HYDROGEOLOGY AND WATER QUALITY NEAR A SOLID- AND HAZARDOUS-WASTE LANDFILL, NORTHWOOD, OHIO

By Jeffrey T. de Roche and Kevin J. Breen

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CONVERSION FACTORS AND ABBREVIATIONS

For the convenience of readers who may prefer to use metric (International System) units rather than the inch-pound units used in this report, values may be converted by using the following factors:

Multiply inch-pound unit	By	To obtain metric unit
inch (in.)	25.4	millimeter (mm)
foot (ft)	0.3048	meter (m)
foot of water (ft of water)	22.4	millimeter of mercury (mm Hg)
foot per day (ft/d)	0.3048	meter per day (m/d)
foot per day (ft/d)	0.000353	centimeter per second (cm/s)
mile (mi)	1.609	kilometer (km)
foot per mile (ft/mi)	0.1894	meter per kilometer (m/km)
foot squared per day (ft ² /d)	0.0929	meter squared per day (m ² /d)
foot squared per day (ft²/d)	0.01075	centimeter squared per second (cm ² /s)
square mile (mi ²)	2.590	square kilometer (km ²)
cubic foot per second (ft ³ /s)	28.32	liter per second (L/s)
gallon per minute per foot per foot ([(gal/min)/ft]/ft)	40.74	liter per minute per meter per meter ([(L/min)/m]/m)

Concentrations of chemical constituents and temperatures of air and waters are given in metric units. Concentration is given in milligrams per liter (mg/L) or micrograms per liter (μ g/L). Milligrams per liter is a unit expressing the concentration of chemical constituents in solution as weight (milligrams) of solute per unit volume (liter) of solution (water). One thousand micrograms per liter is equivalent to one milligram per liter. For concentrations of dissolved solids less than 7,000 mg/L, the numerical value is, for practical purposes, the same as for concentrations in parts per million.

Water and air temperature is given in degrees Celsius (°C), which can be converted to degrees Fahrenheit (°F) by the following equation:

$$^{\circ}F = 1.8(^{\circ}C) + 32$$

<u>Sea level</u>: In this report "sea level" refers to the National Geodetic Vertical Datum of 1929 (NGVD of 1929)—a geodetic datum derived from a general adjustment of the first-order level nets of both the United States and Canada, formerly called "Sea Level Datum of 1929."

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ABSTRACT

Hydrogeology and water quality of ground water and selected streams were evaluated near a landfill in northwestern Ohio. The landfill is used for codisposal of solid and hazardous waste. Water-level and geologic data were collected from 36 wells and 3 surface-water sites during the period November 1983 to November 1985. Waterquality samples were collected from 18 wells and 3 surface-water sites during this same period.

The primary aquifers in the area are the Greenfield Dolomite and underlying Lockport Dolomite of Silurian age. These bedrock carbonates are overlain by two clay tills of Wisconsinan age. The tills are capped by a glacial lake clay. The tills generally are saturated, but do not yield sufficient water to be considered an aquifer. Two wells in the study area yield water, in part, from discontinuous deposits of outwash sand and gravel at the lower till-bedrock interface.

Regional ground-water flow is from southwest to northeast; local flow is influenced by a ground-water mound centered under the northernmost cells of the landfill. Water levels in wells penetrating refuse within the landfill and the presence of leachate seeps indicate that the refuse is saturated. Head relations among the landfill, till, and dolomite aquifer indicate a vertical component of flow downward from the landfill to the dolomite aquifer. Water levels near the landfill fluctuate as much as 14 feet per year, in contrast to fluctuations of less than 3 feet per year in wells upgradient of the landfill.

Ground waters from wells completed in the dolomite aquifer and glacial till were found to have major-ion concentrations controlled, in large part, by reaction with calcite, dolomite, and other minerals in the aquifers. Only minor departures from equilibrium mineral saturation were noted for ground water, except in wells affected by cement/grout contamination. Molal ratios of calcium:magnesium in ground water suggest a similar chemical evolution of waters throughout the dolomite aquifer in the study area. Stableisotope ratios of oxygen and hydrogen indicate the source of water in the till unit and dolomite aquifer is atmospheric precipitation.

Elevated levels of total dissolved solids, boron, ammonia, and iron in the leachate and in wells downgradient of the landfill may indicate mixing of ground water with leachate. Oxygen and hydrogen stable-isotope ratios were used to differentiate waters from the glacial till and dolomite aquifer. Isotope ratios also show a shift off the local mixing line for leachate and for a well just downgradient from the landfill. This shift to heavier values of δD in the well water may be indicative of leachate mixing with ground water.

The effect of this mixing denoted by hydrologic, isotopic, and chemical-quality data is limited mostly to elevated levels of the common ions. Analysis did not indicate significant levels of toxic metals or organic contaminants except phenol, which was present at concentrations of from 1 to 5 micrograms per liter in six wells. Analysis of water-quality data from nearby streams suggests that surface leaching from the landfill does not significantly affect stream-water quality, but may contribute to higher levels of trace metals in the streambed sediments.

INTRODUCTION

Ground water is an important resource for rural residents near Northwood, Ohio. The presence of a solid- and hazardous-waste landfill near Northwood and a lack of current ground-water data for the surrounding area created a need for a study of the hydrogeology and water quality. This study was conducted in cooperation with the City of Northwood, Ohio, and presents findings from data collected from November 1983 through November 1985.

Purpose and Scope

The purpose of this report is to describe the hydrogeology and the chemical quality of ground water and surface water near the landfill. The evaluation was made by (1) review of available hydrogeologic and chemical-quality data; (2) mapping of the glacial overburden and underlying dolomite aquifer from well logs and geologic borings; (3) measurement of hydraulic head in the glacial overburden and dolomite aquifer; (4) collection and analysis of water-quality data from the landfill, glacial overburden, and dolomite aquifer; and (5) collection and analysis of water- and sediment-quality data from local streams.

Location and Setting

The project area (fig. 1) is located in Wood County in northwestern Ohio and encompasses an area of approximately 10 mi² (square miles). The area includes parts of the City of Northwood (population 6,000) and the Village of Walbridge (population 3,000). Land use in the area is a mixture of light and heavy industry, transportation, housing, and open areas used for agriculture.

The climate is temperate. Average annual temperature for the 1951-80 period was 10.8 °C (51.5 °F). For the same period, monthly average precipitation ranged from a

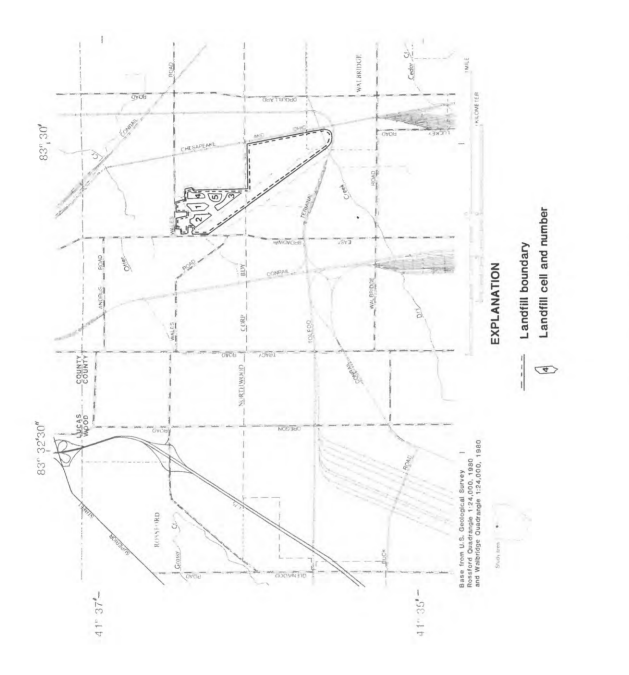


Figure 1.--Location of study area.

high of 3.50 in. (inches) for June to a low of 1.81 in. for February. The 30-year average annual precipitation was 32.29 in. (National Oceanic and Atmospheric Administration, 1982).

Topography in the area is flat and nearly featureless. Drainage is provided by roadside ditches, Otter Creek, and Dry Creek. Most soils belong to the Toledo soil association (U.S. Department of Agriculture, 1966) and are developed in lake-deposited silts and clays.

Description of the Landfill

The landfill site (fig. 1) covers approximately 160 acres and is primarily used for disposal of municipal and commercial refuse. The northern part of the landfill is divided into five separate excavations, or cells. Before construction of the landfill, a railroad switching station known as Outer Yard occupied much of the site. Most of the track has been removed since landfilling started on the original 20-acre site in 1972.

Records indicate that cells 1 through 4 use the local natural clay deposit as liner material and contain primarily municipal and commercial refuse (Waste Management, Inc., written commun., no date). However, a Resource Conservation and Recovery Act (RCRA), Part B application indicates that past landfilling included disposal of wastes now defined as hazardous (Dames and Moore, 1983). The excavation depth below grade for pits 1 through 4 is reported to be 35 to 40 feet (John Barbush, Waste Management, Inc., oral commun., 1984).

Cell 5, which was permitted to receive hazardous waste, also uses the local clay till as a liner. Depth of fill below grade is 35 feet. Site delivery records for 1982 and 1983 indicate the hazardous-waste cell contains primarily heavy-metal sludges, wastewatertreatment sludges from electroplating operations, and air-pollution-control sludges or dust. Records also show soluble cyanide salts, DDT, toluene, 1,1,1,- trichloroethane, and 2,4-D were deposited in the hazardous-waste cell.

Cells 1 through 5 have all been filled and completed and are covered with clay caps. Cells 1, 2, and 4 are equipped with methane-venting wells that may be used for leachate observation. Cell 5 is equipped with a leachate collection and monitoring system.

Landfilling of solid waste expanded into the southern section of the site during the early 1980's. Currently, disposal in the southern section is limited to solid waste; no hazardous wastes are permitted. The investigation centers primarily on the northern section of the site because cells have been in place longer and because of the nature of the waste in cell 5.

Previous Investigations

Most publications on the hydrogeology of Wood County are regional or countywide in scope and provide minimal information on water quality. The regional subsurface geology has been investigated and summarized by Kahle and Floyd (1972) and Janssens (1977). Reports relating geology to land-use planning for Wood County have been done by Forsyth (1968) and Nielsen (1977).

Studies by the Ohio Department of Natural Resources (1970), Norris and Fidler (1969, 1971a, 1971b), and Norris (1974) discuss the regional hydrogeology of northwestern Ohio. A report by Glaze (1972) provides information on the hydrogeology of northern Wood County, and a subsequent study by Paulson (1981) reviews the ground-water resources of Wood County. A recent synopsis of ground-water resources in northwestern Ohio and southern Michigan by the Toledo Metropolitan Area Council of Governments (1982) provides information on the hydrogeology of selected northwestern Ohio counties.

Site-specific reports on the landfill area have been prepared by Bowser-Morner Testing Laboratories (1980) and Dames and Moore (1983, 1984). These reports address the area's geology and ground-water occurrence but provide only minimal information on ground-water quality.

