Measuring experimental cyclohexane-water distribution coefficients for the SAMPL5 challenge

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Small molecule distribution coefficients between immiscible nonaqueuous and aqueous phases—such as cyclohexane and water-measure the degree to which small molecules prefer one phase over another at a given pH. As distribution coefficients capture both thermodynamic effects (the free energy of transfer between phases) and chemical effects (protonation state and tautomer effects in aqueous solution), they provide an exacting test of the thermodynamic and chemical accuracy of physical models without the long correlation times inherent to the prediction of more complex properties of relevance to drug discovery, such as protein-ligand binding affinities. For the SAMPL5 challenge, we carried out a blind prediction exercise in which participants were tasked with the prediction of distribution coefficients to assess its potential as a new route for the evaluation and systematic improvement of predictive physical models. These measurements are typically performed for octanol-water, but we opted to utilize cyclohexane for the nonpolar phase. Cyclohexane was suggested to avoid issues with the high water content and persistent heterogeneous structure of watersaturated octanol phases, since it has which has greatly reduced water content and a homogeneous liquid structure. Using a modified shake-flask LC-MS/MS protocol, we collected cyclohexane/water distribution coefficients for a set of 53 druglike compounds at pH 7.4. These measurements were used as the basis for the SAMPL5 Distribution Coefficient Challenge, where 18 research groups predicted these measurements before the experimental values reported here were released. In this work, we describe the experimental protocol we utilized for measurement of cyclohexane-water distribution coefficients, report the measured data, propose a new bootstrap-based data analysis procedure to incorporate multiple sources of experimental error, and provide insights to help guide future iterations of this valuable exercise in predictive modeling.

Keywords: partition coefficients; distribution coefficients; blind challenge; predictive modeling; SAMPL

I. INTRODUCTION

Rigorous assessment of the predictive performance of physical models is critical in evaluating the current state of physical modeling for drug discovery, assessing the potential impact of current models in active drug discovery projects, and identifying limits of the domain of applicability that require new models or improved algorithms. Past iterations of the SAMPL (Statistical Assessment of the Modeling of Proteins and Ligands) experiment have demonstrated that blind predictive challenges can expose weaknesses in computational methods for predicting protein-ligand binding affinities and poses, hydration free energies, and host-guest binding affinities [1–4]. In addition, these blind challenges have contributed new, high-quality datasets to the community that have enabled retrospective validation studies and

²⁷ data-based parameterization efforts to further advance the ²⁸ current state of physical modeling.

By focusing community effort on the prediction of hydra-30 tion free energies in the first few iterations of this challenge, 31 the SAMPL experiments have now brought physical modeling approaches to the point where they can reliably identify 33 erroneous experimental data [5]. While hydration free energy 34 exercises have shown their utility in improving the state of ₃₅ physical modeling, they are laborious, require specialized 36 equipment no longer found in modern laboratories, are (at 37 least using traditional protocols) limited in dynamic range, 38 and are of questionable applicability in their ability to mimic 39 protein-to-solvent transfer. As a result, no experimental lab-40 oratory has emerged to provide new hydration free energy 41 measurements to sustain this aspect of the SAMPL challenge. 42 We sought to replace this component of the SAMPL challenge 43 portfolio with a new physical property that was easy to mea-44 sure, accessible to multiple laboratories, had a wide dynamic 45 range (in a free energy scale), and better mimicked physical 46 and chemical effects relevant to protein-to-solvent transfer 47 free energies, but was still free of the conformational sam-48 pling challenges protein-ligand binding affinities present. As 49 the measurement of partition and distribution coefficients 50 is now widespread in pharma (due to its relevance in opti-51 mizing lipophilicity of small molecules), we posited that a 52 blind challenge centered around the prediction of distribu-

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provide such a challenge.

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While the measurement of octanol/water distribution coef- 115 property-based log P prediction methods used 96,000 expermental measurements [9]), a number of previously-reported complications in the physical simulation of 1-octanol sugested that this might be too complex for an initial distribution coefficient challenge [10-13], despite some recent reports of success [14]. In particular, water-saturated octanol is very wet, containing 47 ± 1 mg water/g solution [15], and forms complex microclusters or inverse-micelles that create heterogeneous environment that persist for long simulation times [10–13]. For the inaugural distribution coefficient challenge in SAMPL5, we therefore chose to measure cyclohexane/water distribution coefficients. The water content of water-saturated cyclohexane is much lower than watersaturated octanol—0.12 mg water/g solution, approximately 400 times smaller [16–18], and possesses no long-lived heterogeneous structure [19].

The number of freely available sources of cyclohexanewater partition is very limited, and for the purpose of the SAMPL5 distribution coefficient challenge[20], blind data was required. As part of an internship program at Genentech arranged by the coauthors, the lead author was dispatched to work out modifications of a high-throughput shake-flask protocol [21] currently in use for octanol/water distribution coefficient measurements. In particular, the ow dielectric constant of cyclohexane (2.0243) compared o 1-octanol (10.30) [22] and cyclohexane's surprising ability to dissolve laboratory consumables presented some unexpected challenges. In this report, we describe the modified protocol that resulted, and provide suggestions on how it an further be refined for future iterations of the distribution coefficient challenge. Of 95 lead-like molecules with diverse functional groups selected for measurement, we report 53 log D measurements that passed quality controls that were used in the SAMPL5 challenge.

To ensure the reported experimental dataset is useful in assessing, falsifying, and improving computational physical models of physical properties, we require a robust approach to estimating the experimental error (uncertainty in experimental measurements). We explored several procedures for propagating known sources of error in the measurement process into the final reported log distribution coefficients, and report those efforts here. Our primary approach features parametric bootstrap, which allows the use of a physical nodel of the data generating process to sample additional ealizations of the data, using distributions specified in the nodel. These additional realizations are new data points, over which estimates can be calculated. We compared this to nonparametric bootstrap, which can be useful if a physical model can not be constructed. This method generates new data points as well, but it constructs them from selection with replacement from the existing data. We also calculated

tion coefficients—which face many of the same physical and and data. We hope that future efforts to measure cyclohexanechemical effects (such as protonation state [6, 7] and tau- 112 water distribution coefficients can benefit from the model tomer issues [8]) observed in protein-ligand binding—might 113 we have developed, so that this work will also be useful for 114 future challenges.

All code used in the analysis, as well as raw and processed ficients is commonplace (a 2008 benchmark of structure- and data, can be found at https://github.com/choderalab/ 117 sampl5-experimental-logd-data.

Theory of distribution coefficients

The distribution coefficient, D, is a measure of preferential distribution of a given compound (solute) between two im- $_{121}$ miscible solvents at a specified pH, usually specified as $\log D$ in its base-10 logarithmic form,

$$\log D_{\text{solvent1/solvent2}}^{\text{pH}} = \log_{10} \frac{[\text{Solute}]_{\text{solvent1, pH}}}{[\text{Solute}]_{\text{solvent2, pH}}}$$
 . (1)

123 Typically, one solvent is aqueous and buffered at the specified pH (e.g. Tris pH 7.4), while the other is apolar (e.g. 1octanol). At the given pH, the solute may populate multiple protonation or tautomeric states, but the total concentration 127 summed over all states is used in the calculation of concen-128 trations in Equation (1). The total salt concentration of the aqueous phase can also play a role, in case salts can provide 130 stabilization of an ionic state of the ligand in the aqueous phase [23]. Because of this, care must be exercised when comparing distribution coefficients obtained under different 133 experimental conditions.

