

1 **Hexahydro-1,3,5-trinitro-1,3,5-triazine (RDX) Degradation by**  
 2 ***Acetobacterium paludosum***

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8 **Key words:** ammonium, autotrophic, homoacetogens, hydrogen, nitrous oxide, RDX

9 **Abstract**

10 Substrates and nutrients are often added to contaminated soil or groundwater to enhance bioremediation.  
 11 Nevertheless, this practice may be counterproductive in some cases where nutrient addition might relieve  
 12 selective pressure for pollutant biodegradation. Batch experiments with a homoacetogenic pure culture of  
 13 *Acetobacterium paludosum* showed that anaerobic RDX degradation is the fastest when auxiliary growth  
 14 substrates (yeast extract plus fructose) and nitrogen sources (ammonium) are not added. This bacterium  
 15 degraded RDX faster under autotrophic (H<sub>2</sub>-fed) than under heterotrophic conditions, even though het-  
 16 erotrophic growth was faster. The inhibitory effect of ammonium is postulated to be due to the repression  
 17 of enzymes that initiate RDX degradation by reducing its nitro groups, based on the known fact that  
 18 ammonia represses nitrate and nitrite reductases. This observation suggests that the absence of easily  
 19 assimilated nitrogen sources, such as ammonium, enhances RDX degradation. Although specific end  
 20 products of RDX degradation were not determined, the production of nitrous oxide (N<sub>2</sub>O) suggests that  
 21 *A. paludosum* cleaved the triazine ring.

22 **Abbreviations:** RDX – hexahydro-1,3,5-trinitro-1,3,5-triazine

24 **Introduction**

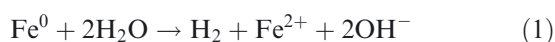
25 The explosive RDX (hexahydro-1,3,5-trinitro-1,3,5-  
 26 triazine) is a toxic and persistent groundwater  
 27 contaminant found at many military installations  
 28 (Held et al. 1997; Schmelling et al. 1997). The U.S.  
 29 EPA has classified RDX as possible human car-  
 30 cinogen, and RDX is also toxic to the neurosystem  
 31 and to other mammals, algae, invertebrates, and  
 32 fish (McLellan et al. 1992; Testud et al. 1996).  
 33 Possibly more toxic than RDX are some potential  
 34 degradation metabolites, such as the nitroso  
 35 heterocyclic compounds MNX (1,3-dinitro-5-nitr-  
 36 oso-1,3,5-triacyclohexane), DNx (1,3-dinitroso-5-  
 37 nitro-1,3,5-triazacyclohexane), and TNx  
 38 (1,3,5-trinitroso-1,3,5-triazacyclohexane), as well

as potential ring fission products 1,1- and 1,2-  
 39 dimethylnitrosamine, azoxymethane, and  
 40 hydrazine, which are known to be mutagens, car-  
 41 cinogens, or both (Fiala 1977; Greenhouse 1976;  
 42 McCormick et al. 1981; Skopek et al. 1978). The  
 43 toxicity of RDX and its potential metabolites is a  
 44 major driving force for the remediation of con-  
 45 taminated sites.

46  
 47 One emerging strategy that holds great poten-  
 48 tial for treating RDX-contaminated groundwater  
 49 is the use of zero-valent iron (Fe<sup>0</sup>) in permeable  
 50 reactive barriers (PRBs) (Hundal et al. 1997).  
 51 Previous studies showed that indigenous aquifer  
 52 microorganisms or mixed cultures from anaerobic  
 53 digesters can enhance both the rate and extent of  
 54 RDX transformation in Fe<sup>0</sup> systems (Oh et al.

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2001; Wildman & Alvarez 2001). This enhancement was postulated to be due, in part, to cathodic hydrogen production during anaerobic Fe<sup>0</sup> corrosion:



Apparently, hydrogen has a biostimulatory effect and is used as an electron donor to drive the bacterial reduction of RDX (Adrian et al. 2003; Beller 2002). Hydrogen is also a common electron donor in anaerobic systems, which adds relevance to the study of RDX degradation by hydrogenotrophs.