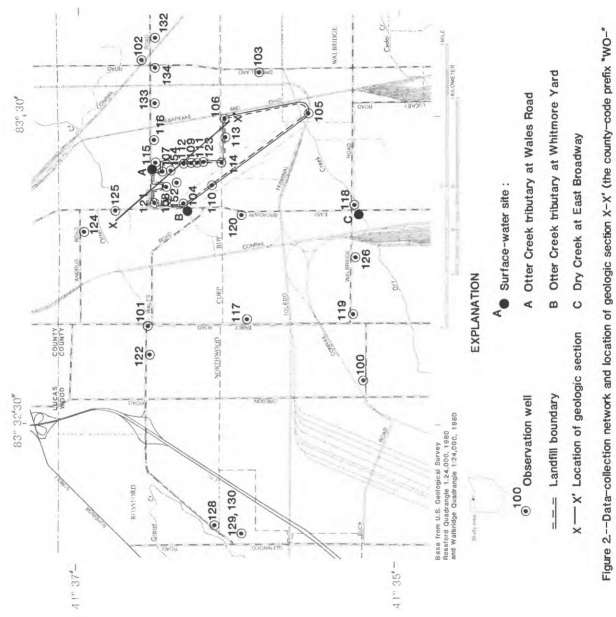
Methods of Study

The data-collection network (fig. 2, table 1) consisted of 36 wells and 3 surfacewater sites. Two wells were equipped with hourly water-level recorders, and a continuous precipitation recorder was installed on site. Water-level measurements were obtained bimonthly from the well network. Additional hydrologic and geologic information was obtained from logs and laboratory tests of 27 borings (Bowser-Morner Testing Laboratories, 1980; Dames and Moore, 1983, 1984).

Of the 36 wells, 22 are domestic or commercial wells that are cased into bedrock and are open hole below. Ten are specially constructed monitoring wells cased into bedrock, screened and sandpacked in the upper zones of bedrock, and grouted with a cement or cement/bentonite mixture. Two piezometers (113 and 123)¹ are cased, grouted, and screened and sandpacked in the overlying till, and two wells (152 and 154) are finished within the northern cells of the solid-waste landfill to vent methane gas.

Chemical-quality data were collected from April 4, 1984, through July 11, 1985, from three surface-water sites and 17 ground-water wells. Water samples were analyzed

¹For the sake of simplicity, the county prefix "WO-" has been omitted from local well numbers in the text and many of the illustrations in this report. Local numbers are given in full in the tables.



has been omitted from well numbers).

Table 1.-Records of selected wells near Northwood, Ohio

[Producing zone: D, dolomite; T, till. Casing type: S, steel; P, PVC. Dashes indicate data not available or not applicable.]

Elevation Well Producing Year Casing Casing of land Depth surface of well number Latitude Longitude zone comtype diameter (inches) (feet) (feet) pleted 41° 35' 12" 83° 32' 09" 4.25 139 WO-100 D 1977 S 621 41°36'31" 83° 31 ' 42" WO-101 D S 6.00 617 250 41°36'35" 83° 29' 34" 1977 WO-102 D S 6.00 614 149 41° 35' 51" 83° 29' 39" WO-103 D 1972 S 6.25 615 250 WO-104 41°36'20" 83° 30' 41" D 1983 S 7.00 617 155 WO-105 41° 35' 32" 83° 29' 58" D 1983 P 4.00 618 100 WO-106 41° 36' 04" 83° 30'01" 1983 P 4.00 94 D 616 41° 36' 26" 4.00 83° 30 "29" 1983 P 123 WO-107 D 619 41°36'25" 83° 30 ' 35" WO-108 1983 P 100 D 4.00 610 41°36'16" 83° 30 '23" WO-109 D 1983 P 4.00 619 109 WO-110 41° 36' 08" 83°30'34" D 1983 P 4.00 618 120 41°36'14" WO-111 83° 30 ' 23" D 1983 Ρ 4.00 617 110 41°36'18" WO-112 83° 30' 23" D 1983 P 4.00 613 109 WO-113 41° 36' 03" 83° 30' 10" 1980 P 1.25 Т 616 49 41°36'05" WO-114 83° 30 ' 23" 4.25 D 1974 S 617 200 P 41°36'30" 83° 30' 23" D 1983 2.00 613 83 WO-115 41°36'30" 83° 30' 12" 1983 P 2.00 90 WO-116 D 615 WO-117 41°36"35 83° 31 ' 39" DT 1946 S 4.50 619 102 WO-118 WO-119 41° 35' 15" 83° 30' 43" 6.00 1974 618 D S 160 41° 35' 15" 83° 31 ' 37" DT 1958 S 4.25 621 132 41° 35' 57" 83° 30 ' 47" WO-120 D 1946 S 4.00 616 84 41° 36' 29" WO-121 83° 30 ' 44" D 6.00 188 ---S 616 WO-122 41°36'31" 83° 31 '58" D S 8.00 617 330 41°36'11" WO-123 83° 30' 22" T 1984 P 2.00 616 59 WO-124 41°36'55" 83° 30' 58" D S 5.50 616 5.50 WO-125 41°36'44" 83° 30' 46" D S 150 616 -41° 35' 15" 83° 31'09" WO-126 _ 4.00 130 D S 622 WO-128 WO-129 41° 36' 06" 83° 33' 21" D _ S 6.00 615 132 41° 35 56" 83° 33' 24" D _ S 6.25 615 130 41° 35' 56" 83° 33' 24" WO-130 D _ S 4.50 615 90 41°35'40" 83° 32' 22" WO-131 D -S 8.00 620 620 41° 36' 29" 83° 29' 22" WO-132 D S 6.00 208 613 41°36'29" 83° 29' 54" WO-133 1958 4.00 D S 615 145 41° 36' 29" 41° 36' 23" 6.00 83°29'37" WO-134 D S 615 109 WO-1521 83° 30' 30" 1983 -P 655 45 41º 36'23" 83° 30' 30" WO-1541 P -1983 8.00 655 49

¹Methane-venting wells

for water characteristics, major ions, trace constituents, nutrients, base/ neutral- and acidextractable organic compounds, volatile organic compounds, and stable isotopes of oxygen and hydrogen. In addition, streambed materials were analyzed for trace metals and base/neutral- and acid-extractable organic compounds. Blanks, spikes, and duplicate samples were submitted to the laboratory for quality assurance and quality control. Water and sediment samples were analyzed by U.S. Geological Survey National Water Quality Laboratory. Isotope samples were analyzed by the U.S. Geological Survey's Isotope Fractionation group in Reston, Va.

Sampling procedures for observation wells were designed to obtain a representative sample from the aquifer and to minimize the introduction of any foreign substance that might affect ambient or native water quality. All observation wells were pumped until a minimum of three casing volumes was purged from the well. Domestic wells were pumped until the volume of the pressure tank plus three casing volumes had been purged. During purging, pH, conductance, temperature, and dissolved oxygen were monitored by use of a flow chamber. After purging the wells, samples were collected when readings became stable.

Most of the monitoring wells within the landfill perimeter were installed by the landfill's contractor and are of PVC construction equipped with dedicated submersible pumps and PVC lift lines. Wells outside of the landfill generally were domestic wells cased with steel and open hole into the dolomite aquifer. In wells not equipped with a dedicated pump, a Johnson-Keck² SP-81 submersible pump with interchangeable EPDM and Viton stators was used for sampling. When sampling for inorganic constituents, 10 percent acetone solution and distilled water were used to clean the pump. The acetone solution was pumped through the pump and sample lines, and was followed by distilled water to flush the acetone. The power line, pump housing, and exterior of the sample line also were cleaned before being lowered into the well.

When sampling for base/neutral- and acid-extractable organic compounds, a solution of 75 percent hexane and 25 percent alcohol was used to clean the interior and exterior parts of the sampling pump. Pump materials for organic sampling were limited to Teflon, Viton, and stainless steel. For all observation-well sampling, the pump was cleaned before it was lowered into the well and immediately after it was removed. For sampling of volatile organic compounds, a Teflon bladder pump was used. A detergent wash and distilled-water rinse were used to clean the pump before and after use.

Bed material was obtained from local streams and analyzed for organic constituents and trace metals. To increase the recovery of fine sediments, pools and low-velocity reaches of the streams were chosen as sampling sites. Streambed penetration during sampling was generally 4 in. or less.

²Use of firm, brand, or trade names in this report is for identification purposes only and does not constitute endorsement by the U.S. Geological Survey.

Bed-material samples for trace-metal analysis were collected with a U.S. Geological Survey RBM80 sampler or a plastic scoop. All sampling equipment and containers were cleaned with a 10 percent nitric acid solution, followed by distilled water, and then by a native-water rinse. Samples were separated after freeze drying, and the less-than-25- μ m (micrometer) fraction was analyzed.

Bed-material samples for organic constituents were collected with an RBM80 sampler, a stainless-steel scoop, and stainless-steel sieves. All sampling equipment was cleaned with a 75 percent hexane and 25 percent alcohol solution, followed by distilled water and native-water rinses. Samples were wet sieved in the field through 90- and 63- μ m sieves, and the less-than-63- μ m fraction (medium silts and smaller) was analyzed.

Analysis of organic constituents in water and sediments was done by gas chromatography/mass spectrometry. Samples were analyzed quantitatively for base/neutral- and acid-extractable organic compounds and qualitatively for all other methylene-chlorideextractable organics. All samples were analyzed at the U.S. Geological Survey National Water-Quality Laboratory.

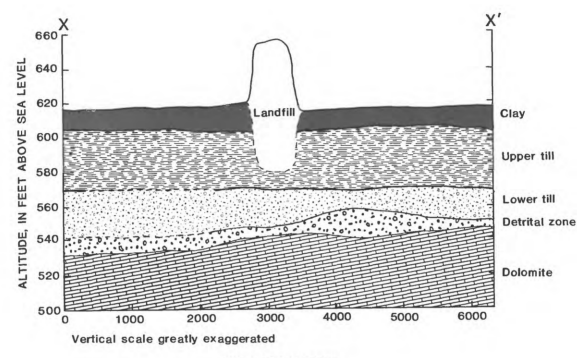
Acknowledgments

The authors are grateful for the information and assistance provided by Wood County Health Department, Toledo Metropolitan Area Council of Governments, the City of Northwood, and Waste Management, Inc. Appreciation is expressed also to Bihn Well Drilling and to the private landowners and homeowners who offered their wells for water-level measurement and chemical-quality sampling. In the interpretation of isotopic results, the assistance of Carol Kendall, U.S. Geological Survey, Reston, Va., also is gratefully acknowledged.

HYDROGEOLOGY

Unconsolidated Deposits

The unconsolidated deposits in the study area (fig. 3) consist of a glacial-lake clay deposit underlain by two tills of Wisconsinan age. The upper till, into which the landfill cells are excavated, is rich in clay, whereas the lower till contains a relatively high percentage of sand and pebbles (Forsyth, 1968). Underlying the lower till is a detrital or broken-rock zone composed of sand, gravel, boulders, rock fragments, and clay. This characteristic layering of the unconsolidated deposits has been correlated over much of northern Ohio by Forsyth (1960).



DISTANCE, IN FEET

EXPLANATION

Geologic contact, dashed where approximately located.

Figure 3.-Generalized geologic section X-X'.

Examination of test borings (Bowser-Morner Testing Laboratories, 1980; Dames and Moore, 1983, 1984) shows the glacial lake deposit is a dark-brown silty clay containing traces of sand and gravel that ranges from 6 to 15 ft in thickness. This horizon is oxidized and mottled, and exhibits vertical fractures that extend into the underlying upper till. A seasonal water table is present in this layer.

The upper till, formally called the Hayesville Till (White, 1961), is an olive-gray clay containing silt, sand, and traces of gravel. This horizon ranges from 35 to 42 ft in thickness and contains occasional thin seams (0 to 2 ft) of sand or sand and gravel. Sand or gravel zones generally are saturated; however, yields are not sufficient to constitute a usable water supply.

