



A Fixed-Charge Model for Alcohol Polarization in the Condensed Phase, and Its Role in Small Molecule Hydration

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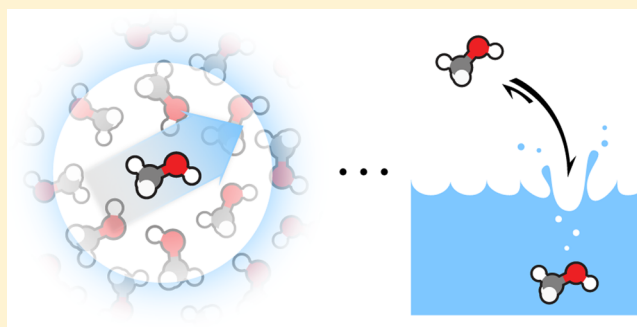
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S Supporting Information

ABSTRACT: We present a simple optimization strategy for incorporating experimental dielectric response information on neat liquids in classical molecular models of alcohol. Using this strategy, we determine simple and transferable hydroxyl modulation rules that, when applied to an existing molecular parameter set, result in a newly dielectric corrected (DC) parameter set. We applied these rules to the general Amber force field (GAFF) to form an initial set of GAFF-DC parameters, and we found this to lead to significant improvement in the calculated dielectric constant and hydration free energy values for a wide variety of small molecule alcohol models. Tests of the GAFF-DC parameters in the SAMPL4 blind prediction event for hydration show these changes improve agreement with experiment. Surprisingly, these simple modifications also outperform detailed quantum mechanical electric field calculations using a self-consistent reaction field environment coupling term. This work provides a potential benchmark for future developments in methods for representing condensed-phase environments in electronic structure calculations.



INTRODUCTION

Computer simulations exist as a way to connect theories of the microscopic world to experimental results. They help validate our understanding of microscopic phenomena and propose new directions for investigation. Such simulations employ molecular models, and the development of molecular models and their component parameters has a long history.^{1–7} One of the primary motivations in their continued development is the increased ability of a molecular model to quantitatively reproduce and/or predict experimental properties of interest.

Our depiction of molecules in simulations has been grounded by several experimental observables. These include densities, enthalpies, liquid structure, molecular geometry, phase coexistence boundaries, and more.^{8–12} In the case of water, anomalous properties such as the temperature of maximum density provide key motivation.^{13–15} Whether the simulation is classical fixed-charge, polarizable, or even quantum mechanical (QM), models used often are unable to reproduce all the observables of interest and, thus, must strike a balance among fitting the selected properties, avoiding too many poorly defined parameters, and keeping the computational cost low. Generalized models, such as OPLS-AA,³ GAFF,⁶ and CGenFF,⁷ typically focus on neat liquid density and heats of vaporization as experimental targets for optimization of Lennard–Jones (LJ) parameters coupled with partial charges assigned using fits to HF/6-31G* electrostatic potentials. While models made from these parameter sets can

be used to study neat liquids, one of the main intents is for the representation of molecules in other environments, such as a ligand interacting with a protein or a solute transferring between different chemical phases. When using such a model outside of a neat liquid, it is understandable that one may observe a systematic error in a new property of interest. In the case of molecular transfer, researchers have explored a variety of approaches to reduce such systematic errors, and these generally amount to fitting the models to get the answer desired,⁵ sometimes coupled to hydrated environment QM calculations,¹⁶ and other times through statistical means by introducing additional water-mixing parameters to the LJ potential terms.¹⁷ For researchers interested in simultaneous reproduction of experimental observables in multiple disparate environments, which is sometimes considered impossible for classical fixed-charge models,^{5,18} more complex polarizable models and corrections have shown promise.^{19–22}

We are interested in both the neat liquid and air-to-water transfer properties of molecules with hydroxyl moieties. Recent studies on the hydration of alcohols using common force field parameters have shown systematic errors in ΔG_{hyd} ^{17,23–28} and

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other thermodynamic quantities, in particular the static dielectric constant, $\epsilon(0)$.²⁹ Are systematic errors in $\epsilon(0)$ and ΔG_{hyd} coupled to one another in some way? It has been suggested that functional groups in typical force fields are underpolarized to some degree,^{16,18,21,23,25,27–30} though this should not be too surprising given that partial charges are typically fit to reproduce a gas-phase HF/6-31G* electrostatic potential, despite this charge model's fortuitous propensity for overpolarizing the gas-phase electric field.^{2,6,31,32} For solution-phase simulations, it would be preferable to have an accurate degree of polarization for the condensed-phase fluid encoded within the atomic partial charges of the molecule while maintaining the typical neat liquid density and enthalpy of vaporization (ΔH_{vap}). Recent studies have begun using the experimentally measured $\epsilon(0)$ of a neat liquid as a fitting target for systematically scaling the degree of polarization of classical fixed-charge models.^{33,34} This provides an orthogonal approach to the practice of setting partial charges with a QM electrostatic potential and optimizing LJ parameters to fit experimental observables. As it relies on experimental quantities to govern the magnitude of the partial charges, it can potentially provide an independent assessment of the quality of a given QM calculation used to derive partial charges. Here, we extend such work with the goal of addressing the following questions: (1) is it possible to simultaneously capture the density, ΔH_{vap} , and $\epsilon(0)$ for simple alcohol solvents; (2) how transferable are the resulting hydroxyl parameters to other more complex alcohols; and (3) if an alcohol model reproduces the neat liquid $\epsilon(0)$, how does that affect the hydration of that alcohol as a solute?

METHODS

General Simulation Protocol. In this work, both extended molecular dynamics simulations and alchemical free energy calculations were used to calculate neat liquid thermodynamic properties and air-to-water transfer free energies for a series of 41 small molecules containing hydroxyl moieties for which both experimental quantities were available. Simulations were performed using GROMACS 4.5.5^{35–37} for the neat liquid molecular dynamics and a development version of GROMACS 4.6 for alchemical free energy calculations on these alcohol systems. Hydration free energies were computed using Thermodynamic Integration (TI), following a combination of previous protocols.^{24,25,38} We note deviations from previous protocols below.

Initial conformations for starting molecules were taken from structures provided in previous studies,²⁵ excepting the case of methanol subject to the neat liquid dielectric optimization, which was constructed and minimized using the structure editor in the UCSF Chimera visualization system.³⁹ The GAFF small molecule parameters⁶ and AM1-BCC partial charges^{40,41} were assigned to each structure using the Antechamber package (Amber 11 version).⁴² After setup, the resulting structures and topologies were converted to the GROMACS format using the ACPYPE python script.⁴³ For neat liquid calculations, cubic simulation boxes of 256 like small molecules were constructed with GROMACS utilities and subject to sets of 150 ps of constant-pressure and constant-temperature equilibration until the target state-point was achieved. Again, methanol in the neat liquid optimization was an exception in needing a large cubic simulation box containing 450 molecules in order to accommodate box size fluctuations along the density optimization pathways. For hydration free energy calculations, single copies of the small molecules were solvated in rhombic-

dodecahedral TIP3P¹⁰ water boxes with at least 1.2 nm of space between any solute atom and a box edge.

