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Approximation of Biodegradation Rate Constants for Monoaromatic Hydrocarbons (BTEX) in Ground Water

by Todd H. Wiedemeier, Matthew A. Swanson, John T. Wilson, Donald H. Kampbell, Ross N. Miller, and Jerry E. Hansen

Abstract

Two methods were used to approximate site-specific biodegradation rates of monoaromatic hydrocarbons (benzene, toluene, ethylbenzene, and xylene [BTEX]) dissolved in ground water. Both use data from monitoring wells and the hydrologic properties of the aquifer to estimate a biodegradation rate constant that can be used in ground water solute fate and transport models. The first method uses a biologically recalcitrant tracer in the ground water associated with the hydrocarbon plume to normalize changes in concentration of BTEX under anaerobic conditions; attenuation of the tracer is attributed to dilution, sorption, and/or volatilization. Attenuation of BTEX in excess of the attenuation of the tracer is attributed to biodegradation, although other processes may affect the observed rate. The second method assumes that the plume has evolved to a dynamic steady-state equilibrium. A one-dimensional analytical solution to the advection-dispersion equation is used to extract the rate of attenuation that would be necessary to produce a steady-state plume of the configuration found at the site. Attenuation is attributed largely to biodegradation because the analytical solution removes the effects of sorption and dispersion and volatilization is assumed to be minimal.

Neither method fully accounts for the effects of continuing dissolution of BTEX in the source area or nonlinear sorption. Therefore, the rates cannot be attributed fully to biodegradation, but still are useful for ground water contaminant fate and transport modeling. The methods were applied to a data set from a JP-4 jet fuel spill at Hill Air Force Base, Utah. In estimates along two separate flow paths, natural attenuation rates for BTEX ranged from 0.006 to 0.038 day⁻¹, with most rates near 0.02 day⁻¹. The rate for benzene ranged from 0.025 to 0.038 day⁻¹. The rates of attenuation of individual BTEX compounds as estimated by the two methods were in close agreement. For an individual compound, the rate estimated using the second method was at most 36 percent greater than; but usually within 20 percent of, the rate estimated using the first method, suggesting that intrinsic bioremediation was the dominant process that attenuated BTEX.

Introduction

As a management strategy, intrinsic remediation is gaining increased acceptance for use at sites contaminated with petroleum products. "Intrinsic remediation" refers to a management strategy that relies on natural attenuation mechanisms to remEDIATE contaminants dissolved in ground water and to control receptor exposure risks associated with contaminants in the subsurface. "Natural attenuation" refers to the actual physical, chemical, and biological processes that facilitate intrinsic remediation. Mechanisms for natural attenuation of fuel hydrocarbons include advection, dispersion, dilution from recharge, sorption, volatilization, and biodegradation. Of these processes, biodegradation is the only mechanism working to transform contaminants into innocuous by-products. Intrinsic bioremediation occurs when indigenous microorganisms work to bring about a reduction in the total mass of contamination in the subsurface without the addition of nutrients. Patterns and rates of intrinsic remediation can vary markedly from site to site depending upon governing physical and chemical processes.

At sites where petroleum products (e.g., gasoline, jet fuel, or diesel fuel) have been released and contribute to ground water contamination, many hydrocarbon compounds will partition into ground water; however, benzene, toluene, ethylbenzene, and xylene (BTEX) are generally the focus of site cleanup actions due to regulatory or
risk concerns. In addition, as shown by the work of Smith et al. (1981), Bruce et al. (1991), and Cline et al. (1991), despite making up no more than 20 percent of the mass of fuel mixtures, BTEX compounds account for 82 percent of the dissolved compounds that partition from fresh JP-4 jet fuel into ground water and at least 98 percent of the dissolved compounds that partition from fresh gasoline into ground water. Therefore, understanding and quantifying the natural attenuation of the BTEX compounds is often the focus of site remediation for risk reduction.