Among the potential hydrogen-utilizers that could enhance RDX removal are homoacetogenic bacteria. Homoacetogens are strict anaerobes that can use H<sub>2</sub> and CO<sub>2</sub> for growth and have been found to colonize the Fe<sup>0</sup> layer in flow-through columns treating RDX (Oh & Alvarez 2002). In theory, homoacetogens could also commensally support heterotrophic activity in anaerobic systems by producing acetate. Increased heterotrophic activity due to higher availability of such a C source might be beneficial for RDX removal, especially if RDX is utilized as an N source by heterotrophs. Homoacetogens have also been implicated in RDX degradation by methanogenic sludge (Adrian & Lowder 1999) and other mixed cultures (Oh & Alvarez 2002), and recently, a pure homoacetogenic culture that degrades RDX was isolated (Adrian & Arnett 2004). However, the ability of homoacetogens to degrade RDX under autotrophic (H<sub>2</sub>-fed) and nitrogen-deficient conditions that are likely to be encountered in groundwater as well as in and around Fe<sup>0</sup> barriers has not been previously reported.

Adding nutrients such as ammonia to contaminated sites or bioreactors is a common biostimulation practice. However, nutrient addition can have a detrimental effect if it inhibits bacteria adapted to oligotrophic environments (Morgan & Watkinson 1992) or if it stimulates the degradation of carbon compounds other than the target pollutants. Whether ammonium enhances or hinders RDX degradation by homoacetogens has not been previously investigated.

This paper is the first to report RDX degradation by the homoacetogenic species *Acetobacterium paludosum*. Emphasis was placed on (1) comparing RDX degradation under heterotrophic versus autotrophic conditions; (2) characterizing

RDX degradation rates and products (including the potential for RDX mineralization); and (3) determining the effect of an easily assimilated nitrogen source, such as ammonium, on RDX degradation. This information contributes to our understanding of microbial niches in RDX contaminated environments.

## Materials and methods 112

### Culture conditions 113

*Acetobacterium paludosum* (ATCC # 51793), isolated by Kotsyurbenko, et al. (Kotsyurbenko et al. 1995) from sediment of a marsh 100 km north of Moscow, Russia, was utilized because of its ability to grow at environmentally relevant temperatures (≤20 °C) as well as its ability to be cultured more easily than other homoacetogenic bacteria (Sherburne 2003). Bacteria were routinely cultured in closed 25 ml Balch anaerobic culture tubes (18 × 150 mm, Bellco Glass Co., Vineland, NJ) capped with 20 mm butyl rubber septum stoppers (Bellco Glass, Co., Vineland, NJ) under anaerobic conditions in liquid ATCC 1019 *Acetobacterium* medium with a headspace consisting of N<sub>2</sub>/CO<sub>2</sub> (95/5, v/v) (Balch et al. 1977). In experiments conducted under autotrophic conditions, fructose and yeast extract were omitted from the medium and a headspace consisting of H<sub>2</sub>/CO<sub>2</sub> (80/20, v/v) was used. All experiments were conducted at room temperature (i.e., 20 °C), which is the optimal growth temperature for *A. paludosum* (Kotsyurbenko et al. 1995).

### Comparison of RDX degradation under autotrophic versus heterotrophic conditions 136

RDX biodegradation was compared under autotrophic and heterotrophic conditions to evaluate the effect of alternative carbon sources on bacterial performance. Degradation assays were conducted in 25 ml Balch anaerobic culture tubes amended with 6 ml of autoclaved ATCC 1019 medium (containing or omitting organic carbon), 1.5 ml of liquid cell culture (washed twice and resuspended in HEPES buffer), and RDX (approximately 3 mg l<sup>-1</sup>). The liquid cell culture was taken from stock *A. paludosum* grown at 20 °C in ATCC 1019 medium (containing fructose as carbon source)