For all neat liquid calculations, a set of five 50 ns production simulations were performed for each molecule with a unique topology in order to obtain reported $\epsilon(0)$ values and associated standard errors. These calculations were performed at 1 atm and the target temperature from the experimental dielectric constant determination, usually 298.15 K unless otherwise noted. Specific temperatures are listed in the associated Supporting Information. A time step of 2 fs was used with the leapfrog integration algorithm for the equations of motion, and hydrogen atom bond lengths were constrained using the LINCS algorithm.⁴⁴ Isotropic pressure and temperature coupling were accomplished using the Parrinello–Rahman barostat⁴⁵ and Nose–Hoover thermostat^{46,47} with time constants of 10 and 1 ps, respectively. Smooth particle-mesh Ewald⁴⁸ with a real-space cutoff of 1.2 Å, spline order of 4, and energy tolerance of 10^{-5} was used for long-ranged electrostatics corrections. LJ interactions were switched off smoothly between 8 and 10 Å, and long-range energy and pressure corrections were applied to these interactions as implemented in GROMACS.⁴⁹

Hydration free energy calculations were performed over a series of separate λ windows as described in previous work,²⁴ with the primary difference being that these simulations were performed at constant pressure as in the neat liquid calculations, but with the Langevin thermostat as in previous work. Performing the calculations with the Parrinello–Rahman barostat was found to result in numerically equivalent results to previous protocols, while eliminating the need for an extensive constant pressure equilibration and average volume rescaling at the beginning of each λ window. For our set of 41 alcohols, TI was used to determine both the electrostatic and nonpolar components of solvation rather than the Bennett acceptance ratio (BAR) for technical reasons in the development version of GROMACS. For the SAMPL4 calculations, our standard BAR protocol was restored with the use of GROMACS 4.6.2 beta 2. The electrostatic component was calculated from the free energy of turning on the solute partial charges in water, less the free energy of the same charging process in vacuum. The nonpolar component was calculated from the free energy for decoupling the uncharged solute LJ interactions from the surrounding water as in previous work.²⁴ This overall protocol was validated against previous calculations on these 41 small hydroxyl containing solutes²⁵ and found to be reliable and equivalently accurate.

Optimization Process for Hydroxyl Groups. Starting with a general model for a solute one would use in a typical alchemical free energy calculation, an iterative optimization process was developed to alter the molecular topology in order to improve agreement with dielectric properties of the neat liquid. This process is a variation of previous work³³ with a focus placed specifically on small molecules containing hydroxyl groups. The simplest of such solutes is methanol, and Figure 1 shows an example optimization flow process for methanol.

The goal of this process is to retain or improve the density and ΔH_{vap} agreement that general molecular models are developed upon, while simultaneously improving the agreement with experimental $\epsilon(0)$ values. For simplicity, a three-stage procedure was employed with a linear interpolation in each stage used to target optimal values for these properties. After employing an initial 1 ns simulation to determine the starting model's density, the hydroxyl oxygen σ value was scaled

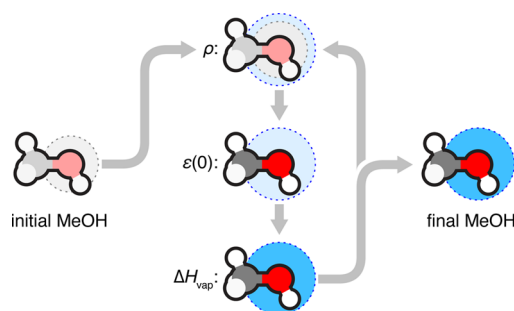


Figure 1. Illustration of the iterative optimization process for a methanol model using experimentally measured data. First, the size (LJ σ) is scaled to match experimental density for the neat liquid. This is followed by charge-scaling, to match the experimental dielectric constant, and well-depth (LJ ϵ) scaling, to match the experimental ΔH_{vap} . The resulting model is tested in a final set of simulations to see how well it matches these three properties, and it is either accepted or returned to the optimization pipeline for further refinement.

up or down by 5% if the density was either greater than or less than the target density window (791.4 kg/m^3 to $\pm 1.0 \text{ kg/m}^3$) respectively. The density of a 1 ns simulation of the resulting model was calculated, and a linear interpolation was performed between these densities to determine a σ value that should correspond to the model having the target density. This process was repeated until the model consistently reported the experimental density. This led to the second stage process of optimizing the $\epsilon(0)$ value (33 to ± 4), calculated using the standard dipole fluctuation formulas,⁵⁰ using similar linear interpolations over 15 ns simulations. In order to maintain charge neutrality, all the partial charges were scaled by $\pm 5\%$, effectively scaling the dipole moment of the methanol model in order to raise or lower the $\epsilon(0)$ value. This is illustrated in Figure 1 as a saturation of atom and bond color on the methanol molecule.

The third stage involved linear interpolation of ΔH_{vap} values to estimate an optimal LJ ϵ well depth for the hydroxyl oxygen atom. The ΔH_{vap} was calculated from the difference in per-molecule total energies from 10 ns single methanol gas phase and 500 ps liquid phase simulations,

$$\Delta H_{\text{vap}} = E_{\text{gas}} - E_{\text{liq}} + RT - E_{\text{pol}} \quad (1)$$

E_{pol} is the energetic term corresponding to the work required to polarize the methanol molecule from the gas phase dipole moment to the liquid phase dipole moment,¹⁴

$$E_{\text{pol}} = \frac{N}{2} (\mu_{\text{gas}} - \mu_{\text{liq}})^2 / \alpha_{\text{gas}} \quad (2)$$

Here, μ_{gas} and α_{gas} were 1.7 D and 3.29 \AA^3 respectively for gaseous methanol.⁵¹ To determine μ_{liq} from a flexible, yet fixed-charge methanol model, we averaged individual μ_{liq} values of each molecule from the last frame of the liquid simulation. Note that, as the neat liquid optimization system contains 450 alcohol molecules, the E_{liq} for the model is able to converge to $\pm 0.1 \text{ kcal/mol}$ over this relatively short optimization stage. As in the other stages, this process was iterated until the model reported ΔH_{vap} within the desired tolerance (8.95 kcal/mol to $\pm 0.59 \text{ kcal/mol}$). It should be noted that we only use ΔH_{vap} in this optimization of methanol and do not calculate it for other solutes later in this study. Experimental ΔH_{vap} , μ_{gas} , and α_{gas} properties are not available for all subsequent molecules investigated, making such comparisons problematic.

After this series of parameter adjustments, a final set of simulations was performed to check if all of these neat liquid properties were within the specified error tolerances for each of their state points. If any of the three properties degraded due to correlations between the parameters and properties not directly associated in the optimization procedure, the series of stages was iterated until the set of properties converged. It should be noted that the experimental values need not be at the same state point. In the case of methanol, the ΔH_{vap} value used was measured at 298.15 K, while the density and $\epsilon(0)$ were measured at 293.15 K.⁵¹ The adjusted topology will simply describe a classical molecular model that captures these experimental quantities at these differing state points. This topology will also not be the only set of parameters to capture the target property values, just the first set found that satisfies the criteria.

