Carbon and Chlorine Isotope Fractionation Patterns Associated with Different Engineered Chloroform Transformation Reactions

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Supporting Information

ABSTRACT: To use compound-specific isotope analysis for confidently assessing organic contaminant attenuation in the environment, isotope fractionation patterns associated with different transformation mechanisms must first be explored in laboratory experiments. To deliver this information for the common groundwater contaminant chloroform (CF), this study investigated for the first time both carbon and chlorine isotope fractionation for three different engineered reactions: oxidative C–H bond cleavage using heat-activated persulfate, transformation under alkaline conditions (pH ~ 12) and reductive C–Cl bond cleavage by cast zirconium iron, Fe(0). Carbon and chlorine isotope fractionation values were −8 ± 1‰ and −0.44 ± 0.06‰ for oxidation, −57 ± 5‰ and −4.4 ± 0.4‰ for alkaline hydrolysis (pH 11.84 ± 0.03), and −33 ± 11‰ and −3 ± 1‰ for dechlorination, respectively. Carbon and chlorine apparent kinetic isotope effects (AKIEs) were in general agreement with expected mechanisms (C–H bond cleavage in oxidation by persulfate, C–Cl bond cleavage in Fe(0)-mediated reductive dechlorination and E1cb elimination mechanism during alkaline hydrolysis) where a secondary AKIEC2 (1.00045 ± 0.00004) was observed for oxidation. The different dual carbon-chlorine (Δδ13C vs Δδ37Cl) isotope patterns for oxidation by thermally activated persulfate and alkaline hydrolysis (17 ± 2 and 13.0 ± 0.8, respectively) vs reductive dechlorination by Fe(0) (8 ± 2) establish a base to identify and quantify these CF degradation mechanisms in the field.

1. INTRODUCTION

Chloroform (CF) is both an anthropogenic environmental contaminant widely distributed around the world as well as a natural compound formed in various aquatic and terrestrial environments.1–3 CF of anthropogenic origin has been extensively used as degreasing agent and as a precursor to Teflon and various refrigerants and was historically used in medicine as anesthetic. It is formed as oxidation byproduct during drinking water treatment and may form as a daughter product of carbon tetrachloride (CT) dehalogenation at contaminated sites. As a result, CF is one of the most frequently detected volatile organic compounds (VOCs) in groundwater.4 Taking into account its high ecotoxicity,5 CF prominently ranks among the halogenated VOCs on the Agency for Toxic Substances and Disease Registry priority list of hazardous substances.6

Aerobic and anaerobic cometabolic biodegradation processes of CF have been described.8 However, CF cometabolic degradation is restricted by several environmental factors such as the presence of other specific compounds that inhibit CF degradation, the availability of the substrate or the toxicity of derived metabolites.9 Reductive dechlorination of CF via dehalorespiration by two Dehalobacter sp. strains (CF50 and CF51) has recently been described in laboratory studies10–14 and proposed as anaerobic bioremediation strategy. However, this strategy is only applicable to contaminated sites in the absence of its parent compound, i.e., CT, which has been shown to strongly inhibit CF dehalorespiration in an enrichment culture containing Dehalobacter spp.15 In turn, CF itself is a strong inhibitor of chlorinated ethene- or ethane-degrading cultures even when present at low concentrations.16–19 These interdependencies make the remediation of sites contaminated with...
several chlorinated compounds particularly challenging so that multiple-stage remediation strategies are warranted in which inhibitors like chloromethanes are removed upfront. Abiotic reactions bear potential to accomplish such an initial removal. Naturally occurring ion-bearing minerals like goethite and iron sulfide under low-redox environments have been demonstrated to be involved in the reductive dechlorination of CF. However, due to the very restricted natural attenuation conditions for CF and its complex distribution in the subsurface as a dense nonaqueous phase liquid (DNAPLs), more efficient engineered remediation strategies have been proposed to increase CF removal in the environment. As a result of the high oxidation state of carbon in CF, its degradation by in situ chemical oxidation (ISCO) is in general much less effective than for chlorinated ethers using common oxidants such as peroxide, or Fenton’s Reagent. However, thermally activated PS was recently shown to be a better option for efficient CF oxidation with the advantage that under thermal activation, the strongly oxidizing sulfate radical and other reactive intermediates (i.e., hydroxyl radicals, or reducing radicals such as superoxide radicals, O$_2^-$) can be generated at neutral pH.