150	with a headspace consisting of 20 ml of H <sub>2</sub> /CO <sub>2</sub>	199	
151	(80/20, v/v) gas mixture. The headspace for the	200	
152	degradation assays also consisted of 20 ml of H <sub>2</sub> /	201	
153	CO <sub>2</sub> (80/20, v/v) gas mixture. A third treatment set	202	
154	was used to investigate the growth of <i>A. paludosum</i>	203	
155	using RDX as the sole source of carbon. The	204	
156	headspace consisted of H <sub>2</sub> /N <sub>2</sub> (5/95 v/v). Two to	205	
157	four replicates were studied for each set. Controls	206	
158	without bacteria were also monitored to obtain a	207	
159	baseline for comparing RDX degradation and	208	
160	acetate production.	209	
161	The tubes were capped and crimped with	210	
162	20 mm butyl rubber stoppers, covered in alumi-	211	
163	nium foil to prevent RDX photolysis, and rotated	212	
164	continuously on a Roto-Torque Heavy Duty		
165	Rotator (Cole-Parmer Instrument Co., Vernon	<i>Effect of ammonium on RDX degradation</i>	213
166	Hills, IL) at 20 °C. Liquid samples (0.7 ml were		
167	collected with sterile disposable syringes, filtered	Similar assays were conducted in 25 ml Balch	214
168	using 0.2 µm syringe filters, and analyzed by high	anaerobic culture tubes with washed cells to	215
169	pressure liquid chromatography (HPLC). The	determine if ammonium (an easily assimilated	216
170	optical density of each tube was recorded	nitrogen source) inhibits RDX degradation. The	217
171	throughout the experiment to determine bacterial	autotrophic medium consisted of one of four	218
172	growth. Each time the reactors were spiked with	substrate combinations: (1) RDX (3 mg l <sup>-1</sup> ) but	219
173	RDX, an additional 20 ml of H <sub>2</sub> /CO <sub>2</sub> headspace	no ammonium; (2) ammonium (1.0 g l <sup>-1</sup> ) but no	220
174	(80/20, v/v) was added to prevent electron donor	RDX, (3) RDX and ammonium; and (4) neither	221
175	and carbon source depletion.	ammonium nor RDX. The headspace consisted of	222
		20 ml of an H <sub>2</sub> /CO <sub>2</sub> (80/20, v/v) gas mixture.	223
176	<i>Evaluation of <sup>14</sup>C-RDX mineralization under</i>		
177	<i>autotrophic conditions</i>	<i>Production of N<sub>2</sub>O from RDX degradation</i>	224
178	Serum bottles (120 ml) were prepared with 54 ml		
179	of ATCC 1019 medium omitting yeast extract and	The production of nitrous oxide (N <sub>2</sub> O) during	225
180	fructose. NaHCO <sub>3</sub> (3 g/l <sup>-1</sup> ) was added to provide	RDX degradation (Figure 1) was investigated to	226
181	a source of inorganic carbon. Six milliliters of pure	determine if RDX ring cleavage occurred. <i>A. pa-</i>	227
182	culture were washed twice and resuspended in	<i>ludosum</i> incubations were prepared with RDX	228
183	HEPES buffer (pH 7) before transfer to each	(approximately 2.5 ml l <sup>-1</sup> , and N <sub>2</sub> O concentra-	229
184	treatment to obtain a 10% (v/v) bacteria/medium	tions in headspace samples (100 µl) were deter-	230
185	concentration. Two sets were prepared in tripli-	mined by gas chromatography. These incubations	231
186	cate: (1) <i>Acetobacterium paludosum</i> in HEPES	were prepared in duplicate 100 ml sealed glass	232
187	buffer and (2) a no-bacteria control consisting of	serum bottles containing ATCC medium 1019,	233
188	5.8 ml of HEPES buffer with 0.1 ml/l <sup>-1</sup> Kathou®	which included yeast extract, fructose (1.3 ml of	234
189	CG/ICP biocide (5-Chloro-2-methyl-3(2H)-isot-	20% solution), and ammonium chloride, and were	235
190	hiazolone and 2-Methyl-3(2H)-isothiazolone	sparged with H <sub>2</sub> /CO <sub>2</sub> headspace (80/20, v/v).	236
191	solution; Sigma-Aldrich, St. Louis, MO). <sup>14</sup> C-	Controls without RDX were also prepared to	237
192	ring-labeled RDX (PerkinElmer Life Sciences,	determine baseline N <sub>2</sub> O production levels.	238
193	Boston, MA) and unlabeled RDX were added to		
194	obtain the initial conditions of 1 µCi total radio-	<i>Analytical methods</i>	239
195	activity and 3 mg l <sup>-1</sup> The activity of the radioac-		
196	tive stock solution was 0.084 µCi µl <sup>-1</sup> . Each bottle	Analysis of RDX and its nitroso derivatives	240
197	held a small test tube with 2 ml of 0.5 N NaOH to	MNX, DNX, and TNX was performed using a	241
198	trap <sup>14</sup> CO <sub>2</sub> . Contents were sparged for 10 min	Hewlett Packard 1100 Series HPLC equipped with	242
		a 250 × 4.6 mm Supelcosil™ LC-18 column, herein	243