Application of Optimized Hydroxyl Parameters to New Model Molecules. While the $\epsilon(0)$ optimization procedure could adjust a classical molecular model to reproduce experimental properties for an entire set of molecules, it is not particularly efficient. In our tests developing an optimization approach, we found that this simple linear approach usually only required three to four $\epsilon(0)$ calculations to converge. As converging on $\epsilon(0)$ is the slow step of the process, we actually found the simple line-search strategy³³ to be more efficient than grid-based approaches that required more $\epsilon(0)$ calculations to map out parameter space. One could possibly employ an even more efficient search strategy such as that used in the ForceBalance tool,⁵² but optimization would still require a computationally intensive search for each molecule of interest or a combined optimization over a set of small molecules. Rather than pursue such a specialized route, we chose to test if the knowledge gained from the hydroxyl optimization for methanol was transferrable to other small molecules containing hydroxyl moieties.

To apply the methanol hydroxyl optimization in a general manner, we assume that the hydrogen bonding between hydroxyls in neat methanol is representative of hydroxyl hydrogen bonding in other systems. Additionally, we assume that intramolecular perturbations of the electric field about a given hydroxyl group that would alter this behavior are treated by the semiempirical or quantum mechanical approach used in the setting of the partial charges for the model. Given these assumptions, we adopt new LJ σ and ϵ values, taken from the methanol optimization, for each hydroxyl oxygen atom. For the partial charges, we scale all hydroxyl group atoms by the amount observed for methanol, as shown in Figure 2. Any non-hydrogen atoms bonded to the α carbon are then treated as neutralization sites (the NC site in Figure 2A) to compensate for any charge difference introduced by this charge scaling. If there are multiple non-hydrogen atoms bonded to the α carbon (Figure 2B), the charge difference between the original and polarized set of hydroxyl atoms is distributed evenly among these bonded atoms.

RESULTS AND DISCUSSION

Dielectric corrected methanol provides general hydroxyl optimization rules. We applied the described optimization protocol to alter the force field parameters for the hydroxyl oxygen in a methanol molecule using the GAFF small molecule parameters and AM1-BCC partial charges. The results of this procedure are shown in Tables 1 and 2.

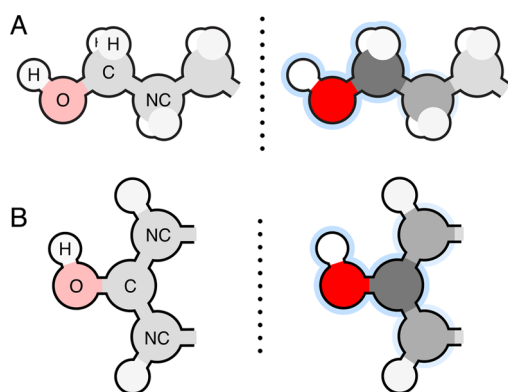


Figure 2. Illustration of the application of optimized methanol polarization to a (A) primary alcohol and (B) phenolic alcohol. The partial charges on all labeled atoms are scaled by the scaling factor from the methanol optimization with the exception of the NC atoms. In the primary alcohol, the single NC atom acts as a neutralization site. In the phenolic alcohol, the charge difference due to scaling of the hydroxyl group is split over the two NC atoms.

Table 1. Calculated Properties for the Original and Altered Methanol Models Alongside Experimental Values

| | ρ^a (kg/m ³) | ΔH_{vap}^b (kcal/mol) | $\epsilon(0)^a$ |
|-----------------------------|-------------------------------|--------------------------------------|-----------------|
| starting CH ₃ OH | 791.6 | 10.95 | 20.7 |
| final CH ₃ OH | 792.4 | 8.96 | 34.6 |
| experiment | 791.4 | 8.95 | 33.0 |

^aAt 293.15 K. ^bAt 298.15 K.

Table 2. LJ Parameters and Charge Scaling Values for the Hydroxyl Oxygen in the Initial and Altered Methanol Models

| | σ (Å) | ϵ (kcal/mol) | q scaling |
|-----------------------------|--------------|-----------------------|-------------|
| starting CH ₃ OH | 3.066 47 | 0.210 40 | 1.000 00 |
| final CH ₃ OH | 3.219 90 | 0.202 07 | 1.209 05 |

Table 1 shows how the properties for the methanol models successfully balance within the predefined experimental constraints. The density is sitting just within our predefined tolerance (1 kg/m³) of the experimental value while the other properties sit well within the $k_B T$ window for the ΔH_{vap} and $\Delta\epsilon(0) = 4$ window. The biggest change is seen for the $\epsilon(0)$ value in moving a quantity that has a 40% discrepancy down to a less than 5% difference. To accomplish this, the dipole moment of the methanol model needed an $\sim 20\%$ increase in magnitude (see Table 2). These modifications are similar to recently proposed modifications derived from matching Kirkwood–Buff integrals, though not quite of the same magnitude,⁵³ and also qualitatively similar to recent changes in the GROMOS force field for alcohols.⁵⁴ Our changes indicate that the condensed-phase polarization of methanol is underaccounted for in the AM1-BCC charge model and the HF/6-31G* basis-set electrostatic potential upon which it is fit,⁴¹ even given this basis set's 10% to 15% overpolarization relative to the gas phase expectation that is considered fortuitous for condensed-phase simulations.^{2,31,41} The experimental liquid-phase dipole moment for methanol is currently unknown, so while it is expected to be greater in magnitude than the gas-phase dipole moment, it is unclear if the optimized liquid state electric field is a truer representation of that in real methanol than the field used in other models.

We cannot expect other properties, such as the density, to remain static in the face of such a large polarization of the liquid-state methanol molecule. Thus, the hydroxyl oxygen σ expands to counter the increase in density that occurs with such a polarization. This increase in size simultaneously decreases the strength of alcohol model pair interactions and increases the molecular volume. While this change in oxygen atom size acts to counter the increased polarization, this does not mean that the structure of the liquid remains static. Figure 3 shows

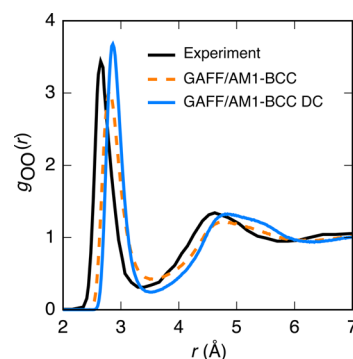


Figure 3. Oxygen–oxygen radial distribution functions for the starting GAFF/AM1-BCC methanol model (orange dashes), the final dielectric corrected (DC) model (blue), and neutron-diffraction studies (black).^{55,56} The topology adjustments in the final DC model make the liquid more structured, but they do little to the radial location of the existing methanol structure of the starting model.

the $g_{\text{OO}}(r)$ for methanol before and after the optimization process, alongside a similar curve from neutron-diffraction studies.^{55,56} We see an approximately 0.05 Å shift of peak maxima to longer distances with the new parameters, in poorer agreement with experimentally supported structural data. We also see that the peaks and troughs are amplified, indicating an increase in the structuring of the liquid, in better agreement with the experimentally supported structural data. These small positive and negative changes, coupled with the fact that the liquid structure cannot be unambiguously determined from experiments without using model fits, make it difficult for us to state if the liquid structure is better or worse relative to experiment. We can state that while the liquid density remains the same, the new parameters act to increase the structure of the liquid. If further changes in the structure are desired, one could consider including target data along with condensed-phase geometry optimizations in the fitting process.^{57,58}