Values of vertical hydraulic conductivity from triaxial chamber tests (Bowser Morner Testing Laboratories, 1980) performed on two samples were 8.5×10^{-5} ft/d (feet per day). Examination of two core samples of upper till taken on site (Dames and Moore, 1984) revealed vertical fractures at a depth of 45 ft, which is within 10 ft of the upper-till/ lower-till contact (R. Pavey, Ohio Department of Natural Resources, Division of Geological Survey, oral commun., 1986). Because of the high resistance of low-K soils to ground-water flow, the effects of nonuniformities such as stratified permeable sand layers, fractures, and fissures become very important, as these provide the low-resistance path to leachate flow (Mixon and others, 1987). During cell excavation (fig. 3) the upper till is removed to within 10 to 15 ft of the contact with the lower till.

The lower or Millbrook Till (White, 1961) is a 15- to 25-ft-thick section of gray clay with silt, sand, and gravel. In situ and laboratory testing (Bowser-Morner Testing Laboratories, 1980; Dames and Moore, 1983) indicates that vertical hydraulic conductivity of the lower till is in the range of 8.5×10^{-5} to 1.2×10^{-4} ft/d. The contact between the upper and lower tills is characterized by sand and gravel zones, mottling, and increased water content. Zones of sand and (or) gravel are found throughout the entire thickness of the lower till and range in thickness from 0.08 to 1.7 ft (Dames and Moore, 1984). The lower till and associated sand zones generally are saturated, but do not yield sufficient water to be considered an aquifer. The composition of the sand zones is variable and ranges from silty/clayey sand to clean sand in composition and from fine to coarse in texture. There are no data to suggest that the zones of sand and gravel are laterally continuous in the sense of a readily identifiable bed or horizon; however, their prevalence and relatively higher permeability than the surrounding clay till may allow these sand zones to act as pathways for lateral and vertical flow and recharge to the underlying dolomite aquifer.

Samples of upper and lower clay till from cores collected during drilling of well 115 were analyzed for mineralogic composition by X-ray diffraction. The samples, which represent 39- to 40.5-ft and 59- to 60.5-ft depths, were found similar in mineralogy and

consisted of muscovite, chlorite, quartz, calcite, ankeritic dolomite, and feldspar. The tills produce carbon dioxide gas when reacted with hydrochloric acid, and, therefore, may be termed calcareous clay tills.

The contact between the lower till and the underlying dolomite aquifer is characterized by discontinuous deposits of outwash sand and gravel and rock fragments. These deposits, which typically range in thickness from 2 to 29 ft, commonly are found in preglacial drainage valleys (Paulson, 1981), and, where saturated, are a good source of water for small domestic supplies (Forsyth, 1968). This zone commonly is noted by local drillers and identified as broken rock or gravel. The maximum thickness of this deposit noted in the study area is 18 ft in boring CB-23 (Dames and Moore, 1984). The deposit consists of clay, sand, gravel, and rock fragments and is located in a preglacial drainage valley just east of the landfill.

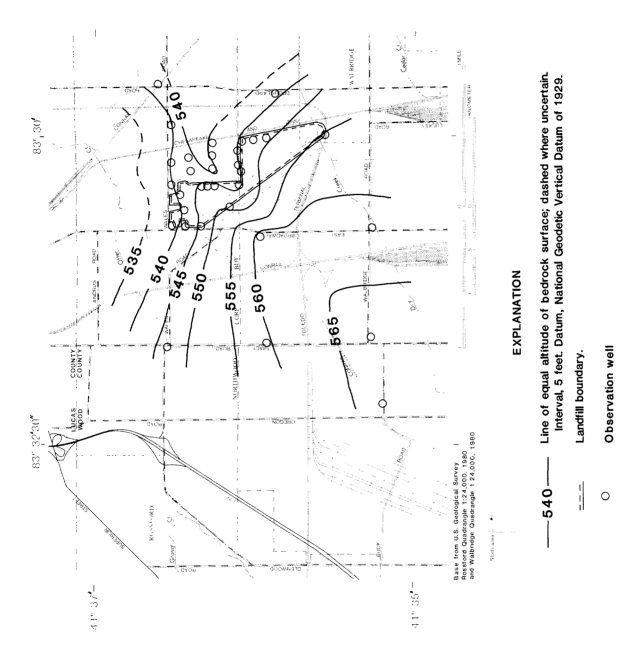
Bedrock

The study area is underlain by the Greenfield Dolomite of Silurian age. Underlying the Greenfield and also of Silurian age is the Lockport Dolomite. The landfill lies near the gradational contact between the Greenfield Dolomite and the Lockport Dolomite.

The Greenfield Dolomite is considered to be a biohermal and biostromal facies of the Lockport (Janssens, 1977). The Greenfield Dolomite is characterized as being brown, microcrystalline, medium-bedded, and stromatolitic, with solution channels, bituminous partings, and stylolites. The Lockport is a gray to blue-gray vuggy dolomite. Although the dolomites underlying the landfill have characteristics of both the Lockport and Greenfield, they generally are assigned to the Greenfield Dolomite based upon the presence of stromatolites and stylolites. Because of the facies changes, the contact between the Lockport and Greenfield is considered to be gradational, and the units cannot be differentiated easily in the subsurface (Janssens, 1977). Wells within the study area most likely produce water from both units.

The complex geology is a result of the Silurian reef complex located in northern and central Wood County (Mesolella, 1978). The Silurian reef complex is greatly variable in thickness and exhibits many changes in facies between adjacent strata, which reflects deposition in an environment that fluctuated between supratidal and intertidal. Evidence of repeated subaerial exposure—an important factor influencing the porosity of carbonate rocks—has been noted in local quarries where the Greenfield and Lockport Dolomites are exposed (Kahle and Floyd, 1972; Kahle, 1978).

The bedrock surface in the vicinity of the landfill (fig. 4) is that of a ridge and a valley trending northeast. Joints and fractures in the carbonate rocks are avenues for carbonate solution and have influenced the present position of bedrock valleys.





Rowland and Kunkle (1970) reported that northeast-trending joints are particularly permeable. Borings near the landfill indicate the presence of joints and fractures (both sealed and open), solution vugs, and cavities or voids.

Permeability in the Greenfield Dolomite is predominantly the result of secondary openings—joints, fractures, and solution cavities. The high degree of aquifer heterogeneity caused by differing depositional environments, subaerial erosion, and formation of secondary openings produces a wide range of water-yielding characteristics. Specific capacity per foot of aquifer penetration ranges from 0.0028 to 0.3103 [(gal/min)/ft]/ft (gallons per minute per foot per foot) for domestic and small commercial wells in the study area. This range of values compares with the median value for "Area 3" of 0.0038 [(gal/min)/ft]/ft (Ohio Department of Natural Resources, 1970). Area 3 includes the entire eastern half of Wood County, including the Northwood area. Wells having the highest specific capacities either produce water from the lower till-bedrock interface, or, because of their greater depth, may encounter a greater number of fractures.

Ground-Water Levels and Flow

Assuming isotropic conditions within an aquifer, ground water generally moves from areas of high hydraulic head to areas of low hydraulic head along paths perpendicular to potentiometric contours. Because flow in the dolomite aquifer is by means of secondary openings, conditions within the aquifer are not isotropic and, therefore, differ with direction.

If fracture spacing is sufficiently dense, flow in fractured media may resemble flow in a granular porous media. If fracture spacings are irregular in a given direction, the media will exhibit spatial trends in heterogeneity (Freeze and Cherry, 1979). Thirteen out of 15 bedrock cores indicate the presence of highly fractured and vuggy zones in the upper bedrock at depths of 70 to 80 ft below land surface. Deeper cores indicate that highly fractured zones also are present from 80 to 100 ft deep.

Most monitoring wells near the landfill are screened in these highly fractured zones. Reported transmissivities (H. B. Eagon, Jr., Eagon and Associates, Inc., oral commun., November 19, 1985) range from $0.83 \text{ ft}^2/\text{d}$ (feet squared per day) to $14.34 \text{ ft}^2/\text{d}$, with a mean value of approximately 5 ft²/d. These values were obtained from eight wells that penetrate only the upper 20 to 30 ft of the dolomite aquifer and are not corrected for partial penetration. Core logs indicate that the fractured zone is widespread and may react like a porous medium; however, the range in transmissivities would indicate that conditions are heterogeneous, and that certain areas can transmit water at a higher rate than others.

In the study area, regional ground-water flow in the confined dolomite aquifer is primarily from southwest to northeast, as shown by the potentiometric map. The potentiometric map (fig. 5) may be viewed as showing the sum of many possible directions of ground-water flow through the various interconnected joints and solution channels.

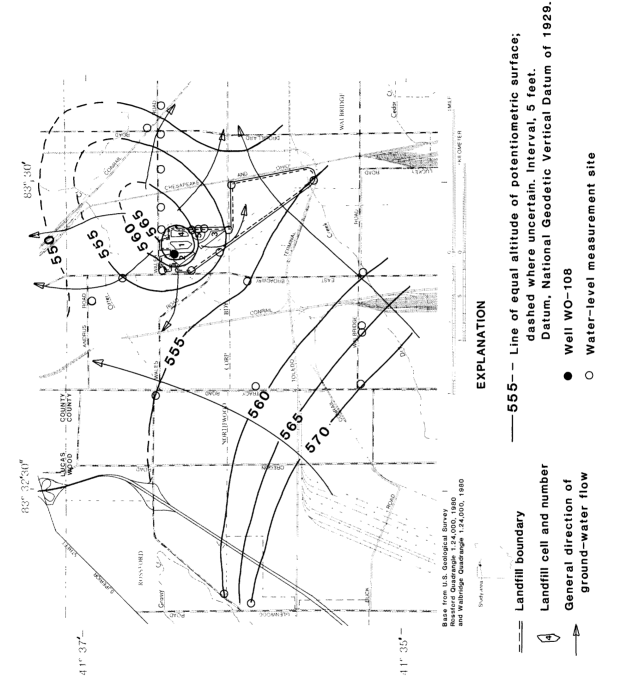
The dominant local feature on the potentiometric map (fig. 5) is the ground-water mound centered under the northern section of the landfill. The mound was first noted in March 1984. Subsequent water-level measurements confirmed that it is persistent and fluctuates seasonally. The mound appears to extend asymmetrically in the direction of regional flow in the dolomite aquifer. The geometry of the surface suggests that the area of recharge is approximately centered on landfill cell 1.

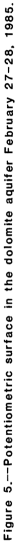
The potentiometric map shows aquifer reaction to recharge by snowmelt and by 1.26 in. of precipitation that occurred during the preceding 5-day period. Weather conditions, combined with constricted drainage in Otter Creek where it enters a culvert under Wales Road, caused surface flooding in the area. The restricted surface waters rose to an altitude of approximately 613.3 ft, as indicated by high-water marks on the surface casings of wells 112 and 115. Flooding was localized and confined to overtopping of natural and manmade drainages near the landfill as opposed to a large area of inundation. The surface waters receded over a 12-day period.