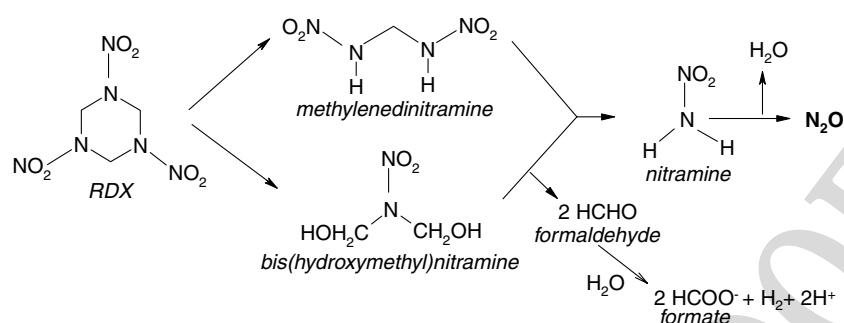


Figure 1. Production of nitrous oxide (N<sub>2</sub>O) and formaldehyde from RDX and hypothetical transformation to formate (Hawari et al. 2000; Oh et al. 2001; Zhao et al. 2002, 2003b).

referred to as the HPLC-1 method. The mobile phase consisted of deionized water and methanol (4:6, v/v) at a flow rate of 1 ml min<sup>-1</sup> (UV detection was at 240 nm. <sup>14</sup>C-RDX and <sup>14</sup>C-metabolites (e.g., methanol and formate) were analyzed by HPLC using a radioactivity detector (Radiomatic, Series A-500, Packard Instrument Co., Downers Grove, IL), herein referred to as the HPLC-RAD method. Analysis for <sup>14</sup>C-formaldehyde was performed using the HPLC-RAD method after derivatization using EPA method # 8315A (omitting the extraction by methylene chloride, due to the small volume of sample used). RDX mineralization was determined from trapped <sup>14</sup>C-CO<sub>2</sub> in the small tubes containing 0.5 N NaOH. Half a milliliter of sample from each of the small tubes was mixed with 9.5 ml of LSC cocktail (Ultima Gold) and was counted on a Beckmann LS 6000IC liquid scintillation counter (Beckman Instr. Inc., Fullerton, CA).

Nitrous oxide analysis was performed using a Hewlett Packard 5890 Series II gas chromatograph instrument with an electron capture detector and a HayeSep Q capillary column (Valco Instruments Co. Inc., Houston, Texas).

Acetate was measured using a Hewlett Packard 1100 Series HPLC equipped with a 150 × 6.5 mm Alltech IOA-2000 Organic Acids column (Deerfield, IL), herein referred to as the HPLC-2 method. The isocratic mobile phase consisted of 0.001 N sulfuric acid in distilled water at a flow rate of 1.0 ml min<sup>-1</sup>. Detection was spectrophotometric at 210 nm, which resulted in a level of detection of less than 2.5 mM.

Bacterial growth was determined by measuring optical density at 660 nm (OD<sub>660</sub>) using a Milton-Roy Spectronic 401 (Milton-Roy Co., Rochester,

New York). The limit of detection was approximately 0.001 absorbance units.

## Results and discussion

### Comparison of RDX degradation under autotrophic versus heterotrophic conditions

Homoacetogens such as *Acetobacterium paludosum* are strict anaerobic mixotrophs that can use H<sub>2</sub> and CO<sub>2</sub> for growth and the production of acetate (Kotsyurbenko et al. 1995). While these bacteria have received considerable attention for their participation in municipal wastewater treatment, our understanding of their role of in aquifer bioremediation is very limited. Thus, experiments were conducted to determine if *A. paludosum* could degrade RDX under environmentally relevant conditions; i.e., when easily assimilated organic carbon sources are absent and H<sub>2</sub> (commonly present in anaerobic systems and Fe<sup>0</sup> barriers) might be the prevalent electron donor.

Experiments were performed to compare RDX degradation by *A. paludosum* under autotrophic versus heterotrophic conditions. No significant RDX removal was observed in abiotic controls, indicating that RDX disappearance was due to biodegradation. Treatments containing *A. paludosum* and RDX as the sole carbon source (i.e., no CO<sub>2</sub>, yeast extract, nor fructose present) degraded approximately 70% of the initial amount of RDX (approximately 3 mg l<sup>-1</sup>) after 9 days incubation (data not shown). Faster degradation was observed in treatments incubated under autotrophic conditions (containing CO<sub>2</sub>) or heterotrophic conditions (containing fructose), where



314 all the RDX was removed within three days.  
 315 Apparently, the presence of alternative (inorganic  
 316 or organic) carbon sources enhanced bacterial  
 317 growth and RDX degradation.

