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Muddana, H. S., Varnado, C. D., Bielawski, C. W., Urbach, A. R., Isaacs, L., Geballe, M. T., & Gilson, M. K. (2012). Blind prediction of host-guest binding affinities: A new SAMPL3 challenge. *Journal of Computer-Aided Molecular Design*, 26(5), 475-487. <https://doi.org/10.1007/s10822-012-9554-1>

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Published in final edited form as:

J Comput Aided Mol Des. 2012 May ; 26(5): 475–487. doi:10.1007/s10822-012-9554-1.

Blind prediction of host-guest binding affinities: A new SAMPL3 challenge

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Abstract

The computational prediction of protein-ligand binding affinities is of central interest in early-stage drug-discovery, and there is a widely recognized need for improved methods. Low molecular weight receptors and their ligands—i.e. host-guest systems – represent valuable test-beds for such affinity prediction methods, because their small size makes for fast calculations and relatively facile numerical convergence. The SAMPL3 community exercise included the first ever blind prediction challenge for host-guest binding affinities, through the incorporation of 11 new host-guest complexes. Ten participating research groups addressed this challenge with a variety of approaches. Statistical assessment indicates that, although most methods performed well at predicting some general trends in binding affinity, overall accuracy was not high, as all the methods suffered from either poor correlation or high RMS errors or both. There was no clear advantage in using explicit vs. implicit solvent models, any particular force field, or any particular approach to conformational sampling. In a few cases, predictions using very similar energy models but different sampling and/or free-energy methods resulted in significantly different results. The protonation states of one host and some guest molecules emerged as key uncertainties beyond the choice of computational approach. The present results have implications for methods development and future blind prediction exercises.

INTRODUCTION

The ability to accurately predict the binding affinity of a ligand to a target protein could dramatically reduce the burden of chemical synthesis and experimental affinity assays to a more manageable number of compounds and, more importantly, accelerate the discovery of needed medications. While computational screening methods such as docking and empirical scoring methods are routinely used in lead identification and enrichment [1], there is still a need for higher-level computational methods able to reliably rank candidate ligands by affinity; i.e., by their binding free energy [2, 3]. Such methods would be valuable in a secondary, higher-precision stage of virtual screening, as well as in the computer-guided optimization of lead compounds.

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One of the central challenges in developing such methods is validation. For one thing, testing detailed computational methods on protein-ligand systems can be quite time-consuming. Furthermore, because proteins are large and flexible, it is difficult to be sure that a calculation has sampled all thermodynamically relevant conformations; i.e., that the calculation has converged. One valuable alternative is to test whether the method can accurately reproduce experimental solvation free energies of small molecules, as discussed elsewhere in this issue. However, solvation free energies can be measured only for molecules that are volatile enough to be detectable in the gas phase, and such molecules tend to be relatively small and nonpolar. As a consequence, solvation free energies cannot be used to test how well a solvation model handles water in the complex microenvironment of a binding pocket. They also do not provide strong tests of the ability of a model to capture interactions between solutes in the aqueous environment.

Host-guest systems are attractive, though largely untapped, test cases for detailed models of binding. Although host molecules are not proteins, molecular recognition events in host-guest and protein-ligand systems are governed by the same statistical mechanical rules and driven by the same interaction forces [4]. In particular, since the binding pockets of hosts are well solvated, the intricacies of hydration can play an equally important role in determining their binding properties. Nonetheless, host molecules are dramatically simpler than proteins, because they typically comprise only a few hundred atoms and tend to be more rigid than proteins. As a consequence, it is dramatically easier to achieve adequate conformational sampling for them compared to proteins [5, 6]. In recent years, cucurbiturils have emerged as particularly appealing test cases, because they exhibit a wide range of affinities for varied guests in aqueous solution [7-9], with maximal affinities rivaling those of the tightest-binding protein-ligand systems [10-12]. In addition, a growing repertoire of cucurbituril derivatives offers the possibility of further diversity in terms of chemistries, molecular flexibility, etc. [13-16] The growing interest in these host-guest systems for applications in drug delivery [17, 18] and chemical sensors [19] is also generating rapid growth in the volume of published binding affinity data.

In addition to the selection of suitable test systems, the format of a validation study is of critical importance. In particular, efforts such as CASP [20], the pK_a Cooperative [21], and SAMPL [22] have proven the value of blinded prediction challenges for objective validation of computational methods. Here we summarize the procedure and results of the first blinded prediction challenge for host-guest binding affinities, which was included as a new component of this year's SAMPL. Other papers in this issue provide detailed reports from the participating groups. This host-guest affinity challenge bridges the scale and complexity of the other SAMPL prediction challenges, including prediction of tautomers, solvation free energies, and protein-ligand affinities. The primary goals of this challenge are to define the state-of-the-art, learn from each other's experiences, identify problem areas requiring further improvement, create improved methods of modeling host-guest binding, and gain knowledge that will advance methods of predicting protein-ligand affinities.

