# Bayesian analysis of isothermal titration calorimetry data

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[JDC: Author order has not been finalized yet, but I wanted to make sure I listed everybody who made substantial contributions to this work.]

Isothermal titration calorimetry (ITC) is the only experimental technique able to reliably measure both the free energies of macromolecule-ligand association as well as its decomposition into enthalpic and entropic contributions in a single experiment. Due to the way in which the thermodynamic parameters are extracted from the data, errors in the free energy, enthalpy, and entropy of binding can differ in magnitude, and are generally highly correlated. Additionally, the use of multiple measurements to improve statistics, measure affinities of stronger ligands by competing off weaker ones, and separate proton uptake effects from binding by titrating in multiple buffers can complicate the propagation of these uncertainties to the physical quantities of interest. Here, we present a simple Bayesian framework for computing the full posterior distribution of all thermodynamic parameters and other quantities of interest from one or more ITC experiments, allowing their uncertainties and correlations to be quantitatively assessed. Use of this Bayesian approach leads to uncertainties that can be orders of magnitude larger than those typically reported when data are re-analyzed, but which more accurately represent the true variability in experiments from laboratory to laboratory. The framework is general and flexible, and further allows the modeling of new experiments in a way that aids the experimenter in selecting experimental parameters that will maximize the expected information gain. A Python implementation suitable for use with most popular calorimeter data formats is freely available online at http://www.simtk.org/home/bayesian-itc.

# I. INTRODUCTION

Isothermal titration calorimetry (ITC) [1] has proven to be powerful technique for measuring the free energy of association of two soluble species, most notably finding application in providing a quantitative measure of the binding affinity of small molecule ligands to biological macromolecules such as proteins and RNA [2, 3]. [JDC: Add some 16 references to reviews and applications to biomoleculecular interactions, such as those by Ernesto Friere.] From a single experiment, estimates of the free energy ( $\Delta G$ ), enthalpy  $(\Delta H)$ , and entropy  $(\Delta S)$  of binding can be simultaneously extracted, providing both a direct assessment of binding affinity as well as insight into the nature of the thermodynamics of binding. Beyond this, ITC also allows the study of more complex reactions, such as competitive binding reactions [2, 4], binding events in the presence of changes in protonation [5, 6] or tautomeric [7] states, and in certain cases, even kinetics of binding [8]. Recently, several groups have argued that ITC could play a more central role in lead optimization efforts in drug discovery [9, 10].

In a typical ITC experiment, small quantities of a titrant

30 (such as a ligand dissolved in buffer) contained in a syringe
31 are injected into a sample cell containing the *titrate* (often a
32 macromolecule in identical buffer), and the quantity of heat
33 liberated or absorbed as a result of each injection is mea34 sured. Given a model of the heat liberated from each injec35 tion due to the thermodynamics of binding, the thermody36 namic parameters of interest are then extracted from a fit of
37 the evolved heat per injection to the binding model [1]. Only
38 in the case that the reaction is *isenthalpic* (in which no heat
39 is produced or consumed) can no useful measurement can
40 be made with ITC.

Because various effects contribute to variation in the ex-<sub>42</sub> perimental operation—e.g. error in titrant or titrate concen-43 trations, unintended variation in injection volume, noise in the measured heat signal—the reported thermodynamic 45 quantities of interest will be determined only up to some 46 degree of uncertainty or error. While some properties of 47 this noise can be assessed by careful calibration runs, how the measurement error propagates into the thermodynamic quantities will depend on the actual binding characteristics of the system under study and the experimental protocol used to make the measurements. Further complicat-52 ing this is the problem that some quantities, such as the 53 degree to which the quantity of titrant actually injected in 54 the first injection is diminished (the so-called "first injection 55 anomaly" [11, 12]), are simply unknown. While good protocols attempt to minimize the sources of these errors [12–15], 57 they cannot eliminate them. With the Bayesian approach we describe below, however, we can at least attempt to account 59 for how these various sources of error affect the determined

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60 parameters.

# ditional nonlinear fitting procedures and their limitations?]

Here, we present a simple Bayesian formalism for inferring the full posterior distribution of the thermodynamic approach provides numerous advantages over more traditional approaches based on nonlinear least-squares error fitting [1, 16, 17]. Unlike least-squares methods, the Bayesian approach provides the full posterior distribution of the inferred parameters, from which joint distributions of the parameters of interest, asymmetric confidence intervals, and measures of covariance of estimates that do not depend on linearity assumptions and asymptotically normal error estimates can be extracted.

selects among models—such as the binding stoichiometry —the Bayesian method can assess the weight of evidence for each model and, given that model, what the unknown thermodynamic parameters are. Multiple experiments on 137 sample preparation [?]. identical or multiple titrants and macromolecules under different experimental conditions can be analyzed simultaneously to provide the best estimate of complex quantities of interest, like differences in binding affinities or entropies between lead candidates. Design of additional experiments can be aided by prior experimental data, knowledge of binding affinities, or even probable ranges, and the experimental protocol expected to yield the largest information gain selected. Finally, instead of using ad hoc subtraction techniques to deal with reference spectra or use of instrument noise reference values, calibration data or runs with blanks can be included directly in the inference procedure.

This paper is organized as follows: Section II describes the isothermal titration calorimetry experiment in detail. In Section III, we describe the Bayesian formulation for the posterior distribution of unknown thermodynamic parameters given the experimental data. An efficient scheme for sampling from the posterior and computing useful quantities from this sample is described in Section IV. We then illustrate and validate the Bayesian analysis scheme by applying it to fully independent replications of an experiment in which MgCl<sub>2</sub> is titrated into a cell containing the chelator EDTA to study the thermodynamics of the Mg<sup>2+</sup>:EDTA reaction in Section VII. Use of the Bayesian method to inform experimental design is described in Section VI, and various common binding models (which can be "pluggedinto" the Bayesian formalism) are provided in Appendix A. A free Python implementation of the Bayesian methodology described here, along with all datasets used in Section ??, is available from http://www.simtk.org/home/ bayesian-itc.

