Exploring the Utility of Compound-Specific Isotope Analysis for Assessing Ferrous Iron-Mediated Reduction of RDX in the Subsurface

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Abstract

Subsurface contamination with the explosive hexahydro-1,3,5-trinitro-1,3,5-triazine (RDX) at ordnance production and testing sites is a problem because of the persistence, mobility, and toxicity of RDX and the formation of toxic products under anoxic conditions. While the utility of compound specific isotope analysis for inferring natural attenuation pathways from stable isotope ratios has been demonstrated, the stable isotope fractionation for RDX reduction by ironbearing minerals remains unknown. Here, the N isotope fractionation of RDX during reduction by Fe(II) associated with Fe minerals and natural sediments was evaluated and applied to the assessment of mineral-catalyzed RDX reduction in a contaminant plume and in sediment columns treated by *in-situ* chemical reduction. Laboratory studies revealed that RDX was reduced to nitroso compounds without denitration and the concomitant ring cleavage. Fe(II)/iron oxide mineral-catalyzed reactions exhibited N isotope enrichment factors, ε_N , between – $6.3\pm0.3\%$ to $-8.2\pm0.2\%$ corresponding to an apparent ¹⁵N kinetic isotope effect of 1.04–1.05. The observed variations of the δ^{15} N of ~15‰ in RDX from groundwater samples suggested an extent of reductive transformation of 85% at an ammunition plant. Conversely, we observed masking of N isotope fractionation after RDX reduction in laboratory flow-through systems, which was presumably due to a limited accessibility to reactive Fe(II).

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Introduction

The synthetic cyclic N-nitramine munition compound, hexahydro-1,3,5-trinitro-1,3,5-triazine (RDX), has been extensively used in military operations.^{1,2} RDX poses both chronic and acute toxicity threats to aquatic and terrestrial organisms and is considered a possible human carcinogen.^{3–5} The persistence, high solubility, and low volatility of RDX have led to widespread contamination in the subsurface;^{6–9} namely in soil, sediment, and water at military training installations, production facilities, and munition disposal sites.^{10,11} Contamination problems are often aggravated under anoxic conditions through the formation of equally mobile nitroso intermediates, which are more toxic to some terrestrial biota than RDX.^{12,13} Due to the long time-scales over which (bio)degradation of RDX occurs and complicating factors such as multiple contamination sources, dilution and sorption processes, and subsurface heterogeneity, the application of compound specific isotope analysis (CSIA) has been proposed for the assessment of RDX degradation in subsurface environments.^{14,15}

CSIA is used to characterize changes to the stable isotope ratios of elements in nitro-containing munition compounds.^{14–20} These changes are associated with a particular (bio)chemical degradation pathway based on the assumption that kinetic isotope effects for the reacting bonds cause the observable stable isotope fractionation.^{16,19–25} For RDX, variations of ¹⁵N/¹⁴N, ¹³C/¹²C, and ¹⁸O/¹⁶O have been associated with biodegradation under aerobic and anaerobic conditions and with abiotic alkaline hydrolysis reactions in both laboratory and field observations.^{14,15,17,18,26} Assignment of the initial bond cleavage reactions in RDX to the different transformation pathways, however, is particularly challenging regardless of whether transformation occurs biologically or abiotically. The first of two principal routes of RDX transformation, sequential

reduction of $-NO_2$ groups, leads to nitroso-products (Scheme 1, top row). The second route of transformation is denitration (Scheme 1, top row to second row), which can be initiated by reduction, oxidation, or deprotonation of RDX or nitroso intermediates. Denitration gives rise to transient intermediates, triggering the decomposition of RDX into small molecules including NO_2^- , N_2O , NH_4^+ , HCHO, and HCOOH (Scheme 1, second row).²⁷

Anaerobic biodegradation of RDX typically leads to the sequential partial-nitro-group-reduction to nitroso intermediates (hexahydro-1-nitroso-3,5-dinitro-1,3,5-triazine (MNX, Scheme 1), 1,3dinitroso-5-nitro-1,3,5-triazine (DNX), and hexahydro-1,3,5-trinitroso-1,3,5-triazine (TNX).^{18,28} This process is accompanied by an N–O bond cleavage that gives rise to substantial N isotope fractionation with N isotope enrichment factors, ε_N , of -4.7‰ to -9.9‰,¹⁷ whereas C isotope enrichment factors, ε_{C} , are much smaller (-2.7‰).¹⁷ An alternative anaerobic biodegradation pathway involves cleavage of the 1,3,5-triazine ring without formation of similarly large quantities of stable nitroso-intermediates, but with removal of nitrite in the denitration reaction. This causes even larger N isotope enrichment (-12‰) which presumably reflects contributions of both NO₂ reduction and denitration(Scheme 1).^{17,29} Biodegradation under aerobic conditions and alkaline hydrolysis pathways have both been suggested to denitrate RDX to a dinitro-1,3,5triazacyclohex-1-ene intermediate (3 in Scheme 1).^{17,18,26,27,30–32} While the enzymatic reaction involves an oxidation of the methylene carbon, the hydrolysis pathway proceeds as a basecatalyzed deprotonation and elimination of nitrite. These mechanistic differences are also manifested in distinct C and N isotope fractionation behavior, with ε_N between -2.1‰ and -2.4‰ as well as negligible $\varepsilon_{\rm C}$ for enzymatic oxidation vs. $\varepsilon_{\rm N}$ of -7.8‰ and $\varepsilon_{\rm C}$ -5.3‰ for the HNO₂ elimination.17,18,26

