# ENVIRONMENTAL FATE STUDIES ON CERTAIN MUNITIONS WASTEWATER CONSTITUENTS

Phase IV - Lagoon Model Studies

Final Report

By

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The major loss processes for the transformation of TNT were phototransformation and biotransformation. A first-order, depth-independent rate constant for the indirect photolysis of TNT in mid-March was measured at 4.3 cm d<sup>-1</sup> which yield an estimated half-life of eight days in a body of water 50 cm deep. A major stable TNT phototransformation product was 2-amino-4,6-dinitrobenzoic acid. Biotransformation occurred in the lagoon with a pseudo-first-order rate constant of 0.010d<sup>-1</sup> and a second-order rate constant of 8.95 x 10<sup>-9</sup> ml cell<sup>-1</sup> d<sup>-1</sup>. The half-life of TNT resulting from biotransformation was calculated to be 69 days. The major biotransformation products were 2-amino-4,6-dinitrotoluene and 4-amino-2,6-dinitrotoluene.

The major transformation process for RDX was found to be photolysis. RDX transformed with first-order rate constants ranging from 0.016 cm d<sup>-1</sup> in winter to 0.076 cm d<sup>-1</sup> in summer. The half-life of RDX was estimated to range from over 2000 days in winter to 456 days in summer in a lagoon 50 cm deep.

Although photolysis was found to be the major transformation process for TNT and RDX, the rate of transformation was slowed due to the high absorptivity of the lagoon waters, whose absorbances ranged from 3.0 in winter to 9.0 in summer. These high absorbances allow little light to penetrate for a significant distance below the lagoon surface. The light absorption, however, heats the lagoon water, thus accelerating the dissolution process of solid TNT and RDX on the lagoon sediment surface. Thus, TNT and RDX concentrations in the lagoon water were found to increase from the winter to the summer months.

A computer model was constructed to simulate the lagoon enviroment. Using the laboratory-developed transformation and dissolution rate constants, the persistence of TNT and RDX can be estimated as a function of time from initial inputs of lagoon depth and aqueous and sediment-surface munition concentrations.

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#### EXECUTIVE SUMMARY

This study was conducted to investigate the persistence of 2,4,6trinitrotoluene (TNT) and hexahydro-1,3,5-trinitro-1,3,7-triazine (RDX) in waste disposal lagoons at Louisiana Army Ammunition Plant. The waste disposal lagoons are complex environments due to a heterogeneous distribution of solid TNT and RDX munitions on the lagoon sediment surface. These solids serve as a constant input source of munitions to the lagoon waters through dissolution. Flux rates for the dissolution of munitions were measured at  $1.08 \times 10^{-4}$  cm sec<sup>-1</sup> for TNT and 7.75 x  $10^{-5}$  cm sec<sup>-1</sup> for RDX at 25°C in the lagoon.

The major loss processes for the transformation of TNT were phototransformation and biotransformation. A first-order, depthindependent rate constant for the indirect photolysis of TNT in mid-March was measured at 4.3 cm d<sup>-1</sup> which yields an estimated half-life of eight days in a body of water 50 cm deep. A major stable TNT phototransformation product was 2-amino-4,6-dinitrobenzoic acid. Biotransformation occurred in the lagoon with a pseudo-first-order rate constant of  $0.010d^{-1}$  and a second-order rate constant of  $8.95 \times 10^{-9}$ ml cell<sup>-1</sup> d<sup>-1</sup>. The half-life of TNT resulting from biotransformation was calculated to be 69 days. The major biotransformation products were 2-amino-4,6-dinitrotoluene and 4-amino-2,6-dinitrotoluene.

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## CONTENTS

EXECUTIVE SUMMARY	í
LIST OF ILLUSTRATIONS	v
LIST OF TABLES	vii
ACKNOWLEDGMENTS	x
I. INTRODUCTION	1
II. SITE DESCRIPTION	3
III. CHEMICALS IDENTIFIED IN LAGOON NO. 9	10
IV. SITE EVALUATION	19
V. TRANSPORT AND TRANSFORMATION STUDIES	21
A. Physical Transport	21
<ol> <li>Water Balance</li></ol>	21 21 23
in Sediment Cores	26 27 28
B. Photochemistry	33
<ol> <li>TNT Photochemical Rate Constant- General Remarks</li> <li>2. Effect of Depth on Photolysis Rate Constant</li> </ol>	36
of TNT in LAAP Water	40 43
for Weather Conditions 5. Effect of Time of Year on Photochemical Rate	47
Constant of TNT in LAAP Water 6. UV-Visible Spectra of LAAP Water 7. Experimental Studies of the Photolysis	48 51
of TNT in LAAF Water	53 56

## CONTENTS (Concluded)

1.

Ņ

-

þ

С. В	lotransformation	59
2	<ul> <li>Bacterial Count</li> <li>Biotransformation Screening Tests</li> <li>a. Aerobic TNT Biotransformation</li> <li>b. Anaerobic TNT Biotransformation at High and Low Concentrations of TNT</li> <li>d. TNT Metabolites</li> <li>e. Biological Oxygen Demand of the Lagoon Water</li> <li>g. RDX Aerobic Biotransformation</li> <li>b. Detailed Biotransformation Rate Study</li> </ul>	59 61 62 64 66 67 70 70 71 73
VI. DEVELOPM	ENT OF THE SIMULATION MODEL	78
A. G	eneral Equations	78
B• P	rogram Structure	79
C. S	imulation Results	81
2	<ul> <li>Simulation of the Water Surface</li> <li>TNT Simulation</li> <li>a. Case I: Complete Mixing</li> <li>b. Case II: Zero Mixing</li> <li>RDX Simulation</li> </ul>	81 84 84 90 93
VII. DISCUSSI	ON	96
VIII. LITERATU	RE CITED	144
APPENDIX A: An	alysis of LAAP Lagoon No. 9 Sediment	100
	ss Transfer ProblemsAssumptions d Solutions	105
Fa	I Computer Program for Modeling the te of TNT and RDX in Lagoons containing mition Wastes	130
DISTRIBUTION LI	ST	146

iv

## ILLUSTRATIONS

1.	Lagoon Area at Louisiana Army Ammunition Plant	4
2.	Diagram of Lagoon No. 9 and TNT-RDX Distributions (%) in the Sediment	7
3.	HPLC Profile of Lagoon Water (Sampled 7/22/82)	11
4.	HPLC Profile of the Methylated Ether II Fraction of Lagoon Water	14
5.	HPLC Profile of Lagoon Water Resolving 2-Amino- 4,6-Dinitro Benzoic Acid and Other Components	17
6.	Estimation of Lagoon Depth Profile Based on Variable Seepage Rates	24
7.	Lagoon Water Rests on Sediment	29
8.	TNT (•) and RDX (0) Flux for LAAP Lagoon No. 9 Sediment	37
9.	Absorption Spectrum for Lagoon 9, Site 1 Water, Filtered, 0.1-cm Cell	52
10.	Aerobic Biotransformation of TNT and RDX in Lagoon Water	63
11.	Anaerobic Biotransformation of TNT and RDX in Lagoon Water	65
12.	Radioautograph of Thin-layer Chromatogram of TNT Metabolites in Lagoon Water after 19 Days of Incubation	69
13.	Lineweaver-Burke Plot of TNT Biotransformation Rate in Lagoon Water	76
14.	Simulation Flow Chart of the Lagoon Model	80
15.	Program Interactions	82

h

## ILLUSTRATIONS (Concluded)

. 7

16.	Approximate Water Contour Lines During the Simulation	83
17.	Simulated TNT Concentration with Respect to Time of Year	85
18.	Simulated TNT Concentration as a Function of Time of Year	86
19.	Simulated TNT Concentration from the Case of Sole Input from Dissolution	88
20.	Concentration-Time-Dependent Profile of TNT Measured in Lagoon No. 9	89
21.	Simulation Results for TNT in Compartments 1 and 13 for the Case of Zero-Mixing	91
22.	Comparison of the Simulation Results from the Case with Dissolution As the Only Input Source of TNT to Measured Data from the Lagoon	92
23.	Simulation Results for RDX Concentration in LAAP Lagoon Water	94
A-l.	HPLC Profile of Lagoon No. 9 Sediment Extract	102
B-1.	Lagoon Water Rests on Sediment	106
B-2.	Boundary Layer Approximation in Sediment/ Water Phase	110
в-3.	TNT (•) and RDX (°) Flux Data Compared with Curve Fit (Equation 62)	129

vi

## TABLES

· ·

· · · ·

- 1

1.	Lagoon Water Temperatures (°C) for the Bottom, Middle, and Surface on December 9, 1981	9
2.	Concentrations of TNT and RDX in Water at Lagoon Center and Southeast Edge As a Function of Time of Day	9
3.	Concentrations (ppm) of Munitions Found in Water Samples on Various Dates	12
4.	Levels of 2-Amino-4,6-dinitrobenzoic Acid Observed in Lagoon No. 9 on Various Sampling Dates	18
5.	Absorbance Values of Lagoon No. 9 Water at 290 and 400 nm at Various Times of the Year	18
6.	Average Monthly Precipitation and Evaporation Data from 1979 through 1981 at Red River Weather Station, Restin, LA	22
7.	Monthly Precipitation and Evaporation Data from January 1982 through August 1982 at Red River Weather Station, Restin, LA	22
8.	Solubility of TNT and ROX in Water at 10, 20, and 30°C	23
9.	Measured and Calculated Solubilities (mg liter <sup>1</sup> ) of TNT and RDX at 283, 293, and 303°K	26
10.	Sorption Partition Coefficients for TNT and RDX in LAAP Lagoon No. 9 Sediment	27
11.	Sediment Core Sample Dry Weight and Percent TNT and RDX Weights	27
12.	TNT and RDX in Sectioned Cores from LAAP Lagoon No. 9	30
13.	TNT and RDX Flux Measurements from LAAP Lagoon	25

.

•

## TABLES (Continued)

1

14.	Depth for 99% Absorption of Light of Specific Wavelengths for LAAP Water	39
15.	Photolysis of 4.97 ppm TNT in LAAP Water in an MGR Apparatus and in Sunlight	43
16.	Photolysis of 4.97 ppm TNT in LAAP Water As a Function of Depth Using a Xenon Lamp	45
17.	Sunlight Photolysis of 4.97 ppm TNT in LAAP Water and Actinometers	46
18.	Variation in TNT Photolysis Rates in LAAP Water As a Function of Time of Year	50
19.	Absorption Coefficient of December and July LAAP Waters at Selected Wavelengths	51
20.	Sunlight Photolysis of TNT in LAAP Water	55
21.	Photochemical Rate Constants for RDX in July LAAP Water as a Function of Season	58
22.	Total Aerobic and Anerobic Bacteria in Lagoon 9 Water Collected in December	60
23.	Total Aerobic and Anaerobic Bacteria in Lagoon 9 Sediment Collected in December	60
24.	Bacterial Plate Count of LAAP Lagoon Water Collected in July	61
25.	RDX Concentration on Different Days after Anaerobic Incubation in LAAP Lagoon Water	71
26.	Anaerobic Transformation of RDX with Selected Organisms and Yeast Extract Concentrations	72
27.	Transformation Rates Determined for Various TNT Concentrations	75
28.	Initial TNT Concentrations and Biotransformation Rates over 5 Days	77

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## TABLES (Concluded)

• •

•

•••••

5

A-1.	Percent Composition of Munition Components in LAAP Lagoon No. 9 Sediment	103
A-2.	Percent Water in Lagoon No. 9 Sediment Samples	104
B-1.	Calculated Flux Values Considering All Parameters $(N_1^*)$ , No Chemical Reaction $(N_2^*)$ , No Convection $(N_3^*)$ , No Reaction or Convection $(N_4^*)$ , and Their Ratios to $(N_1^*)$	113
B-2.	Flux Calculations for Sample Lagoon Problem	120
B-3.	Total Mass per Unit Area Transferred During a Time Interval for Sample Lagoon Problem	120
B-4.	Parameters for Analyzing Flux Data	28

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#### I. INTRODUCTION

The U.S. Army is currently involved in assessing the overall hazard of munitions-related warer pollutants to aquatic and mammalian systems. Part of this assessment involves the loss and movement (fate) of militarily unique chemicals when they are discharged into various aquatic environments. Munitions have been produced and handled for many years with minimal pollution-abatement controls, but it has now become important to estimate the environmental persistence of selected chemicals used in present and past military operations.

Environmental persistence can be estimated through the use of computer modeling, in which the loss and movement processes as well as critical environmental parameters can be described by mathematical equations and approximations. In most environmental situations or in cases of low chemical concentrations, simple first-order or pseudofirst-order rate expressions and equilibrium constants adequately describe the loss and movement processes. Once the rate and equilibrium constants are known for the dominant processes, environments can be simulated under any number of specified conditions. Thus, it becomes possible to estimate the concentration of a chemical as a function of time when the chemical is subjected to various environmental stresses. In a previous study (Spanggord et al., 1981), we applied the modeling concept to munitions-related chemicals in the Holston River and generated concentration- time-dependent profiles that allowed the prediction of chemical concentrations 20 km from the discharge point. Actual data on samples taken from the river verified the applicability of the modeling concept.

On the basis of that experience, this study was undertaken to estimate the persistence of 2,4,6-trinitrotoluene (TNT) and hexahydro-1,3,5-trinitro-1,3,5-triazine (RDX) in disposal lagoon environments. The lagoon disposal of munitions wastes has been practiced for many

years and the Army would like to dispose of these lagoons in a safe manner. The studies reported herein describe the major transport and transformation processes that affect the persistence of TNT and RDX in lagoons. We also developed a computerized simulation model to describe the loss and movement of these chemicals as a function of time, concentration, and location in the lagoon.

#### **II. SITE DESCRIPTION**

This investigation covered evaluation of the persistence of TNT and RDX in the waste disposal lagoons at Louisiana Army Ammunition Plant (LAAP) in Minden, Louisiana. LAAP is a load, assemble, and pack facility that generates munitions wastewaters during shell-washout operations in the loading/unloading process. The wastewaters are transported by truck to disposal lagoons, where natural evaporation concentrates the residues.

We conducted a field study at LAAP on December 7-11, 1981. At that time, no wastes had been transported to the lagoons since May and a new carbon-column waste-treatment facility was to be installed prior to further waste disposal.

The lagoon disposal area at LAAP is outlined in Figure 1. Lagoon No. 9 was selected for our investigations mainly because it is one of the larger lagoons and because previous reports had indicated that TNT and RDX were still being found in the lagoon waters and sediments (Wentzel, et al., 1981).

Lagoon No. 9 is a nearly square, flat-bottomed reservoir measuring 56.7 m  $\times$  52.1 m (186 ft  $\times$  171 ft). During the week of December 7-11, 1981, it was filled to a depth of 60 cm (2.0 ft) with a dark tea-colored water. Water input into the lagoon is from seasonal rains, and water output is due to seepage and evaporation. Trees and shrubbery line the lagoon enbankments, but no plant or aquatic life was evident in the lagoon water.

To begin our field study, a tow line was secured across the middle of the lagoon and the center position was tagged. We used an inflatable raft to transport monitoring and sampling equipment to the center of the lagoon for sample collection and monitoring measurements.



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## FIGURE 1 LAGOON AREA AT LOUISIANA ARMY AMMUNITION PLANT

Water samples were collected with a variable-water-depth sample collector in 50-ml amber glass jars with Teflon-lined caps. We analyzed these samples on site, using a high-pressure liquid chromatograph (HPLC). We collected other water samples in 4-liter amber glass bottles and transported them by air overnight to SRI for total microbial counts. Sediment samples, removed from the lagoon with a hand corer, were collected at various locations and transported to SRI for analysis. These samples indicated a heavy clay bottom that was difficult to penetrate. Except for sorption properties, we did not evaluate the morphological characteristics of the sediment.

To determine the ability of light to penetrate into the lagoon water, we placed photochemical "trees" at various depths in the lagoon. These trees consisted of aluminum rods, cross bars, and flotation supports and were equipped with sealed PNA/PYR<sup>\*</sup> actinometer solutions attached to the ends of the crossbars. After selected time intervals, the PNA/PYR tubes were removed from the apparatus, wrapped in aluminum foil, and transported to SRI for analysis.

Meteorological conditions that were measured periodically at the site were temperature, pressure, wet and dry bulb temperatures (relative humidity), and wind velocity. In the lagoon water, we measured temperature, pH, and dissolved oxygen. LAAP personnel obtained a water sample in March 1982. During a second visit to the site on May 25, 1982, we collected water and sediment samples. In addition, sample bottles were left with plant personnel to collect samples at two-week intervals and to mail them to SRI. Depth at the center of the lagoon was measured during the December visit and again on July 24, 1982, to estimate water loss over an 8-month period. The lagoon was completely dry in late August.

<sup>&</sup>quot;PNA/PYR, p-nitroacetophenone/pyridine, sunlight actinometer developed at SRI.

The yearly meteorological conditions for the lagoon site were estimated from daily data collected by the Red River experimental weather station at Restin, LA, approximately 30 miles east of LAAP. The data collected at this site include daily maximum and minimum air and soil temperatures, 24-hour pan-evaporation, and 24-hour precipitation. We averaged the data collected from this station for three years (1979 through 1981) to estimate water balances within the lagoon at LAAP.

A graphic sketch of Lagoon No. 9 is shown in Figure 2. A concrete ramp in the southeast corner of the lagoon serves as an access point for vehicles to dump liquid and solid wastes. Analysis of sediment samples (Appendix A) collected on May 25, 1982, from a  $5 \times 5$  sectioned matrix of the lagoon showed very high concentrations of munitions that decreased as a function of distance from the dump ramp (Figure 2). When the lagoon dried up in late August 1982, pure munitions compounds were visible on the sediment surface near the ramp. These solids serve as the primary input source of munitions to the lagoon waters. The concentration of munitions also decreased as a function of depth in the sediment. The distribution of munitions in small sediment cores taken from various locations in the lagoon is described in Section V.A.4.

Because the lagoon is a highly colored body of water, it absorbs much of the sunlight energy and converts it to heat, thus establishing a thermocline in the lagoon water. Temperature fluctuations during the course of a day are shown in Table 1 for the bottom, middle, and surface of the lagoon. Over the nine-month course of this study, the lagoon temperatures ranged from 10°C in December to 35°C in May and 41.5°C in August. This temperature change affects rates of dissolution, diffusion, and microbial activity as well as solubility.

The lagoon waters appeared to be well mixed in a vertical direction, as evidenced by similar concentrations of TNT, RDX, and dissolved oxygen measured at the surface and at depths of 25 and 50

E	1.9 x 10 <sup>-3</sup> 1.3 x 10 <sup>-3</sup>		1.1 × 10 <sup>-3</sup> 4.9 × 10 <sup>-4</sup>		× 10 <sup>-3</sup> × 10 <sup>-4</sup>	2.2 × 10 <sup>-4</sup> 0.6 × 10 <sup>-4</sup>	TNT RDX
D	0.049 0.066	0.012 0.161		10 <sup>-4</sup> 10 <sup>-3</sup>		$1.2 \times 10^{-3}$ $  < 1.0 \times 10^{-3}$	TNT RDX
с	0.17 1.03	0.002 0.004		004 10 <sup>-4</sup>	1.5 x 10 <sup>-</sup> 2.0 x 10 <sup>-</sup>		TNT RDX
8	1.44 4.67	0.14 1.74		)40 ?94	8.3 x 10 <sup>-</sup> 7.9 x 10 <sup>-</sup>	(	TNT RDX
A	27.5 8.73	29.7 6.55	     1.   0.1	48 89	0.042	0.002	TNT RDX
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FIGURE 2 DIAGRAM OF LAGOON NO. 9 AND TNT-RDX DISTRIBUTIONS (%) IN THE SEDIMENT

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cm. Horizontal mixing, however, was not complete; concentrations of TNT and RDX were higher in the south bank waters than those in the center of the lagoon (Table 2). The higher concentrations near the edge of the lagoon may be the result of the water there being warmer, which increases the rates of dissolution and diffusion of munitions from the near-shore sediments relative to bulk diffusion rates in the main lagoon.

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## Table 1

## LAGOON WATER TEMPERATURES (°C) FOR THE BOTTOM, MIDDLE, AND SURFACE ON DECEMBER 9, 1981

Time	Bottom (50 cm)	<u>Middle (25 cm)</u>	<u>Surface (0 cm)</u>
0820	11.6	11.8	11.9
0925	11.6	11.9	12.1
1020	11.8	12.0	14.0
1120	11.7	12.0	13.5
1240	11.7	11.7	14.0
1350	12.0	12.0	16.5
1440	12.0	12.3	17.0
1550	12.0	13.8	16.0

#### Table 2

#### CONCENTRATIONS OF TNT AND RDX IN WATER AT LAGOON CENTER AND SOUTHEAST EDGE AS A FUNCTION OF TIME OF DAY

	<u>TNT</u>	(ppm)	RDX (ppm)	
Time of Day	Edge	Center	Edge	Center
0800	8.34	4.81	12.7	12.9
0900	7.11	4.35	13.2	12.5
1000	7.00	4.95	13.6	12.9
1100	7.62	5.02	13.0	13.4
1200	6.80	4.88	12.7	12.6
1300	6.54	4.78	12.9	13.2
1400	7.27	4.99	12.8	12.9
1 500	7.28	4.71	12.8	13.2

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#### III. CHEMICALS IDENTIFIED IN LAGOON NO. 9

Water samples were collected from the lagoon during site visits in December 1981, May 1982, and July 1982. In addition, samples were sent to SRI by LAAP personnel at two-week intervals from June 1982 until the lagoon became dry in late August 1982.

The samples were analyzed by HPLC, using direct aqueous injection of the sample after the addition of an internal standard (3,5-dinitrotoluene).

The following conditions were employed:

Instrument:	Spectra-Physics Model 3500B Liquid Chromatograph
Column:	Waters Assoc C <sub>18</sub> Radial-Pak A
Solvent:	30% B in A to $80%$ B in A. A = H <sub>2</sub> O;
	B = methanol/acetonitrile (50/50)
Flow Rate:	2.0 ml min <sup>-1</sup>
Detection:	UV @ 254 mm

A typical HPLC profile appears in Figure 3. The chemicals identified in this profile and confirmed by probe-mass spectrometry were octahydro-1,3,5,7-tetranitro-1,3,5,7-tetrazocine (HMX), RDX, 1,3,5-trinitrobenzene (TNB), 3,5-dinitroaniline (DNA), TNT, 2-amino-4,6-dinitrotoluene (2-A-4,6-DNT), and 4-amino-2,6-dinitrotoluene (4-A-2,6-DNT). Concentrations of each chemical in water samples at various dates are shown in Table 3.

Investigations were also conducted to identify stable munitionsrelated chemicals in the lagoon water. A 400-ml sample of lagoon water was extracted with benzene (2  $\times$  400 ml) (benzene fraction) and ether (2  $\times$  200 ml) (ether I fraction); then the water was acidified to pH l with HCl and re-extracted with ether (2  $\times$  200 ml) (ether II fraction). Each fraction was dried over anhydrous sodium sulfate and rotary-evaporated to dryness. The residues were dissolved in a small amount of ether for chromatographic analysis.





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#### Table 3

## CONCENTRATIONS (ppm) OF MUNITIONS FOUND IN WATER SAMPLES ON VARIOUS DATES

Date Sampled	Day	HMX	RDX	TNB	4-A-2,6-DNT	2-A-4,6-DNT	TNT	DNA
12/08/81	0		13.2	0.33	1.10	1.50	6.98	1.0
3/15/82	74	2.87	10.9	0.58		1.01	5.49	<0.10
5/25/82	145	3.15	18.3	0.65	0.91	1.33	19.8	0.54
6/09/82	160	4.31	22.4	0.91	2.35	1.33	11.5	0.54
6/23/82	174	4.40	20.9	0.75	2.31	1.39	5.65	0.68
7/07/82	188	5.71	28.9	0.84	0.83	1.48	6.31	0.99
7/22/82	203	4.03	17.1	0.20	0.42	0.77	2.81	0.73
8/04/82	216	6.36	42.1	4.17	0.58	1.37	12.3	0.87
8/20/82	232	6.03	12.6	2.06	0.65	0.83	0.50	0.34
9/03/82	246	5.07	5.6	<0.10		1.16	0.20	<0.10

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We identified TNT, 2-amino-4,6-dinitrotoluene, 4-amino-2,6dinitrotoluene, RDX, and trace amounts of 1,3,5-trinitrobenzene and 2,4,6-trinitrobenzyl alcohol by thin-layer chromatographic (TLC) analysis on silica gel, using benzene as a solvent and an ethylenediamine/DMSO spray reagent. In the ether I fraction, only one spot migrated, which coincided with RDX. In the ether II fraction, several spots were observed close to the origin but could not be identifed by cochromatography with standards or color development with the spray reagent.