MATERIALS AND METHODS

Challenge design

Information about the challenge was posted on the SAMPL website, sent to many commercial, industry, and academic researchers, and posted on the computational chemistry list (CCL; <http://www.ccl.net>). All interested parties, not only those to whom announcements were sent, were welcomed to take part by registering at the website (<http://sAMPL.eyesopen.com>). The data sets were provided for download there in SDF or PDB format. Participants were explicitly informed that the conformation, tautomer, and protonation states provided were not necessarily optimal or complete. Experimental

conditions were specified, including the temperature, salt concentration, and pH at which the binding affinities were determined. In addition, a previous study by Ma et. al. [15] was brought to the attention of all participants to ensure that every one was aware of relevant published data for host **H1**. The host **H1** system (below) was posted on the website on September 23rd, 2010, and the hosts **H2** and **H3** systems were posted on February 20th, 2010. Unfortunately, some participants were unaware that there were two sets of data and therefore submitted predictions for only host **H1** or only hosts **H2** and **H3**; a few did submit predictions for all three systems. The final submission deadline was June 30th, 2011. As in previous SAMPL challenges, the experimental values were not released to the participants until after the submission deadline. The outcome of the challenge and its implications were discussed at a workshop on August 1-2, 2011 at Stanford University, and all participants were invited to submit papers describing their calculations. The manuscripts, which were all reviewed by other SAMPL participants, are also published in this special journal issue.

Host-guest systems and structures

Crystallographic structures of hosts **H1**, **H2** and **H3** were obtained from the literature [15] or Cambridge Structure Database (CSD) [23], and are shown in Figure 1, along with the chemical structures of the guest molecules, **G1** - **G9**. These crystal structures of the hosts were provided to the participants along with modeled 3D coordinates of the guest molecules. Identifying the conformations, protonation state, and tautomer state of the hosts, guests and complexes, in solution was considered as part of the prediction challenge, and the modeled coordinates of the guest molecules were provided to the participants with the disclaimer that these are only meant to serve as potential starting geometries. The identifiers used here for some of the guests differ from the potentially ambiguous names used when originally identifying the compounds to the SAMPL3 participants. The original notation was: h1.guest(1-7) for guests **G1-G7**, h2.guest1/h3.guest1 for guest **G8**, and h2.guest2/h3.guest2 for guest **G9**. Note that other papers in this special issue may use the original notation to identify the guest molecules.

Experimental details

Drs. Lyle Isaacs (host **H1** [15]) and Adam Urbach (hosts **H2**, **H3** [24]) provided the unpublished experimental binding affinities of 11 new host-guest complexes. The data are summarized in Table 1. Binding affinities of the **H1** systems were measured at 298 K in 20 mM sodium phosphate buffer at pH 7.4, in one or more of the following ways: 1) ¹H NMR spectroscopy - titration of a fixed concentration of the guest with an increasing concentration of the host; 2) UV/Vis competition assay - titration of a fixed concentration of the host and a dye with an increasing concentration of guest; 3) ¹H NMR competition assay. The binding affinity reported for **G1** is that of a racemic mixture. A detailed description of the experimental protocols applied to similar host-guest systems can be found in reference [7]. Binding affinities of **G8** and **G9** with hosts **H2** and **H3** were determined using isothermal titration calorimetry (ITC) at 300 K in 10 mM sodium phosphate buffer at pH 7.0. Synthesis and characterization of compounds **G8** and **G9** will be reported elsewhere. Titration experiments were carried out on a VP-ITC calorimeter (Microcal, Inc.). The concentration of the host in the sample cell was between 0.062 and 0.174 mM, and the guest concentration was between 0.5 and 2.81 mM. The titration consisted of 27 consecutive injections of 10 μ L with at least 200 s interval between injections. Heats of dilution, measured by titrating beyond saturation, were subtracted from each data set. All solutions were degassed prior to titration. The data were analyzed using the one-set-of-sites model in the Origin software; this is a simple binary equilibrium model that assumes no interaction between complexes. A high affinity of these guests (**G8** and **G9**) with hosts **H2** and **H3** suggests inclusion complexation [25]. In all cases (hosts **H1**, **H2**, and **H3**), a 1:1 stoichiometry of binding was observed.

Computational methods

Participants employed a range of different methods, as detailed in other articles in this issue. Table 2 lists the methods by name, and categorizes them as follows:

