Benchmarking Atomistic Simulations against the ThermoML Database: **Neat Liquid Densities and Static Dielectric Constants**

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Useful atomistic simulations in the condensed phase require accurate depictions of solvent. While experimental measurements of fundamental physical properties offer a straightforward approach for evaluating forcefield quality, the bulk of this information has been tied up in non-machine-readable formats. This inaccessibility hinders reproducible benchmarks of forcefield accuracy. Here we examine the feasibility of benchmarking atomistic forcefields against the NIST ThermoML database of physicochemical measurements, which aggregates thousands of experimental measurements in a portable, machine-readable, and self-annotating format. We present a detailed benchmark of the generalized Amber small molecule forcefield (GAFF) using the AM1-BCC charge model against measurements (specifically liquid densities and static dielectric constants at ambient pressure) extracted from ThermoML, and discuss the extent of available data for neat liquids. We show that empirical polarizability models correct systematic biases inherent in predicting dielectric constants with fixedcharged forcefields. [JDC: Will need to be convinced!]

Keywords: molecular mechanics forcefields; forcefield parameterization; forcefield accuracy; forcefield validation; mass density; static dielectric constant

I. INTRODUCTION

Recent advances in hardware and molecular dynamics software have provided routine access to atomistic simulations at the 100 ns timescale and beyond. Leveraging these advances in combination with GPU clusters, distributed computing, or custom hardware has brought the microsecond and milliseconds within reach. These dramatic advances in sampling, however, have revealed forcefields as a critical barrier for truly predictive simulation.

of numerous benchmarks [1] and enhancements [2–4], with key outcomes including the ability to fold fastfolding proteins, improved fidelity of water thermodyservables. Although small molecule forcefields have also been the subject of benchmarks [5] and improve-27 ments [6], such work has typically focused on small perturbations to specific functional groups. For example, a recent study found that modified hydroxyl nonbonded parameters led to improved prediction of static dielectrics and hydration free energies [6]. There are also outstanding questions of generalizability of param-33 eters. Will changes to a specific chemical moiety be com-

34 patible with seemingly unrelated improvements? Ad-35 dressing these questions requires agreement on shared 36 benchmarks that can be easily replicated with proposed 37 forcefield enhancements.

A key barrier in forcefield development is that many 39 experimental datasets are heterogeneous, paywalled, 40 and unavailable in machine-readable formats (although 41 notable counterexamples exist, e.g. RCSB [7], Free-42 Solv [8] and BMRB [9]). While this inconvenience 43 is relatively minor for benchmarking a single target 44 (e.g. water), it becomes prohibitive for studies span-Protein and water forcefields have been the subject 45 ning chemical space. To ameliorate problems of data 46 archival, the NIST Thermodynamics Research Center 47 has developed a IUPAC standard XML-based format— 48 ThermoML [10]—for storing physicochemical measurenamic properties, and improved prediction of NMR ob- 49 ments, uncertainties, and metadata. Experimental re-50 searchers publishing measurements in several journals 51 (J. Chem. Eng. Data, J. Chem. Therm., Fluid Phase 52 Equil., Therm. Acta, and Int. J. Therm.) are now guided 53 through a data archival process that involves sanity 54 checks and archival at the TRC (http://trc.nist. 55 gov/ThermoML.html).

> Here we examine the ThermoML archive as a poten-57 tial source for neat liquid density and static dielectric 58 constant measurements, with the goal of developing a 59 standard benchmark for validating these properties in 60 fixed-charge forcefields of drug-like molecules. These 61 two observables provide sensitive tests of forcefield ac-62 curacy that are nonetheless straightforward to calculate. 63 Using the ThermoML data, we evaluate the generalized 64 Amber small molecule forcefield (GAFF) [11] with the 65 AM1-BCC charge model [12, 13] and identify systematic 66 biases that might be improved upon.