[JDC: Should we describe ITC generally, or focus on the [JDC: Do we need to include a paragraph describing tra- 116 MicroCal instrument(s) in particular?] In performing an 117 isothermal titration calorimetry (ITC) experiment with a modern titration microcalorimeter, two identical cells are isolated from the environment by a thermal jacket, as dequantities of interest (free energy, entropy, and enthalpy 120 picted in Figure 1. [JDC: How is the VP-ITC thermal jacket of binding) as well as any unknown instrument parame- 121 thermostatted to the sample and reference cell temperaters from the collected calorimetry data. The Bayesian 122 tures? Is it to minimize heat flow between cells and jacket?] One cell, the sample cell, is filled with a solution containing one or more molecular species dissolved in buffer (the 125 titrate). This usually consists of a macromolecule to which the binding of some ligand is to be assessed, but may also include a weak ligand to be displaced by a stronger ligand, as in the case of competitive binding experiments [18]. The other cell, the reference cell, is filled with a solution of iden-130 tical heat capacity to the titrate. A syringe with automatic injection control, mounted above the sample cell, contains Additionally, Bayesian methods allow for true binding 192 the titrant, which usually consists of a ligand in buffer. In model selection; instead of jointly inferring a parameter that order to minimize heats of mixing upon injection that will obscure the binding signal, the titrant and titrate buffers must be identical. Because of this, common protocols recommend dialysis of both against a common buffer during

> Initially, both cells and the syringe are thermostatted to the desired experimental temperature. Throughout the experiment, the reference cell is slowly heated by application of constant known power to a resistive heating element. This applied reference power is small—generally in the range of 2–30  $\mu$ cal/s for a  $\sim$  1.4 ml sample cell [19]—to ensure the total change in sample temperature over the duration of the 145 experiment (generally tens of minutes to hours) is minimal (much less than one degree Celsius). To maintain identical temperatures in both sample and reference cells, the sample cell is heated by a separate resistive heating element 149 slaved to a highly accurate sensor that measures the thermal differential between sample and reference cells, applying heat to the sample cell to minimize the thermal differ-152 ence and making accurate measurements of the power re-153 quired to do so.

> During the course of the experiment, a series of N injec-155 tions are performed in which a known volume of titrant is inise jected into the sample cell. The volume of the injection may  $_{157}$  vary with injection number, and the volume of injection nis denoted here by  $\Delta V_n$ . Many protocols, for example, recommend a reduced volume on the first injection due to the belief (now invalidated [12]) that some titrant is inevitably lost from the syringe needle prior to the first injection [19]. 162 The majority of instruments (such as the MicroCal VP-ITC) utilize perfusion-type configurations, in which a quantity of liquid equal to the injection volume flows out of the sample cell into an inactive tube, where it no longer contributes to the detected temperature differential [13, 19]. The interval of time  $\tau_n$  between the beginning of injection n and the beginning of the next injection (or the end of the recording period) is chosen to be large enough to allow adequate mixing and exceed all relevant binding (and, if present, reorganization) kinetics, as well as allow the feedback heating mechanism sufficient time to equalize the temperature between

173 the sample and reference cells prior to the next injection. If these criteria are not met, it becomes difficult (if not impossible) to deconvolute the heat signals from sequential injec-175 tions. 176

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While the experiment is running, the temperature difference between the reference and sample cells is sampled at discrete subsecond intervals, converted into power units, and used to drive the heater of the sample cell. This differential applied power between the reference and sample cells (which should average to zero in the absence of titrant injections) is averaged over fixed time intervals  $au_{
m filter}$  and stored to disk, along with readings of the cell temperatures and performance of the instrument.

If the binding reaction is either endothermic (consumes heat) or exothermic (produces heat), the temperature of the sample cell will deviate from that of the reference cell, and the integrated difference in power applied to bring the sample cell to the same temperature as the reference cell will provide a measure of the quantity of heat absorbed during the binding reaction. The various thermodynamic quantities of interest are then extracted from a fit of the evolved heat per injection to the binding model. Should the reaction be isenthalpic (in which no heat is produced or consumed), no useful measurement can be made with ITC. The power applied to the reference cell is chosen to minimize the duration of the experiment while ensuring that the temperature rise over the course of the experiment is minimal.

[JDC: This section contains old material that will be moved and reworked.]

We concern ourselves with M molecular species denoted  $X_m$ . Most commonly, M=2, where one species is a macromolecule and the other species is a ligand or binding partner whose affinity for the macromolecule is to be determined.

A sample containing one or more of the species  $X_m$  is prepared in buffer and loaded into a sample cell of volume  $V_0$ , where the concentration of each species  $X_m$  is initially  $[X_m]_0$ . Most often, the sample cell solution contains only a macromolecule, but a weak ligand of known affinity might also present in order to allow the affinity of very strong ligands to be determined. Another solution containing one or more species  $X_m$  (usually, the ligand of interest) is prepared in identical buffer at concentrations  $[X_m]_s$ , and loaded into 215 the injector syringe.

# **Evolved heat due to binding**

For a simple association reaction of two components P $_{218}$  and L without cooperativity,

$$P + L \stackrel{\Delta G_a}{\rightarrow} PL$$

where  $\Delta G$  denotes the free energy of binding, the dissocia- $_{220}$  tion constant,  $K_d$ , is given by

$$K_d \equiv \exp[-\beta \Delta G_a] (1 \text{ M}) = \frac{[P][L]}{[PL]} (1 \text{ M}).$$
 (1)

where [X] denotes the concentration of species X. We note 233 into the protein solution (as most solutions are nonideal), that  $K_d$  is expressed in units of *molarity* (mol/L).

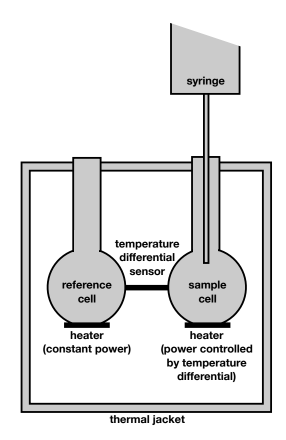


FIG. 1. Schematic diagram of an isothermal titration calorimeter. [JDC: The top of this figure needs to be clipped off.]