BTEX in ground water may be attenuated through the processes of dispersion, dilution from recharge, sorption, volatilization, and biodegradation. Abiotic chemical transformation of BTEX compounds in ground water is not an important process, and volatilization is not likely to remove significant amounts of dissolved volatiles, as shown by Chiang et al. (1989). In most hydrogeologic systems, biodegradation is the most important process that removes BTEX from ground water. BTEX biodegradation processes in ground water include aerobic respiration, denitrification, iron or manganese reduction, sulfate reduction, and methanogenesis (Jamison et al. 1975; Atlas 1981, 1984, and 1988; Young 1984; Bartha 1986; Wilson et al. 1986 and 1990; Barker et al. 1987; Baedecker et al. 1988; Lee 1988; Chiang et al. 1989; Cozzarelli et al. 1990; Leahy and Colewell 1990; Alvarez and Vogel 1991; Evans et al. 1991a and 1991b; Edwards et al. 1992; Edwards and Grbic-Galic 1992; Thierrin et al. 1993; Davis et al. 1994; Lovley et al. 1995). The patterns and rates of intrinsic remediation can vary markedly from site to site, depending upon the relative importance of the various processes.

Three lines of evidence have been used to document intrinsic bioremediation: (1) documented loss of contaminants at the field scale; (2) chemical and geochemical analytical data, including accumulation in ground water of the intermediates and by-products of anaerobic metabolism of BTEX, and the depletion of electron acceptor concentrations in ground water; and (3) microbiological laboratory data, such as the results of microcosm studies, plate counts, or cultures.

The first line of evidence requires a statistically significant historical data base resulting from several sampling efforts. Often, historical data are not available. Microbiological laboratory methods, such as microcosm studies, are expensive and time consuming. Moreover, biodegradation rates determined from individual microcosms may not be applicable throughout the plume (i.e., at the field scale). Therefore, under the time and budget constraints of most site investigations, chemical and geochemical data often play an important role in describing intrinsic bioremediation processes and quantifying intrinsic bioremediation rates, especially at the field scale.

To evaluate the contribution of natural attenuation to the management of risk associated with a release of petroleum hydrocarbons, it often is necessary to calculate site-specific biodegradation rates, particularly if ground water solute fate and transport modeling is part of the assessment. A site-specific biodegradation rate approximated using one or both of the methods presented herein can be useful for modeling, as many numerical and analytical model codes allow the use of a first-order contaminant decay constant. Models incorporating a site-specific decay rate in conjunction with dispersivity and retardation coefficients may be employed for the following: to predict the future extent and concentration of dissolved contaminant plumes by modeling the effects of advection, dispersion, sorption, and biodegradation; to assess the possible exposure of potential downgradient receptors to contaminant concentrations that exceed levels intended to be protective of human health and the environment; and to provide technical support for selection of the intrinsic remediation option as the best remedial alternative, as appropriate.

This paper compares two independent methods that may be used to approximate site-specific rates of biodegradation using monitoring well data and hydrologic properties of the aquifer. The first method involves the use of a tracer (1,3,5-trimethylbenzene) that sorbs more strongly than the BTEX compounds, is less volatile than the BTEX compounds, and is recalcitrant to anaerobic biodegradation at many sites. Attenuation of the tracer at a site can be attributed to dilution, sorption, or volatilization. Attenuation of BTEX compounds in excess of the attenuation of the tracer cannot be attributed to dilution, sorption, and/or volatilization. Changes in the concentration of the conservative tracer are used to correct for changes in the concentration of contaminants caused by the effects of dispersion, dilution from recharge, volatilization, and sorption. The corrected concentrations reveal the contribution of biodegradation, along with the effects of continuing source dissolution and any additional (nonlinear) sorption that may occur in the contaminant plume and the source area.

The second method, proposed by Buscheck and Alcantar (1995), assumes that the plume has evolved to the point that it is being destroyed by natural attenuation as rapidly as it is being produced by dissolution of soluble hydrocarbons from the fuel spill. In this situation, the plume has reached a dynamic steady-state equilibrium. Buscheck and Alcantar (1995) apply a one-dimensional analytical solution (Bear 1979) to interpret the steady-state configuration of the contaminant plume along a flow path. By removing the effects of dispersion and sorption, their analysis extracts an approximate rate of biodegradation that would be necessary to produce a steady-state plume of the configuration actually found at the site.