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Filter	Mass Density	Static Dielectric
0. Single Component	130074	1649
1. Druglike Elements	120410	1649
2. Heavy Atoms	67897	1567
3. Temperature	36827	962
4. Pressure	13598	461
Liquid state	13573	461
6. Aggregate T, P	3573	432
7. Density+Dielectric	245	245

TABLE I. Number of ThermoML measurements matching sequentially applied filters.

II. RESULTS

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Neat Liquid Measurements in ThermoML

We performed a number of sequential queries to sum-70 marize the ThermoML content relevant for benchmarking organic molecule forcefields. Our aim is to explore 72 neat liquid data with functional groups relevant to drug-73 like molecules. We therefore applied the following se-74 quence of filters: has either density or static dielectric 75 measurements, contains a single component, contains 101 76 only druglike elements (H, N, C, O, S, P, F, Cl, Br), has 102 low heavy atom count (≤ 10), has ambient temperature [K] (270 $\leq T \leq$ 330), has ambient pressure [kPA] ₁₀₃ $_{80}$ m^{-3} (a proxy for liquid state). After applying these fil- $_{105}$ mark of electrostatic models. We therefore compare sim-81 ters, we also round all pressures within this range to ex-82 actly one atmosphere. We also round temperatures to one decimal place. These approximations are motived 84 by common data entry errors; for example, an exper-85 iment performed at water's freezing point at ambient 110 tematically underestimates the dielectric constants for $_{\text{86}}$ pressure might be entered as either 101.325 kPA or 100 $_{_{111}}$ nonpolar organics, with the predictions of $\epsilon\approx1.0\pm0.05$ $_{87}$ kPA, with a temperature of either 273 K or 273.15 K. The $_{112}$ being substantially smaller than the measured $\epsilon \approx 2$. Be-88 application of these filters (Table I) leaves 245 conditions 89 for which both density and dielectric data are available. 114 tronic polarization, we added a simple empirical correc-90 The functional groups present are summarized in Table 115 tion for polarization [17] that is based on counting the 91 II.

Benchmarking GAFF against ThermoML: Mass Density

Mass density has been widely used for parameter-94 izing and testing forcefields, particularly the Lennard Jones parameters [15, 16]. We therefore used the present ThermoML compilation as a benchmark of the GAFF 117 A similar polarization correction was used in the develagreement of 4 % on a different benchmark set.

Functional Group	Counts
1,2-aminoalcohol	4
1,2-diol	3
alkene	3
aromatic compound	1
carbonic acid diester	2
carboxylic acid ester	4
dialkyl ether	7
heterocyclic compound	3
ketone	2
lactone	1
primary alcohol	19
primary aliphatic amine (alkylamine)	2
primary amine	2
secondary alcohol	4
secondary aliphatic amine (dialkylamine)	2
secondary aliphatic/aromatic amine (alkylarylamine)	1
secondary amine	3
sulfone	1
sulfoxide	1
tertiary aliphatic amine (trialkylamine)	3
tertiary amine	3

TABLE II. Functional group counts present in the dataset. The number of unique compounds is 44. Functional group classification was performed using checkmol version 0.5 [14].

C. Benchmarking GAFF / AM1-BCC against ThermoML: Static Dielectric

As a measure of the electronic medium, the static di- $(100 \le P \le 102)$, and has density greater than 300 kg $_{104}$ electric constant of neat liquids provides a critical bench-106 ulations against the measurements in our ThermoML 107 compilation. Overall, we find the dielectric constants to be qualitatively reasonable, but with clear deviations 109 from experiment. In particular, GAFF / AM1-BCC syscause this deviation likely stems from the lack of elec-116 elements in a molecule:

$$\frac{\alpha}{\mathring{\mathbf{A}}} = 1.53n_C + 0.17n_H + 0.57n_O + 1.05n_N + 2.99n_S + 2.48n_P + 0.22n_F + 2.16n_{Cl} + 3.29n_{Br} + 5.45n_I + 0.32$$

$$\tag{1}$$

From the polarizability, one can correct the static dielectric using the following equation (from ref. [18]):

$$\epsilon_{corrected} = \epsilon_{MD} + 4\pi N \frac{\alpha}{\langle V \rangle}$$

