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Assessing the Conformational Equilibrium of Carboxylic Acid via Quantum Mechanical and Molecular Dynamics Studies on Acetic Acid

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ABSTRACT: Accurate hydrogen placement in molecular modeling is crucial for studying the interactions and dynamics of biomolecular systems. The carboxyl functional group is a prototypical example of a functional group that requires protonation during structure preparation. To our knowledge, when in their neutral form, carboxylic acids are typically protonated in the syn conformation by default in classical molecular modeling packages, with no consideration of alternative conformations, though we are not aware of any careful examination of this topic. Here, we investigate the general belief that carboxylic acids should always be protonated in the syn conformation. We calculate and compare the relative energetic stabilities of syn and anti acetic acid using ab initio quantum mechanical calculations and atomistic molecular dynamics simulations. We focus on the carboxyl torsional potential and configurations of microhydrated acetic acid from molecular dynamics simulations, probing the effects of solvent, force field (GAFF vs GAFF2), and partial charge assignment of acetic acid. We show that while the syn conformation is the preferred state, the anti state may in some cases also be present under normal NPT conditions in solution.

1. INTRODUCTION

The carboxyl functional group, –COOH, is widespread in nature and highly biochemically relevant. It is present in amino acids that compose proteins, fatty acids of cell membranes, and naturally occurring organic compounds (e.g., niacin, citric acid, biotin). This group is very common in medicinal compounds, found in over 450 marketed drugs including nonsteroidal anti- inflammatory drugs (e.g., aspirin, ibuprofen), antibiotics (e.g., penicillin), and cholesterol-lowering statins (e.g., atorvastatin (Lipitor)). The presence of the hydrophilic carboxyl moiety on organic compounds can confer high solubility in water, which can be important to consider when designing new chemical reactions or developing new medicinal compounds. This group can also have important implications for pharmaceutical drugs; for example, drugs with a carboxyl functional group can be more metabolically unstable6 or have more difficulty diffusively crossing membranes. Given the carboxyl group's ubiquitous presence in nature and its importance as a functional group, understanding its conformational preferences in various settings is fundamental for the design, modification, and property prediction of new and existing molecules.

The preferred orientation of hydroxyl in the carboxyl functional group in solution is a matter of some debate, even for acetic acid, an archetypal carboxylic acid. Given a typical pKa of less than 5, the carboxyl group will usually be in the unprotonated, anionic form at neutral pH when exposed to the environment. However, the pKa may be significantly shifted as part of a ligand in a protein binding pocket or on protein side chains involved in reaction mechanisms. The two equilibrium conformations of the protonated carboxyl group are denoted syn (Figure

1a), where the OCCOCH dihedral angle is defined here to be 0°, and anti (Figure 1b), where the same dihedral angle is defined here as 180°. It is widely believed that the preferred conformation of carboxyl is the syn arrangement, from which there is a large energetic penalty to reach the anti arrangement. The reasoning behind this idea lies in the perceived extra stability of intramolecular hydrogen bonding that occurs in the syn structure. This belief is supported by a number of experimental and theoretical studies done in gas phase, and there is no doubt that this is the preferred conformation in the gas phase.⁷⁻¹¹

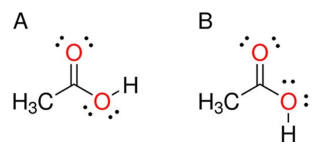


Figure 1. Lewis structure of acetic acid in (a) syn and (b) anti conformations.

The orientational preference of COOH is considerably more complex outside of gas phase. While some workers remain convinced that syn will be more stable, a variety of evidence indicates that this may not always be the case. A recent review article¹² discusses the competition between intramolecular and intermolecular hydrogen bonds in solution, stating that an intramolecular hydrogen bond may be disrupted in protic solution, such as water, when the increase in internal energy is offset by two or more solute—solvent intermolecular hydrogen bonds. Another study found that the carboxyl group has no strong preference, kinetically and thermodynamically, for the syn (or anti) conformation in proton transfer catalysis. ¹³ The anti state may also be important to consider when calculating solvation free energies. ¹⁴ In addition, the anti conformation is not insignificantly represented in structures from the Cam-bridge Structural Database, ¹⁵ supported by related crystallographic and theoretical charge density studies. ^{16,17} The carboxyl group may also be strongly influenced by its surroundings, such as within a protein binding site, to prefer either the syn or anti state. In general, the local environment plays a large role in the conformational state of the carboxyl group, and the preferred orientation is not always obvious.

Past work investigating acetic acid in solvent predominantly considers the syn state, such as in studies characterizing hydrogen-bonding interactions of acetic acid microhydrates using DFT-B3LYP calculations18 or assessing the dimer form in various stages of hydration theoretically^{19,20} and experimentally.²¹ One recent work examining solvent stabilization using DFT-ωB97X-D calculations²² indicates that water may modulate the conformational preferences of acetic acid; however, to our knowledge there has not been a systematic investigation of the preferred conformational state of the carboxyl group in solution. We believe that this collection of evidence on the orientational preference of COOH in solvent lacks a clear, definitive answer on whether both conformational states of the carboxyl group may reasonably be populated in normal aqueous solution when this group is in its neutral protonation state.

In this work, we aim to understand the relative conformational stability and energetic barrier for carboxyl functional group interconversion in both gas and aqueous phases. We present

our investigation on monomeric acetic acid using both ab initio quantum mechanical (QM) calculations and atomistic molecular dynamics (MD) simulations.

2. METHODS

Past gas phase QM studies clearly indicate a preference for the syn structure of the COOH group such as in acetic acid. However, classical all-atom MD simulations show that both are equally favorable in solution, at least with the energy model ("force field") employed. ¹⁴ This could be a real effect of water on the conformational preferences, or a limitation of the force field employed. Therefore, we need to examine a more intermediate region between gas phase QM and solution phase atomistic MD to settle the issue more definitively. Specifically, we look at QM in implicit solvent as well as QM data of snapshots pulled from MD simulations; on the MD end, we consider the effects of force field as well as solvation state.

