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Uncertainty in protein–ligand binding constants: asymmetric confdence intervals versus standard errors

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Abstract

Equilibrium binding constants (K_b) between chemical compounds and target proteins or between interacting proteins provide a quantitative understanding of biological interaction mechanisms. Reported uncertainties of measured experimental parameters are critical for decision-making in many scientifc areas, e.g., in lead compound discovery processes and in comparing computational predictions with experimental results. Uncertainties in measured K_b values are commonly represented by a symmetric normal distribution, often quoted in terms of the experimental value plus–minus the standard deviation. However, in general, the distributions of measured K_b (and equivalent K_d) values and the corresponding free energy change ΔG_b are all asymmetric to varying degree. Here, using a simulation approach, we illustrate the effect of asymmetric K_b distributions within the realm of isothermal titration calorimetry (ITC) experiments. Further we illustrate the known, but perhaps not widely appreciated, fact that when distributions of any of K_b , K_d and ΔG_b are transformed into each other, their degree of asymmetry is changed. Consequently, we recommend that a more accurate way of expressing the uncertainties of K_b , K_d , and ΔG_b values is to consistently report 95% confidence intervals, in line with other authors' suggestions. The ways to obtain such error ranges are discussed in detail and exemplifed for a binding reaction obtained by ITC.

Keywords Isothermal titration calorimetry · Confdence intervals · Standard error · Log-normal distribution · Dissociation constant · Binding constant

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Introduction

Interactions between biomolecules are central to many areas of biomedicine. Protein–protein interactions (Typas and Sourjik [2015;](#page-9-0) Pierce et al. [1999](#page-8-0)) are important, e.g., in immunological antibody–antigen binding reactions (Dam et al. [2008](#page-8-1)), or gene regulatory protein–nucleic acid interactions (Wells et al. [1980](#page-9-1); Buurma and Haq [2007](#page-7-0); Salim and Feig [2009](#page-8-2)). Furthermore, interactions of small molecular weight compound with proteins are fundamental to the action of many metabolic enzymes and their regulators, and in drug discovery during the search for lead compounds, as well as in the fnal characterization of promising therapeutic drugs (Geschwindner et al. [2015](#page-8-3); Renaud et al. [2016;](#page-8-4) Ladbury et al. [2010](#page-8-5)).

Numerous techniques are used to determine biomolecular interactions (Renaud et al. [2016](#page-8-4); Ciulli [2013](#page-8-6)), such as the inhibition of enzymatic activity (Smirnovienė et al. [2017](#page-8-7)), surface plasmon resonance (Myszka and Rich [2000](#page-8-8); Patching [2014;](#page-8-9) Olaru et al. [2015](#page-8-10)), isothermal titration calorimetry (ITC) (Krimmer and Klebe [2015](#page-8-11); Callies and Daranas [2016](#page-7-1); Falconer [2016;](#page-8-12) Vega et al. [2016](#page-9-2); Chaires [2008](#page-7-2); Leavitt and Freire [2001](#page-8-13)), thermal shift assay (diferential scanning fuorimetry) (Pantoliano et al. [2001;](#page-8-14) McDonnell et al. [2009;](#page-8-15) Yanchunas et al. [2005](#page-9-3); Cimmperman and Matulis [2011;](#page-7-3) Cimmperman et al. [2008](#page-8-16)) and numerous others (Renaud et al. [2016\)](#page-8-4). All these techniques are expected to provide comparable values of intermolecular interaction afnities provided that measurements are feasible at nearly identical conditions.

