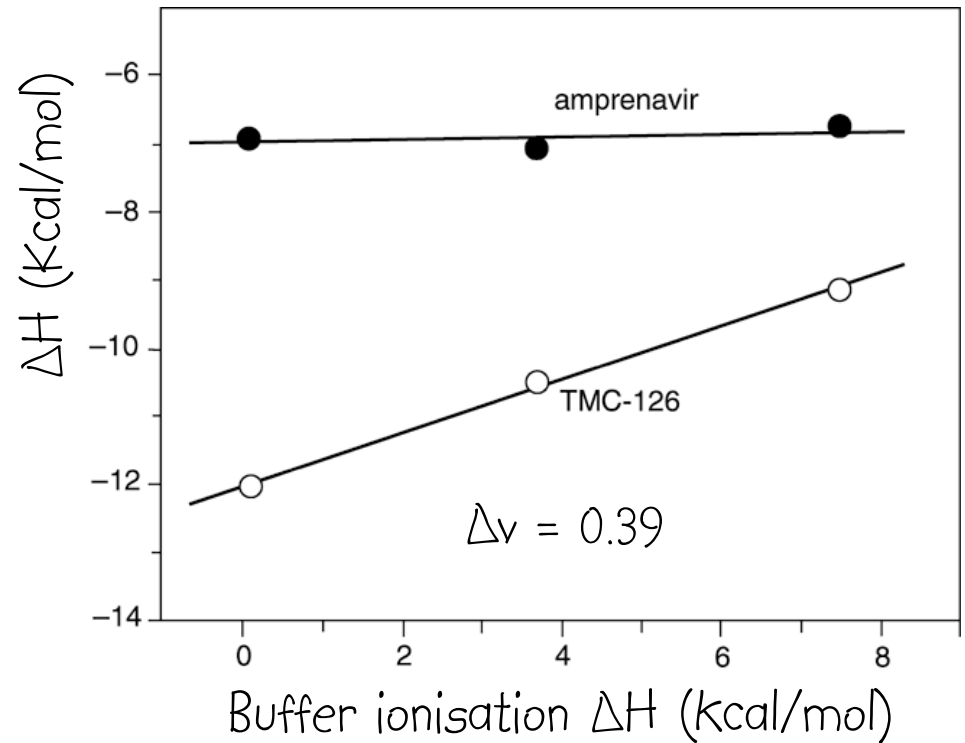
- ΔC_p can be obtained from the variation of ΔH with temperature.
- Rigid body type interactions have a constant typically -ve ΔC_p^{bind}
- Unlike ΔH , ΔC_p^{bind} can be deconvolved to indicate properties of the ligand binding footprint, (ΔASA , non polar, polar etc.)
- Non-constant or large values for ΔC_p^{bind} indicate coupled events such as ligand induced structuring (NDP folding and binding) or conformational change (domain movements)



- Molecular recognition via coupled folding and binding in a TPR domain. Cliff MJ et al, *J Mol Biol.* 2005 346:717-32
- Vega et al., *Biochim Biophys Acta.* 2016;1860(5):868-878