In addition to liquid structure, experimental diffusion constant data are available for methanol at standard conditions, 2.28×10^{-5} cm²/s.⁵⁹ We computed the neat liquid diffusion constant for this methanol model before and after the optimization process. The original model has a diffusion constant of $(3.8 \pm 0.2) \times 10^{-5}$ cm²/s, which is considerably faster than experiment. This fast dynamic behavior is quite similar to that seen with the OPLS-AA force field.⁶⁰ After optimization, the diffusion constant drops to $(1.8 \pm 0.2) \times 10^{-5}$ cm²/s, slower than, but considerably closer to, the experimental value. This change is similar to that seen for the evolution of SPC to SPC/E water, where the additional polarization in SPC/E lowered the diffusion constant closer to the experimental value.^{61,62} This improvement comes despite dynamical performance not being considered in the optimization process. As suggested for the model liquid structure, using the diffusion constant, possibly in place of ΔH_{vap} , could be an

alternate optimization strategy to further the agreement in dynamical properties of general alcohols.

The scaling values listed here were used as general rules in the development of a simple tool for converting such hydroxyl groups in GROMACS topologies into similar dielectric corrected (DC) forms. The source code for perl and python versions of this utility is provided in the Supporting Information for interested readers.

New hydroxyls improve dielectric properties of other alcohols. The optimization improves the dielectric properties of methanol directly but is difficult to apply prospectively to new molecules. If improvement of the $\epsilon(0)$ is desired for models of other molecules with hydroxyl groups, it would be advantageous to not need to undergo such a process for each new molecule. In many cases, such an effort would be impossible due to the lack of relevant experimental data. This is a key concern for researchers wanting to use new molecular models as a step in a computational chemical process for pharmaceutical development, or in any other discovery setting. We therefore decided to test how transferrable these adjustments are to other more complex small molecules. This set, listed in the Supporting Information, consists of methanol and 40 additional molecules for which the experimental $\epsilon(0)$ and ΔG_{hyd} are known.

Figure 4A and 4B show the before and after application of DC hydroxyl modifications results for the calculation of the

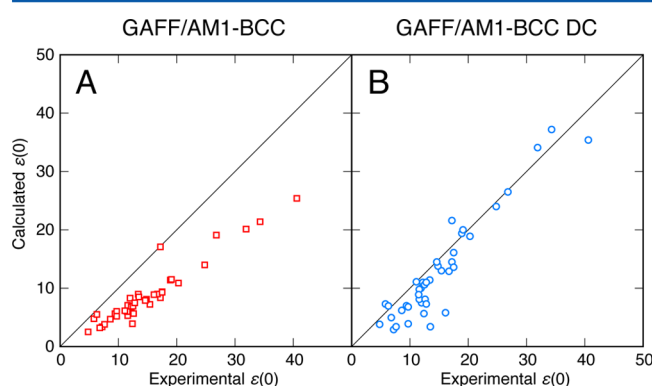


Figure 4. Scatterplots of the calculated $\epsilon(0)$ values versus experimental quantities for a set of 41 small molecules using (A) the GAFF small molecule parameters and AM1-BCC partial charges and (B) those same models after applying the methanol dielectric correction to all hydroxyl groups. For the GAFF/AM1-BCC models, the data are well-correlated ($R^2 = 0.91$) to an underpredicting slope of 0.65(3). The DC models show a similar correlation ($R^2 = 0.89$) but have a slope of 1.07(6), indicating improved agreement with experiment.

neat liquid $\epsilon(0)$ values at the experimentally measured temperatures. One striking feature of the original general alcohol models is the tight correlation but shallow slope for the calculated $\epsilon(0)$ versus experimental measurements. Regression analysis shows the model $\epsilon(0)$ to typically be about two-thirds of the experimental value. Applying the DC methanol modifications to all hydroxyl groups corrects this systematic underprediction, while not degrading the neat liquid densities.

The general trend for the change in $\epsilon(0)$ values when applying the DC methanol modifications was an approximately 50% increase, closing the gap between the calculations and experimental measurements. There were a few exceptions, the most glaring were cyclohexanol and cycloheptanol. In these neat liquids, the $\epsilon(0)$ changed in the opposite manner, an

average 50% decrease. It should be noted that the densities for the starting GAFF/AM1-BCC model liquids were already 10–20 kg/m³ less than experiment, and these values do not get “corrected” by the hydroxyl modifications. This indicates a potential issue may reside in the cycloalkane angle and/or torsion parameters, altering the liquid-state dipole orientation distributions in a manner inconsistent with the trends seen for other small molecules. Regardless, such occurrences were the exception rather than the norm, as only 4 of the 41 molecules showed a statistically significant decay of $\epsilon(0)$ away from the experimental value, with the noted cycloalcohol pair being the worst offenders. The majority of these alcohols showed dramatic enhancement, indicating that the DC methanol modifications are surprisingly transferable for the purpose of improving model $\epsilon(0)$ values.

ΔG_{hyd} trends couple with $\epsilon(0)$. One of the primary motivations behind improving the dielectric properties of model solutes is the potential for improvement in force field depictions of such solutes in other applications such as hydration and binding free energy calculations. Solvation (and desolvation) of ligand molecules is a critical aspect of ligand binding processes, and accurate predictions of ΔG_{hyd} are necessary for molecular modeling to be a useful contributor to pharmaceutical design and development. Transfer processes are governed by the dielectric permittivity of the mediums in implicit molecular models, and we expect an experimentally relevant depiction of the solute and environment interactions will be necessary for explicit models to be quantitative. The dielectric correction process does not use ΔG_{hyd} information for optimization, but it does target the quality of the model by polarizing hydroxyl groups to a level relevant to a condensed-phase hydrogen-bonding environment. Here, we test the DC hydroxyl modifications by comparing alchemical hydration free energy calculations on the original GAFF/AM1-BCC and modified forms. When performing ΔG_{hyd} calculations on a polarized model, it has been suggested that a work term for the polarization of the molecule (E_{pol}) be included in final result similar to how it is applied in the ΔH_{vap} calculation.^{27,63} Through personal correspondence with the suggesting authors, we were advised *not* to include such a term when using the AM1-BCC partial charges. Here, we follow this general advice; however, because we are systematically polarizing the hydroxyl groups beyond the AM1-BCC level, we include an additive *per-hydroxyl* E_{pol} term for polarizing methanol from the AM1-BCC partial charge level to that at the end of the optimization, 0.409 kcal/mol per-hydroxyl. This term is taken to be additive with the number of hydroxyl groups because we are only modifying the parameters of hydroxyl groups—we leave the remaining model parameters untouched. Note that the magnitude of this potential E_{pol} term is less than 1 kT, the chosen tolerance for the ΔH_{vap} optimization. This indicates that the new parameters are a valid fit to the computed ΔH_{vap} whether or not the E_{pol} term is included. The Supporting Information contains results with and without the E_{pol} term. We observe a net benefit whether we apply the E_{pol} term or not, though we see the best performance using this correction. This work does not resolve questions on whether or not such a term should be applied in general,^{24,27,63,64} but it shows how a simplified consideration of this term can influence hydration free energy calculations. Plots and error analysis of all these cases are provided in the Supporting Information for readers interested in additional comparison clarifications.