Reporting the errors of association reactions must be done consistently and accurately to enable reliable interpretation and reuse of results (Jarmoskaite et al. [2020\)](#page-8-17). In most scientifc literature, the uncertainty (i.e., the repeatability) of a binding equilibrium measurement is expressed as $\pm x$ (or $x\%$), accounting for the standard deviation or error of the measured value, with the underlying assumption that the measured affinity values are distributed randomly according to a normal (i.e., Gaussian) statistical distribution (Krimmer and Klebe [2015](#page-8-11); Schnapp et al. [2016](#page-8-18); Ladbury and Doyle [2004](#page-8-19); Lafont et al. [2007](#page-8-20); Dullweber et al. [2001;](#page-8-21) Rühmann et al. [2015;](#page-8-22) Gaspari et al. [2016;](#page-8-23) Pulido et al. [2015](#page-8-24); Hörtner et al. [2007](#page-8-25); Rechlin et al. [2017](#page-8-26); Cheng et al. [2017](#page-7-4); Huschmann et al. [2016](#page-8-27); Guan et al. [2013;](#page-8-28) Ren et al. [2014;](#page-8-29) Krishnamurthy et al. [2007\)](#page-8-30). For a normal distribution, the reported $\pm x$ -value typically corresponds to the 68.3% symmetric confidence interval $(Cl_{s68.3})$. In some cases, the reported symmetric uncertainty is estimated from analysis of several repeated measurements via the appropriate Student's *t*-distribution. However, the uncertainty is frequently retrieved only from the ftting program used to estimate the parameter from a single set of experimental data (Brautigam et al. [2016](#page-7-5)).

The equilibrium binding constant K_b (also referred to as the association or afnity constant) is inversely related to the dissociation equilibrium constant $K_d = 1/K_b$ and is related to the change in the standard Gibbs energy upon binding, $\Delta G_{\rm b}$, (often written with the naught symbol, $\Delta G_{\rm b}^{\rm o}$, which is omitted here for terms of simplicity) by the equations: $\Delta G_{\rm b}$ = – RT ln $K_{\rm b}$, or $K_{\rm b}$ = e^{- $\Delta G_{\rm b}/RT$}. These logarithmic, exponential, and reciprocal relationships do not preserve the shape of distribution of values. For example, a continuous probability distribution of a random variable, whose logarithm is normally distributed follows a log-normal distribution, taking only positive, real values. Therefore, if a normal distribution is assumed for ΔG_b , K_b must have a log-normal distribution.

The inherent asymmetry in parameter distribution has consequences for the reporting of errors. Symmetric $\pm x$ -value is not an appropriate way of accurately expressing the uncertainty for ΔG_b , K_b and related values. Asymmetric *F*-statistics-associated CIs have been introduced as

an elegant approach to report error ranges and the propagation of CIs in non-linear, asymmetrical variance spaces, accounting for non-normal distributed parameter distributions, including ΔG_b , K_b and K_d (Kemmer and Keller [2010](#page-8-31); Krainer and Keller [2015;](#page-8-32) Krainer et al. [2012;](#page-8-33) Broecker et al. [2011](#page-7-6); Johnson [1983](#page-8-34)).

In this work, we explore in detail the unrealistic nature of the assumption of a normal distribution of $K_{\rm b}$, $K_{\rm d}$ and other thermodynamic quantities through various examples and evaluate a practical approach, based on previously established procedures (Kemmer and Keller [2010;](#page-8-31) Krainer and Keller [2015;](#page-8-32) Krainer et al. [2012](#page-8-33); Broecker et al. [2011](#page-7-6)), to obtain estimates of the asymmetric *F*-statistics-associated CIs that are both simpler to treat consistently in transforming between different expressions for the affinity and better refect the real uncertainties in the data.

Results and discussion

We use simulated distribution curves to show how the mathematical relationships between the parameters $K_{\rm b}$, $K_{\rm d}$, and ΔG_b affect the asymmetry of their distributions.