The ether II fraction was treated with diazomethane and subjected to HPLC analysis under the following conditions:

Column: Water's C<sub>18</sub> Radial Pak A Solvent: 0-99% methanol in water with a 10 min delay and a 50 min linear gradient Flow rate: 2.0 ml min<sup>-1</sup> Detection: UV @ 254 nm A typical HPLC chromatogram appears in Figure 4. Components from the HPLC profile in Figure 4 were collected and evaluated by probe mass spectrometry. The fraction 57.34 component was identified as the methyl ether of 3,5-dinitrophenol. Fraction 54.31 was tentatively identified as methyl 2-hydroxy-4,6-dinitrobenzoic acid, methyl ether and methyl ester, but could not be confirmed due to the lack of a reference standard. Fraction 54.31 also contained a component that was identified as methyl 2-amino-4,6-dinitrobenzoic acid, methyl ester, and was confirmed by the synthesis of a reference standard in our laboratory. The other component could not be identified due to multicomponent spectra and elevated background signals. Fraction 43.50 was identified as methyl 2,4-dinitrobenzoic acid, methyl ester.

Thus, the majority of chemicals in the LAAP lagoon water are related to the photochemical transformation of TNT with the exception of 2-amino-4,6-dinitrotoluene and 4-amino-2,6-dinitrotoluene, which arise from microbial sources. Based on the work of Kaplan et al. (1975), who showed that many of the identified components are the result of photochemical processes, Scheme I indicates the probable route of formation of the identified products.



Scheme I



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FIGURE 4 HPLC PROFILE OF THE METHYLATED ETHER II FRACTION OF LAGOON WATER

A key step in this scheme is the series of oxidative-reductive disproportionations leading to 2-amino-4,6-dinitrobenzoic acid, as shown in Scheme II.



Scheme II

All the chemicals identified in the lagoon water were also found in the sediment. In addition, the explosive, tetryl, was found in the sediment.

Because 2-amino-4,6-dinitrobenzoic acid appeared to be a major transformation product in the lagoon water, we developed an HPLC procedure to monitor this compound. The following conditions were employed:

```
Column: 250 mm × 4.6 mm Alltech C_{18} 10 µm
Solvent: 40% B in A + 50% B in A in 10 min after a
5-min hold. A = 0.1M phosphate buffer (pH 7)
with 0.05 M tributylammonium phosphate (TBAP);
B = 0.005 M (TBAP) in methanol.
Flow rate: 1.8 ml min<sup>-1</sup>
Detector: UV @ 254 nm
Retention time: 3.62 min
```

A typical chromatographic profile of lagoon water appears in Figure 5. Lagoon waters that were collected throughout this study were analyzed for 2-amino-4,6-dinitrobenzoic acid. The results (Table 4) suggest that 2-amino-4,6-dinitrobenzoic acid is a major transformation product in lagoon waters.

The lagoon waters were found to be nearly saturated with oxygen. Measurements taken with a dissolved oxygen probe during the December field trip indicated 9.0 ppm  $0_2$  at the surface and 8.8 ppm near the bottom sediment (50 cm deep). In such a highly aerated environment, nitrite is readily oxidized. In fact, when the lagoon water collected in December 1981 was analyzed for nitrite and nitrate according to the method of Thayer (1979), only nitrate was found (16.7 ppm); the pH of the lagoon waters ranged from 7.3 to 7.8, and the total dissolved solids concentration was 540 ppm.

Absorbance measurements were obtained at 290 and 400 nm as indicators of the light-absorbing properties of the lagoon water as a function of time of year. These values, reported in Table 5 along with the measured water volume at the sampling date, indicate that the UV-absorbing materials are concentrated as the lagoon waters evaporate, as evidenced by a 3-fold increase in the absorbance between December and August with the concurrent decrease in volume by a factor of three over the same period. This has the effect of reducing the penetrability of light into the lagoon water in the summer months compared to winter.



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## Table 4

Date of Sample	2-Amino-4,6-Dinitrobenzoic Acid (mg 1 <sup>-1</sup> )			
12/18/81	28.9			
03/15/82	24.3			
03/15/82	24.3			
05/25/82	30.7			
06/09/82	35.2			
06/23/82	43.9			
07/07/82	45.5			
08/04/82	49.9			
08/20/82	45.7			

## LEVELS OF 2-AMINO-4,6-DINITROBENZOIC ACID OBSERVED IN LAGOON NO. 9 ON VARIOUS SAMPLING DATES

#### Table 5

ABSORBANCE VALUES OF LAGOON NO.9 WATER AT 290 AND 400 nm AT VARIOUS TIMES OF THE YEAR

Absorbance	(1-cm cell)	Lagoon Volume,	
290 nm	400 nm		
3.19	1.41	1772	
3.21	1.38		
4.68	2.01	1270	
5.28	2.25		
6.09	2.56		
9.78	4.27	532	
8.62	3.74		
9.44	4.21		
	290 nm 3.19 3.21 4.68 5.28 6.09 9.78 8.62	3.19       1.41         3.21       1.38         4.68       2.01         5.28       2.25         6.09       2.56         9.78       4.27         8.62       3.74	

#### IV. SITE EVALUATION

On the basis of on-site observations and analysis of the lagoon waters, transformation products, and sediments, we conclude that the lagoon is a complex environment governed by chemical inputs that are dependent on lagoon water temperature and transformation rates, which in turn are dependent on time of year, natural sensitizers, and depth at which measurements are made.

Near the dump ramp we found levels of TNT and RDX in the sediment (15 and 8%, respectively) that were significantly greater than amounts that would result from sorption alone. This suggested that particulate munitions were present; in fact, when the lagoon surface became visible, chunks of pure TNT appeared on the surface sediment. The percent of munitions in the sediment decreased as a function of distance from the dump ramp. Therefore, chemical input to the lagoon is controlled by diffusion of chemical through the sediment and dissolution of pure chemicals from the sediment surface to the bulkwater phase. Because the bottom sediment is a heterogeneous system, we could only estimate chemical inputs based on limited laboratory investigations.

Transformation products found in the lagoon suggested that both biological and photochemical transformations were occurring in the lagoon (see above). Most of the light is absorbed in the first few millimeters of the lagoon; consequently, good vertical mixing is required to renew depleted chemicals as rapidly as they are transformed. If good mixing is achieved, then a photochemical rate constant can be estimated as a function of depth. Moreover, because absorption of the sunlight by the lagoon produces radiant heat, lagoon water temperature varies widely throughout the year; for example, it was 10°C in December, 41.5°C in July. This increase in temperature could have a large impact on microbial populations and hence on

biotransformation rates; in addition, it could accelerate diffusion and dissolution rates of chemicals from the bottom sediment.

Estimation of the concentration of a chemical at any particular time is further complicated by the water balance within the lagoon. Water enters the lagoon through precipitation and is lost through evaporation and seepage. Therefore, an understanding of water input and output rates is an integral part of this fate assessment.

On the basis of the above evaluations, we perceived a number of processes that were affecting the concentration of chemicals in the lagoon environment. Processes that would increase concentration of a chemical in the lagoon water include 1) water evaporation and seepage, 2) dissolution of a pure chemical from the sediment surface, and 3) diffusion of munitions from the sediment. Processes that would decrease the concentration in the water include 1) precipitation, 2) photochemical transformation, and 3) biological transformation. We therefore performed laboratory investigations of these processes to provide rate and equilibrium constants that are applicable to the lagoon environment. From these data, we constructed a computer model to simulate the processes in the lagoon and compared the simulation data with actual data obtained for the lagoon.

#### V. TRANSPORT AND TRANSFORMATION STUDIES

#### A. Physical Transport

#### 1. Water Balance

The volume of water in the lagoon is controlled primarily by precipitation, evaporation, and vertical seepage. Surface runoff or horizontal seepage was considered to have little impact on the net volume changes due to the flat bottom and banking of the lagoon.

We created a water-volume balance expression using:

$$\frac{dV}{dt} = R_p - R_{ev} - R_l , \qquad (1)$$

where  $R_p = r_p A$  $R_{ev} = r_{ev} A$  $R_{\ell} = r_{\ell} A$ 

and  $r_p$  represents precipitation,  $r_{ev}$  represents evaporation,  $r_{f}$  represents leaching (all in cm day<sup>-1</sup>), and A represents the surface area of the lagoon. Average values of  $r_{ev}$  and  $r_p$  were obtained from three years of weather data (1979-1981) collected by the Red River weather station near LAAP; these are shown in Table 6. Similar data for 1982 through August are presented in Table 7.

Leaching is usually estimated by a difference determination of volumes when the volume loss from evaporation is known. During the December field study we calculated the leaching rate to be 0.17 cm day<sup>-1</sup>. However, we used this parameter as a variable in the model to adjust the overall water balance to approximate that observed in the lagoon.

## Table 6

#### Precipitation Evaporation (cm/day) (cm/day) Month 1 0.28 0.13 0.30 2 0.20 3 0.46 0.35 4 0.34 0.45 5 0.61 0.52 6 0.28 0.67 7 0.39 0.69 0.12 8 0.66 9 0.18 0.50 0.44 10 0.38 0.25 0.20 11 12 0.16 0.17

### AVERAGE MONTHLY PRECIPITATION AND EVAPORATION DATA FROM 1979 THROUGH 1981 AT RED RIVER WEATHER STATION, RESTIN, LA

#### Table 7

### MONTHLY PRECIPITATION AND EVAPORATION DATA FROM JANUARY 1982 THROUGH AUGUST 1982 AT RED RIVER WEATHER STATION, RESTIN, LA

Month	Precipitation (cm/day)	Evaporation (cm/day)		
Jan	0.31	0.13		
Feb	0.26	0.15		
Mar	0.19	0.29		
Apr	0.27	0.34		
May	0.17	0.59		
June	0.22	0.61		
July	0.25	0.62		
Aug	0.17	0.62		

Using the data in Tables 6 and 7, we plotted the depth of the lagoon as a function of time of year for various seepage rates (Figure 6). From depth measurements taken at various time points in the lagoon we found that a seepage rate of 0.12 cm day<sup>-1</sup> gave the best approximation to the actual depths. This seepage rate projected the lagoon to go dry on Day 234 (August 22); the lagoon actually went dry during the following week. We therefore used the depth profile defined by the 0.12 cm day<sup>-1</sup> seepage rate curve to determine the water volume on any particular day and to define the depth that was critical in estimating the magnitude of the photochemical rate constant at any particular time.

### 2. TNT and RDX Solubility

The aqueous solubilities of TNT and RDX were determined separately at 10, 20, and 30°C and simultaneously at 20°C by the method of May et al. (1978). Briefly, glass beads were coated with TNT or RDX and packed into a column. Then water was percolated through the column. After an equilibration period, the concentration of TNT or RDX was measured in the column effluent. As a check on this method, the solubilities of TNT and RDX were measured at 20°C by the method of Campbell (1930), in which the solids are equilibrated with water in a thermostated bath and filtered. The solubilities using the two methods are shown in Table 8.

#### Table 8

#### SOLUBILITY OF TNT AND RDX IN WATER AT 10, 20, AND 30°C

	TNT	(mg/lite	r)	RDX		
Method	10°C	20°C	30°C	10°C	20°C	30°C
May et al. Campbell	67 ± 1	$104 \pm 2$ 113 ± 8	165 ± 2	21.9 ± 0.3	$38.4 \pm 0.4$ 39 ± 2	67.0 ± 0.8


FIGURE 6 ESTIMATION OF LAGOON DEPTH PROFILE BASED ON VARIABLE SEEPAGE RATES

Solubilities were measured by the May method first at 10°, then 20°, then 30°C, and finally again at 10°C. No significant differences were noted between the two measurements at 10°C. The similarity of the measurements at 10°C demonstrates the repeatability of the method after equilibrating the column and showed that the column was not exhausted. Concentration versus flow-rate measurements were made to check that the saturation capacity of the column had not been exceeded. Finally, a column containing both RDX and TNT was used to measure the solubility of one compound in the presence of a saturated solution of the other compound. Solubilities obtained for RDX and TNT independently were in agreement within the standard error of the values for RDX and TNT in the presence of the other compound.

The results of the two methods are in agreement within the error of each measurement. The Campbell solubility values show a greater error than those using the May method, perhaps because the Campbell method requires more handling of the samples.

Because the temperature of the lagoon water ranged from 10° to 41°C, we developed mathematical expressions to use in the computer modeling to correlate solubility with temperature. These expressions are shown in Equations 2 and 3:

> $S_{TNT} = 5.567 \times 10^7 \exp(\frac{-3860^{\circ}K}{T}) \mu g m 1^{-1}$ ; (2)  $S_{RDX} = 4.974 \times 10^7 \exp(\frac{-4795^{\circ}K}{T}) \mu g m 1^{-1}$

(3)

The measured and calculated solubility values for TNT and RDX at 10°, 20°, and 30°C (283, 293, and 303°K) are shown in Table 9.

MEASURED AND CALCULATED SOLUBILITIES (mg liter<sup>-1</sup>) OF TNT AND RDX AT 283, 293, AND 303<sup>°</sup>K

Water Temperature (°K)	Т	NT	RDX			
(°K)	Measured	Calculated	Measured	Calculated		
283	67	66.4	21.9	21.8		
293	104	105.7	38.4	38.9		
303	165	163.3	67	66.7		

#### 3. Sediment Sorption

It is generally accepted that adsorption of organics on soils and sediments usually fits the empirical Freundlich isotherm (Equation 4):

$$C_{g} = KC_{w}^{n}$$
, (4)

where  $C_s$  is the equilibrium concentration on the sediment (µg g<sup>-1</sup>),  $C_w$ is the equilibrium concentration in the aqueous phase (µg ml<sup>-1</sup> = µg g<sup>-1</sup>), and K and n are constants. Thus, a plot of the logarithms of  $C_s$ versus  $C_w$  has a slope of n and an intercept of K. If n is equal to 1, the adsorption isotherm is linear and K is equal to  $K_s$ . If pure TNT and RDX are present, as in the LAAP lagoon,  $C_w$  is the saturated concentration regardless of  $C_s$  and the Freundlich isotherm may not hold.

We performed several isotherm measurements and fitted the data to both the linear  $(C_g = K_p C_w)$  and nonlinear  $(\log C_g = \log K_p + n \log C_w)$ cases. Very high values of  $K_p$  were obtained from the nonlinear equation and were rejected in favor of more reasonable values obtained from the linear equation. These  $K_p$  values, along with the calculated  $K_{oc}$  values (based on a measured organic carbon content of 1.5%), appear in Table 10.

# SORPTION PARTITION COEFFICIENTS FOR TNT AND RDX IN LAAP LAGOON NO. 9 SEDIMENT

Chemical	к <sub>р</sub>	Koc
TNT	7 ± 2	470 ± 130
RDX	4 ± 7	270 ± 470

# 4. Distribution of TNT and RDX in Sediment Cores

Several core samples were taken throughout the lagoon during the December 1982 visit to determine the horizontal and vertical distribution of RDX and TNT in the lagoon sediment. A 51-cm core sediment sample from the southcast corner of the lagoon was cut into 5-cm segments. Each segment was analyzed for percent TNT, RDX, and dry sediment. The results are presented in Table 11.

#### Table 11

#### SEDIMENT CORE SAMPLE DRY WEIGHT AND PERCENT TNT AND RDX WEIGHTS

Core Segment (cm from top)	% Dry Weight	7 TNT <sup>3</sup>	% RDX <sup>a</sup>
5	71.5	9.34	12.6
5 - 10	76.8	2.74	1.28
10 - 15	79.4	2.38	0.83
15 - 20	80.3	1.25	0.39
20 - 25	83.2	0.14	0.004
25 - 31	82.8	0.010	0.005
31 - 36	82.3	0.0052	0.0065
36 - 41	82.3	0.0052	0.0065
41 - 46	83.2	0.023	0.029
46 - 51	74.2	0.04	0.04

<sup>a</sup>Based on dry weight of sediment.

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Saturated sediment is represented by a concentration of >0.60%for TNT and >0.033% for RDX. Although the TNT and RDX concentrations in the sediment decrease with depth, the data suggest that the amounts exceed the absorption capacity for TNT and RDX at depths less than 25 cm and greater than 50 cm, respectively. The high amounts of TNT and RDX in the surface sediments suggest that the sediment is a source of these munitions in the lagoon water.

To determine the horizontal distribution of TNT and RDX, we took another set of core samples--three from the southeast (SE) corner and one each from the northwest (NW), southwest (SW), and northeast (NE) corners of the lagoon. The cores were about 12 cm in depth and were cut into 1-cm segments, which were extracted and analyzed by HPLC for their RDX and TNT content. The results are presented in Table 12.

Only the top and bottom three sections of each core were analyzed. Concentrations of TNT and RDX were highest in the SE cores, which are near the dump ramp, and they rapidly decrease with depth, as shown by comparing the top three sections with the bottom three. The TNT and RDX concentrations in the NW, NE, and SW cores are low and uniform throughout the core as compared with the SE cores.

#### 5. Mass Transfer Rate From Sediment

The presence of sorbed and solid munitions chemicals in the LAAP sediment presents a complex problem in establishing the mass transfer rate of chemicals to the bulk-water phase. The processes that may be occurring in a sediment pore, described pictorially in Figure 7, include diffusion, seepage, sorption-desorption equilibria, transformation, and dissolution of solid munitions. These processes can be described mathematically and their sum can be used to compute the overall mass transfer rate.

Diffusion is generally accepted as a second-order differential expression relating changes in the chemical concentration in the sediment  $(C_{gw})$  with respect to depth (z) times a diffusion constant  $D_1$ . Seepage is a first-order differential expression relating



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FIGURE 7 LAGOON WATER RESTS ON SEDIMENT

Pond and sediment are considered to be semi-infinite.

concentration to depth times a seepage velocity factor, v. The sorption-desorption equation was defined in Equation 4. The only transformation process believed to be occurring in the sediment is

### Table 12

TNT AND RDX IN SECTIONED CORES FROM LAAP LAGOON NO. 9

		SE-B	SE-C	NW	SW	NE
		_		-1、		
		T	NT (μg g	-)		
12					0.05	
11	501	6.4	367	0.04	0.55	0.03
10	525	71	38	0.05	0.23	0.1^
9		109	<b>28</b> 0	0.02	0.08	
3	91	0.4	5.5	0.14	0.16	0.009
2	27	1.2	2.9	0.09	0.28	0.03
1	26	0.7	1.4	0.06	0.13	0.05
		RI	DX (μg g	-1)		
12					0.10	
11	<b>89</b>	3.0	88	0.01	0.01	0.02
10	48	22	226	0.02	0.02	0.01
9		51	88	0.01	0.01	
3	4.7	0.2	2.1	0.05	0.05	0.00
2	1.3	0.3	0.8	0.05	0.05	0.02
1	1.3	0.3	0.9	0.01	0.01	0.04

Section Identification

<sup>a</sup>Sample 12 is at surface, sample 1 is lowest below surface.

biotransformation. Because of the high concentrations of munitions in the sediment, biotransformation is believed to be a zero-order process  $(k_o)$  for TNT (see Biotransformation, Section V.C).

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To account for the dissolution, we envision solid TNT/RDX dispersed homogeneously throughout the sediment. The dissolution rate of any solid TNT or RDX,  $\frac{dw}{dt}$ , can be written as Equation 5:

$$\frac{dw}{dt} = k_r (C_{sat} - C_i) , \qquad (5)$$

where  $k_r$  is an experimentally determined dissolution coefficient,  $C_{sat}$  is the saturation concentration of the dissolving species in the liquid phase, and  $C_i$  is the actual concentration of the dissolving species at the solid/liquid interface.<sup>\*</sup> If diffusion from the interface into the surrounding liquid is fast compared to dissolution, then  $C_i$  is approximately equal to the concentration of the dissolved species in the surrounding liquid. In this limit,

$$\frac{dw}{dt} = k_r (C_{sat} - C_{sw}) \qquad (6)$$

Therefore, in the sediment, the change in chemical concentration with time  $(\frac{\partial C}{\partial t})$  is equal to the sum of the dissolution rate (Equation 6), diffusion rate  $(D_1 - \frac{\partial^2 C_{gw}}{\partial z^2})$ , and seepage rate  $(\frac{\partial C_{gw}}{\partial z})$  minus the biotransformation rate  $(k_o, \text{ zero-order process in the sediment)}$  and the amount of chemical that is sorbed to the sediment  $(K_p n C_{gw}^{n-1} - \frac{\partial C_{gw}}{\partial t})$  or

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<sup>\*</sup>This is a standard "concentration driving force" model that has been adopted by many workers (Wurster and Kildsig, 1965).

$$\frac{\partial C_{sw}}{\partial t} = k_{dis} [C_{sat} - C_{sw}] + D_1 \frac{\partial^2 C_{sw}}{\partial z^2} + v \frac{\partial C_{sw}}{\partial z} - k_o - K_p n C_{sw}^{n-1} \frac{\partial C_{sw}}{\partial t}$$
(7)

Rearranging Equation 7 yields Equation 8:

$$(1 + K_{p}nC_{sw}^{n-1}) \frac{\partial C_{sw}}{\partial t} = k_{dis}[C_{sat} - C_{sw}] + D_{1}\frac{\partial^{2}C_{sw}}{\partial z^{2}} + v\frac{\partial C_{sw}}{\partial z} - k_{o}$$
(8)

where  $C_{sw}$  = chemical concentration in the sediment/water phase

 $K_p$  = sediment partition coefficient ( $C_s = K_p C_{sw}^n$ )

t = time

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- $D_1$  = diffusion coefficient in the sediment/water phase
  - v = seepage velocity
  - z = distance from the sediment
- k<sub>o</sub> = zero-order rate constant for biotransformation in sediment.

Similarly, for the free-water phase, the change in concentration of a chemical with time can be described by

$$\frac{\partial C_{w}}{\partial t} = k_{dis} [C_{sat} - C_{w}] + D_{2} \frac{\partial^{2} C_{w}}{\partial z^{2}} + v_{\overline{\partial z}}^{\partial C_{w}} - \Sigma k_{i} C_{w} , \qquad (9)$$

where  $C_w$  = concentration of chemical in the free-water phase t = time  $D_2$  = diffusion coefficient in the free-water phase v = seepage velocity  $\Sigma k_i C_w$  = first-order rate expression for the transformation process.

Solutions to these equations and the assumptions that are used to arrive at solutions appear in Appendix B. However, laboratory studies were necessary to determine  $k_{dis}$  so that a flux from the lagoon sediment could be computed.

#### 6. Determination of TNT and RDX Flux Values for Sediment from LAAP Lagoon No. 9

From the sediment analyses and the visual observations of pure TNT and RDX on the sediment surface, we recognized that the bottom sediment was the predominant input source of TNT and RDX to the lagoon waters during the period of our study. Therefore, we designed experiments to determine the flux of these chemicals to the bulk-water phase.

To determine the fluxes, 460 g of LAAP Lagoon No. 9 sediment containing 12% TNT and 9% RDX was placed in a 1000-ml beaker (surface area, 780 cm) equipped with a water-inlet port directly above the sediment and a water-outlet port near the top of the beaker. The beaker was wrapped in aluminum foil. Water was pumped into the beaker from a Milton-Roy precision pump at various flow rates and stirred slowly with an overhead stirrer. The effluent was collected to verify total water volumes passing through the beaker. TNT and RDX in the effluent were monitored until equilibrium values were obtained (usually in 3 to 4 days). We calculated the flux, N, from Equation 10:

$$N = \frac{CQ}{A} , \qquad (10)$$

where C = concentration of chemical at equilibrium

Q = flow rate through the beaker

A = surface area of the sediment.

