Theoretical and Experimental Studies of the Dechlorination Mechanism of Carbon Tetrachloride on a Vivianite Ferrous Phosphate Surface

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Supporting Information

ABSTRACT: Chlorinated organics are the principal and most frequently found contaminants in soil and groundwater, generating significant environmental problems. Over the past several decades, Fe-containing minerals naturally occurring in aquatic and terrestrial environments have been used as natural electron donors, which can effectively dechlorinate a variety of chlorinated organics. However, a full understanding of the reaction mechanism of the dechlorination pathway cannot be obtained by experimental investigations alone, due to the immeasurability of chemical species formed over a short reaction time. In this report, we describe experiments and density functional theory (DFT) calculations carried out to investigate the complex reduction pathway of carbon tetrachloride (CT) on a vivianite (FeII3(PO4)2·8H2O) surface. Our results indicate that chloroform (HCCl3) and formate are the primary transformation products. The experimental results reveal that the reduction kinetics of CCl4 can be dramatically accelerated as the pH is increased from 3 to 11. On the basis of the DFT calculations, we found that HCCl3 can be formed by *CCl3 and :CCl3−* on a deprotonated vivianite surface (an adsorbate on vivianite is denoted using an asterisk). In addition, :CCl3−* can be nonreductively dechlorinated to form :CCl2−* followed by sequential nucleophilic attack by OH−*, resulting in the formation of :CCl(OH)−* and :C(OH)2−*, which are responsible for production of CO and formate, respectively. The results obtained from this study can facilitate the modeling of systems of other halogenated species and minerals, which will provide fundamental insight into their corresponding reaction mechanisms.

I. INTRODUCTION

Chlorinated organics are widely used for industrial purposes1 and are commonly found as principle contaminants in soil and groundwater.2−4 It is well-known that most chlorinated organics are highly toxic and are persistent and widely spread in natural environments due to their limited water solubility and downward migration.5 Because of their harmful effects, maximum concentrations of chlorinated organics in water used for various purposes have been set by the World Health Organization; however, they have still been detected in a third of US hazardous waste sites owing to past inappropriate disposal practices.5 As a result, remediation technologies that can treat a variety of chlorinated organics have been developed extensively over the past few decades. Among these, soil and soil minerals, particularly Fe-containing minerals existing in natural environments, have attracted attention as natural attenuators, which are utilized as efficient electron donors under iron-reducing environments.4,6,7 Indeed, a variety of Fe-containing minerals have shown a remarkable reduction capacity toward various chlorinated organics. For example, Lee and Batchelor reported the reductive degradation of chlorinated ethylenes (tetrachloroethylene (PCE), trichloroethylene (TCE), cis-dichloroethylene, and vinyl chloride) by pyrite, magnetite, and green rust.8 Additionally, Amir and Lee9 demonstrated an enhanced reactivity of nanomackinwite for PCE, and Choi et al.10 reported the reductive dechlorination of...
carbon tetrachloride (CT) and 1,1,1-trichloroethane by iron sulfide.

Recently, the monoclinic octa-hydrated ferrous phosphate vivianite (Fe\(^{II}\)(PO\(_4\))\(_2\)·8H\(_2\)O) has been rehighlighted as a potential reactive Fe-containing mineral due to its adsorption capacity, leading to the immobilization of cobalt, nickel, strontium, and arsenic, as well as its reduction capacity, leading to the reduction of oxidized uranium and CCl\(_4\). In our previous study, the transformation of CCl\(_4\) into CO and formate by vivianite and surface-stabilized carbene complexes, respectively, was suggested as the main reaction process. However, the difficult experimental detection of the intermediates has always prevented a full understanding of the dechlorination pathway due to the extremely short reaction time, low concentration of products, and immearusability of chemical species. Therefore, a novel approach should be carried out to investigate the complex surface reactions of CCl\(_4\) and its intermediates with a vivianite surface.

Now regarded as an indispensable supplement to experiments, density functional theory (DFT) calculations are being used in a wide range of applications to aid in the interpretation of experimental observations and in the investigation of the full atomistic details of reaction pathways based on accurate quantum mechanical energetics. The most beneficial feature of DFT studies is that they provide relative energies of intermediate states based on first-principles quantum mechanics, which can be unbiasedly utilized to determine the feasibilities of possible reaction pathways. Although several previous studies have investigated Fe-containing minerals using DFT calculations to understand the bulk and surface structures of hematite, magnetite, siderite, and vivianite. Also, electrocatalytic activity and water oxidation on Fe-containing minerals were studied using DFT calculation. However, to the best of our knowledge there have been no DFT studies on the catalytic activity between vivianite and chlorinated compounds.