Case 1: Transformation of the distribution of $K_{\rm b}$ **and Δ***G***b values assuming that each has a normal distribution**

Here, we show how transformations of K_b and ΔG_b values results in an asymmetry of their distributions. For this, we assumed that either K_b or ΔG_b values are normally distributed and generated a set of 10,000 random K_b 's with mean value 2×10^7 M⁻¹ and standard deviation of ± 0.6 \times [1](#page-2-0)0⁷ M⁻¹ (Fig. 1a) and a set of 10,000 random $\Delta G_{\rm b}$'s with mean value -41.67 kJ/mol and standard deviation of \pm 0.77 kJ/mol (Fig. [1d](#page-2-0)). Note that K_b =2 × 10⁷ M⁻¹ corresponds to ΔG_b =− 41.67 kJ/mol at 25 °C. Each set of values is then converted to the other representation through the appropriate logarithmic or exponential formula (Fig. [1](#page-2-0)b, e), and in both cases to the dissociation constant K_d (Fig. [1c](#page-2-0), f). These transformations show that the Gibbs energy change, and the binding and dissociation constants do not follow a normal distribution at the same time. The transformation between K_b and ΔG_b in either direction introduces a similar amount of skew into the distribution, and that the transformation between K_b and K_d may impart even greater skew.

When considering the kinetics of the interactions, inaccuracies in the reported parameters (most probable value and uncertainty) may be even more severe. The rate constants are related with the Gibbs energy variation between the reactants and the transition state through an exponential function, and this energy variation is usually larger than that observed between the reactants and the products of the

Fig. 1 Illustrating the effect of transformation of K_b and ΔG_b values into each other and into K_d distributions of protein–ligand binding. **a** Distribution of 10,000 random values of K_b generated for a normal distribution with mean 2×10^7 M⁻¹ and standard deviation of $\pm 0.6 \times 10^7$ M⁻¹. **b** Distribution of ΔG_b values calculated from the 10,000 K_b values in (a). **c** The distribution of dissociation constants K_d corresponding to data in (**a**). **d** Distribution of 10,000 random values of the Gibbs energy of binding generated for a normal distribu-

tion with mean−41.67 kJ/mol and a standard deviation of 0.77 kJ/ mol. **e** Distribution of K_b values calculated from the ΔG_b values in (**d**). **f** The dissociation constant calculated from (**d**). The logarithmic transformation introduces a skew of 1.3 (**b**) and the exponential transformation a skew of 0.9 (**e**) from the normal distributions. The greatest skewness arises from the transformation of the normally distributed K_b to K_d [skew is equal to 4.3 in (**c**)]

transformation. This situation is analyzed in the Supplementary Material.

Confdence intervals are a consistent and inter‑convertible way to accurately represent measurement uncertainty

The previous illustration shows that it is not warranted to use a symmetric $\pm x$ -value as an accurate expression of the uncertainty for all three quantities ΔG_b , K_b and K_d describing the same physical equilibrium. For consistent reporting of uncertainties in experimental data, it is always necessary to state the range of uncertainty $[x_{low}, x_{high}]$ which makes apparent any degree of asymmetry. Asymmetry in any skewed distribution is more evident toward its extremes; thus, the use of a central 68.3% confdence interval does not efectively describe the asymmetry. Consequently, it is advantageous for clear description to report the larger 95% confidence interval (Cl_{95}) , which in any case typically better represents the range in which the true value of the parameter is likely to be found.

Given that a normally distributed ΔG_b is transformed into a log-normal distribution for K_b , one could imagine that given some CI for ∆*G* (symmetrical), it might be required to calculate the 95% confdence interval of this log-normal distribution to obtain uncertainties in K_b . However, as the individual values for the quantities are correctly transformed by the exponential, logarithmic and reciprocal relationships, it is simple to directly convert the lower and upper values of any confdence interval between representations (this is rigorously true for one-to-one mono-parametric conversions). For example, using the ΔG_b =− 41.67 ± 0.77 kJ/mol from Case 1 gives a CI₉₅ = [-43.18, -40.16] for ΔG_b (as 95%) CI limits are $1.96 \times \sigma$ for a normal distribution in the limit of a large number of data points). The upper limit of the CI interval for $K_b = \exp(-\Delta G_{b, lower}/RT) = \exp(43.18/2.47896)$ $=3.68\times10^{7}$ M⁻¹. Similar transformation for the lower limit gives a CI_{95} =[1.09, 3.68]×10⁷ M⁻¹ (which can be seen to match with the distribution in Fig. [1e](#page-2-0) for which the CI_{95} is $[1.09, 3.69] \times 10^7$ M⁻¹). Thus, consistently reporting CI₉₅ makes it easy to transform between representations preserving all information regarding uncertainties in experimental values. This procedure also eliminates the possibility of getting error intervals with negative values for the equilibrium constants.