With each injection, the volume of buffer in the sample  $_{
m 224}$  cell increases, so that we can write the volume  $V_n$  after in- $_{225}$  jection n as

$$V_n = V_0 + n\Delta V. (2)$$

 $_{226}$  The total quantity (number of moles) of protein P and lig- $_{\it 227}\,$  and  $L_n$  in the cell after injection n is easily seen to be

$$P = V_0 [P_T]_0$$

$$L_n = n \Delta V [L_T]_s$$
(3)

228 Conservation of mass gives us the constraints

$$P = V_n ([P]_n + [PL]_n)$$

$$L_n = V_n ([L]_n + [PL]_n)$$
(4)

229 Combining Eqs.

With each injection, three effects will contribute to the (1)  $_{231}$  true quantity of heat  $q_n^*$  liberated by injection n: (1) the asso- $_{232}$  ciation of P with L, the (2) the dilution of ligand and buffer 234 and (3) the mechanical heat produced due by the injection 235 and stirring. We subsume the latter two components into a 263 (up to an irrelevant normalization constant independent of  $_{236}$  single term  $\Delta H_0$ , and write

$$q_n^* = V\Delta H ([PL]_n - [PL]_{n-1}) + \Delta H_0$$
 (5)

 $_{237}$  where  $\Delta H$  is the enthalpy change for the reaction P+L 
ightarrow $^{238}$  PL, and  $\Delta H_0$  is the heat of dilution and stirring.

Given  $K_d$ , we can determine the concentration of bound ligand after n injections,  $[PL]_n$ , from  $V_n, L_n$ , and P by (see 241 Appendix ??)

$$V_n [PL]_n = \frac{1}{2} \left\{ (V_n K_d + L_n + P) - \left[ (V_n K_d + L_n + P)^2 - 4L_n P \right] \right\}^{1/2}$$

$$- \left[ (V_n K_d + L_n + P)^2 - 4L_n P \right]^{1/2}$$
(6) 274

The *measured* heat  $q_n$  is assumed to differ from the *actual*  $_{243}$  heat  $q_n^*$  by a normally-distributed error:

$$p(q_n|q_n^*,\sigma) = \frac{1}{(2\pi)^{1/2}\sigma} \exp\left[-\frac{(q_n - q_n^*)}{2\sigma^2}\right]$$
 (7)

 $_{
m 244}$  where we have introduced the nuisance parameter  $\sigma$  to 245 quantify the magnitude of this error. Because each heat <sub>246</sub> measurement is assumed to be independent, the likelihood  $_{\mbox{\tiny 247}}$  of observing the data  ${\bf q}$  given the actual evolved heats  ${\bf q}^*$  is

$$p(\mathbf{q}|\mathbf{q}^*,\sigma) = \prod_{n=1}^{N} \frac{1}{(2\pi)^{1/2}\sigma} \exp\left[-\frac{(q_n - q_n^*)}{2\sigma^2}\right]$$
(8)

In the absence of any calibration information, we can as- $_{ t 250}$  sign the noise parameter  $\sigma$  a Jeffreys prior

$$p(\sigma) \propto \sigma^{-1}$$
 (9)

Putting this all together, we construct the complete pos-252 terior probability density function

$$p(\Delta G, \Delta H, \Delta H_{\text{dil}}, \sigma | \{q_n\}_{n=1}^{N})$$

$$\propto \left\{ \prod_{n=1}^{N} \frac{1}{(2\pi)^{1/2} \sigma} \exp\left[ -\frac{(q_n - q_n^*)^2}{2\sigma^2} \right] \right\} \sigma^{-1}$$

$$\propto \sigma^{-(N+1)} \exp\left[ -\frac{1}{2\sigma^2} \sum_{n=1}^{N} (q_n - q_n^*)^2 \right]$$
(10)

where the true heats per injection  $q_n^*$  are a function of  $\Delta G$ ,  $_{^{254}}$   $\Delta H$ , and  $\Delta H_0$ , as given by Eq. ??.

# **BAYESIAN FORMULATION**

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In Bayesian inference, we wish to infer the posterior dis-256 tribution of some unknown parameters  $\Theta$  given observed data  $\mathcal{D}$  and any prior information  $\mathcal{I}$ . The posterior quantifies our complete state of knowledge about how well vari- 311 ous choices for the *true values* of the unknown parameters  $\Theta$  312  $p(\theta|\mathcal{I})$  to the unknown thermodynamic parameters  $\theta$ . are supported by the available data and prior information. 313 <sub>262</sub> By Bayes' theorem, this posterior probability can be written <sub>314</sub> rameters, we choose to assign flat priors to binding affinities

$$p(\Theta|\mathcal{D}, \mathcal{I}) \propto p(\mathcal{D}|\Theta) p(\Theta|\mathcal{I})$$
 (11)

The conditional probability  $p(\mathcal{D}|\Theta)$ , termed the *likelihood*, describes the probability of observing data  $\mathcal{D}$  given underlying model parameters  $\theta$ . Due to various sources of random error, each realization of the same protocol of the experiment will generate a different set of observed data  $\mathcal{D}$ , <sub>270</sub> sampled from this distribution  $p(\mathcal{D}|\boldsymbol{\theta})$ . In practice, while we  $_{271}$  may not know  $p(\mathcal{D}|\boldsymbol{\theta})$  exactly, we can often come up with an extremely good model for this distribution if we know in detail what the experiment is measuring.

The quantity  $p(\theta|\mathcal{I})$ , termed the *prior*, expresses our state of knowledge of  $\theta$  before accounting for the data  $\mathcal{D}$ . This distribution characterizes whatever prior information  $\mathcal{I}$  we 277 may have, be it physical considerations or prior experimental data.