AM1-BCC forcefield (Fig. 1). Overall, the densi- 118 opment of the TIP4P-EW water model [18]; however, ties show reasonable accuracy (RMS percent error: 3 % 119 the need is much greater for the nonpolar organics, as \pm 0.1%), consistent with previous studies [5] reporting 120 the missing polarizability is the dominant contribution 121 to the static dielectric constant. In the case of water,

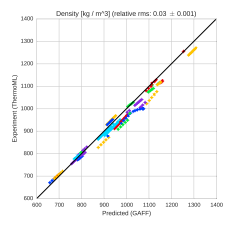


FIG. 1. Comparison of liquid densities between experiment and simulation. Liquid density measurements extracted from ThermoML are compared against densities predicted using the GAFF / AM1-BCC small molecule fixed-charge forcefield. Color groupings represent identical chemical species. Simulation error bars represent one standard error of the mean, with the number of effective (uncorrelated) samples estimated using pymbar. Experimental error bars indicate the standard deviation between independently reported measurements, when available, or author-reported standard deviations in ThermoML entries; for some measurements, neither uncertainty estimate is available. See **Section B** for further discussion of error.

122 the Sales polarizability model predicts a dielectric correction of 0.52, while 0.79 was used for the TIP4P-EW model. For comparison, we also applied the same empirical correction to the VirtualChemistry dataset [5, 19] and saw similarly improved agreement with experiment 127 for both the GAFF and OPLS forcefields (Fig. 7).

III. DISCUSSION

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A. Fitting Forcefields to Dielectric Constants

Recent forcefield development has seen a resurgence of papers fitting dielectric constants as primary data 6, 21]. However, a number of authors have pointed out potential challenges in constructing self-consistent 133 fixed-charge forcefields [22, 23]. 134

Interestingly, a recent work by Dill [22] pointed out that, for CCl₄, reasonable choices of point charges are 136 incapable of recapitulating the observed dielectric of = 2.2, instead producing dielectric constants in the range of $1.0 \le \epsilon \le 1.05$. This behavior is quite general: a fixed charge monopole force field predicts $\epsilon \approx 1$ for sev-145 unanticipated consequences.

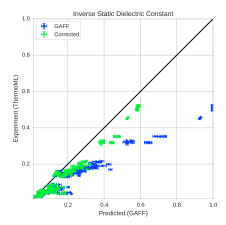


FIG. 2. Measured (ThermoML) versus predicted (GAFF / AM1-BCC) inverse static dielectrics (a). Simulation error bars represent one standard error of the mean estimated via block averaging with block sizes of 200 ps [20]. Experimental error bars indicate the larger of standard deviation between independently reported measurements and the authors reported standard deviations; for some measurements, neither uncertainty estimate is available. See section B for further discussion of error. The inverse dielectric $\frac{1}{\epsilon}$ is plotted instead of ϵ because $\frac{1}{\epsilon}$ is directly proportional to energy in continuum dielectric models: e.g. $U(r) = \frac{1}{4\pi\epsilon} \frac{q_1 q_2}{r} \propto \frac{1}{\epsilon}$.

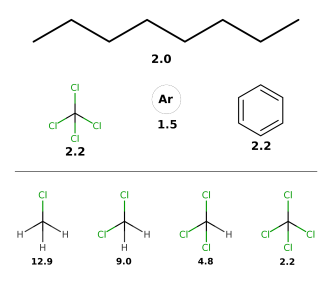


FIG. 3. (a). Measured static dielectric constants of various nonpolar or symmetric molecules [?]; fixed-charge forcefields give $\epsilon \approx 1$ for each species. (b). A congeneric series of chloromethanes show dielectric constants between 2 and 13.