For the SAMPL5 challenge, we concern ourselves with 135 the cyclohexane-water distribution coefficient, where phosphate-buffered saline (PBS) at pH 7.4 is used for the 137 aqueous phase:

$$\log D_{\text{chx/wat}}^{\text{pH 7.4}} = \log_{10} \frac{[\text{Solute}]_{\text{cyclohexane}}}{[\text{Solute}]_{\text{PBS, pH 7.4}}} \quad . \tag{2}$$

138 Another commonly reported value is the partition coefficient 139 P, which quantifies the relative concentration of the neutral $_{\mbox{\tiny 140}}$ species in each phase, again usually specified in \log_{10} form,

$$\log P_{\text{chx/wat}} = \log_{10} \frac{[\text{Solute}]_{\text{cyclohexane}}^{\text{neutral}}}{[\text{Solute}]_{\text{PBS, pH 7.4}}^{\text{neutral}}} \quad . \tag{3}$$

For ligands with a single titratable site and known p K_a , one $_{142}$ can readily convert between $\log P$ and $\log D$ for a given pH (see, e.g. [23]), but ligands with more complex protonation state effects or tautomeric state effects make accounting for the transfer free energies of all species significantly more 146 challenging.

EXPERIMENTAL METHODS

In the following sections we describe how we measured 110 the arithmetic mean and standard error of the measured 149 cyclohexane/water distribution coefficients for the 53 compounds displayed in Figure 1. The compound selection pro- 203 lenge. After the quality control filtering phase (Section II C), cedure is described in Section II A.

and the procedure is schematically summarized in Figure 2. 209 stereochemistry.

The measured data was subjected to a quality control procedure that eliminated measurements thought to be too unreliable for use in the SAMPL5 challenge (Section II C). Re- 210 maining data were analyzed using a physical model of the experiment by means of a parametric bootstrap procedure. We compared this approach to a nonparametric bootstrap 212 data without bootstrap analysis. In Section II D, we describe each approach. The results for each approach can be found in Table I.

Compound selection

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Compounds were initially selected from a database of 221 9115 lead-like molecules available in eMolecules that were present in the Genentech chemical stores in quantities of over 2 mg, with molecular weights between 150-350 Da. The 224 lower bound on molecular weight was chosen to increase the likelihood of detectability by mass spectrometry, and the pper bound to limit molecular complexity.

We initially chose approximately 88 compounds based on 228 several criteria: 176

- First, we selected 8 carboxylic acid compounds. These were of potential interest for the purpose of the challenge, since it was suspected these could potentially partition along into the cyclohexane phase together with water or cations [23].
- MoKa 2.5 was used to calculate cLogP, cLogD, and pKa values [24, 25]. This version of MoKa was trained with Roche internal data to improve accuracy. We selected 20 compounds with predicted pKa values that would potentially be measurable with a Sirius T3 instrument (Sirius Analytical) so validation with an orthogonal technique (electrochemical titration) could be performed in the future.
- The remaining compounds were divided into 10 equalsize bins that spanned the predicted dynamic range of $\log P$ values (-3.0 to 6.6), and 6 compounds were drawn from each bin, to a total of 60.

This set of 88 total was later reduced to 64 due to the unvailability of some compounds or the inability to detect molecular fragments by mass spectrometry at the time of neasurement. This selection was expanded to include 31 compounds used as internal standards for the previously developed octanol/water assay protocol [21], bringing the total number of compounds for which measurements were performed to 95. These compounds were randomly assigned numerical SAMPL_XXX designations for the SAMPL5 blind chal-

204 the resulting data set contained 53 compounds, which are Distribution coefficient measurements utilized a shake- 205 displayed in Figure 1. Canonical isomeric SMILES represenflask approach based on a LC-MS/MS technique previously 206 tations for the compounds can also be found in Table S1. developed for 1-octanol/water distribution coefficient mea- 207 These were generated using OpenEye Toolkits v2015. June by surements [21]. The approach is described in Section II B, 208 converting 3D SDF files, after manually verifying the correct

Shake-flask measurement protocol for cyclohexane/water distribution coefficients

We adapted a shake-flask assay method from an original approach, and the arithmetic mean and standard error of the 213 octanol/water LC-MS/MS protocol [21] to accommodate the 214 use of cyclohexane for the nonaqueous phase. Our modified protocol is described here, and the procedure is explained schematically in Figure 2.

> The $\log D$ is estimated by quantifying the concentration of a solute directly from two immiscible layers, present as an emulsion in a single vial. Capped glass 1.5 mL auto-injector vials with PTFE-coated silicone septa were used for partitioning, as cyclohexane was found to dissolve polystyrene 96-well plates used in the original protocol.

> For each individual experiment, 10 µL of 10 mm compound in dimethyl sulfoxide (DMSO)² and 5 µL of 200 µм propanolol in acetonitrile (an internal standard) were added to 500 µL cyclohexane³, followed by the addition of 500 µL of phosphate buffered saline (PBS) solution⁴. The ionic components of the buffer were chosen to replicate the buffer conditions used in other in-vitro assays at Genentech. Unlike the original protocol, neither phase was presaturated prior to pipetting.

The solute was allowed to partition between solvents ²³² while the mixture was shaken for 50 minutes using a plate shaker⁵ at 800 RPM, while the vials were mounted in a vial holder and taped down to the sides of the vial holder⁶. The 235 two solvents were then separated by centrifugation for 5 minutes at 3700 RPM in a plate centrifuge, using the plate rotor⁷, with the vials seated in the same vial holder.

Aliquots were extracted from each separated phase using a standard adjustable micropipette, and transferred into a 384-well glass-coated polypropylene plate for subsequent quantification⁸. Cylcohexane wells were first prepared with 242 45 μL of 1-octanol⁹ per well. 5 μL of cyclohexane was ex-

¹ Shimadzu cat. no. 228-45450-91

² DMSO stocks from Genentech compound library

³ ACS grade >99%, Sigma-Aldrich cat. no 179191-2L, batch #00555ME

⁴ 136 mM NaCl, 2.6 mM KCl, 7.96 mM Na2HPO4, 1.46 mM KH2PO4, with pH adjusted to 7.4, prepared by the Genentech Media lab

⁵ Thermo Fisher Scientific, Titer Plate Shaker, model: 4625, Waltham, MA, USA

⁶ Agilent Technologies, Vial plate for holding 54 x 2 mL vials part no. G2255-68700

⁷ Eppendorf, Centrifuge 5804, Hamburg, Germany

^{8 384-}well glass coat plate:Thermo Scientific, Microplate, 384-Well; Webseal Plate; Glass-coated Polypropylene; Square well shape; U-Shape well bottom; 384 wells; 90uL sample volume; catalog number: 3252187

⁹ ACROS Organics, 1-octanol 99% pure, catalog number: AC150630010, Geel, Belgium

on two separate mass spectrometry measurements¹²:

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• The solute is analyzed to identify and select parent 295 SAMPL5_058, SAMPL5_060, and SAMPL5_061. and daughter ions, and optimize ion fragment parameters¹³.