318 In theory, *A. paludosum* could metabolize RDX  
 319 by transforming it to formate (Figure 1), which is  
 320 a known growth substrate (Kotsyurbenko et al.  
 321 1995). However, the observed RDX degradation  
 322 in the absence of alternative carbon sources does  
 323 not necessarily imply that this bacterium metabo-  
 324 lized RDX, because H<sub>2</sub> that was present in the  
 325 headspace could have served as an electron donor  
 326 in the initial (reductive) transformation of RDX.  
 327 Furthermore, an internal storage of carbon pres-  
 328 ent in the (heterotrophically grown, then washed)  
 329 bacteria could have also served as the electron  
 330 donor for RDX transformation. No detectable  
 331 growth of *A. paludosum* was observed when RDX  
 332 was provided as the sole C source. This suggests  
 333 that this bacterium did not metabolize RDX-der-  
 334 ived carbon, which does not necessarily rule out  
 335 RDX utilization as an N source.

336 The autotrophic and heterotrophic treatments  
 337 were respiked with RDX, and the concentration  
 338 versus time data were fit by an exponential decay  
 339 model (i.e.,  $C = C_0 e^{-kt}$ ) using SigmaPlot 8.0  
 340 software (Figure 2). RDX degradation was faster  
 341 under autotrophic (H<sub>2</sub> and CO<sub>2</sub>-fed) conditions  
 342 (96% removal within 10 days) than under hetero-  
 343 trophic (yeast extract plus fructose-fed) conditions  
 344 (73% removal), even though the latter contained a  
 345 higher biomass concentration due to higher cell

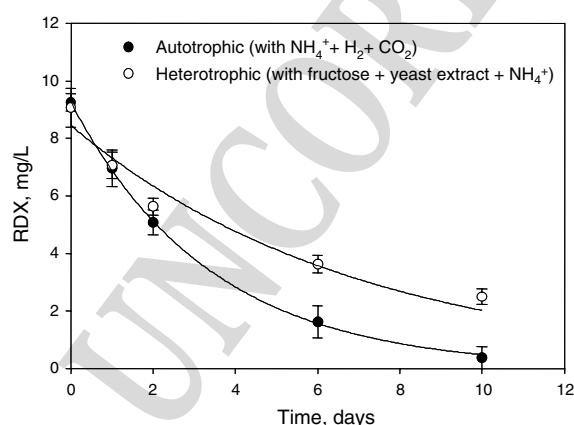


Figure 2. Effect of carbon source on RDX removal by *A. paludosum*. Initial OD<sub>660</sub> was 0.077 ± 0.004 for the autotrophic treatment and 0.510 ± 0.039 for the heterotrophic treatment. Error bars represent 95% confidence intervals.

yield and faster growth under heterotrophic con-  
 ditions (0.510 ± 0.039 versus 0.07 ± 0.004  
 OD<sub>660</sub>). The faster removal for the autotrophic  
 treatment is accentuated when normalized first-  
 order decay coefficients ( $k$ ) are considered. The  
 specific  $k$  value was six times higher for the auto-  
 trophic than the heterotrophic treatment  
 (1.67 ± 0.04 versus 0.28 ± 0.05 (day\*OD<sub>660,0</sub>)<sup>-1</sup>).

These experiments suggest that hydrogen is a  
 better electron donor than fructose and yeast  
 extract for promoting RDX degradation by  
*A. paludosum*, even though the latter are the rec-  
 ommended carbon sources for the growth medium  
 (Balch et al. 1977). The lower RDX removal effi-  
 ciency for the heterotrophic treatments is coun-  
 terintuitive because heterotrophic conditions  
 resulted in faster growth and higher acetate pro-  
 duction after three days (i.e., 30.4 mM acetate  
 (heterotrophic) and 5.9 mM acetate (autotrophic),  
 corresponding to normalized values of  
 153 mM\*(OD<sub>660,0</sub>)<sup>-1</sup> and 31 mM\*(OD<sub>660,0</sub>)<sup>-1</sup>,  
 respectively). Further research will be needed to  
 determine if this observation reflects differences in  
 catabolic activities inherent to autotrophic versus  
 heterotrophic metabolism. For example, auto-  
 trophic metabolism generates more reducing  
 power (for CO<sub>2</sub> fixation) leaving the potential for  
 more electrons to be diverted towards RDX  
 reduction. It may also be possible that the avail-  
 ability of easily assimilated organic carbon sources  
 hinder RDX degradation due to metabolic flux  
 dilution (Lovanh & Alvarez 2004).