Alternatively, CF alkaline hydrolysis has recently been proposed as a remediation technology based on its occurrence in drainage trenches filled with concrete-based construction wastes. For the sustainable use of this new remediation strategy, identifying and assessing the performance of CF degradation reaction by alkaline hydrolysis, as well as understanding the underlying mechanism, is important.

Finally, CF reductive dechlorination by zerovalent metals has been studied at laboratory scale. Nevertheless, this remediation strategy has been successfully proven at field sites contaminated by chlorinated ethers using permeable reactive barriers with micro/macro-scale Fe(0) or Fe(0) nanoparticle injection.

Improved methods are needed to delineate the relative efficacy of the above-mentioned CF remediation approaches in the field. During the last decades, compound-specific isotope analysis (CSIA) has evolved as a tool to monitor transformation reactions and to quantify the progress of natural and enhanced remediation of organic contaminants. Molecules with light isotopes in the reactive position typically react slightly faster than molecules containing the heavy ones leading to a kinetic isotope effect (KIE). As a consequence, the heavier isotopes (e.g., $^{13}$C and $^{37}$Cl) usually become enriched in the remaining substrate. For a given reaction, quantification of the extent of contaminant transformation based on stable isotope ratios requires the experimental determination of isotopic fractionation ($\epsilon$, see Materials and Methods).

Isotopic fractionation values for transformation reactions need to be known for very practical reasons: (i) to understand what changes in isotope values can be expected in the field at all, and whether this holds promise to qualitatively detect degradation; (ii) to understand what mechanism lies behind the isotope effect, in order to subsequently chose an appropriate $\epsilon$ value for quantification in the field.

In order to gain insight into the underlying reaction mechanism, apparent kinetic isotope effects (AKIEs) can be derived from determined $\epsilon$ values taking into account which of the atoms in the target molecule are expected to be present at the reactive position. Comparison of the observed AKIEs to the theoretical maximum KIEs ("semiclassical Streitwieser Limits") associated with breakage of chemical bonds, enables interpretation of occurring transition state(s) of a bond cleavage in terms of (i) primary isotope effects affecting the atoms present in the reacting bond, (ii) secondary isotope effects affecting atoms located adjacent to the reacting position. Often, however, it is uncertain whether additional factors exert an influence on observable isotope fractionation such as (iii) masking due to rate-limitation in mass transfer, and (iv) superimposed isotope effects of multiple reaction steps typical of enzyme catalysis or multistep chemical reactions.

When observable isotope fractionation of a single element varies between experiments, it is, therefore, often uncertain whether this is due to a different mechanism, or whether these other factors are responsible. Dual-element isotope plots, that is, graphs in which changes in isotope values of one element are plotted against those of a second offer a more reliable distinction between reaction mechanisms than $\epsilon$ values alone.

Such information can be highly valuable in field situations. Nondestructive abiotic natural processes, such as sorption, volatilization or diffusion strongly affect concentrations of a contaminant, but generally do not cause significant isotopic fractionation. Temporal or spatial shifts in isotope ratios, in contrast, are highly indicative of degradation and can, therefore, better monitor the success of remediation strategies at contaminated sites than mass balances alone. Dual (or multi) isotope patterns, finally, can even be used to derive the relative contribution of different reaction mechanisms and then to quantify the efficiency of each of them in the field—provided that $\epsilon$ values of these processes have previously been characterized in laboratory experiments.