We present an overview of our approach then discuss the methods in further detail. A torsion drive was conducted on acetic acid over the aforementioned dihedral angle. We conduct restrained geometry optimizations using two different QM methods, each with and without the presence of implicit solvent. Then, we carry out a set of geometry optimizations on pentahydrated acetic acid with varied water configurations obtained from MD simulations. We compared these energies for both syn and anti structures.

On the MD side, we compute a series of free energy landscapes, also known as potentials of mean force (PMFs), from driving the relevant torsion in acetic acid. We evaluated the sensitivity of these one-dimensional free energy surfaces to the force field (GAFF or GAFF2), partial charge assignment, and solvation state. We consider the force field because this factor is likely to vary among users running MD simulations. The partial charge set assigned to a solute depends on the initial conformation and is typically fixed throughout MD simulations, so we investigate potential implications of choosing one set or another. Finally, we compare the results of gaseous and aqueous phases to shed light on how reasonably the syn and anti states may be occupied in either scenario.

2.1. Ab Initio Torsion Drive of Acetic Acid. Acetic acid configurations of the carboxyl O+C+O+H dihedral angle were generated and used as input for both QM torsion drives and MD umbrella sampling simulations. The dihedral angle was rotated using VMD23 in 15° increments from 0° to 360°, yielding 24 total conformations.

The QM torsion drives were run using Turbomole version 7.124 with two different levels of theory: HF/6-31G* and TPSSh-D3BJ/def2-TZVP. The former method, using Hartree–Fock²⁵ with the Pople 6-31G* basis set,^{26,27} was chosen for consistency with the methods often employed in parametrization of force fields used for molecular simulation.²⁸ This low level method also provides historical perspective contributing to the strong bias favoring the syn conformation of the carboxyl group. Taking a more rigorous approach, we employed the TPSSh hybrid functional^{29,30} with Grimme's D3 dispersion correction³¹ and Becke–Johnson damping,³² in combination with the Karlsruhe triple- ζ basis set def2-TZVP.33 We chose the TPSSh functional because prior work indicates it is a suitable approach for treating the molecular dipole moments and polarizabilities of these hydrogen-bonding systems.^{34,35} We also run calculations in implicit solvent with each of the aforementioned methods using the conductor-like screening model (COSMO) with outlying charge corrections.^{36–39}

- Ab Initio Geometry Optimizations from Molecular Dynamics Configurations. We sample various configurations of water molecules around acetic acid by running separate MD simulations of the syn and anti conformations in a box of TIP3P water molecules. The structures were solvated using Antechamber⁴⁰ within a cubic box with TIP3P waters⁴¹ such that the minimum distance between the solute and the edge of the periodic box was 12 Å. Dynamics were run using GROMACS version 5.0.4 with the leapfrog stochastic dynamics integrator and a 2 fs time step. We use a Langevin thermostat for the temperature at 298.15 K with a frictional constant of 2.0 ps⁻¹. The pressure was maintained at 1 atm using the Parrinello-Rahman pressure coupling scheme with a time constant of 10 ps-1 and an isothermal compressibility of 4.5×10^{-5} bar⁻¹. All bonds involving hydrogen atoms were constrained using the LINCS algorithm.42 The systems were simulated with 2500 steps of steepest descent minimization, 50 ps constant volume and temperature (NVT) equilibration, 5 ns of constant pressure and temperature (NPT) equilibration, and then 5 ns of NPT production. Trajectory snapshots were extracted of the most similar configurations for acetic acid and its five closest waters using a root-mean-square deviation clustering of geometries with a 2 Å cutoff. This yielded 14 snapshots for the pentahydrated syn conformation and 17 snapshots for the pentahydrated anti conformation. Each snapshot was MM- optimized via OpenEve's OEChem Python Toolkit⁴³ using the MMFF94S force field^{44–49} and, then, subsequently QM- optimized using Turbomole version 7.124 with COSMO-TPSSh-D3BJ/def2-TZVP. ^{29–33,36–39}
- 2.3. **MD Simulations with Umbrella Sampling along the Carboxyl Dihedral Angle.** We used umbrella sampling⁵⁰ molecular dynamics to compute a potential of mean force (PMF) to analyze the free energy landscape projected onto this one-dimensional coordinate. We compared MD results with the GAFF51 and GAFF252 classical all-atom force fields, with partial charges assigned by the AM1-BCC^{53,54} approach. We consider effects of the solute partial charges in the MD simulations by carrying out MD simulations with AM1-BCC charges assigned from the syn configuration as well as charges assigned from the anti configuration. Energetics were examined in gas phase, then in solvent using explicit TIP3P water molecules.⁴¹

These simulations were run using GROMACS version 5.0.4.55 Each acetic acid configuration generated in VMD was set with partial charges from the AM1-BCC charge model53,54 on the syn (0°) conformation as implemented in OpenEye's Python toolkits.⁴³ The partial charges of the solute depend on initial configuration, so we also consider the anti (180°) conformation for computing partial charges. The OCCOH dihedral angle was restrained in both gas phase and aqueous MD simulations, using a harmonic force constant of 300 kJ/mol·(rad²) (approximately 0.022 kcal/mol·(deg²)).

For the gas phase simulations, the reference temperature of 298.15 K was maintained using Langevin dynamics with a frictional constant of 1.0 ps⁻¹. Maintaining the GROMACS parameters described earlier, the systems underwent steepest descent minimization over 2500 steps, NVT equilibration for 50 ps, and NVT production for 1 ns.

For the explicit solvent simulations, the solvation parameters and other MD simulation settings were maintained as described earlier in the section, 2.2 Ab Initio Geometry Optimizations from Molecular Dynamics Configurations. These systems were simulated with 2500 steps of steepest descent minimization, 50 ps NVT equilibration, 50 ps NPT equilibration, and 5 ns NPT production. The configurations with dihedral angle around 270° seemed not converged, so six conformations were extended 5 ns for a total of 10 ns each: 65°, 90°, 105°, 255°, 270°, 285°. However, there was little to no change in the resulting PMFs.

Analysis of all umbrella sampling simulations was completed with the MBAR algorithm56 to produce the potentials of mean force (PMFs) for rotation of the carboxyl dihedral angle.

3. RESULTS AND DISCUSSION

Results from both QM and MD approaches support former work and the general understanding that syn is favored in gas phase. They also indicate that the anti conformation may also be populated to a significant extent in water. We address our QM results first and then discuss MD results.