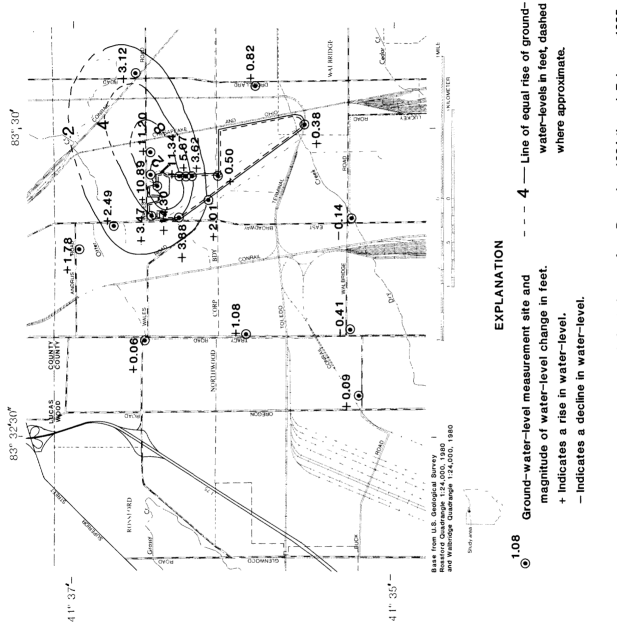
As a result of flooding, surface waters rose to 0.7 ft above the top of the protective casing of well 108. Similar flooding occurred in April 1985, and well 108 again was submerged. The flooding of well 108 may have contributed to the ground-water mound; however, the magnitude of the effect is not known.

Ground-water levels near the northernmost landfill cells fluctuate as much as 14 ft per year (figs. 6A and 6B), in contrast to fluctuations of less than 3 ft per year in wells upgradient from the landfill. Reaction to precipitation and snowmelt is much more pronounced near the landfill, as shown by water-level increases from December 1984 to February 1985 (fig. 6A). Water-level fluctuations (fig. 6B) during the period February to March 1985 show the general decrease in water levels as the mound dissipates. The high permeability of landfilled wastes, the proximity of the cell base to the fractured till zone near the upper/lower till contact, and the fractured nature of the dolomite surface are factors likely contributing to the large water-level fluctuations near the center of the northernmost landfill cells.

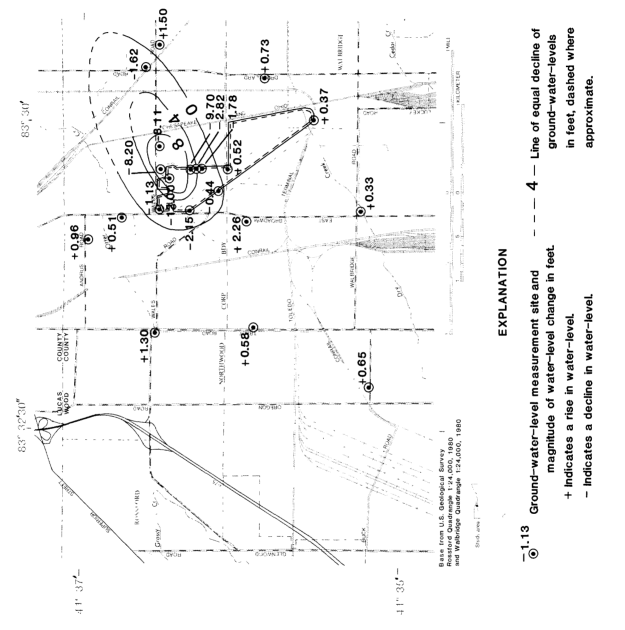
Figure 7 shows the relation between precipitation and water levels in wells completed in the till and underlying confined aquifer. Well 113 is screened, sand packed (25 to 49 ft below land surface), and grouted in the upper-till/lower-till interface, whereas well 121 is cased in the dolomite and is open hole below. Well HY-2 is located approximately 29 miles southeast of the study area, is cased in the dolomite aquifer, and is open hole below. All wells were equipped with water-level gages and digital recorders that recorded water level on an hourly basis.













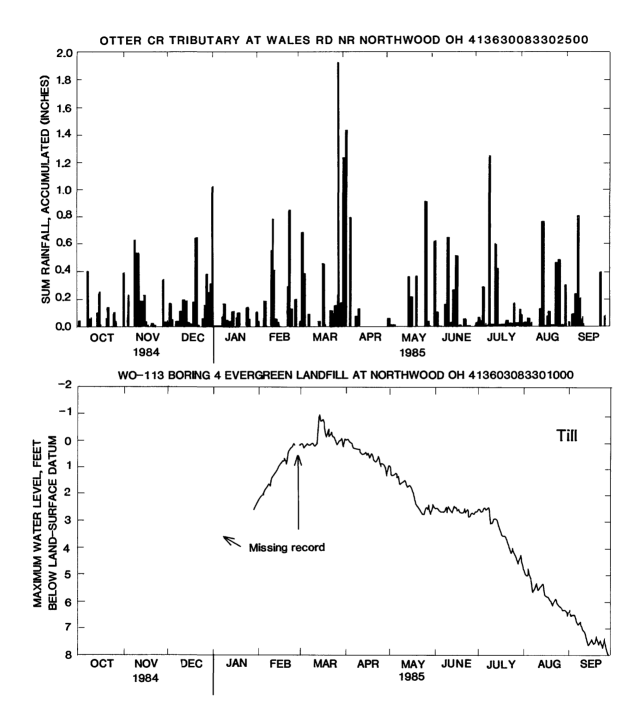


Figure 7.-- Hydrographs showing relation between water levels in wells WO-113 (till), WO-121 (carbonate), and HY-2 (carbonate) and daily precipitation for 1985 water year.

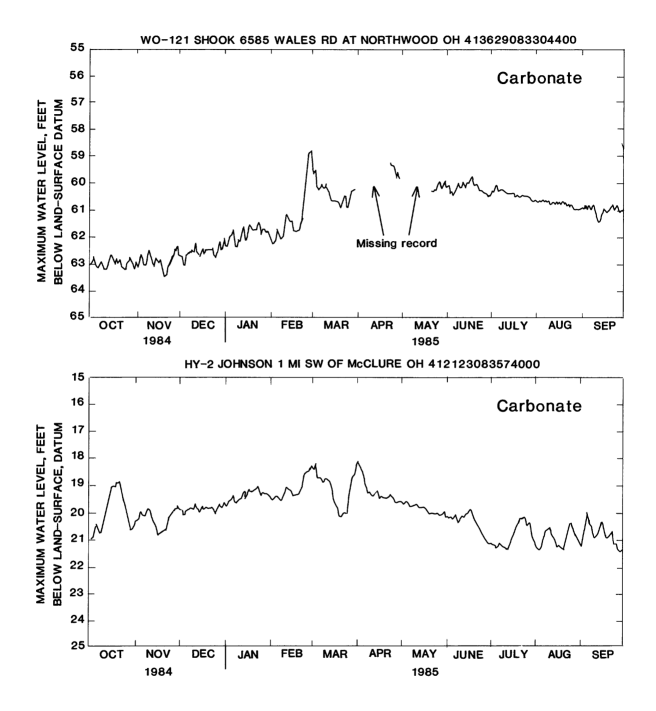


Figure 7.-- Hydrographs showing relation between water levels in wells WO-113 (till), WO-121 (carbonate), and HY-2 (carbonate) and daily precipitation for 1985 water year-Continued.

Water levels in well 113 (Boring B-4, Bowser Morner Testing Laboratories, 1980) reached a maximum of 0.96 ft above land surface on March 15, 1985, in response to precipitation on an already saturated surface. The high water levels reflecting periods of recharge from January through May are replaced by downward-trending water levels during summer and fall that reflect the moisture deficit in the overlying clays and tills. The reaction of well 113 to precipitation indicates communication between the surface and the upper/lower till, perhaps by means of fractures, sand seams, or other permeable deposits.

By comparing the hydrographs of confined wells HY-2 and 121, it is possible to match up a series of barometric fluctuations, even though HY-2 was affected by pumping from October through December 1984 and June through September 1985. Aside from the effects of pumping, the most significant event is the rise in water level from February through March 1985. Water levels rose 2.91 ft in well 121, as compared with 1.05 ft in HY-2. Records of wells LU-1 and S-3, located in confined dolomite aquifers in adjacent Lucas and Sandusky Counties, show increases of less than 0.5 ft for this same period (Shindel and others, 1986), which indicates that water-level increases in 121 are abnormal. Climatological data (National Oceanic and Atmospheric Administration, 1985) indicate that precipitation is comparable for all these areas.

Long-term fluctuations also may be seen in the hydrograph comparison of 121 and HY-2. After spring recharge the hydrograph of well 121 shows a sustained higher water level as compared with well HY-2. Because of this transient response, it is likely that the net water-level increase of approximately 2 ft reflects an increase in storage in the aquifer due to recharge.

Water-level rises in a confined aquifer do not necessarily indicate an increase in storage, and may be caused by atmospheric pressure fluctuations. As atmospheric pressure increases, the water levels in wells tapping confined aquifers are lowered. The ratio of water-level changes in the well to the inverse of water-level changes in a water barometer 1s called the barometric efficiency (BE) of the aquifer (Walton, 1970).

The BE of the dolomite aquifer underlying the landfill was calculated (fig. 8) from barometric and water-level data collected during the period January 25 to 26, 1985. If the barometric efficiency of an aquifer is known, the barometric pressure needed to cause a corresponding increase in water levels within the aquifer can be calculated using the equation:

$$\Delta Bp = \frac{\Delta W(100)}{BE}$$

where BE is barometric efficiency of aquifer, in percent;

 ΔW is change in water level resulting from a change in atmospheric pressure, in ft; and

 Δ Bp is change in atmospheric pressure, in ft of water.

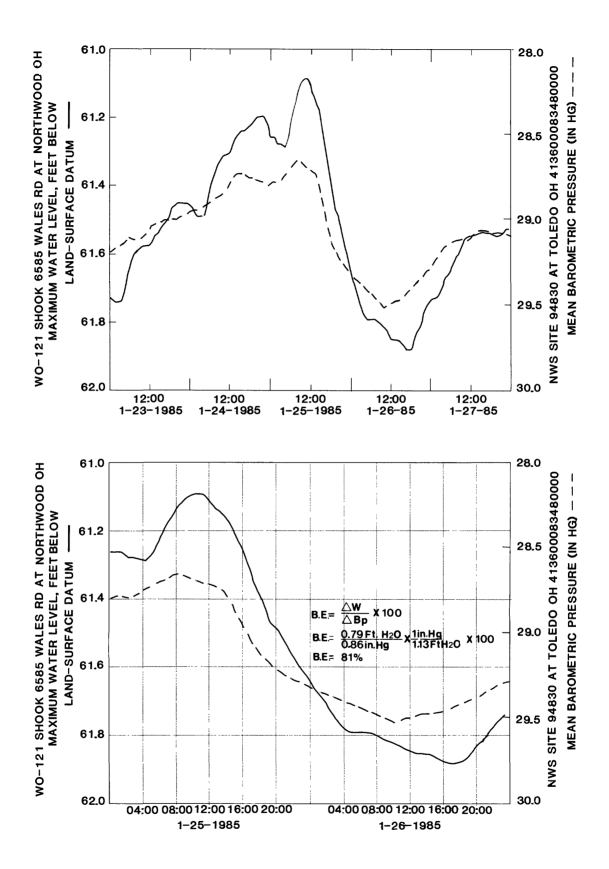


Figure 8.-- Effect of atmospheric-pressure fluctuation on water levels in well WO-121.