In an ITC experiment, the data  $\mathcal{D}$  consists of a series of measurements of the differential power applied to the sample cell, given a protocol describing the concentrations of various species in the sample cell and syringe, the injection volumes and times, and the time between injections. The unknown parameters  $\theta$  consists of the thermodynamic <sub>285</sub> parameters of interest such as the free energy of binding  $\Delta G$ , and its decomposition into enthalpic  $\Delta H$ , and en-<sub>287</sub> tropic  $-T\Delta S$  components. Also included in  $\theta$  are any unknown quantities that play a role in determining the distribution of observed data  $\mathcal{D}$ , such as the heat of dilution, heat due to stirring, magnitude of measurement errors, and so on. The prior information  $p(\theta|\mathcal{I})$  includes any prior infor-292 mation we might have from calibration runs or prior experimental data.

The posterior represents the joint distribution of all parameters—not just the ones we might care about at any moment. As a result, we can marginalize the posterior by integrating out parameters we don't care about, expressing the posterior distribution of only those of interest. For example, we may only be interested in the distribution of the free energy of binding  $p(\Delta G|\mathcal{D},\mathcal{I})$ , or perhaps the joint distribution of the enthalpic and entropic contributions  $p(\Delta H, -T\Delta S | \mathcal{D}, \mathcal{I})$ . Useful statistical quantities like means, (co)variances, and confidence intervals then be extracted from these distributions. The remaining parameters, which have been integrated out, are referred to as nuisance variables.

Below, we describe the various components necessary to model the posterior  $p(\theta|\mathcal{D},\mathcal{I})$  in a way that accounts for 309 many aspects of experimental error.

# Prior for thermodynamic parameters

Given our prior information  $\mathcal{I}$ , we must first assign a prior

In the absence of any previous knowledge about the pa-

 $\Delta G$  and enthalpies of association  $\Delta H$ :

$$p(\Delta G, \Delta H | \mathcal{I}) \propto 1$$
 (12)

<sup>316</sup> A uniform prior is appropriate because the quantities  $\Delta G$ and  $\Delta H$  can be of either sign, and of any value. (The entropic contribution to the binding affinity,  $-T\Delta S$ , is implicitly determined in terms of  $\Delta G$  and  $\Delta H$  by the definition of the Gibbs free energy  $\Delta G = \Delta H - T\Delta S$ .)

If we do have prior information about how, say, 322 the binding free energy must be within a given range  $\Delta G_{
m low}, \Delta G_{
m high}$  (e.g. as determined by another experimental technique), we can write

$$p(\Delta G, \Delta H | \mathcal{I}) \propto \begin{cases} 1 & \text{if } \Delta G \in [\Delta G_{\text{low}}, \Delta G_{\text{high}}] \\ 0 & \text{otherwise} \end{cases}$$
 (13)

Alternatively, if another technique has determined the binding affinity to be  $\Delta G^* \pm \delta \Delta G$ , where the error bars denote the standard error, we can use a normal prior

$$p(\Delta G, \Delta H | \mathcal{I}) \propto \delta \Delta G^{-1} \exp \left[ -\frac{1}{2} \frac{(\Delta G - \Delta G^*)^2}{(\delta \Delta G)^2} \right]$$
 (14)

Finally, if we already have collected data  $\mathcal{D}^*$  from a separate ITC experiment involving one or more of the thermodynamic parameters under study, we can simply use the posterior  $p(\theta|\mathcal{D}^*,\mathcal{I})$  as a suitable prior for  $\theta$ . This can be particularly useful when combining data from multiple experiments on a set of ligands, for example, to determine relative binding affinities or in the analysis of competition experiments where a weaker ligand is competed of by multiple 336 stronger ligands in separate experiments.

# Measurement error in evolved heat

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The calorimeter makes numerous measurements each second of the temperature difference between the reference and sample cell. This temperature difference is translated into power units and used to control the resistive element heating the sample cell. The resulting quantity is termed the differential power, by convention negative when an exothermic reaction in the sample cell results in less power applied to the sample cell than the reference cell, and positive when an exothermic reaction requires more heat to be applied to the sample cell than the reference cell to minimize the temperature differential.

Many sequential differential power measurements are av-350 eraged over a larger time window—termed the filter period  $au_{
m filter}$  —to produce an estimate of the true average differential power applied to the sample cell over this time. Because the true temperature differentials are very small, both the measured temperature differential (and hence reported differential power) and the true applied power will contain random errors. (We presume the instrument has been correctly calibrated so that bias is minimal compared to the 402 <sub>360</sub> ferential power measurements will be normally distributed <sub>405</sub> include the injection duration.]

about the true average differential power regardless of the 362 actual distribution of random errors in the individual measurements. Additionally, the filtering windows do not overlap, and so the random errors in different filtered power measurements can further be assumed to be independent.

The kinetics of injection, mixing, and binding are not relevant in determining the binding thermodynamics, provided 368 the time between injections is long enough for the sample cell to reach the same temperature as the reference cell before the beginning of the next injection (or the end of the measurement period). This is generally satisfied provided the timescale for binding/unbinding kinetics and any 373 associated conformational changes is much shorter than the time between injections. The total quantity of heat absorbed or emitted from the sample cell as a result of each injection is therefore a sufficient statistic for determination of the thermodynamic parameters of binding. We denote the measurement of the heat liberated (or absorbed) as a result  $_{379}$  of injection n by

$$q_n = \sum_{t=1}^{T_n} \Delta t \, P_{nt} \tag{15}$$

where the sum runs over all of the  $T_n$  differential power  $_{381}$  measurements  $P_{nt}$  taken starting with the beginning of in- $_{382}$  jection n and ending just prior to injection n+1 (or when 383 the recording period terminates).