- Overall method
 - Pathway methods [2] compute binding free energies in terms of the free energy changes associated with a series of small steps along a path, such as a path linking the free and bound states. The pathway approaches used here include thermodynamic integration (TI) [26, 27], Bennett acceptance ratio (BAR) [28], Wang-Landau [29], binding energy distribution analysis method (BEDAM) [30], and orthogonal space random walk (OSRW) [31].
 - End-point methods compute binding free energies through analysis of only the free and bound states of the system, rather than connecting them by a path. The end-point methods used here include MM/PBSA [32], MM/GBSA [32], solvated interaction energy (SIE) [33], and Mining Minima (M2) [34, 35].
- Potential energy as a function of conformation
 - Most of the methods used a generalized force field, OPLS [36, 37], GAFF [38], MMFF94 [39], CHARMM [40], or CGenFF [41], as an efficient method of estimating the potential energy of a molecule or complex as a function of conformation. Partial atomic charges, which are a critical force field component, were in turn generated by several different approaches [42-44], but most commonly AM1-BCC [45, 46].
 - One participant avoided the need to generate force field parameters, with their potentially non-optimal parameters and simplified functional form, by using an electronic structure method, density functional theory [47, 48] with the B3LYP functional [49, 50].
- Solvation free energy as a function of conformation
 - All end-point methods used an implicit solvent model based on continuum electrostatics, such as PBSA [51], GBSA [52], PCM [53], and FiSH [54]. In addition, the BEDAM pathway method used the similar but more intricate AGBNP2 model [55, 56]. Note, however, that different implementations of nominally similar continuum electrostatic models might yield significantly different results if they use different atomic radii, and the use of different numerical solvers may also be relevant; in particular, the SIE method uses a boundary element solver instead of the more commonly used finite difference solver.
 - All pathway methods other than BEDAM used the TIP3P [57] explicit water model in the context of molecular dynamics (MD) simulations.
- Conformational sampling method
 - All pathway methods used MD as the conformational sampling method.
 - Several end-point methods, generated and processed conformations by using a docking program to fit the guests into the crystal structures of their respective hosts.
 - The MM/GBSA and MM/PBSA methods sampled multiple conformations from the Boltzmann distribution with MD using explicit solvent.

- The M2 method used Tork [58], an aggressive conformational search algorithm, along with an implicit solvent model, to identify many local energy minima for the free and bound structures.
- Configurational entropy; i.e., the entropy associated with the host and guest, but not the solvent
 - The SIE method did not use any specific method to account for changes in configurational entropy on binding, although the <1 scaling parameter applied to the overall binding energy might be considered to account crudely for an entropic penalty proportional to energy.
 - The pathway methods include entropy implicitly but, in principle, fully, through their free energy approaches.
 - A number of end-point methods used the rigid-rotor/harmonic oscillator (RRHO) approximation.
 - The M2 method applied the harmonic approximation with mode scanning (HA/MS) to each energy well, and also accounted for the entropy associated with the existence of multiple energy wells. The HA/MS method is similar to RRHO, but it accounts for some anharmonicity in each energy well by scanning energies along eigenvectors of the Hessian associated with low eigenvalues (force constants).

RESULTS

Overview of host-guest predictions

A total of 19 submissions were provided by 10 different research groups. Scatter plots of calculated versus experimental binding free energies are shown in Figure 2, and statistical measures of accuracy are provided in the last two columns of Table 2. Because not all groups made predictions for all systems, and because the host **H1** systems are quite different physically from the host **H2** and host **H3** cases, we present the error statistics separately for these systems in Table 2, under the headings Host1 and Host2/Host3. For those methods which were applied to all three hosts, error statistics across all three hosts are also provided at the bottom of Table 2.

Overall, the predicted binding affinities suffer from high root-mean-square errors (RMSE), low correlations, or both. The RMSEs vary from 1.4 to 45.2 kcal/mol, and the correlation coefficients (R^2) vary from 0.01 to 0.94. The correlation coefficients tend to be higher for the host **H2** and host **H3** cases than for host **H1**, but the significance of this pattern is not strong, because the number of data points for **H2** and **H3** systems is only four, compared to seven data points for **H1** system. Moreover, since only a subset of methods was applied to the hosts **H2** and **H3**, it is not clear whether the higher correlations are indicative of the efficacy of these particular methods or of the relatively tractability of the systems. More generally, it is difficult to determine the basis for the differences among the various results with any confidence. One reason is that the submissions varied non-systematically in their choice of force-field parameters and atomic partial charges, solvation models, and conformational sampling methods.

The solvated interaction energy (SIE) model has the lowest RMSE, with better correlation ($R^2 = 0.77$) for the hosts **H2** and **H3** than for the host **H1** ($R^2 = 0.44$). Interestingly, this method yielded similar affinities when host **H1** was treated as uncharged (submission 5). Interestingly, other methods that used the same GAFF/AM1-BCC force field parameters as the SIE calculations (submissions 6, 7, 13 and 14) were not as accurate. More generally, we

could not identify a clear advantage for any particular charge set, force-field, solvation model, or free-energy calculation method. Indeed, seemingly similar methods like TI, BAR, and FEP, combined with similar parameters, produced quite different results, suggesting unexpectedly that the details of the free-energy method might have contributed to the differences among these predictions. For example, submissions 13 and 14 used identical energy models, but their respective RMSEs were 4.0 and 11.3 kcal/mol and their correlation coefficients were 0.34 and 0.81. In principle, the differences in computed free energies are attributable to some combination of differences in energy model and differences in conformations sampled. However, in the absence of information about the predominant conformations found by the various methods, or of a detailed breakdown of the binding free energies into terms like the changes in internal energy, solvation energy, and entropy on binding, it seems impossible to delineate the chief sources of variation among the different predictions. It is also unclear whether or how the correction for standard concentration [59], which normally yields a constant offset, was accounted for in some submissions.

Host1 predictions

The protonation states of the host **H1** systems under the experimental conditions are uncertain and might change on binding. In particular, the mutual proximity of the four carboxyl groups on the host may reduce their acidity, allowing some of them to be protonated; and several of the guests contain anilino groups of uncertain pK_a . Most of the calculations are based on the standard pK_a s of the chemical groups involved and assume no change in protonation on binding.