Reported carbon isotope effects during CF transformation are, however, scarce in the literature. Chan et al. reported a carbon isotope fractionation value of $-27.5 \pm 0.9\%$ during dehalorespiration of CF by a mixed culture containing Dehalobacter sp. strain CF50. In comparison, a much lower $\epsilon_C$ value of $-4.3 \pm 0.4\%$ was reported by Lee et al. for the same dechlorination reaction by a mixed consortium containing another Dehalobacter sp. strain, UNSWDBH, whereas isotope fractionation in CF abiotic reductive dechlorination by microsized Fe(0) was found to be indistinguishable from that of the first experiment ($-29 \pm 2\%$). Significantly, larger carbon isotopic fractionation was observed for CF alkaline hydrolysis at pH ranging from 11.8 to 12.7 ($-53 \pm 3\%$). To the best of our knowledge, chlorine isotope fractionation during any CF transformation mechanism has not been reported so far. Specifically, understanding whether different reaction mechanisms lead to characteristic patterns in dual C–Cl isotope plots is still limited even for chlorinated ethers and, to our knowledge, is currently nonexistent for chlorinated methanes. Hence, there is a need to explore dual element CSIA for defined reactions under controlled laboratory conditions to pave the path for the interpretation of isotope data in field studies.

Therefore, the goal of this study was to determine carbon and, for the first time, chlorine isotope fractionation patterns for different transformation processes of CF in important abiotic engineered reactions in order to explore the ability of CSIA to identify these processes at field sites. The selected chemical reactions were: oxidative C–H bond cleavage by radicals produced from PS activation, alkaline hydrolysis of chloroform at pH 12 and C–Cl bond cleavage in reductive dechlorination by cast milli-sized Fe(0).
2. MATERIAL AND METHODS

2.1. Experimental Setup. All the experiments were conducted in duplicate using glass vials completely filled with aqueous solution without headspace to avoid partitioning of chlorinated volatile compounds into the gas phase. For the experiments with heat-activated PS and alkaline hydrolysis, 21 mL vials sealed with PTFE-coated rubber stoppers and aluminum crimp seals were used, whereas the Fe(0) experiments were performed using 42 mL clear glass vials capped with PTFE-coated rubber stoppers and plastic screw caps. A list of chemicals and additional experiment details is available in the Supporting Information (SI).

For the thermally activated PS experiments, the vials were filled with a pH 7 buffer solution and 0.5 mL of solutions with variable concentrations of PS were added to achieve initial PS-to-CF molar ratios of 5/1, 10/1, or 40/1. The experiments were performed in a thermostatic water bath at 50.0 °C. The vials were placed in the dark at 4 °C until analysis. Control experiments without iron were also carried out. The experiments were performed at room temperature (∼25 °C) and they lasted 51 h. Reactions were stopped by filtration through 0.2 μm filters at different time intervals and samples for analysis were stored frozen in 10 mL vials covered with aluminum foil.

2.2. Analytical Methods. Detailed descriptions of analytical methods are available in the SI. Briefly, concentration measurements of chlorinated compounds were performed by headspace (HS) using GC/MS as explained elsewhere,24 except for the samples from Fe(0) experiments for which GC/TOF/MS was used. Chloride anion concentrations were analyzed by high-performance liquid chromatography. Carbon isotopic analyses of CF and some detectable volatile daughter products were performed using two different GC/IRMS systems located at the University of Neuchâtel (GC/IRMS-1)50 and at the Scientific and Technological Centers of the University of Barcelona (GC/IRMS-2).24 Chlorine isotope CF analyses were performed using a GC/qMS system from the University of Neuchâtel as explained elsewhere50 or a GC/IRMS system from the Institute of Groundwater Ecology of the Helmholtz Institute for Environment and Health in Neuchâtel.

2.3. Kinetic Analysis. The rates of CF degradation were calculated using the following equation:

[k'] = (C0 / C) / t

where [k'] is the pseudo-first-order rate constant, C0 is the initial concentration of CF, C is the concentration of CF at time t, and t is the reaction time. The units of [k'] are d−1. The rate constants were calculated using linear regressions in Sigma Plot 10.0 for Windows. Dashed lines represent 95% CI of regression. For CF dechlorination with Fe(0), [k'] was calculated omitting data after 30 days when the disappearance of CF almost stopped. In the lower panels, error bars of individual data points indicate standard deviations of the measurements. In most cases, error bars are smaller than the symbols.