Δv (protonation change) of binding

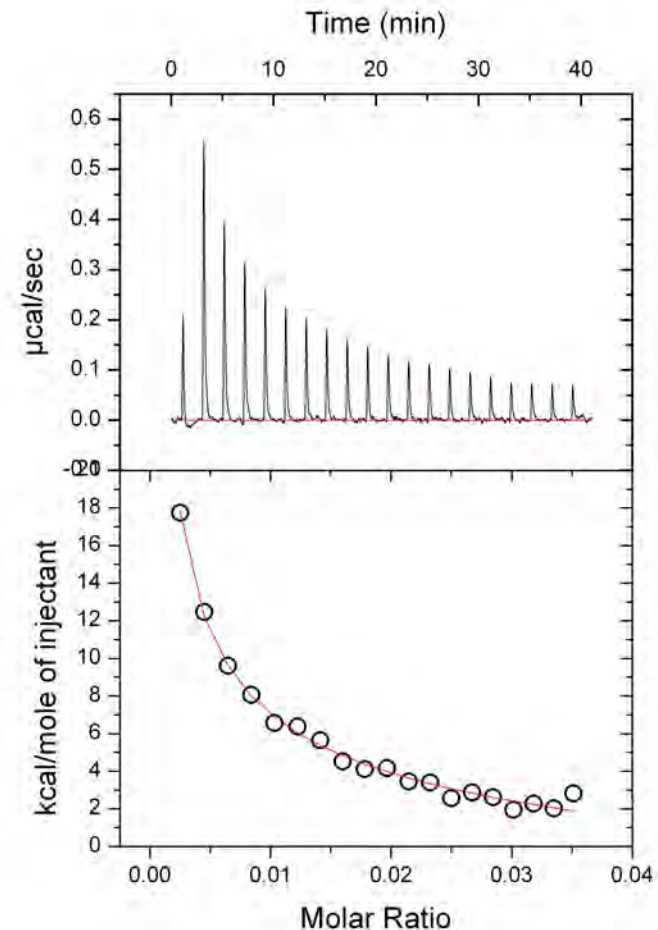
- Titrations in different buffers indicates the net flux of protons.
- Differences in Δv on mutation of protein or between ligands could indicate different binding sites or changes to binding mode.



Overcoming drug resistance in HIV-1 chemotherapy: the binding thermodynamics of Amprenavir and TMC-126 to wild-type and drug-resistant mutants of the HIV-1 protease. Ohtaka et al., Protein Sci. 2002 8:1908-16.

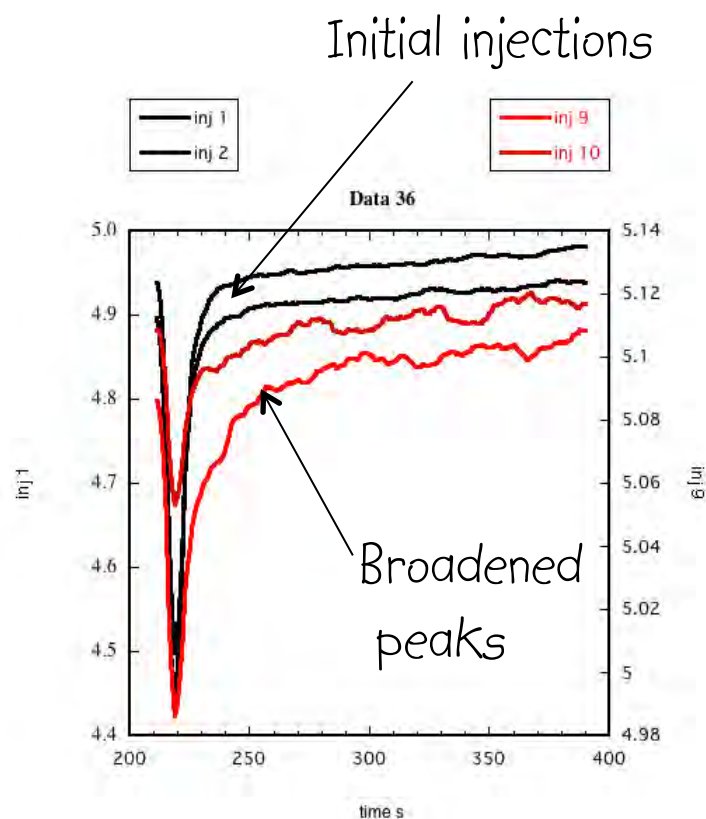
Oligomerisation processes (self association)

- Simply titrate (dilute) protein into buffer (fully recoverable)
- Fit, in this case, to dimer dissociation model
- Suitable for μM k_d 's
- More complex dissociations possible, hetro-dimer, tetramer etc



Binding kinetics from standard ITC binding experiments?