In this paper, we describe a combined experimental and DFT study carried out to examine the reductive dechlorination mechanisms of CCl\(_4\) on a vivianite surface. We experimentally studied the reductive dechlorination of CCl\(_4\) by vivianite, the effect of pH on the CCl\(_4\) degradation, and the intermediate and final product distributions during the dechlorination process. Using DFT, we investigated the energetics of possible intermediate states and of the dominant reaction mechanism, which explain the experimentally observed results.

II. METHODS

**Chemicals.** The chlorinated organics used in this study, CCl\(_4\) (≥99.5%), chloroform (HCCl\(_3\)) (≥99.5%), dichloromethane (CH\(_2\)Cl\(_2\)) (≥99.5%), and methyl chloride (CH\(_3\)Cl) (≥99.9%), were all purchased from Sigma-Aldrich. Ammonium ferrous sulfate hexahydrate (99%, Sigma-Aldrich), sodium acetate anhydrous (>99%, Aldrich), and sodium phosphate dibasic anhydrous (99%, Junsei) were used for the synthesis of vivianite. Hexane (99.9%, Merck) and methanol (≥99.9%, Sigma-Aldrich) were used as an extractant for the chlorinated organics and as a solvent for the preparation of stock solutions, respectively. A SCOTTY gas mixture (1.01% of methane (CH\(_4\)), carbon monoxide (CO), carbon dioxide (CO\(_2\)), ethane, acetylene, and ethylene in N\(_2\) (Supelco) was used to analyze the nonchlorinated transformation products. Formic acid (≥95%, Sigma-Aldrich) and sodium chloride (≥99%, Sigma-Aldrich) were used for the analysis of formate and chloride ion, respectively.

Decarated deionized water (DDW) was prepared by purging ultrapure water (18 M\(_2\)cm) with N\(_2\) for 4 h, and was subsequently stored under anaerobic conditions. The organic buffers used in this study were citric acid monohydrate and trisodium citrate dihydrate (for pH 3 and 5), 1,4-piperazine diethanesulfonic acid (PIES, for pH 7), 2-(cyclohexylamino)ethanesulfonic acid (CHES, for pH 9) and 3-(cyclohexylamino)-1-propanesulfonic acid (CAPS, for pH 11). All of the buffers (0.03 M) were adjusted to the exact required pH using sodium hydroxide aqueous solution (1 M). Unless otherwise stated, all of the chemicals used in this study were used as received without further treatment.

**Synthesis of Vivianite.** Vivianite was synthesized by following the method previously described. An aqueous solution of Fe(NO\(_3\))\(_2\)·6H\(_2\)O (80 g/L) and a solution containing CH\(_3\)OONa (13.33 g/L) and Na\(_2\)HPO\(_4\) (66.67 g/L) were mixed under continuous stirring for 2 days in an anaerobic chamber (Coy Laboratory Products Inc.). The bluish-gray precipitate was continuously washed with DDW to remove remnant iron species during its synthesis. The iron contents in the aqueous phase were monitored after each washing process until the supernatant solution was free of iron. The washed vivianite was then centrifuged and dried under anaerobic conditions, after which it was analyzed by X-ray diffraction (XRD). The XRD results of the synthesized vivianite agreed well with the corresponding Joint Committee on Powder Diffraction Standards (JCPDS) diffraction data.

**Experimental Procedures.** Batch experiments were conducted to investigate the degradation kinetics of CCl\(_4\) by vivianite. Borosilicate glass vials (23.4 mL) sealed with a three-layered septum system (PTFE film, PTFE film-lined rubber septum, and lead foil) were used as batch reactors. The buffer solutions (23.3 mL), prepared using a previously described method, were transferred to vials each containing the same amounts of vivianite (0.234 g). Each vial was spiked using a gastight syringe (Hamilton) with 100 μL of CCl\(_4\) stock solution (23.4 mM) prepared in methanol, resulting in an initial CCl\(_4\) concentration of 0.1 mM. The vials were then taken out from the anaerobic chamber and placed on a tumbler (7 rpm) for vertical mixing. To investigate the effect of pH on the dechlorination of CCl\(_4\) by vivianite, five different buffer solutions (pH 3, 5, 7, 9, and 11) were used. Controls (buffer solutions without vivianite) at the different pHs were also prepared to evaluate potential losses during the reaction. All of the experiments in this research were conducted in duplicate at 26 ± 0.5 °C.