3.1. **Ab Initio Torsion Drive of Acetic Acid.** Our QM calculations in gas phase and implicit solvent show that syn is highly favored in the gas phase but the difference becomes less significant in solvent. From the torsion drive obtained via ab initio QM calculations, the syn-anti energy difference is 7.14 kcal/mol with the basic HF/6-31G* method and decreases to 5.24 kcal/mol with the higher level of theory using the TPSSh functional (Figure 2). With COSMO, a similar trend is seen in which the higher level of theory yields a smaller energy difference between the syn and anti structures. With either level of theory, adding implicit solvent significantly lowers the relative energy difference between syn and anti from 5–7 to 2–3 kcal/mol. A 5–7 kcal/mol difference is large enough that such configurations would occur only extremely rarely, whereas 2–3 kcal/mol is enough that such conformations will occur sporadically in solution (3–7% of the time) and could easily be stabilized by interactions with a nearby receptor or other biomolecule with a strain energy no larger than that reported in many binding interactions, ^{57,58} making it potentially relevant functionally.

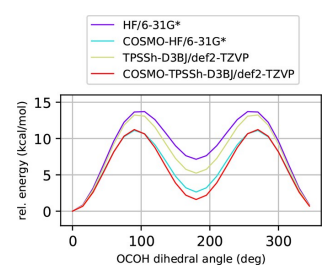


Figure 2. QM torsion drive of acetic acid carboxyl dihedral angle for HF and TPSSh methods. In each case, implicit solvation with COSMO reduces the energy barrier and the relative minima energy to 5–7 and 2–3 kcal/mol, respectively.

We now turn our focus to the energy barrier from the syn state to the anti state. This feature is not particularly critical in molecular simulation, as in most cases systems will be at equilibrium given sufficient relaxation time and sampling. That being said, the energy barrier has

implications for inter- conversion between the two states. One conformation may be more structurally relevant than the other in certain scenarios, and a modeler may wish to achieve an accurate representation of the populations of both conformations. The barrier associated with the rotation of the carboxyl dihedral angle determines how easy it is to interconvert between and sample different conformations. From our QM results, we see a large energetic cost, or barrier, of 13–14 kcal/mol separating the syn form from the anti form in gas phase. Solvation with COSMO reduces this barrier height to around 11 kcal/mol.

Overall, the relative energy difference between the syn and anti conformations of acetic acid appears not very large, especially in the aqueous conditions relevant to biochemistry. The relative energy comparisons from the QM torsion drives are summarized in the top four lines of Table 1. Note that, from our QM results, these are relative energies rather than relative free energies; with MD in the following section; we obtain relative free energies.

Table 1. Summary of Relative Energy Differences between Syn and Anti Conformations of Acetic Acid as Well as Free Energy Barriers of Interconversion^a

method	solvation	minima ^b	barrier
HF/6-31G*	gas	7.1	13.7
HF/6-31G*	COSMO	2.8	11.2
TPSSh/def2-TZVP ^c	gas	5.2	13.2
TPSSh/def2-TZVP	COSMO	1.6	11.2
vac_SC_GAFF	gas	6.2 ± 0.2	12.7 ± 0.3
vac_SC_GAFF2	gas	5.9 ± 0.2	11.7 ± 0.3
vac_AC_GAFF	gas	3.4 ± 0.2	11.0 ± 0.3
vac_AC_GAFF2	gas	3.3 ± 0.2	10.1 ± 0.3
sol_SC_GAFF	TIP3P	-0.8 ± 0.1	7.0 ± 0.2
sol_SC_GAFF2	TIP3P	-0.7 ± 0.1	6.4 ± 0.2
sol_AC_GAFF	TIP3P	-1.3 ± 0.1	6.7 ± 0.2
sol_AC_GAFF2	TIP3P	-1.4 ± 0.1	6.1 ± 0.2

^aThe first four lines are results from QM torsion drives, and the last eight from umbrella sampling are from atomistic molecular dynamics simulations. Energies are listed in units of kilocalories per mole

^bAll relative energy differences are taken with respect to acetic acid's syn conformation.

^cDispersion corrections added with all TPSSh calculations in this work. See details in text.

3.2. Ab Initio Geometry Optimizations from Molecular Dynamics Configurations. To rule out the possibility that stabilization of the anti form in the torsion drive is due to implicit solvent model alone, and to determine whether explicit water might provide additional stabilization, we examined acetic acid with explicit water molecules. We first examined trihydrated syn and anti acetic acid (details in the Supporting Information). However, recent work on the microhydration of acetic acid suggests that the particular arrangement of water molecules may be important when comparing energetic stabilities of acetic acid conformations.²² Given that we are interested in solution-phase behavior, the actual solution-phase geometry of water molecules around acetic acid then becomes very important. In order to reduce any artificial effects of water placement, we sample various conformations of water molecules around acetic acid by running molecular dynamics simulations for each of the syn and anti forms. Both simulations were run using the syn charges for context as these are predominantly used in present-day molecular simulations. Configurations of acetic acid with its five nearest waters were clustered by root-mean-square deviation of geometries. The most common arrangements were extracted for QM optimization in implicit solvent using the method COSMO- TPSSh-D3BJ/def2-TZVP.

The violin plots in Figure 3 display the distributions for the relative energies of the syn (left side) and anti (right side) pentahydrated configurations of acetic acid. Here, we see that the distribution for the syn configurations skews toward lower energies compared to the anti configurations. However, the energy values of the extrema are quite similar, and the population of the anti form at low energies is nonnegligible.

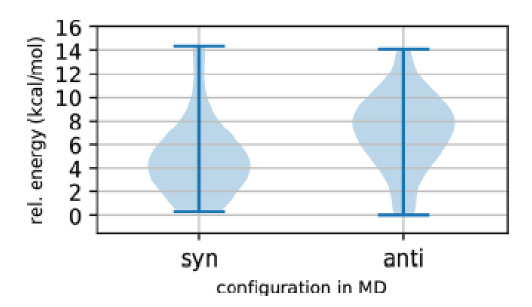


Figure 3. Violin plots for relative energy distributions of pentahydrated syn and anti conformations of acetic acid. The data represent COSMO-TPSSh-D3BJ/def2-TZVP energies of configura- tions taken from MD simulations of the syn form (14 snapshots) and the anti form (17 snapshots).

