Systematic Microsolvation Approach with a Cluster-Continuum Scheme and Conformational Sampling

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Abstract

Solvation is a notoriously difficult and nagging problem for the rigorous theoretical description of chemistry in the liquid phase. Successes and failures of various approaches ranging from implicit solvation modeling through dielectric continuum embedding and microsolvated quantum chemical modeling to explicit molecular dynamics highlight this situation. Here, we focus on quantum chemical microsolvation and discuss an explicit conformational sampling ansatz to make this approach systematic. For this purpose, we introduce an algorithm for rolling and automated microsolvation of solutes. Our protocol takes conformational sampling and rearrangements in the solvent shell into account. Its reliability is assessed by monitoring the evolution of the spread and average of the observables of interest.

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1 Introduction

Solvation can, depending on the type of solvent into which a molecular system is immersed, strongly modulate the properties of a solute. For example, the kinetics of reactions in solution can be affected by the solvent. To that end, experimental studies have investigated the interactions between solute and solvent (see, for examples, Refs. 3–5). A solvent can take the role of an observer in which it assists the course of a reaction solely through non-covalent intermolecular interactions. It can also actively participate in a reaction in such a way that well-structured intermediates including solvent molecules are formed. Therefore, the accurate theoretical description of chemical processes requires not only the elucidation of all relevant intermediates and elementary reactions (see Refs. 6–8) but also adequate modeling of the reactants’ environment.

There are three main approaches of including solvation effects in theoretical models: implicit solvation, hybrid cluster-continuum schemes, and explicit solvation. They differ in accuracy and computational cost. In the following, the different approaches are briefly reviewed mainly in the light of their applicability to the study of chemical reaction mechanisms.

Molecular dynamics simulations can describe a solute’s dynamic surrounding at a given temperature. Particularly suited for the solvation of reactive species are \textit{ab initio} molecular dynamics simulations\textsuperscript{9} as they do not require the choice of some hard-wired interaction potentials, but rely on the exactly known expression for the non-relativistic electromagnetic interactions of elementary particles. However, as the configuration space can become very large, computational costs of carrying out first-principles calculations grow rapidly. As a result, they cannot be performed for every intermediate in large reaction networks.\textsuperscript{7,8,10–13} This issue can be overcome by the application of a reactive force-field.\textsuperscript{14–16} Unfortunately, next to the reduced accuracy, force-field parameters will, in general, not be available for any type of system which limits their applicability. For that reason, hybrid quantum-mechanical/molecular-mechanical (QM/MM) approaches have been frequently applied to explore complex systems with many degrees of freedom such as reactions in aqueous solution (e.g., Ref. 17) and reviews by
Senn and Thiel\textsuperscript{18,20}. Many studies have been devoted to studying the effect of the number of explicitly treated solvent molecules on chemical reactivity and optical properties (for examples, see Refs. \textsuperscript{21–26}). Recently, Boereboom et al.\textsuperscript{27} explored multiscale approaches for the description of a reversible and highly solvent-sensitive nucleophilic bond formation reaction. These studies demonstrate that, while the size of the solute affects the number of QM water molecules necessary to achieve convergence, additional system-specific properties such as solvent polarity, electronic structure of the solute, and solute-solvent interactions determine the required number of solvent molecules.

An implicit solvent model simplifies the interactions between the solute and the solvent by describing the solvent as a polarizable medium with a solvent specific dielectric constant.\textsuperscript{28–30} The solute is then placed in a cavity formed by this medium and the interaction between the solute and solvent is calculated at the cavity boundaries. There exist many implicit solvent models including the polarized continuum model (PCM),\textsuperscript{28} the conductor-like screening model (COSMO),\textsuperscript{31} integral equation formalism (IEFPCM),\textsuperscript{32} and COSMO for ‘real solvents’ (COSMO-RS),\textsuperscript{33} and the reference interaction site model (RISM).\textsuperscript{34} Implicit solvent models remain popular mainly for three reasons: (i) implicit solvation is, by far, the computationally cheapest approach of modeling solvation effects, (ii) most quantum chemistry programs allow for the activation of an implicit solvation model in a straightforward fashion, and (iii) a continuum model can provide efficient access to free energies, even if the electrostatic contribution to the enthalpy is negligible.\textsuperscript{35–37}

However, implicit solvent models fall short in many practical cases. First, they are known to describe strong (directed) interactions between solute and solvent (e.g., hydrogen bonds) poorly. This is often the case when the solute is charged (or contains a charged moiety) and the solvent is polar. Second, they will fail if solvent molecules can react with the solute. This includes cases in which intermediates are formed, protons are shuttled, or functional groups such as aldehydes and ketones undergo tautomerization.

In 2015, Plata and Singleton’s study of the Morita-Baylis-Hillman reaction highlighted issues of implicit solvation models for this specific reaction.\textsuperscript{38} Recent work on this reaction by Basdogan and Keith\textsuperscript{39} demonstrated the necessity of explicit solvation when studying organic reaction mechanisms.
However, the importance of explicit solvation has been shown numerous times for chemical reactions (see, e.g., Refs. 27,40–43). Explicit solvation is also critical for physico-chemical properties (such as pK\textsubscript{a} values of organic compounds\cite{44}), spectroscopic properties (such as NMR chemical shifts\cite{45,46}), and absorption spectra\cite{21,26,47–49}.

A popular attempt to remedy the shortcomings of implicit solvent models is the introduction of explicit solvent molecules to the system\cite{50–55}. The goal of such hybrid cluster-continuum schemes is to model short-ranged interactions explicitly and long-range effects through the continuum model surrounding the cluster (for a recent review see Ref. 56). However, unless one knows the solute’s local solvent environment \textit{a priori}, this approach has multiple pitfalls: First, it is unclear how many solvent molecules need to be added to describe solvent effects to the desired accuracy. Second, the manual process of adding solvent molecules to selected regions around a solute molecule is often guided by ad hoc assumptions. Third, due to the unfavorable scaling of most quantum chemical methods often only one (rarely a few) low-lying solute-solvent configurations are taken into consideration. In the light of the high dimensionality and rugged nature of the potential energy surface (PES) of solvent-solute clusters, it is unlikely to find a representative configuration through manual exploration.

Recent studies attempted to tackle the above-mentioned issues of the static quantum chemical approach. In an extended study of solute-solvent complexes that applied global structure optimization techniques, Li and Hartke\cite{57} explored extreme effects of solvent molecules on chemical reactions (in particular, on barrier heights) that will hardly be seen if one departs from the most likely distribution of solvent molecules around the solute. However, without a measure that informs about the probability of forming such solute-solvent structures, it remains uncertain whether extreme cases are likely to play a role in a reaction in solution. It is therefore mandatory to provide some way of configurational sampling to acquire information on the density of structures in a relevant energy range. In their work on the Morita-Baylis-Hillman reaction, Basdogan and Keith\cite{39} found that a stochastic computational filtering procedure using a global optimization code can help identify low energy structures. They justified the number of solvent
molecules in the solvation model by comparing computational results with experiment. In this way, they favored a model consisting of five methanol molecules over a model with ten. Kildgaard et al. presented a stochastic algorithm for the generation of hydration clusters of sulfuric acid. Their algorithm places water molecules in selected orientations around a solute in an iterative fashion. However, they have not attempted to obtain a distribution over conformations but only the conformation with the lowest free energy was sought. Moreover, the algorithm is not applicable to any choice of solvent but has been tailored for the construction of water clusters.

Considering the shortcomings of current approaches, a method is sought that fulfills the following key requirements:

1. **Computational Feasibility**: a cost-effective model is required that strikes the right balance between accuracy and computational feasibility.

2. **Systematic Improvability**: the accuracy of the model should be adaptable to the computational resources available and, in principle, be systematically improvable so that the physically correct description of solvation in the proper thermodynamic ensemble is recovered.

3. **Universal Applicability**: the model should be applicable to any solute-solvent combination and even solvent mixtures.

4. **Full Automation**: the approach should not require any human intervention to be unbiased and to avoid wasting human time.