Values for the fluxes of TNT and RDX at various flow rates are shown in Table 13. We would expect that as the flow rate increased, the concentration of chemical would decrease and the flux would increase. The changes in concentration between runs A and B for TNT and B and C for RDX do not follow the proper concentration trend. Also, the changes in flux between runs D and E for TNT and RDX do not follow the proper flux trend. These observations cannot be explained by any mass transfer model; however, we cannot be certain whether any of the data points are more or less valid than others and must

# TNT AND FLUX MEASUREMENTS FROM LAAP LACOON NO. 9 SEDIMENT

RDX Flux (N) (10 <sup>-9</sup> g cm <sup>-2</sup> sec <sup>-1</sup> )	2.3	3.2	3.8	5.2	3.4
RDX (C) (μg ml <sup>-1</sup> ) (	5.6	5.0	5.5	3.5	2.0
TNT Flux (N) (10 <sup>-9</sup> g cm <sup>-2</sup> sec <sup>-1</sup> )	6.8	12.7	12.9	20.5	9.8
ТNT (C) (нg ml <sup>-1</sup> ) (	16.9	6.91	18.5	13.8	5.7
Run Code Water Flow Rate (Q) TNT (C) (Number) (ml min <sup>-1</sup> ) (μg ml <sup>-1</sup> )	1.9	3.0	3.3	7.0	8.1
Run Code (Number)	A (2)	B (5)	c (3)	D (4)	E (1)

allow for the imprecision of the data in our interpretations. A plot of TNT and RDX flux versus concentration appears in Figure 8.

From this plot and linear least-squares analysis (Appendix B) we calculated the TNT flux to be  $1.08 \times 10^{-4}$  cm sec<sup>-1</sup> and the RDX flux to be  $7.75 \times 10^{-5}$  cm sec<sup>-1</sup> in the lagoon. The first-order dissolution rate constants were calculated to be  $1.18 \times 10^{-5}$  cm sec<sup>-1</sup> for TNT and  $1.16 \times 10^{-5}$  cm sec<sup>-1</sup> for RDX (see Appendix B).

#### B. Photochemistry

Photochemical transformation of TNT in the lagoon environment was evident from the identification of photolytic transformation products and the deep orange-red color of the water that is reminiscent of the "pink water" at TNT production facilities. Because lagoon water absorbs all UV light near the surface, the photochemical rate constant has a depth dependence of the form  $k'_p = k_p/D$ . Moreover, the value of  $k_p/D$  also is subject to variations owing to changes in light intensity with changes in cloud cover and season.

#### 1. TNT Photochemical Rate Constant--General Remarks

Usually, photochemical kinetic studies are performed using solutions that absorb less than 5% of the incident light. The amount of light absorbed at wavelength  $\lambda$  by a solution is given by the Beer-Lambert Law (Equations 11 and 12):

$$I_{\lambda} = I_{0}(1 - 10^{-A})$$
 , (11)

$$A = \varepsilon_{\lambda} C \ell + \alpha_{\lambda} \ell , \qquad (12)$$

where  $I_0$  and  $I_{\lambda}$  are the incident and absorbed light intensities, respectively, A is the absorbance,  $\ell$  is the path length of light in solution, C is the concentration of chemical,  $\varepsilon_{\lambda}$  is the molar absorption coefficient of a chemical at wavelength  $\lambda$ , and  $\alpha_{\lambda}$  is the





absorbance of solution without added chemical at wavelength  $\lambda$  in a l-cm cell. The  $\alpha_\lambda$  term is useful for characterizing a natural water because the molar concentration of humic acids and their  $\varepsilon_\lambda$  values in water are difficult to determine. Because  $I_\lambda$  is a function of pathlength 1, the value for  $I_\lambda$  can be calculated for any pathlength in solution.

If the solution absorbs less than 2% of the incident light (A < 0.02), Equation 11 takes the form of Equation 13:

$$I_{\lambda} = 2.3I_{0}A$$
 , (13)

and the rate constant for photolysis,  $k_{p}$ , is given by Equation 14:

$$k_{\rm p} = 2.3 \phi \epsilon_{\lambda} I_{\rm o} \ell \qquad (14)$$

where  $\phi$  is the photolysis reaction quantum yield (Mabey ot al., 1982).

However, if the solution absorbs a significant amount of light due to some combination of long pathlength and/or large  $\varepsilon C$  or  $\alpha_{\lambda}$ values, then the photolysis rate constant will show a dependence on pathlength due to attenuation of  $I_0$  with increased absorbance, and the resulting equation for  $k_p$  is more complicated.

The solar photolysis rate constant  $k_{pE}$  for chemicals in aquatic systems in optically thin solutions (i.e., near the surface) can be calculated from Equation 15:

$$k_{\rm pE} = b\phi \Sigma \varepsilon_{\lambda} I_{\lambda} , \qquad (15)$$

where b is a unit conversion factor and  $\varepsilon_{\lambda}I_{\lambda}$  values for each wavelength interval are summed over the entire solar spectral region. If the water has a large  $\alpha_{\lambda}$  value, Equation 15 becomes more complex because the  $\alpha_{\lambda}$  values decrease as wavelengths become longer, and therefore longer wavelength light penetrates natural waters to greater depths than does shorter (more energetic) wavelength light.

Table 14 shows the  $(\alpha_{\lambda} + \epsilon_{\lambda}C)$  values at selected wavelengths for the LAAP water collected in December 1981 and the depths at which 99% of incident light at each wavelength is absorbed. These data clearly show that light of wavelengths below 500 nm will not penetrate below the top few centimeters of LAAP water. The absorbance of the water is almost entirely due to substances other than TNT ( $\alpha_{\lambda} \gg \epsilon_{\lambda}C$ ); absorbance of 6 ppm TNT in a 1.0-cm cell is 0.02 AU at 320 nm ( $\epsilon_{\lambda} =$  $840 \text{ M}^{-1} \text{ cm}^{-1}$ ), which is negligible compared to the 2.60 AU measured for the water at 320 nm.

#### Table 14

# DEPTH FOR 99% ABSORPTION OF LIGHT OF SPECIFIC WAVELENGTHS FOR LAAP WATER

Wa	velength,	elength, $(\alpha_{\lambda} + \epsilon_{\lambda}C)^{a}$			
λ,	(nm)	(cm <sup>-1</sup> )	D <sup>99%</sup> A (cm)		
	300	3.32	0.60		
	320	2.60	0.77		
	340	2.32	0.86		
	360	2.10	0.95		
	380	1.93	1.04		
	400	1.74	1.15		
	420	1.47	1.36		
	440	1.23	1.62		
	460	0.95	2.11		
	480	0.74	2.71		
	505	0.58	3.45		
	585	0.28	7.04		
	655	0.07	27.00		

<sup>a</sup>Absorbance measured in 1.0-cm cell; light absorp ion is entirely due to substances other than TNT in the water (see text). The TNT photolysis rate constant for use in modeling the fate of TNT in LAAP water must account for the loss of TNT in the entire volume of the lagoon. Although the mass of TNT undergoing photolytic transformation per unit time is dependent only on light intensity and spectral distribution entering the water, the rate of change of TNT <u>concentration</u> is dependent on volume. For photolysis of TNT in solution out of the photic zone, the rate constant must then be an inverse function of depth. The theory and validation of the procedure for estimating the photolysis rate constant in LAAP water where the photic zone is shallow compared to the total depth of the water are developed and discussed in the next section.

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# 2. Effect of Depth on Photolysis Rate Constant of TNT in LAAP Water

To estimate a rate constant for TNT in LAAP water, we have developed an expression that relates the TNT photolysis rate constant to lagoon depth (D). This expression is valid only if vertical mixing in the lagoon is substantially faster than the photolysis rate of TNT near the surface so that the TNT concentration is the same throughout the lagoon. The expression also assumes that TNT decomposes by indirect photolysis; the data in Table 14 clearly show that substances other than TNT account for the major absorption of light, and our previous studies have demonstrated that indirect photolysis of TNT in natural waters is more rapid than direct photolysis (Spanggord et al., 1981). Light energy is absorbed by substances present in LAAP water (humic and fulvic acids, TNT reaction products, and other chemical contaminants in the lagoon water) and is then transferred to or induces reaction of TNT. TNT reacts even though it does not absorb light directly.

The proposed mechanism for TNT photolysis is described by Equations 16 to 19 below:

$$S + h_{U} + S^{*}$$
, (16)

$$s + s^{k_2}$$
, (17)

$$\overset{k_3}{s} + TNT \rightarrow S + TNT , \qquad (18)$$

where S is the sensitizer, S\* and TNT\* are excited states, and f is the fraction of TNT converted to products. The rate of photoabsorption, R (Equation 16), per unit volume is shown in Equation 20:

$$R = I_{O}A/V = I_{O}/D , \qquad (20)$$

where  $I_0$  is the incident light intensity (Einstein's per unit area per second), A is the exposed surface area, V is the volume of the container, and D is the average depth.

The quantum yield,  $\phi$ , for TNT loss is defined in Equation 21:

$$\phi = \phi_{s*} k_3 [TNT] f / (k_3 [TNT] + k_2) , \qquad (21)$$

where  $\phi_{S*}$  is the quantum yield for production of S<sup>\*</sup>. In dilute solutions of TNT, where  $k_2 \gg k_3$ [TNT],

$$\phi = \phi_{S \neq k_3}[TNT]f/k_2 \qquad (22)$$

The loss rate of TNT can be defined as the TNT quantum yield times the rate of light absorption of the water body. Then,

$$d[TNT]/dt = \phi R = I_0 \phi_{S*} k_3 [TNT] f/k_2 D$$
, (23)

$$d[TNT]/[TNT] = fI_0 \phi_S k_3/k_2 D$$
 (24)

On rearrangement and integration,

$$\ln([TNT]_{0}/[TNT]_{t}) = fI_{0}(k_{3}/k_{2})\phi_{s}t/D = k_{p}t$$
(25)

and

$$k_{p} = fI_{o}(k_{3}/k_{2})\phi_{S*}/D$$
 (26)

 $I_o$  and D depend on sky conditions and lagoon geometry, respectively. D can be calculated for water in a container with vertical sides and a horizontal bottom.

By making the substitution  $k'_p = I_0 f(k_3/k_2)\phi_{S*}$  into Equation 26, we obtain Equation 27, which states that  $k_p$  will vary inversely with depth:

$$k_{pD} = k_{p}^{*}/D \qquad (27)$$

For containers of depth  $D_1$  and  $D_2$ , respectively, relative photolysis rate constants can be calculated from Equation 28:

$$\frac{k_{pD_{1}}}{k_{pD_{2}}} = \frac{D_{2}}{D_{1}} .$$
 (28)

Using Equation 28, we can calculate the TNT photolysis rate constant in the lagoon from the sunlight photolysis rate constant measured for TNT in LAAP water in a shallow container. This is the only practical way to measure  $k_p$  because TNT in LAAP lagoon water photolyses too slowly in the lagoon or in containers of the same depth for convenient measurement.

#### 3. Experimental Verification of Equation 27

To demonstrate the validity of Equation 27, we first photolyzed LAAP water with a constant intensity source in a merry-go-round apparatus (MGR) to determine the TNT concentration-time dependence of Equation 25. If Equation 25 holds, then a first-order plot ( $\ln[TNT]$ ) versus time) will be linear. For these experiments, LAAP water containing 5 ppm TNT was irradiated in 13 × 100-mm borosilicate tubes with monochromatic light at 313 or 366 nm at room temperature.

The data for these experiments are summarized in Table 15 along with data for the same solution in tubes exposed to sunlight on January 22, 1982.

#### Table 15

#### PHOTOLYSIS OF 4.97 ppm TNT IN LAAP WATER IN AN MGR APPARATUS AND IN SUNLIGHT

Light Source	Duration of Experiment (min)	Final TNT Concentration (ppm)	$\frac{10^3 k_p (min^{-1})}{10^3 k_p (min^{-1})}$
313 nm	110	3.29	4.30 ± 0.07
366 nm	240	2.34	$3.17 \pm 0.11$
Sunlight (Jan. 22)	80	2.49	8.51 ± 0.23

First-order treatment of these data gave linear plots, as expected. Similar first-order behavior was observed in all other experiments.

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We then verified Equations 27 and 28 by irradiating LAAP water in cylinders of different depths using a solar simulator light source--an intensity-regulated, 300-watt xenon lamp. Radiation from the lamp was filtered through water to remove infrared wavelengths and then through borosilicate glass to remove high-energy UV light. Cylinders were filled to depths of 2.0, 4.2, and 7.0 cm and irradiated from above. The sides and bottoms of the cylinders were blackened to reduce reflection and to prevent light from entering through the sides. Data from these experiments are shown in Table 16.

The observed photolysis rate constants are in the ratio 4.05:1.75:1.0; the expected values are 3.5:2.1:1.0. Stirring had no `ffect on the photolysis rate constant: when the stirrer was turned off after 240 min of irradiation of the dish containing 4.2 cm LAAP water, the rate constant did not change. We conclude that natural mixing of the LAAP water in our experiments was sufficiently fast to keep the mixture homogeneous and not allow depletion of TNT in the upper irradiated layers, which would slow down the reaction.

Duration of Experiment (min)	Water Depth (cm)	Final TNT Concentration (ppm)	$\frac{10^3 \text{ k}_p (\text{min}^{-1})^b}{2}$
236	2.0	1.81	4.21 ± 0.09
542	4.2	1.81	1.82 ± 0.02
1075	7.0	1.33	$1.04 \pm 0.03$

#### PHOTOLYIS OF 4.97 ppm TNT IN LAAP WATER AS A FUNCTION OF DEPTH USING A XENON LAMP<sup>a</sup>

<sup>a</sup>300-W controlled-current lamp.

 $b_r^2 = 0.99$ .

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Finally, we exposed LAAP water to similight to measure TNT rate constants that could be adjusted for depth. Actimometer solutions of PNA/PYR were exposed to sunlight in parallel studies so that corrections for sky conditions could be made (discussed in the next section). These data permit calculation of the TNT photolysis rate constant in the LAAP lagoon for any water depth.

To measure the depth-independent photolysis rate constant k' for TNT exposed to sunlight, LAAP water filtered through a 0.45-micron filter was placed in opaque glass cylinders whose tops were exposed to sunlight on March 12 and 15, 1982. On the first day the container was filled to a 4.2 cm depth and on the other day the depth was 3.5 cm. The experimental conditions and results are given in Table 17, uncorrected for sky conditions.

If we assume that an insignificant amount of TNT photolysis occurred before the experiment began and after the experiment was ended on each day, the measured rate constant  $k_p$  (in m<sup>-1</sup>) can be readily converted into a 24-hour day-averaged rate constant  $k_p$  (in

 $d^{-1}$ ) by multiplying the observed rate constant by the fraction of a 24-hour day that the irradiation period represented. Multiplication of the latter by the depth value then gives a value of 3.0 cm  $d^{-1}$ , which is the depth-independent rate constant,  $k_p^*$ , used in Equation 26 for photolysis of TNT in LAAP water under weather conditions of mid-March. To obtain a rate constant for a specific depth,  $k_p^*$  is divided by the lagoon depth. We must also correct  $k_p^*$  for sky conditions during the experiments.

#### Table 17

SUNLIGHT PHOTOLYSIS OF 4.97 ppm TNT IN LAAP WATER<sup>a,b</sup> AND ACTINOMETERS

Date Time	3-12-82 8:45-16:00	3-15-82 9:40-15:30
Sky Conditions	bright, sunny w/ low clouds	cloudy + hazy, windy
Depth, cm	4.2	3.5
Time, min	435	350
$k_{p} \times 10^{3}, m^{-1}$	1.67 ± 0.045	$2.51 \pm 0.041$
t 1 <sub>/2</sub> h	6.9	4.6
k <sub>p</sub> d <sup>-1</sup> (day averaged) <sup>c</sup>	0.72	0.89
k' cm d <sup>-1</sup>	3.02	3.12
k' cm d <sup>-1</sup> , p clear sky conditions	4.14	4.27

<sup>a</sup>Photolyses monitored using PNA/PYR actinometer [PYR] =  $1.25 \times 10^{-3}$  M; k<sub>p</sub> (PNA) =  $4.8 \times 10^{-3}$  m<sup>-1</sup>.

<sup>b</sup>All rate constants are uncorrected for sky conditions.

 $c_{k_p}$  (day averaged) was calculated by multiplying the observed photolysis rate constant expressed in units of  $d^{-1}$  by the fraction of a 24-hour day that the irradiation represented.

# 4. Correction of Measured Rate Constants for Weather Conditions

Sunlight photolysis rate constants are dependent on weather conditions as well as the diurnal and seasonal light intensity and spectral variation. Diurnal and seasonal variations in light intensity (in terms of photon flux) and distribution are included in procedures to calculate direct photolysis rate constants of chemicals in optically thin solutions (Mill et al., 1982). These rate constants are estimated for clear-sky conditions and can be calculated as a 24-hour day-averaged rate constant. Corrections for weather conditions (overcast or cloudy cover) must be made using field data; they were performed as follows.

During the LAAP field study and in our outdoor studies at SRI, we exposed PNA/PYR actinometer solutions to sunlight simultaneously with our LAAP water/TNT experiments. The photolysis rate constants for this actinometer are available for clear-sky conditions as a function of season (Dulin and Mill, 1982). Rate constants calculated for PNA/PYR, with [PYR] =  $1.25 \pm 10^{-3}$  M, were  $4.3 \times 10^{-3}$  m<sup>-1</sup> for January and  $9.6 \times 10^{-3}$  m<sup>-1</sup> for late April [winter and spring season dates, respectively, using the solar program of Zepp and Cline (1977)]. By interpolation, we estimate that an average rate constant of about  $6.6 \times 10^{-3}$  m<sup>-1</sup> would then be expected for photolysis of PNA/PYR in mid-March. Because the actual rate constant measured for photolysis of PNA/PYR on both days was  $4.8 \times 10^{-3}$  m<sup>-1</sup>, or 0.73 of that expected for clear-sky condition, we conclude that 73% of the clear-sky irrad-jance was available during our experiments with LAAP water.

The measured rate constant, k', should therefore be divided by 0.73 to obtain the clear-sky rate constant. The depth-dependent rate constant for photolysis of TNT in LAAP water in mid-March under clear-sky conditions is 3.1/0.73 = 4.3 cm d<sup>-1</sup>. For LAAP water of 50-cm depth, the rate constant of  $8.6 \times 10^{-2}$  d<sup>-1</sup> corresponds to a half-life of 8 days.

The above calculations assume that the 0.73 weather correction factor,  $f_w$ , obtained from the actinometer experiment, applies to light that induces photolysis of TNT in LAAP water. Although we know that this assumption is not strictly correct, it is sufficiently accurate for estimating environmental photolysis rate constants of chemicals that absorb in the same region of the spectrum. The assumption may not hold in the case of LAAP water where light absorption by products occurs into the visible region.

#### 5. Effect of Time of Year on Photochemical Rate Constant of TNT in LAAP Water

Because concentrations of TNT were found to persist in the LAAP water at all times of year, in part due to dissolution of TNT out of the sediments (see Section V.A.5), modeling of the fate of TNT in the pond water required information on how the photolysis rate constant of TNT in the water will vary with the time of year (i.e., seasonal rate constants are needed). Equation 25 shows that the photolysis rate constant of TNT is seen to be a function of the incident light intensity and the reaction quantum yield,  $\phi$ . Although we have no information on the rate of photolysis of TNT in LAAP water as a function of season, the relative rate constants for different times of the year can be calculated if we assume that  $L_{\lambda}$  values of Mabey et al. (1982) are useful approximations of  $L_{\lambda}$ . Then

$$k_{pE}(1)/k_{pE}(2) = \frac{\Sigma L_{\lambda 1}}{\Sigma L_{\lambda 2}}$$
, (29)

where 1 and 2 denote different times of the year. This expression assumes that the quantum yield for photolysis of TNT in LAAP water is the same at all seasons. Use of  $L_{\lambda}$  values to estimate the  $I_{\lambda}$  values for this calculation is not strictly correct because  $L_{\lambda}$  values are day-averaged photon flux values for light passing through optically thin water, and therefore they are corrected for path length and the refractive index of water. The actual flux values that should be used in the calculation are the flux values incident on the surface of the water with no path length correction because all light is absorbed by the water. This flux is referred to as  $W_{\lambda}$  by Zepp and Cline (1977), However, since the day-averaged analogs of  $W_{\lambda}$  are not known, we have used the  $L_{\lambda}$  values as approximations of the  $I_{\lambda}$  values needed for the calculation; comparison of  $W_{\lambda}$  and  $Z_{\lambda}$  values listed by Zepp and Cline for the season dates at 40° latitude indicates that the flux values corrected for path length are larger than the uncorrected values by 10-20% over a wide wavelength range.

The calculation of the relative photolysis rates of TNT in LAAP water also requires the knowledge or an assumption that the absorption of light above a certain wavelength will not photolyze TNT due to energy limitations. Therefore, we used several wavelength-cutoff regions to calculate the ratio in Equation 28 for the summer and winter solstices and the ratio for the September and March dates during which outdoor experiments were performed (Table 18). The data in the table show that photolysis of TNT in LAAP water will vary by a factor of about 3 between winter and summer seasons if only sunlight variations are considered; as discussed in other sections, the depth of the water, weather conditions, and possible changes in the nature of the lagoon water itself (absorbance and sensitizing abilities) will also affect the photolysis rates of TNT in the LAAP water throughout the year.

### VARIATION IN TNT PHOTOLYSIS RATES IN LAAP WATER AS A FUNCTION OF TIME OF YEAR<sup>a</sup>

	$\Sigma^{\alpha}\lambda^{L}\lambda^{L}\lambda^{L}\lambda^{2}$				
Wavelength <sup>D</sup> Cutoff	Summer season Winter season	September <sup>d</sup> March			
400 mm	3.3	1.8			
500 nm	2.9	1.8			
600 nm	2.8	1,8			

<sup>a</sup>Calculated using  $\alpha_{\lambda}$  values for LAAP water collected in July.

 ${}^b\alpha_\lambda$  and  $L_\lambda$  values above this wavelength not used in calculation of rate constant ratio.

 $^{c}L_{\lambda}$  value used for summer and winter solstices at 40° latitude.

 $d_{L_{\lambda}}$  values used were interpolated between season dates.

#### 5. UV-Visible Spectra of LAAP Water

As discussed above, the photic zone of the December LAAP water was thin (1-2 cm) compared to the total depth of the water (~60 cm) measured during the December site visit. In subsequent trips in May and July we found that the water depth was decreasing; therefore the absorption spectrum of the water was measured before use in our experiments. The UV spectrum of LAAP water in December is shown in Figure 9. The spectrum of the May LAAP water was nearly the same as that of the December water, whereas the absorption coefficients of July water were 1.7 to 1.5 times greater than those of the December water (Table 19). From the information and analyses obtained in our field and laboratory studies, it is impossible to determine how the constituents of the water affecting the UV spectra changed as a function of the time of year, although it is clear that the factor of 1.5-1.7 shown in Table 19 is less than a 3-fold increase expected from a simple concentration effect, considering that the water depth of  $\sim 60$ cm in December decreased to approximately 18 cm in July.

### Table 19

	α <sub>λ</sub> ,	α <sub>λ</sub> (July)		
λnaa	Dec LAAP	July LAAP	$\alpha_{\lambda}$ (Dec)	
300	3.38	5.82	1.72	
400	1.76	3.02	1.72	
500	0.66	1.06	1.62	
600	0.22	0.34	1.53	

#### ABSORPTION COEFFICIENT<sup>a</sup> OF DECEMBER AND JULY LAAP WATERS AT SELECTED WAVELENGTHS

<sup>a</sup>Absorbance of the natural water in a 1.0-cm cell.



FIGURE 9 ABSORPTION SPECTRUM FOR LAGOON 9, SITE 1 WATER, FILTERED, 0.1-cm CELL

#### 7. Experimental Studies of the Photolysis of TNT in LAAP Water

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The photolysis of TNT was studied in different LAAP water samples collected during the field trips in December 1981, May 1982, and July 1982; no additional TNT was added to these samples for our experiments. The finding that the character of the lagoon itself and the LAAP water were different on each visit (see Section IV) required that photolysis of TNT be studied in each water so as to obtain the most relevant rate constant data that could be used in the lagoon-modeling efforts. The results from the several studies are summarized in Table 20. The studies are listed in the chronological order in which they were performed.