Figure 1. CF degradation kinetics (upper panels) and changes in C and Cl isotope ratios (lower panels) during oxidation by thermally activated PS with an initial PS/CF molar ratio of 40/1 (A), alkaline hydrolysis (B) and dechlorination by Fe(0) (C). Data from duplicate experiments (A and B parallel series) and from control (CTRL) experiments are shown. In the upper panels, the error bars show the uncertainty in concentration measurements. In some cases, error bars are smaller than the symbols. $k'_{obs}$ values were obtained from curve fittings according to eq 65 (see Figure S1). Fits were conducted with linear regressions in Sigma Plot 10.0 for Windows. Dashed lines represent 95% CI of regression. For CF dechlorination with Fe(0), $k'$ was calculated omitting data after 30 days when the disappearance of CF almost stopped. In the lower panels, error bars of individual data points indicate standard deviations of the measurements. In most cases, error bars are smaller than the symbols.
Zentrum München (GC/IRMS-3). An interlaboratory comparison demonstrating excellent agreement between the two analytical methods is provided in Heckel et al.67

2.3. Isotope Data Evaluation. Bulk carbon and chlorine ε values were obtained from the slope of the linearized Rayleigh equation for a closed system:36

\[
\ln \left( \frac{\delta + 1}{\delta_0 + 1} \right) = \epsilon \times \ln f
\]

(1)

where \(\delta_0\) and \(\delta\) are isotope values in the beginning (0) and at a given time \(t\), respectively, and \(f\) is the fraction of substrate remaining at time \(t\). Isotope signatures were reported in per mil (‰) using the delta notation relative to international standards, that is, Vienna PeeDee Belemnite for carbon (\(\delta^{13}C_{VPDB}\)) and the international Standard Mean Ocean Chloride (SMOC) for chlorine (\(\delta^{37}Cl_{SMOC}\)):

\[
\delta(\text{‰}) = \left( \frac{R}{R_{\text{std}}} - 1 \right) 
\]

(2)

Figure 2. Logarithmic plots according to Rayleigh eq (eq 1) of carbon (left panels) and chlorine (right panels) isotope ratios during CF oxidation by thermally activated PS (A and B), alkaline hydrolysis (C and D) and dechlorination by Fe(0) (E and F). Data from duplicate experiments were used for estimating \(\epsilon_C\) and \(\epsilon_{Cl}\). Dashed lines represent 95% CI of the linear regression. Error bars display the uncertainty calculated by error propagation including uncertainties in concentration and isotope measurements. In some cases, error bars are smaller than the symbols.
where $R$ and $R_{oai}$ are the isotope ratios of the sample and the standard, respectively.

Errors given for $\epsilon$ values are the 95% confidence intervals (CI) of the slope of the regression line in the Rayleigh plots.

The apparent kinetic isotope effects (AKIEs) were calculated to evaluate the intrinsic isotope effect of the bond cleavage (see equations in the SI).

For dual C–Cl isotope plots, the slope of the correlation trend was determined by linear regression and the uncertainty corresponds to the 95% CI.

3. RESULTS AND DISCUSSION

3.1. Carbon and Chlorine Isotope Fractionation.

Changes in CF concentration and C and Cl isotope ratios of CF during degradation by the three different mechanisms are shown in Figure 1. No CF degradation was observed in the experimental controls (without adding PS, Fe(0) or at neutral pH in the case of the hydrolysis reaction) for any of the three experimental controls (without adding PS, Fe(0) or at neutral pH). The pH was kept at circumneutral values (7.0 ± 0.2) during the course of the experiment. This reaction trend was determined by linear regression and the uncertainty corresponds to the 95% CI.