In this paper, we present a static cluster-continuum scheme that fulfills these requirements. Regarding the first requirement, we apply efficient generalized-gradient-approximation density functional theory with density fitting for the optimization of solvent-solute clusters, but note that semi-empirical quantum chemical methods can very much reduce the increasing computational cost caused by considering solvent molecules explicitly. As these methods struggle to properly describe important interactions in solution such as dispersion and hydrogen-bonding, corrections have been developed to alleviate such shortcomings. Hence, their application for the
conformational sampling of large clusters will be beneficial. We develop our approach at the example of a prototypical system: acetonitrile in the solvents water and dichloromethane (DCM). The corresponding algorithm has been integrated into our Chemoton structure exploration program that is part of the SCINE software suite.

2 Methodology

2.1 Stochastic Generation of Solvation Clusters

The algorithm presented below ensures that solvent molecules are placed equally around any solute in a stochastic fashion. To ensure that requirements 'Universal Applicability' and 'Full Automation' are fulfilled, the generation of solvation clusters may only depend on nuclear coordinates and quantum chemical observables such as the electron density.

1. First, the accessible surfaces of the solute-solvent complex (which consists of the solute only in the beginning) and solvent are identified. For a given molecular structure, an icosahedral mesh consisting of 12 vertices \( s_a \) (called sites) is created around each atom \( a \) with radius equal to its van der Waals radius \( R_a \). With mesh sub-division algorithms, meshes of arbitrary smoothness can be generated.

2. Sites will be marked as covered if a ray originating from the site going in the direction of the site’s surface normal hits another mesh within a distance of \( d_{\text{cutoff}} \). All sites that are buried by meshes from neighboring atoms are removed. The concept of (open and covered) sites is illustrated in Fig. 1.

3. Two open sites \( s_{a,\text{solute}} \) and \( s_{b,\text{solvent}} \) of the solute-solvent complex and the solvent, respectively, are selected at random.

4. If there were no open sites in the solute, i.e., all sites are covered by the solute itself or by solvent molecules, a complete solvation shell had been formed. If more solvent molecules are to be added, the solute-solvent
complex will be considered the new starting cluster and the algorithm continues with step 1.

5. The solute-solvent complex and the solvent molecule are arranged relative to each other in such a way that site $s_{a,\text{solute}}$, atom $a$, site $s_{b,\text{solvent}}$, and atom $b$ are on one axis. The distance between the sites $s_{a,\text{solute}}$ and $s_{b,\text{solvent}}$ is set to $d$.

6. The angle around this axis is chosen randomly under that constraint that atoms do not come too close to each other, i.e., their van der Waals spheres do not overlap.

7. If no orientation can be found for which atoms do not come too close to one another, $d$ will be increased in increments of $d_{\text{inc}}$ and step 6 will be attempted again until a maximum distance $d_{\text{max}}$ will be reached.

8. If it is not possible to place a solvent molecule, the site will be marked as covered and the algorithm continues with step 3.

9. Once a solvent molecule is added to the solute-solvent complex, the algorithm continues with step 2 until the desired number of solvent molecules has been added.

The random selection of sites in step 3 will not be optimal if the solute or solvent features charged polar functional groups or those containing strong H-bond donors or acceptors. Depending on the polarity and charge of the solute and solvent an equal distribution of solvent molecules around the solute may be inefficient. To take this into account, one could augment sites with descriptors based on first principles such as the Laplacian of the electron density,\textsuperscript{65} Fukui functions,\textsuperscript{66} partial atomic charges,\textsuperscript{67,68} atomic polarizabilities,\textsuperscript{69,70} or dual descriptors\textsuperscript{71,72} (see also Refs. 73 to 80 for reviews). As a result, electron-rich oxygen atoms, for example, would be placed in such a way that they are facing electron-deficient hydrogen atoms. Such an approach could reduce the number of steps required in the structure optimization and improve the probability of generating a low-energy configuration. At the same time, care should be taken so that the use of descriptors does not introduce any bias.
If the solvent has multiple relevant conformations (e.g., hexane) or a mixture of solvents is to be modeled, multiple structures (possibly with corresponding statistical weights) can be included in the site selection process in step 3. Although these extensions are straightforward to implement, they are beyond the scope of the present work.

2.2 Hybrid Solvation Model

The generated solute-solvent complexes are embedded in a cavity formed by a continuum with the dielectric constant of the solvent $\epsilon_{\text{solv}}$.[81] For the construction of the cavity, its solvent accessible surface needs to be determined. This can be achieved by probing the complex with a sphere of radius $R_{\text{solv}}$[81] that is specific to the solvent. This procedure was originally developed for single molecules, not for molecular clusters. In the latter case, unphysical cavities can form within the complex (see Fig. 2). In practice, however, these cavities can be easily detected and a larger $R_{\text{solv}}$ can be chosen to prevent their formation. The small deviation from the idealized radius has a negligible effect on the solute-solvent interaction, especially as the border of the cavity should be far from the solute.
Figure 2: Acetonitrile-water cluster surrounded by a dielectric continuum (light blue spheres). A cross-section (b) reveals an unphysical cavity within the cluster.

The resulting cluster structure is then subjected to quantum chemical structure optimization that will lock it into a local minimum on the Born-Oppenheimer PES. One could, however, also optimize the structures after each addition of a solvent molecule.

2.3 Sampling of Solvation Clusters and Convergence

In the canonical ensemble, the importance of a set of nuclear coordinates $x$ (the configuration) of a molecular system at temperature $T$ may be taken as governed by a Boltzmann probability distribution,

$$p(x) = \frac{1}{z} \exp \left\{ -\frac{E(x)}{k_B T} \right\},$$

with the total energy of the configuration $E(x)$, the molecular contribution $z$ to the canonical partition function, and the Boltzmann constant $k_B$. For the sake of simplicity, we assume that the total energy is governed by the electronic energy of each solute-solvent cluster in a dielectric continuum so that we can rate the different configurations of one cluster size according to this quantity (nuclear zero-point and finite-temperature contributions may be added through the standard translation-in-a-box harmonic-oscillator rigid-rotor approximation for the partition function equipped with a dielectric-
continuum solvation model for assessing the change in free energy associated with embedding a solute-solvent cluster into the liquid phase).

$M$ samples drawn from this distribution then yield a configuration average of an observable $\langle O \rangle$,

$$\langle O \rangle = \frac{1}{M} \sum_{i=1}^{M} O(x_i).$$

(2)

A strategy now needs to be devised which ensures that sufficiently many relevant configurations are generated so that the configurations are assigned the proper weight $p(x_i)$ when calculating expectation values $\langle O \rangle$ for observables $O$,

$$\langle O \rangle = \sum_{i=1}^{M} p(x_i) O(x_i),$$

(3)

with the weights normalized according to Eq. (1) by

$$z \approx \sum_{j=1}^{M} \exp \left\{ -E(x_j)/(k_B T) \right\}.$$  

(4)

This is non-trivial (especially for systems with many floppy degrees of freedom such as solute-solvent clusters). For this reason, the inclusion of solvent molecules is often avoided or only the lowest-total-energy configuration $x_{\text{minimum}}$ is considered,

$$\langle O \rangle \approx O(x_{\text{minimum}}).$$

(5)

Such an approximation of $\langle O \rangle$ by a single sample will only be reasonable if it can be guaranteed that there exists a sufficiently large energy gap between this single configuration and all others higher in energy. In general, this assumption will not be justified for rugged PESs such as those created by explicit solvation.

To ensure that our approach is systematically improvable (requirement 2) yet computationally feasible (requirement 1), we present a procedure that allows one to a) systematically determine a minimum number of solvent molecules required for modeling the solvent effect to sufficient accuracy and b) approximate the distribution over configurations of solute-solvent clusters to obtain reliable ensemble properties at a given temperature.
We now define the (electronic) interaction energy $E_{\text{inter}}$ between the solute and its environment in the following way:

$$E_{\text{inter}} = E_{\text{total}} - E_{\text{solute}} - E_{\text{solvent}},$$  \hspace{1cm} (6)

where $E_{\text{total}}$ is the total electronic energy of the solute-solvent complex, $E_{\text{solute}}$ is the electronic energy of the solute, and $E_{\text{solvent}}$ is the electronic energy of the solvent molecules including the implicit solvent model (see Fig. 3 for a graphical representation of these energy components).