The rationale for the sequence is as follows. The studies in March (Expts. 2-4) using the December LAAP water were conducted after the effect of depth on the photolysis rate constant was found to be important (see Section V.B.3). In the late summer, Experiments 6 and 7 were conducted simultaneously using the December and July waters after it was found that the TNT rate constant measured in the July LAAP water in September (Expt. 5) was nearly the same as that measured in March (Expt. 3), which was unexpected because photolysis rates are more rapid in summer than in spring (see Section V.B.5). When these experiments showed that the photolysis rates of TNT were different in the two waters at the same time of year. Experiments 8-10 were conducted in which the absorbances of December, May, and July waters were made the same by dilution; the intent of these experiments was to determine whether the absorbances of waters were somehow responsible for the different rate constants observed, or whether the ability of the individual waters to promote TNT photolysis were indeed different among the water samples after allowing for the different absorbances. Unfortunately, the error in the measured rate constants is too large to clearly distinguish between the two effects.

From the data presented in Table 20, several conclusions can be offered that are important in estimating the photolysis rate constants of TNT in LAAP water. On the basis of the results of Experiments

8-10, the ability of the LAAP water to promore photolysis of TNT appears to be the same per unit absorbance of the water, although because of the error in the rate constant data, we cannot entirely rule out the possibility that the difference in the rate constants in Experiments 6 and 7 is not due to a slight effect of the concentration of natural substances, as would be observed by their absorbances.

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SUNLICHT PHOTOLYSIS OF THT IN LAAP WATER

P	0.6 <sup>e</sup>	0.6 <sup>e</sup>	0.3 <sup>f</sup>	0.4 <sup>f</sup>		0.118	0.118	0.16 <sup>8.h</sup>	0.16 <sup>g</sup> .h	0.16 <sup>8.h</sup>	
10 <sup>3</sup> k a.d (aín <sup>-1</sup> )	16.3 ± 0.6 <sup>e</sup>	16.3 ± 0.6 <sup>e</sup>	11.3 ± 0.3 <sup>f</sup>	11.1 ± 0.4 <sup>f</sup>	1	3.27 ± 0.11 <sup>8</sup>	3.27 ± 0.11 <sup>8</sup>	3.13 ± 0.168.h	3.13 ± 0.16 <sup>8.h</sup>	3.13 ± 0.16 <sup>8.h</sup>	
$10^{3} k_{p} \times D^{c}$ (ce min <sup>-1</sup> )	1.56	1.56	8.78	7.01	6.81	3.27	2.13	2.59	2.45	2.17	
10 <sup>3</sup> k (min <sup>-</sup> f)	0.56 ± 0.13	0.56 ± 0.13	2.51 ± 0.04	1.67 ± 0.04	2.27 ± 0.05	1.09 ± 0.05	0.71 ± 0.08	0.74 ± 0.25 <sup>h</sup>	0.70 ± 0.49 <sup>h</sup>	0.62 ± 0.31 <sup>h</sup>	
Water Depth, D (cm)	2.8	2.8	3.5	4.2	3.0	3.0	3.0	3.5	3.5	3.5	
Temp (°C)	13-19	13-19	14-21	15-26	19-25	16-24	16-24	19-25	19-75	19-25	
Time/Duration of Expt.	13:00-17:00, 240 min.	13:00-17:00. 240 min.	9:40-15:30, 350 min.	9:00-16:00, 420 min.	10:00-16:00, 360 min.	10:00-17:00, 400 min.	10:00-17:00, 400 min.	12:15-17:00, 400 min.	12:15-17:00, 400 min.	12:15-17:00,	400 min.
(TNT) (ppm)	5.0	5.0	5.0	5.0	0.8	1.8	0.8	1.9	0.5	11.4	
Water	Dec. LAAP <sup>8</sup>	Dec. LAAP <sup>a</sup>	Dec. LAAP <sup>a</sup>	Dec. LAAP <sup>a</sup>	July LAAP	Dec. LAAP <sup>b</sup>	July LAAF	Dec. LAAP <sup>b</sup>	July LAAP adjusted	May LAAP	
Expt. No. (Date)	(1/26)	(3/12)	(3/13)	(21/0)	5. (9/18)	(10/28)	(10/28)	(1/11)	(1/11)	(1/11)	
Ex		2.	ъ.	4.	5.	6.	7.	<b>.</b>		10.	

Mater from Site 1.

<sup>b</sup>Water from Site 2.

<sup>C</sup>Depth-independent rate constant. <sup>d</sup>PNA/PTR actinometer photolysis rate constant. <sup>d</sup>PNA = 1.0 × 10<sup>-5</sup> M; PTR = 5.0 × 10<sup>-3</sup> M. <sup>f</sup>PNA = 1.0 × 10<sup>-5</sup> M; PTR = 1.25 × 10<sup>-3</sup> M. <sup>B</sup>PNA = 1.0 × 10<sup>-5</sup> M; PTR = 1.24 × 10<sup>-4</sup> M. <sup>h</sup>Rate conr.ant obtained with first three time points; later time point showed slower loss of TNT/actinometer in late afternoon (>15:00).

The data in Table 20 also show that the photolysis rate constant for TNT in the LAAP waters is independent of TNT concentration, as would be expected if the photolysis of TNT were a first-order process (Equation 26). However, we have insufficient data with which to evaluate the effect of changes in the properties of the lagoon water on the photolysis rate constant.

# 8. RDX Photolysis Rate Constant

Concurrent with the TNT studies, we performed calculations to estimate the RDX photolysis rate constant. The rate of RDX photolysis can be described by Equation 30 for wavelength  $\lambda$ :

$$\frac{dC}{dt} = \frac{\phi I_{o\lambda} \left(1 - 10^{-(\alpha_{\lambda} + \epsilon_{\lambda} C)\ell}\right) \epsilon_{\lambda} C}{D(\alpha_{\lambda} + \epsilon_{\lambda} C)}, \quad (30)$$

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where D = depth (cm)

 $I_0 \lambda$  = incident light intensity

 $\alpha_{\lambda}$  = absorpt vity of the water

 $\epsilon_{\lambda}$  = molar absorptivity coefficient for RDX at  $\lambda$ 

l = solution path length

In LAAP water,  $\alpha_{\lambda} >> e_{\lambda}C$ . Therefore, Equation 30 above reduces to Equation 31:

$$-\frac{dC}{dt} = \frac{\phi I_{o\lambda}(1 - 10^{-\alpha}\lambda^{R})\varepsilon_{\lambda}C}{D\alpha_{\lambda}} \qquad (31)$$

Since the exponential function  $10^{-\alpha}\lambda^{1}$  is also very small compared to 1 in waters where  $\alpha_{\lambda}$  is large, Equation 31 reduces to Equation 32:

$$-\frac{\mathrm{d}C}{\mathrm{d}t} = \frac{\Phi^{\mathrm{I}}_{\mathrm{o}\lambda}\varepsilon_{\lambda}^{\mathrm{c}}}{\mathrm{D}\alpha_{\lambda}} \qquad (32)$$

Integration of Equation 32 gives a first-order rate expression where  $k_p$  is equal to the expression shown below at wavelength  $\lambda$ :

$$k_{p\lambda} = \frac{\phi I_{o\lambda} \varepsilon_{\lambda}}{D \alpha_{\lambda}} \qquad (33)$$

The environmental rate constant can now be summed over all wavelengths in the solar spectral region and the photochemical rate constant can be described as shown in Equation 34:

$$k_{pE} = \frac{\phi \Sigma \varepsilon_{\lambda} I_{o\lambda}}{D \alpha_{\lambda}} \qquad (34)$$

Using the molar absorptivity values measured for RDX from 299 nm ( $\varepsilon = 164$ ) to 400 nm ( $\varepsilon = 0.02$ ), the absorptivity of the LAAP water ( $\alpha_{\lambda}$ ) over the same spectral range, and the quantum yield ( $\phi = 0.16$ ), we calculated the photochemical rate constant ( $k_{pE}$ ) at various depths of the LAAP water using a modified version (GC SOLAR) of the computer program developed by Zepp and Cline (1977) for calculating the intensity of ultra-violet radiation (297.5-380 nm) reaching the earth's surface as a function of time of day, longitude, latitude, and season. These data are shown in Table 21 for the four seasons. Also shown are the depth independent rate constants for RDX computed for LAAP waters collected in December and July. From the rate constants shown in parentheses, a comparison of the seasonal effects on photo-lysis can be made; however, these values are not environmentally revelant due to changes in the water's composition as a function of time of year.

PHOTOCHEMICAL RATE CONSTANTS FOR RDX IN LAAP WATERS AS A FUNCTION OF SEASON AND DEPTH

July LAAP Water Depth (cm) Fall Winter Spring Summer Surface<sup>a</sup> 390 200 710 930 3.8 5.0 15 2.0 1.0 1.9 30 1.0 0.52 2.5 0.26 1.3 60 0.51 0.95 90 0.34 0.17 0.63 0.83 Depth Independent Rate (30) (16) (56) 76 Constant (cm d<sup>-1</sup>) in July LAAP -------------Water<sup>b</sup> \_\_\_ ---------Depth Independent Rate 19 (39) (75) (101) Constant (cm  $d^{-1}$ ) in December ---\_\_\_ \_ \_ \_ LAAP water<sup>b</sup> - - -

Rate Constant ( $\times 10^3 d^{-1}$ ) in

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<sup>a</sup>Rate constant at surface not used

<sup>b</sup>Rate constants in ( ) are for comparison of seasonal effects on photolysis and are not environmentally relevant. The above calculations showed that the light intensity for a particular LAAP water changed by a factor of five going from winter to summer months. The depth-independent rate constant, however, changed by only a factor of three due to the greater absorptivity of the LAAP water in the summer ( $\alpha_{\lambda} = 9.0$  at 300 nm) than in the winter ( $\alpha_{\lambda} = 3.0$ at 300 nm).

To determine how well the calculated depth-independent rate constant compared to a measured value, we performed an KDX sunlight photolysis experiment in a 19 × 3-cm wide crystallizing dish (darkened on sides and bottom) over a seven-day period (starting 9/18/82) using lagoon water collected on 24 July 1982 and filtered through a 0.25-um filter. A plot of 1n RDX versus time gave a straight line, from which a first-order rate constant (slope) was calculated to be 4.77 × 10<sup>-5</sup> min<sup>-1</sup>. In the same experiment, TNT photolyzed with a pseudo-firstorder rate constant of 2.4 ± 0.3 × 10<sup>-3</sup> min<sup>-1</sup>, or approximately 50 times faster than RDX. The depth independent rate constant was calculated to be 0.20 cm day<sup>-1</sup>.

This value is about three times larger than the calculated value for July LAAP water. Using the GC SOLAR program, we calculated the RDX photolysis rate constant in pure water to be 2.0 cm  $4ay^{-1}$  during the summer, which is in good agreement with an experimental value of 2.2 cm day<sup>-1</sup> measured in the summer. Therefore, the differences in the measured and calculated photolysis rate constants for LAAP water could be due to sensitization reactions that are not understood at this time. For the modeling study, we used the measured rate constant in LAAP water corrected by the light intensity factors for the various seasons.

#### C. Biotransformation

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#### 1. Bacterial Count

The identification of aminodinitrotoluenes in the lagoon water suggested that biotransformation was occurring in the lagoon water and
sediment. To obtain an indication of aerobic and anerobic microbial populations in the lagoon and sediment, bacterial plate counts were performed on waters and sediment collected during the December field study. In the lagoon water (Table 22), the aerobic bacterial counts averaged 1.7  $\times$  10<sup>6</sup> CFU ml<sup>-1</sup> and the anaerobic bacteria averaged 1.7  $\times$ 10<sup>3</sup> CFU ml<sup>-1</sup>. In the sediment (Table 23), the averages were 1.1  $\times$  10<sup>7</sup> CFU ml<sup>-1</sup> for the aerobic bacteria and 1.5  $\times$  10<sup>5</sup> CFU ml<sup>-1</sup> for the anaerobic bacteria.

#### Table 22

Sample Site	Aerobic Bacteria (CFU ml <sup>-1</sup> )	Anaerobic Bacteria (CFU ml <sup>-1</sup> )
1	$1.6 \times 10^{6}$	$2.9 \times 10^{3}$
2	$1.8 \times 10^{6}$	$1.1 \times 10^{3}$
3	$1.7 \times 10^{6}$	$0.48 \times 10^{3}$
4	$1.7 \times 10^{6}$	$2.2 \times 10^{3}$

#### TOTAL AEROBIC AND ANEROBIC BACTERIA IN LAGOON 9 WATER COLLECTED IN DECEMBER

#### Table 23

# TOTAL AEROBIC AND ANAEBOBIC BACTERIA IN LAGOON 9 SEDIMENT COLLECTED IN DECEMBER

Sample Site	Aerobic Bacteria (CFU ml <sup>-1</sup> )	Anaerobic Bacteria (CFU ml <sup>-1</sup> )
1	$0.51 \times 10^{7}$	$1.4 \times 10^{5}$
2	$1.4 \times 10^{7}$	$1.9 \times 10^{5}$
3	$0.76 \times 10^{7}$	$0.71 \times 10^5$
4	$1.9 \times 10^{7}$	$2.0 \times 10^5$

The tea-colored water and codiment contained unexpectedly high numbers of aerobic bacteria. The numbers of anaerobic bacteria were substantially smaller, reflecting the high dissolved-oxygen content (7 to 8 ppm) of the water in the winter. Total aerobic bacteria in a water sample collected on 25 May 1982 averaged  $1.9 \times 10^6$  CFU ml<sup>-1</sup>.

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A lagoon water sample collected on 24 July 1982 was also evaluated for viable cells by bacterial plate counting. Because the lagoon water temperature had risen above 35°C, plates were incubated at 37°C and 25°C. The number of bacterial colonies formed at each temperature is shown in Table 24. These results indicate that more bacteria will grow at 25°C than at 37°C. However, the bacteria that do not grow at 37°C are still viable.

Table 24

# BACTERIAL PLATE COUNT OF LAAP LAGOON WATER COLLECTED IN JULY

Incubation	Average Colonies Formed
Temperature	(CFU m1 <sup>-1</sup> )
25°C	$2.60 \times 10^{6}$
37°C	1.19 × 10 <sup>6</sup>

These counts do not differ significantly from those found in December 1981 and May 1982, thus suggesting that the microbial population within the lagoon remains fairly constant although populations of specific organisms may change.

### 2. Biotransformation Screening Tests

Because of the presence of substantial microbial populations in the water, we initiated screening tests for the biotransformation of TNT and RDX in lagoon water under aerobic and anaerobic conditions.

Lagoon waters from the four sites (north, south, east and west quadrants) were mixed in equal parts and three preparations of 2-liter sumples were each placed in 4-liter bottles. The three preparations were 1) water alone, 2) water plus 50 ppm sterile yeast extract, and 3) water plus 1% dry weight of bottom sediment. For the anaerobic incubation, 1 liter of each of the above solutions was placed in a 1liter Erlenmeyer flask, which was flushed with  $N_2$  gas and sealed with a rubber stopper. Flushing with  $N_2$  gas was repeated after each sampling. The samples were analyzed periodically for total organic carbon (TOC), TNT, and RDX. The initial concentrations were 75 ppm TOC, 5.0 ppm TNT, and 13.0 ppm RDX. No additional TNT or RDX was added to the water.

#### a. Aerobic TNT Biotransformation

After 1 day of incubation, the bacterial population in the lagoon water under aerobic conditions increased slightly to  $3 \times 10^{\circ}$  CFU ml<sup>-1</sup>, then gradually decreased to  $1.2 \times 10^{6}$  CFU ml<sup>-1</sup> after 16 days of incubation, and remained constant at  $1.1 \times 1^{6}$  CFU ml<sup>-1</sup> from 34 days to 70 days. The TOC remained constant (75 ppm) over this period.

In the lagoon water alone, the TNT concentration slowly decreased from 5 ppm to 2 ppm over 90 days of incubation (Figure 10). The transformation followed first-order kinetics. The pseudo-first-order rate constant was calculated to be  $4.1 \times 10^{-4}$  hr<sup>-1</sup>, and the secondorder rate constant was calculated to be  $3.7 \times 10^{-10}$  ml cell<sup>-1</sup> hr<sup>-1</sup>, based on microbial counts of  $1.1 \times 10^{6}$  CFU ml<sup>-1</sup>.

The addition of yeast extract to the lagoon water accelerated the transformation rate by increasing the microbial population. The microbial count increased from 0.3 to  $1.6 \times 10^7$  CFU ml<sup>-1</sup> after one day of incubation. However, it decreased rapidly to  $1.6 \times 10^6$  CFU ml<sup>-1</sup> after 16 days, and the transformation rate also decreased. The rate accelerated again when yeast extract was added at Day 41 (Figure 10).

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In the lagoon water with added sediment, the total concentration of TNT increased to 74 ppm because the bottom sediment contained large amounts of solid TNT. Total TNT in this water decreased from 74 ppm to 59 ppm during the 90 days of incubation in a similar fashion to



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TNT in water alone; ▲ RDX in water alone; ○ TNT in water plus yeast extract;
 ▲ RDX in water plus yeast extract;
 ♦ 50 ppm yeast extract

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that observed in lagoon water alone. Bacterial counts averaged  $3 \times 10^6$  CFU ml<sup>-1</sup> in the water-sediment mixture.

# b. Anaerobic TNT Biotransformation

Under anaerobic conditions in the lagoon water alone, TNT was transformed at a slower rate than under aerobic conditions (Figure 11). This may be a result of a lower population of anaerobic microorganisms. The pseudo-first-order rate constant was calculated to be  $1.6 \times 10^{-4} \text{ hr}^{-1}$ .

The addition of yeast extract to the water also accelerated the disappearance of TNT. Yeast extract was added at Day 0, Day 41, and Day 76; there was a significant drop in TNT concentration after each addition.





In the sediment-added lagoon water, TNT decreased from 74 ppm to 57 ppm during 76 days of incubation. Therefore, it appears that anaerobic biotransformations in anaerobic waters are similar to the aerobic biotransformation of TNT.

# c. <u>Aerobic TNT Biotransformation at High and Low</u> <u>Concentrations of TNT</u>

The biotransformation of TNT was investigated at high and low concentrations of TNT and in the presence of a high microbial population. The lagoon water (stored at < 10 °C) was exposed to room temperature for one week, centrifuged, and resuspended in one-tenth of the volume of lagoon water in two bottles. Pure TNT was added to one bottle to bring the aqueous concentration to ~50 ppm without changing the other chemical components of the lagoon water. The other bottle contained the original TNT concentration of lagoon water (4.2 ppm). The average bacterial counts in these waters were originally 7.8  $\times 10^{6}$ CFU ml<sup>-1</sup>, but they decreased to 4.0  $\times 10^{6}$  CFU ml<sup>-1</sup> after 25 days of incubation.

The TNT in the water with a low concentration of TNT transformed with a pseudo-first-order rate constant of  $1.9 \times 10^{-3} \text{ hr}^{-1}$  (4.6 × 10<sup>-2</sup> day<sup>-1</sup>; correlation coefficient,  $r^2 = 0.98$ ). The second-order rate constant was calculated to be  $3.2 \times 10^{-10}$  ml cell<sup>-1</sup> hr<sup>-1</sup> (7.7 × 10<sup>-9</sup> ml cell<sup>-1</sup> day<sup>-1</sup>), based on a median bacterial count of  $6.0 \times 10^6$  CFU ml<sup>-1</sup>. This rate constant is close to that observed in the original water ( $3.7 \times 10^{-10}$  ml cell<sup>-1</sup> hr<sup>-1</sup>).

The TNT in the water with a high concentration of TNT decreased from 48 ppm to 33 ppm in 25 days. If the data are treated as pseudofirst-order, the rate constant, calculated by a least-squares method, is  $6.3 \times 10^{-4}$  hr<sup>-1</sup> ( $1.5 \times 10^{-2}$  day<sup>-1</sup>; r<sup>2</sup> = 0.95). This rate constant is only one-third of that observed in the water with a low TNT concentration. If the data are treated as following zero-order kinetics, which is expected for high chemical substrate biotransformation, the zero-order rate constant is  $2.5 \times 10^{-2}$  ppm hr<sup>-1</sup> ( $6.0 \times 10^{-1}$  ppm day<sup>-1</sup>;

 $r^2 = 0.96$ ). Because the correlation coefficients are nearly equal, more data points are required to prove the zero-order kinetic behavior. However, since the calculated pseudo-first-order rate constant is much lower than that observed in the low-TNT-concentration experiment, a zero-order kinetic process may better describe the transformation rate at high TNT concentrations (such as water in the sediment). The calculated rate constant for 6 × 10<sup>6</sup> cell ml<sup>-1</sup> is 4.1 × 10<sup>-9</sup> ppm cell<sup>-1</sup> hr<sup>-1</sup> (9.8 × 10<sup>-8</sup> ppm cell<sup>-1</sup> day<sup>-1</sup>).

#### d. TNT Metabolities

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During the laboratory biotransformation study of TNT in lagoon water, we observed that the concentrations of 2-amino-4,6-dinitrotoluene (2-A-4,6-DNT) and 4-amino-2,6-dinitrotoluene (4-A-2,6-DNT) remained nearly constant throughout the study. To investigate whether 2-A-4,6-DNT and 4-A-2,6-DNT are products of lagoon water biotransformation, we centrifuged lagoon water and resuspended the precipitate-containing microorganisms in 1/20th the volume of lagoon water. Ring-labeled  ${}^{14}$ C-TNT (1.0 µCi in 50 ml) and unlabeled TNT (final concentration 40 ppm) were added to the water and the water was incubated in the dark.

After 3 days and 19 days of incubation, the water was extracted with ethyl acetate, then acidified to pH 2, and re-extracted with ethyl acetate. In the 3-day sample, more than 90% of the radioactivity was found in the neutral fraction. The extracts were dried, concentrated, and chromatographed by TLC using silicia gel plates and chloroform as the developing solvent. The plates were analyzed by radioautography. Most of the activity was associated with TNT; however, radioactive 2-A-4,6-DNT and 4-A-2,6-DNT were produced and identified by their yellow color and by co-chromatography against authentic standards.

In the 19-day sample, the distribution of the radioactivity among the neutral, the acid extract, and the remaining aqueous phase was

80%, 16%, and 4%, respectively. The radioautograph-TLC plate developed in chloroform, in benzene:ethyl acetate:acetic acid (6:3:1), and in chloroform:methanol (6:1) solvent systems showed the major activity at spots with  $R_f$  values corresponding to TNT, 4-amino-2,6-dinitrotoluene, and 2-amino-4,6-dinitrotoluene (Figure 12). The activity at spots corresponding to 2,4-diamino-6-nitrotoluene and 2,6-diamino-4nitrotoluene were insignificant. Several minor spots with weak activity had  $R_f$  values similar to those for trinitrobenzyl alcohol and trinitrobenzaldehyde. However, further identification is needed. The acid extract showed several radioactive components with low  $R_f$  values.

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FIGURE 12 RADIOAUTOGRAPH OF THIN-LAYER CHROMATOGRAM OF TNT METABOLITES IN LAGOON WATER AFTER 19 DAYS OF INCUBATION

Samples: N = neutral extract; A = acidic extract; r = reference chemicals.

Compounds: 1 = TNT; 2 = 2,2',6,6'-tetranitro-4-azoxytoluene; 3 = 4-amino-2,6-dinitrotoluene; 4 = 2-amino-4,6-dinitrotoluene; 5 = 2,6dinitroaniline; 6 = 2,4-dinitroaniline; 7 = trinitrobenzyl alcohol; 8 = trinitrobenzaldehyde; 9 = 1,3,5-trinitrobenzene; 10 = picric acid. The extracts were streaked on preparative TLC plates. Several unknown compound spots were scraped from the plate and subjected to probe mass spectrometric analysis; however, they could not be identified because of large amounts of interfering phthalates and binders from the TLC plates.

On the basis of this study, we conclude that the formation of aminodinitrotoluenes in lagoon water is the result of microbial processes.

### e. Biological Oxygen Demand of the Lagoon Water

The biological oxygen demand of the LAAP lagoon water was measured as an indicator of the oxygen-utilization rate of the microbes. The refrigerated lagoon water was left at room temperature overnight, then shaken to saturate the water with air, and poured into four biological oxygen demand (BOD) test bottles, which were then incubated at 20°C in the dark. The dissolved oxygen (DO) concentration was monitored with a DO meter after 0, 3, 5, 7, and 10 days of incubation. The DO decreased at each sampling point, yielding a first-order rate constant of  $9.6 \times 10^{-2} \text{ day}^{-1}$ . We therefore expect that in the LAAP lagoon (DO = 8 ppm) the microbes will use oxygen at a rate of 0.77 ppm day<sup>-1</sup>.