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Nitro group reduction



Partial nitro group reduction / denitration



H-atom transfer / denitration



Alkaline hydrolysis (HNO2 elimination)



Scheme 1. Top: Principal routes of RDX transformation through sequential reduction to MNX (hexahydro-1-nitroso-3,5-dinitro-1,3,5-triazine), DNX (1,3-dinitroso-5-nitro-1,3,5-triazine), and TNX (hexahydro-1,3,5-trinitroso-1,3,5-triazine) vs. denitration/ ring cleavage of RDX (or nitroso-intermediates MNX, DNX, TNX) to intermediates including 4-nitro-2,4-diazabutanal (1) and methylenedinitramine (2), formaldehyde, nitrite, ammonia, and nitrous oxide. Bottom four rows: Initial reactions for RDX transformation through nitro group reduction to nitroso compounds and denitration routes initiated by partial nitro group reduction, oxidation (H atom transfer), and alkaline hydrolysis.²⁷

Despite the isotopic elucidation of RDX transformation reactions as well as applications of CSIA to infer the extent of RDX degradation processes at contaminated sites,^{14,15,33} the role of abiotic, mineral-catalyzed reduction of RDX for CSIA has remained largely overlooked. In anoxic groundwater, RDX undergoes abiotic reduction mediated by Fe(II) associated with iron minerals, such as iron oxides, iron sulfides, and clay minerals,^{31,34,35} or Fe(II) produced via *in situ* redox manipulation.³⁶ While nitro group reduction to form nitroso intermediates likely occurs in the anoxic subsurface,³⁵ there is a lack of knowledge regarding N and C isotope enrichment factors to assess this degradation pathway. Previous applications of CSIA to assess RDX transformation in contaminated subsurface environments all focused exclusively on enzyme-catalyzed reactions.^{14,15,33} From a mechanistic perspective, it is also unclear if any isotope fractionation associated with Fe(II)-catalyzed processes is caused by the reduction of the NO₂ functional groups of nitramines, which would cause isotope effects comparable to the abiotic reduction of aromatic NO₂ moieties.^{16,19,20,22,23,37–40} Alternatively, denitration and ring cleavage reactions could also contribute to observable stable isotope fractionation, as has been hypothesized for anaerobic RDX biodegradation.¹⁷

The goal of this work was to assess the utility of CSIA for evaluating abiotic reduction of RDX in anaerobic environments. We hypothesize that the mobility of RDX in the subsurface, its high reactivity towards reduction by mineral-bound Fe(II), and the large isotope effect associated with abiotic NO₂ reductions favor the observation of substantial N isotope fractionation and thus provide a sensitive probe for quantifying abiotic RDX transformation. The specific objectives of this study were to (1) quantify stable isotope enrichment factors for RDX reduction under controlled laboratory conditions for different Fe(II)/Fe mineral combinations, including natural

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sediments, under a range of conditions, (2) assess the contribution of -NO₂ group reduction vs. denitration to N isotope enrichment through a combination of isotopic analyses with a semiquantitative evaluation of reaction product formation, and (3) evaluate the diagnostic value of CSIA-based information for inferring the extent of abiotic RDX reduction. To assess the diagnostic value, we analyzed δ^{15} N trends of RDX in groundwater affected by a contaminant plume from the Iowa Army Ammunition Plant (IAAAP, Middletown, IA). In a second application, we used the CSIA-based approach to monitor RDX transformation in laboratory column reactors where reactive Fe(II) for RDX reduction was generated *in situ* through iron mineral reduction with dithionite.^{20,41}

Material and Methods

Materials

The synthesis of RDX, sources of chemicals and synthetic minerals, and collection and processing of aquifer materials from the Twin Cities Army Ammunition Plant (TCAAP) and soils from Tinker Air Force Base (Tinker AFB) are provided in the Supporting Information (SI).

Batch experiments for evaluating RDX reduction kinetics

All batch experiments were conducted on an end-over-end rotator (Glas-Col) inside an anaerobic glove bag (Coy) with an atmosphere of N₂ (97%)/H₂ (3%). Serum bottles (36 mL) containing 35 mL of deoxygenated (sparged 1 h per L with 99.99% N₂; Matheson) ultrapure water (MilliporeSigma, 18.2 M Ω ·cm), to which the desired minerals and 10 mM NaHCO₃ (to represent the buffer present in groundwater) were added, were prepared in a glove bag. Mineral loadings were 0.5 g/L for goethite, 5 g/L for green rust, 0.5 g/L for hematite, 2 g/L for magnetite, 0.45 g/L