Alkaline Hydrolysis. Carbon and chlorine isotope values remained constant ($-41.8 \pm 0.5^{\circ}\text{C}$ and $-2.6 \pm 0.4^{\circ}\text{C}$, respectively, both $n = 5$) in control vials at neutral pH. Under alkaline conditions (the pH remained constant 11.84 ± 0.03 over the duration of the experiment), a 85% decrease in CF concentration within approximately 35 days was observed (Figure 1B). Alkaline hydrolysis induced a significant isotope effect, resulting in $\delta^{13}\text{C}$ and $\delta^{37}\text{Cl}$ values up to $+70.6 \pm 0.3^{\circ}\text{C}$ and $+5.7 \pm 0.4^{\circ}\text{C}$, respectively, after 85% CF removal (SI Figure S3C, D). An $\epsilon_{C}$ of $-57 \pm 5^{\circ}\text{C}$ (Figure 2C) and $\epsilon_{\text{Cl}}$ of $-4.4 \pm 0.4^{\circ}\text{C}$ (Figure 2D) were determined. So far, the only reported carbon isotope fractionation value for CF alkaline hydrolysis has not been reported up to now. The reaction followed pseudo-first-order kinetics ($R^{2} = 0.92$, SI Figure S1) with a $k'$ of 0.052 ± 0.008 d$^{-1}$, which is in agreement with a previously reported rate constant of 0.047 ± 0.004 d$^{-1}$ obtained at a similar pH 11.9 ± 0.1. No particular attempts were made to identify potential products of CF degradation, such as carbon monoxide (CO), formate (HCO$_3^{-}$), and chloride (Cl$^{-}$). In our previous work, excellent chlorine balances were achieved in similar experiments, indicating that CF was completely dehalogenated without accumulation of chlorinated intermediates.$^{24}$

Fe(0) Dechlorination. CF in the controls without Fe(0) at pH 6.3 ± 0.2 did not show any changes in carbon and chlorine isotope composition ($\delta^{13}\text{C} = -47.8 \pm 0.5^{\circ}\text{C}$, $n = 4$ and $\delta^{37}\text{Cl} = -3.2 \pm 0.2^{\circ}\text{C}$, $n = 6$, respectively). In the presence of milliliter-sized Fe(0), CF isotope signatures of both elements showed significant changes leading up to values of $\delta^{13}\text{C} = +55.9 \pm 0.5^{\circ}\text{C}$ and $\delta^{37}\text{Cl} = +1.7 \pm 0.1^{\circ}\text{C}$, respectively, after 84% CF removal (SI Figure S3E, F). Isotope fractionation values of $\epsilon_{C} = -33 \pm 11^{\circ}\text{C}$ and $\epsilon_{\text{Cl}} = -3 \pm 1^{\circ}\text{C}$ were determined (Figure 2E and F). The obtained $\epsilon_{C}$ was not significantly different from $\epsilon_{C} = -29 \pm 2$% reported recently after 50% CF dechlorination by commercial microsized Fe(0).$^{25}$ Chlorine isotope fractionation associated with this reaction has not been reported yet. The pH did not vary significantly over the duration of the experiment (6.2 ± 0.2). The degradation kinetics followed a pseudo-first-order rate law at the beginning of the reaction but after 30 h the disappearance of CF almost stopped (Figure 1C). For Fe(0)-mediated dechlorination of chlorinated ethenes, iron surface passivation due to reactive site saturation by iron hydroxide precipitates has been suggested as the cause of increased reaction half-lives and deviations from pseudo-first-order kinetics at later stages of a reaction.$^{10}$ The obtained $k'$ was 0.07 ± 0.01 h$^{-1}$ ($R^{2} = 0.93$, SI Figure S1), which corresponds to a $k_{\text{obs}}$ of 2.1 ± 0.4 × 10$^{-2}$ L m$^{-2}$ d$^{-1}$ (see SI).

DCM and free chloride were detected as final products in Fe(0) experiments, whereas no compounds different from CF appeared in the control experiments without iron. The yield of DCM, defined as the moles of product formed per mole of CF transformed (DCM/([CF$_{0}$ − CF$_{t}$])), where subscripts 0 and t indicate initial time and different sampling times, respectively) ranged from 0 to 2.4% over time, showing that accumulation of DCM only accounted for a small fraction of the initial CF. DCM was depleted in $^{13}\text{C}$ compared to the initial isotopic composition of the substrate (CF). DCM showed a trend toward higher $\delta^{13}\text{C}$ values, reflecting the enrichment trend of
similar AKIE value (1.008) was obtained for 1,1,1-TCA KIE associated with oxidative cleavage of a C bond, indicating that the observed fractionation was dominated by the hydrogen isotope fractionation during CF oxidation with PS, as there is not primary chlorine isotopic effect and isotope-sensitive branching from the parent compound or intermediates, such as a dichloromethyl radical (SI Figure S4), to DCM. 