We used a flow rate of 0.2 mL/min, mobile phase of 296 water/acetonitrile/formic acid (50/50/0.1 v/v/v) and 1.5 minutes run time. All parameters were automati- 297 pounds, the fragment identification LC-MS/MS procedure did not yield high intensity fragments, and these could therefore not be measured using the MRM ap- 301 proach.

ent ions and daughter ions of the solute identified in 306 quantification. the previous step. The mass/charge (m/z) intensity (proportional to the absolute number of molecules) is quantified as a function of the retention time¹⁴. Infor-³⁰⁷ mation on the gradient can be found in Supplementary Table 1 of Lin and Pease 2013 [21].

Highest m/z intensity fragments were selected using 5 mм solutions consisting of 50% DMSO, 50% acetonitrile.

one aliquot was prepared, and replicate MRM measurements were performed 3 times per aliquot. The $\log D$ can be calculated from the relative MRM-signals, obtained by integrating the single peak in the MRM-chromatogram, using Equa-281 tion (4).

$$\log \mathrm{D}_{\mathrm{chx/wat}}^{\mathrm{pH\,7.4}} = \log_{10} \frac{\mathrm{MRM\,signal}_{\mathrm{cyclohexane}}/\left[d_{\mathrm{chx}}v_{\mathrm{inj,\,chx}}\right]}{\mathrm{MRM\,signal}_{\mathrm{PBS,\,pH\,7.4}}/v_{\mathrm{inj,\,PBS}}} \quad . \tag{4}$$

²⁴³ tracted from the top phase by micropipette and mixed with ²⁸² The cyclohexane signal is normalized by the dilution fac-45 μ L of octanol in the 384 well plate. 50 μ L of aqueous so- 283 tor of our cyclohexane aliquots, $d_{chx}=0.1$, and the injeclution was subsequently extracted from the bottom phase. 284 tion volume $v_{\sf inj,\,chx}$. As the PBS aliquots were not diluted, The octanol dilution was performed mainly to prevent ac- $_{285}$ this is only normalized by the injection volume $v_{\rm inj,PBS}$. Excumulation of cyclohexane on the C18 HPLC columns¹⁰ that 286 periments were carried out independently at least in dupliwere used. For the aqueous (bottom) phase, the aliquot of 287 cate, repeated from the same DMSO stock solutions. In-50 μL was transferred directly into the 384-well plate, into 288 jection volumes of the MRM procedure were 1 μL for cywells that did not contain octanol. The 384-well plates were 289 clohexane (diluted in octanol), and 2 µL for PBS samples. sealed with using glueless aluminum foil seals¹¹, and frag- 290 For one set of measurements, we carried out 2 additional ment concentrations assayed using quantitative LC-MS/MS. 291 repeat experiments with 2 µL injections for cyclohexane Measuring solute distribution into the two phases depends 292 (diluted in octanol), and 1 µL for PBS. This set included 293 SAMPL5_003, SAMPL5_005, SAMPL5_006, SAMPL5_011, ²⁹⁴ SAMPL5_027, SAMPL5_049, SAMPL5_050, SAMPL5_055,

Quality control

In order to eliminate measurements thought to be too cally stored for further MRM analyses. For several com- 298 unreliable for the SAMPL5 challenge, we utilized a simple quality control filter after MRM quantification. Compounds where the integrated MRM signal within either phase varied between replicates or repeats by more than a factor of 302 10 were excluded from further analysis. We additionally removed compounds that exceeded the dynamic range of the • A separate mass spectrometer is employed using 304 assay because they did not produce a detectable MRM sigmultiple-reaction monitoring (MRM) to select for par- 305 nal in either the cyclohexane or buffer phases during the

Bootstrap analysis

Since our ultimate goal is to compare predicted distribution coefficients to experiment to evaluate the accuracy of 310 current-generation physical modeling approaches, it is critical to have an accurate assessment of the uncertainty in From each solvent phase in the partitioning experiment, 312 the experimental measurement. Good approaches to uncertainty analysis propagate all known sources of experimental error into the final estimates of uncertainty. To accomplish this, we developed a parametric bootstrap model [26] of the experiment based on earlier work [27], with the goal of propagating pipetting volume and technical replicate errors 318 through the complex analysis procedure to estimate their impact on the overall estimated $\log D$ measurements.

> Bootstrap approaches provide new synthetic data sets, denoted as realizations, sampled using some function of the observed data that approximates the distribution that the observed data was drawn from. For each compound that was measured, suppose our data set provides N independent repeats (from the same stock solution, typically 2 or 4), and 3 technical replicates for each repeat (quantitation experiments from each repeat, typically 3). Each realization of the bootstrap process leads to a new synthetic data set, of the same size, from which a set of synthetic distribution coefficients can be computed for the realization. We applied two additional approaches for comparison to assess the performance of our parametric bootstrap method (Section II D 1). One features a nonparametric bootstrap ap-

¹⁰ Waters Xbridge C18 2.130 mm with 2.5 m particles

¹¹ Agilent cat no 24214-001

¹² All LC solvents were HPLC-grade and purchased from OmniSolv (Charlotte, 327 NC, USA)

¹³ This was done using a Shimadzu NexeraX2 consisting of an LC-30AD(pump), SIL-30AC (auto-injector), and SPD-20AC(UV/VIS detector) with Sciex API4000QTRP (MS)

¹⁴ This was done using a Shimadzu NexeraX2 consisting of an LC-30AD(pump), SIL-30AC (auto-injector), and SPD-20AC(UV/VIS detector) with Sciex API4000 (MS)

proach (Section II D 2), which does not include any physical 382 details. The other is a calculation of the arithmetic mean and standard error that is limited to the observed data (Section IID3).

Parametric bootstrap

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We used a parametric bootstrap [28] method to introduce random bias and variance into the data, based on known experimental sources. This procedure allows us to use a model to propagate known uncertainty throughout the procedure [28]. This allows us to better estimate the distribution that the observed data was drawn from, so that more accurate estimates of the means and sample variance can be obtained.

Uncertainties in pipetting operations were modeled based on manufacturer descriptions [29, 30], following the work of Hanson, Ekins and Chodera [27]. Technical replicate variation was modeled by calculating the coefficient of variation (CV) between individual experimental replicates. We then took the mean CV of the entire data set, which was found to be \sim 0.3. As a control, we verified that the CV did not depend on the solvent phase that was measured. We included this in the parametric model by adding a signal imprecision, modeled by a normal distribution with zero mean, and a standard deviation of 0.3. We perform a total of 5 000 realizations of this process, and calculate statistics over all realizations, such as the mean (expectation) and standard deviation (estimate of standard error) for each measurement.

Nonparameteric bootstrap

A traditional nonparametric Monte Carlo procedure was 407 calculation (Section IID3) are plotted in Figure 3c. applied to resample data points[26]. This approach can es- 408 ment, to generate a new set of data points with size equal 411 to the observed data set. Nonparametric bootstrap can be a 412 useful approach if larger amounts of data are available, and implemented the procedure in two stages:

- 1. A set of N repeats is drawn with replacement from the 417 original set of measured repeats.
- 2. For each of the repeats, we similarly draw a set of 3 420 technical replicates from the original set of technical 421 replicates.