- ITC measures the rate of heat production (differential power) and so potentially has kinetic information.
- When k_{obs} ($k_{on} + k_{off}$) becomes slower than the instrumental response the ITC peaks become broader
- Normally this occurs as the system 'saturates' and k_{on} becomes slow due to limited free sites
- KinITC software*



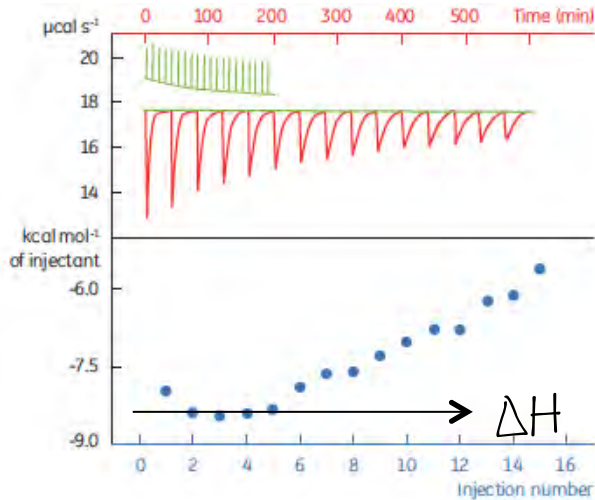
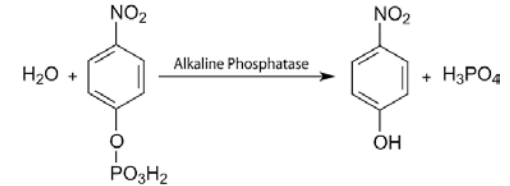
* <https://www.affinimeter.com/site/kinitc-2/>

Enzyme / Reaction Kinetics by ITC

A universal* method ?

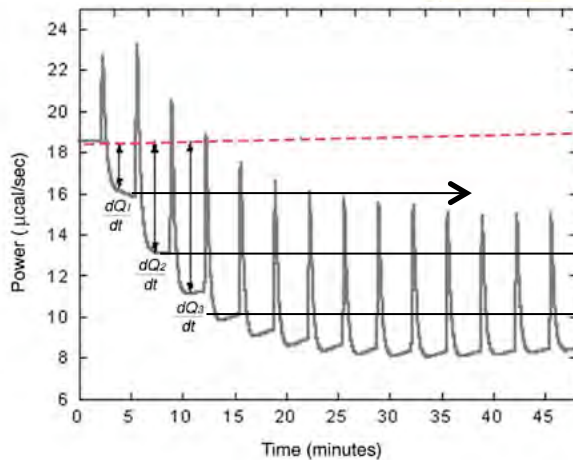
Non-optical, no assay development or coupled reactions

Hydrolysis of pNPP by PPI- γ phosphatase



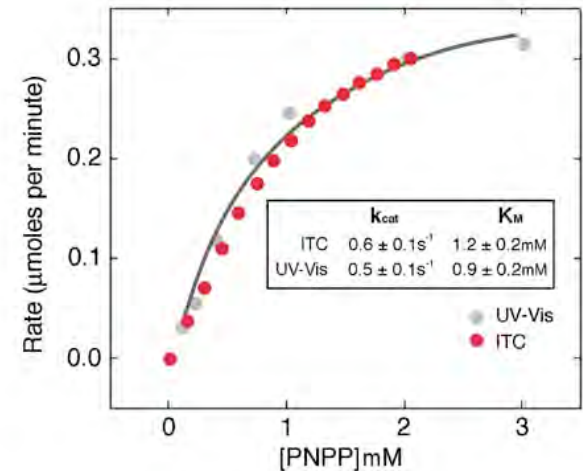
μM enzyme long duration complete turnover