for FeS. Mineral loadings were selected in preliminary experiments to obtain the desired extent of reaction in 2-8 hours. The pH of the solution was adjusted to 6.5, 7.0, or 7.5 using HCl or NaOH. For synthetic Fe(III) minerals (goethite and hematite), the mixed-valent mineral (magnetite), and natural materials (TCAAP sediment and TAFB soil), 1 mM aqueous FeCl₂ was initially supplied in the solution to generate reduction equivalents as Fe(II)/mineral associations.^{35,42} For FeS and green rust, no aqueous Fe(II) was added. The reactors were mixed and rotated at 40 rpm either for 24 h (goethite, magnetite, and hematite, and the natural materials with the Fe(II)) or 1 h (FeS and green rust) prior to initiation of RDX reduction. The amount of Fe(II) adsorbed to the Fe minerals was determined by the difference of the Fe(II) aqueous concentration before and after 24 h equilibration. Reactions were started by adding aliquots from a methanolic RDX stock solution to obtain an initial concentration of $250 \pm 50 \mu$ M. To quantify the kinetics of RDX reduction, 0.5 mL samples were withdrawn from the reactors at selected time intervals, and the suspensions were filtered through 0.2 µm PTFE syringe filters into 2 mL amber vials. At each sampling interval, the aqueous Fe(II) concentration was measured by analyzing 0.1 mL of the filtered aliquot with the ferrozine method.⁴³ Except for FeS and green rust suspensions, the Fe(II) concentration in each reactor was readjusted to 1 mM by adding FeCl₂ from an aqueous stock solution. The pH of each reaction solution was also readjusted to the initial value of 6.5, 7, or 7.5 with NaOH or HCl.⁴² All laboratory reduction experiments were run in duplicate. A set of experiments was performed with 20 µM of TNX under the same reaction conditions using Fe(II)/goethite, Fe(II)/magnetite, and green rust.

Batch reactors for product analysis and CSIA

To identify reaction products from RDX reduction, eight replicate reactors were prepared with Fe(II)/goethite (pH 6.5, 7.0, and 7.5), Fe(II)/magnetite (pH 7.5), green rust (pH 7.5), and FeS (pH 6.5, 7.0, and 7.5) using the procedures above. At each sampling time, a reactor was sacrificed through filtration of 20 mL of the reactor contents followed by acidification of filtrate to pH < 4 with 0.2 mL of 1 M HCl (trace metals purity) for the analysis of RDX, nitroso intermediates, HCHO, and NH₄⁺.

Samples for CSIA of RDX were prepared in sets of eight replicate reactors that were sacrificed at different time points. One additional set of reactors for the Fe(II)/goethite system at pH 7.5 was used to evaluate effects of natural organic matter on RDX isotope fractionation to which Elliot Soil Humic Acid (ESHA) was added to 10 mg-C/L together with the Fe(II). After filtering and acidifying the contents of each reactor, a subsample was taken to measure the remaining RDX concentration by HPLC. The remaining volume of filtered, acidified samples was then extracted via solid phase extraction (SPE).¹⁷ Briefly, Supelclean™ ENVI-Chrom SPE cartridges (Sigma Aldrich) were conditioned sequentially with 3 mL of ethyl acetate, methanol, and ultrapure water prior to loading 20 mL of sample onto each cartridge by gravity. The cartridges were then vacuum-dried and eluted with two 3 mL portions of ethyl acetate, such that minimal water was present in the eluent. Samples were then dried with sodium sulfate, filtered with Teflon syringe filters (0.2 µm), and evaporated to 0.2 mL for stable isotope analyses.

Extraction of groundwater samples and preparation for CSIA

Seven groundwater samples (Figure S1) were collected from monitoring wells along a subsurface plume at the Line 800 site of the Iowa Army Ammunition Plant (IAAAP,

Middletown, IA). Each sample contained approximately 4 L of groundwater. Details of the well locations, sampling method, and water chemistry are described in the SI (Section S2). Concentrations of RDX and the nitroso intermediates measured via gas chromatography-mass spectrometry from separate samples collected at the same time were provided by Jacobs Engineering Group (Table S1). SPE in preparation for isotope analysis was performed as described above, except the sample from each well was split into three ~1.3 L portions that were passed through a separate SPE cartridge. After elution, the three extracts were combined and processed as described below.

Reduction of RDX via in situ chemical reduction (ISCR) of a sediment packed column

The column setup and ISCR procedure followed the protocols of Berens et al.²⁰ Briefly, a borosilicate glass column (Kimble FLEX-COLUMNS®, I.D. 2.5 cm) was wet-packed with 4.8 cm (~39 g) of TCAAP sediment inside an anaerobic glove box. A flow adapter (Kimble CHROMAFLEX®, I.D. 2.5 cm) was secured to the end of the column to prevent sediment migration and leaching. The column was then saturated with a 10 mM NaCl solution to evaluate the total pore volume, porosity, density, and other physical properties of the column (Table S2). To minimize particle release/transport during the column experiments, the ionic strength of all further feed solutions was adjusted by adding 10 mM NaCl.²⁰ A sequence of three experiments at constant volumetric flow of 0.5 mL/min was conducted unless indicated otherwise. First, a 0.115 M NaBr tracer solution was introduced in a step input to determine hydraulic properties of the column (Table S2). Tracer concentrations were measured with a bromide ion selective electrode (Thermo Scientific Orion) after a 3-fold dilution of the samples. Thereafter, the column was conditioned with 10 mM NaHCO₃ buffer for 10 pore volumes to remove the residual NaBr. The

second experiment consisted of an RDX pulse from a 500 mL reservoir with 200 μ M of RDX in 10 mM NaHCO₃ buffer at pH 7.5. The breakthrough of RDX through the column with untreated aquifer material was used to determine the extent of RDX sorption in the column. After flushing the column with NaHCO₃ buffer for 10 pore volumes, the TCAAP sediment (3.0% total iron) was then reduced by feeding a combined solution of 5.5 mM Na₂S₂O₄ and 22.2 mM K₂CO₃ into the column at 0.25 mL/min for ~18 hours. The exact duration of Na₂S₂O₄ input during each reduction cycle was selected to target a reduction of one-tenth of the total iron content. This was determined using eq 1 which shows that 2 mols of Fe(II) are generated by every 1 mol of S₂O₄²⁻.

