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Carbon Dioxide, Bicarbonate and Carbonate Ions in Aqueous Solutions at Deep Earth Conditions

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We investigate the effect of pressure, temperature and acidity on the composition of water-rich carbon-bearing fluids at thermodynamic conditions that correspond to the Earth’s deep Crust and Upper Mantle. Our first-principles molecular dynamics simulations provide mechanistic insight into the hydration shell of carbon dioxide, bicarbonate and carbonate ions, and on the pathways of the acid/base reactions that convert these carbon species into one another in aqueous solutions. At temperature of 1000 K and higher our simulations can sample the chemical equilibrium of these acid/base reactions, thus allowing us to estimate the chemical composition of diluted carbon dioxide and bicarbonate ions as a function of acidity and thermodynamic conditions. We find that, especially at the highest temperature, the acidity of the solution is essential to determine the stability domain of \( \text{CO}_2 \) vs \( \text{HCO}_3^- \) vs \( \text{CO}_3^{2-} \).

INTRODUCTION

Aqueous electrolyte solutions are an important component of geological fluids in the mid and deep Earth’s crust and in the upper mantle, and water pockets may be present even in the transition zone at depths of more than 600 Km. The chemical balance among different forms of ions in aqueous solution, and their pairing activity determine the formation and dissolution of minerals. At the extreme thermodynamic conditions of the Earth’s deep crust and upper mantle water is supercritical and its different structural and physical properties, e.g. static dielectric constant and ionic conductivity, affect the relative stability and the structure of solvated species.

The Deep Carbon observatory (https://deepcarbon.net/) identified carbon dioxide dissolution and hydration in geological fluids as one of the Earth’s most important reactions, as it affects the global carbon cycle. Yet, the current molecular understanding of carbon dissolved in water-rich fluids at deep crust and upper mantle conditions is still limited in terms of both experiments and theoretical models. On the one hand, in experiments it is possible to probe solutions at extreme conditions by Raman spectroscopy, but the interpretation of such spectra is difficult and controversial. On the other hand, there are very few theoretical studies addressing carbonates in supercritical water by first principles: the former addressing higher pressures and the latter focusing on the occurrence of carbonic acid as opposed in carbon dioxide-rich fluids.

The presence of aqueous electrolyte solutions along the Continental (hot) and Oceanic (cold) geotherms in the crust and upper mantle is significant at temperature up to 1400 K and pressure up to 14 GPa (Figure 1). Aqueous solutions may be also found in the “Wet Peridotite” region, at even higher temperatures, beyond 1600 K. Supercritical water behaves very differently as a solvent than at normal conditions. For example, the static dielectric constant of water varies from \( \varepsilon_w \sim 80 \) at normal conditions to \( \varepsilon_w \sim 15 \) at 1 GPa and 1000 K, and the autoionization constant of water \( (K_w) \) increases by several orders of magnitude as a function of temperature and pressure. The equations of state of minerals dissolved in supercritical water at geochemically relevant conditions are usually predicted using models fitted on experimental data, such as the Helgeson-Kirkham-Flowers (HKF) model or its recent development, the Deep Earth Water (DEW) model. Both HKF and DEW models provide an estimate of the free energy of solvation of ions \( (\Delta G_s) \) as a function of the dielectric constant of water \( \varepsilon_w \) and of an electrostatic Born parameter.
specific to each ion, such that:

$$\Delta G = -\frac{N_A z^2 e^2}{8\pi\epsilon_0 \epsilon_r} \left( 1 - \frac{1}{\epsilon_w} \right),$$

(1)

where $z$ is the ion charge, $e$ the electron charge, $r_0$ effective radius of the ion, $\epsilon_0$ the vacuum permittivity and $N_A$ is the Avogadro number. While the DEW model is considered predictive and has wide use in the geochemistry community, its parameters are fitted to experiments at mild conditions, and the validity of the extrapolations to extreme conditions lacks compelling verification. In fact, experimental data of $\epsilon_w$ for water at elevated temperatures are currently limited to 550°C and 0.5 GPa.\textsuperscript{13,15}

First principles molecular dynamics (FPMD) simulations, based on density functional theory (DFT), proved valuable to make up for the experimental gap providing a parameter-free estimate of the dielectric constant of supercritical water for pressure up to 10 GPa.\textsuperscript{10} This achievement suggests that a systematic use of FPMD will allow geochemists to extend aqueous geochemical models to a broader range of thermodynamic conditions. In addition, recent refinements of the standard Born models of solvation\textsuperscript{16,17} call for a more accurate description of the solvent polarization in the presence of ions, which can be provided by FPMD simulations. Ultimately, molecular simulations can provide a direct prediction of the stability domain of aqueous electrolytes at geologically relevant thermodynamic conditions. Especially in the cases of anomalous behavior, atomistic insight from FPMD simulations would rationalize experimental observations, made possible by the development of techniques to probe liquids at extreme conditions, such as high-pressure Raman\textsuperscript{1} and NMR.\textsuperscript{18}

