Quantum chemistry structures and properties of 134 kilo molecules

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Computational de novo design of new drugs and materials requires rigorous and unbiased exploration of chemical compound space. However, large uncharted territories persist due to its size scaling combinatorially with molecular size. We report computed geometric, energetic, electronic, and thermodynamic properties for 134k stable small organic molecules made up of CHONF. These molecules correspond to the subset of all 133,885 species with up to nine heavy atoms (CONF) out of the GDB-17 chemical universe of 166 billion organic molecules. We report geometries minimal in energy, corresponding harmonic frequencies, dipole moments, polarizabilities, along with energies, enthalpies, and free energies of atomization. All properties were calculated at the B3LYP/6-31G(2df,p) level of quantum chemistry. Furthermore, for the predominant stoichiometry, C₇H₁₀O₂, there are 6,095 constitutional isomers among the 134k molecules. We report energies, enthalpies, and free energies of atomization at the more accurate G4MP2 level of theory for all of them. As such, this data set provides quantum chemical properties for a relevant, consistent, and comprehensive chemical space of small organic molecules. This database may serve the benchmarking of existing methods, development of new methods, such as hybrid quantum mechanics/machine learning, and systematic identification of structure-property relationships.

Design Type(s) in silico design • data integration
Measurement Type(s) Computational Chemistry
Technology Type(s) quantum chemistry computational method
Factor Type(s) level of theory
Sample Characteristic(s)

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Background & summary

The goal of computationally designing novel materials and molecules with desired physicochemical properties is yet to be achieved. High-throughput screening represents the most straightforward approach towards materials design. However, it presupposes that all assumptions and approximations inherent to the employed modeling techniques are applicable to the entire chemical compound space, which is the space populated by all stable molecules or materials. Furthermore, due to the combinatorial scaling of chemical space with molecular size, it is difficult to explore or even navigate. Conclusive insights about the domain of applicability (transferability) are lacking even for the most popular first principle quantum chemistry methods. For example, the reliability and accuracy of density functional theory is known to dramatically depend on chemical composition and atomistic configurations, highlighting the importance of reliable experimental or high-level quantum chemistry state-of-the-art results. Unfortunately, the systems reported are typically small, which implies the existence of severe selection bias. One can therefore question how representative they are. The problem of representative diversity has triggered the design of special purpose chemical space libraries for method validation or molecular design.

Here, we report molecular structures and properties obtained from quantum chemistry calculations for the first 134k molecules of the chemical universe GDB-17 database, covering a molecular property set of unprecedented size and consistency. The data-set corresponds to the GDB-9 subset of all neutral molecules with up to nine atoms (CONF), not counting hydrogen. The molecular size distribution of all 134k molecules is shown in Fig. 1. This data set contains small amino acids, such as GLY, ALA, as well as nucleobases cytosine, uracil, and thymine. Also pharmaceutically relevant organic building blocks, such as pyruvic acid, piperazine, or hydroxy urea are included. Among the 134k molecules, there are 621 stoichiometries, among which C_7H_{10}O_2 dominates with 6,095 constitutional isomers for which atomization energies and radii of gyration also are on display in Fig. 1.

For all 134k molecules, we have calculated equilibrium geometries, frontier orbital eigenvalues, dipole moments, harmonic frequencies, polarizabilities, and thermochemical energetics corresponding to atomization energies, enthalpies, and entropies at ambient temperature. These properties have been obtained at the B3LYP/6-31G(2df,p) level of theory which forms the basis for the more accurate state-of-the-art G4 methods which are on par with experimental accuracy. For the 6,095 constitutional isomers of the predominant stoichiometry, C_7H_{10}O_2, we report the energetics at the significantly more accurate G4MP2 level of theory.