$$S_2 O_4^{2-} + 2Fe^{3+} + 2H_2 O \to 2Fe^{2+} + 2SO_3^{2-} + 4H^+$$
(1)

As shown previously by Berens et al.,²⁰ changes of hydraulic properties after dithionite treatment were smaller than variations of porosities and dispersion coefficients among different columns, and iron reduction was incomplete either due to inaccessibility or reactivity constraints. In the third series of experiments, three pulses of RDX (210 μ M, 36 pore volumes) were delivered to the column followed by ~10 pore volumes of NaHCO₃ buffer to remove any residual RDX and then Na₂S₂O₄ and K₂CO₃ to re-reduce the Fe in between each of the cycles.

Column effluent for RDX concentration and stable isotope analysis was collected into test tubes every 0.6 pore volumes using a Bio-rad fraction collector. The sample collection was terminated when effluent RDX concentration attained the influent values or remained constant. The samples were immediately capped for quantification of RDX and nitroso intermediates. For RDX isotope ratio measurements, four consecutive samples were combined to generate sufficient volume (20 mL). The combined samples were re-analyzed to determine the RDX and then processed by SPE as described above.

Analytical methods

Aqueous concentrations of RDX, MNX, DNX, TNX were measured by high pressure liquid chromatography (HPLC) with UV-detection (details in Section S3).⁴² Formaldehyde⁴⁴ and ammonium⁴⁵ were quantified after derivatization by liquid chromatography with UV and fluorescence measurement, respectively (Section S3).

¹⁵N/¹⁴N ratios of RDX were determined by large volume injection coupled to gas chromatography isotope ratio mass spectrometry (GC/IRMS) consisting of a Trace GC and a Delta V IRMS (Thermo Scientific). Analyses followed procedures established previously^{17,19,26} and instrumental parameters are given in the SI. Method quantification limits were derived with the moving mean approach⁴⁶ and corresponded to injection of 3 nmol N (Figure S2a, details in SI). The δ^{15} N of RDX agreed well with typical N isotope signatures determined for RDX from different manufacturers and from various type of explosives.⁴⁷

Measurement of ${}^{13}C/{}^{12}C$ ratios was strongly amplitude-dependent and did not allow for an accurate determination of $\delta^{13}C$ for RDX with our standard material (Figure S2b). Due to this finding as well as the observation of chromatographic interferences of unknown, carbon-containing compounds in samples from RDX reduction experiments and in field samples, no $\delta^{13}C$ are reported here.

Analysis of reaction kinetics and predominant RDX reaction pathways

Pseudo-first order rate constants of RDX degradation were determined by linear regression of $ln(c/c_0)$ vs. time (t) for initial experiments, and non-linear fitting when considering RDX degradation and formation/decay of intermediates and products. Reported errors are 95%

confidence limits propagated from the errors associated with duplicate regressed rate constants. Reaction pathways of RDX were assessed on a carbon mass basis to account for any formation of denitration/ring cleavage products during the reduction of RDX (Schemes 1).³¹ Concentrations of the carbon-containing analytes RDX, MNX, DNX, TNX, and formaldehyde, as well as a hypothetical C-containing product (C-Unidentified) were used to quantify losses of total C as the reduction of RDX proceeded. The model was expressed as sets of differential equations, with different sets of equations used to test whether ring cleavage occurred in RDX or the nitroso compounds (Section S4). The assumption was used that all reactions followed pseudo-first order kinetics, with reductant concentration being constant. Differential equations were solved in MATLAB (MathWorks, Version 9.5) using the initial C-based concentration of RDX (C-RDX) and fit to experimental data via a least-squares regression.

Modeling RDX reactive transport in dithionite-treated sediment columns

Simulations of RDX and Fe(II) concentrations in sediment columns were performed with Aquasim (Version 2.0) using the saturated soil-column compartment.⁴⁸ Model parameters for the different series of experiments are listed in Tables S2 and S3 (Section S5). Sediment porosity and dispersivity were determined from the NaBr tracer experiment. Sorption of RDX to sediment column material was quantified based on its retardation relative to the non-sorbing NaBr tracer assuming linear sorption behavior and instantaneous localized sorption equilibrium according to eqs. 2 and 3,

$$\frac{d[RDX]_s}{dt} = -k_{sorption} \cdot \left([RDX]_{s,eq} - [RDX]_s \right)$$
(2)

$$K_d = \frac{[RDX]_{s,eq}}{[RDX]_{aq}} \tag{3}$$

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where k_{sorption} is the rate constant of the adsorption process, and $[\text{RDX}]_{s,eq}$ and $[\text{RDX}]_{aq}$ are the concentration of adsorbed RDX in equilibrium and aqueous, dissolved species, respectively. K_d is the equilibrium sorption coefficient of RDX. RDX reduction by Fe(II) in sediment columns after ISCR was assessed with eqs. 4 and 5,

$$\frac{d[RDX]_{aq}}{dt} = -v_{RDX} \cdot k_{red} \cdot [RDX]_{aq} \cdot [Fe^{II}]_s$$
(4)

$$\frac{d[Fe^{II}]_s}{dt} = -v_{red-equiv} \cdot k_{red} \cdot [RDX]_{aq} \cdot [Fe^{II}]$$
(5)

where v_{RDX} is the solid-water ratio used for conversion of sorbed to aqueous concentrations, and k_{red} is the apparent RDX reduction and Fe(II) oxidation rate constant. The $v_{red-equiv}$ parameter describes the amount of Fe(II) oxidized per amount of RDX reduced and was also used for a qualitative assessment of Fe(II) availability for reductive RDX transformation after ISCR treatment, and the consequences thereof for the N isotope fractionation of RDX.