Figure 5A and 5B show the results of ΔG_{hyd} calculations before and after the application of the DC hydroxyl

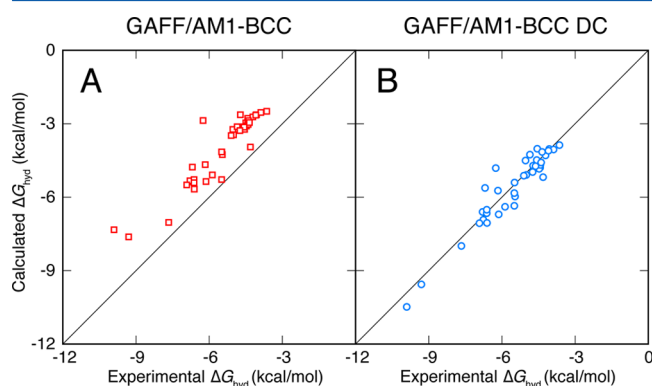


Figure 5. Scatterplots comparing the calculated to experimental ΔG_{hyd} for 41 alcohols depicted using the (A) GAFF and AM1-BCC parameters and (B) the DC hydroxyl modified forms. The GAFF/AM1-BCC mean signed error (MSE) of 1.43 kcal/mol is essentially eliminated (MSE of -0.07 kcal/mol) when using the DC hydroxyl modifications, and the root-mean-square error also drops from nearly $3k_{\text{B}}T$ (1.52 kcal/mol) down to under $k_{\text{B}}T$ (0.45 kcal/mol). The correlation of the data with experiment improves slightly as well, going from an R^2 of 0.87 to an R^2 of 0.91 when using DC hydroxyls.

modifications. Figure 5A highlights a key concern in hydration calculations of alcohols using GAFF and AM1-BCC partial charges. There is a nearly 1.5 kcal/mol systematic underestimation of the ΔG_{hyd} . When applying the DC hydroxyl modifications, this systematic error disappears, and we are left with accurate calculation results that agree extremely well with experiment. These results indicate that the neat liquid methanol environment from the optimization appears to be a good proxy for an aqueous hydrogen-bonding environment and quantitatively better than the essentially gas-phase environment in an AM1-BCC calculation. While we only have two solutes in this set with multiple hydroxyl groups (1,2-ethandiol and 1,3-propanediol), the DC hydroxyl modifications also appear to avoid potential issues related to nonadditivity of local polarization, taking ΔG_{hyd} values that are nearly 2 and 3 kcal/mol too unfavorable compared to values that are within 0.5 kcal/mol of experiments.

Environment is important at both the classical and quantum levels. As the methanolic environment provided by the DC hydroxyl modifications is beneficial for both neat liquid $\epsilon(0)$ and ΔG_{hyd} values calculated for general alcohols, we are interested in determining if the new LJ parameters for the hydroxyl oxygen atom could be used in conjunction with a QM method that better treats the condensed-phase environment's effect on the solute electric field and resulting partial charges. A better accounting of the condensed-phase environment could result in a truer representation of the electrostatics of the solute molecule. Ideally, we could set the partial charges for a given solute containing hydroxyl groups by fitting to a QM electrostatic potential calculated using a self-consistent reaction field (SCRF) or similar environment term, rather than using the modified AM1-BCC partial charges. To test this, we applied partial charges from MP2/cc-pVTZ SCRF calculations²⁴ to a small subset of phenolic and linear alcohol solutes and compared calculated $\epsilon(0)$ and ΔG_{hyd} values with similar results from GAFF/AM1-BCC and the DC modified models. The results of these calculations are shown in Figure 6A and 6B.

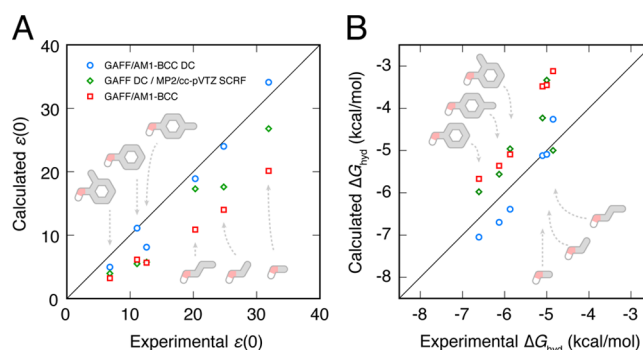


Figure 6. Comparison of calculated (A) $\epsilon(0)$ and (B) ΔG_{hyd} with experiment when using GAFF/AM1-BCC (red squares), the DC hydroxyl form (blue circles), and MP2/cc-pVTZ SCRF with the DC hydroxyl oxygen LJ parameters (green diamonds). By using the new LJ parameters, the neat liquid densities for methanol, ethanol, propan-1-ol, phenol, *o*-cresol, and *p*-cresol do not degrade when using a condensed-phase quantum calculation. The resulting $\epsilon(0)$ and ΔG_{hyd} are still not quite to the accuracy level of the simple approach outlined in this work.

It is apparent from the improved $\epsilon(0)$ values that the use of an SCRF polarizes the alcohols more than the semiempirical AM1-BCC approach. Still, the more complex QM method does not result in dielectric constants as accurate as our simple application of the polarization changes from a neat liquid optimization of methanol. Similarly, the hydration calculation shows improvement when using the more detailed QM method, but agreement with experiment is still not quite to the level of that from our DC hydroxyl modifications. While this is somewhat disappointing from the perspective of the higher-level QM approach, it does indicate that using the SCRF improves both dielectric and hydration properties. Additionally, the neat liquid optimization rules appear to provide a potential target benchmark for future development of environment terms in QM electric field calculations.

Blind predictions in the SAMPL4 experiment show general improved agreement with experiment. Our approach of correcting dielectric properties proved successful for a moderately sized set of small molecules, but our real goal is an approach which is transferable and predictive. Thus, it would be helpful to have blind studies to evaluate if this DC hydroxyl modification can be used in predictive situations. To this end, we contributed ΔG_{hyd} results both without and with the DC modifications to the SAMPL4 hydration prediction event.^{65–67} The results of these contributions are summarized in Table 3. For the actual SAMPL4 event, the E_{pol} contribution to ΔG_{hyd} was not included, but the results with this term are included here for completeness. The reported errors in Table 3 were computed as the standard deviation in the mean over 1000 bootstrap trials, with experimental values resampled with

Table 3. Analysis of Contributed Hydration Predictions Using Versions of GAFF/AM1-BCC Parameters in TIP3P Water for the 18 Hydroxyl Containing Solutes in the SAMPL4 Experiment

| force field | RMSE (kcal/mol) | MSE (kcal/mol) | AUE (kcal/mol) |
|-------------------------|-----------------|------------------|-----------------|
| unmodified | 1.77 ± 0.37 | 0.80 ± 0.39 | 1.40 ± 0.27 |
| DC, no E_{pol} | 1.38 ± 0.25 | -0.65 ± 0.30 | 1.02 ± 0.22 |
| DC | 1.51 ± 0.29 | -0.15 ± 0.35 | 1.12 ± 0.24 |

added Gaussian noise drawn from a distribution with a standard deviation equal to the experimental uncertainty.