3.3. *MD Simulations with Umbrella Sampling along Carboxyl Dihedral Angle.* Our above QM calculations study only conformational energies, not free energies, so we computed the one-dimensional free energy landscape (the potential of mean force, or PMF) of rotating the acetic acid dihedral angle with classical molecular dynamics. The MD results in gas phase and in explicit solvent are in qualitative agreement with our QM data and indicate that water substantially increases the stability of the anti conformation. We considered various force fields, partial charge sets, and solvation states for a total of eight PMFs. Atomic partial charges are held fixed within our simulations, as is typical in MD, but these charges are sensitive to the molecular conformation when assigning charges, so we assigned charges using both conformations. Hereafter we use the notation SC for acetic acid partial charges obtained from the syn conformation and AC for charges obtained from the anti conformation. Error bars on the PMFs are obtained from the MBAR estimator.56 We present a comprehensive comparison in Figure 4 and in Table 1 and discuss each of these three factors (force field, charge set, and solvation state) separately.

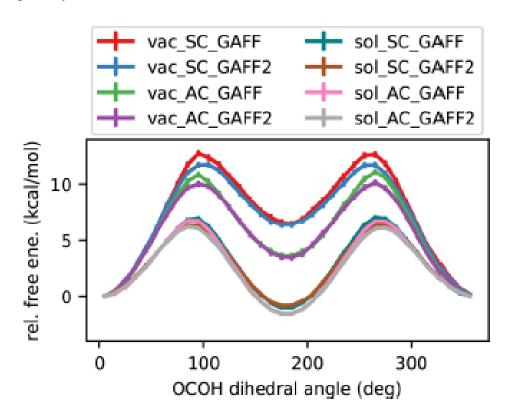


Figure 4. PMFs of rotating the acetic acid carboxyl dihedral angle. We consider variations on the force field (GAFF, GAFF2), solute AM1- BCC partial charges (starting from syn or anti), and solvation state (gas phase, explicit TIP3P waters).

Considering the GAFF and GAFF2 force fields, the PMFs are in good agreement with each other in both gas and aqueous phases as well as with either SC or AC (Figures 4, 5). We observe consistent relative free energies between the syn and anti minima. In gas phase, for the SC solute, the syn structure is favored in free energy by 6.2 ± 0.2 kcal/mol with GAFF and

 5.9 ± 0.2 kcal/mol with GAFF2 (Figure 5). In aqueous phase, the anti structure is favored in free energy by -0.7 ± 0.1 kcal/ mol with GAFF and -1.4 ± 0.1 kcal/mol with GAFF2 (Figure 4, teal vs brown). These qualitative conclusions are the same when considering the AC solute. Thus, GAFF and GAFF2 give very similar results for the conformational equilibrium of acetic acid which holds true regardless of the partial charge set. Overall these results, at least within the classical framework, indicate that explicit solvent provides approximately 5-8 kcal/ mol of stabilization of the anti conformation relative to the syn conformation. This trend is in the same direction as that provided by COSMO implicit solvent but provides further stabilization. (Error bars shown here are obtained from the MBAR estimator.)

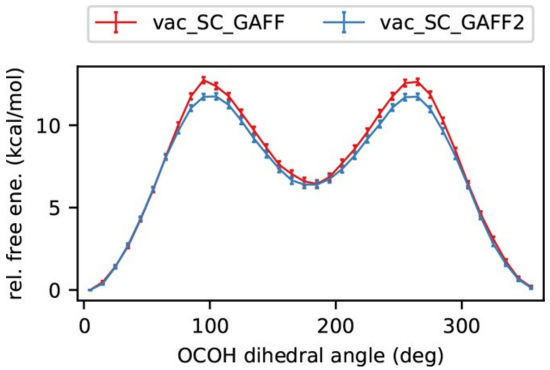


Figure 5. Comparison of GAFF and GAFF2 force fields in PMFs of rotating the acetic acid carboxyl dihedral angle. Both are in strong agreement with each other. The PMFs displayed in this figure came from gas phase simulations with syn charges. Similar conclusions were drawn for PMFs from aqueous simulations and from using anti charges (Figure 4).

We also compare the two force fields in terms of the conformational transition barriers. We note that the GAFF barrier height is higher than the GAFF2 barrier in each pairwise combination of the two force fields with various solvent and charge models. The barrier height differences are 0.9 ± 0.4 kcal/mol in gas phase (compare barrier heights in Figure 4 for red vs blue and for green vs purple). The rotational barriers differ by 0.6 ± 0.2 kcal/mol in aqueous phase (compare barrier heights in Figure 4 for teal vs brown and for pink vs gray). For both gaseous and aqueous states, the effects of the partial charges on the PMFs are stronger than those of the force field. For example, in Figure 4, the red and green curves are more distinct from each other, while the red and blue curves are more similar. Since the partial charges of the solute may affect the PMFs more so than the force field, as shown here, one should carefully consider other likely conformations when assigning partial charges. Next we further investigate the solute partial charge sets.

There is a pronounced difference in the PMFs depending on the conformation used to charge acetic acid (Figure 6). Charges are typically fixed throughout a molecular dynamics simulation, meaning that initial charge assignment is important for capturing correct energetics throughout a simulation. The free energy difference between the syn and anti structures is notably larger in gas phase than in water. When we use the syn form to obtain AM1-BCC charges (SC), the gas phase PMFs are higher in energy for both the barrier height and the two minima (Figure 7a) compared to using the anti form to obtain AM1-BCC charges (AC). Qualitatively, the SC set is slightly stronger in magnitude than the AC set, meaning a slightly stronger polarization along the bonds of the carboxyl group; this is consistent with the intramolecular hydrogen bonding aspect of the syn conformation. The stronger SC partial charges contribute to increased stabilization of the lower- energy syn structure in gas phase, which results in a greater free energy difference and barrier height compared to AC. On the other hand, in water, (Figure 7b), syn and anti are closer in relative free energy for SC than for AC. In this setting, syn is higher in energy than anti. Once again, the stronger SC partial charges contribute to increased stabilization of syn, in this case via more stabilizing interactions with the solvent. Here, the two minima are closer in free energy. Therefore, we see again that the relative free energies at the minima are governed more strongly by solute charges than by force field.