FPMD describes accurately the properties of water and ice at high pressure\textsuperscript{10,11,19,22} and it has been employed in the past to study the solvation structure of various aqueous species\textsuperscript{22}, from monoatomic anions and cations\textsuperscript{24,26} to complex molecules of biological importance.\textsuperscript{27} FPMD is particularly useful to study reactions in solutions and at conditions at the limit of experimental possibilities, not only at high pressure and temperature but also in the presence of strong static electric field.\textsuperscript{28,29} Specifically relevant to the present study, FPMD has already been used to resolve the solvation shell of carbon dioxide and the conversion of CO$_2$ into bicarbonate in high-pH environment.\textsuperscript{30,31} showing that this computational technique is able to quantitatively predict free energies of reaction as well as reaction barriers, provided that suitable correction terms are calculated from higher-level gas phase quantum chemical calculations.\textsuperscript{32} Using parameter-free FPMD simulations a pioneering work by Pan and Galli\textsuperscript{6} showed that, contrary to conventional models, at Upper Mantle conditions (1000 K and 11 GPa) CO$_2$ is not the major carbon species, but it transforms into CO$_3^{2-}$ and HCO$_3^-$. Intermediate pressures, corresponding to the boundary between the Lithosphere and the Upper Mantle were studied to unravel the abundant occurrence of H$_2$CO$_3$. However, a systematic study of the carbon species in water-rich fluids at these conditions is still lacking, and the effects of the acidity of the solutions remain largely unexplored. In this work, we perform extensive FPMD simulations to investigate the properties of dissolved carbon in geological fluids at the deep crust and upper Mantle conditions as a function of temperature, pressure and acidity (Figure 1). To set different pH conditions we performed simulations of ~1 molar solutions with different initial conditions, namely 2Na$^+$+CO$_3^{2-}$, Na$^+$+HCO$_3^-$ and CO$_2$. Low-temperature simulations shed light on the solvation shell of the three carbon species as a function of pressure. At temperatures at and above 1000 K the carbon species are highly reactive and FPMD simulations allow us to map the average composition of the solutions at both acidic and basic conditions as a function of pressure and temperature. Furthermore, we obtain direct insight into the hydration and dehydration reaction pathways of carbon solutes, without the bias of either a priori assumptions or the choice of specific reaction coordinates.

**METHODS AND MODELS**

FPMD simulations were carried out using the Quickstep approach, implemented in the CP2k code.\textsuperscript{33} We employ the Perdew-Burke-Ernzerhof (PBE) generalized gradient approximation (GGA) for the DFT exchange and correlation functional.\textsuperscript{34} Valence Kohn-Sham orbitals are represented on a Gaussian triple-$\zeta$ basis set with polarization, optimized for molecular systems in gas and condensed phase.\textsuperscript{35} and core states are treated implicitly using Geodecker-Teter-Hutter pseudopotentials.\textsuperscript{36} Semilocal GGA functionals do not provide a good description of dispersion forces, and it is well established that including van der Waals corrections improves the description of water at ambient temperature.\textsuperscript{37,38} However, the effect of vdW corrections, and manybody effects in general, in supercritical water is less drastic and it has not been studied in detail for the PBE functional.\textsuperscript{38,39} In fact, the PBE functional has been extensively used to study the properties of supercritical water\textsuperscript{10,11,21} and of hydrocarbons at deep Earth conditions as well.\textsuperscript{40} Although it is known that GGA may not be accurate in the description of solvated doubly-charged anions because of the charge delocalization error,\textsuperscript{41} it was recently shown that PBE performs better under extreme conditions than under ambient conditions in terms of equation of state of water, as well as in predicting static and electronic dielectric constants.\textsuperscript{10,11} Pan and Galli also showed that PBE provides very similar results for the structure and the speciation of CO$_2$ and CO$_3^{2-}$ in supercritical water.\textsuperscript{6,7}
When carbon species are dissolved in water, the following hydration/dehydration reactions and their equilibrium constants determine the relative composition of the solution:

\[
\begin{align*}
\text{CO}_2 + 2\text{H}_2\text{O} & \rightleftharpoons \text{HCO}_3^- + \text{H}_3\text{O}^+ \\
\text{HCO}_3^- + \text{H}_2\text{O} & \rightleftharpoons \text{CO}_3^{2-} + \text{H}_3\text{O}^+ \\
\text{HCO}_3^- (\text{aq}) & \rightleftharpoons \text{CO}_2 + \text{OH}^- \\
\text{CO}_3^{2-} + \text{H}_2\text{O} & \rightleftharpoons \text{HCO}_3^- + \text{OH}^- 
\end{align*}
\] (2)