The uncertainty of AKIE and ε values corresponds to the 95% CI. In all cases, AKIE and ε were combined. Lines are linear regressions of the data sets with 95% CI (dashed lines). Error bars show uncertainty in isotope measurements. Note that error bars of δ13C values are smaller than the symbols.

The CF from which it was formed (SI Figure S3E). The DCM-related isotope fractionation εsubstrate-product was estimated as −19 ± 3‰ using the fitting parameter, D(Δ31Cl) = +13 ± 2‰ (R² = 0.62) (see equations in the SI). This discrepancy between the product-related and the substrate-related isotope fractionations (εCF–DCM = −19 ± 3‰ vs εC = −33 ± 11‰) is likely attributable to the formation of other products including isotope-sensitive branching from the parent compound or intermediates, such as a dichloromethyl radical (SI Figure S4), to DCM. However, due to the lack of DCM isotope signatures at early stages of reaction, such interpretations must be conducted with caution.

3.2. Mechanistic Considerations. For further elucidation of the reaction mechanisms, AKIE values were calculated using eq S16, to characterize the isotope effect of the cleavage of the chemical bond at the reactive positions. Table 1 summarizes the obtained results and proposed reaction pathways for the three studied reactions shown in Figure 3 and discussed in detail in the SI.

The AKIECl for the oxidative reaction was 1.008 ± 0.001, which is within the range of reported carbon AKIEs for oxidative C–H bond cleavage for both abiotic (1.008–1.015) and microbial oxidation reactions (1.001–1.044) (SI Table S2), indicating that the observed fractionation was dominated by the isotope associated with oxidative cleavage of a C–H bond. A similar AKIECl value (1.008) was observed for 1,1,1-TCA oxidation by thermally activated PS from which it was suggested that the first reaction step was the rupture of the C–H bond and the abstraction of the hydrogen atom from the molecule by the attack of any of the radicals formed after persulfate activation. 

The secondary AKIECl estimated in the present experiments (1.0045 ± 0.0004) also points to an oxidative cleavage of a C–Cl bond and thus a primary chlorine kinetic isotope effect is not expected. A reaction pathway involving the cleavage of the C–H as the rate-limiting step is proposed (Figure 3A and SI). In order to track more confidently the proposed mechanism, hydrogen isotope fractionation during CF oxidation with thermally activated PS might be further measured.

During alkaline hydrolysis, CF is abiotically dechlorinated to carbon monoxide and formate. A stepwise elimination mechanism (E1cb) has been proposed for this reaction. This mechanism consists of the rapid, reversible, base-catalyzed deprotonation of the molecule with the formation of a trichloromethyl carbanion (:CCl3−), followed by the rate-determining unimolecular loss of a chloride ion to produce the reactive intermediate dichlorocarbene (:CCl2), which is then rapidly transformed into carbon monoxide and formate (SI Figure S4). If this is the case, the deprotonation step is reversible and the loss of a chloride ion is the rate-determining step, both carbon and chlorine primary isotope effects in the CF molecule are expected during this process. In the present experiments, the AKIECl of 1.061 ± 0.006 was obtained for alkaline hydrolysis, which is consistent with the Streitwieser limit for a primary carbon KIE in C–Cl bonds (1.057, SI Table S2) and is equivalent, within the given 95% CI, to the value previously found by Torrené et al. (1.056 ± 0.003). The AKIECl was calculated as 1.0133 ± 0.0004, which is equal to the maximum expected KIECl for cleavage of a C–Cl bond (1.013, SI Table S2), indicating the involvement of a C–Cl bond. In principle, the Cl kinetic isotope effect estimated in the present study is, therefore, consistent with the occurrence of a carbanion mechanism.
(Figure 3A) but also with a C−Cl bond cleavage via a concerted one-step $S_{n,2}$ nucleophilic substitution mechanism. Nevertheless, based on energy considerations, the 1$\text{E}_{\text{Cl}}$ mechanism seems more plausible for this reaction (see the SI for further discussion). Further deuterium-exchange experiments might be performed to confirm the existence of a carbanion intermediate as a way to further corroborate the occurrence of the stepwise elimination mechanism.$^7$