This yields a sample data set with the same size as the originally observed data (N repeats, with 3 replicates each). We perform a total of 5 000 realizations of this process, and calculate statistics over all realizations, such as the mean (expectation) and standard deviation (estimate of standard error) for each measurement.

Arithmetic mean and sample variance

We calculated the arithmetic mean over all replicates and ³⁸⁴ repeats, and estimated the standard error from the total of 6 or 12 data points, to compare to our bootstrap estimates. 15

Kernel densities

As a visual guide, in Figure 3 data are plotted on top of an estimated density of points. This density was calculated using kernel density estimation [31], which is a nonparametric way to estimate a distribution of points using kernel functions. Kernel functions assign density to individual points in a data set, so that the combined set of data points reflects ³⁹³ a distribution of of the data. We used the implementation available in seaborn 0.7.0 [32]. We used a product of Gaussian kernels, with a bandwidth of 0.4 for $\log D$ and 0.3 for the standard error. To prevent artifacts such as negative density estimates for the standard errors, they were first transformed by the natural logarithm \ln , and the results were then con-399 verted back into standard errors by exponentiation.

DISTRIBUTION COEFFICIENTS

The $\log D$ values and their uncertainties for the 53 small 402 molecules that passed quality controls are presented in Table I. In the following two sections, we describe the differences between the analysis results in more detail.

Mean and standard errors in $\log D$

The results from the arithmetic mean and sample variance

Despite the compound selection effort, the distribution timate the distribution that the observed data was drawn $_{409}$ of data along the $\log D$ -axis is less dense in the region -1 to from by resampling from the observed data with replace- $_{\scriptscriptstyle 410}$ 0 log units. The data outside this region seems to be centered around -2 log units, or around 1 log unit. We could attribute this distribution of data to coincidence, though this way warrant future investigations into systematic errors. Usa detailed physical model of the experiment is absent. We 44 ingthe arithmetic mean of the combined repeat and replicate measurements (Section II D 3) the distribution coefficients measured spanned from -3.9 to 2.5 log units.

> The $\log D$ measurements distribution appears bimodal along the uncertainty axis. A subset of mostly negative $\log D$ values (Figure 3c) has a smaller estimated standard deviation, though this is not the case for the majority of negative $\log D$ values. The average standard error, rounded to 1 significant 422 figure, is 0.2 log units for the arithmetic mean calculation.

¹⁵ For the purpose of the D3R/SAMPL5 workshop, we originally erroneously reported the standard deviation $\sqrt{3}$ instead of the standard error $\sqrt{3}$. The factor of $\sqrt{3}$ corrects the sample standard deviation across all MRM measurements for the correlation between the 3 replicate measurements belonging to a single independent experimental repeat.

Bootstrap results

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Estimates of the $\log D$ span the range between -3.9 to 2.6 $\,^{_{478}}$ duce the bandwidth of the method. log units, using either of the two bootstrap approaches (Sec- 479 estimated standard errors. When applying our bootstrap procedures (Section II D 1 and Section II D 2), we see an upwards shift in the uncertainties, compared to the sample variance calculations. The nonparametric approach yields an average uncertainty of 0.3 log units. The parametric approach yields an average uncertainty of 0.4 log units. The parametric bootstrap suggests that by propagating errors such as the cyclohexane dilution, and the replicate variability into the model, some of the observed low uncertainties might be an artifact of the low number of measurements. This suggest used. that simply calculating the arithmetic mean, and standard error of all measured data might not reliably capture the error in the experiments. We also note that for certain compounds, 493 bootstrap distributions exhibit multimodal character and as such, standard errors might not accurately capture the 494 the supplementary information.

Using the parametric scheme, we see an average shift of uncertainties to larger values compared to the nonparametric 499 bootstrap. The density estimate suggests we should expect a 500 462 point).

DISCUSSION

Solvent conditions

It is important to consider the fact that different cosolvents 518 466 may have on the measured values. The solutions contained approximately 1% DMSO, as well as approximately 0.5% ace- 519

476 solvents such as DMSO and acetonitrile. This would make 477 experiments much more laborious, and would therefore re-

One of the things we could not completely account for in tion II D 1 and Section II D 2). The $\log D$ estimates do not differ 480 the model was the exact ratio of cyclohexane and PBS sosignificantly from the arithmetic mean calculations. The dif- 481 lution. We attempted to account for volumetric errors from ference between the results is seen when we compare the 482 pipetting by performing independent repeat experiments, although this may still leave some systematic error. Cyclo-484 hexane is also a volatile compound, especialy compared to water. For comparison, the vapor pressure of cyclohexane 486 is 97.81 torr [33], versus 23.8 torr for water [34]. It is possible that evaporation may occur, which could lead to a systematic ⁴⁸⁸ overestimation of the cyclohexane volume. For future investigations, it may be fruitful to study the evaporation so that 490 this can be accounted for in the model of the experiment, and to take note of systematic bias in the pipette models

Compound detection limits

Calculations using COSMO-RS software[35] suggested a full extent of the experimental uncertainty. We provide the $_{ t 495}$ systematic underestimation of $\log D$ values in the negative bootstrap sample distributions of the parametric model in $_{496}$ log unit range, in particularly past a $\log D$ of -2. Without further experimental investigation, we can not draw definite conclusions as to whether this is the case, or if so, where the source of the systematic error lies.

One possibility that may cause an artificial reduction of lower bound to the error that we have now incorporated into $_{\scriptscriptstyle 501}$ the dynamic range—especially at high log D values—is the the analysis. Not every compound shows the same increase potential for MS/MS detector saturation at high ligand conuncertainty, though if we compare the two bootstrap ap- 503 centrations. Previous work (Figure 2 from [21]) examined proaches, results are similar above this empirically observed 504 detector saturation effects, finding it possible to reach sufower bound. The nonparametric approach returns higher ы ficiently high compound concentrations (generally ≥10 µм) uncertainties for some data on average, but estimates lower 506 that MRM is no longer linear in compound concentration for uncertainties for some as well. It can be concluded that the 507 that phase. This work also found that different compounds error would typically be underestimated without the use of 508 reach detector saturation at different concentrations [21], in a bootstrap approach. Without a physical model, a nonpara- 509 principle requiring an assessment of detector saturation to metric approach might still underestimate errors due to the 510 be performed for each compound. While we could not delimited sample size for each measurement (either 2 or 4 fully 511 duce obvious signs of detector saturation in our LC-MS/MS independent repeats, and a total of 3 replicates per data size chromatograms, these effects could be mitigated by performing a dilution series of the aliquots sampled from each phase of the partitioning experiment to ensure detector response is linear in the range of dilutions measured. This may also reveal whether compound dimerization may be a complicating factor in quantitation.