Using 81 percent as the calculated BE of the aquifer and the water-level increase of 2.91 ft in well 11 during February 1985, an impossible barometric-pressure change of 26.5 ft of water or 594 mm (millimeters) Hg would have been required to cause a 2.91-ft water-level fluctuation. Considering normal yearly barometric fluctuations for the study area are usually less than 3.4 inches of water (6.35 mm Hg), it is likely that water-level increases in the study area are a result of recharge to the aquifer, not barometric-pressure changes.

Vertical head gradients near the landfill (fig. 9) indicate that water is moving downward from the landfill to the dolomite aquifer through a continuously saturated section of till. Wells 113 and 123, which are completed in the upper/lower till and the lower till, respectively, have consistently higher heads than 112, which is completed in the dolomite. The direction of ground-water flow is from high head to low head; thus, the flow near the landfill is downward toward the dolomite aquifer.

Downward vertical flow toward the dolomite aquifer also is indicated by head conditions in the landfill's northern solid-waste cells. Fluid levels in the methane-venting wells were measured twice and showed a rise similar to that in other wells in the study area.

Well	Date	Altitude of fluid level (ft above sea level)
WO-152	1-29-85	614.67
	4-03-85	615.21
WO-154	1-29-85	610.37
	4-03-85	611.12

Although it is possible that these fluid levels represent perched zones within the refuse, the presence of seepage zones at the edge of the landfill would indicate that the fluid surface is fairly uniform. A leachate seep emerges at an altitude of 614 ft above sea level at the landfill toe near well 107, and there is a seepage face at an approximate altitude of 614 ft along the eastern boundary of the landfill.

These measurements indicate a fluid level within the landfill at or near the original land surface of 615 ft above sea level—very similar to levels in well 113. Comparison of fluid levels within the landfill with wells 123 and 112 indicates a downward hydraulic gradient from the landfill to the till and to the dolomite aquifer. The position of the potentiometric mound (fig. 9) suggests that the landfill is the area of recharge (downward flow) and the cause of the mound.

Water levels in wells 113 and 123 indicate that parts of both the upper and lower till are saturated. Similar conditions likely exist beneath the landfill. Rapid transmission of

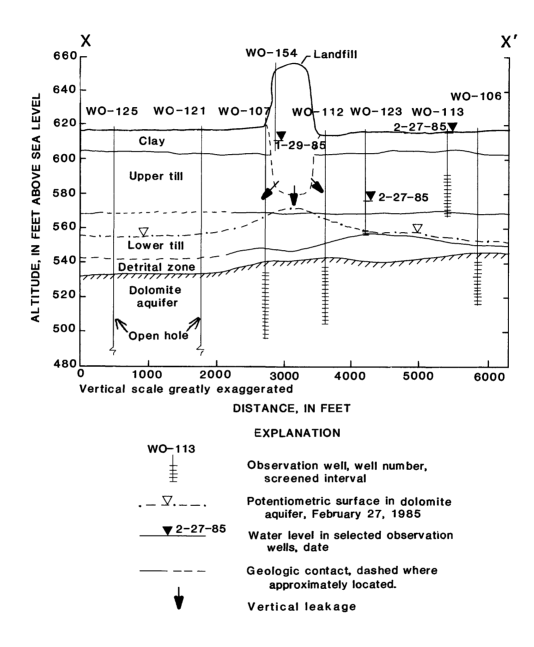


Figure 9.-- Hydrogeologic section X-X' showing head relations between the landfill, till, and dolomite aquifer.

recharge to the dolomite aquifer may be encouraged by desiccation cracks formed by drying and shrinkage of clays and by fractures within the till.

WATER QUALITY

The water-quality sampling program was designed to assess the current water quality in the study area and to determine if the landfill was affecting ground-water quality or surface-water quality. Water-quality samples (fig. 10) were collected from 18 wells, 3 surface-water sites, and 2 methane-venting wells. To assess seasonal water-quality variations, several wells were sampled more than once during the period April 1984 to July 1985. Water samples were analyzed for physical properties; major, minor, and trace constituents; nutrients; organic constituents; and stable isotopes. In addition, streambed sediments were analyzed for trace metals and base/neutral- and acid-extractable organic compounds. Table 2 (at back of report) presents results of analyses of the ground water.

Ground Water

The chemical composition of ground water in the dolomite aquifer differs considerably throughout the study area, as seen from the wide range in concentration of many individual constituents (table 3). Ground water from the dolomite aquifer is very hard, has a median pH of 7.55 and a median specific conductance of 1,025 μ S/cm (microsiemens per centimeter at 25 °C). The most abundant anions are sulfate and bicarbonate, with median values of 425 mg/L (milligrams per liter) and 141 mg/L, respectively. The major cations are calcium, with a median concentration of 100 mg/L, and magnesium, with a median concentration of 41 mg/L. Chloride concentrations range from 3.3 to 26 mg/L, with a median value of 10 mg/L. Sodium has a median concentration of 61 mg/L. Potassium is present in amounts ranging from 1.9 to 3.6 mg/L.

Iron and manganese range widely in concentration, iron from 3 to 2,400 μ g/L (micrograms per liter), and manganese from 1 to 550 μ g/L. Ground water in well 112 exceeds the Ohio Environmental Protection Agency (1978, 1980) drinking-water standard (table 2) for both iron and manganese. The dominant nitrogen species is ammonia, with a median concentration of 0.4 mg/L. Total organic carbon ranges from 0.7 to 8.7 mg/L, with a median concentration of 1.3 mg/L.

Phenols with concentrations ranging from 1 to 5 μ g/L were detected in four wells. Waters in wells 107 and 111 exceed the Ohio Environmental Protection Agency drinking-water standards for phenol. In all other wells tested, no organic constituents were above detection levels. Analysis of ground-water samples for 26 VOCs (volatile organic compounds), 43 base/neutral-extractable compounds, and 11 acid-extractable organic compounds (table 4) detected 2 μ g/L of di-n-butyl phthalate in well 107 and 8 μ g/L of 2-ethylhexyl phthalate in well 112.

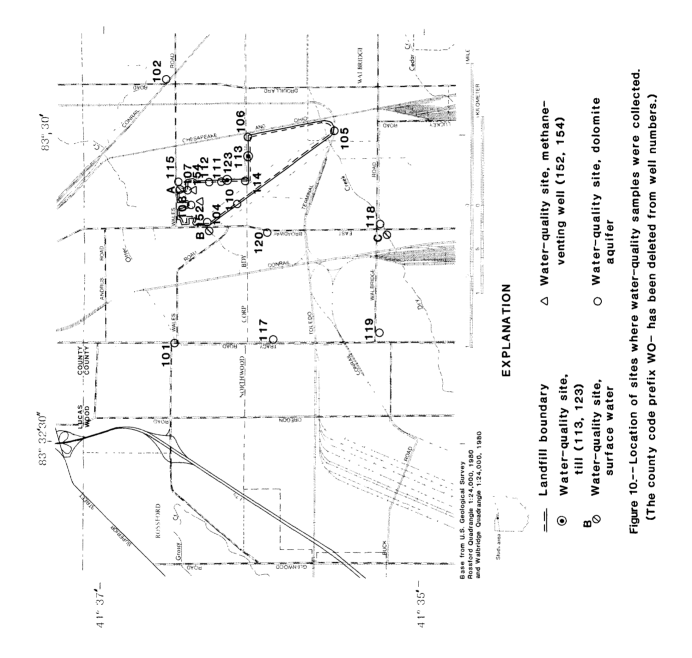


Table 3.--Statistical summary of ground-water-quality data for wells in the dolomite aquifer

	Number					
	of	ŊĊ	er			
Constituents	obser	1				
or	va-	-	5		Minimum	Maximum
properties	Unit tions	ns tions	s Mean	Median	value	value
Alkalinity, field (as CaCO3)		2 32	159	116	30	492
Carbon, organic, total			1.68	1.3	۲.	8.7
Cyanide, total		11 0	1	1		ł
Methylene blue active substances			.03	.02	.01	.07
Nitrogen, nitrite, dissolved (as N)			.01	.01	.01	.05
Nitrogen, NO2 + NO3, dissolved (as N)	mg/L 2	7 1	ł	ł	<0.1	.16
Nitrogen, ammonia, dissolved (as N)	mg/L 2	7 27	.42	-4	.28	.63
ammonia + organic, e	mg/L 2	7 27	. 98	۲.	.30	6.20
organic, dis	mg/L 2	7 27	.57	.31	00.	5.80
Nitrogen, ammonia, dissolved (as NH4)	mg/L 2	7 27	.54	.52	.36	.81
Oxygen, dissolved			6.		<0.1	3.9
pH			T.T		6.6	11.5
Phenols, total			1.7		-1	S
Specific conductance at 25 degrees C	us/cm 3	32 32			667	1,560
					<0.5	. 6
Chloride. dissolved					3.3	26
Fluoride. dissolved					.2	2.1
Hardness (as CaCO3)					220	730
Hardness. noncarbonate (as CaCO3)					35	550
					ŝ	2.400
Magnesium, dissolved					5	82
Manganese, dissolved	uq/L 2	29 29	58	ъ		550
					1.9	3.6
Solids, sum of constituents, dissolved		27 27			500	1,200
Solids, residue at 180 deg. C, dissolved					546	1,250
_ !					3.8	20
Sodium, dissolved					27	130
Strontium, dissolved			• •		14,000	28,000
Sulfate, dissolved					200	750
Chromium, dissolved					10	10
Boron, dissolved					240	830
Aluminum, dissolved		0 16			10	300
Arsenic, dissolved					ħ	1
Cadmium. dissolved						16
Calcium, dissolved	mg/L 3				56	170
Lead, dissolved					Ļ	9
Lithium, dissolved	ng/L 1	12 12			16	44
Selenium. dissolved					1	ł
Oxygen demand. chemical	ma/L 1	6	14		10	34
		1	1			

[Dashes indicate value not calculated.]

Purgeable organic compounds, 1 total recoverable

- Benzene Bromoform Carbon tetrachloride Chlorobenzene Chloroethane 2-Chloroethyl vinyl ether Chloroform Chloromethane Dibromochloromethane Dichlorobromomethane 1,1-Dichloroethane 1,2-Dichloroethane 1,1-Dichloroethylene
- 1,2-trans-Dichloroethylene 1,2-Dichloropropane 1,3-Dichloropropene Ethylbenzene Methylbromide Methylene chloride 1,1,2,2-Tetrachloroethane Tetrachloroethylene Toluene 1,1,1-Trichloroethane 1,1,2-Trichloroethane Trichloroethylene Vinyl chloride

Base/neutral-extractable compounds²

Acenaphthene Acenaphthylene Anthracene Benzo (a) anthracene Benzo (b) fluoranthene Benzo (k) fluoranthene Benzo (g,h,i) perylene Benzo (a) pyrene 4-Bromophenyl phenyl ether Butyl benzyl phthalate bis (2-chloroethoxy) methane bis (2-chloroethyl) ether bis (2-chloroisopropyl) ether 2-chloronaphthalene 4-chlorophenyl phenyl ether Chrysene Dibenzo (a,h) anthracene 1,2-Dichlorobenzene 1,3-Dichlorobenzene 1,4-Dichlorobenzene Diethyl phthalate Dimethyl phthalate

Di-n-butyl phthalate 2,4-Dinitrotoluene 2,6-Dinitrotoluene Di-n-octylphthalate bis (2-Ethylhexyl) phthalate Fluoranthene Fluorene Hexachlorobenzene Hexachlorobutadiene Hexachlorocyclopentadiene Hexachloroethane Indeno (1,2,3-cd) pyrene Isophorone Naphthalene Nitrobenzene n-Nitrosodimethylamine n-Nitrosodi-n-propylamine n-Nitrosodiphenylamine Phenanthrene Pyrene 1,2,4-Trichlorobenzene

Acid-extractable compounds²

4-Chloro-3-methylphenol 2-Chlorophenol 2,4-Dichlorophenol 2,4-Dimethylphenol 4,6-Dinitro-2-methylphenol 2,4-Dinitrophenol 2-Nitrophenol 4-Nitrophenol Pentachlorophenol Phenol 2,4,6-Trichlorophenol

¹Analyses for selected purgeable organic compounds were performed on ground water from wells WO-102, WO-104, WO-107, WO-111, WO-112, WO-113, WO-115, WO-119, and WO-123. The detection level for purgeable organic compounds is 3 μ g/L. (Analysis by gas chromatograph with mass-specific detector.)