In the case of reductive dechlorination by Fe(0), an AKIE$_C$ of 1.034 ± 0.012 was obtained, which is similar to the value of 1.030 ± 0.007 obtained by Lee et al.$^29$ and within the AKIE$_C$ range for the reductive cleavage of C−Cl bonds reported in the literature (1.003−1.060) (SI Table S2). In fact, most AKIE$_C$ values for reductive dehalogenation fall in the range of 1.027 and 1.033, which corresponds to about 50% bond cleavage when considering a Streitwieser limit for a C−Cl bond of 1.057 for complete bond cleavage in an infinitely late transition state.$^30$ Regarding AKIE$_{Cl}$, a value of 1.008 ± 0.001 was calculated, which is also about 50% of the Streitwieser limit for KIE$_{Cl}$ in C−Cl bonds (1.013).$^31$ Similar AKIE$_{Cl}$ values, ranging from 1.008 to 1.016 for abiotic reductive dechlorination and from 1.004 to 1.011 for biotic reductive dechlorination, have been previously reported for chlorinated methanes, ethenes and ethanes (SI Table S2). Therefore, both C and Cl-AKIEs pointed to cleavage of a C−Cl bond in the first rate-limiting step, which is compatible with the two-step pathway that is commonly hypothesized for this reaction (see SI). The first step may, for example, involve the transfer of a single electron from the metal surface causing the removal of a chlorine atom and the formation of a dichloromethyl radical (CHCl$_2$) (Figure 3A).

### 3.3. Dual Element Isotope Plot.

Figure 3B shows the dual C−Cl isotope plot for the reactions of this study. A linear correlation between $\Delta$δ$^{13}$C and $\Delta$δ$^{37}$Cl was observed for the three studied transformation mechanisms ($r^2 \geq 0.92$). A comparison of the slopes ($A = \Delta$δ$^{13}$C/ $\Delta$δ$^{37}$Cl) for the regression lines was performed by analysis of covariance (ANCOVA). Statistical significance was accepted at the $p < 0.05$ level. There is no significant statistical difference between oxidation by thermally activated persulfate (17 ± 2) and alkaline hydrolysis (13.0 ± 0.8) slopes (ANCOVA, $p = 0.2$). In contrast, these results differ significantly (ANCOVA, $p < 0.0001$) from the slope observed during CF reductive dechlorination by Fe(0) (8 ± 2).

Hence, although different mechanisms are involved in CF degradation by oxidation by thermally activated PS (cleavage of a C−H bond) and by alkaline hydrolysis (cleavage of a C−Cl bond), the obtained $A$ values for both degradation reactions are not significantly different. This may be explained by considerations for carbon and chlorine isotope effects. Carbon, as expected, the obtained AKIE$_C$ value for CF degradation by oxidation with heat-activated PS is smaller than for alkaline hydrolysis. The higher the mass of the bonding partner, the greater is typically the primary kinetic isotope effect.$^37$ Hence it can be explained that carbon isotope fractionation associated with C−Cl bond cleavage is greater than in C−H bond cleavage since the carbon atom is bound to a heavier atom (chlorine vs hydrogen). Chlorine. This difference, however, is matched by similar differences in chlorine isotope fractionation.

On the one hand, C−Cl bond cleavage involves a primary AKIE$_{Cl}$, which is clearly greater than a secondary AKIE$_{Cl}$ next to a reacting C−H bond. On the other hand, this primary AKIE$_{Cl}$ is "diluted" in $F_{Cl}$ due to the intramolecular competition between three chemically equivalent C−Cl bonds ($\Delta = 3$ in eq S16), whereas the simultaneous secondary AKIE$_{Cl}$ of three Cl atoms are not diluted ($\Delta = 1$). By coincidence, the interplay of these factors results in a similar reduction of carbon as of chlorine isotope fractionation so that similar $\Lambda$ are obtained. This unexpected result restrains the use of C−Cl isotope plots for distinguishing these reactions in the field and highlights the need to apply this approach with precaution and using complementary tools for identification of degradation mechanisms in the field (e.g., complementary hydrogen isotope analysis).