Neither method estimates a true biodegradation rate because they do not account for some of the processes affecting plume formation and attenuation. For example, unless the contaminant source has been removed completely, contaminants will continue to enter ground water due to direct dissolution into ground water or dissolution into infiltrating precipitation. Thus, without accounting for the mass entering the system, one is prevented from determining a true biodegradation rate,
or volatilization are removed. In addition, if background even if the effects of sorption, dispersion, dilution, and/or volatilization are removed. In addition, if background measurements of aquifer organic carbon are used to calculate retardation factors (assuming linear sorption) for the contaminants of interest, any nonlinear sorption or desorption within the plume and source area is not accounted for. Another factor preventing calculation of a true biodegradation rate is that the tracer method does not account for the loss of mass due to aerobic biodegradation, and the method of Buscheck and Alcantar (1995) does not account for mass lost to aerobic biodegradation at the upgradient margin of the plume.

The two methods were applied to data from the petroleum, oils, and lubricants (POL) bulk storage facility at Hill Air Force Base (AFB), Utah. Jet fuel (JP-4) was released into the soil and shallow ground water during fuel transfer operations at the POL facility.

**Site Description**

Hill AFB is located on a bench of the Wasatch Mountains on the edge of the Great Salt Lake Basin. Figure 1 shows the POL facility and the immediately adjacent area. Surface topography at the site slopes to the southwest. The shallow sediments underlying the site consist of light reddish-brown to dark gray, cohesive clayey silts to silty clays. This unit is 4- to 15-feet (1.2 to 4.6 m) thick and is underlain by poorly to moderately sorted, yellowish-brown to reddish-brown, silty fine-grained sands that coarsen downward into a 3- to 22-foot-thick (0.9 to 6.7 m) sequence of moderately sorted, medium- to coarse-grained sands. Underlying the sands is a sequence of competent, thinly interbedded clay to silty clay and fine- to very fine-grained, clayey sand and silt of unknown thickness. This sequence of interbedded clay and fine-grained sand and silt acts as an effective barrier to the downward migration of water and contaminants.

The water table aquifer is present in the medium- to coarse-grained sands described above, with the water table present between 5 and 20 feet (1.5 and 6.1 m) below ground surface. Ground water flow is to the southwest with an average gradient of 0.046 foot per mile (0.9 to 6.7 m) sequence of moderately sorted, medium- to coarse-grained sands. Underlying the sands is a sequence of competent, thinly interbedded clay to silty clay and fine- to very fine-grained, clayey sand and silt of unknown thickness. This sequence of interbedded clay and fine-grained sand and silt acts as an effective barrier to the downward migration of water and contaminants.

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The concentration of total organic carbon (TOC) measured in four soil samples ranged from 0.069 to 0.094 percent. Because of the low TOC concentration and clay mineral content observed in the shallow saturated zone, retardation of the BTEX compounds is not likely to be an important process affecting solute transport at this site.

General ambient ground water geochemistry at the site is conducive to intrinsic bioremediation. Total alkalinity at the site is fairly high, ranging from 349 to 959 milligrams per liter as calcium carbonate. This amount of alkalinity is sufficient to buffer potential changes in pH caused by biological production of carbon dioxide. Ground water pH at the POL facility ranges from 6.3 to 8.3 standard units. The average temperature of ground water is 64 degrees Fahrenheit (18 degrees Celsius). The oxidation/reduction potential (ORP) at the POL ranges from 274 mV to -190 mV. Areas at the site with low ORPs coincide with areas of BTEX contamination and areas with low concentrations of dissolved oxygen, nitrate, and sulfate and elevated concentrations of iron (II) and methane. This is strong evidence that dissolved BTEX at the site is subjected to a variety of biodegradation processes, including aerobic respiration, denitrification, iron reduction, sulfate reduction, and methanogenesis.

Figure 1 shows the approximate extent of mobile light nonaqueous phase liquid (LNAPL) at the site. This LNAPL is composed of weathered JP-4 released from the POL facility and is up to 4-feet (1.2 m) thick. Measured residual total BTEX concentrations in soil decrease rapidly outside the area of mobile LNAPL contamination. The highest dissolved BTEX concentration observed in ground water at the site was 21,000 μg/L. Figure 1 shows the distribution of total BTEX dissolved in ground water in July 1994. BTEX contamination is migrating to the southwest, in the direction of ground water flow. Figure 1 includes data collected from 12 monitoring wells in the source area north of 6th Street in January 1994. These

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**Figure 1. Total BTEX isopleth map.**

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188 • SUMMER 1996 GWMR
wells cover a small area relative to the extent of the dissolved plume, and these data represent the only data collected in this area in 1994. Data collection points shown on the map with no data represent locations sampled during other sampling events. Dissolved BTEX concentrations observed at these locations during previous and subsequent sampling events were taken into consideration when preparing Figure 1.