We characterize the measurement noise  $\epsilon$  in the reported 385 differential power (resulting from both errors in the mea-386 sured temperature differential and errors in the true applied <sub>387</sub> power averaged over many samples during  $au_{
m filter}$ ) by a nor- $_{388}$  mal distribution of width  $\sigma$ :

$$\epsilon_t \sim \mathcal{N}(0, \sigma^2)$$
 (16)

The use of the normal distribution for noise  $\epsilon_t$  is, again, justi-390 fied by the central limit theorem. The noise in the measured  $q_n$  heat  $q_n$  will therefore also be normal, but with a variance that depends on the number of filtering periods  $T_n$  between  $_{393}$  injections n and n+1:

$$q_n \sim \mathcal{N}(q_n^*, T_n \sigma^2)$$
 (17)

In the absence of any calibration information, we can as- $_{395}$  sign the noise parameter  $\sigma$  a Jeffreys prior [20]

$$p(\sigma) \propto \sigma^{-1}$$
 (18)

 $_{396}$  If we do have calibration data on the magnitude of  $\sigma$ , we can 397 condition the prior on this data. For example, from a cal- $_{398}$  ibration run  $\mathcal{D}_{\mathrm{cal}}$  consisting of measuring the reported dif- $_{\mbox{\scriptsize 399}}$  ferential power  $P_{{\rm cal},n}$  over N filter periods where no ligand  $_{ t 400}$  is injected, we can update the prior on  $\sigma$  using the same as- $_{ t 401}$  sumption of normality for the  $P_n$  as above to obtain:

$$p(\sigma^2|\mathcal{D}_{\text{cal}}) \propto \sigma^{-(N+1)} \exp\left[-\frac{1}{2\sigma^2} \sum_{n=1}^N P_{\text{cal},n}^2\right]$$
 (19)

[JDC: We should allow more complex error models to be magnitude of these random errors.) Because of the filter- 403 included and, ideally, automatically selected. Look at Joel's ing stage, the central limit theorem demands the filtered dif- 404 paper on this topic [21]. Ideally we can extend his models to

# C. Errors in sample preparation

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Because the preparation of the sample and titrant will in-408 evitably errors in the concentrations of the solutions, we can account for these errors as well during our analysis to ensure that the reported binding affinities reflect these experimental uncertainties as well. In particular, errors in quantities ike the reported initial concentrations of macromolecule or igand will have a direct effect on binding affinities. [JDC: How much will a 1% error in initial concentrations affect the the reported binding affinities – is this directly proportional?] We can therefore include an error model to describe the uncertainties in the true initial concentrations given the reported initial concentrations.

Suppose, for example, we determine that the solution loaded into the sample cell had a concentration of macromolecule  $[M]_0 \pm \delta[M]_0$ , where the standard deviation  $\delta M$  was estimated by careful propagation of error during the sample preparation process (from estimates of known pipetting error magnitudes, known analytical balance accuracies, and reported compound purities). We can model the true initial concentration of macromolecule  $[M]_0^{\ast}$  by a normal distribution:

$$[M]_0^* \sim \mathcal{N}([M]_0, (\delta[M]_0)^2)$$
 (20) <sub>463</sub>

# D. Volume displacement by shaft and stirring paddle of syringe assembly

Insertion of the shaft and stirring paddle of the injector 430 syringe assembly into the sample cell displaces a small but significant volume which must be accounted for in order to avoid subsequent errors in the measured binding affinity [13]. For the MicroCal VP-ITC instrument, this volume has been estimated at  $0.044\pm0.005\,\mathrm{mL^1}$  though Calorimetry Sciences Corp. recommends performing a separate calibration procedure for their instruments [13].

To account for this reduction in sample cell volume, we provide two options: 439

1. If calibration data is available (of the form  $\Delta V_{disp}^{
m cal}$   $\pm$  $\delta \Delta V_{disp}^{\mathrm{cal}}$ ), a normal prior can be assigned to the displacement volume  $\Delta V_{disp}$  of the assembly:

$$\Delta V_{disp} \sim N(\Delta V_{disp}^{\rm cal}, (\delta \Delta V^{\rm cal})^2)$$
 (21

2. If no calibration data is available, or the calibration data is distrusted, the displacement volume can be inferred completely by assigning a broad uniform prior:

$$\Delta V_{disp} \sim U(0, V_{cell})$$
 (22)

446 Note that, in traditional ITC data analysis, the site parameter n is partially able to absorb errors in the stated cell volume, allowing correct binding affinities and enthalpies to be obtained at the expense of deviations from the expected n=1for 1:1 complexation reactions [13].

# The "first injection anomaly"

A commonly encountered problem in ITC experiments 453 performed according to manufacturer-recommended pro-454 tocols (such as the VP-ITC User Guide [19]) is the observa-455 tion that the first injection will yield an integrated heat of 456 magnitude smaller than expected, a phenomenon termed 457 the "first injection anomaly" [12]. Widespread practice has been to attempt to minimize the effect of this phenomenon 459 on the fit to the binding equations by performing a small 460 initial injection that is discarded before analysis, attributing the "first injection anomaly" to a loss of titrant from the sy-462 ringe needle prior to the first injection.

This phenomenon is now understood to be the result of 464 the "purge-and-refill" syringe preparation procedure suggested by instrument manufacturers [12]. As a result of this 466 procedure, the worm screw in the syringe that converts rotary stepper motion to vertical motion to expel the syringe contents must change direction from "fill" to "purge" during the first injection, resulting in an expulsion of titrant that is 470 smaller than expected. The anomaly can be eliminated by simply issuing the syringe a short "plunger down" command prior to loading the syringe into the apparatus [12]. This 473 modified procedure is highly recommended over the com-474 mon practice of discarding the initial injection, especially because such a procedure will change the observed binding behavior of subsequent titrant injected into the sample cell because some titrate is already complexed with titrant.

However, it is recognized that many datasets have already been collected using a protocol which results in a 480 first-injection anomaly. In order to permit analysis of these datasets within our framework, we provide the option of 482 inferring the quantity of first injection volume shortfall (21) 483 through the following procedure. We introduce two nui $v_{
m out}$  sance parameters:  $v_{
m out}$ , denoting volume of titrant lost external to the sample chamber or a shortfall of injection volume due to work gear reversal, and  $v_{\rm in}$ , denoting volume of titrant lost within the sample chamber prior to the start 488 of the first injection. We apply the reasonable constraint that the total volume of this loss must be less than the total 490 volume of the first injection,  $\Delta V_1$ . Other than this, we ap-<sub>491</sub> ply no additional knowledge of what the magnitude of this loss may be, assigning a uniform prior to  $v_{\text{out}}$  and  $v_{\text{in}}$  in this range. Their actual values will be inferred—and marginalized out, if they are not of interest—during the analysis from the magnitudes of heat liberated or consumed from all the 496 injections.