Of the 47 small molecules in the full SAMPL4 set, 18 contained at least one hydroxyl group. Since there are a diverse set of functional groups represented in the SAMPL molecule sets, we would not expect the application of these DC hydroxyl modifications to result in perfect agreement with experimental values. However, the ~ 0.3 kcal/mol improvement shown in Table 3 in hydroxyl containing solutes shown does shift the overall set of predictions closer to experiment by 0.1 to 0.2 kcal/mol. We ran a paired *t*-test on GAFF/AM1-BCC versus GAFF/AM1-BCC DC and found that the latter is significantly better ($p = 0.008$). It is interesting to note from the Mean Signed Errors (MSE) that the DC hydroxyl modifications without the E_{pol} term improve the overall results by countering the systematically hydrophobic nature of the general force field, possibly skewing the predictions too hydrophilic. The inclusion of the E_{pol} term tempers this effect, yet still results in similar overall prediction quality.

While the aim of the SAMPL experiment is to inform rather than act as a competition between groups, we note that the overall performance of this simple force field + hydroxyl adjustments submission was regularly among the top three predictions across the various performance metrics.⁶⁷ These results indicate that further force field improvements using this DC strategy applied to other functional groups could be an effective route toward more accurate predictions of hydration from theoretical methods that rely upon the quality of small molecule force field parameters.

CONCLUSIONS

We have shown a method to incorporate experimental dielectric response information on neat liquids in classical molecular models of alcohol. By optimizing the LJ parameters and polarization of methanol to capture the density, $\epsilon(0)$, and ΔH_{vap} , we were able to determine simple hydroxyl modulation rules that can be applied in a general manner to other alcohols. We tested how these DC hydroxyl modifications affect the $\epsilon(0)$ for 41 small molecule alcohols, and we found that these changes are transferable and lead to substantially improved model property agreement with experiment. We applied these modifications in the air-to-water transfer of these molecules as well as in a blind prediction event and found that these changes improve agreement with experiment by eliminating an observed systematic hydrophobic bias in the hydration of GAFF/AM1-BCC alcohols. These simple modifications outperform detailed QM calculations using an SCRF environment coupling term and provide a potential benchmark for future developments in QM calculation methods.

These findings make for several clear observations. First, the relative simplicity and general nature of this effort to correct perceived flaws in existing classical molecular mechanics force fields indicates that there is room for improvement in the LJ and partial charge representations of standard chemical groups. Here, we employed an optimization strategy that utilized accepted experimentally measured thermodynamic properties as target quantities. This work on the simplest of alcohols resulted in a transferable approach that simultaneously improved dielectric and hydration properties of more complex alcohols.

Additionally, the improvement in ΔG_{hyd} results indicates that the methanolic environment from the hydroxyl optimization is a good analogue for the aqueous hydrogen-bonding condensed-

phase environment. Using such an environment when optimizing LJ and partial charge parameters helps resolve systematic errors in hydroxyl group hydration seen when using accepted classical force fields. We suspect a similar improvement in hydration could occur for other polar functional groups, such as amines and carbonyls, following optimization, as their polar neat liquid environments might also be reasonable analogues to polar aqueous environments.

While the optimization process applied here is specific to the originating force field, it appears to be general for a given charge model basis. We have applied a similar optimization to models using the OPLS-UA force field for alcohols, which are fit to a slightly less detailed basis set,⁵⁷ and arrived at similar end-point alterations to those seen here. This indicates that further development in condensed-phase QM electric field calculation methodology, possibly in conjunction with experimentally grounded efforts similar to those presented here, can potentially lead to even more quantitative and insightful future classical molecular simulations.

ASSOCIATED CONTENT

Supporting Information

Full tables of $\epsilon(0)$ and ΔG_{hyd} results, the source code for a perl program to apply DC modifications to hydroxyl groups in GROMACS topologies, and relevant structure and topology information for molecules investigated in this study can be found in the related Supporting Information. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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Notes

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REFERENCES

- (1) Jorgensen, W. L.; Tirado-Rives, J. The OPLS Force Field for Proteins. Energy Minimizations for Crystals of Cyclic Peptides and Crambin. *J. Am. Chem. Soc.* **1988**, *110*, 1657–1666.
- (2) Cornell, W. D.; Cieplak, P.; Bayly, C. I.; Gould, I. R.; Merz, K. M., Jr.; Ferguson, D. M.; Spellmeyer, D. C.; Fox, T.; Caldwell, J. W.; Kollman, P. A. A second generation force field for the simulation of proteins and nucleic acids. *J. Am. Chem. Soc.* **1995**, *117*, 5179–5197.
- (3) Jorgensen, W. L.; Maxwell, D. S.; Tirado-Rives, J. Development and Testing of the OPLS All-Atom Force Field on Conformational Energetics and Properties of Organic Liquids. *J. Am. Chem. Soc.* **1996**, *118*, 11225–11236.