$$\text{Rate} = (1/\Delta H) * dQ/dt$$



nM enzyme short duration initial rate of turnover based on ΔH

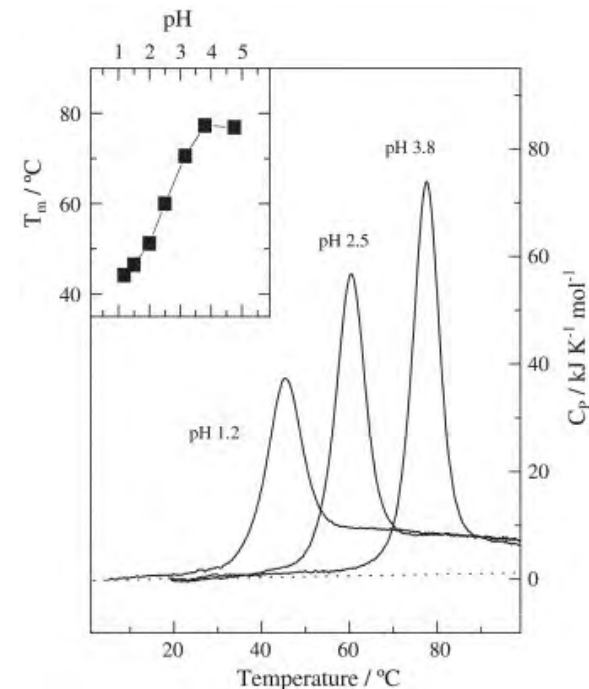
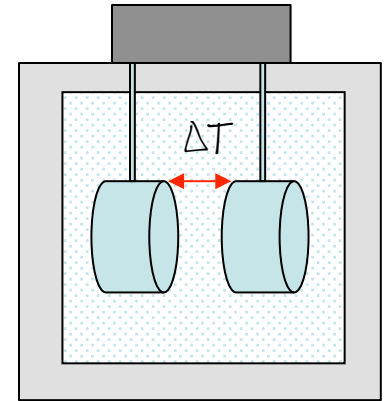
dQ/dt



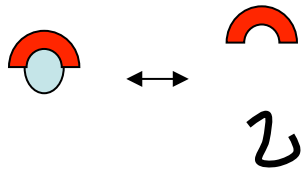
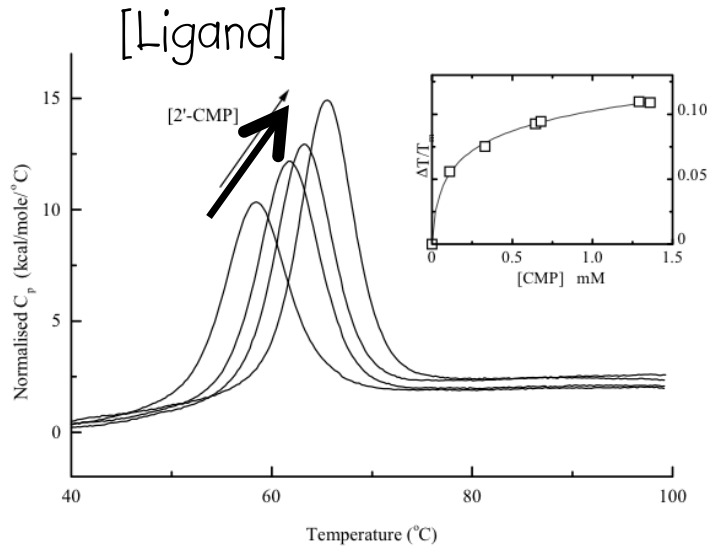
DSC

Typical DSC Experiment

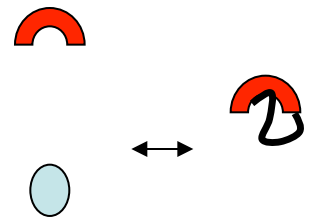
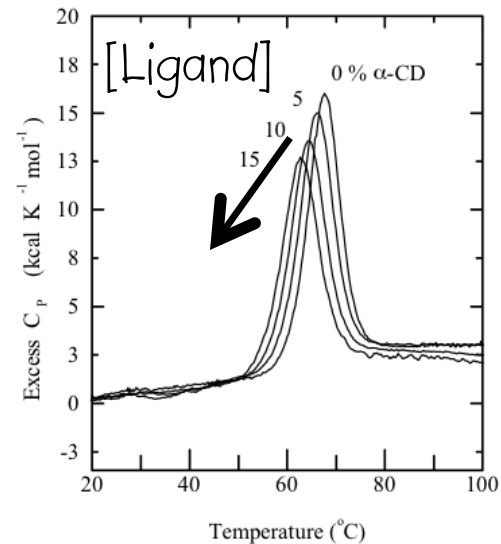
- CAP DSC instrument; robotic loading and operation
- Active cell volume $\sim 130 \mu\text{L}$
- 360 μL of a 0.2 – 1 mg ml^{-1} macromolecule
- Temperature range (5–125°C) and scan rate (0.1–2.5 °C/min) variable
- Gives thermal stability (T_m) and enthalpy under different conditions or for mutants