The logarithmic relationship between K_b and ΔG_b , is also important for other practical purposes such as calculating averages from a set of determinations. The mean of several ΔG_b values can be calculated using the arithmetic average, while the mean of several K_b values should be calculated using the geometric average. Alternatively, the value to report for K_b can be calculated from the obtained arithmetic average of ΔG_b values. That way, the correspondence between ΔG_b and K_b averages is maintained in the same way that the correspondence between confdence interval limits is maintained.

Case 2: Simulated error distributions for ITC measurements of 1:1 binding

The preceding hypothetical illustrations (Case 1) leave open the question of whether the uncertainty in any parameter might be expected to have a strongly asymmetric or near normal distribution, and, if so, under what circumstances. To investigate these issues, we simulated the impact of binding affinity changes on the variability of ITC experiments.

In common with many methods for obtaining binding constants, ITC experiments are performed as a titration where successive injections of one reactant species into a fxed amount (or concentration) of the other species leads to progressive saturation of a binding site. A signal monitors formation of the bound form and the values for the signal (transformed into heat, Q) are fit to an equation describing the relationship between the heat released or absorbed and the thermodynamic parameters K_b (or ΔG_b), the enthalpy change ΔH_b and the apparent stoichiometry, that are, thus, determined as parameters of the ft. The measurement errors are typically propagated to the ftted parameters in a way that depends on the equation describing the titration and a range of experimental variables, e.g., the number of injections, the fnal degree of saturation and concentrations of reactants.

The measurement's variability for the heat of each injection is random normally distributed (as it arises from the combined random efects of mechanical variability of the injection volume, electrical noise and how these impact upon the software process for integrating the heat signal for each injection).

To simulate the effect of measurement variation, random values from an appropriate normal distribution are added to the theoretically expected injection heat values for an ideal 1:1 binding reaction. The simulations used here model a VP-ITC instrument (MicroCal/Malvern). Tellinghuisen has determined that for this instrument (when set for high maximum injection heats) the standard deviation of measurements (when including the efect of subtracting blanks or heats of dilution) is approximately constant at 3 μJ for injections of heat $<$ 500 μ J (Tellinghuisen [2005\)](#page-8-35). Recently, alternative error models with a smaller constant standard deviation of 0.5–0.9 μJ plus an injection-heat-dependent term of 0.002–0.01 μJ per μJ (not including effects of subtraction) have been proposed (Tellinghuisen, [2017](#page-8-36),[2018](#page-8-37)). We use the original injection-heat-independent error model here, but note that it may overestimate the error for an optimally set up instrument for a low-heat biochemical reaction; however, the absolute value of the error is not critical as it is the proportional error (the signal to noise) that infuences the shape of the parameter error distribution. The variation of the fitted parameters, K_b (and the corresponding ΔG_b) and ΔH_b across 10,000 Monte Carlo simulated 1:1 binding reactions of fxed stoichiometry are shown in Fig. [2.](#page-4-0) The simulations performed here calculate the dilution efects on reactant concentrations occurring during the titration and modifcation of injection heats due to the volume displaced from the reaction cell following a discrete inject step model (or instantaneous injection model) (Tellinghuisen [2003](#page-8-38)). The simulated experiments are then ftted using an unweighted least-squares ft to the Wiseman equation (Wiseman et al. [1989](#page-9-4)) following typical experimental data analysis practice.