Experimental design considerations

In order to adjust our experimental setup, we had to switch tonitrile. Further work would benefit from a comparison with $_{\scriptscriptstyle 520}$ away from using polystyrene 96 well plates, as these were experiments starting from dry stocks, and thereby not adding 521 dissolved by cyclohexane. We attempted the use of glass extra solvents. This would eliminate DMSO and acetonitrile, 522 inserts, and glass tubes but these were too narrow and proby dispensing compound directly into either cyclohexane, or $_{\scriptscriptstyle 523}$ vided insufficient mixing when shaken. We switched to glass the mixture of cyclohexane and PBS. In this case, care should 524 vials because their larger diameter provides improved mixbe taken that all compound is dissolved. If found to be nec- 525 ing when shaken. For future work, we would recommend essary, we could then consider starting all experiments from 526 the use of glass coated plates, which have the automation dry compound stocks, to entirely eliminate effects from co- 527 advantages of the plates used in the original protocol [21].

Plate seals need to be selected carefully. We experimented 586 ent volumes of 10 mm ligand stocks. This would help detect

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umulation of unknown origin at the end of each UV chro- 597 more on these. matogram. Accumulation was reduced by injecting less cyclohexane. As a result, we diluted the cyclohexane with 1octanol for the experiments described here, and ran blank 598 injections containing ethanol between batches of 64 measurements to ensure the column was clean.

Another change to the protocol that we would like to conider for future measurements is to optimize the time spent equilibrating the mixture. In this work, we separated phases ia centrifugation and sampled aliquots for concentration neasurement within minutes. The post-centrifugation time prior to sampling aliquots could be extend to 24 hours to allow for more equilibration for the solute between phases. This may have a downside, since we would have to consider the effects that may follow if compounds prefer to be in the interface-region between cyclohexane and water, or water and air. These could cause high local concentrations, introducing a dependency of the results on exactly which part of the solution aliquots are taken from. We can get around this by only taking samples from the pure cyclohexane and aqueous regions, avoiding the interfaces. This way, we still get the ight distribution coefficients for partitioning between bulk phases even if some compound is lost to the interfaces.

It may be worthwhile to consider other effects of pipetting perations on the procedure. Some compounds could potentially stick to the surface of pipettes, or glass surfaces. This could adversely affect our measurements by changing local concentrations.

While cyclohexane and water have much lower mutual sol- 624 proaches can be taken to resolve part of this issue. Among the ubility than octanol, it is still possible that this affects the 625 options are the use of statistical tests, such as the bootstrapmeasurement.

difficulties with respect to interpretation of the data. This 628 as a parametric bootstrap) or extract uncertainty already was mainly because we could not separate out many effects 629 available in the data (such as nonparametric bootstrap). The from the data that could affect the interpretation. We ob- 630 parametric approaches can be improved in terms of the physserved that for the many effects used to explain discrepan- 631 ical models that are used to analyze the data. These models cies between model and experiment, none of those could 632 should ideally include all known sources of error, such as dimerization, the transfer of water or ions into the cyclohex- $_{\scriptscriptstyle 635}$ many other conditions that could affect the results. ane phase, from changes in the $\log D$ values based on the $_{ text{ iny 636}}$ molecular properties itself.

assays are carried out at multiple final concentrations of 639 incorporate data and propagate uncertainty from multiple

with silicone sealing mats, but these absorbed significant 587 dimerization issues, and may help account for issues with quantities of cyclohexane. We also had to discontinue use of 588 detector oversaturation. Note that the absolute errors in aluminum seals that contained glue, since the glue is soluble 589 these stock volumes will not be critical, since the measurein cyclohexane and would contaminate LC-MS/MS measure- 590 ments rely on the relative measurement between the two ments. In the end, we used aluminum PlateLoc heat seals 591 phases. We could build models that allow for extrapolation and glass coated 384 well plates to circumvent these issues. 592 to the infinite dilution limit, which should then provide sim-Sensitivity also suffered due to the need to dilute cyclohex- 593 pler test cases for challenge participants to reproduce. On ne in octanol to prevent its accumulation on C18 columns 594 the opposite end, it may be useful to even investigate wavs sed in the LC-MS/MS phase of the experiment. Trial injec- 595 to design an experimental set that represents these type of ions on a separate system and chromatograms showed ac- 596 issues, such as compound dimerization, so that we can focus

Uncertainty analysis

We hope the experience from this challenges will lay the groundwork for improving the reliability of data sets regarding the physical properties that we as a modeling community rely on. Many computational studies are limited in the amount of high-quality experimental data that they have access to. Unfortunately, most data is taken straight from literature tables, without much thought being spent on the data collection process. By performing the experimental part of the SAMPL5 challenge we were in the position to provide new data to the modeling community, with an opportunity to decide on an analysis strategy that suits modeling applications. This not only allows for blind validation of physics-based models, but also a re-evaluation of the exact properties a data set should have to provide utility to the modeling community. An important fact that we feel needs reemphasizing is that experimental data is limited in utility by the method that was used to analyze it.

Among the lessons learned from this challenge, we would recommend that future challenges would also feature a rigorous statistical treatment of the experimental analysis procedure, ideally going beyond these initial efforts. One crucial part of the analysis procedure is obtaining not only accurate estimates of the observable, but also its uncertainty. We also consider that assay results might be less variable if 622 As indicated in our data set, standard error estimates from re presaturated water and cyclohexane before mixing them. 623 small populations may underestimate the error. Several apping methods we applied in this work. These can help us both The computational end of the challenge featured some 627 propagate information on uncertainty into the model (such asily be tested with the current state of the data set. It may 👸 pipetting errors, evaporation of solvent, errors in integration e possible to discombobulate matters such as compound 🚳 software, fluctuations in temperature, pressure and likewise

Another approach would be to perform statistical inference on the data set, to provide uncertainty estimates from For future challenges, we would recommend that these 638 the data itself. The model structure can provide ways to the ligand in the assay. This could be achieved using differegree experiments. Common parameters, such as variance in measurements between experiments could be inferred from com- $_{_{693}}$ ane/water $\log D$ measurements in the same manner as the bining the entire data set into one model. When prior knowl- 694 original octanol/water assays, though further optimizations edge on the experimental parameters is available, a Bayesian 695 are needed to reach the same level of throughput. Cyclomodel can be used to effectively infer this type of uncertainty 696 hexane did pose several challenges for experimental design, from the data, and use it to propagate the error into $\log D_{_{697}}$ such as the need for different container types, and the poestimates. Distinctions could be made between an objec- 698 tential accumulation of substrate on reversed phase HPLC tive treatment of the problem, or an empirical Bayesian ap- 699 columns. proach, where prior parameters are derived from the data. 700 One could use a maximum a posteriori (MAP) probability ap- 701 and dimerization might need to be accounted for in order proach to obtain an estimate of one of the modes of the 702 to reproduce experiments. This challenge taught us conparameter distribution. This has obvious downsides when $_{703}$ siderations that should be made on the experimental side. posterior densities are multimodal, and in such a case, one 704 Cases where dimerization were pointed out as possible reamay wish to estimate the shape of the entire posterior distri- 705 son for discrepancy between experiment and model, could bution instead. An approach like Markov chain Monte Carlo $_{706}$ only be hypothesized from the modeling end and not tested could provide such estimates, and will allow for calculation 🔞 experimentally. Issues with detector saturation could also of credible intervals. MCMC methods can be computationally 708 be affecting the overall quality of the data set. Future experiintensive compared to MAP, though if the resulting posterior 709 ments would benefit from more rigorous protocols, such as fortunately, we were unable to construct a Bayesian model 711 experimental components. of the experiments within our time constraints. We would encourage future challenges to make an attempt at creating 713 general, use physical models of experiments in the analysis a Bayesian model, since this would allow for robust inference of all experimental parameters.