To probe these reactions for diluted solutions, we prepared cubic simulation cells containing 63 water molecules and one solvated species (CO\(_2\), HCO\(_3^-\), or CO\(_3^{2-}\)) with edge length \(L = 12.37\) \(\text{Å}\) with periodic boundary conditions. Charge neutrality is achieved by compensating the net negative charge of the anions by a corresponding number of sodium cations. This initial condition corresponds to total densities between 1.03 and 1.09 \(\text{g/cm}^3\). To perform simulations at higher pressure, we reduced the box linear size to 11.76 and 11.14 \(\text{Å}\). These systems correspond to solution with different molar concentrations: specifically 0.88 \(M\), 1.02 \(M\), and 1.20 \(M\) respectively. For each system, we performed a set of MD simulations at \(T = 500, 1000,\) and 1600 K. The MD equations of motion are integrated with the velocity Verlet integrator with a timestep of 0.25 fs. To perform simulations in the constant temperature constant volume canonical ensemble (NVT) we used the stochastic velocity rescaling thermostat with a coupling constant \(\tau = 0.5\) ps. Each model was first equilibrated for 5 ps at the target temperature, and the analysis was performed on data obtained from 50 ps-long production runs. The overall project amounted to 27 runs for a total simulation time of \(\sim 1.5\) ns.

If the reactions in Eq. 2 can be considered at equilibrium over time scales of several tens of picoseconds, we can exploit direct FPMD simulations to estimate the chemical balance of the solvated species at given thermodynamic conditions. Since the products of these reactions are hydronium and hydroxide ions, from sufficiently long simulations we can also obtain the equilibrium concentrations of these species, which allow us to roughly estimate the pH and pOH of the solution. In principle, to fix the pH, one would have to perform grand canonical (GC) ensemble simulations with a fixed chemical potential for either H\(_3\)O\(^+\) or OH\(^-\). However, the
standard approach to GC simulations, i.e. Widom particle insertion, is extremely costly and practically unfeasible for condensed phases, especially with first-principles simulations. Yet, here we argue that it can be avoided by fixing the conjugate thermodynamic variable of the chemical potential, i.e. the number of particles in the system. This idea works in the same way as fixing the volume (or the density) of a system determines the equilibrium pressure, and fixing the energy determines the equilibrium temperature. Therefore, in reactive simulations we can achieve the goal of mapping the relative abundance of the solute species as a function of temperature, pressure and acidity. Using the concentrations of H$_3$O$^+$ and OH$^−$ averaged over a production run, and the experimental value of the autoionization constant of supercritical water, we estimate the acidity of the solution as in the textbook case of weak polyprotic acids and bases. We define the acidity/basicity of the solution as the difference between pH and pOH (see Supporting Information). However, given the small size of the simulation cell, the high solute concentration and the limited accessible timescale, this estimate is hampered by large uncertainties. More accurate estimates of acidity and solutes stability may be achieved either simulating larger systems, or computing directly redox potentials by free energy calculations, for example exploiting the Born-Haber cycle, which is, however, beyond the scope of this work.

RESULTS AND DISCUSSION

Solvation of Carbon Species at 500 K

We carried out a first set of low-temperature simulations to characterize the solvation shell of the different carbon species as a function of pressure at $T = 500$ K. As expected, these systems show no reactivity on the timescale of our production runs, thus no information about the predominant species can be extracted from these simulations. Geochemical models and previous experiments suggest that at relatively low temperature and pressure, the major dissolved carbon species is CO$_2$(aq). Nevertheless, the metastability of all three carbon solutes allows us to shed light into the structure of their hydration shell, which has an essential role in the formation of minerals, such as calcite or dolomite, at mild conditions, and for carbon geosequestration. Further studies using enhanced sampling methods would be required to estimate the relative stability of dissolved carbon species and the composition of solutions in the colder layers of the Earth’s crust.