**Analytical Procedures.** CCl\(_4\) and the other chlorinated transformation products were quantified using a gas chromatograph (GC) equipped with electron capture detector (ECD, GC-2010 Plus, Shimadzu, Co.) and HP-5 column. At each sampling time, the vials were centrifuged at 3000 rpm for 5 min and a 100 μL portion of the supernatant was transferred to a 2 mL GC vial containing 0.9 mL of hexane with an internal standard (i.e., 0.25 mM 1,2-dibromopropane). The GC vials were shaken at 150 rpm using an orbital shaker for 1 h to extract the chlorinated organics, and the extractant (1 μL) was automatically injected into the injector (200 °C) with a split ratio of 30:1. The temperatures of the detector and oven were kept constant at 230 and 100 °C, respectively.

The nonchlorinated hydrocarbon transformation products were analyzed using a GC equipped with a flame ionization detector.
The vivianite (010) surface was cleaved by a although it underestimates the band gap due to the self-

properties such as lattice parameters of the vivianite structure functional is regarded to be reliable in predicting equilibrium contribution of the cavitation energy.

The DFT optimized bulk structure for vivianite is shown in Figure 1a. The simulation cell consists of three iron atoms, two phosphate groups, and eight water molecules that coordinate to the iron atoms. The optimized cell parameters are \( a = 9.99 \, \text{Å}, b = 12.84 \, \text{Å}, c = 4.77 \, \text{Å}, \) and \( \beta = 103.82^\circ \). The experimental values of \( a = 10.021 \, \text{Å}, b = 13.441 \, \text{Å}, c = 4.721 \, \text{Å}, \) and \( \beta = 102.84^\circ \).

The vivianite has a layered structure in which stacking occurs along the \( b \)-direction, where relatively weak hydrogen bonds between the H\(_2\)O ligands connect each layer. Therefore, the most stable surface of vivianite is the (010) surface, which is shown in Figure 1b. On the surface, two distinct types of iron centers are exposed: (1) Fe A sites, which are iron atoms coordinated by two H\(_2\)O molecules, four phosphate oxygen atoms, and one iron atom, and (2) Fe B sites, which are iron atoms coordinated by four H\(_2\)O molecules and two phosphate oxygen atoms. We found that the energy states near the Fermi level are mostly contributed by Fe 3d states with negligible contribution by phosphate groups (Figure S1, Supporting Information). Since the reduction chemistry is dominated by the valence states, we think that the primary role of phosphate group is to structurally stabilize the ferrous ion within the crystal surface where the CCl\(_4\) uptake an electron.

In our calculations, we fully relaxed the positions of the nuclei in the one-layer slab model of vivianite with the (010) surface, which was employed to investigate the CCl\(_4\) dechlorination pathway (the \( a, c, \) and \( \beta \) cell parameters were fixed as their bulk values, which were optimized using DFT). In addition, we assumed that the sequential transformations of chemical species were occurred on vicinal Fe centers, not only for the brevity of our calculations with limited size of the simulation cells but also for considering that the reaction intermediate radicals (e.g., \(^{12}\text{CCl}_3\)) would have not enough lifetime to migrate to non-neighboring Fe centers. However, we note that there could be a possibility that the reactions undergo at distant Fe centers, which might slightly change the reaction energetics.

### III. RESULTS AND DISCUSSION

Experimental Results on the Reductive Degradation of CCl\(_4\) by Vivianite. Figure 2 shows the dechlorination kinetics of CCl\(_4\) by vivianite at pH 7. The control solutions without vivianite showed no significant change in the CCl\(_4\) concentration over a period of 40 d, indicating that CCl\(_4\) loss by sorption in the batch reactor or volatilization during the sampling process was not significant in this study. In contrast with the control solutions, approximately 90.3% of the initial CCl\(_4\) was removed by vivianite in 40 d. Over the course of the CCl\(_4\) reduction, HCCl\(_3\) and formate gradually increased, finally reaching levels of 0.051 and 0.016 mM at the last sampling.