**Approximate Biodegradation Rate Constant Calculation Using a Conservative Tracer**

To determine approximate biodegradation rate constants, measured concentrations of dissolved BTEX were corrected for the effects of dispersion, dilution from recharge, volatilization, and sorption using a tracer. One tracer that has proved useful in some, but not all, ground water environments is trimethylbenzene (TMB). The three isomers of this compound (1,2,3-TMB, 1,2,4-TMB, and 1,3,5-TMB) have Henry’s law constants and soil sorption coefficients similar to (although somewhat higher than) those of the BTEX compounds (Table 1). Also, the TMB isomers generally are present in sufficient quantities in fuel mixtures to be readily detectable in ground water in contact with a fuel spill. Finally, they often are recalcitrant to biodegradation in the anaerobic portion of a plume; 1,3,5-TMB was found to be the most recalcitrant TMB isomer at the POL site and was used in the following calculations. Other compounds of potential use as conservative tracers are the tetramethylbenzene isomers, provided they are present in concentrations that are detectable throughout most of the plume.

The corrected concentration of a compound is the concentration of the compound that would be expected at one point (B) located downgradient from another point (A) after correcting for the effects of dispersion, dilution from recharge, volatilization, and sorption between points A and B. One method of calculating the corrected concentration (Wilson et al. 1994; Wiedemeier et al. 1995a) is given by:

\[ C_{B,corr} = C_B \left( \frac{TMB_A}{TMB_B} \right) \]

where
- \( C_{B,corr} \) = corrected concentration of compound of interest at Point B
- \( C_B \) = measured concentration of compound of interest at Point B
- \( TMB_A \) = measured concentration of trimethylbenzene at Point A
- \( TMB_B \) = measured concentration of trimethylbenzene at Point B.

Because trimethylbenzene is slightly more hydrophobic than BTEX, and therefore has a higher soil sorption coefficient, there likely is preferential sorption of TMB relative to the BTEX compounds. In addition, TMB may not be entirely recalcitrant under some anaerobic conditions, and it appears to degrade rapidly under aerobic conditions. The degree of recalcitrance of TMB is site specific, and the use of this compound as a tracer must be evaluated on a case-by-case basis. However, if any TMB mass is lost to the processes of biodegradation or preferential sorption, Equation 1 is conservative because the calculated mass losses and the attenuation rate constants calculated on the basis of those losses will be lower than the actual losses and attenuation rates.

Seven points along two flow paths parallel to the direction of ground water flow were chosen for comparison of corrected and observed BTEX concentrations (Figure 1). Table 2 presents BTEX and TMB data for these points. One flow path (points A, B, and C) coincides with the highest observed BTEX concentrations, and thus delineates the centerline of the dissolved BTEX plume. The other flow path (points D, E, F, G, and H) is located off the centerline of the plume, closer to the plume periphery. Table 2 also shows tracer-corrected BTEX concentrations. In conjunction with the aforementioned geochemical data, the calculations presented in this table imply that biodegradation of BTEX compounds is an important natural attenuation process at this site.

An approximate first-order biodegradation rate can be calculated if it can be shown that the corrected contaminant distribution approximates a distribution resulting from a first-order process. Figure 2 is a log-linear plot of tracer-corrected dissolved BTEX concentrations vs. downgradient travel time along the centerline of the ground water flow path (flow path ABC). Figure 3 is a log-linear plot of tracer-corrected dissolved BTEX concentrations vs. downgradient travel time along the flow path DEFGH. Figures 2 and 3 show that natural attenuation (largely due to biodegradation) along these flow paths approximates a first-order process.