Figure 2(a,d,g) shows that the first solvation shell of CO$_2$ is rather unstructured, giving rise to a broad first peak in the carbon-oxygen radial distribution function (RDF) ($g_{CO}(r)$), as well as in the oxygen-hydrogen RDF ($g_{O-H}(r)$, where $O_H$ indicates the oxygen atom bonded to carbon). Also the oxygen-oxygen RDF exhibits a broad first peaks and weak structuring at low pressure, while the peak gets better defined and shifts to shorter distance at higher pressure (Fig. 2(i)). The reason is that CO$_2$ is apolar and does not form hydrogen bonds with the surrounding water molecules. This behavior, observed in both experiments and simulations at ambient conditions, is retained at 500 K for pressures ranging from 1.1 to 5.1 GPa. The effect of increasing pressure is to shift the first broad peak of $g_{CO}(r)$ toward smaller distances, indicating a spatial contraction of the first solvation shell, but no significant change in the number of nearest neighbors, which is defined as the integral of the first peak of the $g_{CO}(r)$.

The carbon-oxygen RDFs of anionic species, HCO$_3^−$, and CO$_3^{2−}$, (Figure 2(c,e)) exhibit a sharper first peak that shifts toward smaller distance as a function of pressure. Both anions form hydrogen bonds with water, as it appears from the well-defined structure of the $g_{OC OH}(r)$ in Fig. 2(h,i). The OH group of HCO$_3^−$ donates one H-bond, and each oxygen accepts on average 2.2 H-bonds from the neighboring water molecules. This number is not sensitive to pressure, but the overall hydrogen-bonding structure of the solvation shell undergoes significant changes when the pressure is increased from 2.7 to 6.2 GPa. At high pressure a larger number of water molecules from the second solvation shell enters the radius of the first shell of hydrogen-bonded molecules, disrupting the local order of the hydrogen-bonded network. This effect can be observed also for CO$_3^{2−}$: the double-charged anion accepts on average 2.8 H-bonds per oxygen, leading to a total number of ~9 neighbors, in agreement with former FPMD simulations, but smaller than that estimated at ambient conditions with empirical potentials. Increasing pressure causes weakening of hydrogen bonding, as showcased by the lower peak of the $g_{OH}(r)$ (Figure 2(h,i)), but an overall increase of the neighboring water molecules results in a more disordered and dynamic solvation shell.

Solvation of Carbon Species at 1000 and 1600 K

High temperature and high pressure enhance the rate of autoionization in water, thus making the solvent increasingly reactive. At $T = 1000$ K and pressures above 11 GPa, water rapidly dissociates and recombines

1 The number of hydrogen bonds has been calculated according to the same geometric criteria used in Ref. 54: two molecules are considered hydrogen bonded if the oxygen-oxygen distance is lower than 3.3 Å, the oxygen-(donor)hydrogen distance is lower than 2.4 Å and the H-O···O angle is smaller than 30°.
through a bimolecular mechanism that produces short-lived hydronium-hydroxide ion pairs, and it eventually turns into an ionic fluid. Conversely, at lower pressure along the Hugoniot compression curve, nearly no autoionization was observed in pure water over the typical FPMD simulation timescale of few tens ps. In our simulations at $T = 1000$ K we also do not see spontaneous water autoionization up to $P \sim 4$ GPa, whereas at higher pressure transient $H_2O^+/OH^-$ pairs occur spontaneously. Nevertheless, in all the runs, except for those starting with $CO_2$ at $P = 2$ and 3.9 GPa, the enhanced reactivity of supercritical water engenders fast interconversion among carbon dioxide, bicarbonate and carbonate ions with the consequent release of either hydronium or hydroxide ions, according to the reactions in Eq. 2.

![Figure 3](image-url)

**Figure 3.** Mole percents of $CO_2$ (green), $CO_3^{2-}$ (orange), and $HCO_3^-$ (blue) calculated as functions of time at $T = 1000$K with $CO_2$ (a), $CO_3^{2-}$ (b), and $HCO_3^-$ (c) as the starting solvated species. Here the mole percent relative to $CO_2$ actually considers also the presence of $H_2CO_3$ because carbonic acid appears in amounts comparable to $CO_2$. The reaction $CO_2 + H_2O \rightleftharpoons H_2CO_3$ does not alter the pH of the solution.