Setting aside the calculations for a moment, one may observe that the experimental affinity trends of the host **H1** cases are intuitively reasonable. Thus, the host, whose charge may be as great as -4, binds the neutral guest **G4** with lowest binding affinity (-4.17 kcal/mol), the +1 guests (**G1**, **G2**, **G3**, **G5**, and **G7**) with intermediate affinity (-6 to -8 kcal/mol), and the +2 charge guest **G6** (at pH 7.0 based on standard pK_a 's) with highest binding affinity (-10.7 kcal/mol). The sizes of the guests correlate with their charges, so the weakest affinity also corresponds to the smallest guest and the greatest affinity to the largest guest. A majority of the submissions capture this general trend, but they also tend to invert the affinity rankings within the group of +1 guests (**G1**, **G2**, **G3**, **G5**, and **G7**), resulting in poor overall correlation coefficients. Two submissions (7 and 14), captured the relative affinities of +1 charged guests somewhat better, resulting in an overall correlation coefficient of 0.8. However, these methods had RSMEs greater than 10 kcal/mol. The large mean error of submission 14 results primarily from a large error for guest **G6** (+2); the authors retrospectively attribute this to a setup error (W. Yang, personal communication). It is not clear why these methods ranked the +1 guests relatively well, while others using similar force-field parameters showed anti-correlation for the +1 guests. The fact that host **H1** is flexible enough to undergo significant conformational changes might play a role, as different methods might have sampled different conformers.

Host2/host3 predictions

The protonation states of hosts **H2** and **H3** and their guests are relatively unambiguous, and the molecules are also rather rigid. Both hosts were studied with the same set of guests. The key trend observed in the experimental data is that the guests bind more tightly to the larger host (host **H3**) than to the smaller one (host **H2**), and a majority of the submissions captured this trend. Additionally, guest **G8** binds host **H2** more tightly than guest **G9** does, while the converse is true in the case of host **H3**. Only 4 of the 12 submissions captured this pattern, but the differences in measured binding free energies were less than 1.5 kcal/mol, which is likely well within the range of uncertainty of the energies of current force fields.

More broadly, the RMSEs of the computed binding affinities submitted range from 1.4 and 45.2 kcal/mol, with correlation coefficients between 0.14 and 0.94. Encouragingly, 9 of the 14 submissions show correlations greater than 0.7, despite large RMSEs. Although the SIE model has the lowest RMSE, a method based on a quantum mechanical (QM) energy model has the highest correlation coefficient, 0.94, and clearly captures the key trends in the data. Interestingly, when changes in entropy are dropped from this method, the correlation coefficient drops to 0.64. Also, calculations using the same basic approach as the quantum method, but now using MMFF with AM1-BCC charges (submissions 15 and 16), show similarly high correlation coefficients, greater than 0.9. Much as observed in the host **H1** submissions, some binding affinities computed with the same force field and similar continuum solvent models as those of the relatively successful SIE approach incurred significantly larger RMSEs and lower correlation coefficients, and indeed anti-correlated with the measured affinities.

Discussion

This paper summarizes the results of the first blind prediction challenge for host-guest binding affinities. The availability today of host-guest systems with binding affinities similar to those of protein-ligand systems provides an important opportunity to test and ultimately improve physics-based models of binding, in a setting where computational demands are less limiting than they are in the case of proteins. In particular, the challenge of conformational sampling is dramatically reduced for relatively rigid, low molecular weight hosts. New host-guest affinities were measured to enable the present study, and a number of research groups had the courage to submit blind predictions. The participants used a wide variety of approaches ranging from simple docking to extensive molecular dynamics based free-energy calculations.

Sources of accuracy and error

Given the relative simplicity of the host-guest molecular systems, it is somewhat surprising that none of the predictions agree particularly well with the experimental data. In addition, the various methods provide a rather wide spread of results. One fundamental question is whether or not the various methods sampled essentially the same set of conformations. If they did – and one may hope this is the case, given the small sizes of these systems – then the differences among the methods should be attributable primarily to differences in the energy models they used. On the other hand, models with similar energy functions were observed to yield significantly different results (e.g., methods 13 and 14), suggesting significant differences in conformational sampling. It is also striking that the SIE approach [33] provided some of the most accurate results, despite the fact that conformational sampling in this method is limited to extensive sampling of the ligand, while the receptor is kept rigid except for a final energy-minimization step. There was no clear advantage to using more elaborate sampling methods, such as molecular dynamics simulations or the Tork conformational search method [58]. One possible explanation of this broad observation is that conformational sampling might amplify the effects of errors in the energy models by putting too much statistical weight on unrealistic conformations. It is also worth noting that only the SIE method applies empirical weighting coefficients to its computed energy terms. It is intriguing that these coefficients appear to work reasonably well, despite having been fitted based on protein-small molecule data [33].