First-order decay is described by the relationship:

\[ C = C_0 e^{-kt} \]

where
- \( C \) = contaminant concentration at time \( t \)
- \( C_0 \) = initial contaminant concentration
- \( k \) = first-order decay constant (total attenuation rate).

### Table 1: Soil Sorption Coefficients and Henry’s Law Constants for BTEX and TMB

<table>
<thead>
<tr>
<th>Compound</th>
<th>Soil Sorption Coefficient ( K_{oc} ) (L/Kg)</th>
<th>Henry’s Law Constant ( K_H ) (atm-m³/mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Benzene</td>
<td>79</td>
<td>0.005</td>
</tr>
<tr>
<td>Toluene</td>
<td>190</td>
<td>0.007</td>
</tr>
<tr>
<td>Ethylbenzene</td>
<td>468</td>
<td>0.007</td>
</tr>
<tr>
<td>Xylene</td>
<td>400</td>
<td>0.006</td>
</tr>
<tr>
<td>1,3,5-trimethylbenzene</td>
<td>676</td>
<td>0.006</td>
</tr>
<tr>
<td>Trimethylbenzenea</td>
<td>800</td>
<td>0.007</td>
</tr>
</tbody>
</table>

| *a* From Wiedemeier et al. (1995a) |
| *b* Average value for all isomers |

**SUMMER 1996 GWMR = 189**
Table 2
Benzene, Toluene, Ethylbenzene, Xylene, and TMB Concentrations

<table>
<thead>
<tr>
<th>Sample Location</th>
<th>Travel Distance (m)</th>
<th>Travel Time (days)</th>
<th>1,3,5-TMB Corrected Benzene (pg/L)</th>
<th>Toluene (pg/L)</th>
<th>Ethylbenzene (pg/L)</th>
<th>Xylene (pg/L)</th>
<th>TMB Corrected Total BTEX (pg/L)</th>
<th>Total TMB-Corrected BTEX (pg/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flow Path ABC (Plume Centerline)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>A</td>
<td>0</td>
<td>0</td>
<td>417</td>
<td>5600</td>
<td>5600</td>
<td>5870</td>
<td>5870</td>
<td>955</td>
</tr>
<tr>
<td>B</td>
<td>67</td>
<td>50</td>
<td>485</td>
<td>4260</td>
<td>4260</td>
<td>3910</td>
<td>3910</td>
<td>816</td>
</tr>
<tr>
<td>C</td>
<td>335</td>
<td>250</td>
<td>396</td>
<td>6</td>
<td>7</td>
<td>18</td>
<td>22</td>
<td>103</td>
</tr>
<tr>
<td>Flow Path DEFGH (Plume Periphery)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>D</td>
<td>0</td>
<td>0</td>
<td>650</td>
<td>941</td>
<td>941</td>
<td>2800</td>
<td>2800</td>
<td>505</td>
</tr>
<tr>
<td>E</td>
<td>143</td>
<td>106</td>
<td>125</td>
<td>458</td>
<td>458</td>
<td>10</td>
<td>52</td>
<td>454</td>
</tr>
<tr>
<td>F</td>
<td>311</td>
<td>232</td>
<td>144</td>
<td>7</td>
<td>6</td>
<td>10</td>
<td>9</td>
<td>23</td>
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<tr>
<td>G</td>
<td>341</td>
<td>254</td>
<td>78</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>4</td>
</tr>
<tr>
<td>H</td>
<td>416</td>
<td>310</td>
<td>2</td>
<td>0</td>
<td>c</td>
<td>3</td>
<td>c</td>
<td>2</td>
</tr>
</tbody>
</table>

Notes:
- TMB concentrations increased slightly between points A and B and points E and F. This suggests that 1,3,5-TMB is entirely recalcitrant between these points and that observed decreases in BTEX concentrations are due entirely to biodegradation.
- Benzene and ethylbenzene are more recalcitrant than 1,3,5-TMB between points D and E.
- 1,3,5-TMB is not recalcitrant between points G and H.

Substituting the TMB-corrected concentration, $C_{B,corr}$ at a downgradient point (B) for C in Equation 2, and the measured concentration $C_A$ at an upgradient point (A) for $C_d$, this relationship becomes:

$$C_{B,corr} = C_A e^{-\lambda t}$$  \hspace{1cm} (3)

where

- $C_{B,corr}$ = TMB-corrected contaminant concentration at time $t$ at downgradient point B
- $C_A$ = measured contaminant concentration at upgradient point A
- $\lambda$ = approximate first-order biodegradation constant.