The spontaneous hydration/dehydration reactions among $CO_2$, $HCO_3^-$ and $CO_3^{2-}$ in unbiased FPMD simulations allows us to explore the reaction mechanisms at the atomic scale. Figure 4 shows the representative molecular pathways of the water-assisted transformation of $CO_2$ into $HCO_3^-$, $HCO_3^-$ into $CO_3^{2-}$, and of $HCO_3^-$ into $CO_3^{2-}$. The formation of bicarbonate from carbon dioxide neither require the presence of a free hydroxide ion nor has $H_2CO_3$ as an intermediate. The reaction occurs via the nucleophilic attack of a water molecule to $CO_2$. The oxygen atom of the $H_2O$ molecule attacks the carbon of $CO_2$, while one of its protons is released into the solution, initially forming a $H_2O^+$ ion with a neighboring water (Fig. 4a). Eventually the proton diffuses in solution via Grotthuss mechanism. Also the reverse reaction, from $HCO_3^-$ to $CO_2$, proceeds along a water-mediated pathway, and we never observe direct breaking of the C-OH bond. The OH group of $HCO_3^-$ accepts a proton from a neighboring water molecule and detaches from the carbon atom as a water molecule (Figure 4b). The $H_2O$ that donated the proton diffuses in solution as a hydroxide ion. Both the conversion of $HCO_3^-$ into $CO_3^{2-}$ (Figure 4c) and the reverse reaction (not shown) occur via direct proton exchange with a neighboring water molecule. It is important to stress that all these reactions do not require the direct interaction between the carbon species and either $OH^-$ or $H_3O^+$. Identifying these mechanistic reaction pathways is an essential step to eventually implement enhanced sampling methods to refine the calculation of reaction free energies.

Besides the reactions given in Eq. 2 we observe that the configuration of $HCO_3^-$ is highly dynamic, in the sense that the OH site swaps from one oxygen to another. This mechanism can be tracked by monitoring the length of the C-O bond length ($r_{CO}$) for each oxygen atom in $HCO_3^-$. The average $r_{CO}$ for the protonated oxygen atom is larger ($\sim 1.4$ Å) than the other two C-O bonds ($\sim 1.28$ Å). Figure 5 shows that over a trajectory of a solution mostly containing $HCO_3^-$ the "long" C-O bond evolves from one site to another through rapid transitions. Figure 5 shows how this swapping mecha-
FIG. 4. Atomistic representation of the mechanism of the reactions that transforms CO$_2$ into HCO$_3^-$ (a), HCO$_3^-$ into CO$_2$ (b) and HCO$_3^-$ into CO$_2^-$ (c). The first and the third reactions are mediated by water in the first solvation shell and release one hydronium ion in solution, while the second one produce one hydroxide ion.

The mechanism happens: a water molecule hydrogen bonded to an oxygen of the bicarbonate donates a proton to the ion forming transient neutral carbonic acid (central panel in Fig. 4). The abundance of H$_2$CO$_3$ at these conditions is actually relevant and comparable to that of CO$_2$ in agreement with extensive simulations recently performed by Stolte and Pan. In the following analysis we group the concentrations of CO$_2$ and H$_2$CO$_3$ into a single neutral contribution, as the relative abundance of one of these two neutral species with respect to the other does not affect the acidity of the solution.

At higher temperatures ($T=1600$ K) the solutions exhibit further increased reactivity than those at 1000 K, with frequent transitions from one species to another, thus providing more accurate statistic on the equilibrium composition of the fluid. In these high temperature runs the solvent remains mostly molecular, and the main transition mechanisms are the same as those described in detail for the simulations, involving neutral water molecules as facilitators of the reactions. However, in the high pressure runs ($P=10$ GPa) we observed a more significant ionic character of supercritical water, possibly enhanced by the presence of bicarbonate and carbonate ions.

Comparing the carbon-oxygen radial distribution functions at different pressure and temperature (Figure S2 in the Supporting Information), we observe that the effect of the pressure at high temperature is similar to that discussed for the runs at 500 K, and the structure of the first solvation shell of CO$_2$, HCO$_3^-$ and CO$_2^-$ does not change significantly with temperature.

FIG. 5. (a) Carbon-oxygen distances calculated over the productions runs starting with solvated CO$_2$ at P=6.9 GPa and T=1000 K. The predominant species at these conditions is HCO$_3^-$ but the position of the OH group frequently changes from one oxygen site to another through a reaction mediated by the nearest neighboring molecules, with the occurrence of H$_2$CO$_3$(b). In the presence of HCO$_3^-$ the average $r_{CO}$ for the protonated oxygen atom is larger ($\sim 1.4$ Å) than the other two C-O bonds ($\sim 1.28$ Å), while in the case of H$_2$CO$_3$, the average $r_{CO}$ for the two protonated oxygen atoms is slightly reduced ($\sim 1.35$ Å). Accordingly, we used black arrows to highlight the three different steps for such swapping mechanism. Blue arrows indicate the hydrogen atoms involved in the process.