Suppose, for example, that one attempts to fit forceeral nonpolar or symmetric molecules, but the measured 147 field parameters to match the static dielectric constants dielectric constants are instead $\epsilon \approx 2$ (Fig. 3). While 148 of CCl₄, CHCl₃, CH₂Cl₂, and CH₃Cl. In moving from this behavior is well-known and results from missing 149 the tetrahedrally-symmetric CCl₄ to CHCl₃, it suddenly physics of polarizability, we suspect it may have several 150 becomes possible to achieve the observed dielectric con-151 stant of 4.8 by an appropriate choice of point charges.

152 However, the model for CHCl₃ uses fixed point charges 203 to account for both the permanent dipole moment and the electronic polarizability, whereas the CCl₄ model ₂₀₄ contains no treatment of polarizability. We hypothesize that this inconsistency in parameterization may lead to 157 strange mismatches, where symmetric molecules (e.g. 158 benzene, CCl₄) have qualitatively different properties than closely related asymmetric molecules (e.g. toluene, $CHCl_3$). 160

As a possible real-world example, we imagine that 161 162 the missing polarizability could be important in accurate transfer free energies involving low-dielectric solvents. Using the Onsager model for the transfer free energy of a dipole (Eqn. 2) gives an error of $\Delta\Delta G =$ $\Delta G(\epsilon = 2.2) - \Delta G(\epsilon = 1)$ of 2 kcal / mol for the trans-167 fer of water ($a=1.93~\text{Å}~\mu=2.2\text{D}$) into a low dielectric 168 medium such as tetrachloromethane or benzene.

$$\Delta G = -\frac{\mu^2}{a^3} \frac{\epsilon - 1}{2\epsilon + 1} \tag{2}$$

Given their ease of measurement and direct connec-170 tion to long-range electrostatic interactions, static dielectric constants are potentially usable as primary data for forcefield parameterization efforts. Although this will require the use of forcefields with explicit polarizability, the inconsistency of fixed-charge models in low-dielectric media is sufficiently alarming to motivate further study of polarizable forcefields. In particular, continuum methods [24–26], induced dipole methods 178 [27, 28], and drude methods [29, 30] have been maturing rapidly. Finding the optimal balance of accuracy 180 and performance remains an open question; however, 181 the use of experimentally-parameterized direct polarization methods [31] may provide polarizability physics at a cost not much greater than fixed charge forcefields.

B. ThermoML as a Data Source

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sity and dielectric measurements present in ThermoML 239 ment during equilibration, applied every 25 steps. Par-[10, 32, 33] as a target for molecular dynamics force- 240 ticle mesh Ewald [39] was used with a long-range cutoff field validation. While densities and dielectric constants 241 of 0.95 nm and an long-range isotropic dispersion corhave been widely used in forcefield work, several as- 242 rection. [JDC: Can we report the automatically-selected pects of ThermoML make it a unique resource for the 243 PME parameters?] Simulations were continued until forcefield community. First, the aggregation, support, 244 density standard errors were less than 2×10^{-4} g / mL, and dissemination of ThermoML is supported by NIST, 245 as estimated using the equilibration detection module in whose mission makes these tasks a long-term priority. 246 pymbar 2.1 [40]. Trajectory analysis was performed us-Second, ThermoML is actively growing, through part- 247 ing OpenMM [37] and MDTraj [38]. Density data was nerships with several journals—new experimental mea- 248 output every 250 fs, while trajectory data was stored evsurements published in these journals are critically ex- 249 ery 10 ps. amined by the TRC and included in the archive. Finally, the files in ThermoML are machine readable via a formal XML schema, allowing facile access to thousands of 250 measurements. In the future, we hope to examine addi-201 tional measurement classes, including both mixture and 251 202 two-phase data.