Ligand induced stabilisation or destabilisation

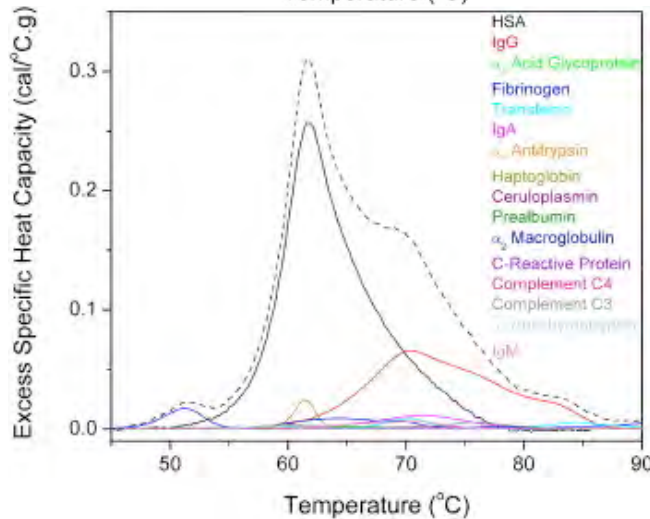
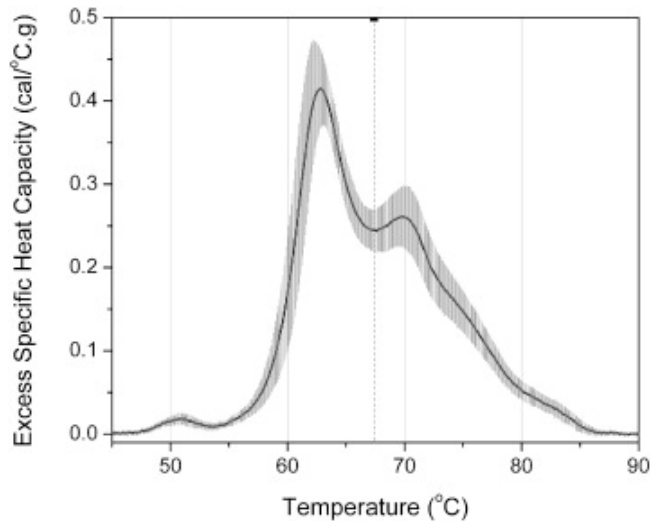


- Mass action increases or decreases the stability as ligands bind to native or denatured state