Funding future challenges

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The execution of this work would not have been possi-720 ble without the resources provided by Genentech. Access 721 666 a rich library of compounds onsite allowed us to select 722 dataset that was both challenging and useful for the pur- 723 ing the information gain by using the model. oses of the SAMPL challenge. At the same time, the instrunentation provided us with the bandwidth to perform many measurements. Rapid redesign of experiments by trial and 🛛 🖂 libraries, and the equipment to perform the experiments is error, as a result of the difficulties with cyclohexane compatibility of laboratory consumables and equipment, would not 728 rations with Genentech scientists were important in solving have been possible without the expertise shared by Genen- 729 many technical challenges. The collaboration between industech scientists and the opportunities to do many measure- 730 try and academics was not only fruitful, but fundamental in 675

tinued collaboration between industry and academia. Aca- 733 hard to come by without industry resources. At the same demic groups can partner with industry groups to pair avail- 734 time, the need and expertise in investigating these challengable skilled academic labor (graduate students and postdoc- 735 ing physical chemical problems provided by the community, and compound libraries. The graduate student industry in- 737 tial in turning this challenge into a success. We welcome ternship model proved to be a particularly successful ap- 738 such future efforts and collaborations, as it is apparent that proach, with measurements for a blind challenge providing 739 both experimental and computational approaches for oba well-defined, limited-scope project with clear high value to 740 taining $\log D$ estimates for small molecules, would benefit the modeling community.

CONCLUSION

The experimental data provided by this study was very 743 ward. We showed that it was possible to perform cyclohex- 147 grated MRM data including excluded data points are avail-

Many details, such as protonation states, tautomer states, complicated, a MAP estimate can give poor results. Un- 700 measurements at multiple concentrations, and models of all

We recommend that future challenges, and experiments in of experimental uncertainty. These should be part of the analysis procedure, but also in experimental design. These will reveal abnormalities in data more clearly.

We recommend that future challenges look into the use of bootstrap models such as those considered here. Additionally, the use of Bayesian inference methods, that allow the incorporation of prior information should lead to a more robust estimate of experimental uncertainty. They will allow for joint inference on multiple experiments, thereby increas-

Lastly, the sponsoring of this internship by Genentech was ₇₂₅ fundamental to generating this data. Access to compound crucial to the design and execution of a study. Close collaboestablishing standardized challenges for the modeling field. Future iterations of this challenge would benefit from con- T32 The amount of data we were able to gather would have been oral researchers) with specialized measurement equipment 736 and the forum provided by the SAMPL challenge was essen-₇₄₁ from further optimization.

SUPPLEMENTARY INFORMATION

Canonical isomeric smiles for each of the measured comuseful for hosting the first small-molecule distribution coeffi- 744 pound are available in Table S1. An sdf file containing all comcient challenge in the context of SAMPL. It revealed that $\log D_{745}$ pounds, including the measured distribution coefficients is prediction, as well as measurement, is not always straightfor- 746 available as part of the supplementary information. Inteable as part of the supplementary information. Bootstrap 763 distributions from the parametric bootstrap samples for each compound are provided. We also include a csv file containing a full list of SAMPL5_XXX identifiers and canonical isomeric smiles, including unmeasured compounds.

VII. FINANCIAL SUPPORT

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CONFLICT OF INTEREST STATEMENT

DLM and JDC are members of the Scientific Advisory Board 761 762 for Schrödinger, LLC.

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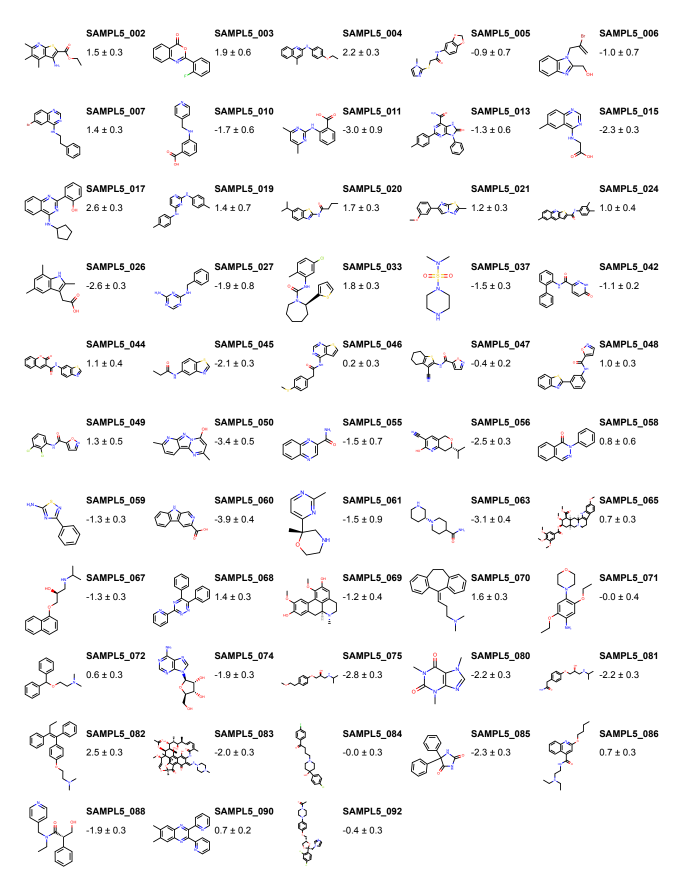


FIG. 1: Molecules and corresponding measured log distribution coefficients for measurements that passed quality controls. Log D measurements are reported as expectation ± standard errors, calculated using our parametric bootstrap method (Section | | D).

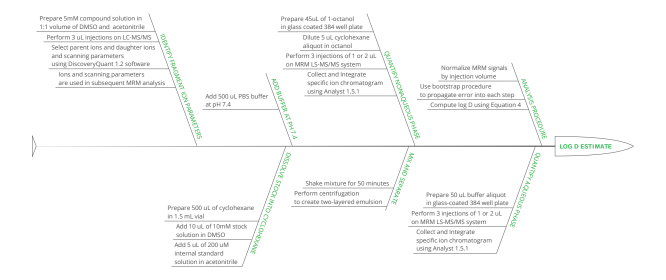
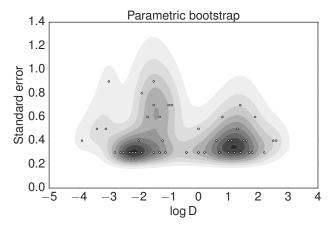


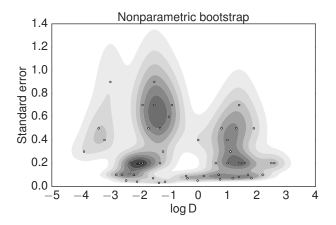
FIG. 2: Illustration of the shake-flask procedure used for cyclohexane-water distribution coefficient measurements.