The simulations show that for all three true K_b values, the expected variation in the observed value is asymmetrically distributed. Furthermore, increasing the magnitude of K_b increases both the range of variation and the positive skew of the distribution of the K_b values (Table [1\)](#page-4-1), i.e., there is an increasing tendency to observe more frequently higher than lower affinity values as K_b increases; thus, the higher K_b values become more likely to occur. For this particular experimental scenario, we fnd that there is an 8% probability of observing a K_b less than half, and a 9% chance of more than twice the true value when $K_b = 2 \times 10^5 \text{ M}^{-1}$. Those probabilities, respectively, increase to 18% and 24% when the assumed $K_b = 2 \times 10^7$ M⁻¹ (and the probability of an observation more than $4 \times$ the true value is 12%).

The simulated scenario in Fig. [2](#page-4-0) has relatively low injection heats of approximately 1/3rd of the average for protein–ligand interactions in a recent large-scale study (Scheuermann and Brautigam [2015\)](#page-8-39) and with high proportional measurement errors. These values have been chosen to clearly illustrate the various efects on the distributions of increasing afnity. Since the asymmetry increases with proportional measurement error (Tellinghuisen [2017](#page-8-36)), observed asymmetries would be smaller for most experimental cases where those errors are smaller. Even in this simulated experimental scenario, the resulting ΔG_b distribution has little asymmetry except at the highest affinity and conversely the enthalpy change is only appreciably asymmetric and has greater uncertainty at the lowest affinity. In particular, we note that in this experimental scenario even with its quite large proportional measurement ∆*G*_b is near normally distributed for reasonably optimal experiments

Fig. 2 Monte Carlo simulation modeling of variation in the ftted parameters of ITC experiments for three different binding affinities **a**–**c** 2×10^5 M⁻¹, **d**–**f** 2×10^6 M⁻¹ and **g**–**i** 2×10^7 M⁻¹. For each binding affinity, 10,000 simulated data sets were created. All simulations were for a 1:1 binding reaction with molar enthalpy change

 ΔH_b =− 10 kJ/mol, 20×15 µL injections of 200 µM ligand into 20 μM protein leading to a fnal ligand:protein ratio of 2.5:1. The simulations are modelling a VP-ITC instrument and a measurement error of standard deviation of 3 μ J per injection. Simulated datasets where a satisfactory ft could not be obtained are excluded

Table 1 Statistical parameters for simulated ITC data at three different binding affinities

Starting K_h (M ⁻¹)	K_{h} Skew	$K_{\rm h}$ (M^{-1}) CI_{95}	$\Delta G_{\rm h}$ (kJ/mol)	$\Delta G_{\rm b}$ Skew	$\Delta G_{\rm h}$ (kJ/mol) $CI_{\alpha5}$	$\Delta H_{\rm h}$ Skew	$\Delta H_{\rm h}$ (kJ/mol) CI_{95}
2×10^5	1.3	$[0.78, 5.8] \times 10^5$	-30.26	-0.2	$[-32.9, -27.9]$	-0.7	$[-13.2, -8.0]$
2×10^6	1.5	$[0.78, 6.5] \times 10^6$	-35.97	-0.4	$[-38.9, -33.6]$	-0.2	$[-11.1, -9.1]$
2×10^7	23	$[0.51, 22] \times 10^7$	-41.67	-1.1	$[-47.6, -38.3]$	-0.1	$[-10.8, -9.3]$

Data correspond to the distributions in Fig. [2.](#page-4-0) The 95% confidence intervals CI_{95} are determined directly from central 95% of values of the distributions

(Wiseman parameters C of 4 and 40, respectively for the frst two simulations) for less optimal experiments (Wiseman parameter of 400 for the last simulation) skewness can be readily seen. This can most likely be attributed to the fact that for a 20-point ITC experiment with the parameters used for the simulation essentially only 3–4 datapoints fall

into the transition region and any noisy data here will have a bigger infuence leading to larger deviations and long tailing to higher K_b values. Also seen is an inverse relationship between the variations in measured affinity and enthalpy, this is expected due to the shape of the titration curve changing appreciably with binding affinity under the experimental conditions we have simulated with fxed concentration. At high C, almost all the ligand binds in the early injections of a titration giving several injection heats with high signal to noise and, thus, lowering the observed variation in the enthalpy measurement. An inverse relationship between the uncertainties in affinity and enthalpy is generally expected for ITC data. Under the experimental conditions that we have simulated with fxed concentrations, the shape of the titration curve changes appreciably with binding affinity.