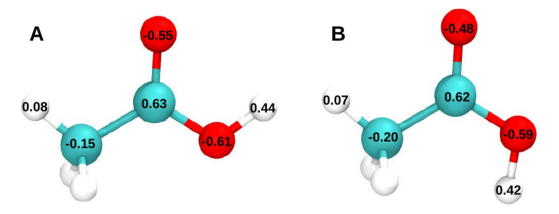


Figure 6. AM1-BCC charges generated for (a) syn and (b) anti configurations of acetic acid.

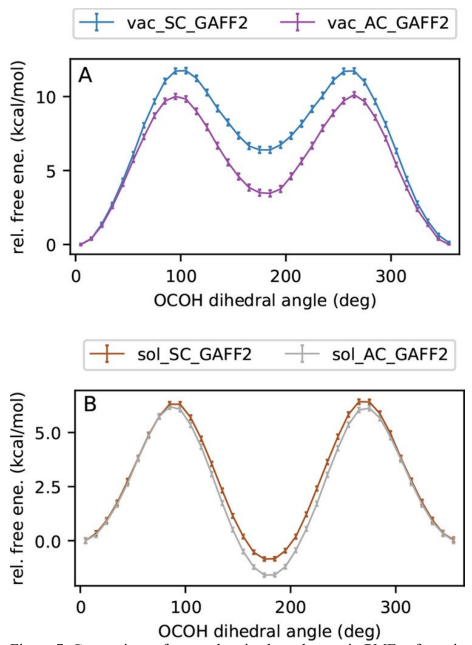


Figure 7. Comparison of syn and anti solute charges in PMFs of rotating the acetic acid carboxyl dihedral angle. In each situation with anti charges (A) and syn charges (B), the AC set more strongly stabilizes the anti conformation than the SC set.

We take a final look at the MD PMFs in the lens of gaseous versus aqueous phases. These results are in harmony with earlier work on ibuprofen (a carboxylic acid) which found that the syn conformation was favorable in vacuum but the anti conformation was slightly preferred in water. ¹⁴ The major takeaway from the aqueous phase PMFs is that the anti conformation of acetic acid is the lower free energy state in solution due to an increased ability to form stabilizing interactions with the solvent. This conclusion qualitatively parallels the result obtained with COSMO-QM calculations on microhydrated acetic acid which showed that the anti conformation

is lower in energy than the syn conformation by about 1.6 kcal/mol, at least for certain arrangements of water molecules.

Overall, the MD results are qualitatively consistent with QM calculations in determination of relative energy differences of the minima and energy barriers for conformational inter- conversion. The SC charge set seems better than the AC set in reproducing the relative energy differences obtained with QM DFT in gas phase and in implicit solvent, consistent with our previous practice of considering this conformation more important when assigning charges.

To summarize our PMF results, we considered the effects of force field, partial charge set, and solvation state on the relative minima free energies as well as on the transition barriers between the two minima. The force fields GAFF and GAFF2 yielded generally similar results to each other. The PMFs in both gas phase and aqueous phase revealed strong dependence on solute charges, especially at the minima. More specifically, the set of partial charges assigned to acetic acid is sensitive to the orientation of O–H in the carboxyl group, leading to variations of up to several kcal/mol in the free energy difference between the syn and anti structures. Lastly, the dihedral rotation free energy barriers between the syn and anti conformations are more dependent on the charge set than the force field in gas phase simulations, while they are more influenced by the force field in aqueous phase simulations. All eight PMFs, obtained from permutation of the force field, solute charges, and solvation state are summarized in Figure 4 and Table 1.

4. CONCLUSIONS

Our results call into question the conventional wisdom that carboxylic acids will almost always be in the "more stable" syn conformation in biomolecular systems. Typically, the increased stability of the syn form is understood to be from the stabilizing intramolecular interaction between the hydrogen atom in the hydroxyl group and the carbonyl oxygen. This idea is in tune with gas phase results we present in this work. However, in aqueous phase, we conclude that the anti state may nearly be as populated as the syn state due to stabilizing interactions from the solvent. Thus, for MD studies that involve a carboxylic acid or other functional group with possible intramolecular hydrogen bonds, it may be necessary to ensure sufficient sampling of all potentially relevant conformations in solution. This can be challenging given the particularly large barrier associated with rotation of the carboxylic acid torsion.

Our findings also have implications for partial charge calculations for parameter assignment for MD simulations. Carboxylic acids are a case in which neither partial charge set adequately represents the electrostatics of the solute as it samples various conformations. When generating an empirical force field, such as for a small molecule ligand, charges are typically computed for a particular given conformation. These fixed charges are then used for scenarios involving conformational change. In this work, we observe that different solute charges may lead to deviations in relative free energies to as large as 3 kcal/mol. Interconversion is not expected to be frequent, given that the torsional barrier is at least 6 kcal/mol. For that reason, one may wish to treat syn and anti conformation charges individually, though this could present difficulties in cases that interconversion is needed for convergence (e.g., a carboxylic acid in a binding site where one conformation forms better contacts than the other). As an alternative approach, the use of polarizable charges may provide a more holistic picture of the carboxyl group's variable nature.

The carboxyl conformational equilibrium has implications for several other types of studies. Hydration free energy calculations may lead to results which depend substantially on the starting conformation. For example, kinetic trapping into one particular conformation can lead to computed hydration free energies which are sensitive to starting conformation and vary by more than 2 kcal/mol because of large torsional barriers. ¹⁴ This work also informs efforts to accurately calculate pKa values for ionizable side chains in proteins, i.e., aspartate and glutamate. ^{59–69} An accurate insight into the preferred aqueous phase structure of the carboxyl group is important for catalysis, with impacts in atmospheric science and industrial processes. 70 Further impact may be in crystal engineering and drug cocrystallization, in which the carboxyl group is often used to promote aqueous solubility. ⁵ Theoretical studies on proton transfer such as on solvated acetic acid 71 or on green fluorescent protein ^{72,73} typically employ the syn conformation due to its expected energetic preference; however, it is worth investigating possible adaptations of carboxyl groups to their local environments. Being aware of the carboxyl moiety's nuanced conformational preferences in different environments may thus lead to better insight for calculated properties, reactivity, and molecular design.