#### f. RDX Aerobic Biotransformation

The original concentration of RDX in the lagoon water was 13 ppm. No loss of RDX was observed during the aerobic experiment-whether RDX was in the lagoon water alone, with added yeast extract, or with 1% of bottom sediment (Figure 10). Therefore, it appears that RDX is biologically persistent in the lagoon water under aerobic conditions.

# g. RDX Anaerobic Biotransformation

Under anaerobic conditions, no change in RDX concentration was observed in the lagoon water alone or in water plus sediment. RDX (13 ppm) in lagoon water with 50 ppm of added yeast extract did not show any loss of RDX after 46 days. An additional 50 ppm of yeast extract was added on Day 46 and 100 ppm on Day 76. After 83 days of incubation, the RDX concentration remained at 13 ppm and then dropped to 2.9 ppm at Day 90 and 1.4 ppm at Day 92. Enthaps the repeated addition of yeast extract acclimated RDX-utilizing organisms.

On Day 92, the microorganisms in the lagoon water were inoculated into basal salts medium (BSM) plus 500 ppm of yeast extract and 9.1 ppm of RDX. The RDX concentration decreased to 0.6 ppm after 5 days of anaerobic incubation. The organisms were transferred to a second flask containing BSM, 50 ppm yeast extract, and 9.4 ppm of RDX; the RDX was transformed to 3.4 ppm after 4 days of incubation.

The RDX-acclimated organisms were then inoculated into two flasks: one contained BSM plus 50 ppm yeast extract and 8.0 ppm RDX, the other contained lagoon water (12.9 ppm RDX) and 50 ppm of yeast extract. The results of the anaerobic incubation are shown in Table 25.

#### Table 25

RDX CONCENTRATION ON DIFFERENT DAYS AFTER ANAEROBIC INCUEATION IN LAAP LAGOON WATER

Day	RDX (ppm)
0	12.4
5	9 <b>.8</b>
10	5.8
17	2.5
20	<0.1

When the above organisms were inoculated into BSM with 8.0 ppm RDX and 50 ppm of yeast extract, the RDX was transformed to nondetectable levels (<0.01 ppm) within 5 days.

In a second experiment, the above organisms were inoculated into samples of lagoon water with 50 ppm or 300 ppm yeast extract. A third flask was inoculated with RDX-transforming organisms [obtained from the HMX waste-line sediment at Holston Army Ammunition Plant (HAAP)], lagoon water, and 50 ppm of yeast extract. The RDX concentration was monitored over a 20-day period. The results (Table 26) indicate that the anaerobic biotransformation of RDX can be accelerated by extra organic nutrients (300 ppm yeast extract) or by a different source of RDX-transforming organisms.

## Table 26

#### ANAEROBIC TRANSFORMATION OF RDX WITH SELECTED ORGANISMS AND YEAST EXTRACT CONCENTRATIONS

Source of	Yeast Extract	RDX C	oncent	ratio	n (pp	m) at Day
Organisms	<u>   (ppm)    </u>	0	7	13	17	
LAAP Lagoon Water	50	13.0	11.3	7.0	5.3	3.6
LAAP Lagoon Water	300	12.7	11.1	8.4	7.8	<0.1
HAAP Sediment	50	12.6	11.2	4.8	1.0	<0.1

For comparison, RDX-tranforming organisms from the \_AAP water and the HAAP sediment were each inoculated into flasks containing BSM with 9.0 ppm RDX and 50 ppm yeast extract; the RDX disappeared within three days. This result and the data in Table 26 show that the anaerobic biotransformation of RDX is much slower in lagoon water than in BSM even in the presence of large yeast extract concentrations (300 ppm) or with an alternate source of RDX-transforming organisms. These data suggest that lagoon water components have an inhibitory effect on RDX biotransformation.

To test the possibility that the slow RDX anaerobic biotransformation in lagoon water was due to the lack of mineral nutrients rather than to inhibition by some chemical present in the lagoon water, we investigated the biotransformation of RDX in lagoon water with yeast extract and BSM components. The results showed that the addition of mineral salts to lagoon water collected either in December or May did not significantly accelerate the RDX biotransformation, suggesting that the lack of salts is not responsible for the inhibitory effect of RDX anaerobic biotransformation in lagoon water.

#### 3. Detailed Biotransformation Rate Study

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The biotransformation of TNT depends on enzymes resulting from microorganisms utilizing organic nutrients other than TNT for growth. Under these conditions, the biotransformation rate will follow Michaelis-Menton kinetics at different concentrations of TNT. The transformation rate,  $\frac{dc}{dt}$ , can be represented as shown in Equation 35.

$$\frac{dC}{dt} = v = V_m C/(K_m + C) , \qquad (35)$$

where C is the TNT concentration,  $V_m$  is the maximum transformation rate at a given enzyme concentration or microbial population, and  $K_m$ is the Michaelis constant, which is the TNT concentration that yields half the maximum transformation rate.

When the TNT concentration is very low,  $K_m$  is much greater than C and the transformation rate expression reduces to Equation 36:

$$\frac{dC}{dt} = \frac{V_{m}C}{K_{m}} = k_{b}C , \qquad (36)$$

which is the pseudo-first-order transformation rate expression.

In environments where the TNT concentration is high, such as near the sediment in the LAAP lagoon, C is much greater than  $K_m$  and Equation 35 reduces to Equation 37:

$$\frac{dC}{dt} = V_{m} , \qquad (37)$$

which is a zero-order rate expression with respect to TNT. When  $V_m$  is divided by the bacterial concentration, the maximum rate per organism can be obtained.

Equation 35 can be rewritten as shown in Equation 38, which is the Lineweaver-Burke equation. A plot of 1/v versus 1/C yields a straight line, from which the values  $K_m/V_m$  (slope) and  $1/V_m$  (y-intercept) can be determined.

$$\frac{1}{v} = \frac{K_{\rm m}}{V_{\rm m}C} + \frac{1}{V_{\rm m}} .$$
 (38)

To determine  $K_m$  and  $V_m$ , we incubated LAAP lagoon water with 100 ppm of yeast extract for two days to generate a high cell population. The water was dispensed into five flasks and TNT was added to give concentrations ranging from 2.39 to 37.7 ppm. TNT was monitored for 24 hr and the transformation rates were determined. These data are shown in Table 27.

# Table 27

# TRANSFORMATION RATES DETERMINED FOR VARIOUS TNT CONCENTRATIONS

Initial TNT Concentration (ppm)	Transformation Rate, (ppm_day <sup>-1</sup> )
2.3	0.37
5.5	0.75
10.0	1.19
18.7	1.55

Using the Lineweaver-Burke method, we plotted the above data (Figure 13) and calculated  $K_m$  and  $V_m$ . Linear regression of the plot gave values of  $K_m = 19.3$  ppn and  $V_m = 3.5$  ppm day<sup>-1</sup>. The initial microbial population was  $1.0 \times 10^8$  CFU ml<sup>-1</sup> and, therefore, the maximum rate per organism will be  $3.5 \times 10^{-8}$  ppm cell<sup>-1</sup> day<sup>-1</sup> (1.5 ×  $10^{-9}$  ppm cell<sup>-1</sup> hr<sup>-1</sup>). The value is slightly lower than the 9.8 ×  $10^{-8}$  ppm cell<sup>-1</sup> day<sup>-1</sup> obtained with centrifuged lagoon water mico-organisms and 50 ppm TNT determined in a previous experiment. This may be the result of a modification of the mixed culture population when grown on yeast extract.

The above results indicate that the biotransformation of TNT will follow pseudo-first-order kinetics at TNT concentrations less than 19 ppm and zero-order kinetics at TNT concentrations greater than 19 ppm.

A second experiment was conducted with lagoon water microorganisms incubated without the addition of yeast extract. From the lagoon water collected in May, we prepared high cell populations by incubating the water at 20° to 25°C in the dark over a three-week period, centrifuging the microorganisms, and resuspending the cells in 1/3Sth of the volume of lagoon water. This water, contairing 2.1 ×  $10^7$  CFU ml<sup>-1</sup>, was dispensed into flasks and TNT was added to give concentrations ranging from 8 to 47 ppm.



FIGURE 13 LINEWEAVER-BURKE PLOT OF TNT BIOTRANSFORMATION RATE 'N LAGOON WATER

The biotransformation of TNT was followed over five days. The initial TNT concentrations and biotransformation rates are shown in Table 28.

# Table 28

# INITIAL TNT CONCENTRATIONS AND BIOTRANSFORMATION RATES OVER 5 DAYS

Initial TNT Concentration (ppm)	Biotransformation Rate (ppm day <sup>-1</sup> )
8.04	0.33
10.7	0.40
16.3	0.51
21.3	0.55
34.7	0.61
47.3	1.02

Using the Lineweaver-Burke plotting method, we calculated  $K_m$  (the Michaelis Constant) to be 19.1 ppm and  $V_m$  (the maximum transformation rate at a given enzyme concentration or microbial population) to be 1.1 ppm day<sup>-1</sup>. The maximum rate per organism is  $5.3 \times 10^{-8}$  ppm cell<sup>-1</sup> day<sup>-1</sup> (2.2 × 10<sup>-9</sup> ppm cell<sup>-1</sup> hr<sup>-1</sup>), which is close to the value obtained for the yeast extract-generated lagoon water organisms.

#### VI. DEVELOPMENT OF THE SIMULATION MODEL

#### A. General Equations

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A simulation model was developed based on a mathematical description of material balance. The basic equation for material balance is shown in Equation 39:

$$\frac{dM}{dt} = \sum_{i} I_{i} - \sum_{i} O_{j} , \qquad (39)$$

where M = total amount of chemical in the system

 $I_1 = total input of chemical to the lagoon$ 

 $0_{j}$  = total removal of chemical from the lagoon.

The change in total mass, dM/dt, can also be related to the chemical concentration, C, and volume, V, of the lagoon as shown in Equation 40:

$$\frac{dM}{dt} = C\frac{dV}{dt} + V\frac{dc}{dt} \qquad (40)$$

Thus, from Equations 39 and 40, Equation 41 can be formulated:

$$V\frac{dc}{dt} = \sum_{i} I_{i} - \sum_{j} O_{j} - C\frac{dV}{dt} \qquad (41)$$

Becaus, no munitions wastes have been dumped since early in 1981, the only input source of munitions arises from the diffusion of accumulated chemicals in the sediment layer  $(D_{dif})$  and the dissolution of solid chemicals lying on the bottom sediment surface  $(D_{dis})$ . Therefore, the chemical input can be described as shown in Equation 42:

$$\Sigma I = \Sigma (D_{dif_{i}} + D_{dis_{i}})$$

$$i i = 1$$

$$(42)$$

Because the bottom sediment and surface are heterogeneous with respect to the amounts of TNT and RDX present, the bottom sediment was divided into 25 compartments ( $30' \times 30'$ ) from which the diffusion flux and dissolution rate were calculated individually (i = 1 to 25).

Chemicals are removed from the lagoon through transformation processes (photolysis and biotransformation) and seepage. The seepage transport process has been accounted for in the calculation of net diffusion flux  $(D_{dif})$ ; therefore the total removal of chemical can be expressed as Equation 43:

$$\sum_{j=1}^{\Sigma} 0_{j} = \sum_{k=1}^{\Sigma} 2^{k} \chi^{2}, \qquad (43)$$

where  $k_{\ell}$  represents the transformation rate constants.

Equation 41 can now be written as Equation 44:

$$v_{dt}^{dC} = \sum_{i=1}^{25} (D_{dif} + D_{dis_i}) - \sum_{k \in V} CV - C_{dt}^{dV} \qquad (44)$$

A numerical integration scheme, using Euller's prediction and correction method, was used to integrate Equation 44 over selected time intervals to provide simulation results.

# B. Program Structure

A flow chart of the computer program is shown in Figure 14. System parameters, transformation rate constants, weather data, lagoon descriptions, and initial conditions are read in when the program is





initiated. The main program is linked to a number of subroutines such as VOLUME and SURFACE, which calculate the depth above each sediment compartment and the number of compartments covered by water. This number and the corresponding locations of sediment covered by water are then used separately in FLUX and FLUX 1 subroutines to calculate the diffusion flux and dissolution rate of chemicals from the bottom sediment, respectively. Material balance is performed in the subroutine COMPS when Euller's prediction and correction method is employed to predict the chemical concentration in the lagoon. The above procedures are repeated at every time step until the simulation period is complete or the lagoon goes dry. The status of the lagoon is obtained at specified print-out time intervals, and a plotting subroutine GRAPH 4 is used to plot the simulation results. The program interactions are shown in Figure 15, and a complete listing of the program appears in Appendix C.

# C. Simulation Results

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#### 1. Simulation of the Water Surface

The effect of seepage rate on the lagoon depth was discussed in Section V.A.1; an estimated rate of 0.12 cm day<sup>-1</sup> gave good agreement with measured values. Changes in the water surface were also simulated, as shown in Figure 16. As the lagoon dries, the southeast corner is the first to dry out and the lagoon recedes to the northwest corner. Observations of the water remaining in the lagoon in July and August were in good agreement with the model simulation.



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FIGURE 15 PROGRAM INTERACTIONS



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# FIGURE 16 APPROXIMATE WATER CONTOUR LINES DURING THE SIMULATION

# 2. TNT Simulation

#### a. Case 1: Complete Mixing

Five scenarios were investigated in which the assumption was made that the lagoon is a completely mixed reactor. Figure 17A shows the TNT concentration as a function of time using inputs from diffusion flux and dissolution, a photolysis rate constant of 0.78 cm day<sup>-1</sup>, and a biotransformation rate constant of 0.012 day<sup>-1</sup>. The results indicate that the TNT concentration will increase until Day 200, with the peak concentration being 17.5 ppm. After Day 200, TNT concentration decreases due to decreases in the lagoon depth, which increases the magnitude of the photolysis rate constant.

To test the impact of the photolysis rate constant, a simulation was run under conditions identical to those above except that the photolysis rate constant was reduced by half--to 0.39 cm day<sup>-1</sup>. Figure 17B shows the simulation result. In this case, the TNT concentration builds up faster than in the previous case and reaches a maximum of 23 ppm on Day 200. After Day 200, the TNT concentration decreases at a slower rate than before.

The photolysis rate constant is also seasonally dependent due to high light intensity in the summer months relative to that in the winter months. The magnitude of the intensity change is usually between a factor of 2 to 3. In Figure 17C, we adjusted the photochemical rate constant by a factor of 2, starting in June, and observed an accelerated decrease in TNT concentration in the late summer months. The overall effect appears to be a lowering of the maximum TNT concentration buildup followed by a rapid loss of TNT.

We performed a simulation in which the biotransformation rate constant was doubled for every 10°C rise in lagoon temperature, based on a number of assumptions involving the enzymatic reduction of a TNT nitro group. The result is plotted in Figure 18. In the early part of the simulation, the TNT concentration rises above that observed in Figure 17A due to the low biotransformation rate constant. However,





Curve A:  $k_p = 0.78$  cm day<sup>-1</sup>,  $k_b^1 = 0.012$  day<sup>-1</sup>; input from diffusion and dissolution. Curve B:  $k_p = 0.39$  cm day<sup>-1</sup>,  $k_b^1 = 0.012$  day<sup>-1</sup>; input from diffusion and dissolution. Curve C:  $k_p = 0.78$  cm day<sup>-1</sup> plus radiant correction;  $k_b^1 = 0.12$  day<sup>-1</sup>; input from diffusion and dissolution.



FIGURE 18 SIMULATED TNT CONCENTRATION AS A FUNCTION OF TIME OF YEAR

 $k_p = 0.78$  cm day<sup>-1</sup>;  $k_b^1 = 0.006$  day<sup>-1</sup> and doubled for every 10°C rise in temperature; input from diffusion and dissolution. the effect of biotransformation becomes evident between Days 150 and 200. In the latter phase of the simulation, photolysis becomes the dominant transformation process.

Finally, we considered a case in which dissolution was the only input source of TNT (Figure 19). The simulation results show TNT concentration decreasing in the early stage of simulation and then increasing in May, reaching a maximum at Day 200 and then decreasing as the lagoon dries out.

The concentration-time-dependent profile measured in the lagoon for TNT appears in Figure 20. None of the above scenarios exactly match the measured data. The low concentration observed in March (Day 74) does not appear in any simulation except in the case of the sole input arising from dissolution (Figure 19). The maximum TNT concentration was observed in May (Day 145), whereas the simulation results predicted that the maximum would occur in July. To reach a maximum in May, a large input from the bottom sediment would be expected. The present model probably overestimates the diffusion and dissolution processes in the latter months. Also, the lagoon begins to dry out at the end of June; this process should concentrate chemicals due to volume reduction and concentrate the TNT in July unless the transformation processes are greatly accelerated. Both the photolysis and biotransformation of TNT in lagoon water collected in July give no suggestion of dramatic rate increases compared to other experimental data.

The discrepancy between the observed data and the simulations may be explained by the sampling technique and the lagoon location from which the samples were taken. Because the plant personnel did not have access to the center of the lagoon to collect samples, all sampling done by them was performed at the southeast edge near the dump ramp, where high levels of TNT were found in the sediment. As the lagoon dried, samples were taken progressively closer to the center of the lagoon approaching the northwest corner. Poor mixing of the lagoon water may therefore obscure the simulation results.







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FIGURE 20 CONCENTRATION-TIME-DEPENDENT PROFILE OF TNT MEASURED IN LAGOON NO. 9

# b. Case II: Zero Mixing

If the sampling locations become critical, the assumption of complete mixing may not be a valid representation of the distribution of chemicals in the lagoon. Although the data on dissolved oxygen obtained in December indicated that the lagoon was well-mixed in the vertical direction, discrepancies in TNT concentrations at the edge of the lagoon relative to the center were noted. This suggests that the horizontal mixing of lagoon water may not be complete and may intensify as the lagoon becomes shallow.

We performed simulations for the case of zero-mixing for compartments 1 (dump site) and 13 (center). In the case of zero-mixing, the lagoon waters do not interact with neighboring compartments--only with the bottom sediment directly below the water. The simulation results are shown in Figure 21. In compartment 1, the TNT concentration continuously increases until maximum solubility is reached. In compartment 13, the TNT concentration continually decreases due to low TNT levels in the sediment and transformation rates that surpass those of diffusion and dissolution fluxes. In the real environment, partial mixing probably occurs that dissipates the TNT from compartments 1 to other compartments in the lagoon. These results do imply, however, that sampling from the southeast corner and following the water-coverage profile (Figure 16) for further samples may give skewed versions of the TNT concentration profile that are difficult to model. In this respect, the model simulation may be adequately predicting the loss and movement of TNT in the dissolution-input-only case (Figure 19), when compared to the actual data, that may be higher than in a completely mixed reactor (Figure 20). An overlay of the latter profiles appears in Figure 22.







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FIGURE 22 COMPARISON OF THE SIMULATION RESULTS FROM THE CASE WITH DISSOLUTION AS THE ONLY INPUT SOURCE OF TNT TO MEASURED DATA FROM THE LAGOON

# 3. RDX Simulation

The laboratory studies showed that there would be little biotransformation of RDX in the lagoon and that photolysis, although slow, probably would be the dominant transformation process. Knowing the range of the depth-independent photochemical rate constant for the winter through summer seasons, we derived a linear expression relating the rate constant,  $k_p$ , to the day of the year, D:

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(45)

$$K_p = (3.0 \times 10^{-4} \text{ cm day}^2) (D) + 1.6 \times 10^{-2} \text{ cm day}^{-1}$$
.

Thus, at Day 0 (January 1) the winter rate constant, 0.016 cm day<sup>-1</sup>, is applied to the model and linearly increases to Day 200 (July 19), where the summer rate constant is applied. At each simulation interval, the rate constant is adjusted according to the depth of the lagoon for that time period.

Three scenarios were simulated representing the observed rate constants and rate constants representing 75% and 50% of the observed rate constants. The latter rate constants provide a sensitivity analysis of the impact of the photochemical rate constant on RDX loss in the lagoon. The simulation results are plotted in Figure 23. In each case, the simulated results indicate that the removal of RDX is balanced by the diffusion and dissolution input over the first 150 days. As the water gets warmer, the concentration increases up to Day 225; then the effect of photolysis is noted only when the lagoon is very shallow. Also, halving the photolysis rate constant (Curve C) does not have a large effect on the RDX-loss profile.

Superimposed on these curves are the actual measured values for RDX in the lagoon. Considering the problem of partial mixing, the simulation results and the actual values are comparable except for one data point where a very high level (42.1 ppm) of RDX was observed.



FIGURE 23 SIMULATION RESULTS FOR RDX CONCENTRATION IN LAAP LAGOON WATER

Curve A:  $k'_p = k_p$ ; Curve B:  $k'_p = 0.75 k_p$ ; Curve C =  $k'_p = 0.50 k_p$ . • = Found Value The remaining data are, however, inconsistent with this point. A curve for the profile of RDX can be approximated (dotted line) that shows a peak in RDX concentration in late July and a decrease in late August. These results suggest that the diffusion/dissolution processes become faster than predicted by the model in May and June (Days 145 to 175) and that the loss rate is faster than predicted in late August (Day 245). This is probably a result of (a) warmer waters near the edge of the lagoon where the samples were taken, which could give high concentrations in May-June, and (b) the low water level and the recession of the water's edge from the munitions-enriched sediment area, which could give low concentrations in late August. If complete mixing were to occur, the simulation results might be fairly accurate.

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#### VII. DISCUSSION

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The results of this study show that the lagoons at LAAP are highly complex environments in which the fate of TNT is controlled by photolysis and biotransformation and the fate of RDX probably is controlled by photolysis. If there were no chemical inputs to the lagoon, we would estimate a TNT half-life ranging from 8 days in the winter months to <4 days in the summer and an RDX half-life ranging from 173 days in the summer to >1000 days in the winter in a lagoon 50 cm deep. However, there are chemical inputs to the lagoon from solid and sorbed munitions in the bottom sediment and on the bottom surface. Thus, both diffusion and dissolution kinetics play important roles in the input of chemicals to the lagoon. These processes are complicated by 1) a heterogeneous distribution of munitions in and on the sediment and 2) a temperature-dependence that increases chemical inputs as the lagoon warms. Thus, chemical input is not a constant process, but one that can change dramatically, depending on the time of year and the portion of the lagoon that is covered by water.

Furthermore, each transformation process has additional complications. In the case of biotransformation, the enzymatic reduction of the nitro group should be temperature-dependent, and therefore the biotransformation rate constant could vary by a factor of 8 or more (assuming a doubling of the rate constant for every 10° rise in temperature) between the winter and summer months. Also, we cannot be sure that the population of organisms at 10°C is the same as that at 40°C. The temperature effect could have an impact on the number and type of microorganisms that are available to transform TNT. To average this effect, we used a rate constant determined at 25°C as an average value in one example and doubled the rate constant for every 10°C rise in temperature in another example. The results of these

examples indicated that changes in the biotransformation rate constant did not have a major impact on the loss profile of TNT because the photochemical transformation process overwhelmed any such changes.

There is also a high degree of uncertainty in the photochemical rate constant as a function of the time of year. Photochemical rate constants measured in lagoon water collected in December showed an inverse relationship with depth, which allows the prediction of a rate constant at any depth, provided that the absorptivity of the water is the same. However, as the lagoon waters evaporate, much of the UVabsorbing materials become concentrated, reducing the ability of light to penetrate into the water. The effect of the more concentrated waters will be to lower the photochemical rate constant, but simultaneously the shallower depth will tend to increase it. Because the photochemical transformation of TNT occurs by indirect mechanisms and is governed by unknown sensitizers (both humic materials and TNT transformation products) in the waters, it is difficult to correlate the photochemical rate constant dependences on the absorptivity and depth of the lagoon water from the experiments that were performed during this study.

Sampling was a problem because of poor access to a center location for the bulk water and because the sites from which the samples were collected changed as the lagoon dried. Finally, the lagoon dried up in late August.

In spite of these difficulties, we were able to develop a simulation model that describes the loss profile of TNT and RDX as a function of time of year. Of primary importance in this model is the dissolution rate of pure chemicals from the bottom sediment. This process is temperature-dependent and therefore we expect the concentrations of TNT and RDX to rise as the lagoon warms in the spring and summer months. In the absence of transformation, dissolution will continue until the solubility limit for each chemical is reached.

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A concentration buildup in the lagoon is counteracted by the photochemical and biological transformation of TNT and the photochemical transformation of RDX. The rate of photochemical transformation of TNT and RDX follows pseudo-first-order kinetics and first-order kinetics, respectively. Thus, for a given depth, TNT will photolyze approximately 50 times faster than RDX. The rate constant is dependent on the solar radiance; therefore, rates will be two to three times faster in the summer than in the winter months for TNT and up to five times faster for RDX.