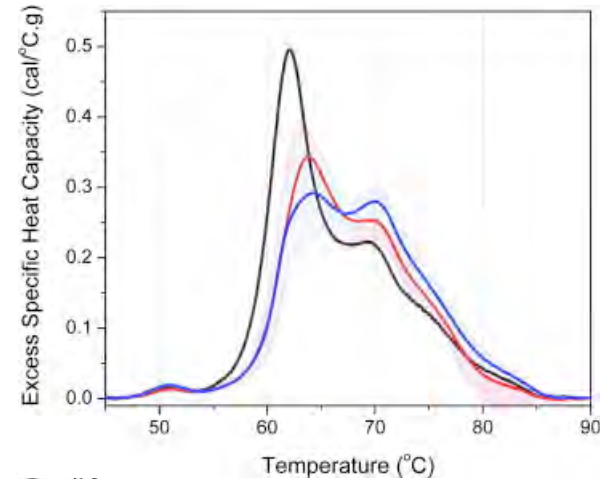


DSC and Complex mixtures

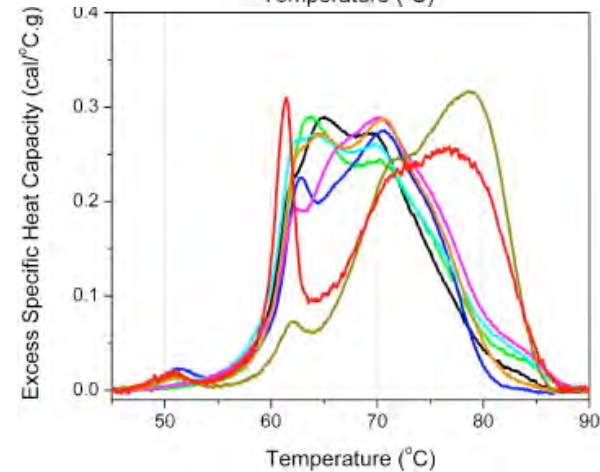
Normal human plasma n=15



Plasma from patients with a range of cancers and diseases



Blue invasive cervical cancer



Dark yellow lyme disease

Summary

- Calorimetry is a label free and very general method for any biomolecular equilibrium not just proteins!
- ITC mainly Binding (K_d , n , ΔH , ΔS , ΔC_p , $\Delta \text{protonation}$)
- DSC mainly thermal stability (T_m , ΔH , ΔC_p , ΔH_{vH} and mechanism, $\Delta \text{protonation}$)
- Changes in T_m indicate ligand binding, mechanism
- Many 'other' uses for ITC and DSC



Enthalpy and Entropy changes are difficult to interpret in isolation since they include contributions from many processes in the system....

Changes in conformation

Changes in dynamics

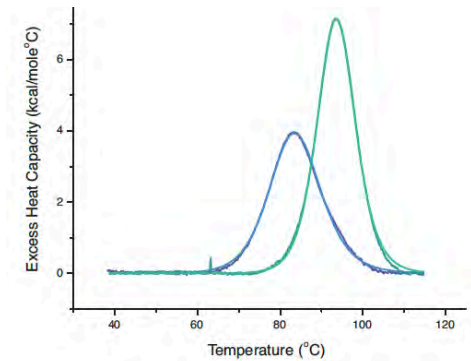
Changes in charge

Contributions of water (Proteins may be big but there is a lot of water!)

Some examples of LMB work employing some calorimetry

Ubiquitin Ser65 phosphorylation affects ubiquitin structure, chain assembly and hydrolysis.

Wauer T, Swatek KN, Wagstaff JL, Gladkova C, Pruneda JN, Michel MA, Gersch M, **Johnson CM**, Freund SM, Komander D. *EMBO J.* 2015 Feb 3;34(3):307-25

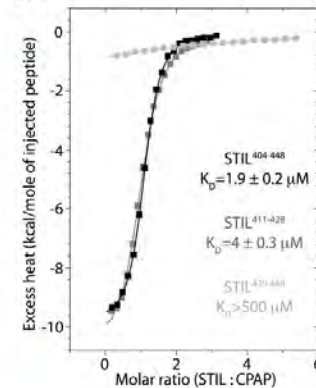
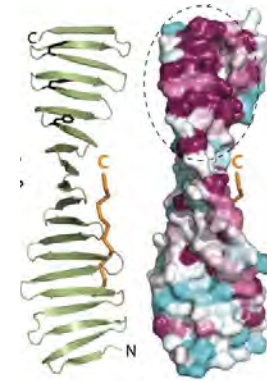


Bacterial actin MreB forms antiparallel double filaments.

van den Ent F, Izoré T, Bharat TA, **Johnson CM**, Löwe J. *Elife.* 2014 May 2;3:e02634

Crystal structures of the CPAP/STIL complex reveal its role in centriole assembly and human microcephaly.

Cottee MA, Muschalik N, Wong YL, **Johnson CM**, Johnson S, Andreeva A, Oegema K, Lea SM, Raff JW, van **Breugel M.** *Elife.* 2013 Sep 17;2:e01071



The hepatitis B virus preS1 domain hijacks host trafficking proteins by motif mimicry.

Jürgens MC, Vörös J, Rautureau GJ, Shepherd DA, Pye VE, Muldoon J, **Johnson CM**, Ashcroft AE, Freund SM, Ferguson N. *Nat Chem Biol.* 2013 Sep;9(9):540-7

Bacterial actin MreB assembles in complex with cell shape protein RodZ.

van den Ent F, **Johnson CM**, Persons L, de Boer P, Löwe J. *EMBO J.* 2010 Mar 17;29(6):1081-90