ASSOCIATED CONTENT

*S Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jcim.8b00835.

Raw data and Python scripts for all figures, configuration input files for all calculations, initial and final coordinates of all structures, Python script for computing free energies from PMFs (ZIP)

Computations and discussion on trihydrated acetic acid, comparison of acetic acid geometries to existing literature, discussion on choice of QM method with calculations on water hexamers (PDF)

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Notes

The authors declare the following competing financial interest(s): D.L.M. serves on the scientific advisory board of OpenEye Scientific Software and is an Open Science Fellow with Silicon Therapeutics.

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REFERENCES

- (1) Ballatore, C.; Huryn, D. M.; Smith, A. B. Carboxylic Acid (Bio)Isosteres in Drug Design. ChemMedChem 2013, 8, 385–395.
- (2) Lou, Y.; Zhu, J. In Bioactive Carboxylic Compound Classes; Lamberth, C., Dinges, J., Eds.; Wiley-VCH Verlag GmbH & Co. KGaA, 2016; pp 221–236.
- (3) David, S. E.; Timmins, P.; Conway, B. R. Impact of the Counterion on the Solubility and Physicochemical Properties of Salts of Carboxylic Acid Drugs. Drug Dev. Ind. Pharm. 2012, 38, 93–103.
- (4) McNamara, D. P.; Childs, S. L.; Giordano, J.; Iarriccio, A.; Cassidy, J.; Shet, M. S.; Mannion, R.; O'Donnell, E.; Park, A. Use of a Glutaric Acid Cocrystal to Improve Oral Bioavailability of a Low Solubility API. Pharm. Res. 2006, 23, 1888–1897.
- (5) Desiraju, G. R. Crystal Engineering: From Molecule to Crystal. J. Am. Chem. Soc. 2013, 135, 9952–9967.
- (6) Lassila, T.; Hokkanen, J.; Aatsinki, S.-M.; Mattila, S.; Turpeinen, M.; Tolonen, A. Toxicity of Carboxylic Acid-Containing Drugs: The Role of Acyl Migration and CoA Conjugation Investigated. Chem. Res. Toxicol. 2015, 28, 2292–2303.
- (7) Nagy, P. I. The Syn-Anti Equilibrium for the COOH Group Reinvestigated. Theoretical Conformation Analysis for Acetic Acid in the Gas Phase and in Solution. Comput. Theor. Chem. 2013, 1022, 59–69.
- (8) Nagy, P. I.; Smith, D. A.; Alagona, G.; Ghio, C. Ab Initio Studies of Free and Monohydrated Carboxylic Acids in the Gas Phase. J. Phys. Chem. 1994, 98, 486–493.
- (9) Sato, H.; Hirata, F. The Syn-/Anti-Conformational Equilibrium of Acetic Acid in Water Studied by the RISM-SCF/MCSCF Method1. J. Mol. Struct.: THEOCHEM 1999, 461–462, 113–120.
- (10) Wiberg, K. B.; Laidig, K. E. Barriers to Rotation Adjacent to Double Bonds. 3. The Carbon-Oxygen Barrier in Formic Acid, Methyl Formate, Acetic Acid, and Methyl Acetate. The Origin of Ester and Amide Resonance. J. Am. Chem. Soc. 1987, 109, 5935–5943.
- (11) Derissen, J. L. A Reinvestigation of the Molecular Structure of Acetic Acid Monomer and Dimer by Gas Electron Diffraction. J. Mol. Struct. 1971, 7, 67–80.
- (12) Nagy, P. I. Competing Intramolecular vs. Intermolecular Hydrogen Bonds in Solution. Int. J. Mol. Sci. 2014, 15, 19562–19633.
- (13) Montzka, T. A.; Swaminathan, S.; Firestone, R. A. Reversal of Syn-Anti Preference for Carboxylic Acids along the Reaction Coordinate for Proton Transfer. Implications for Intramolecular Catalysis. J. Phys. Chem. 1994, 98, 13171–13176.
- (14) Klimovich, P. V.; Mobley, D. L. Predicting Hydration Free Energies Using All-Atom Molecular Dynamics Simulations and Multiple Starting Conformations. J. Comput.-Aided Mol. Des. 2010, 24, 307–316.
- (15) D'Ascenzo, L.; Auffinger, P. A Comprehensive Classification and Nomenclature of Carboxyl-Carboxyl(Ate) Supramolecular Motifs and Related Catemers: Implications for Biomolecular Systems. Acta Crystallogr., Sect. B: Struct. Sci., Cryst. Eng. Mater. 2015, 71, 164–175.
- (16) Pal, R.; Reddy, M. B. M.; Dinesh, B.; Venkatesha, M. A.; Grabowsky, S.; Jelsch, C.; Guru Row, T. N. Syn vs Anti Carboxylic Acids in Hybrid Peptides: Experimental and Theoretical Charge Density and Chemical Bonding Analysis. J. Phys. Chem. A 2018, 122, 3665–3679.