During the workshop discussion, the protonation states of the host **H1** systems emerged as a key source of uncertainty in the calculations. Although the participants were aware of this issue, almost all predictions were based on the simplifying assumption that the protonation states are identical in the bound and unbound conformations, and the charge states were set

based on the standard pK_a of the different chemical groups in solution. In a few cases where the binding free energy computed for the anilino guests was greater than zero, the participants considered alternate protonation states. There are actually two issues here. In addition to uncertainty in the baseline pK_a s of the various ionizable groups, especially the four host carboxyls, it is also possible that protonation states change significantly on binding. Indeed, there is substantial experimental evidence for pK_a shifts induced by cucurbituril binding [17]. In general, it is observed that cucurbiturils favor binding of the protonated form of the guest, and therefore induce increases in pK_a ($pK_a^{\text{bound}} - pK_a^{\text{unbound}}$) ranging up to +4.5 units; negative shifts as great as -1.5 units have also been observed [17]. The positive pK_a shifts of bound guest molecules are for standard, electrically neutral cucurbiturils and, presumably will only be increased for host **H1**, with its four carboxyls. Of particular relevance here is the pK_a shift of the aniline group of 2-aminoanthracene from 4.0 to 7.1 upon binding to cucurbit[7]uril [60]. Also, guests **G2** (procaine) and **G7** (tetracaine) exhibit 1.22 and 1.91 units of pK_a shift, respectively, upon binding to cucurbit[7]uril [61]. Three out of seven guests for the host **H1** system contained an aniline group, and therefore the choice of protonation state based on standard pK_a could have significantly affected the predicted binding affinities. It is important to note that, to the extent that uncertainties in protonation states are sources of error in the present calculations, the present comparisons are not informative regarding the quality of the force fields used and the adequacy of conformational sampling. Thus, for future rounds of host-guest blind predictions, it may be helpful to steer clear of systems posing this issue, or to be sure of including some without it. Since protonation state issues were not prominent for the host **H2** and **H3** cases, it is unfortunate that a number of groups in this initial round were unaware of these systems until it was too late to process them before the workshop. Although the predicted binding affinities for hosts **H2** and **H3** still deviated significantly from the experimental values, the correlation between experiment and computation was better for these systems.

The possibility that protonation states and shifts contributed to the computational errors observed here is not necessarily surprising, but it is somewhat sobering, given that similar issues frequently arise in protein-ligand binding because of the high density of charged amino acids in the catalytic sites of enzymes and other protein binding sites [62, 63]. Enzyme active sites in particular frequently contain catalytic histidine residues whose pK_a s are poised near physiological pH, making their charge states both uncertain and changeable [63]. Although the theoretical framework for coupled protonation and binding equilibria has long been known [64, 65], it is not clear whether current computational methods for modeling protonation equilibria are sufficiently reliable to enable accurate affinity predictions. Prediction of pK_a s based on physical models has a long history and we recommend these excellent reviews on this topic [66-68]. The pK_a Cooperative [21], a blind challenge for structure-based computational methods for predicting pK_a values and electrostatic energies in proteins, is notable in this context.

The heart of these binding affinity calculations lies in the energy model itself, i.e. the force field parameters and the solvation model. All participants in the present study used generalized force-field parameters, such as GAFF [38] with AM1/BCC charges [45, 46], except for one group which used a quantum mechanical Hamiltonian. Interestingly, the quantum mechanical results correlated relatively well with experiment, despite overestimating the affinities considerably. Currently, a large amount of benchmarking data is being made available by computational chemists using high-level quantum mechanical models [69-74], and these are likely to be helpful in improving the current generalized force-fields. For instance, Goodman et al. [75] have recently compared several classical generalized force-fields against some of the above mentioned quantum-mechanical benchmarking data and showed that the mean errors in interaction energies are 2 kcal/mol or more, even for seemingly simple, fragment-like molecules. Current protein force-fields

(OPLS [37] and AMBER [76]) perform significantly better against similar benchmarks, with mean errors in interaction energies less than 1 kcal/mol [72]. It would appear that, although force-fields for proteins are already well optimized, there may be significant room for improvement of the generalized force fields used for the broader chemistries of drug-like ligands and host-guest systems. Indeed, it may well be that force field errors underlie many of the problems observed in the present exercise.

Changes in solvation free energy also contribute significantly to the computed host-guest binding affinities [6, 10]. Interesting, although explicit solvent models may be regarded as the gold standard, predictions here using an explicit solvent model did not outperform those that relied on faster continuum methods. In fact, the best-scoring SIE model used the Poisson-Boltzmann Surface Area (PBSA) implicit solvent model [51]. The existence of other differences among the models makes it impossible to draw definite conclusions about the relative merits of the different solvent models, these broad observations suggest that implicit solvation models will continue to play an important role in models for binding.

Finally, it is of interest to consider whether different computational definitions of the bound state of the various complexes could lead to significant differences among the various calculations and potentially to deviations from experiment. The appropriate definition of the bound state will be unambiguous in systems with a single, deep well in the host-guest potential of mean force [3, 59, 77]. However, the situation can in principle become more complicated, such as if the energy well is shallow or if there are multiple energy wells of similar depths, especially if some of these have the guest positioned outside the cavity of the host.