The photochemical rate constant is inversely proportional to depth; hence, rates of transformation will increase as water is lost to evaporation and seepage. Usually, the lagoon is shallower in summer than in winter; therefore, rates of photochemical transformation are predicted to be greater in the summer. The biotransformation rate constant for TNT is an order of magnitude smaller than the photochemical rate constant; so a comparison of the dissolution flux and photochemical rate constant will indicate whether the TNT or RDX concentration will increase or decrease over any particular time period. Furthermore, if the input source is eliminated (as was observed when portions of the lagoon dried out), the TNT and RDX will transform at rates defined by the depth and mitigated by the absorptivity of the water. Thus, an overall profile of munitions in the lagoon will show a buildup of munitions when the lagoon is full, periods of equilibrium when the dissolution and transformation fluxes are equal, and periods when transformation flux exceeds that of dissolution--usually in the summer months.

The above theme has a number of variations that can be postulated and for which modeling can serve as a tool to predict chemical behavior. For example, if TNT and RDX were uniformly distributed in the bottom sediment, as was observed at the dump ramp, they would reach their solubility limits in the lagoon water. Similarly, if the sediment had a uniform distribution, as was found at the center of the lagoon, the transformation of TNT and RDX could be predicted from the transformation rate constants developed in this study.

In conclusion, the persistence of TNT and RDX in LAAP lagoons is controlled by dissolution and photochemical processes, with dissolution being complicated by a heterogeneous distribution of solid munitions on the bottom surface and photolysis being complicated by changes in absorptivity of the water that affects the photochemical rate constant as a function of time of year. This effect could not be investigated in detail because of project time and money contraints. A simulation model was developed to describe a first approximation of the loss and movement of TNT and RDX in the lagoon water, but vigorous validation could not be achieved due to sampling and mixing constraints within the lagoon.

#### Appendix A

ANALYSIS OF LAAP LAGOON NO. 9 SEDIMENT

#### Sediment Preparation

Twenty-four sediment samples scraped from the bottom surface of Lagoon No. 9 were sieved through a large mesh screen to remove twigs, leaves, and large pebbles. Each sample was homogenized with a mortar and pestle until an even consistency was obtained.

## Moisture Determination

A known weight (~20 g) of sediment was placed in a tared beaker and allowed to dry at 75°C for 48 hr. The percent moisture in the sample was calculated as the dry weight minus the wet weight, divided by the wet weight times 100.

#### Extraction of Sediment

A sample of wet sediment (10-20 g) was placed in a 50-ml screw-cap centrifuge tube and 20 ml of neutral ethyl acetate was added. The tube was vigorously shaken for 2-3 min and then centrifuged at 2000 rpm for 10 min. The ethyl acetate was removed by pipet. This process was repeated twice. The sediment was then acidified by adding 5 ml of 2N HCl and re-extracted with ethyl acetate (3 × 20 ml). The combined extracts were dried over anhydrous sodium sulfate, filtered, and rotary-evaporated to dryness. The residue was dissolved in methanol, and an internal standard was added for HPLC analysis.

## HPLC Analysis

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The HPLC analyses of the sediment extracts were performed under the following conditions:

Instrument:	Spectra-Physics Model 3500B Liquid
	Chromatograph
Column:	250 mm × 4.6 mm Altex RP600-C <sub>18</sub> @ 50°C
Solvent:	A - water, B - 50% methanol: 50%
	acetonitrile
Program:	30% B in A to 50% B in A in 15 min; linear
	gradient
Flow rate:	2.0 ml min <sup>-1</sup>
Detection:	UV @ 254 mm
Retention times:	5.68 min, HMX
	6.65 min, RDX
	10.55 min, 3,5-dinitroaniline
	12.40 min, TNT
	13.77 min, aminodinitrotoluenes
	15.65 min, 3,5-dinitrotoluene (internal
	standard)

A typical chromatographic profile of the sediment extract appears in Figure A-1.

## Results

The results of duplicate analyses of the 24 sediment extracts are presented in Table A-1 (also see Figure 2 in main text). Table A-2 shows the percent water in the sediment samples.



FIGURE A-1 HPLC PROFILE OF LAGOON NO. 9 SEDIMENT EXTRACT

# Table A-1

## PERCENT COMPOSITION OF MUNITION COMPONENTS IN LAAP LAGOON NO. 9 SEDIMENT

Sample	TNT	RDX	HMX	ADNTS	3,5-DNA
A-5	27.5	8.7	2.1	$6 \times 10^{-3}$	
A-4	29.7	6.6	$8.9 \times 10^{-1}$	$5.9 \times 10^{-2}$	
A-3	1.5	$8.9 \times 10^{-1}$	3. 10 <sup>-1</sup>	9.7 × $10^{-2}$	
A-2	$4.2 \times 10^{-2}$	$1.0 \times 10^{-3}$	$8.7 \times 10^{-2}$	$8.3 \times 10^{-2}$	$5.0 \times 10^{-3}$
A-1	$1.8 \times 10^{-3}$	$4.7 \times 10^{-3}$	$1.5 \times 10^{-2}$	$9.2 \times 10^{-3}$	$6.0 \times 10^{-3}$
B5	1.4	4.7	$8.4 \times 10^{-1}$	$8.0 \times 10^{-2}$	$8.0 \times 10^{-3}$
B-4	$1.4 \times 10^{-1}$	1.7	$3.5 \times 10^{-1}$	$3.0 \times 10^{-2}$	
B-3	$4.0 \times 10^{-2}$	$2.9 \times 10^{-1}$	$1.1 \times 10^{-1}$	$3.0 \times 10^{-2}$	$3.3 \times 10^{-2}$
B-2	$8.3 \times 10^{-4}$	$7.9 \times 10^{-3}$	$3.4 \times 10^{-3}$	$1.3 \times 10^{-2}$	9.0 × $10^{-4}$
B-1	$1.4 \times 10^{-2}$	$3.2 \times 10^{-4}$	$1.5 \times 10^{-2}$	$2.1 \times 10^{-2}$	$1.5 \times 10^{-3}$
C-5	$1.7 \times 10^{-2}$	1.0	$2.4 \times 10^{-1}$	$4.2 \times 10^{-2}$	$4.0 \times 10^{-3}$
C-4	$2.1 \times 10^{-3}$	$3.6 \times 10^{-3}$	$3.0 \times 10^{-2}$	$5.9 \times 10^{-3}$	$4.0 \times 10^{-4}$
C-3	$4.4 \times 10^{-3}$	$2.2 \times 10^{-4}$	$1.8 \times 10^{-2}$	$3.1 \times 10^{-2}$	$3.1 \times 10^{-3}$
C-2	$1.5 \times 10^{-3}$	$2.0 \times 10^{-4}$	$1.2 \times 10^{-2}$	9.1 × $10^{-3}$	$2.0 \times 10^{-4}$
C-1	5.7 × $10^{-4}$	$7.0 \times 10^{-5}$	$1.5 \times 10^{-2}$	$2.1 \times 10^{-3}$	$6.0 \times 10^{-4}$
D <b>-</b> 5	$4.9 \times 10^{-3}$	$1.7 \times 10^{-2}$	$6.6 \times 10^{-2}$	$3.3 \times 10^{-2}$	$3.4 \times 10^{-3}$
D-4	$1.2 \times 10^{-2}$	$1.6 \times 10^{-1}$	9.0 × 10 <sup>-2</sup>	$8.2 \times 10^{-3}$	$1.0 \times 10^{-3}$
D-3	$7.3 \times 10^{-4}$	$3.8 \times 10^{-2}$	$8.0 \times 10^{-3}$	$1.3 \times 10^{-2}$	$1.8 \times 10^{-3}$
D-2	$1.1 \times 10^{-3}$	$1.0 \times 10^{-4}$	$1.3 \times 10^{-2}$	$1.0 \times 10^{-2}$	$8.5 \times 10^{-4}$
D-1	$1.2 \times 10^{-3}$	$9.0 \times 10^{-5}$	9.1 × $10^{-3}$	$1.4 \times 10^{-2}$	$1.2 \times 10^{-3}$
E-4	$1.9 \times 10^{-3}$	$1.3 \times 10^{-3}$	$5.0 \times 10^{-3}$	$8.6 \times 10^{-3}$	$1.2 \times 10^{-3}$
E-3	$1.1 \times 10^{-3}$	$4.9 \times 10^{-4}$	$1.4 \times 10^{-2}$	$1.6 \times 10^{-2}$	$2.6 \times 10^{-3}$
i <b>:-2</b>	$1.1 \times 10^{-3}$	$7.5 \times 10^{-4}$	$1.6 \times 10^{-2}$	$1.5 \times 10^{-2}$	9.4 × $10^{-4}$
E	$2.2 \times 10^{-4}$	$6.0 \times 10^{-5}$	$4.0 \times 10^{-4}$	$3.1 \times 10^{-4}$	$6.0 \times 10^{-5}$

<sup>a</sup>ADNTS = aminodinitrotoluenes.

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PERCENT WATER IN LAGOON NO. 9 SEDIMENT SAMPLES

Sample No.	Percent	Water
A-5	51.6	
A-4	33.6	
A-3	34.1	
A-2	43.3	
A-1	48.5	
B-5	41.6	
B-4	32.4	
B-3	31.0	
B-2	33.9	
B-1	42.1	
C-5	33.3	
C-4	38.6	
C-3	30.2	
C-2	32.9	
C-1	58.2	
D-5	35.3	
D-4	48.1	
D-3	38.0	
D-2	39.2	
D-1	43.3	
E-4	32.2	
E-3	29.5	
E-2	30.1	
E-1	45.7	

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#### Appendix B

MASS TRANSFER PROBLEMS--ASSUMPTIONS AND SOLUTIONS

#### Description of Mass Transfer Problems

A body of water containing some concentration of one or more contaminants is in contact with a porous sediment (Figure B-1). Water residing in the pores of the sediment (interstitial water) also contains some concentration of one or more contaminants. The sediment solids may contain the same contaminants, presumed to be at equilibrium with the interstitial water concentration of contaminants. Contaminants will diffuse through the water both in the free-water phase and the sediment/water phase. In addition, chemical reactions (photolysis, biodegradation, hydrolysis) may occur, and contaminants may be transported by bulk convection (seepage of water into the sediment).

Far from the water/sediment interface, one expects the concentrations of contaminants to reach a constant (known) value. Also, on the free-water side of the interface, a region should exist where the macroscale mixing processes of the pond are negligible. With these ideas in mind, one can pose a set of simultaneous differential equations for the concentration profile of a contaminant in the water and sediment/water phases.

The change in concentration of chemical with time  $\begin{pmatrix} \partial C \\ sw \\ \partial t \end{pmatrix}$  is equal to the sum of the diffusion and seepage rates  $\begin{pmatrix} D \\ D \\ 1 \\ \partial z^2 \end{pmatrix}$  and  $v \frac{\partial C }{\partial z}$ , respectively minus the biotransformation rate  $(k_0)$  and the amount of chemical that is sorbed to the sediment:



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FIGURE B-1 LAGOON WATER RESTS ON SEDIMENT

Pond and sediment are considered to be semi-infinite.

$$\frac{\partial C_{sw}}{\partial t} = D_1 \frac{\partial^2 C_{sw}}{\partial z^2} + v \frac{\partial C_{sw}}{\partial z} - k_0 - K_0 C_{sw}^{n-1} \frac{\partial C_{sw}}{\partial t} .$$
(1)

Rearrangement of Equation 1 yields

$$1 + KnC_{sw}^{n-1} \frac{\partial C_{sw}}{\partial t} = D_1 \frac{\partial^2 C_{sw}}{\partial z^2} + v \frac{\partial C_{sw}}{\partial z} - k_0$$

where C<sub>sw</sub> = chemical concentration in the se i w
K = sediment partition coefficient (C<sub>g</sub> = KC<sup>n</sup><sub>Sw</sub>)
t = time
D<sub>1</sub> = diffusion coefficient in the sediment/water phase
v = seepage velocity
z = distance from the sediment

Similarly, for the free-water phase, the change in concentration of a chemical with time can be described by

$$\frac{\partial C_{w}}{\partial t} = D_{2} \frac{\partial^{2} C_{w}}{\partial z^{2}} + v \frac{\partial C_{w}}{\partial z} - \Sigma k_{1} C_{w} , \qquad (3)$$

where C<sub>w</sub> = concentration of chemical in the free-water phase
t = time
D<sub>2</sub> = diffusion coefficient in the free-water phase
v = seepage velocity
Ek<sub>1</sub>C<sub>w</sub> = first-order rate expression for transformation process.

The boundary conditions for these equations are that

$$\phi D_1 \frac{\partial C}{\partial z} = D_2 \frac{\partial C}{\partial z} \quad \text{at } z=0 \quad , \qquad (4)$$

where  $\phi$  is the sediment porosity (void volume fraction) and

$$C_{sw}(0) = C_{w}(0)$$
, (5)

$$C_{sw} (-\infty) = C_{sw,\infty} = known , \qquad (6)$$

$$C_{w}(\infty) = C_{w,\infty} = known , \qquad (7)$$

$$C_{gw}(z) = known at time zero = C_{gw,0}$$
, (8)

$$C_w(z) = known at time zero = C_w.o$$
 (9)

Because of Equations 5 and 6, Equations 2 and 3 must be solved simultaneously and are not amenable to a closed-form analytical solution. The equation could be solved on a digital computer, but the additional complexity would greatly increase the expense of running the overall SRI pond simulation model. Therefore, we have made several simplifying assumptions, described below.

#### Comparison of Diffusion, Convection, Reaction Rates

Because there is a strong incentive to reach a closed-form, analytical solution, the first assumption is that the concentration of chemical is constant everywhere in the free-water phase. This assumption allows Equation 3 to be eliminated.

Further simplifications could be made if either the convective term  $\left(v\frac{\partial C}{\partial z}\right)$  or the reaction term  $(k_0)$  could be neglected. By nondimensionalizing the equations, one can evaluate the relative importance of diffusion, reaction, and convection. Unfortunately, even the single differential equation cannot be solved analytically if there is no boundary on the sediment phase. Therefore, to compare the importance of diffusion, reaction, and convection, we pose a problem where the sediment phase has a boundary layer of finite thickness, l, near the water/sediment interface (Figure B-2). Furthermore, we assume steady-state conditions. This assumption leaves a simple differential equation describing the concentration in the sediment-water phase. In dimensionless form, this equation is

$$\frac{d^{2}C^{*}}{\frac{sw}{dz}} + \xi_{0} \frac{dC^{*}}{\frac{sw}{dz}} - \gamma_{0} = 0 , \qquad (10)$$

where  $\xi_0$  = ratio of convective transport rate to diffusive transport rate

 $\gamma_{0}$  = ratio of chemical reaction rate to diffusive transport rate

with boundary conditions

$$C_{sw}^{*}(0) = \frac{C_{w}}{C_{sw,\infty}} C_{w}^{*} = known \qquad (11)$$

and

$$C_{SW}^{*}(-1) = 1$$
 . (12)

The dimensionless parameters are

$$C_{sw}^{*} = \frac{C_{sw}}{C_{sw,\infty}}$$
(13)



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## FIGURE B-2 BOUNDARY LAYER APPROXIMATION IN SEDIMENT/WATER PHASE

Beyond a depth of  $\ell$ , the sediment is undisturbed by the mass transfer. Hence, flux can be computed as if sediment/water phase extended only to  $z = -\ell$ .

$$z^{\star} = \frac{z}{k} , \qquad (14)$$

$$\xi_{o} = \frac{lv}{D_{1}} , \qquad (15)$$

and

$$\gamma_{o} = \frac{k_{o} k^{2}}{C_{sw, \infty} D_{1}} . \qquad (16)$$

The parameter  $\xi_0$  is a measure of the ratio of convective flux to diffusive flux, and  $\gamma_0$  is a measure of the ratio of the rate of reaction to the rate of diffusion. The flux, N, in dimensional form is

$$N = -\phi D_1 \left(\frac{\partial C}{\partial z}\right)_{z=0} - \phi v C_{sw}(0) \qquad (17)$$

Defining a dimensionless flux,

$$N_1^{\star} = \frac{N!}{\phi D_1 C_{sw,\infty}} , \qquad (18)$$

we find, by solving Equation 10 for  $C_{sw}^{*}$  and substituting into Equation 17, that

$$N_{1}^{*} = \frac{\xi_{0}C_{w}^{*} + \frac{\gamma_{0}}{\xi_{0}}e^{\circ} - \xi_{0} - \frac{\gamma_{0}}{\xi_{0}} - \gamma_{0}}{1 - e^{\circ}} \qquad (19)$$

To evaluate the relative importance of the various phenomena, Equation 8 was solved with  $\gamma_0 = 0$  (no chemical reaction,  $N_2^*$ ), with  $\xi_0 = 0$  (no convection,  $N_3^*$ ), and with  $\gamma_0 = \xi_0 = 0$  (no chemical reaction and no convection). The dimensionless fluxes for each of these cases,  $N_2^*$ ,  $N_3^*$ ,  $N_4^*$ , are given as follows:

$$N_{2}^{*} = \frac{\xi_{0}(1 - C_{w}^{*} e^{\xi_{0}})}{e^{\xi_{0}} - 1}$$
(20)

$$N_3^{\star} = (1 - C_w^{\star}) - \gamma_0/2$$
 (21)

$$N_4^* = (1 - C_w^*)$$
 (22)

To compare  $N_1^*$  to  $N_2^*$ ,  $N_3^*$ , and  $N_4^*$ , we need to pose some reasonable values of the parameters  $C_w^*$ ,  $\xi_o$ , and  $\gamma_o$ . SRI has taken data from field samples of the waste disposal pond and found the following numbers:

$$= 2.2 \times 10^{-6} \text{ cm s}^{-1} \tag{23a}$$

$$k_{0} = 1.2 \times 10^{-5} \ \mu g \ cm^{-3} \ s^{-1} \ (for \ TNT)$$
(23b)  

$$D_{1} = 3.05 \times 10^{-6} \ cm^{2} \ s^{-1} \ (for \ TNT)$$
(23c)  

$$C_{gW_{1}} = 50 \ \mu g \ cm^{-3} \ (for \ TNT)$$
(23d)

$$C_{\pm}^{*} = 0.1$$
 (for TNT) (23e)

Because the boundary thickness, 1, exists only in our model (not in the real pond), reasonable estimates for 2 must be made. Therefore, in Table B-1, the results are presented for various boundary thicknesses. The table shows the dimensionless flux  $(N_1^{\pi})$  and flux values neglecting chemical reaction  $(N_2^*)$ , convection  $(N_3^*)$ , and reaction and convection  $(N_{L}^{*})$ , along with the ratios  $N_{2}^{*}/N_{1}^{*}$ ,  $N_{3}^{*}/N_{1}^{*}$ , and  $N_{L}^{*}/N_{1}^{*}$ . These ratios show how much difference it makes if the convection, the reaction, or both are neglected. It can be seen that  $N_2^{\frac{1}{2}}$  is not much different from  $N_1^*$  and that  $N_3^*$  is not much different from  $N_4^*$ . Hence, the chemical reaction term can be neglected. In contrast,  $N_3^{"}$ can be much larger than  $N_1^*$  and  $N_2^*$  can be much larger than  $N_2^*$ . These results imply that the convective term cannot be neglected. Hence, the reaction can be neglected but convection must be considered. This conclusion was expected from the magnitude of the dimensionless numbers [ $\xi_0 \sim 1$  and  $\gamma_0 < 1$  (Table B-1)].

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CALCULATED FLUX VALUES CONSIDERING ALL FARAMETERS  $(n_1^*)$ , no chemical reaction  $(n_2^*)$ , no convection  $(n_3^*)$ , no reaction or convection  $(n_4^*)$ , and their ratios to  $(n_1^*)$ 

Boundary Layer Thickness, Å (cm)	° °	~	R*	N2 N2	** 3	* \\$ S	N <sup>*</sup> /N <sup>*</sup>	$n_2^*/N_1^* = N_3^*/N_1^* = N_4^*/N_1^*$	N4/N1
0.5	0.355	0.019	0.705	0.714	0*890	0.900	1.013 1.263	1.263	1.276
0.	0.710	0.076	0.514	0.547	0.463	0.901	1.065	1.679	1.753
1.5	1.066	0.171	0.327	0.397	0.397 0.815	006*0	1.215	2.491	2.752
0	1.42	0.304	0.148	0.265	0.748	0.748 0.900	1.793	5.056	6.081

#### Unsteady Solution with Convection, Without Reaction

By neglecting the chemical reaction term, we can develop a closed-form solution for flux without appeal to some arbitrary boundary layer thickness,  $\pounds$ . In this regard, the solution developed below is the most realistic situation for which a closed-form solution can be developed. The solution will apply to TNT and RDX because the RDX was found not to react in the sediment/water phase (i.e.,  $k_o = 0$  for RDX).

Physically, we envision a situation like that in Figure B-1. We retain the assumption that the concentration in the water phase is constant and known. For this situation, the following equations apply in the sediment/water phase:

$$1 + Kn C_{sw}^{n-1} \frac{\partial C_{sw}}{\partial t} = D_1 \frac{\partial^2 C_{sw}}{\partial z^2} + v \frac{\partial C_{sw}}{\partial z} , \qquad (24)$$

with the following boundary conditions:

$$C_{sw}(-\infty,t) = C_{sw,\infty} = known$$
, (25a)

$$C_{gW}(0,t) = C_{W} = known$$
, (25b)

$$C_{sw}(z,0) = C_{sw,\infty} = known \qquad (25c)$$

With n = 1 (linear isotherm assumption) and in dimensionless form, these equations become

$$\frac{\partial C_{gw}^{*}}{\partial t} = \frac{\partial^{2} C_{gw}^{*}}{\partial z} + \beta_{0} \frac{\partial C_{gw}^{*}}{\partial z} , \qquad (26)$$

where  $\beta_0$  = dimensionless ratio of convective transport rate to diffusive transport rate,

$$C_{sw}^{*}(-\infty,t)=0$$
 (27a)

$$C_{sw}^{*}(0,t^{*}) = C_{w}^{*} - 1$$
 (27b)

$$C_{sw}^{*}(z^{*},0) = 0$$
, (27c)

where the dimensionless variables have been defined as follows:

$$C_{sw}^{\star} = \frac{C_{sw}^{-C} - C_{sw,w}}{C_{sw,w}}$$
(28a)

 $t^{*} = \frac{D_{1}t}{z_{0}^{2} (1+K)}$  (28b)

$$z^{\star} = z/z_{o}$$
 (28c)

$$\beta_{0} = \frac{z_{0}^{v}}{D_{1}} \qquad (28d)$$

The quantity  $z_0$  is an arbitrary length scale that, if reasonably chosen, can give some idea of the relative importance of diffusion and convection. (The final answer will not, of course, depend on  $z_0$ .)