Composition of Carbon-Bearing Fluids as a Function of Pressure and Acidity

The relatively high frequency of reactive events allows us to consider the chemical reactions at equilibrium and to estimate the equilibrium molar fraction of solutions at given thermodynamic conditions as the average molar fraction of over a whole production run. It is important to note that, as one can infer from Figure 5, the average compositions of the solution at similar temperature and pressure may differ significantly, depending on the starting solute species. The reason is that the hydronium/hydroxide ions produced in the acid/base reaction among different solutes change the pH of the solutions. The latter can be estimated in the same way as the molar fraction, by averaging the concentration of excess hydronium/hydroxide over a trajectory. Simulations starting with CO$_2$ result in acidic solutions, as the transforma-
tion into HCO$_3^-$ produces hydronium, unless no reaction happens and CO$_2$ remains the only solvated species for the whole duration of the run. According to the same argument, systems starting with CO$_3^{2-}$ can only probe basic conditions, whereas HCO$_3^-$ may transform producing either H$_2$O$^+$ or OH$^-$, thus engendering either acidic or basic equilibrium conditions.

The autoionization constant of supercritical water $K_w$ is extremely sensitive to temperature and pressure and it increases monotonically with either. In the $(P, T)$ range explored in this work it ranges from $K_w = 10^{-7}$ at $P = 1$ GPa and $T = 1000$ K to $K_w \geq 10^{-1}$ at $P = 10$ GPa and $T = 1600$ K. Hence, we cannot use pH alone to define acidity or basicity of a supercritical solution, as the condition of neutrality changes with the change of $K_w$. We then classify the acidity/basicity of the solution in terms of the difference between pOH and pH, which corresponds to the log$_{10}$ of ratio between the equilibrium concentrations of H$_3$O$^+$ and OH$^-$(f = pH – pOH = –log$_{10}$[H$_3$O$^+$]/[OH$^-$.]), computed from the reactive FPMD simulations. pH – pOH = 0 defines neutrality at any temperature and pressure, f = pH – pOH < 0 represents acid conditions and f = pH – pOH > 0 basic conditions. A detailed explanation about the calculation of such ratio is reported in Supplementary Information$^1$.

\[ \text{FIG. 6. Composition of carbon species in aqueous solution as a function of pH and pressure, for } T = 1000\text{K (a) and } T = 1600\text{K (b).} \]

\[ \text{In square brackets is reported the initial carbon species. The top panel contains also the first results at extreme conditions obtained by FPMD simulations.} \]

\[ \text{The mole percent, the pressure and the } f \text{ values can be found in the Supplementary Information in Table S1 and S2.} \]

\[ ^2 \text{For reference, in water at ambient conditions CO}_2\text{ is stable at acidic conditions, for pH<6.5, HCO}_3^-\text{ for pH from 6.5 to 10.5 and CO}_3^{2-}\text{ at basic conditions pH>10.5.} \]
substantial amount of free ions in solution.

CONCLUSIONS

In conclusion, we carried out an extensive set of FPMD simulations of a water-rich carbon-bearing fluid at various temperatures and pressures corresponding to the thermodynamic conditions of the Earth’s deep crust and upper mantle ($2 < P < 10$ GPa and $500 < T < 1600$ K). We systematically probed the effect of preparing the systems with carbon species as solutes, namely CO$_2$, HCO$_3^-$, and CO$_3^{2-}$ at $\sim$ 1 M molar concentration, and we found that in highly reactive conditions, averaging over sufficiently long runs gives an unbiased estimate not only of the composition of the solution, but also of its acidity. This observation allows us to map the composition of geological fluids as a function of temperature, pressure and acidity: this information is essential for geochemical models, as the stability of different forms of carbon solutes impacts the growth and dissolution of calcium, magnesium and iron carbonates, including calcite, aragonite, dolomite, magnesite and siderite, which are present in significant amounts both in the crust and in the mantle.\[5, 59\]

Nevertheless, we find that the equilibrium composition of the solutions depends critically on the initial conditions, which, in turn determine the equilibrium content of H$_3$O$^+$ and OH$^-$ ions. This result highlights the importance of considering acidity, at the same level as temperature and pressure, to predict the composition of geological fluids at deep Earth conditions.

The HKF and DEW models can account qualitatively for the observed trends in the relative stability of different carbon species as a function of pressure and temperature. However, at high temperature (1600 K) and pressure ($\sim 10$ GPa), as water approaches the transition from molecular to ionic, the increasing abundance of free H$_3$O$^+$ and OH$^-$ ions influences the reactivity of the system and favors the formation of CO$_3^{2-}$, at mildly basic conditions.