Figure 1. (a) Optimized structure of bulk vivianite using DFT, and (b) (010) surface slab model of vivianite (left panel). Two distinct Fe\(^{\text{II}}\) sites exist on the surface, which are denoted as Fe A site and Fe B site (see top view of the surface slab model shown in the right panel).
time, while a negligible amount of methane was detected during the reaction. CH2Cl2 and CH3Cl were not detected in this study. We also observed a continuous increase (to ∼0.25 mM) in the amount of chlorine ion in aqueous solution, with 97.6% chlorine mass balance measured at the last sampling time. The total carbon mass fluctuated from 62.6−91.2% during the reaction, indicating that some of the nonchlorinated intermediates may be (1) volatilized during the sampling process, (2) sorbed on the mineral surface, and (3) undetected by the analytical methods used in this study. The formation of HCCl3 and formate accompanying the CCl4 degradation implies that both the hydrogenolysis and the carbene hydrolysis pathways may have occurred simultaneously.30,31 More details regarding the formation of HCCl3 and formate, as well as of the nonchlorinated intermediates on the vivianite surface during the CCl4 degradation, will be discussed in the next section.

![Figure 3a](image1.png)

**Figure 3a.** Effect of the suspension pH (3−11) on the CCl4 degradation kinetics with vivianite. The initial concentration of CCl4 was 0.1 mM and standard deviations obtained from duplicate measurements are indicated as the error bars.

Figure 3. (a) Effect of the suspension pH (3−11) on the CCl4 degradation kinetics with vivianite. The initial concentration of CCl4 was 0.1 mM and standard deviations obtained from duplicate measurements are indicated as the error bars. (b) Change in the observed kinetic constants for the reductive dechlorination of CCl4. (c) Product formation rate constants for HCCl3, formate, and chloride ion as a function of pH.
0.35 eV and that on the Fe B site is 0.28 eV. We note that the desorption of water involves an entropic gain, facilitating the water desorption, because the liquid water molecule has more degrees of freedom of which extent can be estimated as ∼0.1 eV considering the entropy of water (70 J/mol/K) and the entropy of ice (41 J/mol/K).

We investigated the binding affinities of the chlorinated compounds at the reactive site generated on the Fe B site, which has the lower water desorption energy. When CCl₄ is placed near the Fe II in the initial structure of the DFT calculation, it spontaneously decomposes into trichloromethyl radical (•CCl₃), and after performing the geometry optimization steps, one Cl⁻ ion is observed to be left behind on the Fe site. In other words, a dissociate electron transfer step (diss-ET) takes place. Our DFT results showed that the diss-ET energy of CCl₄ is 0.35 eV downhill with a negligible reaction barrier (Figure 4b). We then determined that the adsorbed Cl⁻ is replaced with OH⁻ with a downhill reaction energy of 0.22 eV (evaluated following Scheme S1), which supports the experimental result demonstrating a significant increase in the chlorine ion concentration in the aqueous phase. This is also consistent with a previous discussion suggesting that the reductive deformation of CCl₄ by Fe-containing minerals₄,₃¹−₃₈ is initiated by the first electron transfer and is followed by bond cleavage between carbon and chloride₃₉,₄₀.

After the diss-ET step, hydrolysis of CCl₄ can occur in two ways: trichloromethyl radical (•CCl₃) can be transformed to chloroform (HCCl₃) by abstracting hydrogen (hydrogenolysis)₄₁ or it can be further reduced to form trichlorocarbanion as a surface-stabilized form (:CCl₃⁻*) that is released as a form of HCCl₃ by abstracting a proton (carbenoid formation).₄₂ We have investigated the DFT energetics required for these two pathways.