<sup>&</sup>lt;sup>1</sup> The statement "With allowance for uncertainties . . . this value is likely within 0.01 mL of the true value" [13] is taken to mean that the 95% confidence interval is  $\pm 0.01$  mL, suggesting a standard deviation of  $\pm 0.005$ mL as referenced in the text.

$$p(v_{\rm out},v_{\rm in}) \propto \begin{cases} 1 & \text{if } (v_{\rm out}>0) \cap (v_{\rm in}>0) \\ & \cap (v_{\rm out}+v_{\rm in} \leq \Delta V_1) \\ 0 & \text{otherwise} \end{cases} \tag{23} \qquad p(\Delta H,T\Delta S|\mathcal{D}) = \int d\Delta G \int d\Delta H_0$$

Other effects

JDC: Can we include other things into our model to make 499 analysis more robust?

- Baseline correction/drift
- Better models for heat of mixing/dilution
- Bubbles?

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Multiple experiments and including titrations with

#### SAMPLING FROM THE POSTERIOR

Because the posterior distribution (Eq. 10) is not amenable to direct sampling, we employ a Markov chain Monte Carlo (MCMC) procedure to sample from it for the purposes of computing means, variances, and approximations to the joint and marginal distributions [22].

JDC: Things to discuss here include

- ullet Gibbs sampling for parameters like  $\sigma$
- David's normally-distributed MCMC moves
- Automated step size adjustment during burn-in
  - Correlated moves among parameters for increased acceptance probability
- Gibbs sampling to allow convenient introduction of additional nuisance parameters?
- Assessing sampling burn-in and convergence

# **ANALYSIS**

# [JDC: This section is still under construction.]

In most practical applications, we will want to integrate 523 out parameters that are not of direct interest, producing 558 marginal distributions for the free energy of binding

$$p(\Delta G|\mathcal{D}, \mathcal{I}) = \int d\Delta H \int d\Delta H_0 \cdots \times p(\Delta G, \Delta H, \Delta H_0, \dots | \mathcal{D}, \mathcal{I})$$

or the joint distribution of the enthalpic ( $\Delta H$ ) and entropic  $_{526}$   $(T\Delta S)$  contributions to binding

$$p(\Delta H, T\Delta S | \mathcal{D}) = \int d\Delta G \int d\Delta H_0$$

$$\times \int d\sigma \, p(\Delta H - T\Delta S, \Delta H, \Delta H_0, \sigma | \mathcal{D})$$
(24)

- **Confidence intervals**
- Joint distributions

# **EXPERIMENTAL DESIGN**

# [JDC: This section is still under construction.]

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The Bayesian framework presented here can also be used 532 to aid in the design of new experiments using previously-533 collected experimental data or prior information. A natural way to judge the utility of a given choice of experimental design parameters y is to maximize the expected information sas content (EIC) of the experiment y given prior information  $\mathcal{I}$ :

$$E[I(\boldsymbol{y})] \equiv \int d\boldsymbol{\theta} I(\boldsymbol{y}|\boldsymbol{\theta}) p(\boldsymbol{\theta}|\mathcal{I})$$
 (25)

537 This prior knowledge may simply be the bare prior assumptions outlined above  $p(\theta)$ , or it may be conditioned on additional expectations about the potential range of binding affinity, prior experimental data  $\mathcal{D}$ , or any information about the behavior of the apparatus (such as control experiments). Though prior experimental data on the system of interest will likely be the most informative, any information (provided it is not wrong) can be useful in making the estimation of EIC more precise.

Instead of computing only the mean EIC, our strategy computed the posterior distribution of the EIC based on the given assumptions and data by a double sampling approach, allowing the operator to determine whether one experiment will clearly deliver more information than another experiment. (In fact, the probability that this is the case case

To estimate the posterior p(EIC), the following strategy 554 is used:

- 1. Given the prior information and any data or assumptions to condition on, a proposed true model  $\theta^*$  is sampled from  $p(\boldsymbol{\theta}|\boldsymbol{D}_1)$ .
- 2. A random outcome  $\mathcal{D}_y$  is chosen for one or more experiments  $\boldsymbol{y}$  by sampling from  $p(\mathcal{D}_{\boldsymbol{y}}|\boldsymbol{\theta}^*)$ .
- 3. The information content of the new data  $\mathcal{D}_y$  given prior data  $\mathcal{D}$  and prior information  $p(\boldsymbol{\theta})$  is computed and tallied, and the process repeated.

$$I(\boldsymbol{y}|\boldsymbol{\theta}) = \int d\mathcal{D}_y \, p(\mathcal{D}_y|\mathcal{I}) \, I(\mathcal{D}_y|\mathcal{I})$$

$$= \int d\mathcal{D}_y \, p(\mathcal{D}_y|\mathcal{I}) \, \left( H[p(\boldsymbol{\theta}|\mathcal{I})] - H[p(\boldsymbol{\theta}|\mathcal{D}_y, \mathcal{I})] \right)$$

$$= \tag{26}$$

$$\mathrm{E}\left[I(oldsymbol{y})\right] \equiv \int doldsymbol{ heta} p(oldsymbol{ heta}) I(oldsymbol{y} | oldsymbol{ heta})$$

$$= \int doldsymbol{ heta} p(oldsymbol{ heta} | \mathcal{D}) \tag{27}$$

# VII. ILLUSTRATIVE APPLICATIONS

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# Simple 1:1 complexation of ${ m Mg}^{2+}$ to EDTA

We illustrate the effectiveness of the Bayesian approach 622 in describing the true uncertainty in the experimental mea-1:1 binding of  $Mg^{2+}$  to the chelator EDTA. In order to assess the true variation among measurements, including er- 626 rors in the concentrations of titrant and titrate, the experi- 627 applications appearing in the paper.] ment was repeated from scratch ten times. For each trial, the 628 titrant, MgCl<sub>2</sub>, titrate, EDTA, and buffer, 50 mM Tris-HCl at pH 629 such as the MicroCal Auto-iTC<sub>200</sub> (capable of running 75 8.0, were weighed and dissolved to prepare solutions at the so samples/day, and 384 samples unattended) and the Vivacplanned concentrations; of course, the actual concentration 631 tis MiDiCal II (projected to be capable of running up to 1000 of the solutions in the syringe and sample cell differed due 692 samples/day) on the horizon, it becomes increasingly imto various measurement, preparation, and solution transfer 633 portant that robust, reliable, and automated methods be errors.