Furthermore, FPMD simulations also provide insight into the atomistic mechanisms of the protonation/deprotonation reactions of different carbon species. These reaction pathways are non-trivial and mostly involve neutral water molecules in the first solvation shell of the carbon species, and proton-hopping. For example, we find that CO$_2$ transforms into HCO$_3^-$ via the nucleophilic attack of a neutral water molecule and the release of hydronium. Conversely the reverse reaction proceeds via the electrophilic attack of the OH group and the concurrent release of a water molecule and a hydroxide ion.

CONFLICTS OF INTEREST

The authors declare that they have no competing interests.

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[1] A. B. Thompson, Nature, 1992, 358, 295–302.
[2] O. Tschauner, S. Huang, E. Greenberg, V. B. Prakapenka, C. Ma, G. R. Rossman, A. H. Shen, D. Zhang, M. Newville, A. Lanziriotti and K. Tait, Science, 2018, 359, 1136–1139.
[3] C. A. Eckert, B. L. Knutson and P. G. Debenedetti, Nature, 1996, 383, 313–318.
[4] R. S. Hawke, K. Syassen and W. B. Holzapfel, Rev. Sci. Instrum., 1974, 45, 1598–1601.
[5] C. E. Manning, Annu. Rev. Earth Planet. Sci., 2018, 46, 67–97.
[6] D. Pan and G. Galli, Sci. Adv., 2016, 2, e1601278.
[7] N. Stolte and D. Pan, J Phys Chem Lett, 2019, 10, 5135–5141.
[8] W. B. Holzapfel, The Journal of Chemical Physics, 1969, 50, 4424–4428.
[9] H. Weingärtner and E. U. Franck, Angew. Chem. Int. Edit., 2005, 44, 2672–2692.
[10] D. Pan, L. Spanu, B. Harrison, D. A. Sverjensky and G. Galli, P. Natl. Acad. Sci. USA, 2013, 110, 6646–6650.
[11] V. Rozsa, D. Pan, F. Giberti and G. Galli, P. Natl. Acad. Sci. USA, 2018, 115, 6952–6957.
[12] H. C. Helgeson, D. H. Kirkham and G. C. Flowers, Am. J. Sci., 1981, 281, 1249–1516.
[13] D. A. Sverjensky, B. Harrison and D. Azzolini, Geochim. et Cosmochim. Ac., 2014, 129, 125–145.
[14] N. C. Caciagli and C. E. Manning, Contrib. Mineral. Petr., 2003, 146, 275–285.
[15] I. Martinez, C. Sanchez-Valle, I. Daniel and B. Reynard, Chem. Geol., 2004, 207, 47–58.
[16] I. Nakamura, J. Phys. Chem. B, 2018, 122, 6064–6071.
[17] X. Duan and I. Nakamura, Soft Matter, 2015, 11, 3566–3571.
[18] B. G. Pautler, C. A. Colla, R. L. Johnson, P. Klavins, S. J. Harley, C. A. Ohlin, D. A. Sverjensky, J. H. Walton and W. H. Casey, Angew. Chem. Int. Edit., 2014, 53, 9788–9791.
[19] E. Schwegler, G. Galli and F. Gygi, Phys. Rev. Lett., 2000, 84, 2429–2432.
[20] E. Schwegler, G. Galli, F. Gygi and R. Q. Hood, Phys. Rev. Lett., 2001, 87, 986–4.
[21] E. Schwegler, M. Sharma, F. Gygi and G. Galli, P. Natl. Acad. Sci. USA, 2008, 105, 14779–14783.
[22] N. Goldman, E. J. Reed, I. F. W. Kuo, L. E. Fried, C. J. Mundy and A. Curioni, J. Chem. Phys., 2009, 130, 124517–7.
[23] M. E. Tuckerman, D. Marx and M. Parrinello, Nature, 2002, 417, 925–929.
[24] T. K. Todorova, I. Infante, L. Gagliardi and J. M. Dyke, *J. Phys. Chem. A*, 2008, **112**, 7825–7830.
[25] A. P. Gaiduk and G. Galli, *J. Phys. Chem. Lett.*, 2017, **8**, 1496–1502.
[26] T. A. Pham, S. M. Mortuza, B. C. Wood, E. Y. Lau, T. Ogitsu, S. F. Buchsbaum, Z. S. Siwy, F. Fornasiero and E. Schwegler, *J. Phys. Chem. C*, 2016, **120**, 7332–7338.