For the hydrogenolysis pathway, we calculated the energetics associated with the uptake of one hydrogen atom by •CCl₃ from the water molecule coordinating Fe II (assumed to be the nearest Fe B site in our calculations), accompanied by the oxidation of Fe II. This yields HCCl₃ and two oxidized Fe III centers coordinated to OH⁻ on the surface, resulting in a net change in energy calculated as 1.23 eV downhill (Figure 4c).
For the carbenoid formation pathway, we considered the following sequential reactions: (1) One additional water molecule desorbs from the surface (0.11 eV), $\cdot$CCl$_3^{-*}$ can be decomposed into dichlorocarbene ($\cdot$CCl$_2^{-*}$) and Cl$^{-*}$ on the surface (0.07 eV uphill). Cl$^{-*}$ is then exchanged with OH$^{-*}$ in the solution phase (0.50 eV downhill), where OH$^{-*}$ undergoes a nucleophilic attack on $\cdot$CCl$_3^{-*}$ to form $\cdot$CCl$_3$OH$^{-*}$. The subsequent additional C–Cl bond cleavage is an exothermic process by 0.34 eV, which yields CClOH$^{-*}$ and Cl$^{-*}$. If an electron rearrangement takes place, leaving behind H$^{+}$ and Cl$^{-*}$, CClOH$^{-*}$ is converted to CO$^{-*}$ (1.17 eV downhill), which can be easily released from the surface (0.003 eV uphill). Otherwise, OH$^{-*}$ (which is replaced with Cl$^{-*}$ in a 0.45 eV downhill process) participates in a nucleophilic attack on CClOH$^{-*}$, yielding $\cdot$CCl(OH)$_2^{-*}$ (1.47 eV downhill process). An additional C–Cl bond cleavage of $\cdot$CCl(OH)$_2^{-*}$ yields C(OH)$_2^{-*}$ and Cl$^{-*}$ (0.01 eV downhill), where C(OH)$_2^{-*}$ is desorbed from the surface while proton rearrangement yields the formic acid (HCOOH). (b) Relative energy diagram for the overall nonreductive dechlorination of $\cdot$CCl$_3^{-*}$ to form either CO or HCOOH.

Figure 5. (a) Nonreductive dechlorination (nr-deCl) of $\cdot$CCl$_3^{-*}$. When one water molecule is desorbed from the surface (0.11 eV), $\cdot$CCl$_3^{-*}$ can be decomposed into dichlorocarbene ($\cdot$CCl$_2^{-*}$) and Cl$^{-*}$ on the surface (0.07 eV uphill). Cl$^{-*}$ is then exchanged with OH$^{-*}$ in the solution phase (0.50 eV downhill), where OH$^{-*}$ undergoes a nucleophilic attack on $\cdot$CCl$_3^{-*}$ to form $\cdot$CCl$_3$OH$^{-*}$. The subsequent additional C–Cl bond cleavage is an exothermic process by 0.34 eV, which yields CClOH$^{-*}$ and Cl$^{-*}$. If an electron rearrangement takes place, leaving behind H$^{+}$ and Cl$^{-*}$, CClOH$^{-*}$ is converted to CO$^{-*}$ (1.17 eV downhill), which can be easily released from the surface (0.003 eV uphill). Otherwise, OH$^{-*}$ (which is replaced with Cl$^{-*}$ in a 0.45 eV downhill process) participates in a nucleophilic attack on CClOH$^{-*}$, yielding $\cdot$CCl(OH)$_2^{-*}$ (1.47 eV downhill process). An additional C–Cl bond cleavage of $\cdot$CCl(OH)$_2^{-*}$ yields C(OH)$_2^{-*}$ and Cl$^{-*}$ (0.01 eV downhill), where C(OH)$_2^{-*}$ is desorbed from the surface while proton rearrangement yields the formic acid (HCOOH). (b) Relative energy diagram for the overall nonreductive dechlorination of $\cdot$CCl$_3^{-*}$ to form either CO or HCOOH.
while the resulting OH$^-$ stabilizes the oxidized Fe$^{II}$ center. As summarized in Figure 4c, the DFT energetics of this pathway are (1) 0.07 eV uphill for the second water desorption, (2) 0.32 eV downhill for the adsorption of $^*$CCl$_3$ to form $^*$CCl$_4$, and (3) 0.99 eV downhill for the final HCCl$_3$ formation. This implies that both the hydrolysis and carbeneoid formation pathways are responsible for the HCCl$_3$ production.

By repeating either of the hydrolysis or carbeneoid formation pathways, CH$_2$Cl$_2$, CH$_3$Cl, or methane can be formed, as has been observed in other Fe-containing mineral systems. However, the DFT energy for the diss-ET step to HCCl$_3$ was calculated as 0.37 eV uphill (Figure S2), indicating that further reduction is energetically unfavorable. This supports our experimental results, which show that no significant amount of methane was generated and that CH$_3$Cl and CH$_2$Cl$_2$ were also not detected. It also explains why the dechlorination of HCCl$_3$ hardly occurs in the presence of vivianite.