This report is structured as follows. We first describe the genesis of the results. Thereafter, we discuss the validation of our DFT results by comparison to (i) G4MP2, (ii) G4, and (iii) CBS-QB3 results for
100 molecules, randomly chosen out of the 134k set. This data can serve the development, training and evaluation of inductive statistical data analysis-based machine learning (ML) models\textsuperscript{13}. It might also assist the search and discovery of hitherto unknown trends, structure-property relationships, and molecular materials design\textsuperscript{14,15}.

**Methods**

**Generation of atomic coordinates**

Starting with ref. 11, we use all SMILES\textsuperscript{16} strings for molecules with up to nine heavy atoms. Cations, anions, and molecules containing S, Br, Cl, or I, have been excluded, resulting in 133,885 molecules. 1,705 zwitterions have been kept in the data due to their occurrence in small biomolecules, such as amino acids. Initial Cartesian coordinates for all molecules were generated by parsing the corresponding SMILES strings using Corina (Version 3.491 2013)\textsuperscript{17}. We subsequently carried out geometry relaxations at the PM7 semi-empirical level of theory using MOPAC (Version 13.136L 2012)\textsuperscript{18}. In the PM7 calculations, we invoked tight electronic and geometric convergence thresholds, using *precise* keyword. PM7 equilibrium geometries have subsequently been used as input for B3LYP geometry relaxations using Gaussian 09\textsuperscript{19}. We iteratively refined the electronic and geometry thresholds. For the first iteration, Gaussian 09’s default electronic and geometry thresholds have been used for all molecules. For those molecules which failed to reach SCF convergence *ultrafine* grids have been invoked within a second iteration for evaluating the XC energy contributions. Within a third iteration on the remaining unconverged molecules, we identified those which had relaxed to saddle points, and further tightened the SCF criteria using the keyword *scf*(maxcycle=200, verytight). All those molecules which still featured imaginary frequencies entered the fourth iteration using keywords, opt(calcfc, maxstep=5, maxcycles=1000). calcfc constructs a Hessian in the first step of the geometry relaxation for eigenvector following. Within the fifth and final iteration, all molecules which still failed to reach convergence, have subsequently been converged using opt(calcall, maxstep=1, maxcycles=1000). calcall constructs a Hessian for all steps through the geometry relaxation. After all these measures taken, eleven problematic molecules still failed to converge to a minimal geometry. Out of these eleven molecules, six can be converged with low threshold using the keyword *scf*(maxcycle=200, verytight). All those molecules which still featured imaginary frequencies entered the fourth iteration using keywords, opt(loose), opt(calcfc, maxstep=5, maxcycles=1000). calcfc constructs a Hessian in the first step of the geometry relaxation for eigenvector following. Within the fifth and final iteration, all molecules which still failed to reach convergence, have subsequently been converged using opt(calcall, maxstep=1, maxcycles=1000). In the remaining five there were two near-linear molecules which converged to saddle points with very low imaginary frequencies (\(\omega_0 < 100 \text{cm}^{-1}\)) for the lowest mode. In the *readme.txt* file of this report, all these 11 molecules are specified using their indices in the database. In the case of the 6,095 constitutional isomers of C\textsubscript{7}H\textsubscript{10}O\textsubscript{2}, all molecules converged to local minima during the B3LYP geometry relaxation. To compute atomization energies, we have also performed spin-unrestricted calculations for all atoms with spin-multiplicities 2, 3, 4, 3, 2 for the atoms H, C, N, O, F, respectively.

**Data Records**

Molecular structures and properties are publicly available at Figshare (Data Citation 1) in a plain text XYZ-like format described below. Deposited files include the 133, 885 GDB-1 to GDB-9 molecules (dsgdb9nsd.xyz.tar.bz2), the 6,095 constitutional isomers of C\textsubscript{7}H\textsubscript{10}O\textsubscript{2} (dsC7O2H10nsd.xyz.tar.bz2), the 100 validation molecules (see Table 1) enthalpies of atomization (validation.txt), and atomic reference data (atomref.txt).