Analysis of CSIA data

Nitrogen isotope enrichment factors, ε_N , in batch reactor experiments were evaluated according to the methods described by Pati et al.²¹ from the natural log-linearized form of the correlation between measured isotope ratio and the fraction of remaining RDX. Apparent ¹⁵N-kinetic isotope effects (¹⁵N-AKIE) were derived from ε_N by accounting for the 6-fold isotopic dilution caused by the 6 N atoms in RDX. The fractional extent of RDX transformation, *F*, during abiotic reduction at the Line 800 site of the IAAAP was calculated with eq. 6, where $\delta^{15}N_x$ and $\delta^{15}N_0$ are the RDX N isotope signatures measured at sampling locations *x* and at the operationally defined contamination source.

$$F = 1 - \left(\frac{\delta^{15} N_x + 1}{\delta^{15} N_0 + 1}\right)^{1/\varepsilon_N} \tag{6}$$

Values of δ^{15} N during experiments in dithionite-treated sediment columns were also evaluated for possible ¹⁵N isotope effects.

Results and Discussion

Kinetics and Pathways of RDX reduction

RDX was reduced in all of the tested suspensions as depicted in Figure 1a for Fe(II)/goethite at pH 7 and Figure S3 for Fe(II)/goethite, Fe(II)/magnetite, green rust, and FeS. The pseudo-first order rate constants for RDX reduction by iron minerals in the initial batch experiments are listed in Tables S4 (RDX) and S5 (TNX), and those from the experiments to evaluate product formation are in Table S6. The results are similar, and any differences were attributed to the linear vs non-linear fitting and experimental error. Consistent with past results,^{35,49,50} reaction rates increased with increasing solution pH for all minerals tested. Fe(II)/goethite rate constants were 0.13 ± 0.03 , 0.38 ± 0.16 , and 1.24 ± 0.28 h⁻¹ at pH 6.5, 7, and 7.5, respectively (Table S6), consistent with increased adsorption of Fe(II) on the iron (oxy)hydroxides (the adsorption of Fe(II) by goethite was 0.15 ± 0.07 , 0.28 ± 0.08 , 0.39 ± 0.07 mmol Fe (II)/g goethite at pH 6.5, 7, and 7.5) and more favorable reduction potentials for Fe(II)/mineral systems with increasing pH.⁴⁹

Previously, no degradation of RDX or DNAN by TCAAP sediment (containing predominantly magnetite, 3.03%-wt Fe) was observed in the absence of aqueous Fe(II).^{20,42} Thus, aqueous Fe(II) was constantly supplied to TAFB (containing predominantly hematite, 2.44%-wt Fe) and TCAAP reactors in the same manner as for synthetic materials. The lower rate constants

observed at similar or greater iron mineral loadings for these materials (Table S4) suggests the formation of Fe(II) associations with less reactive minerals than iron oxides (e.g., silicates).^{42,51}



Figure 1. Concentration versus time data for RDX, intermediates, and end products during abiotic RDX reduction in the Fe(II)/goethite reactors at pH 7. (a) Measured concentrations of RDX, nitroso intermediates and final degradation products, (b) total carbon mass balance (initial C from RDX = 600 μ M), (c) concentration of RDX, nitroso intermediates, and products plotted in μ M as carbon (denoted as C-compound). A hypothetical carbon-containing product (C-Unidentified; black solid circle) was introduced to compensate for the incomplete carbon mass balance and was assumed to be a single reaction product in kinetic fitting.

Different minerals can react with RDX through different and/or multiple transformation pathways (Scheme 1). An interpretation of the observable stable isotope fractionation (see below) thus requires an assessment of the principal pathways. For all minerals tested, MNX, DNX, and TNX were observed, indicating reduction of the nitro groups on RDX. Additionally, RDX and/or the nitroso compounds must be reacting via ring cleavage, given the formation of HCHO and a decreasing overall carbon mass balance (Figures 1 and S3). The molar mass balance for each experimental system was converted to a carbon-based mass balance as illustrated in Figure 1b for Fe(II)/goethite at pH 7. C-Unidentified was considered as a single, final product because carbon deficits began during late stages of the reaction (Figure 1b) and C-Unidentified and HCHO appear in parallel. Methylenedinitramine (**2** in Scheme 1) is unlikely responsible for the carbon deficit because it undergoes instantaneous transformation to nitramine, HCHO, N₂O, and NH₄⁺ under anaerobic conditions.³¹ C-Unidentified could be one or more carbon-containing products, but from the prospective pathways, this portion of the mass is an end-product.