Instead of further sequential reductive dechlorinations, we considered a nonreductive dechlorination (nr-deCl) pathway of HCCl$_3$. The chemical equilbrium of HCCl$_3$ can be considered as follows:

$$\text{HCCl}_3(\text{aq}) \rightleftharpoons \text{CCl}_3(\text{aq}) + \text{H}^+(\text{aq}) \quad K_a$$
$$\text{CCl}_3(\text{aq}) \rightleftharpoons \text{CCl}_4^+ \quad K_{bind}$$

where $K_a$ and $K_{bind}$ are equilibrium constants, and $K_a = 10^{-25}$ because the pH value of HCCl$_3$ is 25. This leads to a concentration ratio of the carbeneoid ($^*$CCl$_4^+$) and HCCl$_3$ of $|\text{[CCl}_3]|/|\text{[HCCl}_3]| = K_aK_{bind}|\text{[H}^+]| = 10^{45-25} \exp(-\Delta G_{bind}/RT)$, where $\Delta G_{bind}$ is the binding free energy of the trichlorocarbonanion. The above implies that $|\text{[CCl}_3]|$ becomes substantial at neutral pH only when $\Delta G_{bind} < -1$ eV and exponentially increases when the pH increases. This result provides evidence that the nr-deCl pathway takes place through the formation of $^*$CCl$_4^+$ because the higher pH produces the faster kinetics of CCl$_4$ decomposition observed in our experiment, which are likely due to the higher concentrations of $^*$CCl$_4^+$ present under high pH conditions. Moreover, we note that both diss-ET pathway (Figure 4) and nr-deCl pathway (Figure 5; details will be discussed in below) requires anion exchange steps, suggesting another possible pH effect; the higher pH leads to the faster anion exchange of the surface adsorbed Cl$^-$ into OH$^-$, promoting the CCl$_4$ decomposition kinetics. This explains the experimentally observed pH dependence of the reaction kinetics; (1) the formation kinetics of Cl$^-$ was significantly enhanced at the pH regime of 5–7, which is understandable as the outcome of enhanced kinetics of the surface anion exchange considering that the low point of zero charge (PZC) value of vivianite surface that is 3.5; (2) the formation kinetics of formate was significantly enhanced at the high pH regime of 11, which is understandable as the outcome of the equilibrium shift toward the carbeneoid formation. We note that the two distinct pH effect promoting the surface anion exchange and carbeneoid formation are disadvantageous in elevating the CCl$_4$ decomposition kinetics, and thus the kink behavior observed in the CCl$_4$ decomposition kinetics at the pH regime of 7–9 is conjectured to be manifested by the interplay of two distinct pH effects.

To provide the full atomistic and energetic details of the reaction, we computed the DFT energetics for the nr-deCl pathway starting from $^*$CCl$_3^-$, as shown in Figure 5a. The water desorption energy from the Fe$^{III}$ center where the carbeneoid is located was calculated to be 0.11 eV uphill. We then found that $^*$CCl$_3^-$ can decompose into dichlorocarbene ($^*$CCl$_2^+$) and Cl$^-$ at the Fe$^{III}$ center, for which the DFT energy is only 0.07 eV uphill. This step is followed by the ligand exchange of Cl$^-$ with OH$^-$, which was calculated to be 0.50 eV downhill.

We found that the Cl–C–Cl angle of $^*$CCl$_2^+$ is 109.6$^\circ$, which is nearly identical to the angle of free Cl–C–Cl (110.0$^\circ$) (Figure S3). Considering that CCl$_3$ has a trigonal planar structure with sp$^2$ hybridized bonds and little steric hindrance, as a following step we examined the substitution of Cl by OH, which occurs though the nucleophilic attack of OH$^-$ on the sp$^2$ carbon center of $^*$CCl$_2^+$. This yields Fe$^{III}$ coordinated with CCl(OH) via the formation of an intermediate state of $^*$CCl$_3$(OH)$^-$*. The overall change in energy for this process was calculated by DFT to be 1.31 (=0.97 + 0.34) eV downhill. By redistributing electrons and chemical bonds, CClOH$^*$ can form CO$^*$ with the release of HCl (1.17 eV downhill), which is followed by the release of CO (0.03 eV uphill).