IV. METHODS

ThermoML Processing

ThermoML XML files were obtained from the 206 the NIST TRC. To explore their content, we cre-207 ated a python (version 2.7.9) tool (ThermoPyl: 208 https://github.com/choderalab/ThermoPyL) 209 that munges the XML content into a spreadsheet-210 like format accessible via the Pandas (version 211 0.15.2) library. First, we obtained the XML schema 212 (http://media.iupac.org/namespaces/ 213 ThermoML/ThermoML.xsd) defining the layout of the 214 data. This schema was converted into a Python object 215 via PyXB 1.2.4 (http://pyxb.sourceforge.net/). 216 Finally, this schema and Pandas was used to extract the 217 data and apply the data filters described above.

B. Simulation

Boxes of 1000 molecules were constructed using 220 PackMol [34]. AM1-BCC [12, 13] charges were gen-221 erated using OpenEye Toolkit 2014-6-6 [35], using 222 the oequacpac.OEAssignPartialCharges module 223 with the OECharges_AM1BCCSym. The selected con-224 former was then processed using antechamber in Am-225 berTools 14 [36]. The resulting AMBER files were con-226 verted to OpenMM [37] ffxml forcefield XML files. Sim-227 ulation code used libraries gaff2xml 0.6, TrustButVer-228 ify 0.1, OpenMM 6.2 [37], and MDTraj 1.2 [38]. [TODO: 229 Provide a script to install all of these versions via 230 conda.

Molecular dynamics simulations were performed us-232 ing OpenMM 6.2 [37] using a Langevin integrator (with collision rate 1 ps^{-1}) and a 1 fs timestep; interestingly, ²³⁴ we found that a 2 fs timestep led to insufficient accuracy 235 in equilibrium densities (Table III). [JDC: Cite Langevin 236 integrator used in OpenMM.] Pressure coupling at 1 at-237 mosphere was achieved with a Monte Carlo barostat uti-The present work has focused on the neat liquid den- 238 lizing molecular scaling and automated step size adjust-

V. CONCLUSIONS

 ThermoML is a potentially useful resource for the forcefield community

 We have curated a subset of ThermoML for neat liquids with druglike atoms, with thousands of densities and hundreds of dielectrics

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 Empirical polarization models correct a systematic bias in comparing fixed-charge forcefields to static dielectric constants

VI. ACKNOWLEDGEMENTS

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TABLE III. To probe the systematic error from finite time-step integration, we examined the timestep dependence of butyl acrylate density. The number of effective samples was estimated using pymbar's statistical inefficiency routine [40]. To approximate the timestep bias, we compare the density expectation ($\langle \rho \rangle$) to values calculated with a 0.5fs timestep. We find a 2fs timestep leads to systematic biases in the density on the order of 0.2%, while 1fs reduces the systematic bias to less than 0.1%—we therefore selected a 1fs timestep for the present work, where we aimed to achieve three digits of accuracy in density predictions.

Appendix A: Supplementary Information

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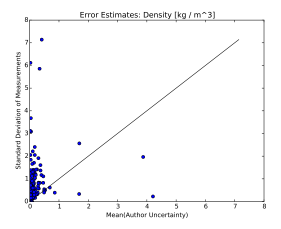
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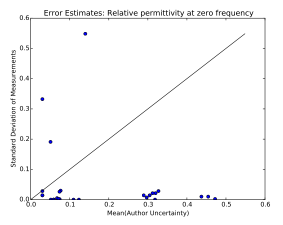
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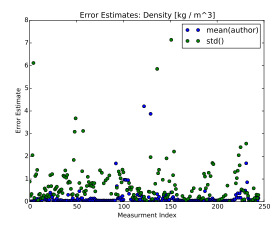
All information below this point will eventually be pulled into a separate SI. This will happen closer to submission, as the formatting may be journal-specific. The references may be split in two as well, depending on journal.