- (17) Medvedev, M.; Bushmarinov, I.; Lyssenko, K. Z-Effect Reversal in Carboxylic Acid Associates. Chem. Commun. 2016, 52, 6593–6596.
- (18) Gao, Q.; Leung, K. T. Hydrogen-Bonding Interactions in Acetic Acid Monohydrates and Dihydrates by Density-Functional Theory Calculations. J. Chem. Phys. 2005, 123, 074325.
- (19) Pasălic, H.; Tunega, D.; Aquino, A. J. A.; Haberhauer, G.;
- Gerzabek, M. H.; Lischka, H. The Stability of the Acetic Acid Dimer in Microhydrated Environments and in Aqueous Solution. Phys. Chem. Phys. 2012, 14, 4162–4170.
- (20) Chocholousŏva, J.; Vacek, J.; Hobza, P. Acetic Acid Dimer in the Gas Phase, Nonpolar Solvent, Microhydrated Environment, and Dilute and Concentrated Acetic Acid: Ab Initio Quantum Chemical and Molecular Dynamics Simulations. J. Phys. Chem. A 2003, 107, 3086–3092.
- (21) Ouyang, B.; Howard, B. The Monohydrate and Dihydrate of Acetic Acid: A High-Resolution Microwave Spectroscopic Study. Phys. Chem. Chem. Phys. 2009, 11, 366–373.
- (22) Krishnakumar, P.; Maity, D. K. Microhydration of Neutral and Charged Acetic Acid. J. Phys. Chem. A 2017, 121, 493–504.
- (23) Humphrey, W.; Dalke, A.; Schulten, K. VMD: Visual Molecular Dynamics. J. Mol. Graphics 1996, 14, 33–38.
- (24) TURBOMOLE, V7.1 2016; a development of University of Karlsruhe and Forschungszentrum Karlsruhe GmbH; available from http://www.turbomole.com.
- (25) Haser, M.; Ahlrichs, R. Improvements on the Direct SCF Method. J. Comput. Chem. 1989, 10, 104–111.
- (26) Hehre, W. J.; Ditchfield, R.; Pople, J. A. Self-Consistent Molecular Orbital Methods. XII. Further Extensions of Gaussian—Type Basis Sets for Use in Molecular Orbital Studies of Organic Molecules. J. Chem. Phys. 1972, 56, 2257–2261.
- (27) Hariharan, P. C.; Pople, J. A. The Influence of Polarization Functions on Molecular Orbital Hydrogenation Energies. Theoret. Chim. Acta 1973, 28, 213–222.
- (28) Cornell, W. D.; Cieplak, P.; Bayly, C. I.; Gould, I. R.; Merz, K. M.; Ferguson, D. M.; Spellmeyer, D. C.; Fox, T.; Caldwell, J. W.; Kollman, P. A. A Second Generation Force Field for the Simulation of Proteins, Nucleic Acids, and Organic Molecules. J. Am. Chem. Soc. 1995, 117, 5179–5197.
- (29) Staroverov, V. N.; Scuseria, G. E.; Tao, J.; Perdew, J. P. Comparative Assessment of a New Nonempirical Density Functional: Molecules and Hydrogen-Bonded Complexes. J. Chem. Phys. 2003, 119, 12129–12137.
- (30) Staroverov, V. N.; Scuseria, G. E.; Tao, J.; Perdew, J. P. Erratum: "Comparative Assessment of a New Nonempirical Density Functional: Molecules and Hydrogen-Bonded Complexes" [J. Chem. Phys. 119, 12129 (2003)]. J. Chem. Phys. 2004, 121, 11507–11507.
- (31) 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.
- (32) Grimme, S.; Ehrlich, S.; Goerigk, L. Effect of the Damping Function in Dispersion Corrected Density Functional Theory. J. Comput. Chem. 2011, 32, 1456–1465.
- (33) 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. Phys. 2005, 7, 3297–3305.
- (34) Laurent, A. D.; Jacquemin, D. TD-DFT Benchmarks: A Review. Int. J. Quantum Chem. 2013, 113, 2019–2039.

- (35) Hickey, A. L.; Rowley, C. N. Benchmarking Quantum Chemical Methods for the Calculation of Molecular Dipole Moments and Polarizabilities. J. Phys. Chem. A 2014, 118, 3678–3687.
- (36) 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, 2, 799–805.
- (37) Klamt, A.; Jonas, V.; Bürger, T.; Lohrenz, J. C. W. Refinement and Parametrization of COSMO-RS. J. Phys. Chem. A 1998, 102, 5074–5085.
- (38) Eckert, F.; Klamt, A. Fast Solvent Screening via Quantum Chemistry: COSMO-RS Approach. AIChE J. 2002, 48, 369–385.
- (39) Sinnecker, S.; Rajendran, A.; Klamt, A.; Diedenhofen, M.; Neese, F. Calculation of Solvent Shifts on Electronic G-Tensors with the Conductor-Like Screening Model (COSMO) and Its Self- Consistent Generalization to Real Solvents (Direct COSMO-RS). J. Phys. Chem. A 2006, 110, 2235–2245.
- (40) Wang, J.; Wang, W.; Kollman, P. A.; Case, D. A. Automatic Atom Type and Bond Type Perception in Molecular Mechanical Calculations. J. Mol. Graphics Modell. 2006, 25, 247–260.
- (41) Jorgensen, W. L.; Chandrasekhar, J.; Madura, J. D.; Impey, R. W.; Klein, M. L. Comparison of Simple Potential Functions for Simulating Liquid Water. J. Chem. Phys. 1983, 79, 926–935.
- (42) Hess, B.; Bekker, H.; Berendsen, H. J. C.; Fraaije, J. G. E. M. LINCS: A Linear Constraint Solver for Molecular Simulations. J. Comput. Chem. 1997, 18, 1463–1472.
- (43) OEChem, version 2.1.1, 2017, OpenEye Scientific Software Inc.: Santa Fe, NM, USA.
- (44) Halgren, T. A. Merck Molecular Force Field. II. MMFF94 van Der Waals and Electrostatic Parameters for Intermolecular Interactions. J. Comput. Chem. 1996, 17, 520–552.
- (45) Halgren, T. A. Merck Molecular Force Field. I. Basis, Form, Scope, Parameterization, and Performance of MMFF94. J. Comput. Chem. 1996, 17, 490–519.
- (46) Halgren, T. A. Merck Molecular Force Field. III. Molecular Geometries and Vibrational Frequencies for MMFF94. J. Comput. Chem. 1996, 17, 553–586.
- (47) Halgren, T. A.; Nachbar, R. B. Merck Molecular Force Field.
- IV. Conformational Energies and Geometries for MMFF94. J. Comput. Chem. 1996, 17, 587–615.
- (48) Halgren, T. A. Merck Molecular Force Field. V. Extension of MMFF94 Using Experimental Data, Additional Computational Data, and Empirical Rules. J. Comput. Chem. 1996, 17, 616–641.
- (49) Halgren, T. A. MMFF VI. MMFF94s Option for Energy Minimization Studies. J. Comput. Chem. 1999, 20, 720–729.
- (50) Torrie, G. M.; Valleau, J. P. Nonphysical Sampling Distributions in Monte Carlo Free-Energy Estimation: Umbrella Sampling. J. Comput. Phys. 1977, 23, 187–199.
- (51) Wang, J.; Wolf, R. M.; Caldwell, J. W.; Kollman, P. A.; Case, D. A. Development and Testing of a General Amber Force Field. J. Comput. Chem. 2004, 25, 1157–1174.
- (52) Case, D. A.; Betz, R. M.; Cerutti, D. S.; Cheatham, T. E., III; Darden, T. A.; Duke, R. E.; Giese, T. J.; Gohlke, H.; Goetz, A. W.; Homeyer, N.; Izadi, S.; Janowski, P.; Kaus, J.; Kovalenko, A.; Lee, T. S.; LeGrand, S.; Li, P.; Lin, C.; Luchko, T.; Luo, R.; Madej, B.; Mermelstein, D.; Merz, K. M.; Monard, G.; Nguyen, H.; Nguyen, H. T.; Omelyan, I.; Onufriev,