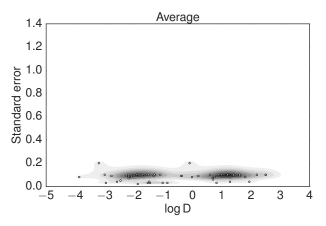
TABLE I: **Log distribution coefficient measurements and standard errors.** Estimates of log distribution functions and their associated standard errors are described for parametric bootstrap (Section II D 1), nonparametric bootstrap (Section II D 2), and arithmetic mean and corrected sample variance (Section II D 3).

	Uncertainty analysis method			
	Вос	otstrap	Arithmetic mean	
Compound ID	Parametric	Nonparametric	Standard error	
SAMPL5_002	1.5 ± 0.3	1.5 ± 0.2	1.4 ± 0.1	
SAMPL5_003	1.9 ± 0.6	1.9 ± 0.5	1.94 ± 0.04	
SAMPL5_004	2.2 ± 0.3	2.2 ± 0.1	2.2 ± 0.1	
SAMPL5_005	-0.9 ± 0.7	-0.9 ± 0.7	-0.86 ± 0.03	
SAMPL5_006	-1.0 ± 0.7	-1.0 ± 0.6	-1.02 ± 0.03	
SAMPL5_007	1.4 ± 0.3	1.39 ± 0.08	1.38 ± 0.09	
SAMPL5_010	-1.7 ± 0.6	-1.7 ± 0.5	-1.7 ± 0.1	
SAMPL5_011	-3.0 ± 0.9	-3.0 ± 0.9	-2.96 ± 0.03	
SAMPL5_013	-1.3 ± 0.6	-1.3 ± 0.5	-1.5 ± 0.1	
SAMPL5_015	-2.3 ± 0.3	-2.3 ± 0.2	-2.25 ± 0.09	
SAMPL5_017	2.6 ± 0.3	2.6 ± 0.2	$2.5~\pm0.1$	
SAMPL5_019	1.4 ± 0.7	1.4 ± 0.7	$1.2~\pm0.1$	
SAMPL5_020	1.7 ± 0.3	1.7 ± 0.1	$1.6~\pm0.1$	
SAMPL5_021	$1.2\ \pm0.3$	1.18 ± 0.07	1.2 ± 0.1	
SAMPL5_024	1.0 ± 0.4	1.0 ± 0.4	$1.0~\pm0.1$	
SAMPL5_026	-2.6 ± 0.3	-2.6 ± 0.1	-2.58 ± 0.04	
SAMPL5_027	-1.9 ± 0.8	-1.9 ± 0.7	-1.87 ± 0.02	
SAMPL5_033	1.8 ± 0.3	1.82 ± 0.07	1.80 ± 0.08	
SAMPL5_037	-1.5 ± 0.3	-1.54 ± 0.07	-1.53 ± 0.03	
SAMPL5_042	-1.1 ± 0.2	-1.13 ± 0.04	$-1.1 \hspace{0.2cm} \pm \hspace{0.2cm} 0.1$	
SAMPL5_044	$1.1~\pm0.4$	$1.1\ \pm0.3$	$1.0~\pm0.1$	
SAMPL5_045	-2.1 ± 0.3	-2.09 ± 0.04	-2.09 ± 0.08	
SAMPL5_046	0.2 ± 0.3	0.19 ± 0.09	$\textbf{0.20} \pm \textbf{0.09}$	
SAMPL5_047	-0.4 ± 0.2	-0.37 ± 0.07	-0.37 ± 0.09	
SAMPL5_048	1.0 ± 0.3	1.0 ± 0.2	$0.9~\pm0.1$	
SAMPL5_049	1.3 ± 0.5	1.3 ± 0.5	1.28 ± 0.04	
SAMPL5 050	-3.4 ± 0.5	-3.4 ± 0.5	-3.2 ± 0.2	
SAMPL5_055	-1.5 ± 0.7	-1.5 ± 0.7	-1.48 ± 0.04	
SAMPL5_056	-2.5 ± 0.3	-2.46 ± 0.05	-2.46 ± 0.05	
SAMPL5_058	0.8 ± 0.6	$0.8~\pm0.5$	0.82 ± 0.03	
SAMPL5_059	-1.3 ± 0.3	-1.34 ± 0.03	-1.33 ± 0.09	
SAMPL5_060	-3.9 ± 0.4	-3.9 ± 0.3	-3.87 ± 0.08	
SAMPL5_061	-1.5 ± 0.9	$-1.5~\pm0.9$	-1.45 ± 0.03	
SAMPL5 063	-3.1 ± 0.4	-3.2 ± 0.4	$-3.0 \ \pm 0.1$	
SAMPL5 065	0.7 ± 0.3	0.7 ± 0.1	0.69 ± 0.07	
SAMPL5_067	-1.3 ± 0.3	-1.3 ± 0.2	-1.3 ± 0.1	
SAMPL5 068	1.4 ± 0.3	1.4 ± 0.2	1.41 ± 0.09	
SAMPL5_069	-1.3 ± 0.4	-1.2 ± 0.3	-1.3 ± 0.1	
SAMPL5_070	1.6 ± 0.3	1.6 ± 0.2	1.61 ± 0.09	
SAMPL5_071	-0.0 ± 0.4	-0.0 ± 0.4	-0.1 ± 0.2	
SAMPL5_072	0.6 ± 0.3	0.6 ± 0.2	0.6 ± 0.1	
SAMPL5 074	-1.9 ± 0.3	-1.9 ± 0.2	-1.9 ± 0.1	
SAMPL5_075	-2.8 ± 0.3	-2.8 ± 0.1	-2.77 ± 0.09	
SAMPL5_080	-2.2 ± 0.3	-2.2 ± 0.1	-2.18 ± 0.07	
SAMPL5_081	-2.2 ± 0.3	-2.2 ± 0.1	-2.19 ± 0.09	
SAMPL5_082	2.5 ± 0.3	2.5 ± 0.2	2.5 ± 0.1	
SAMPL5_083	-2.0 ± 0.3	-2.0 ± 0.2	-1.9 ± 0.1	
SAMPL5 084	-0.0 ± 0.3	-0.02 ± 0.05	-0.02 ± 0.08	
SAMPL5_085	-2.3 ± 0.3	-2.3 ± 0.2	-2.2 ± 0.1	
SAMPL5_086	0.7 ± 0.3	0.7 ± 0.1	0.70 ± 0.06	
SAMPL5_088	-1.9 ± 0.3	-1.9 ± 0.2	-1.9 ± 0.1	
SAMPL5_090	0.7 ± 0.2	0.75 ± 0.2	0.76 ± 0.08	
SAMPL5_090	-0.4 ± 0.2	-0.41 ± 0.09	-0.39 ± 0.09	
3/1111 L3_U3Z	∪.¬ ⊥ ∪.3	-0.41 ⊥ 0.09	_0.39 ± 0.09	



(a) **Parametric bootstrap** (Section II D 1). Standard error estimates calculated by using a parametric bootstrap (circles) and a kernel density estimate (contours) of the entire set.





(b) **Nonparametric bootstrap** (Section II D 2). Standard error estimates calculated using a nonparametric bootstrap (circles), and a kernel density estimate (contours) of the entire set.