Whereas the decay constant in Equation 2 describes the overall attenuation rate due to advection, dispersion, sorption, dilution from recharge, volatilization, and biodegradation, the decay constant in Equation 3 can be attributed largely to biological attenuation. However, as noted before, this is not a true biodegradation rate.

The travel time, $t$, between two points is given by:

$$t = \frac{x}{v_c}$$  \hspace{1cm} (4)

where

- $t$ = travel time between two points
- $x$ = distance between two points
- $v_c$ = retarded solute velocity (where applicable).

Use of the retarded solute velocity is appropriate where the organic carbon content of the aquifer matrix is sufficient to retard the migration of dissolved BTEX compounds. However, in some cases (such as at Hill AFB), retardation of the plume will be so limited that use of the advective velocity will approximate the retarded velocity. Background aquifer organic carbon concentrations may be used to calculate a retardation factor and a retarded solute velocity, but this may not accurately represent the solute velocity in the plume interior. However, as long as the retarded solute velocity is overestimated (i.e., retardation is underestimated),

190 • SUMMER 1996 GWMR
Table 3
Results of Exponential Regression Analyses

<table>
<thead>
<tr>
<th>Compound</th>
<th>Calculated y Intercept (b)</th>
<th>Slope (m)</th>
<th>R²</th>
<th>Approximated Biodegradation Rate Constant (λ, day⁻¹)⁰⁹</th>
<th>Calculated y Intercept (b)</th>
<th>Slope (m = k/v)</th>
<th>R²</th>
<th>Approximated Biodegradation Rate Constant (λ, day⁻¹)⁰⁹</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flow Path ABC (Plume Centerline)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Benzene</td>
<td>9249</td>
<td>-0.028</td>
<td>0.98</td>
<td>0.028</td>
<td>9430</td>
<td>-0.022</td>
<td>0.98</td>
<td>0.038</td>
</tr>
<tr>
<td>Toluene</td>
<td>8229</td>
<td>-0.023</td>
<td>0.98</td>
<td>0.023</td>
<td>8390</td>
<td>-0.018</td>
<td>0.98</td>
<td>0.031</td>
</tr>
<tr>
<td>Ethylbenzene</td>
<td>1075</td>
<td>-0.009</td>
<td>0.99</td>
<td>0.009</td>
<td>1095</td>
<td>-0.007</td>
<td>0.98</td>
<td>0.010</td>
</tr>
<tr>
<td>Xylene</td>
<td>9508</td>
<td>-0.006</td>
<td>0.99</td>
<td>0.006</td>
<td>9693</td>
<td>-0.005</td>
<td>0.99</td>
<td>0.008</td>
</tr>
<tr>
<td>Total BTEX</td>
<td>23,568</td>
<td>-0.010</td>
<td>0.99</td>
<td>0.010</td>
<td>24,028</td>
<td>-0.008</td>
<td>0.99</td>
<td>0.012</td>
</tr>
<tr>
<td>Flow Path DEFGH (Plume Periphery)</td>
<td></td>
<td></td>
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<td></td>
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<td></td>
<td></td>
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<tr>
<td>Benzene</td>
<td>1872</td>
<td>-0.025</td>
<td>0.92</td>
<td>0.025</td>
<td>1633</td>
<td>-0.016</td>
<td>0.89</td>
<td>0.027</td>
</tr>
<tr>
<td>Toluene</td>
<td>1696</td>
<td>-0.026</td>
<td>0.95</td>
<td>0.026</td>
<td>998</td>
<td>-0.017</td>
<td>0.78</td>
<td>0.029</td>
</tr>
<tr>
<td>Ethylbenzene</td>
<td>1243</td>
<td>-0.020</td>
<td>0.89</td>
<td>0.020</td>
<td>1122</td>
<td>-0.015</td>
<td>0.89</td>
<td>0.024</td>
</tr>
<tr>
<td>Xylene</td>
<td>17,126</td>
<td>-0.027</td>
<td>0.87</td>
<td>0.027</td>
<td>5547</td>
<td>-0.015</td>
<td>0.94</td>
<td>0.024</td>
</tr>
<tr>
<td>Total BTEX</td>
<td>17,431</td>
<td>-0.021</td>
<td>0.94</td>
<td>0.021</td>
<td>11,296</td>
<td>-0.015</td>
<td>0.99</td>
<td>0.025</td>
</tr>
</tbody>
</table>