#### Degradation of <sup>14</sup>C-RDX under autotrophic conditions

*A. paludosum* degraded RDX (3 mg l<sup>-1</sup>) within  
 9 days, converting it to soluble radio-labeled  
 metabolite(s) (Figure 3). Less than 1% of the  
 radiolabeled RDX was recovered as <sup>14</sup>CO<sub>2</sub> after  
 20 days, indicating that mineralization did not  
 occur. However, separate experiments indicated  
 that production of N<sub>2</sub>O occurred only in treat-  
 ments containing RDX (Table 1), which is evi-  
 dence of ring fission (Figure 1). Recent studies  
 with another homoacetogen, *Acetobacterium mal-*  
*icum*, also reported RDX ring cleavage but no  
 mineralization (Adrian & Arnett 2004).

Anaerobic production of N<sub>2</sub>O from RDX has  
 also been shown for *Clostridium bifermentans*  
 HAW-1 (Zhao et al. 2003a,b). Recovered N<sub>2</sub>O

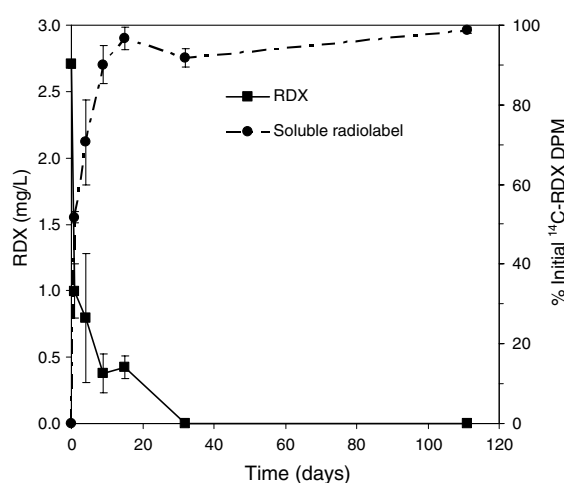


Figure 3. RDX degradation ( $2.7 \text{ mg l}^{-1}$ ,  $1 \mu\text{Ci/bottle}$ ) and soluble  $^{14}\text{C}$ -metabolite(s) formation by  $\text{H}_2$ -fed *A. paludosum* (autotrophic conditions,  $\text{OD}_{660, \text{inoculum}} = 1.27$ ). Error bars represent 95% confidence intervals from the mean of triplicate reactors.

395 accounted for 64% of N-RDX in these experi-  
 396 ments compared to 29.5% reported for  
 397 *C. bifermentans* HAW-1 (Zhao et al. 2003a), which  
 398 suggests a different end-products distribution by  
 399 these two anaerobic organisms. RDX transfor-  
 400 mation by *A. paludosum* was less rapid than that  
 401 reported for *C. bifermentans* HAW-1 (Zhao et al.  
 402 2003a).

403 Attempts to identify the radiolabeled byprod-  
 404 uct(s) after 111 days of incubation were unsuc-  
 405 cessful. Several potential RDX metabolites, that  
 406 have been reported by others (e.g., Adrian &  
 407 Chow 2001; Hawari et al. 2000, 2001; McCormick  
 408 et al. 1981; Oh et al. 2001; Zhao et al. 2003a, b)  
 409 were not detected using the HPLC-1 or HPLC-  
 410 RAD analysis methods used for this research. For  
 411 example, Zhao et al. (2003a) showed *C.*