- Table: Timestep-dependence of density
- Figure: Error analysis for ThermoML dataset
- Table (CSV File): ThermoML Dataset used in present analysis.

Appendix B: Assessment of experimental error in ThermoML measurements







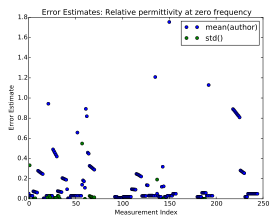


FIG. 4. Assessment of experimental error in ThermoML data. To assess the experimental error in our benchmark set, we consider two orthogonal error measurements. The first is the mean of the uncertainties reported by the measurement au-



FIG. 5. Comparison of simulated and experimental densities for all compounds. Measured (blue) and simulated (green) densities are shown in units of kg/m^3 .

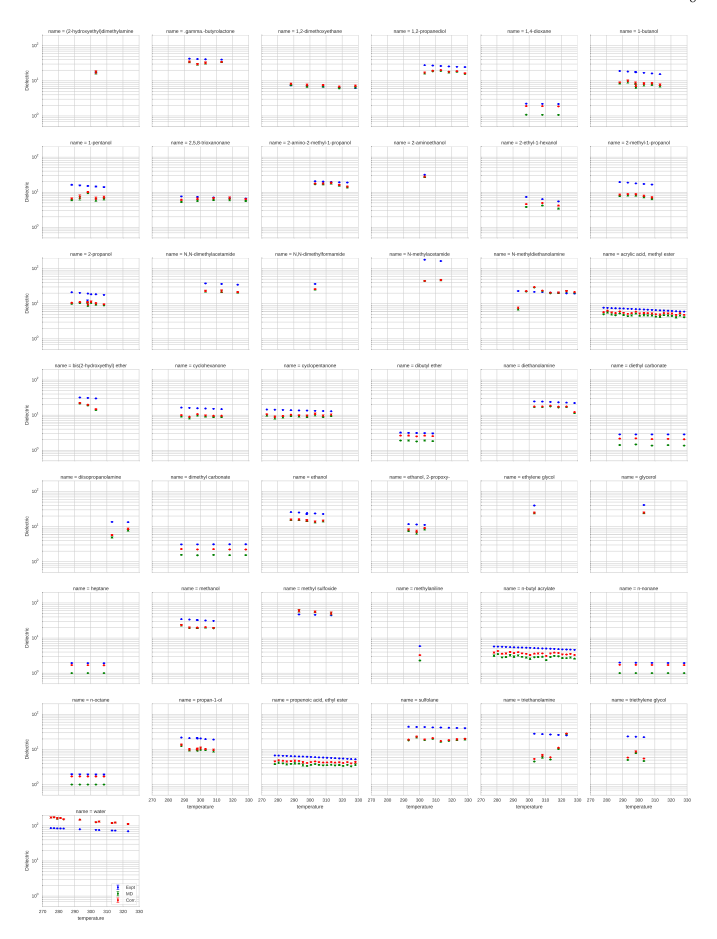
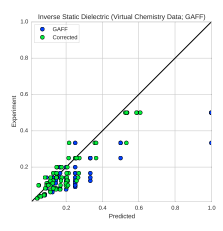


FIG. 6. Comparison of simulated and experimental static dielectric constants for all compounds. Measured (blue), simulated (green), and polarizability-corrected simulated (red) static dielectric constants are shown for all compounds. Note that dielectric constants, rather than inverse dielectric constants, are plotted here.



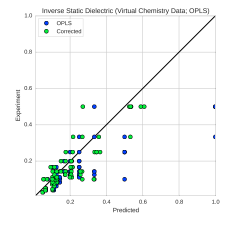


FIG. 7. Comparison of measured and simulated dielectric constants from the virtualchemistry dataset with and without polarizability correction. Measured (blue), MD (green), and MD + polarizability-corrected (red) dielectrics for the Virtual Chemistry dataset [5, 19].

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