If the total electronic energy $E_{\text{total}}$ of the $i$-th cluster had been used in Eqs. (3) and (4), it would be strongly dominated by configurations in which the solvent molecules (most of which will be far away from the solute) are arranged in a stable configuration. This effect would draw the attention from the solute to the solvent, particularly in the case of large clusters. As a remedy, we employ the following expression for the energy in the Boltzmann weighting in Eq. (1):

$$E_{\text{solute+env}} = E_{\text{solute}} + E_{\text{inter}} = E_{\text{total}} - E_{\text{solvent}},$$  \hspace{1cm} (7)

In this way, also the interaction between the solvent molecules and the continuum model are quenched (this assumes that the solute is not in direct contact with the continuum). With this partitioning scheme, however, free energies cannot be straightforwardly calculated because vibrational modes cannot (easily) be attributed to the solute and its direct surrounding (see also the recent comparison of finite-temperature models for solution in Ref. [82] and references therein). We note that a recent study[83] showed that the inclusion of vibrational contributions in the free energy of solvation can have a negligible effect on the accuracy of thermodynamic cycle predictions of pK_a's and reduction potentials.

The complete algorithm then runs as follows:

1. Generate $N_{\text{sample}}$ solute-solvent complexes by adding $n_{\text{solv}}$ solvent molecules to the solute with the procedure detailed in Section 2.1.

2. Optimize the structures of the complexes.
3. Evaluate $\langle E_{\text{inter}} \rangle$ and $\langle E_{\text{distort}} \rangle$ employing $E_{\text{solute-env}}$ for the weights $w$. $E_{\text{distort}}$ is defined as the energy of the solute (without counterpoise correction) from which the energy of the solute optimized in vacuum is subtracted. With $\langle E_{\text{inter}} \rangle$ and $\langle E_{\text{distort}} \rangle$ we have measures quantifying the solute interactions with the solvent and the distortion of the solute induced by the environment, respectively. If the cluster partition function $z$ is dominated by only a few configurations, more samples must be generated in step 1.

4. If $\langle E_{\text{inter}} \rangle$ and $\langle E_{\text{distort}} \rangle$ are sufficiently close to the values of a set of clusters consisting of fewer solvent molecules, terminate. Else, increase $n_{\text{solv}}$ and go back to step 1.

This algorithm is based on first principles and does not depend on the solute’s shape or charge. As a result, it will, in principle, be applicable to any solute, also to those that are strongly interacting with the solvent. In these cases, more solvent molecules may be required until the convergence criteria are met.

Further, when a larger complex is generated, its structure is not based on a cluster from the previous iteration to ensure that no bias is introduced.
3 Computational Details

The protocol described so far was implemented into our program package Chemoton\textsuperscript{13} and carried out in a fully automated fashion. Icosahedral meshes were generated with the C++ library OpenMesh.\textsuperscript{84} We will make Chemoton including the features described in this work available through our SCINE web page.\textsuperscript{64}

All structure optimizations and single-point calculations were carried out with the \textit{Orca} program package (version 4.0.1)\textsuperscript{85} employing the exchange-correlation density functional PBE,\textsuperscript{86} D3BJ dispersion corrections,\textsuperscript{87,88} the def2-TZVPP\textsuperscript{89,90} basis set, and the corresponding density-fitting resolution-of-the-identity approximation for the Coulomb integrals. For structure optimizations and single-point calculations the default convergence criteria were chosen. We note that for the raw-data generation other quantum chemistry packages can be easily interfaced.

For the conformer generation according to the protocol described above the following parameters were employed: $d_{\text{cutoff}} = 5$ Å, the initial setting of $d = 0$ Å, $d_{\text{inc}} = 0.25$ Å, $d_{\text{max}} = 5$ Å, and $N_{\text{sample}} = 100$. Between each iteration, in the protocol specified in Section 2.3, we increased the number of solvent molecules by 5. This choice was made arbitrarily for this work, but can, in principle, be treated as a solute-size-dependent parameter.

The implicit solvent was described with the conductor-like polarizable continuum solvation model (CPCM).\textsuperscript{31,91–93} The solvent probe radius was increased from 1.3 to 1.9 Å to avoid cavities being formed within the solute-solvent complex. To verify that even for small cluster sizes this parameter change does not significantly distort our results we studied an extreme case. We ran single point calculations for acetonitrile in a continuum (i.e., no explicit solvent molecules) with two solvent probe radii: 1.3 (default) and 1.9 Å. With the computational setup described in the paper the electronic energy does not change significantly ($< 10^{-6}$ a.u.) for both solvents. In addition, visual inspection of the cavities showed that they are practically the same.

When calculating the Boltzmann weight of each solute-solvent complex, the temperature was set to $T = 291.15$ K. For the calculation of $E_{\text{solute}}$ and
\(E_{\text{solvent}}\), a basis set superposition error was avoided by attaching the appropriate ghost-basis functions to unoccupied nuclear positions in the energy evaluation.

The RDFs were calculated with a bin size of 1.8 a.u. A cluster’s average solvent density was estimated by dividing the number of solvent molecules by the volume of the convex hull spanned by all atoms.

All results were saved to a Mongo database. Automated data analysis was performed with the Python libraries matplotlib and pandas. The structures obtained are provided in the Supporting Information.

4 Results

We study our protocol at the example of acetonitrile as solute in the solvents DCM and water. We chose these components because neither acetonitrile nor either solvent contains conformational degrees of freedom to be sampled in addition to the configurational ones. Moreover, acetonitrile is soluble but chemically inert in both solvents and contains a polar, hydrogen-bond accepting nitrile group as well as a non-polar methyl group. Our protocol will terminate if the absolute differences with respect to \(\langle E_{\text{inter}} \rangle\) and \(\langle E_{\text{distort}} \rangle\) between the current cluster and the previous one are both below 0.5 kcal/mol (or \(\approx 2.1\) kJ/mol). This threshold value may be adjusted as it depends on the system’s size and the required accuracy.

4.1 Acetonitrile-dichloromethane clusters

In Table 1, the number of successfully optimized acetonitrile-DCM clusters \(N_{\text{clusters}}\) (out of \(N_{\text{sample}} = 100\) generated starting structures) and the number of clusters with significant contributions to the partition function \(n_{\text{sig}} (p(x) > 0.05\), where \(M = N_{\text{clusters}}\)) are given for different cluster sizes. A structure optimization of a cluster is considered successful if the converge criteria are met within 1200 structure optimization steps.

We note that the algorithm from Section 2.3 would have terminated after 15 solvent molecules. First, it can be seen that almost all generated clusters could be optimized successfully. This demonstrates that our algorithm
Table 1: Number of successfully optimized clusters $N_{\text{clusters}}$ and number of significant clusters $n_{\text{sig}} (p(x) > 0.05)$ for clusters containing $n_{\text{solv}}$ DCM molecules. $T$ is the average wall time (in hours, on a 16-core CPU) and $n_{\text{cycles}}$ is the average number of required optimization cycles.

| $n_{\text{solv}}$ | $N_{\text{clusters}}$ | $n_{\text{sig}}$ | $T$ | $n_{\text{cycles}}$ |
|-------------------|------------------------|-------------------|-----|---------------------|
| 5                 | 98                     | 5                 | 2   | 225                 |
| 10                | 99                     | 4                 | 8   | 264                 |
| 15                | 95                     | 6                 | 18  | 254                 |
| 20                | 93                     | 4                 | 35  | 248                 |
| 25                | 83                     | 5                 | 45  | 179                 |

produces starting structures that reproduce proper bonding patterns so that subsequent structure optimizations do not yield dissociating structures such as hydronium ions.