 $_{579}$  MgCl $_2$  was prepared to act as the titrant and 0.05 M solu-  $_{636}$  ware which automatically determines the optimal expertion of EDTA to act as as the titrate. Magnesium Chloride 637 imental design parameters (e.g. concentrations, competi-Hexahydrate (MgCl<sub>2</sub>) was purchased from Fisher Scientific 638 tion assays) for follow-up experiments (within specified re-(Catalog No. BP214-500, Lot No. 006533) and Ethylene- 639 source restrictions) that become too tedious or complex diaminetetraacetic acid, anhydrous (EDTA) was purchased 640 for a human laboratory worker. The Bayesian framework from Sigma-Aldrich (Catalog No. E6758-500G, Batch No. 641 described here provides a natural choice for these high-034K0034). Tris Base was purchased from Fisher Scientific 642 throughput environments, where all experimental data can (Catalog No. BP154-1, Lot No. 082483). Buffer was prepared 643 be simultaneously analyzed and used to decide which by weighing Tris base, adding MilliQ water, and adjusting the 644 follow-up experiments are necessary to confirm or reject hyfinal pH to 8.0. Solutions were prepared by weighing pow- 645 potheses with statistical certainty. der ( $0.1 \, g$  for MgCl<sub>2</sub> and  $0.01 \, g$  for EDTA) and adding the apby buffer, to make a concentrated solution (15 mM for MgCl<sub>2</sub> <sup>648</sup> bayesian-itc. and 1.0 mM for EDTA). The solutions were then further diluted with buffer to prepare the titrant and titrate.

The ITC experiment consisted of a total of 24 injections, 649 with the first programmed to deliver 2  $\mu$ L of titrant (MgCl<sub>2</sub>) and the remaining 23 injections programmed to deliver 12 650  $\mu$ L. Data was collected for 60 s prior to the first injection and and the reference power was fixed at 5  $\mu$ cal/s.

[JDC: More here.]

We have described a simple Bayesian framework for the analysis of isothermal titration calorimetry experiments that significantly extends what was possible with earlier analysis techniques. First, the approach more accurately captures the uncertainties in estimated thermodynamic parameters by allowing rigorous confidence intervals, rather than just standard deviations of the mean, to be estimated from the data. Second, by capturing the full posterior distribution of the thermodynamic parameters, complex (often nonlinear) covariations in the estimated parameters can be accurately assessed. This may, for example, have great utility in assessing whether entropy-enthalpy compensation is present in a system under study in a way that is statistically meaningful. Thirdly, the ability to combine data from multiple experiments conducted under different conditions allows binding affinities and other thermodynamic parameters to be determined much more precisely than before, and their errors to be assessed yet more accurately. Fourth, the ability to use existing information or data to aid in the design of new experiments using a rigorous information the-623 oretic criterion gives the experimenter a powerful new tool surements by studying a simple complexation reaction—the 624 that goes far beyond the 'rules of thumb' that have been the 625 mainstay of the field for many years.

[JDC: Add a paragraph summarizing observations on the

With automated, high-throughput microcalorimeters, available for analyzing the large quantities of ITC data that For each replicate of the experiment, a 0.5 M solution of GBB are becoming available. There is obvious potential for soft-

A Python implementation of this method is freely propriate amount of buffer, neglecting the volume occupied 647 available online at http://www.simtk.org/home/

# IX. ACKNOWLEDGMENTS

The authors are indebted to Sarah E. Boyce (UCSF), Alan P. Graves (GSK), Veena L. Thomas (UCSF), Brian K. Shoichet 300 s for each injection. The injection rate for all injections 652 (UCSF), and Phillip L. Geissler (UCB) for insightful discuswas 0.5  $\mu$ L/s. All experiments were conducted at 298.1 K,  $_{653}$  sions. JDC acknowledges support from a QB3-Berkeley 654 Distinguished Postdoctoral Fellowship. VSP acknowledges 555 support through an NSF grant for Cyberinfrastructure (NSF 656 CHE-0535616). JDC and SEB thank Brian K. Shoichet (UCSF) 658 periments presented here. [JDC: Add appropriate acknowl657 for generous use of his laboratory to conduct some of the ex659 edgments for other authors.]

# **Appendix A: Binding models**

Any reversible association model in which the equilibrium concentrations of the various species  $[X_n]$  can be computed (potentially numerically) in terms of the total quantity of each species in the sample cell can be used in the inference procedure. The inference procedure can be even used to test multiple models simultaneously by along transitions between models: The fraction of samples generated from each model reflects the degree by which that model is preferred (see Bayesian model selection [?]).

In developing a model of binding, we fundamentally must arrive at a way to compute enthalpy potential  $H(\boldsymbol{X})$  as a function of the total quantities of the various species  $X_n$  that are present in the sample cell. Usually, this is obtained by first constructing a physical binding model and identifying both the relevant association constants and other parameters and the enthalpy change parameters associated with these events. Using this information, the enthalpy potential can be expressed in terms of the equilibrium quantities of all species in the sample cell. For a given set of total quantities of each dissociable species, we can solve (potentially numerically) for the equilibrium concentrations of all species and compute the new enthalpy potential.

To illustrate this process, we have worked out the solution of the equilibrium concentrations and enthalpy potential for various binding models in common use below.