- A.; Roe, D. R.; Roitberg, A.; Sagui, C.; Simmerling, C. L.; Botello-Smith, W. M.; Swails, J.; Walker, R. C.; Wang, J.; Wolf, R. M.; Wu, X.; Xiao, L.; Kollman, P. A. AMBER 2016, University of California, San Francisco, 2016.
- (53) Jakalian, A.; Bush, B. L.; Jack, D. B.; Bayly, C. I. Fast, Efficient Generation of High-Quality Atomic Charges. AM1-BCC Model: I. Method. J. Comput. Chem. 2000, 21, 132–146.
- (54) Jakalian, A.; Jack, D. B.; Bayly, C. I. Fast, Efficient Generation of High-Quality Atomic Charges. AM1-BCC Model: II. Parameter- ization and Validation. J. Comput. Chem. 2002, 23, 1623–1641.
- (55) Abraham, M. J.; Murtola, T.; Schulz, R.; Pall, S.; Smith, J. C.;
- Hess, B.; Lindahl, E. GROMACS: High Performance Molecular Simulations through Multi-Level Parallelism from Laptops to Supercomputers. SoftwareX 2015, 1–2, 19–25.
- (56) Shirts, M. R.; Chodera, J. D. Statistically Optimal Analysis of Samples from Multiple Equilibrium States. J. Chem. Phys. 2008, 129, 124105.
- (57) Mobley, D. L.; Dill, K. A. Binding of Small-Molecule Ligands to Proteins: "What You See" Is Not Always "What You Get. Structure 2009, 17, 489–498.
- (58) Sahai, M. A.; Biggin, P. C. Quantifying Water-Mediated Protein-Ligand Interactions in a Glutamate Receptor: A DFT Study. J. Phys. Chem. B 2011, 115, 7085-7096.
- (59) van Vlijmen, H. W. T.; Schaefer, M.; Karplus, M. Improving the Accuracy of Protein pKa Calculations: Conformational Averaging versus the Average Structure. Proteins: Struct., Funct., Genet. 1998, 33, 145–158.
- (60) Li, H.; Robertson, A. D.; Jensen, J. H. Very Fast Empirical Prediction and Rationalization of Protein pKa Values. Proteins: Struct., Funct., Genet. 2005, 61, 704–721.
- (61) Bashford, D.; Karplus, M. pKa's of Ionizable Groups in Proteins: Atomic Detail from a Continuum Electrostatic Model. Biochemistry 1990, 29, 10219–10225.
- (62) Beroza, P.; Case, D. A. Including Side Chain Flexibility in Continuum Electrostatic Calculations of Protein Titration. J. Phys. Chem. 1996, 100, 20156–20163.
- (63) Kilambi, K. P.; Gray, J. J. Rapid Calculation of Protein pKa Values Using Rosetta. Biophys. J. 2012, 103, 587–595.
- (64) Yang, A.-S.; Gunner, M. R.; Sampogna, R.; Sharp, K.; Honig, B. On the Calculation of pKas in Proteins. Proteins: Struct., Funct., Genet. 1993, 15, 252–265.
- (65) Thurlkill, R. L.; Grimsley, G. R.; Scholtz, J. M.; Pace, C. N. pK Values of the Ionizable Groups of Proteins. Protein Sci. 2006, 15, 1214–1218.
- (66) Song, Y.; Mao, J.; Gunner, M. R. MCCE2: Improving Protein pKa Calculations with Extensive Side Chain Rotamer Sampling. J. Comput. Chem. 2009, 30, 2231–2247.
- (67) Warwicker, J. Simplified Methods for pKa and Acid pH- Dependent Stability Estimation in Proteins: Removing Dielectric and Counterion Boundaries. Protein Sci. 1999, 8, 418–425.
- (68) Antosiewicz, J.; McCammon, J. A.; Gilson, M. K. The Determinants of pKas in Proteins. Biochemistry 1996, 35, 7819–7833.
- (69) Antosiewicz, J.; McCammon, J. A.; Gilson, M. K. Prediction of Ph-Dependent Properties of Proteins. J. Mol. Biol. 1994, 238, 415–436.
- (70) Kumar, M.; Busch, D. H.; Subramaniam, B.; Thompson, W. H. Organic Acids Tunably Catalyze Carbonic Acid Decomposition. J. Phys. Chem. A 2014, 118, 5020–5028.
- (71) Gu, W.; Frigato, T.; Straatsma, T. P.; Helms, V. Dynamic Protonation Equilibrium of Solvated Acetic Acid. Angew. Chem., Int. Ed. 2007, 46, 2939–2943.

- (72) Scharnagl, C.; Raupp-Kossmann, R.; Fischer, S. F. Molecular Basis for pH Sensitivity and Proton Transfer in Green Fluorescent Protein: Protonation and Conformational Substates from Electrostatic Calculations. Biophys. J. 1999, 77, 1839–1857.
- (73) Lill, M. A.; Helms, V. Proton Shuttle in Green Fluorescent Protein Studied by Dynamic Simulations. Proc. Natl. Acad. Sci. U. S. A. 2002, 99, 2778–2781.