Table 4
Representative First-Order Biodegradation Rate Constants

<table>
<thead>
<tr>
<th>Reference</th>
<th>Biological Decay Rate (day⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chapelle (1994)</td>
<td>0.01⁸</td>
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<tr>
<td>Wilson et al. (1994)</td>
<td>0.19⁸</td>
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<tr>
<td>Stauffer et al. (1994)</td>
<td>0.01⁹ to 0.019⁹⁰¹</td>
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<tr>
<td>MacIntyre et al. (1993)</td>
<td>0.01 to 0.02⁵¹</td>
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<tr>
<td>MacIntyre et al. (1993)</td>
<td>0.007 to 0.012⁶⁹¹</td>
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<tr>
<td>Wilson et al. (1993)</td>
<td>0.0024 to 0.067⁹¹</td>
</tr>
<tr>
<td>Buscheck et al. (1993)</td>
<td>0.001 to 0.01⁹¹</td>
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Approximate Biodegradation Rate Constant Calculation Using the Method of Buscheck and Alcantar (1995)

Buscheck and Alcantar (1995) derive a relationship that allows calculation of approximate first-order biodegradation rate constants. An important assumption that must be made when using this method is that the contaminant plume has reached a steady-state configuration. This method involves coupling the regression of contaminant concentration (plotted on a logarithmic scale) vs. distance downgradient (plotted on a linear scale) to an analytical solution for one-dimensional, steady-state, contaminant transport that includes advection, dispersion, sorption, and biodegradation. The effects of volatilization on the dissolved BTEX plume are assumed to be negligible. For a steady-state plume, the first-order decay rate is given by (Buscheck and Alcantar 1995):

\[ \lambda = \frac{v_c}{4\alpha_x} \left[ \left( 1 + 2\alpha_x \left( \frac{k}{v_x} \right) \right)^2 - 1 \right] \]  \hspace{1cm} (5)

where

- \( \lambda \) = first-order biodegradation rate constant (approximate)
- \( v_c \) = retarded contaminant velocity in the x-direction
- \( \alpha_x \) = dispersivity
- \( k/v_x \) = slope of line formed by making a log-linear plot of contaminant concentration vs. distance downgradient along flow path.

Figure 4 shows the dissolved BTEX plume in August 1993, July 1994, and September 1995. On the basis of the calculated advective velocity of the ground water,
the contaminant plume at Hill AFB should have migrated approximately 1450 feet (442 m) downgradient between August 1993 and July 1994, and 1700 feet (518 m) downgradient between July 1994 and September 1995. Figure 4 shows that the dissolved BTEX plume has not migrated a significant distance downgradient from the position it occupied in 1993, suggesting that the dissolved BTEX plume has reached a steady-state configuration.

Figures 5 and 6 present log-linear plots of measured BTEX concentrations vs. distance downgradient along flow paths ABC and DEFGH, respectively. Table 2 presents the data that were used to construct these plots. Exponential regressions were completed for each of the BTEX compounds and for total BTEX along each flow path (Figures 5 and 6). The value of k/v determined from the regression analysis is entered into Equation 5 and the approximate biodegradation rate constant, λ, is calculated. A longitudinal dispersivity of 50 feet (15 m) was estimated by using one-tenth of the distance between the spill source and the longitudinal centroid of the plume. This value of dispersivity is consistent with published literature values for these types of sediments. As noted previously, retardation is not important at the Hill AFB site, and the advective ground water velocity was used for these calculations. Table 3 presents the results of the regression analyses for this method.

First-order biodegradation rate constants approximated using the method of Buscheck and Alcantar (1995) range from 0.008 day⁻¹ (xylene) to 0.038 day⁻¹ (benzene) along flow path ABC, and range from 0.024 day⁻¹ (xylene and ethylbenzene) to 0.029 day⁻¹ (toluene) along flow path DEFGH.