(c) **Arithmetic mean and sample variance** (Section II D 3). Standard error estimates calculated using corrected sample variance (circles), and a kernel density estimate (contours) of the entire set.

FIG. 3: Joint kernel density estimates of log distribution coefficient (log D) measurements and measurement error estimates. $\log D$ measurements are plotted with their corresponding estimated standard errors (circles) for the three analysis approaches described in Section II D. A kernel density estimate (contours, described in Section II E) is shown to highlight the differences in error estimates for the different methods.

X. SUPPLEMENTARY INFORMATION

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A. Compound identifiers

TABLE S1: All of the compounds that were selected, and for which $\log D$ was obtained.

Molecule ID Canonical Isomeric SMILES	eMolecules ID (if available)
SAMPL5_002 CCOC(=0)c1c(c2c(c(c(nc2s1)C)C)C)N	1254130
SAMPL5_003 c1ccc2c(c1)c(=0)oc(n2)c3ccccc3F	1231787
SAMPL5_004 CCOc1ccc(cc1)Nc2cc(c3ccccc3n2)C	1221528
SAMPL5_005 Cn1ccnclSCC(=0)Nc2ccc3c(c2)OCO3	1363085
SAMPL5 006 C=C(Cn1c2cccc2nc1C0)Br	2118862
SAMPL5 007 c1ccc(cc1)CCNc2c3cc(ccc3ncn2)Br	1373587
SAMPL5 010 c1cc(cc(c1)NCc2ccncc2)C(=0)O	1491855
SAMPL5_011_Cc1cc(nc(n1)Nc2cccc2C(=0)O)C	542592
SAMPL5_013_Cc1ccc(cc1)c2nc(c3c(n2)n(c(=0)[nH]3)c4ccccc4)C(=0)N	16095985
SAMPL5_015_Cc1ccc2c(ct)c(ncn2)NCC(=0)O	1355949
SAMPL5_017_c1ccc2c(c1)c(nc(n2)c3ccccc3O)NC4CCCC4	2425478
SAMPL5 019 Cc1ccc(cc1)Nc2ccrc(n2)Nc3ccc(cc3)C	43423819
SAMPL5 020 CCCC(=0)Nc1nc2ccc(cc2s1)C(C)C	5522978
SAMPL5 021 Cc1nn2cc(nc2s1)c3cccc(c3)OC	31467689
SAMPL5	16329490
SAMPL5_026_Cc1cc(c2c(c1)c(c([nH]2)C)CC(=0)O)C	703690
SAMPL5_027_clecc(cc1)CNc2ncnc(n2)N	2483781
SAMPL5_033_Cc1ccc(cc1NC(=0)N2CCCCC[C@@H]2c3cccs3]Cl	16363773
SAMPL5_037_CN(c)S(=0)(=0)NICCNCC1	833953
SAMPL5_042_c1ccc(cc1)c2ccccc2NC(=0)c3ccc(=0)[nH]n3	1552842
SAMPL5_044_c1ccc2c(c1)cc(c(=0)o2)C(=0)Nc3ccc4c(c3)scn4	4987019
SAMPL5_045_CCC(=0)Nc1ccc2c(c1)ncs2	12474692
SAMPL5_046_CScIccc(cc1)CC(=0)Nc2c3ccsc3ncn2	12046880
SAMPL5_047_clcnoc1C(=0)Nc2c(c3c(s2)CCCC3)C#N	5627778
SAMPL5_048_c1ccc2c(c1)nc(s2)c3cccc(c3)NC(=0)c4ccno4	5627798
SAMPL5_049 c1cc(c(c(c(1)C1)C1)NC(=0)c2ccno2	5627856
SAMPL5_050_Cc1ccc2c(n1)nn3c2nc(cc30)C	5663556
SAMPL5_055_c1ecc2c(c1)ncc(n2)C(=0)N	3800934
SAMPL5_056 CC(C)[C@@H]1Cc2c(cc(c(n2)0)C#N)C01	
SAMPL5_058_c1ccc(cc1)n2c(=0)c3ccccc3cn2	3730323
SAMPL5_059_c1ccc(cc1)c2nc(sn2)N	711981
SAMPL5_060_c1ccc2c(c1)c3cc(ncc3[nH12]C(=0)O	42618372
SAMPL5_061_CcInccc(n1)[C@@]2(CNCCO2)C	43241882
SAMPL5 063 C1C[C@H](CNC1)N2CCC(CC2)C(=0)N	38498425
SAMPL5_065 COc1ccc2c(c1)[nH]c3c2CC[N@]4[C@@H]3C[C@H]5[C@@H]((C@H]([C@H])([C@H]5C(=0)OC)OC)OC(=0)c6cc(c(c6)OC)OC)OC	
SAMPL5_067_CC(C)NC[C@@H](COc1cccc2clcccc2)O	
SAMPL5_068_c1ccc(cc1)c2c(nnc(n2)c3ccccn3)c4ccccc4	
SAMPL5_069_C[N@]1CCc2cc(c(c-3c2[C@@H]1Cc4c3cc(c(c4)O)OC)O	
SAMPL5 070 CN(C)CCC=C1c2ccccc2CCc3c1cccc3	
SAMPL5_071_CCOc1cc(c(cc1N2CCOCC2)OCC)N	
SAMPL5_072_CN(c)CCOC(c1ccccc1)c2ccccc2	
SAMPL5_074_c1nc(c2c(n1)n(cn2)[C@H]3[C@@H]([C@H]((C@H](03)CO)O)N	
SAMPL5_075_CC(C)NC[C@@H](COc1ccc(cc1)CCOC)O	
SAMPL5_083 C[C@H]/(C=C/C=C(\C(=0)NC\2=C(C3=C(C)=C4C(=C3C(=0)/C2=C/NN5CCN(CC5)C)C(=0)[C@](04)(0/C=C/[C@@H]([C@@H]([C@@H]([C@@H]([C@@H]([C@@H]([C@@H]([C@@H]([C@@H]([C@](C3)C(=0)/C)C)C)C)C)C)C)C)C)C)C)C)C)C)C)C)C)C)C	С
SAMPL5_080_Cn1cnc2c1c(=0)n(c(=0)n2C)C	
SAMPL5_081_CC(C)NC[C@H](COc1ccc(cc1)CC(=0)N)O	
SAMPL5_082 CC/C(=C(\cleccce1)/c2ccc(cc2)OCCN(C)C)/c3ccccc3	
SAMPL5_084_c1cc(ccc1C(=0)CCCN2CCC(CC2)(c3ccc(cc3)Cl)O)F	
SAMPL5 085 c1ccc(cc1)C2(C(=0)NC(=0)N2)c3ccccc3	
SAMPL5_086 CCCCOc1cc(c2cccc2n1)C(=0)NCCN(CC)CC	
SAMPL5_088_CCN(Cc1ccncc1)C(=0)TC@H1(C0)c2ccccc2	
SAMPL5_090 Cc1cc2c(cc1C)nc(c(n2)c3ccccn3)c4ccccn4	
SAMPL5_092_CC(=0)N1CCN(CC1)c2ccc(cc2)OC[C@H]3CO[C@](O3)(Cn4ccnc4)c5ccc(cc5Cl)Cl	
20 OC OCCUPANT OF THE PROCESS OF THE	