Second, Table 1 shows that the number of structure optimizations that failed to converge within 1200 steps increases with the number of solvent molecules. The slow convergence of these structure optimizations can be attributed to the system size and its many soft degrees of freedom. Further, the ratio between the number of significant clusters $n_{\text{sig}}$ and the total number of generated clusters $N_{\text{clusters}}$ is relatively low ($\approx 2 - 5\%$) for all clusters sizes. As described in Section 2.3, the weight of each cluster is calculated from $E_{\text{solute}+\text{env}}$. Since the variance in $E_{\text{solute}}$ is small (see below), it is the interaction energy $E_{\text{inter}}$ that is responsible for the large spread. Therefore, there must be a few configurations that have significantly higher $E_{\text{inter}}$ than others.

In Fig. 1, $E_{\text{inter}}$ (left) and the dispersive component of $E_{\text{inter}}$ (right) of acetonitrile-DCM complexes (white discs) are plotted as a function of the number of solvent molecules. First, it can be seen that $E_{\text{inter}}$ drops from $-26$ to $-90$ kJ/mol with increasing cluster size until clusters consist of ten DCM molecules. After that point, $E_{\text{inter}}$ remains constant (within error bars). A similar trend can be observed for $E_{\text{inter, disp}}$. Visual inspection of the complexes shows that with ten DCM molecules a complete solvation shell can be formed (see the most stable configuration in Fig. 5). This may explain why the interaction energy does not change after that point: once a full solvation shell is formed, hardly any further pronounced interactions between
the solute and the additional solvent molecules emerge for this solute-solvent combination.

Further, Fig. 4 shows that \( E_{\text{inter}} \) and \( E_{\text{inter, disp}} \) of the most stable complexes are located close to the clusters with the strongest solute-solvent interaction (lowest \( E_{\text{inter}} \) and \( E_{\text{inter, disp}} \)). From this finding, we conclude that the stability of the clusters can be attributed to the interaction between solute and solvent, rather than from interactions between solvent molecules.

To study the effect of the environment, we also track the length of the C-N bond in acetonitrile, \( d_{\text{C-N}} \). In Fig. 6, \( E_{\text{distort}} \) and \( d_{\text{C-N}} \) are shown as a function of the number of DCM molecules in the solute-solvent complex. It can be seen, that \( E_{\text{distort}} \), as well as changes in \( d_{\text{C-N}} \), are relatively small for all cluster sizes. In addition, the spread of \( E_{\text{distort}} \) and \( d_{\text{C-N}} \) among clusters of the same size is small (\( \approx 0.4 \) kJ/mol and \( \approx 0.005 \) bohr, respectively). Therefore, in this system, the main contributor to deviations in \( E_{\text{solute+env}} \) among clusters is the interaction energy \( E_{\text{inter}} \), not the stability of the solute. For large molecules that can undergo large conformational changes in solution, this

\[ E_{\text{inter}} (\text{kJ/mol}) \]

\[ E_{\text{inter, disp}} (\text{kJ/mol}) \]

Figure 4: \( E_{\text{inter}} \) (left) and its dispersive component \( E_{\text{inter, disp}} \) (right) of acetonitrile-DCM complexes (white discs) as a function of the number of solvent molecules in kJ/mol. Black discs connected by black solid lines show \( \langle E_{\text{inter}} \rangle \) and \( \langle E_{\text{inter, disp}} \rangle \) (with error bars indicating \( \pm 2\sigma \)). Red discs connected by red dashed lines highlight \( E_{\text{inter}} \) and \( E_{\text{inter, disp}} \) of the most stable complexes.
Figure 5: Ten DCM molecules form a complete solvation shell around one acetoni-trile molecule.

Figure 6: $E_{\text{distort}}$ (left, in kJ/mol) and $d_{C-N}$ (right, in atomic units (a.u.), i.e. bohr in this case) of acetonitrile-DCM complexes (white discs) as a function of the number of solvent molecules. Black discs connected by black solid lines show $\langle E_{\text{distort}} \rangle$ and $\langle d_{C-N} \rangle$ (with error bars indicating $\pm 2\sigma$). Red discs connected by red dashed lines show $E_{\text{distort}}$ and $d_{C-N}$ of the most stable complexes.

will not necessarily be the case. Furthermore, it can be seen that $\langle E_{\text{distort}} \rangle$
converges (within error bars) whereas $E_{\text{distort}}$ of the most stable configuration does not. This showcases the issue of employing the total energy as a measure for computing Boltzmann averages: with an increasing number of explicit solvent molecules, the stability of the environment will dominate over that of the solute.

### 4.2 Acetonitrile-water clusters

As a difficult case for microsolvation, we now turn to water as the medium for acetonitrile. In Table 2, the number of successfully optimized acetonitrile-water clusters $n_{\text{clusters}}$ and the number of clusters with significant contributions to the partition function $n_{\text{sig}} (p(x) > 0.05$, where $M = N_{\text{clusters}})$ are given for different cluster sizes. We note that the algorithm from Section 2.3 would have terminated after 35 solvent molecules. Similar trends as in Table 1 can be identified, however, $n_{\text{sig}}$ is generally lower in water than in DCM. This can be explained by the strong hydrogen-bond network formed around the nitrile group which consists of a specific arrangement of water molecules (see Fig. 7). Out of the optimized clusters, only a small fraction features this particular arrangement.

**Table 2:** Number of successfully optimized clusters $n_{\text{clusters}}$ and number of significant clusters $n_{\text{sig}} (p(x) > 0.05)$ for clusters containing $n_{\text{solv}}$ water molecules. $T$ is the average wall time (in hours, on a 16-core CPU) and $n_{\text{cycles}}$ is the average number of required optimization cycles.

| $n_{\text{solv}}$ | $n_{\text{clusters}}$ | $n_{\text{sig}}$ | $T$ | $n_{\text{cycles}}$ |
|------------------|-----------------------|------------------|-----|---------------------|
| 5                | 97                    | 7                | 1   | 230                 |
| 10               | 98                    | 3                | 3   | 302                 |
| 15               | 98                    | 1                | 4   | 296                 |
| 20               | 99                    | 1                | 7   | 319                 |
| 25               | 99                    | 3                | 12  | 328                 |
| 30               | 98                    | 3                | 16  | 309                 |
| 35               | 96                    | 2                | 24  | 324                 |

In Fig. 8, $E_{\text{inter}}$ and the dispersive component of $E_{\text{inter}}$ (right) of acetonitrile-water complexes (white discs) are plotted as a function of the number of solvent molecules. First, it can be seen that $E_{\text{inter}}$ drops from $-30$ to
Figure 7: Formation of four hydrogen bonds (indicated by yellow dashed lines) between water and the nitrile group of acetonitrile. Among the clusters containing 15 water molecules, this structure has the highest (absolute value of the) interaction energy $E_{\text{inter}}$. 

$-140$ kJ/mol with increasing cluster size until the complex consists of 30 molecules; after that point, $E_{\text{inter}}$ remains constant (within error bars). A similar trend can be observed for $E_{\text{inter, disp}}$. As expected, more water than DCM molecules are required to reach the point of convergence: visual inspection of the complexes consisting of 30 water molecules shows that one to two solvation shells are formed (see the most stable configuration in Fig. 9). The smaller water molecule interacts more strongly with the solute than DCM ($\langle E_{\text{inter}} \rangle$ of $-140$ compared to $-90$ kJ/mol).

An outlier for the cluster size of 15 can be identified in Fig. 8. This particular cluster (shown in Fig. 7) is the only one of that size that features four hydrogen bonds. As a result, the absolute value of the interaction energy is largest and so is the cluster’s weight in the cluster partition function $z$. However, from Fig. 7 it can also be seen that a complete solvation shell has not been formed yet. In Fig. 8 both $E_{\text{inter}}$ and $E_{\text{inter, disp}}$ reflect this fact.