# 1. Simple two-component binding

In a two-component 1:1 complexation reaction, we have reversible association between a macromolecule M and ligand  $_{\it 677}$  L

$$M + L \stackrel{K_a}{\rightleftharpoons} ML$$
 (A1)

where the equilibrium bound and free concentrations are characterized by an association constant  $K_a$ 

$$K_a = \frac{[ML]}{[M][L]} \tag{A2}$$

 $_{679}$  where the total number of moles of macromolecule M and ligand L are given by the constraints

$$M = V([M] + [ML])$$

$$L = V([L] + [ML])$$
(A3)

 $_{ text{\tiny 680}}$  Combining these relations, we obtain a quadratic equation for the complex concentration [ML]

$$ML/V^2 - (M/V + L/V + 1/K_a) [ML] + [ML]^2 = 0$$
 (A4)

where the only solution that satisfies  $0 \leq [\mathrm{ML}] \leq \min\{[\mathrm{M}], [\mathrm{L}]\}$  is

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$$[ML] = \frac{1}{2V} \left\{ (M + L + V/K_a) - \left[ (M + L + V/K_a)^2 - 4ML \right]^{1/2} \right\}$$
 (A5)

The change in molar enthalpy  $\Delta H$  of the association reaction is denoted by

$$M + L \stackrel{\Delta H}{\to} ML$$
 (A6)

If we assign the [ML]=0 state a reference heat potential of 0, we can write the heat potential Q for a solution of M moles of macromolecule and L moles of ligand in a volume V as

$$Q(M, L, V) = \Delta H \cdot V \cdot [ML] \tag{A7}$$

where the concentration [ML] is a function of M, L, V, and  $K_a$  determined from Eq. A5.

# 2. Multiple independent binding sites of equal affinity

A binding model traditionally used in the analysis of macromolecule-ligand association in ITC experiments [?] assumes there are N independent binding sites on the macromolecule M for ligand L, all with equal association constants  $K_a$ :

$$M + L \stackrel{K_a}{\rightleftharpoons} ML$$

$$ML + L \stackrel{K_a}{\rightleftharpoons} ML^2$$

$$\vdots$$

$$ML^{N-1} + L \stackrel{K_a}{\rightleftharpoons} ML^N \tag{A8}$$

Note that, even though traditional analysis allows n to be fractional, N must, by physical necessity, be a whole number. (Fractional deviations from whole numbers in more traditional analysis can be considered as the fitting procedure attempting to correct for errors in the sample cell volume [13].)

All of the equilibrium constants are assumed to be equal:

$$K_a = \frac{[ML^n]}{[ML^{n-1}][L]} , n = 1, \dots, N$$
 (A9)

Again, we also have the conservation of mass equations

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$$M = V([M] + \sum_{n=1}^{N} [ML^{n}])$$
 
$$L = V([L] + \sum_{n=1}^{N} [ML^{n}])$$
 (A10)

To determine the heat potential Q, we further assume that the enthalpy change of each association event is identical

$$ML^{n-1} + L \stackrel{\Delta H}{\rightarrow} ML^n \ , \ n = 1, \dots, N$$
 (A11)

Assigning the fully dissociated macromolecule a reference heat potential of zero, the heat potential can be written:

$$Q = \Delta H(\sum_{n=1}^{N} n \left[ ML^{n} \right]) \tag{A12}$$

If N is unknown, it can be determined by assigning a prior to it (such as  $p(N|\mathcal{I}) \propto N^{-1}$ ) and sampling over a collection of models to determine which model is best supported by the evidence. Once determined, N can be fixed, and the thermodynamic parameters of binding determined given the fixed-N model.

# 3. N-component competitive binding

In an N-component competitive binding scenario, the macromolecule M can associate with any ligand  $L_1,\ldots,L_N$ 

701 The various equilibrium constants are defined as

$$K_{an} = \frac{[ML_n]}{[M][L_n]} \tag{A14}$$

For N>1, there is no longer a simple analytical solution for  $[ML_n]$  given total quantities of macromolecule M and ligands  $L_n$ , so we must solve for the  $[ML_n]$  numerically.

Given the association constants  $\{K_{an}\}_{n=1}^N$  and total quantities of macromolecule M and ligand  $\{L_n\}_{n=1}^N$ , we solve for the equilibrium concentrations [M],  $[ML_n]$ , and  $[L_n]$  given the set of equations:

$$M=V\left([M]+\sum_{n=1}^N[ML_n]\right)$$
 
$$L_n=V([L_n]+[ML_n])$$
 
$$[ML_n]=K_{an}[M][L_n] \tag{A15}$$

subject to the constraints that [M] > 0,  $[L_n] > 0$ , and  $[ML_n] > 0$ . This can be easily done with any number of nonlinear root-finding methods.

We presume that the binding of each species  $L_n$  is governed by a separate enthalpy of association  $\Delta H_n$ :

$$M + L_n \stackrel{\Delta H_n}{\to} ML_n \tag{A16}$$

<sub>709</sub> Assigning the fully dissociated state a reference heat potential of zero, the heat potential can be written:

$$Q = \sum_{n=1}^{N} \Delta H_n \left[ M L_n \right] \tag{A17}$$

# 4. Cooperative binding with a Hill coefficient

For the case where ligand L binds cooperatively to macromolecule M with Hill coefficient n, association is described by 712 the process

$$M + nL \stackrel{K_a,n}{\rightleftharpoons} ML^n \tag{A18}$$

713 where the association constant is defined by

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$$K_a = \frac{[ML^n]}{[M][L]^n} \tag{A19}$$

714 Together with the conservation equations

$$M = V([M] + [ML^n])$$

$$L = V([L] + n[ML^n])$$
(A20)

we can again numerically solve for [M], [L], and  $[ML^n]$ .

If we again define the fully dissociated species as having a heat potential of zero, the heat potential Q can be written

$$Q = \Delta H \left[ ML^n \right] \tag{A21}$$

 $_{777}$  where  $\Delta H$  refers to the enthalpy of association for n ligands. The per-ligand association enthalpy can be obtained by divid-718  $\log \Delta H$  by n.

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