²Analyses for selected base/neutral- and acid-extractable organic compounds were performed on ground water from wells WO-102, WO-107, WO-112, and WO-119. Surface water and bottom material from sites A, B, and C were also analyzed for organic compounds. Detection levels for acid-extractable compounds were 1 μ g/L and 20 μ g/kg in water and bottom material, respectively. Base-neutral-extractable organic compounds were detectable at concentrations greater than 1 μ g/L in water and greater than 10 μ g/kg in bottom material.

The phthalate esters detected are commonly used as plasticizers, and low levels of these compounds are considered to be widespread in the environment (Mayer and others, 1972). Phthalate esters also may appear as artifacts in organics analysis because of low-level contamination of items used in the laboratory process. Because of these factors, the organic compounds detected cannot be considered significant in terms of denoting organic contamination.

The differences in relative concentrations of the common ions are easily seen by means of a trilinear diagram (fig. 11), which graphically compares ground waters on the basis of percentage of total cation and anion charge in a water sample. Water types in the dolomite aquifer range from calcium-magnesium sulfate to calcium-magnesium-sodium sulfate. Upon visual inspection of the cation and anion triangles of the diagram, the well waters may be qualitatively separated into groups:

Cation groups	Anion groups
 WO-110 (grout effect) WO-119 WO-113 WO-123 (till), WO-115 WO-152 (methane) All other wells 	 WO-110 (grout effect) WO-107, WO-113 (till), WO-119 WO-123 (till) WO-112 WO-152 (methane) WO-111 All other wells

In the case of sample 152, the cause of the distinct group is high percentages of bicarbonate and chloride and a relatively low percentage of sulfate. However, the cause of other groups is not so obvious and is due to well construction, aquifer mineralogy, or degraded water quality.

Waters in wells 113 and 123 are chemically different from wells in the dolomite aquifer because they produce water only from the upper/lower and lower till, respectively. They also differ appreciably from each other because of the anomalously high concentration of sodium in 123. A plausible source for the elevated concentration of sodium is sodium-montmorillonite clay, commonly used in bentonite grout and the bentonite plug found above the well screen sand pack in well 123.

There are also significant differences between the April 1985 and July 1985 samples from till well 113. Elevated concentrations of dissolved oxygen, low temperature, and low concentrations of iron, strontium, and manganese (table 2) may indicate that relatively recent recharge affected water quality in April 1985. In contrast, values for the July 1985 sample show much higher concentrations of iron, strontium, and manganese and low dissolved oxygen (table 2). As would be expected from geologic, mineralogic, and permeability data, waters from the till are more mineralized than waters from the dolomite aquifer. Because of the calcareous nature of the till and because the surfacearea-to-volume ratio is greater in fine-grained aquifer materials, more minerals may be

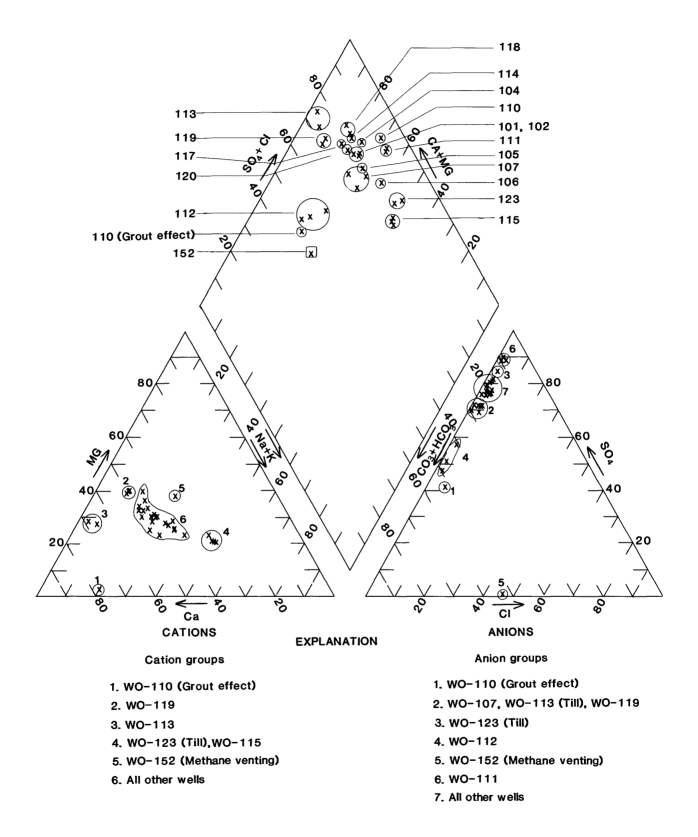


Figure 11.-- Chemical character of ground water and cation and anion groupings of waters at Northwood, Ohio (data in percentages of meq/L: the county-code prefix WO- has been deleted from well numbers).

dissolved into solution. As a result of the differing environment, the ground water from the tills is chemically very different from water in the dolomite aquifer; thus, comparisons between till and bedrock waters can be made only after careful consideration of the individual constituents.

Wells 117 and 119, although finished in the dolomite, also may produce water from the overlying till. The log of well 119 indicates the casing penetrates only 1.5 ft into the bedrock, and the owner reports recurring problems with turbidity during spring recharge and periods of heavy precipitation (Wendell Bihn, Bihn Well Drilling, oral commun., 1984). The log of well 117 indicates that the casing stops 15 ft short of the lower till/ bedrock contact.

Well 118, although not depicted in a separate cation or anion group on the trilinear diagram, has higher concentrations than other dolomite aquifer wells of almost all cations and anions. Because the cations and anions are present in the same relative percentages as other wells in the main group, the more mineralized nature of water from well 118 is not evident. There is no evidence to suggest that the well produces water from the overlying till, and well records indicate a higher specific capacity than most wells in the study area. The more highly mineralized water in well 118 may be related to the geologic strata penetrated.

Waters from well 110 and, to a lesser extent, well 106, are affected by leaching of cement grout and by chemical reactions between the grout and well water in the aquifer near the well screen. Because of the interactions between grout and ground water, many constituent concentration levels are not truly representative of native water quality in the aquifer.

Chemical Reactions in Grouted Wells

Barcelona and Helfrich (1986) have noted that cement-grout contamination in monitoring wells can have dramatic effects on water chemistry. These researchers found pH values greater than 12 and alkalinity dominated by hydroxyl ion in cement-grouted wells. As mentioned previously, two of the monitoring wells at the landfill show the effects of grout contamination. Examples of shifts in constituent concentrations that occurred in a well with elevated pH appear in table 5, as well as some plausible chemical reactions that may have contributed to the changes observed. The table illustrates that complex sequences of chemical reactions are likely taking place around the screen of an improperly grouted well. For this reason, water-quality data obtained from such wells (110, 106) should be interpreted with caution.