**File format**

For each molecule, atomic coordinates and calculated properties are stored in a file named *dataset_index.xyz*. The XYZ format (originally developed for the XMol program by the Minnesota Supercomputer Center) is a widespread plain text format for encoding Cartesian coordinates of molecules, with no formal specification. It contains a header line specifying the number of atoms \(n_a\), a comment line, and \(n_a\) lines containing element type and atomic coordinates, one atom per line. We have extended this format as indicated in Table 2. Now, the comment line is used to store all scalar properties.

| Reference | MAE  | RMSE | maxAE |
|-----------|------|------|-------|
| G4MP2     | 5.0  | 6.1  | 16.0  |
| G4        | 4.9  | 5.9  | 14.4  |
| CBS-QB3   | 4.5  | 5.5  | 13.4  |

*Table 1*. Validation of atomization enthalpies at B3LYP/6-31G(2df,p)-level. For 100 molecules randomly drawn out of the pool of 134k molecules, mean absolute error (MAE), root mean square error (RMSE), and maximal absolute error (maxAE) with respect to more accurate reference methods are reported. All values are in kcal/mol.
Mulliken charges are added as a fifth column. Harmonic vibrational frequencies, SMILES and InChI are appended as respective additional lines.

Properties
All molecular geometries were relaxed, and properties calculated, at the DFT/B3LYP/6-31G(2df,p) level of theory. The list of properties of the 134k molecules is summarized in Table 3. For a subset of 6,095 isomers of C_7H_{10}O_2, energetics (properties 12–16) were additionally calculated at the G4MP2 level of theory. For a validation set of 100 randomly drawn molecules from the 133,885 GDB-9 set, enthalpies of atomization were calculated at the DFT/B3LYP/6-31G(2df,p), G4MP2, G4 and CBS-QB3 levels of theory.

Technical Validation
Validation of geometry consistency
To validate the consistency of the relaxed B3LYP geometries, we have used them to generate the corresponding InChI strings with Corina and Open Babel (Version 2.3.0 2011). InChI corresponds to Workshop 1.020.

Table 2. XYZ-like file format for molecular structure and properties. \( n_a \) = number of atoms.

| No. | Property | Unit | Description |
|-----|----------|------|-------------|
| 1   | tag      | —    | ‘gdb9’ string to facilitate extraction |
| 2   | \( i \)  | —    | Consecutive, 1-based integer identifier |
| 3   | \( A \)  | GHz  | Rotational constant |
| 4   | \( B \)  | GHz  | Rotational constant |
| 5   | \( C \)  | GHz  | Rotational constant |
| 6   | \( \mu \) | D    | Dipole moment |
| 7   | \( \alpha \) | \( \AA^3 \) | Isotropic polarizability |
| 8   | \( \epsilon_{\text{HOMO}} \) | Ha   | Energy of HOMO |
| 9   | \( \epsilon_{\text{LUMO}} \) | Ha   | Energy of LUMO |
| 10  | \( \epsilon_{\text{gap}} \) | Ha   | Gap (\( \epsilon_{\text{LUMO}} - \epsilon_{\text{HOMO}} \)) |
| 11  | \( \langle R^2 \rangle \) | \( a_0^2 \) | Electronic spatial extent |
| 12  | zpve    | Ha   | Zero point vibrational energy |
| 13  | \( U_0 \) | Ha   | Internal energy at 0 K |
| 14  | \( U \)  | Ha   | Internal energy at 298.15 K |
| 15  | \( H \)  | Ha   | Enthalpy at 298.15 K |
| 16  | \( G \)  | Ha   | Free energy at 298.15 K |
| 17  | \( C_v \) | cal/mol\( K \) | Heat capacity at 298.15 K |