### 4. ENVIRONMENTAL SIGNIFICANCE

Carbon and chlorine isotope fractionation of CF during oxidation with heat-activated PS, by alkaline hydrolysis and by reductive dechlorination with Fe(0) was studied in batch experiments in order to explore the potential of CSIA for the identification of reaction mechanisms in the monitoring of remediation strategies at contaminated sites. For the first time, carbon isotope fractionation values (for heat-activated PS) and chlorine isotope fractionation values (for the three reactions) were determined. These new $\varepsilon$ values increase the options of using CSIA for estimating the extent of contaminant degradation at field sites where remediation strategies are implemented that rely on induced abiotic transformations of CF. Based on the obtained $\varepsilon$ values, it is likely that changes in isotope values in the field may be larger than 2%$^\text{o}$ for carbon and 0.4%$^\text{o}$ (for GC/IRMS instruments) or 2%$^\text{o}$ (for GC/qMS instruments) for chlorine - these are the significant levels that have been suggested as reliable indicators of degradation.$^{34,61}$ Even with the relatively small carbon fractionation obtained for CF oxidation by PS ($\sim 8 \pm 1\%$) and chlorine isotope fractionation observed for Fe(0)-mediated reductive dechlorination ($\sim 3 \pm 1\%$), shifts in CF isotopic composition will be already detectable with a reasonable accuracy if the substrate is degraded by 20% and 25−50%, respectively.

Although only the reductive dechlorination showed significantly statistically different C−Cl isotope slope compared with the other two reactions (oxidation and alkaline hydrolysis), the dual isotope approach might still be used to identify different CF degradation mechanisms in the field, which would (or not) necessarily take place at the same time. For example, the coupling of two common treatments—ISCO and in situ bioremediation—has been shown not only to be feasible, but in many cases also to be able to provide a more efficient and extensive cleanup of contaminated sites.$^{72}$ In the case of PS, the anaerobic environment that is created following the consumption of the oxidant is ideal for CF microbial dehalogenation under sulfate reduction conditions to be enhanced. Enhanced CF bioremediation has also been observed when combining Fe(0) and methanogens that use the cathodic hydrogen generated by iron corrosion for cometabolic degradation of CF$^{53−65}$ or even better by dehalorespiring bacteria which are not inhibited at certain concentrations of CF.$^{29}$ Therefore, there would be an increasing number of case studies, where CF degradation due to PS application or Fe(0) barriers should be distinguished from biotic reductive dechlorination in the field. Although chlorine isotope fractionation during biotic CF dechlorination remains to be evaluated in detail, as well as the effect in CF degradation of Fe(0) aging or the presence of Fe(0) impurities such as graphite,$^{39}$ the dual C−Cl slopes obtained in the present work sets the grounds for the potential application of this approach.
for assessing if CF abiotic reductive dechlorination performed by Fe(0)-PRB or naturally occurring iron-bearing minerals would be or not distinguishable from microalgal reductive dechlorination.

Although CF anaerobic biodegradation has been reported to occur mainly via cometabolic dechlorination or by dehalorespiration, an alternative pathway was suggested in the presence of cobalamins and involving CF hydrolysis. The mechanism of this reaction is not well-known, but it presumably involves the cobalamin-catalyzed conversion of CF to a monochlorocarbene, which would be subsequently hydrolyzed to formaldehyde. The produced formaldehyde could then be oxidized to CO or formate and finally to CO₂. The abiotic alkaline hydrolysis reaction characterized in the present study might be used as a reference system for this suggested CF biotic hydrolysis in future dual isotope-CSIA studies.

Finally, due to the significant difference between the C–Cl isotope slope of CF oxidation and reductive dechlorination, the dual isotope approach might be in addition useful for distinguishing between aerobic and anaerobic CF biodegradation pathways. CF aerobic biodegradation only occurs during oxidative cometabolism with other primary substrates such as methane, butane or toluene by oxygenase-expressing microorganisms. The pathway of CF cooxidation starts by insertion of one oxygen atom into the molecule via H abstraction with phosgene as intermediate and final mineralization to chloride and CO₂. The chemical mechanism of CF oxidation is variable among the various existent monoxygenases, but the rate-limiting step is expected to be the cleavage of the C–H bond.

In conclusion, our study established an expedient base with an overview. Biogeochemistry 2002, 60, 121–139. 
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