Grout + CaO + cement + H ₂ O> Ca ²⁺ + 2OH ⁻ PH Ca ²⁺ + Me ²⁺ grout <> Ca ²⁺ grout + Me ²⁺ CaSO ₄ ⁰ (ion pair) + Mg ²⁺ grout <> no reaction likely Ma ²⁺ + CO ₃ ⁻ <> Mg(OH) ₂ (brucite) Mg ²⁺ + 2OH ⁻ <> Mg(OH) ₂ (brucite) 3MGOH ⁺ + 4H3SIO ₄ ⁻ <> Mg(OH) ₂ (brucite) 3MGOH ⁺ + 4H3SIO ₄ ⁻ <> Mg ₂ Si ₄ O ₁₁ .4H ₂ O + 3H ₂ O + OH ⁻ (sepiolite) Ma ⁺ + K ⁴ grout <> Ma ⁺ grout + K ⁺ 2Ma ⁺ + K ⁴ grout <> Ma ⁺ grout + K ⁺ 2Na ⁺ + Me ²⁺ grout <> 2Na ⁺ grout + K ⁺ Ma ⁺ + K ⁴ grout <> 2Na ⁺ grout + K ⁺ Ma ⁺ + K ⁴ grout <> 2Na ⁺ grout + K ⁺ Ma ⁺ + K ⁴ grout <> Ca ^{2²⁺} + CO ₂ (g) Mi ⁺ + CO ₂ (g) <> Ca ^{2²⁺} + H ₂ Mg (OH) ₂ + CO ₂ (g) <> CaCO ₃ CaO + H ₂ O <> Ca ^{2²⁺} + 2OH ⁻ > Ca(OH) ₂ Ca ^{2²⁺} + CO ₃ ^{2⁻} <> CaCO ₃	Characteristics and chemical composition of example native car- bonate ground water un- affected by contamination (median concentrations, table 3)	Plausible reaction at the well screen (not confirmed by analysis of solid phase)	Characteristics and chemical composition of "affected" well water
$ \begin{array}{llllllllllllllllllllllllllllllllllll$	Ш	Grout + CaO + cement + H ₂ O> Ca ²⁺ + 2OH ⁻	= 11.
$ \begin{array}{llllllllllllllllllllllllllllllllllll$	calcium = 100 mg/L	ca ²⁺ + Me ²⁺ grout <> ca ²⁺ grout + Me ²⁺	Calcium = 100 mg/L
		$casO_4^0$ (ion pair) + Mg^{2+} grout <> no reaction likely	
$ \begin{split} Mg^{2^{+}} + 2 OH^{-} <> Mg (OH) 2 (brucite) \\ & 3Mg OH^{+} + 4H_{3}SiO_{4}^{-} <> Mg _{3}Si_{4}O_{11}.4H_{2}O + 3H_{2}O + OH^{-} (sepiolite) \\ & 3Mg OH^{+} + 4H_{3}SiO_{4}^{-} <> Mg _{3}Si_{4}O_{11}.4H_{2}O + 3H_{2}O + OH^{-} (sepiolite) \\ & Na^{+} + K^{+}grout <> Na^{+}grout + K^{+} \\ & 2Na^{+} + Me^{2^{+}}grout <> 2Na^{+}grout + Me^{2^{+}} \\ & Na^{+} + K^{+}grout <> 2Na^{+}grout + K^{+} \\ & Na^{+} + K^{+}grout <> Na^{+}grout + K^{+} \\ & Na^{+} + K^{+}grout <> Ca_{2}^{-} + H^{+} \\ & HCO_{3}^{-} <> Ca_{2}^{-} + H^{+} \\ & HCO_{3}^{-} <> Ca_{2}^{+} + 2OH^{-}> Ca (OH)_{2} \\ & CaO + H_{2}O <> Ca^{2^{+}} + 2OH^{-}> Ca (OH)_{2} \\ & Ca^{2^{+}} + CO_{3}^{2^{-}} <> CaCO_{3} \\ \end{array}$	lagnesium = 41 mg/L	Mg ²⁺ + co ₃ ⁻ <> MgCO ₃	
$\begin{array}{llllllllllllllllllllllllllllllllllll$		Mg ²⁺ + 20H ⁻ <> Mg(OH)2(brucite)	
Na ⁺ + K ⁺ grout <> Na ⁺ grout + K ⁺ $2Na^+ + Me^{2^+}$ grout <> $2Na^+$ grout + Me^{2^+} $Na^+ + K^+$ grout <> Na^+ grout + K ⁺ Na ⁺ + K ⁺ grout <> Na ⁺ grout + K ⁺ $2HCO_3^- <> H_2O + CO_3^{2^-} + CO_2(g)$ 1y $HCO_3^- <> CO_3^{2^-} + H+$ $M_1 CO_3^- <> CO_3^{2^-} + H+$ $M_2 (OH)_2 + CO_2(g) <> MGCO_3 + H_2O$ $CaO + H_2O <> CaCO_3$ $Ca^{2^+} + CO_3^{2^-} <> CaCO_3$		3MgOH ⁺ + 4H ₃ SiO ₄ ⁻ <> Mg ₃ Si ₄ O ₁₁ .4H ₂ O + 3H ₂ O + OH ⁻ (sepiolite)	
$2Na^{+} + Me^{2t}grout <> 2Na^{+}grout + Me^{2t}$ $Na^{+} + K^{+}grout <> Na^{+}grout + K^{+}$ Poly $Na^{+} + K^{+}grout <> Na^{+}grout + K^{+}$ $2HCO_{3}^{-} <> H_{2}O + CO_{3}^{2^{-}} + H^{+}$ $HCO_{3}^{-} <> CO_{3}^{2^{-}} + H^{+}$ $HCO_{3}^{-} <> CO_{3}^{2^{-}} + H^{+}$ $Mg (OH)_{2} + CO_{2} (g) <> MgCO_{3} + H_{2}O$ $CaO + H_{2}O <> CaCO_{3}$ $Ca^{2^{+}} + CO_{3}^{2^{-}} <> CaCO_{3}$	sodium = 61 mg/L	Na ⁺ + K ⁺ grout <> Na ⁺ grout + K ⁺	Sodium = 27 mg/L
Na ⁺ + K ⁺ grout <> Na ⁺ grout + K ⁺ Pol 2HC03 ⁻ <> H ₂ O + CO3 ²⁻ + CO ₂ (g) All Ly HCO3 ⁻ <> CO3 ²⁻ + H ⁺ ity HCO3 ⁻ <> CO3 ²⁻ + H ⁺ Mg (OH) 2 + CO2 (g) <> MgCO3 + H ₂ O CaO + H ₂ O <> Ca ²⁺ + 2OH ⁻ > Ca (OH) 2 Ca ²⁺ + CO3 ²⁻ <> CaCO3		2Na ⁺ + Me ²⁺ grout <> 2Na ⁺ grout + Me ²⁺	
<pre>Iy 2HCO3⁻ <> H₂O + CO3²⁻ + CO2 (g) All ity HCO3⁻ <> CO3²⁻ + H⁺ isent) Mg (OH) 2 + CO2 (g) <> MgCO3 + H₂O CaO + H₂O <> Ca²⁺ + 2OH⁻> Ca (OH) 2 Ca²⁺ + CO3²⁻ <> CaCO3</pre>	Potassium = 2.5 mg/L	Na ⁺ + K ⁺ grout <> Na ⁺ grout + K ⁺	
HCO3 ⁻ <> CO3 ²⁻ + H ⁺ t) Mg (OH) 2 + CO2 (g) <> MgCO3 + H2O CaO + H2O <> Ca ²⁺ + 2OH ⁻ > Ca(OH) 2 Ca ²⁺ + CO3 ²⁻ <> CaCO3	Alkalinity = 116 mg/L	2HCO3 ⁻ <> H ₂ O + CO ₃ ²⁻ + CO ₂ (g)	Alkalinity = 223 mg/L;
Mg (OH) 2 + CO2 (g) <> MgCO3 + H2O CaO + H2O <> Ca ²⁺ + 2OH ⁻ > Ca (OH) 2 Ca ²⁺ + CO3 ²⁻ <> CaCO3	bicarbonate alkalinity	$Hco3^{-} <> co3^{2^{-}} + H^{+}$	primarily carbonate alkalinity with
0H ⁻ > Ca (OH) 2	(carbon dioxide present)	Mg (OH) 2 + Co2 (g) <> MgCo3 + H2O	hydroxy1 contribution
ca ²⁺ + co ₃ ²⁻ <> caco ₃		cao + H2o <> ca ²⁺ + 20H ⁻ > ca(OH) 2	(carbon gloxide absenu)
		ca ²⁺ + co ₃ ²⁻ <> caco ₃	

Table 5.-Chemical changes and plausible reactions in a grouted well at Northwood, Ohio

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Characteristics and chemical composition of example native car- bonate ground water un- affected by contamination	Plausible reaction at the well screen (not confirmed by analysis of solid phase)	Characteristics and chemical composition of "affected" well water
Silica <u>~</u> 8.5 mg/L (as H₄SiO₄)	si(OH)4 + OH ⁻ <> H ₃ SiO ₄ ⁻ + H ₂ O 3MgOH ⁺ + 4H ₃ SiO ₄ ⁻ <> Mg ₃ Si ₄ O ₁₁ •4H ₂ O + 3H ₂ O + OH ⁻ (sepiolite)	H ₃ SiO4 [−]
Aluminum < 20 µg/L	Al (OH) 3am + OH ⁻ <> AlOH4 ⁻ (aq)	Aluminum ~ 300 µg/L
Sulfate = 425 mg/L	ca ²⁺ + so ₄ ²⁻ <> caso ₄ ⁰ Mg ²⁺ + so ₄ ²⁻ <> Mgso ₄ ⁰	Sulfate $\simeq 200 \text{ mg/L}$
	CaSO4 ⁰ (aq) + 2H ₂ 0 <> CaSO4•2H ₂ 0 Sr ²⁺ + SO4 ²⁻ <> SrSO4 (celestite)	
Strontium = 20 mg/L	$sr^{2+} + Me^{2+}grout <> sr^{2+}grout + Me^{2+}$ $sr^{2+} + so_4^{2-} <> srso_4 (celestite)$ $sr^{2+} + co_3^{2-} <> srco_3$	Strontium
Iron ≃ 73.5 µg/L	Fe ²⁺ + 20H ⁻ <> Fe (OH) 2	Iron = $12 \ \mu g/L$

Table 5.-Chemical changes and plausible reactions in a grouted well at Northwood, Ohio-Continued

Controls on Native Ground-Water Quality

The chemistry of ground water is controlled, in part, by the rocks and minerals with which the waters are in contact. The presence and solubility of certain mineral phases may control the concentration of constituents in natural-water systems.

The general chemical composition of the carbonate rocks (primarily the Lockport Dolomite) of northwestern Ohio has been characterized by studies conducted at quarries in Wood County and adjacent Sandusky and Ottawa Counties. Stout (1941) presented numerous chemical analyses of the carbonates of northwestern Ohio. Turner (1977) lists analyses of the Greenfield Dolomite in Lucas County and the Lockport in Lucas and Sandusky Counties. Stith (1983) lists a single analysis of the Lockport Dolomite from Sandusky County. The chemical results for Greenfield and selected samples from the Lockport appear in table 6, and show that the rocks are dolomites containing silica, aluminum, strontium, sodium, potassium, iron, manganese, and sulfur.

Mineralogically, the rocks are comprised of calcite $(CaCO_3)$ and dolomite $(CaMg(CO_3)_2)$. Fluorite (CaF_2) , celestite $(SrSO_4)$, and strontianite $(SrCO_3)$ have been recognized in Lockport rocks of Sandusky County (Carlson, 1986) and Ottawa County (Morrison, 1935). Much of the silica in the carbonate rocks of the midwestern United States is present as chert or chalcedony in layered nodules or bands (Weiner and Koster Van Groos, 1976). Gypsum $(CaSO_4 \cdot 2H_2O)$ and anhydrite $(CaSO_4)$ also have been observed in the Lockport Dolomite and the younger Salina Formation of Silurian age (Carlson, 1986; Janssens, 1977).

The equilibrium chemical model WATEQF (Plummer and others, 1976) was used to compute saturation indices for mineral phases recognized in carbonate rocks of the region, as well as for several related phases including quartz, amorphous silica (silica gel precipitate), and amorphous silica (glass), which has an intermediate solubility between amorphous silica gel and the crystalline-SiO, polymorphs, quartz, and chalcedony.

Mineral-equilibria calculations on well waters at Northwood that had cation-to-anion imbalances of less than 5 percent appear as saturation indices in table 7. This form of the saturation index also has been termed the "disequilibrium index." Jenne and others (1980) observed an increase in the variance of disequilibrium indices about the equilibrium (zero) value as the magnitude of the solubility value increased for a solid phase. A value of 5 percent of the log of the solid's solubility product (± 0.05 (log K)) was found to represent a useful error band for equilibrium with the solid.

Waters with saturation-index values within the error limit are in a state of chemical equilibrium with respect to those minerals. While in this state, there is no tendency for precipitation of a mineral from, or dissolution of a mineral by, these waters. Table 7 also shows the log K values and associated 5-percent limits for selected mineral phases at a temperature of 11 °C, which is generally representative of the ground-water system.

Table 6 <i>Chemical analyses of</i>		samples of Gr	Greenfield in	and Loc. northwe	i and Lockport Dolomites reported in northwestern Ohio	omites re	ported in	in the literature	rature for	r selected	d sites
		[Dashes ir	indicate da	data not a	available o	or not ap	applicable]				
County/quarry	Reference					Weight	percent				
		caco ₃	MgCO3	si02	Fe203	MnO	K ₂ 0	Na ₂ 0	SrO	A1203	S
Greenfield Dolomite											
Lucas/Maumee Stone (average of 11 analyses)	Turner, 1977	54.37*	44.67*	1	ł	ł	ł	ł	ł	1	ł
<u>Lockport Dolomite</u>											
Sandusky/Martin Marietta, Woodville (average of 11 analyses)	Turner, 1977 (Guelph)	55.09*	44.15*	;	1	ł	ł	ł	1	}	ł
Lucas/Maumee Stone (average of 4 analyses)	Turner, 1977 (Guelph)	54.71*	44.46*	!	1	}	1	}	ł	1	ł
Sandusky/Ohio Lime	Stith, 1983	54.5	44.6	0.38	0.060	0.006	0.031	0.030	0.006	0.068	0.021
Ottawa/U.S. Gypsum	Stout, 1941	52.42	44.17	.14	.02	°03	.02	<.01	1.59	.04	.50
Sandusky/Woodville Lime Products	Stout, 1941	53.72	45.38	.10	-02	.025	<.01	<.01	.35	.02	.12
Wood/Pelton	Stout, 1941	53.76	45.42	.14	.12	.02	<.01	<.01	<.01	.05	10.
Wood/Sherman	Stout, 1941	53.67	45.28	.19	.20	.02	<.01	<.01	<.01	.14	.01
Wood/Owens Pit	Stout, 1941	53.70	45.34	.20	.10	.02	<.01	<.01	<.01	.04	.01
Wood/Wood Stone, Bowling Gr.	Stout, 1941	53.79	44.96	.20	.14	.025	<.01	<.01	<.01	.14	.16
Wood/Wood Stone, Bowling Gr.	Stout, 1941	53.85	44.82	.30	.12	.025	<.01	<.01	<.01	.16	.22
Wood/Luckey	Stout, 1941	54.04	45.65	.17	.02	.015	<.01	<.01	<.01	.02	<.01
*Approximated from mole percentages	le percentages.										