Equation 26, subject to the boundary conditions of Equations 27a-27c, can be solved by using LaPlace transformations. One finds

$$C_{sw}^{*}(z^{*},t^{*}) = \frac{(C_{w}^{*}-1)}{2} \left\{ erfc(\eta_{1}) + exp(-\beta_{0}z^{*}) erfc(\eta_{2}) \right\},$$
  
(29a)

and

where

$$\eta_1 = \frac{-z^*}{2t^{*1/2}} - \frac{\beta_0 t^{*1/2}}{2}$$
(29b)

and

$$n_2 = \frac{-z^*}{2t^{*1/2}} + \frac{\beta_0 t^{*1/2}}{2} \qquad . \tag{29c}$$

The flux, N, is given by

$$N = -\phi D_1 \left(\frac{\partial C_{gw}}{\partial z}\right)_{z=0} - \phi V C_{gw}(o) \qquad (30)$$

In dimensionless form,

$$N^{\star} = \frac{z_{o}N}{\phi D_{1}C_{sw,\infty}} = -\left(\frac{\partial C_{sw}}{\partial z}\right)_{z=0} - \beta_{o}C_{w}^{\star} \qquad (31)$$

Differentiating Equation 28 and substituting into Equation 31, one finds

$$N^{*} = \frac{(1 - C_{w}^{*})}{(\pi t^{*})^{1/2}} \exp(\frac{-\beta_{o}^{2} t^{*}}{4}) - \frac{\beta_{c}}{2}(1 - C_{w}^{*}) \operatorname{erfc}(\frac{\beta_{o} c^{*}}{2}) - \beta_{o} C_{w}^{*}$$
(32)

Equation 32 can be cast in a form that explicitly removes  $z_0$  from the equation for the flux. If we define a new dimensionless flux,

$$\widetilde{N} = \frac{N}{\phi v C_{gw,\infty}} , \qquad (33)$$

and a new dimensionless time,

$$\tilde{t} = \frac{v^2 t}{D_1(1+K)} , \qquad (34)$$

then Equation 32 becomes

$$\widetilde{N} = \frac{(1 - C_{w}^{*})}{(\pi \widetilde{t})} \exp(-\widetilde{t}/4) - (\frac{1 - C_{w}^{*}}{2}) \operatorname{erfc}(\frac{\widetilde{t}}{2}) - C_{w}^{*} \qquad (35)$$

## Relationship To SRI Lagoon Model

Equation 32 is the basis for computing how much material transfers into or out of the sediment during a given time interval.<sup>\*</sup> If M is the total mass of material per unit area transported across the water/sediment interface in a time interval,  $\tau$ ,

then

$$\widetilde{M} = \int_{0}^{\tau} \widetilde{N}d\widetilde{t} , \qquad (36)$$

with

$$\widetilde{M} = \frac{vM}{\phi D_1 (1+K)C_{gW,\varpi}}$$
(37)

Note that the z direction was chosen positive in Figure B-1. Therefore N > 0 implies that material moves from the sediment into the water.

$$\tau^* = \frac{v^2 \tau}{D_1(1+K)} .$$
 (38)

Substituting Equation 35 into Equation 36, one finds

$$\widetilde{M} = \left(1 - C_{W}^{\star}\right) \left\{ \operatorname{erf} \left(\frac{\tau^{\star^{1/2}}}{2}\right) - \frac{\tau}{2} \operatorname{erfc} \left(\frac{\tau^{\star^{1/2}}}{2}\right) + \frac{\tau^{\star^{1/2}}}{\sqrt{\pi}} \exp(-\tau^{\star/4}) \right\} - C_{W}^{\star} \tau^{\star} .$$

(39)

The functions erf and erfc are well-known, tabulated integrals.<sup>\*</sup> These functions are available on any computer that has the standard IMSL math library software.<sup>†</sup>

In the SRI lagoon model, one knows  $C_w^{\star}$  and  $C_{sw,\infty}$  at the beginning of a time step (and, by implication, at time zero). During a time step, one allows mass to transfer between the water and the sediment/water phases. The total amount of mass transferred is given by Equation 39. At the end of the time step, the new values of  $C_w^{\star}$  and  $C_{sw},^{\infty}$  are computed based on the addition or depletion of contaminant mass in the water or sediment/water phases. Computations are repeated as desired with these new values of  $C_w^{\star}$  and  $C_{sw},^{\infty}$ .

and

Carslaw, H.S. and J.C. Jaeger (1959) Conduction of Heat in Solids. Oxford-at-the-Clarendon, London.

Abramowitz, M. and I.A. Stegun (1964) Handbook of Mathematical Functions, National Bureau of Standards, Applied Mathematics Series 55, June, 1964.

#### Sample Calculation

The flux, N, and the total amount transformed, M, will be computed for the particular parameters relevant to the TNT and RDX disposal lagoon. The values of the parameters are given by Equations 23a, 23b, 23c, and 23d. In addition, we must specify a value for K and  $\phi$ . For TNT, K  $\stackrel{\sim}{-}$  10; no data are available for RDX. Consequently, the calculations below apply only to TNT. The porosity,  $\phi$ , which has been measured by SRI, is 0.2.

With those values of the parameters, Tables B-2 and B-3 can be generated from Equations 33 and 39. It is worthwhile to record the values of the groups of parameters appearing in Equations 33, 34, and 37. Each group is a "characteristic" value, and we define the following parameters:

$$N_o = \phi v C_{sw,w} = 1.872 \ \mu g \ cm^{-2} \ day^{-1}$$
 (40)

$$M_{o} = \frac{\phi D_{1} (1+K) C_{gW, \infty}}{v} = 154.8 \ \mu g \ cm^{-2}$$
(41)

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FLUX CALCULATIONS FOR SAMPLE LAGOON PROBLEM

	<u> Time</u>		Flux
<u> </u>	<u>t (hr)</u>	<u> </u>	N ( $\mu g \ cm^{-2} \ day^{-1}$ )
0.01	19.8	4.57	8.55
0.04	79.3	2.01	3.77
0.09	178	1.18	2.21
0.16	317	0.769	1.44
0.25	496	0.528	0.989
0.36	714	0.371	0.695
0.49	971	0.263	0.491
0.64	1269	0.184	0.344
1.00	1982	0.080	0.149

## Table B-3

TOTAL MASS PER UNIT AREA TRANSFERRED DURING A TIME INTERVAL FOR SAMPLE LAGOON PROBLEM

<u>Time</u>	Interval	Mass 1	ransferred
<u>τ</u> *	<u>τ (hr)</u>	<u> </u>	$M (\mu g cm^{-2})$
0.01	19.8	0.0961	14.88
0.04	79.3	0.182	28.14
0.09	178	0.257	39.86
0.16	317	0.324	50.10
0.25	496	0.381	58.95
0.36	714	0.429	66.48
0.49	971	0.470	72.78
0.64	1269	0.503	77.89
1.00	1982	0.548	84.86

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$$t_o = \frac{D_1(1+K)}{v^2} = 1982 \text{ hr}$$
 (42)

Equation 42 shows that a steady-state solution for the flux would be inadequate unless time scales of interest were much longer than 1982 hr (~ 83 days). Time steps taken in the SRI lagoon model are variable; however, time steps shorter than 83 days necessitate the use of an unsteady solution as developed herein. Equation 40 states that a characteristic value of the flux is approximately 2  $\mu$ g cm<sup>-2</sup> day<sup>-1</sup>.

Part of the significance of  $N_0$ ,  $M_0$ , and  $t_0$  is that any situation with the same values of  $N_0$ ,  $t_0$ , and  $C_w^*$  will be the same.  $M_0$  is <u>not</u> an independent characteristic value because

$$M_{o} = N_{o} t_{o}$$
 (43)

#### Addition of Dissolution to the Mass Transfer Problem

The above solutions and assumptions were derived for the case where the dissolution rate constant  $(k_r)$  equals zero. We must also account for the case where  $k_r \neq 0$ , which occurs when the lagoon is full or when the lagoon waters cover the area near the dump ramp. As previously developed, the dissolution rate of solid TNT or RDX,  $\frac{dw}{dt}$ , can be written as

$$\frac{dw}{dt} = k_r (C_{sat} - C_i) , \qquad (44)$$

where  $k_r$  is an experimentally determined dissolution rate constant,  $C_{sat}$  is the saturation concentration of the dissolving species in the liquid phase, and  $C_i$  is the concentration of the dissolved species at the solid/liquid interface. If diffusion from the interface to the surrounding liquid is fast compared to dissolution, then Equation 44

can be represented as shown in Equation 45:

$$\frac{dw}{dt} = k_r (C_{sat} - C_{sw}) , \qquad (45)$$

where  $C_{gw}$  is the concentration of TNT or RDX in the sediment/water phase at any depth in the sediment. Therefore, Equation 45 can be incorporated into the transport equation developed in the previous section, as shown in Equation 46:

$$(1 + K)\frac{\partial C_{sw}}{\partial t} = D_1 \frac{\partial^2 C_{sw}}{\partial z^2} + k_{dis} [C_{sat} - C_{sw}] + v \frac{\partial C_{sw}}{\partial z} , \qquad (46)$$

where

$$K = \left(\frac{1-\phi}{\phi}\right) K_{eq} F_{oc} , \qquad (47)$$

and

$$k_{dis} = \frac{Fk_{r}S_{bet}\rho_{bs}(1-\phi)}{\phi}, \qquad (48)$$

and where  $F_{oc}$  is the organic fraction of the sediment solids,  $S_{bet}$  is the surface area of solid per unit mass of solid (typically determined by the BET absorption method),  $\rho_{bs}$  is the mass density of the sediment solids,  $\phi$  is the void fraction and F is the fraction of the solid surface area that is covered by the dissolving species.

A linear isotherm has been assumed in deriving Equation 46, with

$$C_{s} = K_{eq}C_{sw}, \qquad (49)$$

where  $C_s$  is the concentration of TNT or RDX dissolved in the organic

fraction of the sediment solids and  $K_{eq}$  is experimentally determined.

The boundary conditions for Equation 46 are:

$$C_{sw}(-\infty,t) = known = C_{sw,\infty} t > 0$$
 (50a)

$$C_{sw}(0,t) = C_{w} t > 0$$
 (50b)

$$C_{sw}(z,0) = C_{sat}$$
, (50c)

where  $C_w$  is the concentration of TNT or RDX in the bulk-water phase.

The flux, N, is given by

$$N = -\phi D_1 \left( \frac{\partial C_{sw}}{\partial z} \right)_{z = 0} + (1 - \phi) k_r F \left( C_{sat} - C_w \right) - \phi v C_w \qquad .$$
(51)

If Equation 46 can be solved subject to Equations 50a-50c, then the flux can be computed from Equation 51. Equation 46 will describe the flux in the laboratory experiment and in the lagoon. With laboratory data, we can find the value of  $k_r$  and use it to predict the flux in the lagoon.

#### Solution to Flux Equations for Laboratory Experiment

In the previous sections, Equation 46 was solved with  $k_r = 0$ (i.e., no dissolution). In the laboratory experiment, however, v = 0and  $k_r \neq 0$  (i.e., there is no seepage but there is dissolution of solid TNT and RDX). Also in the laboratory,  $C_{sw,\infty} = C_{sat}$ . Consequently, the mass transfer equations become

$$(1 + K) \frac{\partial C_{sw}}{\partial t} = D_1 \frac{\partial^2 C_{sw}}{\partial z^2} + k_{dis} [C_{sat} - C_{sw}] , \qquad (52)$$

with 
$$C_{gw}(-\infty,t) = C_{sat}$$
, (53a)

$$C_{gw}(0,t) = C_{w}$$
, (53b)

and 
$$C_{sw}(z,0) = C_{sat}$$
 (54)

The flux is given by

$$N = -\phi D_1 \left( \frac{\partial C_{gw}}{\partial z} \right)_z = 0 + (1-\phi) Fk_r \left( C_{gat} - C_w \right) \quad . \quad (55)$$

At steady state, we can solve Equation 52 subject to the boundary conditions 53a and 53b. We find

$$\frac{C_{sat} - C_{sw}}{C_{sat} - C_{w}} = \exp\left(\left(\frac{k_{dis}}{D_{1}}\right)^{1/2} z\right) \qquad . \tag{56}$$

The flux, N, is then given by

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$$\frac{N}{N_o} = (1 + \beta_D) (1 - C_w^*) , \qquad (57)$$

with N<sub>o</sub> = C<sub>sat</sub> 
$$(\phi D_1 k_r FS_{bet} \rho_{bs} (1-\phi))^{1/2}$$
, (58)

$$\beta_{\rm D} = \left( \frac{Fk_{\rm r}(1-\phi)}{S_{\rm bet}^{\rho} bs^{\phi \rm D}} \right)^{1/2} , \qquad (59)$$

and 
$$C_w^* = \frac{C_w}{C_{sat}}$$
 - (60)

In Equations 58, 59, and 60,  $\phi$ , D, and C<sub>sat</sub> are known. The surface fraction, F, approximately equals the (known) mass fraction. The product S<sub>bet</sub> $\rho_{bs}$  can be estimated by assuming a reasonable particle size for the sediment solids. If the particle diameter is d<sub>p</sub>, then

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$$S_{bet}^{\rho}_{bs} = \frac{6}{d_p} \qquad (61)$$

Sediment that can be described as fine soil or clay has particle sizes in the range of 10 to 100  $\mu$ m. (The square root dependence of the flux on S<sub>bet</sub> $\rho_{bs}$  de-emphasizes the importance of a proper choice for d<sub>p</sub>.)

Because all parameters in Equations 58, 59, and 60 except  $k_r$  are known or can be reasonably estimated, the laboratory flux data can be used to find a best value of  $k_r$ . To make this procedure a bit more transparent, we rearrange Equations 57 through 62, substitute Equation 61 for  $S_{bet}^{\rho}p_{bs}$ , and obtain

$$N = (C_{sat} - C_w) \left\{ \left| \phi(1-\phi) \left(\frac{\delta D_1}{d_p}\right) k_r F \right|^{1/2} + k_r F(1-\phi) \right\}$$
(62)

In the flux experiments, N and  $C_w$  are measured and a plot of N versus  $C_{sat}-C_w$  will have a slope equal to the bracketed expression in Equation 17.

A least-squares fit of the data to the form y = mx yields the following values for the bracketed expression in Equation 62:



TNT

$$\left(\phi(1-\phi) \left(\frac{6D_1}{d_p}\right) \kappa_r F\right)^{1/2} + \kappa_r F(1-\phi) = 7.752 \times 10^5 \text{ cm sec}^{-1} \qquad .$$
(64)

The values of the known parameters in Equations 63 and 64 are listed in Table B-4 for TNT and RDX. The diffusion coefficient is the average result from Equations 8-34 and 8-36 of Reid and Sherwood.<sup>\*</sup> The void volume fraction,  $\phi$ , was calculated from the dry weight of sediment solids, measured prior to the flux experiments. The saturation concentrations are known from solubility data reported in Section V.A., Physical Transport. The surface fraction, F, is assumed equal to the mass fraction of TNT or RDX in the sediment solids, a quantity measured prior to the flux experiments. The value of d<sub>p</sub> is an estimate.

From Equations 63 and 64 and the values of the parameters in Table B-4, we find that the  $k_r$  is  $1.172 \times 10^{-5}$  cm sec for TNT and  $1.164 \times 10^{-5}$  cm sec<sup>-1</sup> for RDX.

\*Reid, R.D. and T.K. Sherwood (1958) The Properties of Gases and Liquids. McGraw-Hill, New York. The flux derived from Equation 62 with these values of  $k_r$  has been placed on Figure B-3 with the data points to show the goodness of fit. One can see that Equation 62 is within a factor of 2 in predicting the flux over the range studied experimentally. This degree of accuracy can be disturbing unless one allows for the inherent difficulty of performing the flux measurements and of describing analytically the mass transfer in a multicomponent, heterogeneous system. One must also refrain from the temptation to fit the data in Figure B-3 to a straight line of the form y = mx + b. The result of this seemingly intuitive approach is a model that predicts negative flux values, a physically impossible situation in the laboratory experiment.

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## Table B-4

## PARAMETERS FOR ANALYZING FLUX DATA

Parameter	TNT	RDX
φ	0.379	0.579
D <sub>1</sub> (25°C)	$2.13 \times 10^{-5} \text{ cm}^2 \text{ sec}^{-1}$	$2.24 \times 10^{-5} \text{ cm}^2 \text{ sec}^{-1}$
C <sub>sat</sub> (25°C)	131 μg cm <sup>-3</sup>	50.6 µg сm <sup>-3</sup>
F	0.156	0.078
<sup>d</sup> p	50 µma	50 µm.

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FIGURE B-3 TNT (•) AND RDX (0) FLUX DATA COMPARED WITH CURVE FIT (EQUATION 62)

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#### Appendix C

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## SRI COMPUTER PROGRAM FOR MODELING THE FATE OF TNT AND RDX IN LAGOONS CONTAINING MUNITIONS WASTES

This appendix presents the computer program developed to simulate TNT and RDX in a waste-lagoon environment. It encompasses a number of subroutines to determine changes in volume, surface area, and depth of the lagoon and the transport of munitions (flux) from the bottom sediment to the free-water phase. The subroutine COMPS performs the calculation of munition concentration in the bulk-water phase and in the suspended sediment. The data output can be conveniently displayed graphically by the subroutine GRAPH 4.

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	_0009	<u> </u>		
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02100	0020	č	PC PARTITION	
02200	0022	č	SD DENSITY OF SEDIMENT	
02300	0023	č	wku uxidation hate constatnt	
02400	0024	č	WKP PHOTOLYSIS RATE CONSTANT	
02500	0025	c	NKH HIDHULYSIS RATE CONSTANT	
02600	0026	č	WHY VOLATILIZATION RATE CONSTANT	
02700	0027	ć	WKB BIODEGRATION RATE CUNSTANT	
02800	0024	С	PR PRECIPITATION RATE, CM/DAY	
02900	0029	C	EV EVAPORATION RATE, CH/DAY	
03000	0030		SP SEEPAGE RATE, CM/DAY	
03100	0031	С	DIF DIFFUSIVITY	
03200	0032	С	PUEPTH DEPHTH AT THE LOCATION CORRESPONDENTING TO SEDIMENT	
03300	0033		COMPARTMENT	
03400	0034	С	TMASS TUTAL TRANSPORTATION RATE OF THE ACROSSING THE	
03500	0035	С	INTERPHASE	
03600	0036	<u> </u>	NS NUMBERS OF SEDIMENT COMPARTMENTS	
03700	0037	Ç	NW NUMBERS OF SEDIMENT COMPARTMENTS CUVERED WITH FREE WATER	
03800	0038	С	PHI VOID FRACTION	
03802	_ 00.19	<u> </u>	KK is equal to 2, only dissolution process is considered.	
03804	0ú40	c	rk is the dissolution rate constant	
03806	0041	C	ts and sm are the parameters for calculating the solubility	
<u>03808</u>	0042	<u> </u>	ta is the solubility	
U3810	0043		-DIMENSION Y(100), X(100), 1T(3), JA(7)	
03850	0044		COMMON/PAKA/NS,NW,DT,TIME,PRT,T,WT Commun (Const/A)	
03855	0045		COMMUN/CONST/A,B COMMUN/RATE/WKO,WKP,WKH,WKV,WKB	
03800	0048		COMMUN/PHYS1/NJ,ALPHA,TMASS,4C,DIF,PH1,SD,SCUUK,SETL	
03672	0048		<u>COMMON/GRAPH4/SCA, ANG, UX, UX, VX, VX, XH1, XH1, XHA</u>	
03680	0049		COMMUN/PHIS2/PR,EV,SP,VOL,PUEPTH(SU), area, depth(SU)	
03881	0050		COMMUN/CONCN/C, TSS, SSP, CSW(50), CMT(50)	
03883	0051	_	-CHARACTER+80 17,1STR	
03884	0052		_ DATA IT(1)/'SINULATION TIME'/	
03885	0053		_DATA 1T(2)/'DAYS'/,1T(3)/'Tox concentration, ppm'/	
U3886	0054		-DATA 1A/0,0,0,0,1,0,1/	
	0055		LUGICAL TPRINT	
03667				
03888	0056		OPEN(UN11=5,HAME='RDX,DAT',STATUS='OLD')	
	29700	0001		sudfoutine flux]
---	---------	-------------	----------	---
	29600	0002	c	mass traisfer rate coefficient is 0.235 for tht is 15%
	<u></u>		<b>_</b>	_ 11.15 ASSUMED that mass tlansfer coefficient is proportional to
	30000	0004	c	the tht deposition percentage.
	30100	0005		common/concn/c,tss,ssp,csv(50),cmt(50)
	40110	0006		COMMON/PARA/NS.NW.DI.JIME.PRI.I.MI
	30200	<b>UU07</b>	-	COMMUN/PHISZ/PH,EV,SP,VUL,PUEPTH(50),AREA,UEPIH(50)
	30300	0008		CUMMUN/PHISS/UM(50), DMASS, 1MAS(50)
	10310	0009		
_	30320	0010		CUMMUN/Phisi/NJ, ALPHA, TMASS, PC, DIF, PH1, SU, SCOUK, SEIL
	30322	0011		common/solu/ta
	30324	0012		COMBOD/SOLUI/IK.Sm.ts
	30400	0013		DMASS=0.
	30402	0014		it(t .le. 70.) go to 300
	30944	_ 2115_		1111 gt. 70. and t Je. 130.1 90 LO. 350
	30406	0016		if(t .gt, 130, .and, t .le, 210) yo to 400
	30408	0017		11(t .gt. 210.) go to 450
_	30500	0018		temp=263.
	30501	0019		go to 500
	30502	0020	350	temp=293.
	30503	0021		
	30504	V022	400	temp=303.
	30505	0023		ao to 500
	30506	0024	450	
	30507	0025	500	ta=sm*10.**(8.)*exp(-ts/temp)
	30508	0026		do 100 1=1, nj
	10510	0027		$J \equiv N ( (NS+1) = 1)$
_	30600	0028		<pre>dm(j)=rk*cmt(j)*area/1000,*(ta*1000,-c)</pre>
	30610	0029	c	dm(j) is in ug per day
	30700	0030	100	UMASS=DMASS+DM (J)
	30710	0031		RAITE (6,200) UMASS
	30720	0032	200	FORMAT(7H DMASSE,E12.4)
	10800	0033		RETURN
	30900	0034		END

21500	0301		
21000	<b>ύ</b> υί∠		SURFOUTINE FLUX
2/740			THIS SUBRUITINE IS TO CALCULATE THE TUTAL LEANSPORT OF
27800	0004	c	CHEMICAL INKUUGH SEDIMENI INIEKFACE
27810	0005		common/pala/ns.nw.dt.time.plt.t.wt
27906	0006		CUMMON/PHISI/AJ, ALPHA, THASS, PC. DIF, PHI, SD, SCUDR, SETL
28000	0007		COMMUN/CUNCN/C, TSS, SSP, CSN(50), CHT(50)
28010	0008		CUMMUN/PHIS3/DP(50), UNASS, TMAS(50)
20150			
28055	0010		common/water/n(50)
28057	0011		common/solu/te
28070	0012		
20000	0013	1000	format(Sn flux,15)
20100	0014		THASS=0,
28200	0015		
28300	0016		j=N((NS+1)+1)
26310	0017	c	-
27320	0016	5.44	temperature adjustment on solubility
28.130	0019	c	
28340	0020		tb=ta/cs=())+1000.
28350	0021		<u></u>
28400	0022		condl=c/(csw(j)*tb)
24540	0023		TD=SP##2.#DT/(DIF#(1.#ALPHA))
	.0024		
28800	0025		bb#exp(+tg/4.0)
28900	0026		d=1.=erf(td==(0.5)/2.)
29000	0027		tmas()]=(###D/(3,1416#td)##(0,5)=#z/2,0#d=cond)
29010	VU28	c	
29020	0029	C###	ceta in microgram per square centimeter per day
	0030	¢	
29100	0031		petaxpn1#sp#c%w(j)#tD#10,##(-3)
29105	U032		tmas(j)=Deta+tmas(j)
29300	0011		<u>THASSZTHASS+JHAS(J)#ARLA</u>
29400	0034	100	CONTINUE
29500	0035		RETURN
29600	0036		END
-	2/600 2/800 2/800 2/800 2/800 2/800 2/800 2/8055 2/8055 2/8057 2/8057 2/8057 2/8050 2/8050 2/800 2/9010 2/9010 2/9010 2/9010 2/9010 2/9010 2/9010 2/9010 2/9010 2/9010 2/9010 2/9010 2/9010 2/9010 2/9010 2/9010 2/9100 2/9100 2/9010 2/9100 2/9500	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$

1. S. S. S. S.

COMPS :

20410	0058	5	+tmass/vol=cu*(ropr=roev=rosP)/vol)/(l.=tss*pc*j0.**(-1.))	1
20820	<b>UU59</b>	•	1f(c .gt. u.) go to 500	
	0000		CAC0+01+(=\$UBK+C	
20910	0061	<b>\$</b> -	+tmass/vol=C+(ropr=roev=rosp)/vol)/(1,=tss+pc+10.++(-3.))	
26912	0062	500	1F(C .G1. 0.) GU TO 700	
26913	Oub3		<u>11 (01 .89. 1.) GO TO 700</u>	
26914	Ú064		DT=DT/7.	
20910	0065		PF7=PK1/7.	
20918-	_UUDD			
26920	0067	700	CONTINUE	
26930	0068		if(c _gt. ta+1000_) c=ta+1000.	
26950	.0009		1 ± (C) = 0, ) C=0.	
27000	0070		SSP=C/1000.+PC	
27100	0071	С	SSP IN UG/G	
27110	0072		tmac*vol	
27200	0073		RETURN	
27400	U074		END	