In addition, it can be seen that the spread of $E_{\text{inter}}$ is large ($\approx 90$ kJ/mol for clusters consisting of 30 solvent molecules). Considering that all of these structures are minimum energy structures, this finding stresses the impor-
Figure 8: $E_{\text{inter}}$ (left) and its dispersive component $E_{\text{inter}, \text{disp}}$ (right) of acetonitrile-water complexes (white discs) as a function of the number of solvent molecules in kJ/mol. Black discs connected by black solid lines show $\langle E_{\text{inter}} \rangle$ and $\langle E_{\text{inter}, \text{disp}} \rangle$ (with error bars indicating $\pm 2\sigma$). Red discs connected by red dashed lines highlight $E_{\text{inter}}$ and $E_{\text{inter}, \text{disp}}$ of the most stable complexes.

Figure 9: Acetonitrile molecule surrounded by 30 water molecules.

tance of rigorous sampling: an inappropriate selection of solvation clusters (from a manual exploration, for instance) can be detrimental to the signifi-
cance of a theoretical study.

The spread of energies is lower for $E_{\text{inter, disp}}$ than for $E_{\text{inter}}$. Not only is the dispersive component smaller, but dispersive interactions are also not directional, and hence, not dependent on the exact arrangement of the molecules. This is in contrast with non-dispersive interactions such as hydrogen bonds that play a central role in this case.

Figure 10: Most stable acetonitrile-water cluster consisting of 20 water molecules. No hydrogen bonds are formed between the nitrile group and the solvent and the cluster stabilization is solely brought about by the strong network of solvent-solvent hydrogen bonds.

Finally, it can be seen that $\langle E_{\text{inter}} \rangle$ and $\langle E_{\text{inter, disp}} \rangle$ do not coincide with $E_{\text{inter}}$ and $E_{\text{inter, disp}}$ of the most stable complexes, respectively. This difference is more pronounced in water than in DCM (compare Fig. 4) because in the former the interactions between the solvent molecules are stronger. In Fig. 10, the most stable acetonitrile-water complex (consisting of 20 solvent molecules) is shown. This structure is represented by the red point in Fig. 8, left, for 20 solvent molecules. In this structure, no hydrogen bond with the acetonitrile group is formed but the water molecules are arranged such that the hydrogen-bond network of the solvent molecules is optimal. However, such a configuration is solely a result of the limited number of solvent molecules in the solvent shell and would be dissolved in larger clus-
ters, where further water molecules break up the hydrogen network of the 20 water molecules and allow then for hydrogen bonding with the solute. However, this most stable configuration with 20 water molecules would have the highest weight if $E_{\text{total}}$ had been employed in the Boltzmann weighting. This emphasizes again how decisive it is to choose a suitable energy measure for calculating the weights $p(x)$.

![Figure 11: $E_{\text{distort}}$ (left, in kJ/mol) and $d_{\text{C-N}}$ (right, in atomic units (a.u.), i.e. bohr) of acetonitrile-water complexes (white discs) as a function of the number of solvent molecules. Black discs connected by black solid lines show $\langle E_{\text{distort}} \rangle$ and $\langle d_{\text{C-N}} \rangle$ (with error bars indicating ±2σ). Red discs connected by red dashed lines show $E_{\text{distort}}$ and $d_{\text{C-N}}$ of the most stable complexes.](image)

In Fig. 11, $E_{\text{distort}}$ and $d_{\text{C-N}}$ are shown as a function of the number of water molecules in the solute-solvent complex. Compared to Fig. 6, it can be seen that $E_{\text{distort}}$, as well as changes in $d_{\text{C-N}}$, are larger for all cluster sizes. The outlier in $\langle E_{\text{inter}} \rangle$ for clusters with 15 water molecules represents the complex shown in Fig. 7. The strong interaction also causes the strong nitrile bond to be elongated. The presence of additional solvent molecules then reduces the strong solute-solvent interaction. This is why a similar outlier cannot be found for clusters with more than 15 water molecules.

In Fig. 12, the radial distribution function (RDF) $g(r)$ between the nitrogen of acetonitrile and the oxygen of water for two cluster sizes are shown.
together with classical molecular dynamics reference data taken from Ref. 97. For the clusters, the RDF was approximated by taking the weighted average of the RDFs of clusters of the same size. The cluster sizes 30 and 35 were chosen because clusters of this size describe the solute-solvent interaction sufficiently well. It can be seen that clusters’ RDFs compare very favorably with the reference data below $r \approx 8$ a.u. The RDF is close to zero below 5 a.u. to reach its maximum at $\approx 6.5$ a.u. After that the cluster RDFs drop rapidly as they should because the density of explicit solvent molecules vanishes and replaced by implicit solvation of the continuum model in our cluster-continuum approach. It should be noted that for the calculation of the RDF a relatively large bin size of 1.8 a.u. was chosen to ensure that each bin contains sufficiently many solvent molecules. Further, larger clusters and a larger sample size would be required for a quantitative comparison with simulation data, especially for the region of the RDF beyond 8 bohr.

**Figure 12:** Radial distribution function $g(r)$ of the nitrogen-oxygen distance for cluster sizes 30 and 35 (circles) together with classical molecular dynamics reference data (continuous curve) adapted from Ref. 97. Note the matching position of the maxima.
5 Conclusions

In this study, we presented a systematic approach for modeling solvation effects by establishing a fully automated static hybrid cluster-continuum scheme with conformational sampling. Our general algorithm for the stochastic generation of solute-solvent clusters is applicable to any solute or solvent. We established criteria that not only indicate how many configurations need to be generated but also how many explicit solvent molecules need to be added to capture most of the interaction between the solute and its environment. For this, we applied a scheme of weighting sampled configurations to obtain meaningful configuration statistics. We then proposed measures that indicate how many explicitly treated solvent molecules are necessary to accurately describe the solute’s environment. As the required size of the clusters depends strongly on the particular solvent and solute, this needs to be determined in a case-by-case manner (and in an automated fashion) when our hybrid cluster-continuum scheme is employed.

We note that the conformational sampling of the solute molecule can be conveniently separated from that of the solvent shell through, for example, our structure exploration program Chemoton, which first generates solute conformations that can then be solvated. For the sake of efficiency, it is also possible to first solvate one solute conformation and then change this conformation in the solvent cages generated to obtain guess structures to be subjected to structure optimization.

The application of our scheme to model acetonitrile in DCM and water highlighted the shortcomings of standard quantum chemical microsolvation attempts. The results show, as one would have expected, that in solvents with strong intermolecular interactions (e.g., water) a single most-stable cluster is unlikely to be representative. It is not appropriate then to identify only the most stable cluster for calculating ensemble statistics. A fully automated approach for systematically generating and sampling solute-solvent clusters in a static quantum chemical picture will be mandatory. The PES of such clusters is too rugged for a rigorous in-depth manual exploration. This also points toward systematic sampling of configuration space through Monte-Carlo or molecular dynamics algorithms, to which our automated cluster
generation protocol may be seamlessly coupled to eventually allow sampling even under periodic boundary conditions.

As such extensions will require significant computational resources, a combination with approximate interaction models ranging from semi-empirical methods to classical molecular-mechanics force fields and machine learning models is a natural extension (for the fast-growing literature on these schemes, we may refer to references in Refs. [98][101]). However, we emphasize that reliability of such methods which trade accuracy for computational efficiency is best guaranteed if suitable uncertainty quantification schemes (such as those reported by us in Refs. 102–105) are in operation that inform about the range of applicability. Note also that such an approach should then not be considered as some general transferable model, but as a system-focused baseline model whose suitability is quantitatively assessed by uncertainty quantification procedures in a rolling fashion through continuous benchmarking as discussed in Refs. 102 and 105.

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References

[1] Kirchner, B. Theory of Complicated Liquids: Investigation of Liquids, Solvents and Solvent Effects with Modern Theoretical Methods. Phys. Rep. 2007, 440, 1–111.

[2] Reichardt, C.; Welton, T. Solvents and Solvent Effects in Organic Chemistry, 4th ed.; Wiley-VCH: Weinheim, 2010.

[3] Kropman, M. F.; Bakker, H. J. Dynamics of Water Molecules in Aqueous Solvation Shells. Science 2001, 291, 2118–2120.