*bifermentans* HAW-1 transformed RDX tran-  
 412 siently to MNX, DNX, and TNX, which were  
 413 further transformed to methanol, formaldehyde,  
 414 carbon dioxide, and nitrous oxide. However, no  
 415 MNX, DNX, TNX, formaldehyde, formic acid, or  
 416 methanol were detected in our analyses. Whereas  
 417 the radiolabel eluted as one peak (2.8 min) using  
 418 the RDX (HPLC-RAD) method, it eluted as two  
 419 peaks (1.8 and 3.0 min) using the acetate  
 420 (HPLC-2) method, suggesting the presence of two  
 421 compounds that were not acetate (3.5 min).  
 422 Retention times for  $^{14}\text{C}$ -labeled formaldehyde  
 423 (7.0 min), formate (2.7 min), and methanol  
 424 (3.1 min) were also determined with the HPLC-  
 425 RAD method. These elution times suggest that  
 426 formate might have been one of the unidentified  
 427 RDX metabolites (Figure 1). The HPLC-2 meth-  
 428 od was not run with formate to verify this notion.  
 429 However, other previously reported RDX degra-  
 430 dation products such as hydroxylamino metabo-  
 431 lites (Adrian & Chow 2001), methylenedinitramine  
 432 and bis(hydroxymethyl)nitramine (Figure 1) (Ha-  
 433 wari et al. 2000; Oh et al. 2001), are relatively  
 434 short lived (Adrian & Chow 2001; Bhushan et al.  
 435 2002) and are unlikely to persist as long as the  
 436 unidentified metabolites did in this experiment  
 437 (Figure 3). Similarly, no metabolites were identi-  
 438 fied using an Agilent 1100 series liquid chromato-  
 439 graph/mass spectrometer, presumably due to lack  
 440 of sensitivity in full scan mode.

441 The volatility and reactivity of the radiolabeled  
 442 metabolites towards oxygen was also investigated.  
 443 Two 20 ml LSC vials were prepared with 5 ml of the  
 444 medium remaining from the autotrophic experi-  
 445 ment in which *A. paludosum* transformed  
 446  $^{14}\text{C}$ -RDX. Both vials were covered with aluminum  
 447 foil and lightly capped with aluminum foil to pre-  
 448 vent photo-interactions but still allow for volatil-  
 449 ization. One vial was placed in the anaerobic  
 450

Table 1. Production of  $\text{N}_2\text{O}$  by *A. paludosum* incubated with RDX

Treatment	RDX degraded ( $\mu\text{M}$ )	Maximum theoretical $\text{N}_2\text{O}$ produced ( $\mu\text{M}$ ) <sup>a</sup>	Measured $\text{N}_2\text{O}$ produced (mM)	Percentage of Theoretical Maximum $\text{N}_2\text{O}$ produced ( $\mu\text{M}$ ) <sup>b</sup>
With RDX	1.50	4.50	2.9	64%
With RDX, duplicate	1.53	4.59	0.9	18%

<sup>a</sup>Theoretical calculation assumes that 3 M of  $\text{N}_2\text{O}$  gas could be produced from 1 M RDX.

<sup>b</sup>Corrected for background  $\text{N}_2\text{O}$  detected under conditions without RDX.

451 chamber and the other was exposed to air outside  
452 on the lab bench. After one week, the samples were  
453 analyzed by LSC and HPLC. No loss of radioac-  
454 tivity and no changes in HPLC peak elution times  
455 had occurred, indicating that the metabolites were  
456 not volatile and did not spontaneously react with  
457 oxygen.

#### 458 *Effect of ammonium on RDX degradation*

459 Figure 4 shows the degradation of three sub-  
460 sequent spikes of RDX by *A. paludosum* under  
461 autotrophic conditions in the presence and absence  
462 of ammonium ( $1.0 \text{ g l}^{-1} \text{ NH}_4\text{Cl}$ , which is the rec-  
463 ommended concentration for the *A. paludosum*  
464 growth medium). RDX degradation rates  
465 decreased for both treatments upon subsequent  
466 RDX spikes, possibly due to toxicity associated  
467 with RDX biotransformation or to the accumu-  
468 lation of inhibitory metabolites. Ammonium had  
469 no significant effect on the degradation of the first  
470 spike of RDX. Estimated  $k$  values (normalized to  
471 the initial optical density) were  $4.43 \pm 0.71$  and  
472  $4.57 \pm 0.30 \text{ (day*OD}_{660,0})^{-1}$  with and without  
473 ammonium, respectively. However, ammonium  
474 had an inhibitory effect on the degradation of the  
475 second spike of RDX, decreasing the  $k$  value by  
476 about one-half, from  $4.52 \pm 0.57$  to  $2.41 \pm 0.18$   
477  $\text{(day*OD}_{660,0})^{-1}$  (Figure 4). The inhibitory effect  
478 of ammonium was reproducible during the deg-  
479 radation of the third RDX spike, where the  $k$  value  
480 for the treatment without ammonia, ( $1.66 \pm 0.12$   
481  $\text{(day*OD}_{660,0})^{-1}$ ) was significantly higher  
482 ( $p < 0.05$ ) than the value for the treatment with  
483 ammonia ( $0.74 \pm 0.13 \text{ (day*OD}_{660,0})^{-1}$ ).