PRUGRAM SECTIONS

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	Bytes_	Attfibutes
0 SCUDE	688	PIC CON REL LCL SHR EXE RD NUWRT LONG
	158	PIC CON REL LCL SHR NUEXE RD NOWRT LONG
2 SLUCAL	52	PIC CUN REL LCL NUSHR NOEXE RD WRT LONG
3 PARA	28	PIC OVE REL GEL SHE NUEXE RD WET LONG
4 RATE	20	PIC OVE REL GEL SHE NUEXE RD WRI LONG
5 PHISI	46	PIC OVA REL GAL SHA NOLXE AD WAT LUNG
0 PHYS2	420	PIC OVE REL GEL SHR NUEXE RD WRT LUNG
/ PHYS3	404	PIC OVE REL GEL SHE NOEXE RD WET LONG
B CONCN	412	PIC OVE REL GEL SHE NULXE RD WET LONG
9 MASS	4	PIC OVE HEL GEL SHE NUEXE RD WAT LONG
10 RATE1		PIC OVE REL GEL SHE NOEXE RD WET LONG
11 CONST	8	PIC OVE REL GEL SHR NUEXE RD WRT LUNG
12 WATER	200	PIC OVE REL GBL SHR NOEXE RD ERT LONG
13 NEA1H	12	PIC OVE REL GEL SHE NUEXE RD WET LONG
14 SOLU	4	PIC OVR REL GBL SHR NOEXE RD WRT LUNG
15 CONIRL	12	PIC OVE REL GEL SHR NUEXE RD WRT LUNG

lotal Space Allocated

2466

ENTRY PUINTS

Address Type Name

0-00000000

COMPS

25200	0001		
25300	0002		
25400	FORD		SUBRUUTINE COMPS
25500	0004	C	THIS SUBRUUTINE IS TO CALCULATE THE CUNCENTRATION IN FREE WATER
25000	0005	c	AND ON SUSPENDED SULID
45700	UUUD	-	CONKUN/PANA/NS, Na, D1, 11ML, PR1, 1, a1
25800	0007		Charles Foren and and the set of
25900	0008		CUMMUN/KA1E/wKU, whF, whH, why, whB
			CUMMUN/PHYSI/NJ, ALPHA, THASS, PC, DIF, PHI, SD, SCOUK, SETL
			CUMPUN/Phis2/Ph.EV.SP.VUL, PUEPIH(50), drea, depth(50)
	0010		COMMOR/PHYS3/DM(50), DMASS, 1MAS(50)
26100	0011		COMMUN/CUNCH/C,ISS,SSP,CSW(5V),CN1(5V)
	V012_		
20110	0013		CUMHUN/KAILI/AWKP,AWKB
20120	0014		CUMMON/CUNS1/A,B
			COMMON/#A1ER/N(50)
26132	0016		common/weath/ropr,roev,rosp
20134	0017		conmon/solu/ta
Zoldo	0018		Common/contrl/KK,Kl.Kn.
20140	0019		avh=vo]*ju**(3,)/(area*nj)
26160	0020		write(0,400) avh.awko
20170	0021	400	format(150 average cepth=,e10.4.20h a0justeo photolysis rate=,
26180	Ú022	\$	el0.4)
20182	0023	•	if(amod(t,50.) .ne, 0.) go to 900
20184	0024		<u>it(t_,gt_,224_)_a=1</u>
20160	0025	ويعي وي النباس الله والأقومية	
20168	0026		
26189	0027		a#Kb=#Kb=2.++(b)
		0.0.4	
26190	0028	900	wiite(6,800) awkb
26192	0029	800	format(6h_awkp=,e12.4)
20200	0030		SUMK#WKO+@WKD+WKH+WKV+AWKB
26400	0031		C0=C
* 2041U	0032	c	
20420_	_دورو	<u></u>	<u>1, 101 1, 101 1, 116 1</u>
20430	0034	C	
20440	0035		call flux1
26445	0036	<u> </u>	
20450	0037	C***	call tor flux
20455	0038	С	
20457	0039	s^	
20400	0040		do 3u0 i=i.ni
20402	0041		J=N((NS+1)-1)
20405_	0042		
20407	0043		1:(kk ,eq. 2) tt=am(j)
20408	0044		IF(AMUD(I,7.) .NE. 0.) GO TO 300
20470	0045		<u>white(6,200)</u> J.T1
20475	0046	200	
20460	0047	300	FURMAT(20H DIFFUSIUN RATE FRUN, 15, 4H IS , E12, 4) CONTINUE
	V048	300	
20487	0049		TMASS=1MASS+DMASS
26490	0049		lf(Kk .eq. 2) thass=dmass
20490	0051		WRITE(6,350)1MASS
20495	and the second s		
	0052	350	FORMAT (40H TUTAL DIFFUSION HATE THROUGH BOITOM SEDIMENTE, E12.4)
26500	0053	C	
26600		C+++	SS_1N_G/LITER,C_IN_UG/LITER
26700	0055	C	
26710	0056	000	continue
26800	0057		_c=c0+(dt/2)*(~sumk*c0

	24130	KCUU		11(Kn .nt. 2) go to 500
	24737	.0059	_	if(t .qt. 16h.) wkpt=2.4wkpt
	24/40	UÚ61		11(WKPt .9t. WKP+0.) WKPtmwKp+6.
÷	24745	0062		dwkp#awkp+wkpt .
<b></b>	24760	0063		
	24800	0064		write(6,150) n((ns+1)=L),pdepth(1),delta
	24810	0065	200	continue
				<u></u>
i	24850	0067		AwkPaawkP/NJ
i i	24900	0068	150	tormat(15,2e12.4)
	24902	0069	350	continue
	24910	0070		write(6,1000)nj
	24920	0071	1000	tormat(Wh surtace,15)
	25000	<u>072</u>		RETURN
	25100	0073		END

PRUGRAM SECTIONS

•	Name	Bytes	Attributes
v	SCUDE	645	PIC CUN HEL LCL SHR EXE NU NUMPT LUNG
	SPUATA	31	PIC CUN REL LCL SHR NUEXE RD NUWRT LUNG
	SLUCAL	56	PIC CUN REL LCL NUSHR NUEXE RD WRT LUNG
	PARA	28	PIC OVE REL GEL SHE NOEXE RD WET LUNG
	Ph151 Ph152	36	PIC UVE REL GBL SHR NOEXE RD WRT LUNG
	WALER		PIC UVR REL GBL SHR NUEXE RD WRT LUNG
	KAILI	200	PIC OVE REL GEL SHE NUELE RD WET LONG
	CUNTRL	12	PIC OVE HEL GEL SHE NOEXE RU WAT LONG
9	KAIL	20	PIC UVR HEL GOL SHE NOEXE RU WET LUNG PIC UVR REL GUL SHE NOEXE RD WET LONG
	Iotal Space Allocated	1456	PIC OVR REL GBL SHR NOLXE RD WHI LONG

ENTEL PUINTS

Address Type Name

U-UUUUUUUU SURFACE

VARIABLES

	agress	Type	Name	Address	TYpe	hame
	00000004	• • •	ALPHA Del1a	5-00000008 2-00000018	∦\$4 R\$4	AREA DEL1D
9 <b>-</b> (	0000000	R#4 1#4 1#4	EV KK M	2=00000008 8=00000004 4=00000000	1*4 1*4 1*4	1 KL
4 - ( 4 - (	0000000C 0000001C 0000000	R*4 R*4 R*4	PC SCOUR SUM	4-0000014 4-0000018 2-0000010	R#4 R#4 R#4	NJ PH1 SD SUM1

Addr	255	Туре	Name	Address	Туре	Name
7-0000	0004	H=4	A#K5	7-00000000	H#4	A
4-00000		K#4	DIF	3=00000008	R+4	Анкр — D1
2-00000		144	J	2-00000000	144	<u>- K</u>
8-00000		1+4	KN	2-00000024	1+4	ĥ
3-60666		1+4	NS	3-00000004	1+4	Nh
5-00000		H+4	99	3-00000010	R+4	PHT
4-00000		R#4	SETL	5-00000008	H+4	SP
3-0000	014	F # 4	1	3-00000000	R#4	TIME

51HAA	0001		
21200	<b>JUJ</b> 2		ANTALISING BUTING
22000			. SUDIOUTING SUITAGE TO CHECK THE SUBLACE WEICH IN COVERED
55100	0004	ç	THIS SUBFUUTINE IS TO CHECK THE SURFACE WHICH IS COVERED
22200	0005	С	NITH FREE WATER
22300	Uu06	····	COMMUN/PARA/NS.NW.DT.TIME.PR1.1.+T
22400	0007		CUMMUN/PHISI/NJ, ALPHA, IMASS, PC, UIF, PHI, SU, SCOUK, SEIL
22500	00 <b>0</b> 8		CUMMUN/PHIS2/PF,EV,SP,VUL,PDEFIn(50),area,deptn(50)
	<u> </u>		CORBON/NHLEI/D(50)
22507	001U		COMMUN/KAIEI/AWEP,AWEB
22500	0011		common/contil/kk,kl,kn
			COMNUN/RAIL/NKU, WKF, WKH, WKY, WAB
22510	0013	C	
22515	0014	Č+++	WHEN POND IS DRIED
22520	0015		
22525	0016		IF(VOL _GT. 0.) GO TO 300
22530	0017		NJ=0
22531	01B		NEITE(6.400) NJ. YUL
22532	0019	400	FORMAT(15, E12.4)
22535	0020		GO 10 200
22540	0021	300	
22600	0022		SUM#U.
22700	0323		DO 100 J=1, (NS+1)
22800	0024		1=N((NS+1)=J)
22900	0025		N=N(NS+J)
23000	0025		SUM=SUM+(DEP1H(1)=DEP1H(K))+AREA*J*10.**(-3.)
23010	0027		
23100	U028		IF (VOL ,GT, SUM) GO TO 110
23300	0029		DELTA=VOL-SUMI
	0029		
23310			IFIDELTA , LL, Q.) DELTA=VOL
*23400	0031		DELTD=DEL1A/(AREA+(j+1)+10.**(-3.))
23410	0032		
23500	_0033_		<u>GO TO 120</u>
23600	0034	110	SUMIESUM
23800	0035	100	CONTINUE
23501	0036	-	- M3 <del>3</del> 7
23602	0037	c	when need to table
23004	0038	C+++	when pond is tull
23406		C	
23810	0040		if(vol_le.sum) go to 120
23820	0041		delta=vol+sum1
23830	_0042		delta=delta/(area*(j+1)*10,**(-3,))
23840	0043		1
23900	0044	120	CONTINUE
24000	0045	<u> </u>	
24100	0046	C+++	DETERMINE WATER DEFTH ABOVE EACH SEDIMENT COMPARTMENT
24110	0047	с	
2412u	0048_		
24130	0049		AWKP=U.
24140	0050		*KPI=WNP
24200	0051		DG 200 L=1,nj
24400	0052		]=h((NS+1)=L)
24100	0053		PDEPTH(1)=+(DEPTH(1)=DEPTH(M))+DELTD
24710	0054	C	
24720	0055	C	ADJUSTING PHUTULYSIS RATE
24730	0056	č	
24/35		-	wkptzwkp#2.U/Ddepth(1)
	· · · · ·		ĸĸĸ₽₽₽₽₽ĸĊĸĸĸĸ₽₽₽₽₽₩₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽

19400	0001		
19500	0002		SUBRUUTINE VOLUME
TAONT-			THIS SUBRUUTINE IS TO ESTIMATE THE THEE MATEL YOLUME.
19700	0004		CUMMUN/PHIS2/PK, EV, SP, VOL, PULP (H(50), area, geptn(50)
19/02	0005		COBBON/WELET/N(50)
19744	4044		COMMON/PHISI/AJ, ALPHA, THASS/PC, DIF, PHI, SD, SCUUR/SEIL
19706	0007		Common/pais/ns,nw,dt,time,prt,t,wt
19706	0008		common/weath/ropr,roev,rosp
19800	0009		
19900	0010	C+++	CALCULATE THE TOTAL SEEPAGE DURING THE TIME INTERVAL
20000	0011	Ċ	
20100	0012		
20200	0013		rosp=1AREA+SF+10.++(-3.)
20300	0014	С	
20400	0015	C+++	CALCULATE THE TUTAL EVAPORATION DURING THE TIME INTERVAL
20500	0016	С	
20600	0017		ROEV=EV+TAREA+10.++(-3.)
20700	0019	<u> </u>	
20800	0019	C###	CALCULATE THE TOTAL PRECIPITATION DURING THE TIME INTERVAL
20900	UV20	С	
21000	0021		RUPH=PR+TAREA+10, ++ (-3,)
21100	0022	С	
21200	0023	C***	INTEGRATE THE VOLUME BALANCE EWQUATION
21300	0024		
21310	0025		VOL1=VUL
21400	0026	300	VOL=VOL1+DT+(KUPH-KUEV-RUSP)
21402	<u>_0027</u>	حريد في بيد جي	IF(aps((Vol-Vol1)/Vol1) .1t. 0.5) GD 10 200
21404	0028		IF(DT.EQ. 1.) GO TO 200
21406	0029		DT=DT/7.
_21408_	0030		<u>GO TO 300</u>
21409	0031	200	CUNTINUE
21410	0032	с	
21420	0033	C+++	check sulface area
21430	0034	C	
21440	0035		call surface
21500	0036		<u>16(VOL _ LE_ 0_) VOL=0.</u>
21600	0037		RETURN
21700	0038	r	END

1,000	0001		
13700	. ນັບບັນຊີ		
7.1400	UVUJ.		SUBROUTINE INITE
17300	0004	С	THIS SUBROUTINE IS TO INITIALIZE THE SIMULATION SYSTEM
17400	0005		COMMUN/CONCH/C,155,55P,C8w(50),CH1(50)
	2020	·	
17420	0007		common/pata/ns,nw,DT,Tlme,pT1,T,W1
1/430	UUUN		COMMON/MOSS/LR.
17500		C	
17000	0010	C+++	HEAD IN INITIAL CONCENTRATIONS
17700	0011	С	
17800	0012		
17900	0013	100	FURMAT(10E8.2)
17910	0014	C	
			read in initial concentrations in segiment
17430	0016	C	
17935	0017		do 200 i=1,ns
17940	VOIA		READ(5,150)_CSW(1),CMT(1)
17942	0019		wR11E(6,150) Csw(1),CMT(1)
17945	0020	200	continue
17950	0021	150	tormat (8610.4)
18000	0022	С	
1.8100	0023	C***	READ IN INITIAL VOLUME
18200	0024	C	
18210	0025	C	VUL 14 LITER, AREA IN (CN)++2
18300	0026		READ(5,100) VOL, AREA
18310	0027		tm=c+yol
18400	J028		wklTE(6,110)
18500	0029	110	FORMAT(101,200 IN)11AL CUNDITIONS )
18500	0030		WHITE(6,120) YOL
16700	0031	120 #	URMAT(0H VUL=,E12.4)
18500	ÚU32	•	NHITE(6,130)
18900	0033	130	FURMATIBH COMP, JUH CW (UG/L), JUH SS (UG/L), JOH CS (UG/L))
19000	0034		wpite(6,140) C,TSS,SSP
19100	0035	140	FUPMAT(101,3E10,2)
19200	0036		RETURN
19300	0037		ENU
12244	~~ )		

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139

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1-12-03

1300 13000 	0056 0059 	C C C	CUMPARED TO THE ONE BEFORE THE PREVIOUS ONE
13800-	0001		JF (DEPTH(N(M)) .GT. DEPTH(N(M-1))) GU 10 130
13400	0462		
14000			. h(M=1) #h(M)
14100	0064		N(N)=L
14200	0065		
	0060		<u>GD_TU_140</u>
14400	0067	120 N	(k+1)=1
14000	0068	130 C	ONTINUE
14700	0069		K#K+1
14800	0070	410 C	UNTINUE
14900	0071	C	
15040_	0072_	C***	DEFINE ALPHA AND DELTI
15100	0073	С	
15200	0074		ALPHA=1.8+10.++(-3.)+PC+(1PHI)/PHI
15400	0075		wpltt(6,300)
15500	0076	300	FORMAT(1H1,29HINPUTS OF KINETIC INFORMATION // 9H CUMPART-
15600	0077	\$,	4X, 5HPART., 9X, 4H X1, 8X, 7HPHUTUL-, 5X, 7HHYDRUL-, 5X, 8HVULATL-
15700	0^7 <u>8</u>		bx.6HM1CRU. / 3X.4HMENT.6X.6HCOLFF.7X.6HUA11UN.2(6A.4HYSIS.
15800	0079 0080	\$1	x),6X,7HIZATION,8X,6HBIUDEGJ
15900	0080		WRITE(6,125)NS,PC,WKO,WKP,WKH,WKV,WKB
<u> </u>	0082	C 123	FURMAT(/ 15,3X,9(3X,E10.3))
16200	0083	C***	WRITE DEPTH
10200	0084	C	while Derin
16310	0085		write(6,500)
16370	0085	500	format(1h1,7x,4hcomp,2x,5hdeptn/2x,5horder,4x,1h#,3x,2hcm)
16400	0087	300	DO 170 K#1,NS
16500	0089		WRITE(6,160)K,N(K),DEPTH(N(K))
<16500	0089	160	tormat(215,e10.2)
			TOIMBC(TT3)EIN#T3
10/00	0090		UNTINUE
16800	0090		ONTINUE
10/00	0090		UNTINUE
16800	0090		ONTINUE
16800	0090		ONTINUE RETURN END

いおうしじ	0001		
VALUU	0002		SUDNOUIINE INPUT
		<b>.</b>	THIS SUBFUUTINE IS TO DEFINE SIMULATION FAFAMETEPS
08800	0004		CUMPUN/PAHA/NS,Na,DT,TIME,PH1,1,a1
08500	v005		COMMUN/HAIE/nKO,NKP,NKH,NKV,NKE
09000	<u>DODP</u>		COMMON/PHISI/NJ.ALPHA.TMASS.PC.DIF.PHI.SD.SCUUK.SEIL
09100	0007		COMMON/PHISZ/PH, EV, SP, VOL, PUEPHIM(50), area, depth(50)
09110	0008		CORBON/WATES/N(50)
			CUMMUN/KATEL/ANNP, ANND
04130	0010		COMMON/SOLU1/IK,SM,TS
09140	0011		common/contri/kk,ki,kn
09390	_0012		
09400	V013	C+++	REAU TITLE INFORMATION
09500	0v14	c	
09400	0015		<u>HEAD(5,94) TITLE</u>
09610	0016		write(6,90) title
09700	0017	90	FORMAT(8A10)
00100	0018	<u> </u>	
U <b>99900</b>	0019	C+++	KEAD FUN CONTRUL DATA
10000	0010	с	
	.0021		<u></u>
10200	00 <u>2</u> 2	100	format(15,415,1,315)
10210	0U23		write(6,100) ns,time,dt,prt,#t
	0024	<u> </u>	
10400	0025	C+++	READ IN SETTLING AND SCOURING RATE
10500	0026	c	
LUDUU	0027		READ(5,150) SETL, SCOUR
10700	0028		FURMAT(2E10.3)
10800	0029	с	
10900		<u>C+++</u>	<u>READ_1N_SYSTEM_PAHAHETERS</u>
11000	ひいまた	c	
11100	0032		HEAD(5,200)PC,ph1,d1f,6D,wKG,mKP,mKH,mKV,mKB
11110	1033		
11200	0034	200	FORMA1(9f0,4)
11210	0035		Awk8≠≈k8
11400	0010	<u> </u>	
11400	0037	C+++	REAU IN CEPTH DATA
11500	0038	c	
11996			DD 490 J=1, NS
11700	0040	490	HEAD(5,400) DEPTH(J)
11800	0041	400	FURMAT(0E12.4)
11900	<u> </u>	<u> </u>	
12000	0043	C+++	ARRANGE CUMPARTMENTS IN ACCURUANCE WITH DEPTH
12100	0044	c	
12200	0045		
12300	2046		N(K)=1
12400	0047		N(N+1)=2
12500	0048		DD 410 1=2,N8
12000	0049	ç	
12700	0050	ç	CUMPARED TO THE PREVIOUS DEPTH
12500	0051	<u> </u>	
12960	0052	450	IF(DEPTH(1) .GT. DEPTH(N(K))) GO TO 120
13000	0053		Lun (K)
	0054		
13200	0055		N(K+1)#L Mak
13300	0050	140	11- (M .EO. 1) CU TU 130

	UBUUS U115	1F(A400(1,7.) .NF. 0.) GU 10 180
	08010 0116 C	SIUNE 1HE UAIA
	08040 0119 08030 0118 C	X(11)#T
	<u></u>	<u>v(11)mc/1000</u> 11=11+1
		CUNTINUE 
	0110 0124 01200 0125 	5x,2hc=.e12.4) GO TO 100 
	UU211 U127 C UU221 0120 C	DHAN THE CURVE
-	<u>08231 0129 C</u> 06233 0130	FROGEO.
. <u></u>		DU 10 ]=1,100 
	UB239 0133 06240 U134 	IF(FRUG .L1. 20.) FRUG#20. 11#11=1 
	UN241	CALL GRAPH4(X(1),Y(1),11,1,300.,0.,80.,0.,IT(_),IA(1))
-	0#2710138C 0#2#10139C	TERMINATION OF GRAPH
	06291 0140 08293 0141	CALL GRAPH4(X(1),Y(1),-1,1,0.,0.,0.,0.,11(1),1A(1)) CALL GPLUT(0,.0.,999)
	08300 0142 08400 0143	stop End
_		
_		
		•
		142
<b>.</b>		



04000	0058		7=0.
04000	0059	с	,=0,
U 4204		C+++	CALL FUE LIFUI
UAJUO	0061	C	
04400	0062	-	CALL INPUT
	_ [ هي ت		u1b12=D1/2
04600	0064	c	
U4700	0005	C+++	CALL FUR INITIAL CONCERTRATIONS
	_ מקונה	<b>c</b>	
04900	0067		call init!
U491U	0068		e=1.
u4920	0003		
04930	0070		]]#]
05000	0071	100	T#T+DT
05140	0022		<u>1F(T_G1, T1HE) GU_TU_310</u>
05200	0073		T2#T+,0000003
05300	0074		TPRINTE, FALSE.
05400	<u> </u>		
05700	0070	C	
05800	0077	C+++	HEAD IN RAIES OF EVAPORATION, SEEPAGE, AND PRECIPITATION
	<u> </u>	<b>c</b>	1517 NO 3 AND AUDITS -11 CF 000001 - CL 10 140
05910	0079		IF(T .NE. 1, .AND. AMUD(T2, .T) .GTUUUD01) GU TU 160
00000 _00100	0080 0081	150	READ(5,150) EV,SP,PR FORMAT/2610 AN
	0082	<u></u>	_ EOBMAT(3L19.4)
06110	0083	c	detetaination of sculting and setting
00110	0184	ć	determination of schuring and setting
00500	0085	<b>b</b>	RATIO=PR*AREA+N5/VUL+10.++(-3.)
00000	0086		IF(HAT]D .LT. 0.1) GO TO 160
N6740	0007		
00800	0068		SEL=(1+HAIJU-0.1)*SETL
06900	0089	160	CONTINUE
02000	0090	<u> </u>	
07100	0091	C+++	CALL FOP VOLUME
07200	0092		CALL VOLUME
_07402_	0093	<u> </u>	
07204	0094	¢	check to see the pond is dried out or not?
07206	0095	c	
	0090		1E1YOL _G1_ 0.1 GO 10 300
07220	0097		WR11E(6,350)T
07230	0098	350	FORMAT(26H THE POND IS DRIED OUT AT ,F5.1,5H DAYS)
01240	<u></u>		
07250	0100	300	CONTINUE
01300	0101	C C+++	
07400	<u>0102</u> 0103	<u> </u>	CALL FOR CUMPS
07500	0103	~	CALL CUMPS
0/600	_0105	с	CADA CONTO
<u>91790</u> 07600	U106	C+++	PRINT UUT THE SIMULATION RESULTS
0/900	0107	c	LUIUI AAI INC OTAARVIAN KEBARIO
	_V108		<u>11(yol . gt. y.) go to 210</u>
<u>9/349</u> U7920	0109		•rite(b,200)
07920	0110	200	for#at(22h th# pond is dried out)
v7940	0111		
0/950	0112	210	continue
07955	0113		ii(prt .#q. i.) tprint#.true.
08000	0114		<u>11 (10 RIN1) WHITE (6, 170) 1, VOL, AKEA, C</u>

143

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