[4] Laage, D.; Hynes, J. T. Reorientional Dynamics of Water Molecules
in Anionic Hydration Shells. *Proc. Natl. Acad. Sci. U.S.A.* 2007, 104, 11167–11172.

[5] Park, S.; Fayer, M. D. Hydrogen Bond Dynamics in Aqueous NaBr Solutions. *Proc. Natl. Acad. Sci. U.S.A.* 2007, 104, 16731–16738.

[6] Sameera, W. M. C.; Maeda, S.; Morokuma, K. Computational Catalysis Using the Artificial Force Induced Reaction Method. *Acc. Chem. Res.* 2016, 49, 763–773.

[7] Dewyer, A. L.; Argüelles, A. J.; Zimmerman, P. M. Methods for Exploring Reaction Space in Molecular Systems. *WIREs Comput. Mol. Sci.* 2018, 8, e1354.

[8] Simm, G. N.; Vaucher, A. C.; Reiher, M. Exploration of Reaction Pathways and Chemical Transformation Networks. *J. Phys. Chem. A* 2019, 123, 385–399.

[9] Marx, D.; Hutter, J. *Ab Initio Molecular Dynamics: Basic Theory and Advanced Methods*; Cambridge University Press, 2009.

[10] Maeda, S.; Ohno, K.; Morokuma, K. Systematic Exploration of the Mechanism of Chemical Reactions: The Global Reaction Route Mapping (GRRM) Strategy Using the ADDF and AFIR Methods. *Phys. Chem. Chem. Phys.* 2013, 15, 3683–3701.

[11] Zimmerman, P. M. Navigating Molecular Space for Reaction Mechanisms: An Efficient, Automated Procedure. *Mol. Simul.* 2015, 41, 43–54.

[12] Bergeler, M.; Simm, G. N.; Proppe, J.; Reiher, M. Heuristics-Guided Exploration of Reaction Mechanisms. *J. Chem. Theory Comput.* 2015, 11, 5712–5722.

[13] Simm, G. N.; Reiher, M. Context-Driven Exploration of Complex Chemical Reaction Networks. *J. Chem. Theory Comput.* 2017, 13, 6108–6119.
[14] van Duin, A. C. T.; Dasgupta, S.; Lorant, F.; Goddard, W. A. ReaxFF: A Reactive Force Field for Hydrocarbons. *J. Phys. Chem. A* 2001, 105, 9396–9409.

[15] Döntgen, M.; Przybylski-Freund, M.-D.; Kröger, L. C.; Kopp, W. A.; Ismail, A. E.; Leonhard, K. Automated Discovery of Reaction Pathways, Rate Constants, and Transition States Using Reactive Molecular Dynamics Simulations. *J. Chem. Theory Comput.* 2015, 11, 2517–2524.

[16] da Silva, G. C. Q.; Cardozo, T. M.; Amarante, G. W.; Abreu, C. R. A.; Horta, B. A. C. Solvent Effects on the Decarboxylation of Trichloroacetic Acid: Insights from Ab Initio Molecular Dynamics Simulations. *Phys. Chem. Chem. Phys.* 2018, 20, 21988–21998.

[17] Galván, I. F.; Martín, M. E.; Aguilar, M. A. A New Method to Locate Saddle Points for Reactions in Solution by Using the Free-Energy Gradient Method and the Mean Field Approximation. *J. Comput. Chem.* 2004, 25, 1227–1233.

[18] Senn, H. M.; Thiel, W. QM/MM Methods for Biological Systems. *Top. Curr. Chem.* 2007, 268, 173–290.

[19] Senn, H. M.; Thiel, W. QM/MM Studies of Enzymes. *Curr. Opin. Chem. Biol.* 2007, 11, 182–187.

[20] Senn, H. M.; Thiel, W. QM/MM Methods for Biomolecular Systems. *Angew. Chem. Int. Ed.* 2009, 48, 1198–1229.

[21] Neugebauer, J.; Jacob, C. R.; Wesolowski, T. A.; Baerends, E. J. An Explicit Quantum Chemical Method for Modeling Large Solvation Shells Applied to Aminocoumarin C151. *J. Phys. Chem. A* 2005, 109, 7805–7814.

[22] Murugan, N. A.; Jha, P. C.; Rinkevicius, Z.; Ruud, K.; Ågren, H. Solvatochromic Shift of Phenol Blue in Water from a Combined
Car–Parrinello Molecular Dynamics Hybrid Quantum Mechanics-Molecular Mechanics and ZINDO Approach. *J. Chem. Phys.* **2010**, *132*, 234508.

[23] Flaig, D.; Beer, M.; Ochsenfeld, C. Convergence of Electronic Structure with the Size of the QM Region: Example of QM/MM NMR Shieldings. *J. Chem. Theory Comput.* **2012**, *8*, 2260–2271.

[24] Zuehlsdorff, T. J.; Haynes, P. D.; Hanke, F.; Payne, M. C.; Hine, N. D. M. Solvent Effects on Electronic Excitations of an Organic Chromophore. *J. Chem. Theory Comput.* **2016**, *12*, 1853–1861.

[25] Provorse, M. R.; Peev, T.; Xiong, C.; Isborn, C. M. Convergence of Excitation Energies in Mixed Quantum and Classical Solvent: Comparison of Continuum and Point Charge Models. *J. Phys. Chem. B* **2016**, *120*, 12148–12159.

[26] Milanese, J. M.; Provorse, M. R.; Alameda, E.; Isborn, C. M. Convergence of Computed Aqueous Absorption Spectra with Explicit Quantum Mechanical Solvent. *J. Chem. Theory Comput.* **2017**, *13*, 2159–2171.

[27] Boereboom, J. M.; Fleurat-Lessard, P.; Bulo, R. E. Explicit Solvation Matters: Performance of QM/MM Solvation Models in Nucleophilic Addition. *J. Chem. Theory Comput.* **2018**, *14*, 1841–1852.

[28] Miertuš, S.; Scrocco, E.; Tomasi, J. Electrostatic Interaction of a Solute with a Continuum. A Direct Utilization of AB Initio Molecular Potentials for the Prevision of Solvent Effects. *Chem. Phys.* **1981**, *55*, 117–129.

[29] Tomasi, J. Thirty Years of Continuum Solvation Chemistry: A Review, and Prospects for the near Future. *Theor. Chem. Acc.* **2004**, *112*, 184–203.

[30] Tomasi, J.; Mennucci, B.; Cammi, R. Quantum Mechanical Continuum Solvation Models. *Chem. Rev.* **2005**, *105*, 2999–3094.
[31] Klamt, A.; Schüürmann, G. COSMO: A New Approach to Dielectric Screening in Solvents with Explicit Expressions for the Screening Energy and Its Gradient. J. Chem. Soc., Perkin Trans. 2 1993, 799–805.

[32] Tomasi, J.; Mennucci, B.; Cancès, E. The IEF Version of the PCM Solvation Method: An Overview of a New Method Addressed to Study Molecular Solutes at the QM Ab Initio Level. J. Mol. Struct. THEOCHEM 1999, 464, 211–226.

[33] Klamt, A. Conductor-like Screening Model for Real Solvents: A New Approach to the Quantitative Calculation of Solvation Phenomena. J. Phys. Chem. 1995, 99, 2224–2235.

[34] Kovalenko, A.; Hirata, F. Potentials of Mean Force of Simple Ions in Ambient Aqueous Solution. I. Three-Dimensional Reference Interaction Site Model Approach. J. Chem. Phys. 2000, 112, 10391–10402.

[35] Ben-Naim, A. Statistical Thermodynamics for Chemists and Biochemists; Plenum: New York, 1992.

[36] Kim, Y.; Mohrig, J. R.; Truhlar, D. G. Free-Energy Surfaces for Liquid-Phase Reactions and Their Use To Study the Border Between Concerted and Nonconcerted α,β-Elimination Reactions of Esters and Thioesters. J. Am. Chem. Soc. 2010, 132, 11071–11082.