Based on the concentration of carbon-containing species during batch experiments (Figures 1 and S3), potential reaction pathways were tested through kinetic modeling using the equations in Section S4 with the assumption that the nitroso intermediates were sequentially produced $(RDX \rightarrow MNX \rightarrow DNX \rightarrow TNX)$. The differences among the potential reaction pathways (Section S6) were the species from which C-HCHO and C-Unidentified were formed. The formation of these final products was considered to occur from RDX, one of the nitroso compounds, or a combination thereof. The experimental data and the rate constants given by the model with the lowest normalized residuals for the tested pH/mineral combinations are summarized in Figure S3 and Table S6. For Fe(II)/goethite systems, reactions of MNX, DNX, and TNX were more rapid than reaction of RDX. As illustrated in Table S6, the rate constants for the transformation of DNX to both HCHO and C-Unidentified were twice as high as those for RDX or MNX. This modeling outcome is consistent with the limited observed accumulation of transient MNX, DNX, and TNX species. The delayed formation of HCHO and C-Unidentified (Figure 1c) indicates these ring cleavage products form from the nitroso intermediates. For the iron oxides, the data are best fit by Scheme S1a (Tables S6-S8), where HCHO and C-Unidentified are formed from DNX and/or TNX. For FeS, the slower degradation of MNX and accumulation of DNX and TNX resulted in the best fit if ring cleavage of RDX and/or MNX to form HCHO and C-Unidentified were included in the kinetic model (Scheme S1b; Tables S6 and S9).

The goodness of fit for other proposed pathways for the iron oxides and FeS, which had substantially higher residuals, is shown in Tables S7-S9. The results indicate that for the iron oxides, the slowest reaction during the sequential transformation of RDX to TNX is to the formation of MNX from RDX. Moreover, the reaction of RDX only occurs via nitro group reduction, allowing for a mechanistic interpretation of the RDX CSIA results. For FeS, RDX appears to react via a combination of reduction and dinitration/ring cleavage. The slower reaction of the nitroso compounds with FeS leads to their accumulation and affects isotopic analyses (see below).

RDX reduction evaluated by CSIA

Nitrogen isotope fractionation during RDX reduction was assessed for each iron mineral suspension. As shown in Figure 2 and Table 1, the extent of N isotope fractionation was large (up to 30‰) and the N isotope enrichment factors, ε_N , were confined to values between $-6.3 \pm 0.3\%$ and $-8.1 \pm 0.2\%$. Small changes of pH by 0.5 units or the presence of organic matter did not lead to changes of N isotope fractionation in RDX. The identical behavior was observed for Fe(II)-amended TCAAP sediment and TAFB soil ($\varepsilon_N = -8\%$) implying the same abiotic reduction reaction of RDX by mineral associated Fe(II) in all batch experiments. Initial analyses showed that the nitroso intermediates were also retained by SPE and co-eluted with RDX on the GC column, thus introducing interferences to the instrumental signal. Thus, experiments with Fe(II)/goethite at pH 6.5 and with FeS were not included. As a consequence, we were also not able to gather additional insights into Scheme S1b by CSIA from reactors containing FeS.



Figure 2. Nitrogen isotope signatures, δ^{15} N, versus fraction of unreacted RDX (c/c₀) determined in experiments with various Fe(II) containing suspensions. The goethite data represent experiments with Fe(II)/goethite at pH 7.0 and pH 7.5, and Fe(II)/goethite with ESHA at pH 7.5, and all other experiments were carried out at pH 7.5. The curve was fit using the data from each set of conditions to obtain a single ε_N value of -7.4±0.2‰ (individual values in Table 1). The error band shows the 95% confidence interval.

Similar ε_{N} -values of $-9.9 \pm 0.7\%$ were reported for anaerobic RDX biodegradation that was initiated by reduction of a NO₂ moiety,¹⁷ whereas other biological and abiotic transformations of RDX under aerobic conditions, such as denitration or alkaline hydrolysis, led to much smaller ε_{N} -values (Figure S4, Scheme 1).^{17,18,26,28} Based on the investigation of different anaerobic biotransformation pathways, Fuller et al.¹⁷ hypothesized that N isotope fractionation is even larger if denitration and concomitant ring-cleavage reactions were responsible for RDX disappearance. In this interpretation, bond-cleavage reactions would generate cyclic nitramine-N species and give rise to stronger N isotope fractionation with ε_N -values as negative as -12%. Considering the slightly less negative ε_N shown in Table 1, these findings indirectly confirm the interpretation that the initial steps of Fe(II)-catalyzed reduction of RDX are confined to reduction of the NO₂ moiety. This interpretation explained the observed ¹⁵N-AKIE values were between 1.039 and 1.051 for this reaction. These isotope effects are approximately 20% larger than most of those reported for the reduction of nitroaromatic compounds studied in identical experimental systems (¹⁵N-AKIEs <1.040)¹⁹ as well as by other abiotic reductants^{16,39,40} even though ¹⁵N-AKIEs of 1.048 were also found for aromatic NO₂ reduction by stoichiometric magnetite.³⁸ Our data set adds to previous observations¹⁷ that the AKIEs for reduction of NO₂ moieties of cyclic nitramines may exceed those of aromatic NO₂ groups.