- (4) MacKerell, A. D., Jr.; Bashford, D.; Bellott, M.; Dunbrack, R. L., Jr.; Evanseck, J. D.; Field, M. J.; Fischer, S.; Gao, J.; Guo, H.; Ha, S.; et al. All-Atom Empirical Potential for Molecular Modeling and Dynamics Studies of Proteins. *J. Phys. Chem. B* **1998**, *102*, 3586–3616.
- (5) Oostenbrink, C.; Villa, A.; Mark, A. E.; van Gunsteren, W. F. A Biomolecular Force Field Based on the Free Enthalpy of Hydration and Solvation: The GROMOS Force-Field Parameter Sets 53A5 and 53A6. *J. Comput. Chem.* **2004**, *25*, 1656–1676.
- (6) Wang, J.; Wolf, R. M.; Caldwell, J. W.; Kollman, P. A.; Case, D. A. Development and Testing of a General Amber Force Field. *J. Comput. Chem.* **2004**, *25*, 1157–1174.
- (7) Vanommeslaeghe, K.; Hatcher, E.; Acharya, C.; Kundu, S.; Zhong, S.; Shim, J.; Darian, E.; Guvench, O.; Lopes, P.; Vorobyov, I.; et al. CHARMM general force field: A force field for drug-like molecules compatible with the CHARMM all-atom additive biological force fields. *J. Comput. Chem.* **2010**, *31*, 671–690.
- (8) Jorgensen, W. L. Structure and Properties of Liquid Methanol. *J. Am. Chem. Soc.* **1980**, *102*, 543–549.
- (9) Jorgensen, W. L. Transferable Intermolecular Potential Functions. Application to Liquid Methanol Including Internal Rotation. *J. Am. Chem. Soc.* **1981**, *103*, 341–345.
- (10) Jorgensen, W. L.; Chandrasekhar, J.; Madura, J. D.; Impey, R. W.; Klein, M. L. Comparison of simple potential functions for simulating liquid water. *J. Chem. Phys.* **1983**, *79*, 926–935.
- (11) Jorgensen, W. L.; Jenson, C. Temperature Dependence of TIP3P, SPC, and TIP4P Water from NPT Monte Carlo Simulations: Seeking Temperatures of Maximum Density. *J. Comput. Chem.* **1998**, *19*, 1179–1186.
- (12) Abascal, J. L. F.; Sanz, E.; Fernández, R. G.; Vega, C. A potential model for the study of ices and amorphous water: TIP4P/Ice. *J. Chem. Phys.* **2005**, *122*, 234511.
- (13) Mahoney, M. W.; Jorgensen, W. L. A five-site model for liquid water and the reproduction of the density anomaly by rigid, nonpolarizable potential functions. *J. Chem. Phys.* **2000**, *112*, 8910–8922.
- (14) Horn, H. W.; Swope, W. C.; Pitera, J. W.; Madura, J. D.; Dick, T. J.; Hura, G. L.; Head-Gordon, T. Development of an improved four-site water model for biomolecular simulations: TIP4P-Ew. *J. Chem. Phys.* **2004**, *120*, 9665–9678.
- (15) Chatterjee, S.; Debenedetti, P. G.; Stillinger, F. H.; Lynden-Bell, R. M. A computational investigation of thermodynamics, structure, dynamics and solvation behavior in modified water models. *J. Chem. Phys.* **2008**, *128*, 124511.
- (16) Cerutti, D. S.; Rice, J. E.; Swope, W. C.; Case, D. A. Derivation of Fixed Partial Charges for Amino Acids Accommodating a Specific Water Model and Implicit Polarization. *J. Phys. Chem. B* **2013**, *117*, 2328–2338.
- (17) Nerenberg, P. S.; Jo, B.; So, C.; Tripathy, A.; Head-Gordon, T. Optimizing Solute-Water van der Waals Interactions To Reproduce Solvation Free Energies. *J. Phys. Chem. B* **2012**, *116*, 4524–4534.
- (18) Villa, A.; Mark, A. E. Calculation of the Free Energy of Solvation for Neutral Analogs of Amino Acid Side Chains. *J. Comput. Chem.* **2002**, *23*, 548–553.
- (19) Yu, H.; Geerke, D. P.; Liu, H.; van Gunsteren, W. F. Molecular dynamics simulations of liquid methanol and methanol-water mixtures with polarizable models. *J. Comput. Chem.* **2006**, *27*, 1494–1504.
- (20) Anisimov, V. M.; Vorobyov, I. V.; Roux, B.; MacKerell, A. D., Jr. Polarizable Empirical Force Field for the Primary and Secondary Alcohol Series Based on the Classical Drude Model. *J. Chem. Theory Comput.* **2007**, *3*, 1927–1946.
- (21) Leontyev, I. V.; Stuchebrukhov, A. A. Electronic continuum model for molecular dynamics simulations. *J. Chem. Phys.* **2009**, *130*, 085102.
- (22) Ponder, J. W.; Wu, C.; Ren, P.; Pande, V. S.; Chodera, J. D.; Schnieders, M. J.; Haque, I.; Mobley, D. L.; Lambrecht, D. S.; DiStasio, R. A., Jr.; et al. Current Status of the AMOEBA Polarizable Force Field. *J. Phys. Chem. B* **2010**, *114*, 2549–2564.
- (23) Shirts, M. R.; Pitera, J. W.; Swope, W. C.; Pande, V. S. Extremely precise free energy calculations of amino acid side chain analogs: Comparison of common molecular mechanics force fields for proteins. *J. Chem. Phys.* **2003**, *119*, 5740–5761.
- (24) Mobley, D. L.; Dumont, E.; Chodera, J. D.; Dill, K. A. Comparison of Charge Models for Fixed-Charge Force Fields: Small-Molecule Hydration Free Energies in Explicit Solvent. *J. Phys. Chem. B* **2007**, *111*, 2242–2254.
- (25) Mobley, D. L.; Bayly, C. I.; Cooper, M. D.; Shirts, M. R.; Dill, K. A. Small molecule hydration free energies in explicit solvent: An extensive test of fixed-charge atomistic simulations. *J. Chem. Theory Comput.* **2009**, *5*, 350–358.
- (26) Klimovich, P.; Mobley, D. L. Predicting hydration free energies using all-atom molecular dynamics simulations and multiple starting conformations. *J. Comput. Aided Mol. Des.* **2010**, *24*, 307–316.
- (27) Swope, W. C.; Horn, H. W.; Rice, J. E. Accounting for Polarization Cost When Using Fixed Charge Force Fields. II. Method and Application for Computing Effect of Polarization Cost on Free Energy of Hydration. *J. Phys. Chem. B* **2010**, *114*, 8631–8645.
- (28) Fennell, C. J.; Kehoe, C. W.; Dill, K. A. Modeling aqueous solvation with semi-explicit assembly. *Proc. Natl. Acad. Sci. U.S.A.* **2011**, *108*, 3234–3239.
- (29) Caleman, C.; van Maaren, P. J.; Hong, M.; Hub, J. S.; Costa, L. T.; van der Spoel, D. Force Field Benchmark of Organic Liquids: Density, Enthalpy of Vaporization, Heat Capacities, Surface Tension, Isothermal Compressibility, Volumetric Expansion Coefficient, and Dielectric Constant. *J. Chem. Theory Comput.* **2012**, *8*, 61–74.
- (30) MacCallum, J. L.; Tieleman, D. P. Calculation of the Water-Cyclohexane Transfer Free Energies of Neutral Amino Acid Side-Chain Analogs using the OPLS All-Atom Force Field. *J. Comput. Chem.* **2003**, *24*, 1930–1935.
- (31) Kuyper, L. F.; Hunter, R. N.; Ashton, D.; Merz, K. M., Jr.; Kollman, P. A. Free Energy Calculations on the Relative Solvation Free Energies of Benzene, Anisole, and 1,2,3-Trimethoxybenzene: Theoretical and Experimental Analysis of Aromatic Methoxy Solvation. *J. Phys. Chem.* **1991**, *95*, 6661–6666.
- (32) Wang, J.; Cieplak, P.; Kollman, P. A. How Well Does a Restrained Electrostatic Potential (RESP) Model Perform in Calculating Conformational Energies of Organic and Biological Molecules? *J. Comput. Chem.* **2000**, *21*, 1049–1074.
- (33) Fennell, C. J.; Li, L.; Dill, K. A. Simple liquid models with corrected dielectric constants. *J. Phys. Chem. B* **2012**, *116*, 6936–6944.
- (34) Keasler, S. J.; Charan, S. M.; Wick, C. D.; Economou, I. G.; Siepmann, J. I. Transferable Potentials for Phase Equilibria-United Atom Description of Five- and Six-Membered Cyclic Alkanes and Ethers. *J. Phys. Chem. B* **2012**, *116*, 11234–11246.
- (35) Berendsen, H. J. C.; van der Spoel, D.; van Drunen, R. GROMACS: A message-passing parallel molecular dynamics implementation. *Comput. Phys. Commun.* **1995**, *91*, 43–56.
- (36) van der Spoel, D.; Lindahl, E.; Hess, B.; Groenhof, G.; Mark, A. E.; Berendsen, H. J. C. GROMACS: Fast, Flexible, and Free. *J. Comput. Chem.* **2005**, *26*, 1701–1718.
- (37) Hess, B.; Kutzner, C.; van der Spoel, D.; Lindahl, E. GROMACS 4: Algorithms for Highly Efficient, Load-Balanced, and Scalable Molecular Simulation. *J. Chem. Theory Comput.* **2008**, *4*, 435–447.
- (38) Fennell, C. J.; Kehoe, C.; Dill, K. A. Oil/Water Transfer Is Partly Driven by Molecular Shape, Not Just Size. *J. Am. Chem. Soc.* **2010**, *132*, 234–240.
- (39) Pettersen, E. F.; Goddard, T. D.; Huang, C. C.; Couch, G. S.; Greenblatt, D. M.; Meng, E. C.; Ferrin, T. E. UCSF Chimera—a visualization system for exploratory research and analysis. *J. Comput. Chem.* **2004**, *25*, 1605–1612.
- (40) Jakalian, A.; Bush, B. L.; Jack, D. B.; Bayly, C. I. Fast, efficient generation of high-quality atomic charges. AM1-BCC model: I. Method. *J. Comput. Chem.* **2000**, *21*, 132–146.
- (41) Jakalian, A.; Jack, D. B.; Bayly, C. I. Fast, Efficient Generation of High-Quality Atomic Charges. AM1-BCC Model: II. Parameterization and Validation. *J. Comput. Chem.* **2002**, *23*, 1623–1641.
- (42) Wang, J.; Wang, W.; Kollman, P. A.; Case, D. A. Automatic atom type and bond type perception in molecular mechanical calculations. *J. Mol. Graphics Modell.* **2006**, *25*, 247–260.