One additional source of error in this calculation is the use of an assumed dispersivity. However, a sensitivity analysis shows that the model is only slightly sensitive to changes in dispersivity. Increasing dispersivity results in higher calculated attenuation rates. For example, if the dispersivity is assumed to be 1 foot (0.31 m), the calculated attenuation rate for total BTEX along flow path DEFGH is 0.020 day⁻¹, and if the dispersivity is taken as 100 feet (30.5 m), the attenuation rate along DEFGH is 0.030 day⁻¹. Along flow path ABC, the calculated attenuation rate for total BTEX is 0.011 day⁻¹ when dispersivity is 1 foot (0.31 m), and 0.013 day⁻¹ when dispersivity is 100 feet (30.5 m).

Conclusions

Table 3 shows that site-specific first-order biodegradation rate constants approximated using BTEX concentrations corrected using TMB as a tracer are remarkably consistent with the rate constants approximated using the method of Buscheck and Alcantar (1995). For all of the BTEX compounds, the rate constants calculated using a tracer are slightly less than those calculated using the method of Buscheck and Alcantar (1995). One reason for this is that tracer-corrected rate constants do not account for differential sorption or the fact that TMB is not entirely recalcitrant under anaerobic conditions. However, both of these factors make the TMB results more conservative (i.e., a slower rate constant). In both cases, the biodegradation rates are approximate because continuing source dissolution and nonlinear sorption are not accounted for. Once calculated, these site-specific rates can be employed in analytical or numerical ground water contaminant fate and transport models to predict plume behavior, evaluate potential risks to downgradient receptors, and make site management decisions.

Sensitivity analyses suggest that dispersion has a
small influence on the results of the calculations completed using the method of Buscheck and Alcantar (1995) for this site. The aquifer is overlain by a silty clay unit that should restrict recharge and limit volatilization. Furthermore, volatilization is generally not an important process acting to remove BTEX. For example, Chiang et al. (1989) used mass balance considerations to show that less than 5 percent of the mass of BTEX removed from ground water at a Michigan gas plant could be attributed to volatilization.

Approximate site-specific biodegradation rates calculated from the Hill AFB data are within the ranges published in recent literature (Table 4). Calculated degradation rates along the centerline of the Hill AFB plume (flow path ABC) generally are slower than those calculated along a flow path near the periphery of the plume (flow path DEFGH). This is likely due to the continuing introduction of dissolved BTEX in the vicinity of the source and the greater availability of electron acceptors near the margins of the BTEX plume.

Benzen degradation rates at Hill AFB were generally similar to the rates calculated for the other BTEX compounds. This is not consistent with experience at other sites where intrinsic bioremediation of BTEX has been documented. Barker et al. (1987), Cozzarelli et al. (1990 and 1994), and Wilson et al. (1993) found benzene to be more recalcitrant to biodegradation than the other BTEX compounds at the sites they studied. This suggests that pathways and patterns of BTEX biodegradation are site specific, as are the apparent rates of biodegradation. The fate of benzene may be controlled by the dominant terminal electron-accepting process. The dominant electron-accepting process at Hill AFB was sulfate reduction, accounting for 59 percent of the total expressed electron-accepting activity (Wiedemeier et al. 1995c). Methanogenesis accounted for 7 percent, iron reduction for 7 percent, denitrification for 21 percent, and aerobic respiration for 6 percent of the total expressed electron-accepting activity.

Where possible, both methods presented in this paper should be compared to evaluate the natural attenuation of BTEX in a plume. Use of this approach may not be appropriate at sites or in portions of a plume where the tracer is not biologically recalcitrant, or where the BTEX plume has not reached a steady-state configuration. If the plume has not reached a steady-state configuration, then tracer data can be used to provide a conservative estimate of biodegradation rates in the plume. If the tracer is not biologically recalcitrant and the plume is not stable, neither method is entirely appropriate. It may still be possible in this case to derive a rate constant using tracer-corrected data. However, the approach will likely produce unrealistically show rates and should not be used for transport calculations or modeling without a sensitivity analysis.

References


strictly anaerobic conditions. *Applied and Environmental Microbiology* 58, 2663–2666.


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