The inhibition of RDX degradation by 484  
ammonium might be due to its preferential utili- 485  
zation over RDX as a nitrogen source. Whereas 486  
we did not demonstrate *A. paludosum* assimilation 487  
of RDX-derived nitrogen (which would have 488  
required the use of  $^{15}\text{N}$ -labeled RDX), numerous 489  
studies have shown that RDX can serve as a 490  
nitrogen source to bacteria (Beller 2002; Binks 491  
et al. 1995; Coleman et al. 1998; Sheremata & 492  
Hawari 2000) Thus, further research is recom- 493  
mended to test this hypothesis and to evaluate 494  
whether ammonium represses enzymes that initiate 495  
RDX degradation by reducing its nitro groups 496  
(Bhushan et al. 2002), as is the case for ammonia 497  
repression of assimilatory nitrate and nitrite 498  
reductases (Madigan et al. 2000). 499

#### 500 **Summary and conclusions**

501 Little is known about the role of homoacetogenic 501  
bacteria in bioremediation. This study is the first 502  
report of RDX biodegradation by *A. paludosum*, 503  
which degraded RDX under both heterotrophic 504  
and autotrophic conditions that might prevail, 505  
respectively, in bioreactors and in the vicinity of 506  
iron barriers. Although RDX was not mineralized 507  
to  $\text{CO}_2$ , evidence of ring fission (per  $\text{N}_2\text{O}$  accu- 508  
mulation) with possible conversion to innocuous 509  
formate was obtained, and no objectionable het- 510  
erocyclic nitroso derivatives (i.e., MNX, DNx, 511  
and TNx) were detected. However, not all deg- 512  
radation products were identified, which precludes 513  
our full endorsement of this pathway for bio- 514  
remediation purposes. 515

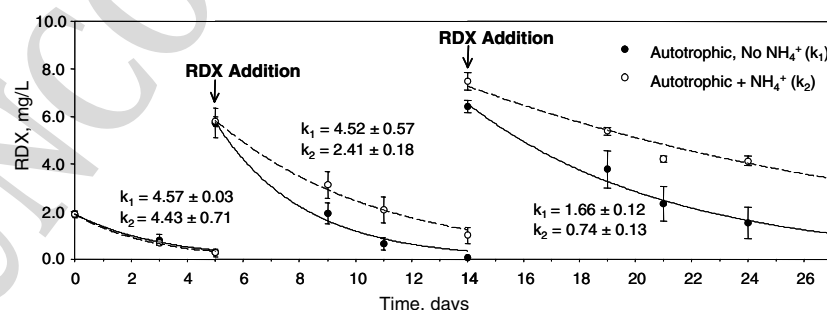


Figure 4. Degradation of three spikes of RDX by *A. paludosum* in the presence or absence of ammonium. The experiment was conducted under autotrophic, anaerobic conditions with an average initial optical density (at 660 nm) of  $0.080 \pm 0.009$  for reactors with ammonium and  $0.073 \pm 0.003$  for reactors without ammonium. The depicted first-order rate coefficients,  $k$ , have units of  $\text{(days*OD}_{660,0})^{-1}$ . Error bars represent 95% confidence intervals from the mean of triplicate reactors.

516 RDX was inhibitory to *A. paludosum* growth,  
517 and its degradation was more efficient under  
518 autotrophic ( $H_2$ -fed), nitrogen deficient condi-  
519 tions, even though faster growth occurred under  
520 heterotrophic (yeast extract plus fructose-fed)  
521 conditions. The addition of ammonia had an  
522 inhibitory effect on RDX degradation, possibly by  
523 relieving selective pressure for the utilization of  
524 RDX as a nitrogen source. Demonstration of  $^{15}N$ -  
525 RDX incorporation into biomass is recommended  
526 for future studies to confirm assimilation of RDX  
527 nitrogen by *A. paludosum*. Nevertheless, these  
528 results suggest that the common practice of bi-  
529 ostimulation through the addition of auxiliary  
530 substrates and nutrients should be carefully eval-  
531 uated on a case by case basis to prevent a coun-  
532 terproductive effect on RDX bioremediation.

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