If one additional nucleophilic attack by OH$^-$ on the sp$^2$ carbon center of $^*$CCl(OH)$^*$ occurs, C(OH)$_2^*$ will be generated via the formation of $^*$CCl(OH)$_2^*$, for which the overall DFT-calculated change in energy is 1.48 (1.47 + 0.01) eV downhill. The C(OH)$_2^*$ can then desorb from the surface, after which proton rearrangement will occur almost immediately within the aqueous medium, yielding HCOOH. The energy required for this process was calculated to be 1.64 eV downhill.

Our DFT study of the energetics shows that the thermodynamic driving force toward the formic acid formation pathway is greater than that of the CO formation pathway (see Figure 5b), which partly explains the propensity for formation of the experimentally observed products (formate production is preferred over CO production on the vivianite surface). However, considering that both pathways are energetically available and consist of downhill processes, it is difficult to find a reasonable rationale for the CO production pathway being virtually switched-off, as was observed in our experiment. We note that the pathways discussed herein are solely based on relative thermodynamic energies; therefore, there could be additional barriers and local minima between neighboring ground states. In particular, we expect that the heterogeneous dissociation of the C–Cl bond, which is required to release HCl from CClOH$^*$ to form CO, would be kinetically unfavored. Moreover, other chemical constituents such as ions in the buffer solution or impurities may help stabilize/destabilize one pathway over the other, implying that determination of the reaction pathways could be complicated by other factors. For instance, we note that more CO was formed when we used the biogenic vivianite than the chemicogenic vivianite, even under the same pH conditions.

IV. CONCLUSION

Because a variety of ferrous iron minerals are known to be electron donors for CCl$_4$ degradation, a thermodynamic and kinetic investigation of the reaction mechanism of the reductive dechlorination of CCl$_4$ by vivianite was carried out in this study. The experimental and the theoretical results represent a comprehensive investigation on the kinetics and reactive pathways of abiotic CCl$_4$ degradation on the surface of vivianite under anaerobic condition. These results indicate that (1) *CCl$_3$ is the key intermediate in the production of HCCl$_3$. 
and formate; (2) $\text{CCl}_3$ abstracts a hydrogen from surface water molecules to generate HCCl (hydrogenolysis pathway); (3) $\text{CCl}_3^\cdot$ is formed on the vivianite surface by further reduction of $\text{CCl}_2^\cdot$ (which also can form HCCl when it is desorbed and protonated (carbene hydrolysis via carbenedium formation pathway)); (4) $\text{CCl}_2^\cdot$ is nonreductively dechlorinated to form $\text{CCl}_1^\cdot$, followed by sequential nucleophilic attack by OH$^\cdot$ yielding $\text{CCl(OH)}^\cdot$ and $\text{C(OH)}_2^\cdot$; and (5) $\text{CCl}_1^\cdot$ and $\text{C(OH)}_2^\cdot$ can respectively form CO and formate when they are desorbed.

The study of the CCl$_4$ reduction pathway on vivianite provides us with significant understanding by analyzing every reduction step. On the basis of the results of the calculated rate constants, which increase with increasing pH, the surrounding pH environment may act as an important rate-determining factor during the remediation process. According to previous studies, vivianite-rich layers formed by precipitation$^{44}$ and in groundwater sediments$^{37}$ where CCl$_4$ is detected often. Therefore, the results obtained from this study indicate that the natural attenuation of CCl$_4$ by reductive dechlorination in the presence of vivianite at subsurfaces may be taking place.

Most studies have provided explanations for reaction mechanisms and pathways using experimental results obtained from partial transformation products, and have carried out analyses that rely on a wide variety of assumptions. In contrast, theoretical results from DFT calculations directly suggest thermodynamically favorable intermediates on mineral surfaces for every single step, and provide a wealth of information on their molecular structures, bonding character, and oxidation states. Recently, large reaction mechanistic studies have been undertaken using computational calculations regarding sorption,$^{18}$ reductive dechlorination,$^{39}$ and oxidative decomposition,$^{60}$ among others.$^{15,51,52}$ We believe that the systematic investigation of the reaction of dechlorination taking place on a vivianite surface presented in this study may provide fundamental aid for the development of an adaptable application for other halogenated species and minerals. We believe that modeling systems and reaction mechanisms on the surfaces of structures serves not only as a supplement for experimental data but also can provide basic knowledge and optimal results through simple control of the studied environments, leading to promising alternative approaches.

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