Table 3. Calculated properties. Properties are stored in the order given by the first column.
The resulting strings have been compared to the InChI strings obtained from the initial Cartesian coordinates (generated by Corina using the original GDB-17 SMILES strings). See Fig. 2 for a flow-chart of this consistency check. Out of the 134k molecules, 3,054 molecules did not pass this test. This is due to the fact that SMILES and InChI representations are not unique because transformation of Cartesian coordinates to string based chemical identifiers is prone to implementation specific artifacts. For molecules with same topology, small differences in interatomic distances, bond and dihedral angles can lead to different molecular graphs encoded by the string. To quantify this artifact, the distribution of Coulomb-matrix distances \( D_{ij} \) in Hartree, using the Manhattan or L1 metric, between Corina generated Cartesian coordinates (see Fig. 2) and B3LYP coordinates is on display in Fig. 3 for all the 3,054 molecules.

Consider, for example, molecule indexed 58 in the 134k set, which is among the 3,054 molecules for which the consistency check failed. Its original GDB-17 SMILES corresponds to NC(=N)C#N, and
Cartesian coordinates can be generated using Corina. When feeding back these coordinates to Open Babel to perform the inverse task of reproducing the initial SMILES string, \([\text{NH}]C(=\{\text{NH}_2\})\#N\) is obtained instead. By contrast, performing first a geometry relaxation of the Corina generated Cartesian coordinates using PM7 followed by B3LYP, and only then parsing through Open Babel, recovers the original SMILES string. In this case, \(D_1\) using the Coulomb-matrices of Corina and B3LYP geometries is rather small (~27 Ha) when compared to distances between geometries of other failed molecules, see distance distribution plot in Fig. 3. By contrast, failed molecules with large \(D_1\) between Corina and B3LYP geometries, see Fig. 3, correspond to molecules for which the B3LYP relaxation induces dramatic distortion with significant alteration and rearrangement of covalent bonding patterns. As a result, parsing back these geometries with Open Babel yields different SMILES strings. Note that all the 6,095 constitutional isomers of \(\text{C}_6\text{H}_4\text{O}_2\) for which G4MP2 calculations have been performed, passed this geometry consistency check, shown as a flow-chart in Fig. 2.

**Validation of quantum chemistry results**

All 134k molecules have been modeled using B3LYP/6-31G(2df,p) based DFT. Previously, B3LYP has been validated for several subsets, containing up to a few hundred small molecules. These benchmarks are of limited use since they are not necessarily sufficiently representative for gauging B3LYP’s performance in general. In the case of DFT’s systematic errors this issue is particularly pertinent\(^\text{22}\). Experimental data assembled in the NIST database is very sparse by comparison to our 134k organic molecules made up of CHONF atoms. Consequently, we have performed additional benchmark calculations for a subset of 100 randomly selected molecules using high level theories G4MP2\(^\text{12}\), G4\(^\text{23}\), and CBS-QB3\(^\text{24,25}\).

The predictive power of the G4MP2 method is widely considered to be on par with experimental uncertainties. For example, comparison to the G3/05 test set\(^\text{26,27}\) with 454 experimental energies (including enthalpies of formation, ionization potentials, electron affinities, proton affinities, and hydrogen bond energies) of small molecules yields MAE, and RMSE of 1.0, and 1.5 kcal/mol, respectively\(^\text{22}\). For the same properties and molecules, the slightly more accurate, and considerably more expensive method G4\(^\text{12,23}\) yields errors of MAE = 0.8 kcal/mol, RMSE = 1.2 kcal/mol. G4MP2 has been shown to deviate only by 1.4 kcal/mol from 261 bond dissociation enthalpies computed with the highly accurate W1w composite procedure\(^\text{28,29}\) for the BDE261 data set\(^\text{28}\). Consequently, we believe these calculations to be sufficiently suitable to validate the quality of the B3LYP energetics predictions. Various resulting deviations are summarized in Table 1. For the 100 molecules, the mean absolute error of B3LYP heats of atomization amounts to no more than 5 kcal/mol.

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Data Citation
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Author Contributions
All authors designed and performed research, and wrote the paper.

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