[37] Ribeiro, R. F.; Marenich, A. V.; Cramer, C. J.; Truhlar, D. G. Use of Solution-Phase Vibrational Frequencies in Continuum Models for the Free Energy of Solvation. J. Phys. Chem. B 2011, 115, 14556–14562.

[38] Plata, R. E.; Singleton, D. A. A Case Study of the Mechanism of Alcohol-Mediated Morita Baylis–Hillman Reactions. The Importance of Experimental Observations. J. Am. Chem. Soc. 2015, 137, 3811–3826.

[39] Basdogan, Y.; A. Keith, J. A Paramedic Treatment for Modeling Explicitly Solvated Chemical Reaction Mechanisms. Chem. Sci. 2018, 9, 5341–5346.
[40] Park, J. M.; Laio, A.; Iannuzzi, M.; Parrinello, M. Dissociation Mechanism of Acetic Acid in Water. *J. Am. Chem. Soc.* **2006**, *128*, 11318–11319.

[41] Michel, C.; Anneau, F.; Delbecq, F.; Sautet, P. C–H versus O–H Bond Dissociation for Alcohols on a Rh(111) Surface: A Strong Assistance from Hydrogen Bonded Neighbors. *ACS Catal.* **2011**, *1*, 1430–1440.

[42] Glaves, R.; Mathias, G.; Marx, D. Mechanistic Insights into the Hydrolysis of a Nucleoside Triphosphate Model in Neutral and Acidic Solution. *J. Am. Chem. Soc.* **2012**, *134*, 6995–7000.

[43] De Wispelaere, K.; Wondergem, C. S.; Ensing, B.; Hemelsoet, K.; Meijer, E. J.; Weckhuysen, B. M.; Van Speybroeck, V.; Ruiz-Martínez, J. Insight into the Effect of Water on the Methanol-to-Olefins Conversion in H-SAPO-34 from Molecular Simulations and in Situ Microspectroscopy. *ACS Catal.* **2016**, *6*, 1991–2002.

[44] Thapa, B.; Schlegel, H. B. Improved pKa Prediction of Substituted Alcohols, Phenols, and Hydroperoxides in Aqueous Medium Using Density Functional Theory and a Cluster-Continuum Solvation Model. *J. Phys. Chem. A* **2017**, *121*, 4698–4706.

[45] Roggatz, C. C.; Lorch, M.; Benoit, D. M. Influence of Solvent Representation on Nuclear Shielding Calculations of Protonation States of Small Biological Molecules. *J. Chem. Theory Comput.* **2018**.

[46] Caputo, M. C.; Provasi, P. F.; Sauer, S. P. A. The Role of Explicit Solvent Molecules in the Calculation of NMR Chemical Shifts of Glycine in Water. *Theor. Chem. Acc.* **2018**, *137*, 88.

[47] Jacob, C. R.; Neugebauer, J.; Jensen, L.; Visscher, L. Comparison of Frozen-Density Embedding and Discrete Reaction Field Solvent Models for Molecular Properties. *Phys. Chem. Chem. Phys.* **2006**, *8*, 2349–2359.

[48] Georg, H. C.; Coutinho, K.; Canuto, S. Solvent Effects on the UV-Visible Absorption Spectrum of Benzophenone in Water: A Combined
Monte Carlo Quantum Mechanics Study Including Solute Polarization. *J. Chem. Phys.* 2007, 126, 034507.

[49] Neugebauer, J. Chromophore-Specific Theoretical Spectroscopy: From Subsystem Density Functional Theory to Mode-Specific Vibrational Spectroscopy. *Phys. Rep.* 2010, 489, 1–87.

[50] Pliego, J. R.; Riveros, J. M. The Cluster-Continuum Model for the Calculation of the Solvation Free Energy of Ionic Species. *J. Phys. Chem. A* 2001, 105, 7241–7247.

[51] Kelly, C. P.; Cramer, C. J.; Truhlar, D. G. Adding Explicit Solvent Molecules to Continuum Solvent Calculations for the Calculation of Aqueous Acid Dissociation Constants. *J. Phys. Chem. A* 2006, 110, 2493–2499.

[52] Coutinho, K.; Georg, H. C.; Fonseca, T. L.; Ludwig, V.; Canuto, S. An Efficient Statistically Converged Average Configuration for Solvent Effects. *Chem. Phys. Lett.* 2007, 437, 148–152.

[53] Bryantsev, V. S.; Diallo, M. S.; Goddard III, W. A. Calculation of Solvation Free Energies of Charged Solutes Using Mixed Cluster/Continuum Models. *J. Phys. Chem. B* 2008, 112, 9709–9719.

[54] Kua, J.; Avila, J. E.; Lee, C. G.; Smith, W. D. Mapping the Kinetic and Thermodynamic Landscape of Formaldehyde Oligomerization under Neutral Conditions. *J. Phys. Chem. A* 2013, 117, 12658–12667.

[55] Kua, J.; Galloway, M. M.; Millage, K. D.; Avila, J. E.; De Haan, D. O. Glycolaldehyde Monomer and Oligomer Equilibria in Aqueous Solution: Comparing Computational Chemistry and NMR Data. *J. Phys. Chem. A* 2013, 117, 2997–3008.

[56] Pliego, J. R.; Riveros, J. M. Hybrid Discrete-Continuum Solvation Methods. *WIREs Comput. Mol. Sci.* 2019, 0, e1440.

[57] Li, Y.; Hartke, B. Assessing Solvation Effects on Chemical Reactions with Globally Optimized Solvent Clusters. *ChemPhysChem* 2013, 14, 2678–2686.

31
[58] Kildgaard, J. V.; Mikkelsen, K. V.; Bilde, M.; Elm, J. Hydration of Atmospheric Molecular Clusters: A New Method for Systematic Configurational Sampling. *J. Phys. Chem. A* 2018, 122, 5026–5036.

[59] Kildgaard, J. V.; Mikkelsen, K. V.; Bilde, M.; Elm, J. Hydration of Atmospheric Molecular Clusters II: Organic Acid–Water Clusters. *J. Phys. Chem. A* 2018, 122, 8549–8556.

[60] Husch, T.; Vaucher, A. C.; Reiher, M. Semiempirical Molecular Orbital Models Based on the Neglect of Diatomic Differential Overlap Approximation. *Int. J. Quantum Chem.* 2018, 118, e25799.

[61] Korth, M.; Pitoňák, M.; Řezáč, J.; Hobza, P. A Transferable H-Bonding Correction for Semiempirical Quantum-Chemical Methods. *J. Chem. Theory Comput.* 2010, 6, 344–352.

[62] Korth, M. Third-Generation Hydrogen-Bonding Corrections for Semiempirical QM Methods and Force Fields. *J. Chem. Theory Comput.* 2010, 6, 3808–3816.

[63] Korth, M.; Thiel, W. Benchmarking Semiempirical Methods for Thermochemistry, Kinetics, and Noncovalent Interactions: OMx Methods Are Almost As Accurate and Robust As DFT-GGA Methods for Organic Molecules. *J. Chem. Theory Comput.* 2011, 7, 2929–2936.

[64] The Reiher research group, SCINE – Software for Chemical Interaction Networks. [http://scine.ethz.ch](http://scine.ethz.ch) (Accessed: 12. September 2018).

[65] Bader, R. F. W. *Atoms in Molecules: A Quantum Theory*; Clarendon Press: Oxford, 1994.

[66] Fukui, K. Role of Frontier Orbitals in Chemical Reactions. *Science* 1982, 218, 747–754.

[67] Mulliken, R. S. Electronic Population Analysis on LCAO–MO Molecular Wave Functions. I. *J. Chem. Phys.* 1955, 23, 1833–1840.
[68] Mulliken, R. S. Electronic Population Analysis on LCAO–MO Molecular Wave Functions. II. Overlap Populations, Bond Orders, and Covalent Bond Energies. *J. Chem. Phys.* **1955**, *23*, 1841–1846.