Error estimations for ΔG_b **and** K_b **using asymmetric profle likelihood Confdence Intervals**

Because the calculation is built into most analysis software, most researchers (Salim and Feig [2009](#page-8-2); Geschwindner et al. [2015](#page-8-3); Renaud et al. [2016;](#page-8-4) Ladbury et al. [2010;](#page-8-5) Ciulli [2013](#page-8-6); Smirnovienė et al. [2017;](#page-8-7) Myszka and Rich [2000](#page-8-8); Patching [2014](#page-8-9); Olaru and Bala [2015](#page-8-10); Krimmer and Klebe [2015;](#page-8-11) Callies and Daranas [2016;](#page-7-1) Falconer [2016](#page-8-12); Vega et al. [2016](#page-9-2); Chaires [2008;](#page-7-2) Leavitt and Freire [2001](#page-8-13)) report the precision of the affinities between molecules using asymptoticsymmetric confidence intervals $(Cl_{S,\alpha})$ calculated using the standard deviation for each parameter, as estimated from the sum of squared residuals, RSS, and the covariance matrix from the ftting analysis, and a chosen confdence level *α* (typically 68.3%) of a *t*-Student distribution. The experimental and simulated results have shown that this symmetric approach does not communicate and accurately preserve information about uncertainty, and that use of asymmetric F -statistics-associated (or profile likelihood) CI_{95} is both simple and preferable. How then can we estimate CI_{95} in practice, e.g., from single experiments? We will analyze this case here in detail, as it is the simplest one. In the case of several experiments, the reasoning will be basically the same—we suggest to perform a global ft with all individual datasets and get the global confdence intervals.

A possible approach is to fit for ΔG_b and use the capabilities of diferent software to obtain the asymptotic-symmetric error $CI_{S,95}$ for the Gibbs energy change. Asymmetric confidence intervals for K_b or K_d can then be obtained by simply transformation of the upper and lower limits of the ΔG_b confidence interval. As we have seen that for titration data ΔG_b is less affected by asymmetry, consequently this approach will be efective in many circumstances and is certainly better than current practice. However, it also fails in some circumstances, e.g., for high affinity interactions as shown in Case 2 or where measurement errors are large [as extensively studied previously (Tellinghuisen [2017\)](#page-8-36)]; so, prior knowledge of the theoretically expected behavior of ΔG_b uncertainty for the experiment scenario is required to apply it. Consequently, this approach cannot be generally recommended.

A second possible approach that is applicable in all circumstances is to use a 'bootstrap' procedure in which values of the residuals of the best ft are randomly selected and added to the ftted value at each titration point and the new set of data points so created reftted. Repeating this re-sampling procedure many times (>1000) yields a distribution of values for each ftted parameter from which the confdence intervals can be obtained. This approach is applicable where the measurement variability is constant throughout the titration and produces $CI_{S,95}$ ranges that are only slightly larger than the true values. Unfortunately, the requisite re-sampling procedure is not widely available in commercial software.

We believe that the most practicable way to express the repeatability for K_b and ΔG_b is through calculation of the profile likelihood confidence intervals $CI_{P\alpha}$ (at statistical signifcance level *α*) (Kemmer and Keller [2010](#page-8-31); Krainer and Keller [2015](#page-8-32); Krainer et al. [2012](#page-8-33); Broecker et al. [2011](#page-7-6); Johnson [1983\)](#page-8-34), which can be determined by an extension of the typical ftting approach. Once the non-linear least squares regression analysis of *N* experimental points has been performed with a model with *P* parameters, the best estimates for the *P* parameters are obtained with an associated residual sum of squares $RSS₀$. Ideally, the *P* parameters could be systematically varied to get a *P*-dimensional contour fulflling the expression:

$$
RSS = RSS_0 \left(1 + \frac{P}{N - P} F_{P,N - P}(\alpha) \right),
$$

where $F_{n,m}$ is the Fisher–Snedecor distribution with $n = P$ and $m = N - P$ degrees of freedom, and the α is the chosen confdence level (Motulsky and Christopoulos [2004](#page-8-40)). Within that *P*-dimensional contour, the diferent possible sets of *P* parameters provide RSS values that are not statistically different (at a confidence level α) from RSS₀. Then, by projecting the *P*-dimensional contour onto the diferent *P* axes, the confdence interval for each parameter can be determined. However, this procedure is not practical if there are more than two ftting parameters. Therefore, very often marginal confdence intervals are determined by varying just one parameter at a time (Kemmer and Keller [2010;](#page-8-31) Bates and Watts [2007\)](#page-7-7). Thus, a given parameter, *p*, is selected and kept fxed at diferent values, while the RSS is minimized over the remaining free parameters, an RSS (*p*) curve (RSS as a function of *p*) with the resulting minimized RSS values (see Fig. [3](#page-6-0), right). The two limiting values for the given parameter *p* defining its profile confidence interval $CI_{P,\alpha}$ will fulfll the expression:

Fig. 3 a A simulated ITC titration curve corresponding to the K_b =2×10⁶ M⁻¹, ΔH =41.8 kJ/mol, and 1:1 binding stoichiometry with random measurement error for each injection. The continuous line corresponds to the best fit considering either K_b or ΔG_b as fitting parameters. **b** RSS dependence on $K_{\rm b}$. **c** RSS dependence on $\Delta G_{\rm b}$. Either K_b or ΔG_b are varied systematically (stepped through fixed values below and above their best estimate) and the remaining parameters are freely adjusted to re-minimize the RSS; then, that minimum RSS is plotted as a function of either K_b or ΔG_b . The minimum in

$$
RSS(p) = RSS_0 \left(1 + \frac{1}{N - P} F_{1, N - P}(\alpha) \right)
$$

These two limiting values defne the interval in which the parameter *p* provides RSS values that are not statistically different (at a confidence level α) from RSS₀. The process can be repeated for each of the other parameters, and all *P* marginal confdence intervals estimated.

In general, RSS is not a symmetric function of the studied parameter with respect to the minimum value $RSS₀$, and the two limiting values satisfying the previous equation defne an asymmetric confdence interval for each parameter. The asymmetry degree and the size of the confdence interval depends on the nature of the parameter considered and the sensitivity of RSS to that parameter.

The procedure is illustrated by analyzing an isothermal calorimetric titration simulated (with $K_b = 2 \times 10^6$ M⁻¹, $\Delta H = 41.8$ kJ/mol, and $n = 1$) with a chosen noise level (Fig. [3\)](#page-6-0) and shown in detail for the Origin software as an example in Supplementary Material (2. Quick calculation of profile likelihood asymmetric confidence intervals $CI_{P,95}$.).

The standard output for the best fit to the titration in Fig. [3](#page-6-0) has the following estimated parameters: $K_b = (2.2 \pm 0.4) \times 10^6$ M⁻¹, $\Delta H = 43.3 \pm 1.2$ kJ/ mol, and $n = 0.99 \pm 0.01$. If the Gibbs energy of

RSS (i.e., $RSS₀$) corresponds to the best fit (all parameters freely varying, including K_b or ΔG_b). The horizontal dotted line is the limit reference value for RSS with 95% confidence, equal to $RSS₀$ (1+(1/ $(N-P)F_{1,N-P}(0.95)$). The intercepts between the RSS curve and the reference RSS value (indicated by arrows) provide the limits for the confdence intervals. The way to get these CIs with the Origin software is shown in Supplementary Material as an example (2. Quick calculation of profle likelihood asymmetric confdence intervals $CI_{p.95}$.)