Implications for protein-ligand modeling

It is interesting to consider how the results of this community exercise bear on the calculation of protein-small molecule binding affinities, particularly in connection with computer-aided drug design. The underlying statistical mechanical principles of binding are expected to be the same for host-guest and protein-ligand systems, and the same physical forces including non-covalent interactions, hydrogen bonding, and solvation, are at play in both settings. However, as noted above, calculations for proteins, rather than hosts, benefit from the availability of more refined force field parameters. On the other hand, it is still necessary to assign force field parameters to small, drug-like molecules, and this could be an important source of error in protein-ligand modeling. Models of both protein-ligand and host-guest systems also encounter the challenge of dealing with protonation states, and issues of solvation models appear similar in both settings. Perhaps the chief difference, then, is that proteins have far more degrees of freedom, so achieving adequate conformational sampling is expected to be far more challenging. It is perhaps relevant that one message of the present study is that more conformational sampling does not necessarily lead to more accuracy. Nonetheless, it is unlikely on physical grounds that calculations without significant conformational sampling can ever be highly accurate. Overall, given the similarities between protein-ligand and host-guest binding, it is hard to imagine that reliable results can be achieved for proteins without also achieving them for host-guest model systems. Because host-guest systems have the added merit of simplicity, they represent informative and tractable test systems for future methods development.

Directions

As evidenced by the special papers in this issue, the results of this blind prediction challenge are leading to re-evaluations of the various methods that will hopefully lead to better results in the future rounds of SAMPL, not only for the host-guest systems but also for protein-ligand binding. The SAMPL3 host-guest binding affinity challenge was the first step in a

collaborative process of confronting experiment with computation, and the blind prediction setting is an excellent way to mimic the challenges faced by today's pharmaceutical industry.

The main goal of this blind prediction challenge was to identify the key problem areas that need further attention. The range of computational methods used here by the participants was informative in assessing the strengths and weaknesses of each of the methods. However, due to the varied modeling choices made by the participants, it was impossible to draw definite conclusions regarding the contribution of the different energy components, solvation, force fields, protonation, and conformational sampling, to the errors in predictions.

Future rounds of host-guest blind predictions would be more informative if participants supplied not only their free energy predictions, but also the coordinates of conformations that were sampled, and, where possible, the energy components underlying their computed free energies. A more detailed analysis of these additional data could shed light on the right direction for future improvements. In addition, one of the major sources of confusion was the choice of ionization states, as the different choices made by some participants masked the effects of other methodological differences, such as the force field and sampling method. It might help to understand this problem in more detail if more participants submitted predictions based on different ionization states. Finally, it was unexpected that seemingly similar approaches using identical energy models resulted in very different predictions. In the future rounds of SAMPL blind prediction challenge, it may be useful for participants to make multiple submissions with controlled variations in the choices, as this would also help understand how different methods performed in the hands of different participants.

Overall, progress on any single front, such as force field parameters or conformational sampling, will likely be insufficient for a marked improvement in the accuracy of predictions. Rather, a concerted effort towards all the challenges discussed here will be required. It will also be important to take advantage of other community prediction exercises, such as the Community Structure-Activity Resource (CSAR) [78, 79]. This effort focuses on protein ligand systems, and in addition to affinity scoring, the CSAR challenges the participants to accurately predict binding poses, which are compared against high-quality crystallographic structures. Another blind challenge that deserves mention here is the pK_a Cooperative [21], which is focused towards advancing the development of structure-based computational methods for computing pK_a values and electrostatic energies in proteins. The continued involvement of researchers across the country and possibly the globe in such prospective prediction challenges will speed progress in molecular modeling and lead to improved tools for computer-aided drug design.

Acknowledgments

This work was supported in part by Grants GM61300 from the NIGMS to M.K.G, CHE-1110911 from the NSF to L.I., Robert A. Welch Foundation (F-1621) to C.W.B, and NSF grant CHE-0748483 to A.R.U. The content is solely the responsibility of the authors and does not necessarily represent the official views of the NIGMS or the National Institutes of Health. We thank Drs. Emilio Gallicchio, Enrico Purisima and Wei Yang for corrections and suggestions.

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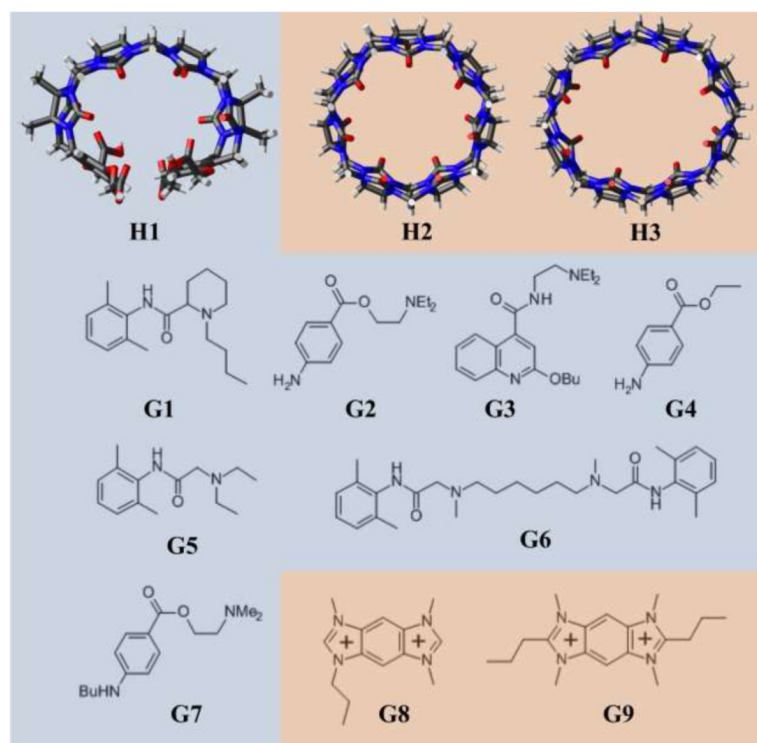