Mineral	ϵ_{N}^{a} (%)	¹⁵ N-AKIE (-)	εc ^a (‰)	¹³ C-AKIE (-)
Green rust	$\textbf{-6.9}\pm0.8$	1.043 ± 0.005	ND^{b}	ND
Hematite	-8.1 ± 0.2	1.051 ± 0.001	ND	ND
Magnetite	-6.3 ± 0.3	1.039 ± 0.002	ND	ND
Goethite pH 7	-7.7 ± 0.4	1.049 ± 0.001	ND	ND
Goethite pH 7.5	-7.3 ± 0.3	1.046 ± 0.002	ND	ND
Goethite pH 7.5-NOM	$\textbf{-6.3}\pm0.6$	1.039 ± 0.003	ND	ND
TAFB soil	-7.9 ± 0.4	1.049 ± 0.002	ND	ND
TCAAP sediment	$\textbf{-8.2}\pm0.2$	1.050 ± 0.001	ND	ND
Anaerobic biodegradation ^c	-9.9 ± 0.7	1.063 ± 0.005	-4.7 ± 1.1	1.005 ± 0.001
Anaerobic biodegradation ^d	-5.0 ± 0.3	1.031 ± 0.002	NA^{g}	NA
Aerobic biodegradation ^e	$\textbf{-2.3}\pm0.5$	1.006 ± 0.003	-0.8 ± 0.5	1.001 ± 0.001
Hydrolysis ^f	-5.3	1.033	-7.80	1.008

Table 1. Bulk N isotope enrichment factors ε_N and ε_C during abiotic reduction, biodegradation, and hydrolysis of RDX

^a bulk N isotope enrichment factor.

^b ND = no data. Analysis for C was affected by interferences. See SI.

^c composited isotope data from RDX anaerobic degradation by multiple anaerobic strains via nitro reduction.¹⁷

^d RDX anaerobic degradation by non-specific sediment strains via nitro reduction.¹⁸

^e composited isotope data from RDX aerobic degradation by multiple anaerobic strains via denitration. ¹⁷

^f isotope data from RDX hydrolysis via ring cleavage. ²⁶

^gNA=data not available

Using CSIA to evaluate RDX transformation in groundwater samples from an ammunition plant

CSIA was applied to evaluate RDX transformation in seven monitoring wells along Line 800 of a subsurface plume at the IAAAP (Figure S1). Concentrations of RDX ranged from <0.01 µM to 29.3 µM and decreased radially from sampling point G-20, where the highest value was measured. Samples from only four monitoring wells exceeded concentrations of 0.1 µM and were suitable for N isotope ratio measurements of RDX. These wells also had low dissolved oxygen, consistent with reducing conditions (Table S1). The G-20 sample was assumed as the reference point for CSIA of RDX because of its upstream location along the hydrologic gradient.^{52,53} It is in close proximity to the source of the plume⁵⁴ and coincides with the highest aqueous RDX concentration and smallest $\delta^{15}N$ (+0.66‰). This operational $\delta^{15}N_0$ is somewhat higher than data from the only study of manufactured RDX, which range from -17% to -4% depending on synthesis method and raw materials.⁴⁷ The δ^{15} N of G-20 could reflect some degradation of RDX given the historic concentrations of RDX >50 µM reported for 1990-2000.54 Figures 3a and b show that decreasing RDX concentrations in the sampling wells considered here also correspond with δ^{15} N values increasing by ~15‰. The presence of typical RDX reduction products, MNX, DNX, TNX in concentrations <1.0 µM (Figure 3c) supported the assumption of reductive RDX transformation in the subsurface.



Figure 3. Evaluation of RDX concentrations and δ^{15} N values with regard to reductive transformation in monitoring wells along Line 800 at IAAAP (map see Figure S1). The samples are sorted according to increasing distance from the operationally defined contamination source well G-20. (a) Aqueous concentrations of RDX at different sampling locations including calculated extent of RDX reduction (eq. 6) based on the ϵ_{N} -value obtained in laboratory batch experiments with Fe-minerals. (b) δ^{15} N values of RDX at different sampling locations. (c) Concentrations of partially reduced RDX reduction products.

Based on the detection of RDX reduction products typically associated with abiotic reduction, we evaluated the extent of RDX transformation relative to the δ^{15} N values measured from sampling well G-20 and the average isotope enrichment factor obtained from the batch experiments ($\epsilon_N = -7.4\%$). Figure 3a shows that calculated RDX concentrations after abiotic reduction with this ϵ_N -value range between 5 and 15 μ M, compared to measured concentrations of 0.1, 0.8, and 3 μ M. These predicted concentrations correspond to an extent of reductive transformation of 54% to 85%. If abiotic reduction was the predominant mode of RDX transformation, our data would imply that the measured, lower concentrations were also the consequence of other non-isotope fractionating processes (e.g., sorption, dilution, volatilization). Given that RDX sorption to the solid matrix is likely negligible (see column study below) and that RDX is largely non-volatile we hypothesize that the observed decreases in concentration were due to dilution.

CSIA-based evaluation of RDX reduction in sediment columns after ISCR treatment

The breakthrough of RDX in columns containing TCAAP sediment before and after the first of three dithionite-treatments during ISCR are shown in Figure 4a. RDX retention was only slightly retarded compared to the NaBr tracer (Figure S5, retardation factor 1.17), indicating minimal sorption of RDX to the sediment matrix. This observation is consistent with historic field observation of high RDX mobility in the subsurface of contaminated sites.⁵⁵ By contrast, RDX transport through dithionite-treated TCAAP sediment was substantially delayed, with complete breakthrough after 30 pore volumes (Figure 4a). The detection of the RDX reduction products MNX and DNX as well as previous observations of DNAN reduction to phenylenediamine in identical experiments, implies that apparent RDX retardation after the first ISCR cycle was due to reduction by ISCR-generated Fe(II). Because of the numerous reaction products formed from RDX reduction including ring-cleavage products that were, in part, not detected, no mass or electron balance was computed. The identical behavior of RDX breakthrough was observed after the second ISCR cycle (Figure S6) whereas RDX concentration only reached 60% of the input concentration after ISCR cycle 3 (see below).