[69] Meister, J.; Schwarz, W. H. E. Principal Components of Ionicity. *J. Phys. Chem.* **1994**, *98*, 8245–8252.

[70] Bultinck, P.; Van Alsenoy, C.; Ayers, P. W.; Carbó-Dorca, R. Critical Analysis and Extension of the Hirshfeld Atoms in Molecules. *J. Chem. Phys.* **2007**, *126*, 144111.

[71] Brown, R. E.; Simas, A. M. On the Applicability of CNDO Indices for the Prediction of Chemical Reactivity. *Theoret. Chim. Acta* **1982**, *62*, 1–16.

[72] Kang, Y. K.; Jhon, M. S. Additivity of Atomic Static Polarizabilities and Dispersion Coefficients. *Theoret. Chim. Acta* **1982**, *61*, 41–48.

[73] Kunz, C. F.; Hättig, C.; Hess, B. A. Ab Initio Study of the Individual Interaction Energy Components in the Ground State of the Mercury Dimer. *Mol. Phys.* **1996**, *89*, 139–156.

[74] Morell, C.; Grand, A.; Toro-Labbé, A. New Dual Descriptor for Chemical Reactivity. *J. Phys. Chem. A* **2005**, *109*, 205–212.

[75] Morell, C.; Grand, A.; Toro-Labbé, A. Theoretical Support for Using the $\Delta f(r)$ Descriptor. *Chem. Phys. Lett.* **2006**, *425*, 342–346.

[76] Ayers, P. W.; Morell, C.; De Proft, F.; Geerlings, P. Understanding the Woodward–Hoffmann Rules by Using Changes in Electron Density. *Chem. Eur. J.* **2007**, *13*, 8240–8247.

[77] Cárdenas, C.; Rabi, N.; Ayers, P. W.; Morell, C.; Jaramillo, P.; Fuentealba, P. Chemical Reactivity Descriptors for Ambiphilic Reagents: Dual Descriptor, Local Hypersoftness, and Electrostatic Potential. *J. Phys. Chem. A* **2009**, *113*, 8666–8667.

[78] Geerlings, P.; De Proft, F.; Langenaeker, W. Conceptual Density Functional Theory. *Chem. Rev.* **2003**, *103*, 1793–1874.
[79] Geerlings, P.; Proft, F. D. Conceptual DFT: The Chemical Relevance of Higher Response Functions. *Phys. Chem. Chem. Phys.* **2008**, *10*, 3028–3042.

[80] Johnson, P. A.; Bartolotti, L. J.; Ayers, P. W.; Fievez, T.; Geerlings, P. *Modern Charge-Density Analysis*; Springer, 2011; pp 715–764.

[81] Pye, C. C.; Ziegler, T. An Implementation of the Conductor-like Screening Model of Solvation within the Amsterdam Density Functional Package. *Theor. Chem. Acc.* **1999**, *101*, 396–408.

[82] Besora, M.; Vidossich, P.; Lledós, A.; Ujaque, G.; Maseras, F. Calculation of Reaction Free Energies in Solution: A Comparison of Current Approaches. *J. Phys. Chem. A* **2018**, *122*, 1392–1399.

[83] Ho, J. Are Thermodynamic Cycles Necessary for Continuum Solvent Calculation of pK\textsubscript{a}s and Reduction Potentials? *Phys. Chem. Chem. Phys.* **2015**, *17*, 2859–2868.

[84] Botsch, M.; Steinberg, S.; Bischoff, S.; Kobbelt, L. OpenMesh 7.1 – A Generic and Efficient Polygon Mesh Data Structure. https://www.openmesh.org, (Accessed: 10. April 2019).

[85] Neese, F. The ORCA Program System. *WIREs Comput. Mol. Sci.* **2012**, *2*, 73–78.

[86] Perdew, J. P.; Ernzerhof, M.; Burke, K. Rationale for Mixing Exact Exchange with Density Functional Approximations. *J. Chem. Phys.* **1996**, *105*, 9982–9985.

[87] Grimme, S.; Antony, J.; Ehrlich, S.; Krieg, H. A Consistent and Accurate Ab Initio Parametrization of Density Functional Dispersion Correction (DFT-D) for the 94 Elements H-Pu. *J. Chem. Phys.* **2010**, *132*, 154104.

[88] Grimme, S.; Ehrlich, S.; Goerigk, L. Effect of the Damping Function in Dispersion Corrected Density Functional Theory. *J. Comput. Chem.* **2011**, *32*, 1456–1465.
[89] Weigend, F.; Ahlrichs, R. Balanced Basis Sets of Split Valence, Triple Zeta Valence and Quadruple Zeta Valence Quality for H to Rn: Design and Assessment of Accuracy. *Phys. Chem. Chem. Phys.* **2005**, *7*, 3297–3305.

[90] Weigend, F. Accurate Coulomb-Fitting Basis Sets for H to Rn. *Phys. Chem. Chem. Phys.* **2006**, *8*, 1057–1065.

[91] Andzelm, J.; Kölmel, C.; Klamt, A. Incorporation of Solvent Effects into Density Functional Calculations of Molecular Energies and Geometries. *J. Chem. Phys.* **1995**, *103*, 9312–9320.

[92] Barone, V.; Cossi, M. Quantum Calculation of Molecular Energies and Energy Gradients in Solution by a Conductor Solvent Model. *J. Phys. Chem. A* **1998**, *102*, 1995–2001.

[93] Cossi, M.; Rega, N.; Scalmani, G.; Barone, V. Energies, Structures, and Electronic Properties of Molecules in Solution with the C-PCM Solvation Model. *J. Comput. Chem.* **2003**, *24*, 669–681.

[94] MongoDB Inc., MongoDB 3.2. www.mongodb.com, (Accessed: 30. September 2016).

[95] Hunter, J. D. Matplotlib: A 2D Graphics Environment. *Comput. Sci. Eng.* **2007**, *9*, 90–95.

[96] McKinney, W. Data Structures for Statistical Computing in Python. Proceedings of the 9th Python in Science Conference. 2010; pp 51 – 56.

[97] Bakó, I.; Megyes, T.; Pálinkás, G. Structural Investigation of Water–Acetonitrile Mixtures: An Ab Initio, Molecular Dynamics and X-Ray Diffraction Study. *Chem. Phys.* **2005**, *316*, 235–244.

[98] Amabilino, S.; Bratholm, L. A.; Bennie, S. J.; Vaucher, A. C.; Reicher, M.; Glowacki, D. R. Training Neural Nets To Learn Reactive Potential Energy Surfaces Using Interactive Quantum Chemistry in Virtual Reality. *J. Phys. Chem. A* **2019**, *123*, 4486–4499.
[99] Lunghi, A.; Sanvito, S. A Unified Picture of the Covalent Bond within Quantum-Accurate Force Fields: From Organic Molecules to Metallic Complexes’ Reactivity. Sci. Adv. 2019, 5, eaaw2210.

[100] Lu, J.; Wang, C.; Zhang, Y. Predicting Molecular Energy Using Force-Field Optimized Geometries and Atomic Vector Representations Learned from an Improved Deep Tensor Neural Network. J. Chem. Theory Comput. 2019, 15, 4113–4121.

[101] Brunken, C.; Reiher, M. Self-Parametrizing System-Focused Atomistic Models. 2019, arXiv: 1908.10492.

[102] Simm, G. N.; Reiher, M. Error-Controlled Exploration of Chemical Reaction Networks with Gaussian Processes. J. Chem. Theory Comput. 2018, 14, 5238–5248.

[103] Weymuth, T.; Proppe, J.; Reiher, M. Statistical Analysis of Semiclassical Dispersion Corrections. J. Chem. Theory Comput. 2018, 14, 2480–2494.

[104] Proppe, J.; Reiher, M. Mechanism Deduction from Noisy Chemical Reaction Networks. J. Chem. Theory Comput. 2019, 15, 357–370.

[105] Proppe, J.; Gugler, S.; Reiher, M. Gaussian Process-Based Refinement of Dispersion Corrections. 2019, arXiv: 1906.09342.