Figure 1. Crystal structures of hosts **H1**, **H2**, and **H3**, and chemical structures of guests **G1-G9**. Host **H1** and its respective guests (**G1-G7**) are shaded in blue; hosts **H2** and **H3** with their respective common guests (**G8** and **G9**) are shaded in orange.

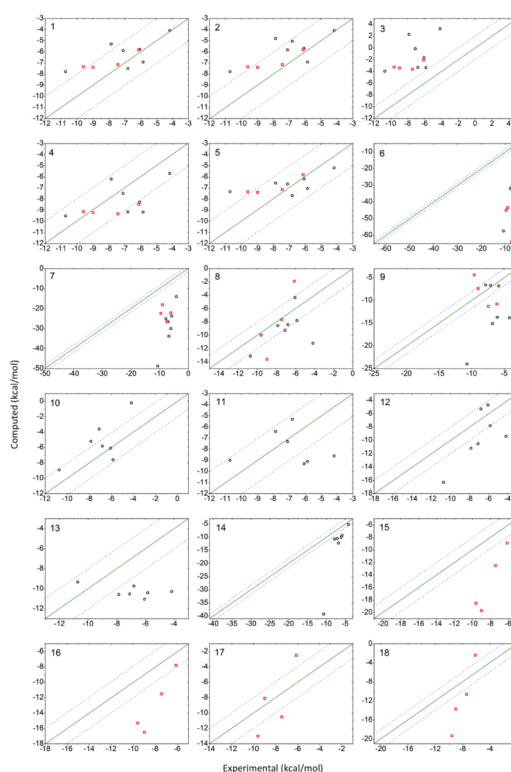


Figure 2.

Individual predictions submitted for the SAMPL3 host-guest blind prediction challenge. The numbers indicated in the top left corner of each plot corresponds to the submission ID given in Table 2. Host1 systems are indicated by black open circles, whereas hosts 2 and 3 are indicated by red open squares. The blue dotted lines indicate an error range of ± 2 kcal/mol.

Table 1

Experimentally measured binding affinities of various host-guest complexes. Measurements are reported as mean \pm S.D.

Host	Guest	K_a (M^{-1})	ΔG (kcal/mol)	ΔH (kcal/mol)	ΔS (cal/mol.K)
H1	G1	$1.8 \pm 0.1 \times 10^4$	-5.84	-	-
	G2	$1.5 \pm 0.1 \times 10^5$	-7.10	-	-
	G3	$9.0 \pm 0.6 \times 10^4$	-6.80	-	-
	G4	$1.1 \pm 0.1 \times 10^3$	-4.17	-	-
	G5	$2.6 \pm 0.1 \times 10^4$	-6.06	-	-
H2	G6	$6.5 \pm 0.7 \times 10^7$	-10.72	-	-
	G7	$5.3 \pm 0.4 \times 10^5$	-7.85	-	-
	G8	$2.9 \pm 0.5 \times 10^4$	-6.12	-8.5 (± 0.5)	-7.9 (± 1.9)
	G9	$2.6 \pm 0.5 \times 10^5$	-7.43	-3.8 (± 0.2)	12.0 (± 1.0)
	G8	$1.0 \pm 0.1 \times 10^7$	-9.60	-7.2 (± 0.3)	8.1 (± 0.9)
H3	G9	$3.6 \pm 0.4 \times 10^6$	-8.99	-4.2 (± 0.1)	16.0 (± 1.0)

Table 2

Summary of predictions submitted for the SAMPL3 host-guest binding affinity challenge.