[27] M. P. Gaigeot and M. Sprik, *J. Phys. Chem. B*, 2003, **107**, 10344–10358.
[28] F. Pietrucci and A. M. Saitta, *Proc Natl Acad Sci USA*, 2015, **112**, 15034–15038.
[29] G. Cassone, J. Sponer, S. Trusso and F. Saija, *Phys. Chem. Chem. Phys.*, 2019, **21**, 21205–21212.
[30] K. Leung, I. M. B. Nielsen and I. Kurtz, *J. Phys. Chem. B*, 2007, **111**, 4453–4459.
[31] S. R. Zukowski, P. D. Mitev, K. Hermansson and D. Ben-Amotz, *J. Phys. Chem. Lett.*, 2017, **8**, 2971–2975.
[32] E. Grifoni, G. Piccini and M. Parrinello, *P. Natl. Acad. Sci. USA*, 2019, **116**, 4054–4057.
[33] J. Vandevondele, M. Krack, F. Mohamed, M. Parrinello, T. Chassaing and J. Hutter, *Comput. Phys. Commun.*, 2005, **167**, 103–128.
[34] J. P. Perdew, K. Burke and M. Ernzerhof, *Phys. Rev. Lett.*, 1996, **77**, 3865–3868.
[35] J. Vandevondele and J. Hutter, *J. of Chem. Phys.*, 2007, **127**, 114105.
[36] S. Goedecker and M. Teter, *Phys. Rev. B*, 1996, **54**, 1703–1710.
[37] I.-C. Lin, A. P. Seitsonen, I. Tavernelli and U. Rotllisberger, *J. Chem. Theory Comput.*, 2012, **8**, 3902–3910.
[38] R. Jonchiere, A. P. Seitsonen, G. Ferlat, A. M. Saitta and R. Vuilleumier, *J. Chem. Phys.*, 2011, **135**, 154503–11.
[39] P. Schienbein and D. Marx, *J. Phys. Chem. B*, 2017, **122**, 3318–3329.
[40] L. Spanu, D. Donadio, D. Hohl, E. Schwegler and G. Galli, *P. Natl. Acad. Sci. USA*, 2011, **108**, 6848–6846.
[41] A. J. Cohen, P. Mori-Sánchez and W. Yang, *Science*, 2008, **321**, 792–794.
[42] G. Bussi, D. Donadio and M. Parrinello, *J. Chem. Phys.*, 2007, **126**, 014101.
[43] B. Widom, *J. Chem. Phys.*, 1963, **39**, 2808–2812.
[44] D. Frenkel and B. Smit, *Understanding Molecular Simulation*, Academic Press, Inc., Orlando, FL, USA, 2nd edn, 2001.
[45] S. D. Hamann and M. Linton, *Trans. Faraday Soc.*, 1969, **65**, 2186–11.
[46] J. Cheng, X. Liu, J. Vandevondele, M. Sulpizi and M. Sprik, *Accounts Chem Res*, 2014, **47**, 3522–3529.
[47] Z. Duan and Z. Zhang, *Geochim. et Cosmochim. Ac.*, 2006, **70**, 2311–2324.
[48] C. Zhang and Z. Duan, *Geochim. et Cosmochim. Ac.*, 2009, **73**, 2089–2102.
[49] J.-M. Huizenga, *Lithos*, 2001, **55**, 101–114.
[50] P. Raiteri and J. D. Gale, *J. Am. Chem. Soc.*, 2010, **132**, 17623–17634.
[51] M. De La Pierre, P. Raiteri, A. G. Stack and J. D. Gale, *Angew. Chem. Int. Edit.*, 2017, **129**, 8584–8587.
[52] Y. Liang, S. Tsuji, J. Jia, T. Tsuji and T. Matsuoka, *Acc. Chem. Res.*, 2017, **50**, 1530–1540.
[53] G. A. Gallet, F. Pietrucci and W. Andreoni, *J. Chem. Theo. Comput.*, 2012, **8**, 4029–4039.
[54] A. Luzar and D. Chandler, *Nature*, 1996, **379**, 55–57.
[55] F. Bruneval, D. Donadio and M. Parrinello, *J. Phys. Chem. B*, 2007, **111**, 12219–12227.
[56] J. D. Gale, P. Raiteri and A. C. T. van Duin, *Phys. Chem. Chem. Phys.*, 2011, **13**, 16666.
[57] P. Raiteri, R. Demichelis and J. D. Gale, *J. Phys. Chem. C*, 2015, **119**, 24447–24458.
[58] A. F. Goncharov, N. Goldman, L. E. Fried, J. C. Crowhurst, I.-F. W. Kuo, C. J. Mundy and J. M. Zaug, *Phys. Rev. Lett.*, 2005, **94**, 85–4.
[59] G. W. Berg, *Nature*, 1986, **324**, 50–51.