The breakthrough of RDX in sediment columns after ISCR could not be rationalized with the simple reactive transport model described by eqs. 2-5 (Section S8), which assume localized sorption equilibrium and that reduction of RDX leads to the corresponding removal of solid-associated Fe(II). As a simplified means of qualitatively assessing RDX breakthrough behavior,

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Figure 4 (a) RDX concentrations at the in- and outlet of columns filled with native TCAAP sediment and after the first of three ISCR cycles. Note the different y-axes for species concentrations. Solid lines represent calculated RDX breakthrough curves, and calculations for breakthrough following ISCR reflect manipulation of $v_{red-equiv}$ to account for the observation of apparent overstochiometric removal of Fe(II) by RDX. MNX and TNX were detected only after ISCR. (b) δ^{15} N values and concentrations of RDX measured in the effluent of sediment columns after ISCR. The lines represent the calculated δ^{15} N values of RDX based on parameters listed in Table S3 and different ¹⁵N-AKIE values for the abiotic reduction of RDX. The ¹⁵N-AKIE value of 1.04 represents the N isotope fractionation observed during the RDX reduction in batch experiments whereas a value 1.002 reflects the N isotope fractionation during reactive transport.

we manipulated the value of $v_{red-equiv}$ to reflect the observation of apparent highly overstochiometric removal of Fe(II) by RDX (Table S3). This apparent parameter exceeded the known amount of reduction equivalents necessary to transform RDX to TNX by a factor of >10 and for complete reduction of all N from RDX to ammonia by a factor of >2.5 (Table S2, Figures S7-S12). Based on these observations, we speculate that the oxidation of Fe(II) by the RDX front caused a limitation of the available Fe(II) for RDX reduction due to formation of aggregates of lower hydraulic conductivity,^{56,57} the generation of phases of lower Fe(II) reactivity,⁵⁸ or a combination thereof. No characterization of the column material after the Fe reduction/reoxidation cycles was performed.

The measured δ^{15} N values of RDX at the column outlet after ISCR cycles 1 and 2 are shown in Figures 4b and S13. δ^{15} N values of RDX were between -4‰ to -7‰ at early stages of the breakthrough because RDX was only partially reduced. δ^{15} N of RDX reached its original value of -9.5‰ once the RDX outlet and input concentrations matched. The extent of N isotope fractionation (~5‰) observed in column experiments was substantially smaller than expected based on the isotope enrichment factors and AKIEs determined in batch experiments as well as that observed at IAAAP groundwater sampling wells (~15‰). The lines in Figure 4b show the calculated δ^{15} N of RDX based on assumptions for different ¹⁵N-AKIEs. An AKIE of 1.04 would correspond to the N isotope fractionation observed in batch experiments. Instead, we find a 20fold smaller fractionation as expressed by a ¹⁵N-AKIE of 1.002 that would best describe our data. This observation, as well as evidence for limited accessibility of RDX to Fe(II) described above, are consistent with substantially masked N isotope fractionation. Modeling approaches incorporating corrections to enrichment factors for potential mass transfer limitations, as demonstrated for diffusion-dominated vapor phase transport,⁵⁹ could potentially offer improved interpretation of observed masking. We maintain, however, that our simplified modeling approach is justified, considering that the column experiments are just one facet to the much broader scope of this study.

Environmental Implications

Results from our laboratory experiments show that abiotic reduction mediated by various iron minerals with Fe(II) gives rise to consistent patterns of N isotope fractionation of RDX due to large ¹⁵N kinetic isotope effects, which often exceed those observed for the same reactions of nitroaromatic compounds. These large isotope effects have two practical consequences. First, shifts of δ^{15} N in RDX can be large as observed in the RDX contaminant plume at the Iowa Army Ammunition Plant. This is consistent with similar extents of N isotope fractionation reported for RDX and other nitro-containing explosives such as TNT and DNT at other contaminated sites.^{16,33} Second, large ε_N values, such as those reported for reductive RDX transformations, lead to conservative estimates of the fractional amount of contaminant conversion. Together with previous evaluations of isotope fractionation associated with biological RDX transformation under anaerobic and aerobic conditions,^{17,18,26,28} our work contributes to an assessment of the relevant RDX degradation pathways via CSIA on the basis of well-defined isotopic enrichment factors.

Experiments in a flow-through dithionite-treated sediment column point to a potential masking of strongly isotope fractionating reactions. The experimental conditions applied to study the RDX breakthrough, namely high RDX concentration pulses to exhaust the available reduction

equivalents, are likely not representative of contaminant-to-reductant ratios in the subsurface. Nevertheless, our data imply that application of CSIA to assess RDX reduction after redoxmanipulation requires further scrutiny to provide equally realistic estimates of transformations as found for DNAN in identical systems.²⁰ This issue is aggravated by the frequently observed challenges of determining ¹³C/¹²C ratios of RDX reliably by CSIA^{17,26,33} that, contrary to CSIA studies for many other subsurface contaminants, makes the identification of RDX reaction pathways through multi-element isotope analysis particularly challenging.

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Supporting Information

Supporting information (SI) available: A detailed report of materials, additional analytical methods, sampling, modeling approaches, kinetics information, and further isotope results.

Competing Interests

The authors declare no competing interests.

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