ID	Method (e/p) ^a	Pot. Energy	Charges	Solv.	Conf.	Config-Entropy	RMSE	R ²
Host1								
1	SIE (e)	GAFF	AM1-BCC	PBSA	Docking	None	1.6	0.42
2	SIE (e)	GAFF	AM1-BCC	PBSA	Docking	None	1.9	0.44
3	SIE (e)	GAFF	AM1-BCC	FISH	Docking	None	6.4	0.22
4	rSIE (e)	GAFF	AM1-BCC	FISH	Docking	None	2.0	0.22
5	SIE (e)	GAFF	AM1-BCC	PBSA	Docking	None	1.5	0.40
5*	rSIE (e)	GAFF	AM1-BCC	PBSA	Docking	None	2.8	0.40
6	MM/CBSA (e)	GAFF	AM1-BCC	GBSA	MD/Expl.	RRHO	31.9	0.58
7	MM/PBSA (e)	GAFF	AM1-BCC	PBSA	MD/Expl.	RRHO	23.4	0.80
8	Szybki (e)	MMFF94	AM1-BCC	Szybki	Docking	RRHO	3.1	0.19
9	M2 (e)	CHARMm	Vcharge	PBSA	Tork	HA/MS	7.5	0.24
10	BEDAM (p)	OPLS	OPLS	AGBNP2	MD/Imp.	Implicit	2.5	0.49
11	TI/BAR (p)	CGenFF	CGenFF	TIP3P	MD	Implicit	2.6	0.01
12	TI (p)	GAFF	RESP	TIP3P	MD	Implicit	3.6	0.51
13	Wang-Landau (p)	GAFF	AM1-BCC	TIP3P	MD	Implicit	4.0	0.34
14	FEP/OSRW (p)	GAFF	AM1-BCC	TIP3P	MD	Implicit	11.3	0.81
Host2/Host3								
1	SIE (e)	GAFF	AM1-BCC	PBSA	Docking	None	1.4	0.77
2	SIE (e)	GAFF	AM1-BCC	PBSA	Docking	None	1.4	0.77
3	SIE (e)	GAFF	AM1-BCC	FISH	Docking	None	5.1	0.44
4	rSIE (e)	GAFF	AM1-BCC	FISH	Docking	None	1.5	0.46
5	SIE (e)	GAFF	AM1-BCC	PBSA	Docking	None	1.4	0.77
5*	rSIE (e)	GAFF	AM1-BCC	PBSA	Docking	None	1.9	0.79
6	MM/CBSA (e)	GAFF	AM1-BCC	GBSA	MD/Exp.	RRHO	45.2	0.57
7	MM/PBSA (e)	GAFF	AM1-BCC	PBSA	MD/Exp.	RRHO	14.8	0.14
8	Szybki (e)	MMFF94	AM1-BCC	Szybki	Docking	RRHO	3.1	0.77
9	M2 (e)	CHARMm	Vcharge	PBSA	Tork	HA/MS	4.1	0.80
15	MM/PBSA (e)	MMFF94	AM1-BCC	PBSA	Docking	RRHO	7.5	0.93

ID	Method (e/p) ^a	Pot. Energy	Charges	Solv.	Conf.	Config. Entropy	RMSE	R ²
16	MM/PBSA (e)	MMFF94	AM1-BCC	PBSA	Docking	RRHO	5.2	0.92
17	DFT/PCM (e)	B3LYP	--	PCM	Docking	None	3.0	0.64
18	DFT/PCM (e)	B3LYP	--	PCM	Docking	RRHO	5.9	0.94
Hosts 1/2/3								
1	SIE (e)	GAFF	AM1-BCC	PBSA	Docking	None	1.5	0.51
2	SIE (e)	GAFF	AM1-BCC	PBSA	Docking	None	1.7	0.53
3	SIE (e)	GAFF	AM1-BCC	FiSH	Docking	None	6.0	0.27
4	rSIE (e)	GAFF	AM1-BCC	FiSH	Docking	None	1.8	0.27
5	SIE (e)	GAFF	AM1-BCC	PBSA	Docking	None	1.5	0.50
5*	rSIE (e)	GAFF	AM1-BCC	PBSA	Docking	None	2.5	0.50
6	MM/GBSA (e)	GAFF	AM1-BCC	GBSA	MD/Expl.	RRHO	37.3	0.24
7	MM/PBSA (e)	GAFF	AM1-BCC	PBSA	MD/Expl.	RRHO	20.7	0.29
8	Szybki (e)	MMFF94	AM1-BCC	Szybki	Docking	RRHO	3.1	0.26
9	M2 (e)	CHARMm	Vcharge	PBSA	Tork	HA/MS	6.5	0.02

^aThe basic approach, end-point (e) or pathway (p), is indicated in parentheses next to the name of each method. Abbreviations: MD/Expl. -- Molecular dynamics simulation with explicit water model. SIE - Solvated interaction energy; rSIE - Rescaled SIE; GAFF - Generalized AMBER force field; AM1 - Austin Model 1; BCC - Bond charge correction; PBSA - Poisson-Boltzmann Surface Area; FiSH - First-shell hydration; GBSA - Generalized-Born Surface Area; MM - Molecular mechanics; MD - Molecular dynamics; RRHO - Rigid rotor harmonic oscillator; M2 - Mining Minima; BEDAM - Binding energy distribution analysis method; HA/MS - Harmonic approximation/Mode scanning; MMFF - Merck molecular force field; CHARMM - Chemistry at Harvard molecular mechanics; TI - Thermodynamic integration; BAR - Bennett acceptance ratio; CGenFF - CHARMM generalized force-field; FEP - Free energy perturbation; OSRW - Orthogonal space random walk; OPLS - Optimized potentials for liquid simulations; AGBNP2 - Analytical Generalized Born plus non-polar 2; RESP - Restrained electrostatic potential; PCM - Polarizable continuum model; DFT - Density functional theory; B3LYP - Becke, three-parameter, Lee-Yang-Parr.

Despite the methodological differences, submission 5 is effectively a linear scaling of submission 5 ($R^2 = 0.998$), with a scaling coefficient of 0.41 and an offset of -2.90 kcal/mol.