- (43) Sousa da Silva, A. W.; Vranken, W. F. ACPYPE - AnteChamber PYthon Parser interface. *BMC Res. Notes* **2012**, *5*, 367.
- (44) Hess, B.; Bekker, H.; Berendsen, H. J. C.; Fraaije, J. G. E. M. LINCS: A Linear Constraint Solver for Molecular Simulations. *J. Comput. Chem.* **1997**, *18*, 1463–1472.
- (45) Parrinello, M.; Rahman, A. Polymorphic transitions in single crystals: A new molecular dynamics method. *J. Appl. Phys.* **1981**, *52*, 7182–7190.
- (46) Nosé, S. A molecular dynamics method for simulations in the canonical ensemble. *Mol. Phys.* **1984**, *52*, 255–268.
- (47) Hoover, W. G. Canonical dynamics: equilibrium phase-space distributions. *Phys. Rev. A* **1985**, *31*, 1695–1697.
- (48) Essman, U.; Perela, L.; Berkowitz, M. L.; Darden, T.; Lee, H.; Pedersen, L. G. A smooth particle mesh Ewald method. *J. Chem. Phys.* **1995**, *103*, 8577–8592.
- (49) Allen, M. P.; Tildesley, D. J. *Computer Simulations of Liquids*; Oxford University Press: New York, 1987.
- (50) Neumann, M. Dipole moment fluctuation formulas in computer simulations of polar systems. *Mol. Phys.* **1983**, *50*, 841–858.
- (51) Lide, D. R., Ed. *CRC Handbook of Chemistry and Physics*, 84th ed.; CRC Press, Inc.: Boca Raton, FL, 2004.
- (52) Wang, L.-P.; Head-Gordon, T.; Ponder, J. W.; Ren, P.; Chodera, J. D.; Eastman, P. K.; Martinez, T. J.; Pande, V. S. Systematic Improvement of a Classical Molecular Model of Water. *J. Phys. Chem. B* **2013**, *117*, 9956–9972.
- (53) Weerasinghe, S.; Smith, P. E. A Kirkwood-Buff Derived Force Field for Methanol and Aqueous Methanol Solutions. *J. Phys. Chem. B* **2005**, *109*, 15080–15086.
- (54) Horta, B. A. C.; Fuchs, P. F. J.; van Gunsteren, W. F.; Hünenberger, P. H. New Interaction Parameters for Oxygen Compounds in the GROMOS Force Field: Improved Pure-Liquid and Solvation Properties for Alcohols, Ethers, Aldehydes, Ketones, Carboxylic Acids, and Esters. *J. Chem. Theory Comput.* **2011**, *7*, 1016–1031.
- (55) Yamaguchi, T.; Hidaka, K.; Soper, A. K. The structure of liquid methanol revisited: a neutron diffraction experiment at -80°C and $+25^{\circ}\text{C}$. *Mol. Phys.* **1999**, *96*, 1159–1168.
- (56) Yamaguchi, T.; Hidaka, K.; Soper, A. K. The structure of liquid methanol revisited: a neutron diffraction experiment at -80°C and $+25^{\circ}\text{C}$. *Mol. Phys.* **1999**, *97*, 603–605.
- (57) Jorgensen, W. L. Optimized Intermolecular Potential Functions for Liquid Alcohols. *J. Phys. Chem.* **1986**, *90*, 1276–1284.
- (58) Jorgensen, W. L.; Briggs, J. M.; Contreras, M. L. Relative Partition Coefficients for Organic Solutes from Fluid Simulations. *J. Phys. Chem.* **1990**, *94*, 1683–1686.
- (59) Derlacki, Z. J.; Easteal, A. J.; Edge, A. V. J.; Woolf, L. A.; Roksandic, Z. Diffusion coefficients of methanol and water and the mutual diffusion coefficient in methanol-water solutions at 278 and 298 K. *J. Phys. Chem.* **1985**, *89*, 5318–5322.
- (60) Wensink, E. J. W.; Hoffmann, A. C.; van Maaren, P. J.; van der Spoel, D. Dynamic properties of water/alcohol mixtures studied by computer simulation. *J. Chem. Phys.* **2003**, *119*, 7308–7317.
- (61) Berendsen, H. J. C.; Postma, J. P. M.; van Gunsteren, W. F.; Hermans, J. In *Intermolecular Forces*; Pullman, B., Ed.; Reidel: Dordrecht, 1981; pp 331–342.
- (62) Berendsen, H. J. C.; Grigera, J. R.; Straatsma, T. P. The Missing Term in Effective Pair Potentials. *J. Phys. Chem.* **1987**, *91*, 6269–6271.
- (63) Swope, W. C.; Horn, H. W.; Rice, J. E. Accounting for Polarization Cost When Using Fixed Charge Force Fields. I. Method for Computing Energy. *J. Phys. Chem. B* **2010**, *114*, 8621–8630.
- (64) Chipot, C. Rational Determination of Charge Distributions for Free Energy Calculations. *J. Comput. Chem.* **2003**, *24*, 409–415.
- (65) Guthrie, J. P. A Blind Challenge for Computational Solvation Free Energies: Introduction and Overview. *J. Phys. Chem. B* **2009**, *113*, 4501–4507.
- (66) Guthrie, J. P. SAMPL4, a blind challenge for computational solvation free energies: the compounds considered. *J. Comput. Aided Mol. Des.* **2014**, DOI: 10.1007/s10822-014-9738-y.
- (67) Mobley, D. L.; Wymer, K. L.; Lim, N.; Guthrie, J. P. Blind prediction of solvation free energies from the SAMPL4 challenge. *J. Comput. Aided Mol. Des.* **2014**. Advance online publication. doi: 10.1007/s10822-014-9718-2.