) interaction is considered as a ftting parameter instead of K_b the best fit provides the following estimated parameters: $\Delta G = -36.2 \pm 0.4$ kJ/mol, $\Delta H = 43.3 \pm 1.3$ kJ/mol, and $n = 0.99 \pm 0.01$.

These uncertainties correspond to the typical standard errors $CI_{S.68.3}$ for the fitting parameters generated from the covariance matrix. The asymptotic-symmetric confdence intervals for a confidence level of $\alpha = 95\%$ CI_{S 95} are calculated as a multiple of these values determined by the *t*-Student distribution. The asymptotic-symmetric and asymmetric profle likelihood 95% confdence intervals are shown in Table [2.](#page-7-8)

As expected from the previous results, the uncertainty in K_b is revealed to be asymmetric by the profile likelihood confidence interval $CI_{P,95}$. Also as expected from the Monte Carlo simulations (in Case 2) at this binding affinity, the $CI_{P.95}$ for ΔG_b is almost symmetric. Indeed, as discussed above in this case of an actual symmetric uncertainty in $\Delta G_{\rm b}$, both the asymptotic-symmetric and profile likelihood methods give the same confdence intervals for both parameters provided that ΔG_b is fitted and then transformed to K_b . However, only the profile likelihood approach produces the same results if K_b is fitted. When using $CI_{P,95}$, there is a perfect correspondence between estimated values and uncertainty interval limits for K_b and ΔG_b ; thus, no matter which parameter is employed as a ftting parameter, the

Table 2 Best fit parameters and uncertainties determined as asymptotic-symmetric $CI_{S.95}$ and profile likelihood CI_{P95} confdence intervals

The directly ftted values are shown in bold

^a Fitting of K_b and calculation of its CI_{P,95} with subsequent transformation to ΔG_b values

^bFitting of ΔG_b and subsequent calculation of K_b

other parameter and its confdence interval can be readily calculated through their mathematical relationship. However, when using $CI_{S,95}$, the estimated values for K_b and ΔG_b are in correspondence, but the uncertainty intervals are not. Furthermore, in some experimental scenarios the upper limit of the affinity may be undefined, e.g., for a very steep titration with few data points contributing to the ft. The profle likelihood confdence intervals will only produce a lower limit in such cases, a highly advantageous presentation over the use of symmetrical intervals. The proile likelihood approach is, thus, more robust in estimating uncertainties and, consequently, the recommended approach.

Conclusions

Although the inherent asymmetry that appears as a result of the measurement error propagating through the data analysis process has been illustrated here with only ITC simulations, they are true for all binding experiments (i.e., to an extent that depends on the nature of the measurement errors and the equations used to analyze each particular method). The presented concepts and procedures for dealing with this asymmetry to determine accurate uncertainties can be extended to any experimental technique used to determine binding afnities or any related quantities (see the distribution of the kinetic rate constant in the supplementary material). The calculation of the confdence intervals can be performed manually following the step-by-step procedure explained above. These can be conveniently carried out in Excel (Kemmer and Keller [2010\)](#page-8-31), but fortunately, commercially available software packages (e.g., Origin, GraphPad, Sedphat) also provide profle likelihood confdence intervals, for all ftting parameters at any confdence level, in a user friendly manner, without the need for complicated calculations. Once the best ft is achieved (with all parameters freely varying), the profle likelihood confdence intervals (at a certain confdence level) for all ftting parameters are readily calculated within just a single step as explained in the Supplementary materials.

Considering the variations in the shape and inherent diference of the uncertainty distribution of the two thermodynamically related parameters K_b and ΔG_b , reporting a symmetric error appears not to be the scientifcally correct way. Reporting 95% confdence intervals removes the artifcial restriction of symmetry and enables more accurate reporting of uncertainty. The statistically sound construction of the profle likelihood confdence intervals, and the perfect agreement shown of CIs obtained when the ftting of ITC data was performed for K_b or ΔG_b and ΔH_b , shows that profle likelihood confdence intervals can be used to report the repeatability of K_b and ΔH_b as retrieved from ITC, or for binding affinities determined by any other method.

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