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Table 7.-Logarithms of pCO2 and saturation indices for selected mineral phases in well waters at Northwood, Ohio

Well	Pro- duc- ing zone	Date	<pre>Percent cation(+) anion(-) imbal- Datel ance</pre>)/) log pCO2 (atm)	Cal- cite	Dolo- mite	Fluo- rite	Celes- tite	Stron- tianite	-dyp-	Anhy- drite	Quartz	Chal- cedony	Silica glass, amor- phous	Silica gel, amor- phous
WO-1 02	D	7-10	+1.1	-2.60	-0.12	-0.44	+0.09	6u 0-	-0.33	-0.99	-1.37	+0.37	-0.17	-0.68	-0.96
WO104	D	7-10	-1.9	-2.77	- 08	35	+ .03	08	34	95	-1.32	+ .30	24	75	-1.03
WO-105	D	4-04	+1.7	-2.93	+ .10	02	60. +	10	04	-1.07	-1.45	+ .31	23	74	-1.02
WO-106	D	4-04	+3.4	-3.99	66. +	+1.71	- 03	05	87	-1.04	-1.43	+ .37	17	68	96
WO-107	Δ	7-10	-1.7	-2.43	90. +	11	+ .04	04	12	98	-1.35	+ .39	15	66	¥6
WO-110	D	4-04	6.1	-2.25	+ .76	+1.17	24	18	+ .50	-1.03	-1.42	+ .11	43	94	-1.22
WO-111	D	7-11	+1.2	-3.08	+ .08	06	16	11	47	68	-1.06	+ .41	13	64	92
WO-112	D	7-11	L. +	-1.22	13	51	33	19	63	81	-1.19	+ .75	+ .21	30	58
WO-113	٤	4-04	+2.2	-2.02	+ .23	05	-1.48	-1.48	-1.94	40	85	+ .53	03	55	81
WO-114	۵	4-04	+3.5	-2.85	+ .05	12	+ .12	03	11	98	-1.37	+ .35	19	70	98
WO-115	D	7-09	+ .3	-3.08	+ .25	+ .36	19	11	+ .18	-1.19	-1.51	+ .29	23	74	-1.04
WO-117	ħ	4-02	+4.4	-2.42	09	38	10	03	30	94	-1.32	+ .41	13	65	92
WO-118	D	4-02	0. +	-2.31	15	44	+ .07	+ .12	35	79	-1.18	+ .37	17	68	- 96
WO-119	DT	7-10	-1.0	-2.16	00. +	14	04	09	31	06	-1.29	+ .49	05	56	84
WO-120	۵	3-27	+3.5	-2.67	10. +	16	+ .02	14	24	-1.00	-1.39	+ .32	22	73	-1.01
WO-123	٤ı	7-11	+5.4	-2.58	+ .19	+ .25	- 99	32	60	67	-1.00	+ .39	32	65	94
logK11oC	υ				-8.41	-16.75	-11.13	-6.60	-9.29	-4.62	-4.23	-4.23	-3.69	-3.18	-2.90
5%logK error limit (absolute value)	error lute	logK error limit (absolute value)	۔ ب		.42	.84	.56	.33	.46	.23	.21	.21	.18	.16	.14
-															

[Producing zone: D, dolomite; T, till]

¹Month-day. All dates are for 1985.

Negative index values greater than the 5-percent error indicate conditions of undersaturation. Under these conditions, mineral phases should be dissolving. Positive index values greater than the 5-percent error indicate conditions of oversaturation. There is a thermodynamic tendency for mineral phases to precipitate from oversaturated waters. From these data, conclusions can be drawn about the controls on water chemistry operating in the native ground-water system. The conclusions are listed in table 8, and selected points are further addressed below.

The chemical characteristics of ground water in carbonate-rock aquifers often are controlled by chemical processes far more complicated than equilibrium mineral solubility. Back and others (1983) explain that as ground water flows in a carbonate-rock aquifer, the water initially dissolves calcite, dolomite, and gypsum or anhydrite at varying rates. Chemical equilibrium (saturation) is first established with calcite and dolomite. The chemistry of the ground-water system is altered by the continued dissolution of gypsum, which increases calcium and sulfate concentrations. Increased calcium and sulfate concentrations lead to the incongruent dissolution of more dolomite accompanied by the formation of calcite. Calcite precipitation removes calcium ion from the ground water and effectively keeps the water undersaturated with respect to gypsum. The net result of this process is to alter calcium:magnesium concentration ratios and increase the concentrations and distinct gypsum undersaturation found in waters of the dolomite aquifer near Northwood.

Ratios of chemical-constituent concentrations such as calcium:magnesium are helpful for indicating similar chemical processes and controls in waters taken from different wells. For this reason and because the carbonate rocks of the study area are not pure mineral phases, the molal ratios of calcium and magnesium were compared between well waters and those determined by chemical analyses of carbonate rocks from selected localities near the Northwood site (table 9). The calcium:magnesium molal ratios appear in table 9, and show that the ratios in the carbonate rocks differ to a slight degree depending on sample location and formation. Ca:Mg ratios in the Greenfield of Lucas County (Turner, 1977) range from 1.015 to 1.043, with an average of 1.025. In the Lockport Dolomite, for which a greater number of samples and sample localities are reported, the Ca:Mg ratio ranges from 0.998 to 1.052. The ground waters in the Northwood area show distinctly higher Ca:Mg molal ratios that range between 1.21 and 1.64 for wells in the dolomite aquifer thought to be unaffected by well grout.

The higher values of Ca:Mg in well waters relative to the rock analyses is likely a result of (1) preferential dissolution and precipitation of calcium carbonate and calcium sulfate mineral phases in the fractures and vugs within the aquifer, or (2) a lack of chemical data for carbonate rocks at the Northwood site, which may indicate higher Ca:Mg mole ratios do exist in the bulk carbonate rocks or in minerals found in vugs and fractures in the dolomite at Northwood. In either case, the carbonate well waters show consistent Ca:Mg ratios, which indicates similar chemical processes and conditions are operating to control native ground-water quality.

Table 8.—Se	lected minerals and solubility	Table 8Selected minerals and solubility control of selected constituent concentrations in well waters at Northwood, Ohio
Mineral	Chemical formula	Conclusions about solubility control
Calcite	CaCO ₃	All waters in equilibrium except at wells 106 and 110 (cement grout effects); calcite considered as control for calcium concentrations.
Dolomite	CaMg (CO3) 2	All waters in equilibrium except those with grout effects. Dolomite considered a control on magnesium concentrations.
Fluorite	CaF2	Fluorite is in equilibrium and is the controlling phase for fluoride concentration, except in waters from till wells 113 and 123, which are undersaturated.
Celestite	SrS04	All waters in equilibrium except at well 113; celestite is a likely control for strontium and a source of sulfate ion.
Strontianite	SrCO3	All waters in equilbrium except till water and grout-affected well water. If the mineral is present, possibly contributes to the control of strontium concentration.
Gypsum	CaSO4 • 2H2O	All wells extremely undersaturated. Gypsum is source for (but not a direct solubility control of) calcium or sulfate.
Anhydrite	caso4	Same as gypsum above, if present.
Quartz	SiO2	All oversaturated. Quartz unlikely, but possible, solubility control for silica.
Chalcedony	SiO2	All in equilibrium except at wells 104, 105, 107, 114, 115, and 120, where slight undersaturation exists. Likely control for silica concentrations.
Amorphous silica glass	SiO ₂ (am)	All well water undersaturated; not considered solubility control for silica, presence is unlikely.
Silica gel	SiO ₂ (am)	Extreme undersaturation, silica control unlikely.

Selected ¹ carbonate rocks			lected ¹ nd waters
County/quarry	Mole ratio of calcium to magnesium	Well number	Mole ratio of calcium to magnesium
Greenfield Dolomite		WO-102	1.49
Lucas/Maumee Stone (average of 11 analyses)	1.025	WO-104	1.45
Lockport Dolomite		WO-105	1.53
	1 007	WO-106 ²	1.67
Lucas/Maumee Stone (average of 4 analyses)	1.037	WO-107	1.56
Ottawa/U.S. Gypsum	1.000	WO-110 ²	2.01
Sandusky/Woodville Lime Products	.999	WO-111	1.60
Sandusky/Martin Marietta,	1.052	WO-112	1.64
Woodville (average of 11 analyses)	1.052	WO-113	2.39
Sandusky/Ohio Lime Company	1.035	WO-114	1.48
Wood/Pelton	.998	WO-115	1.48
		WO-117	1.42
Wood/Sherman	.999	WO-118	1.25
Wood/Owens Pit	.998	WO-119	1.21
Wood/Wood Stone, Bowling Gr.	1.009	WO-120	1.37
Wood/Wood Stone, Bowling Gr.	1.013	WO-123	1.44
Wood/Luckey	.998		

Table 9.-Calcium:magnesium mole ratios in selected carbonate rocks in northwestern Ohio and in ground waters at Northwood, Ohio

¹Ratios in water are from analyses used for table 7; in rock are from analyses in table 6.

²Grout contamination likely.

Although only two wells were sampled, the Ca:Mg molal ratio showed a much higher value in 123 than in 113, which compared favorably with water from the dolomite aquifer. The Ca:Mg ratio in till wells is 1.44 and 2.39 for 123 and 113, respectively.

The till waters have sulfate concentrations nearly double those in the dolomite aquifer waters, and an explanation for this is not readily forthcoming. X-ray-diffraction analyses did not indicate gypsum or other sulfur-bearing phases to be present in the tills at an abundance greater than 5 percent by weight.

Plausible sources of the sulfate are: (1) gypsiferous material in the till matrix; (2) pyrite or other sulfide minerals; (3) hydrogen sulfide gas, which has been oxidized by waters recharging the till; and (or) (4) sulfate retained by anion exclusion processes within the till matrix.

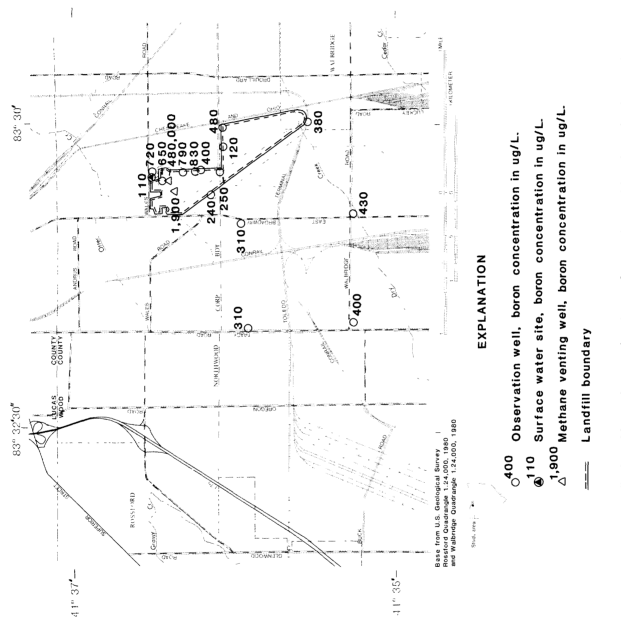
Water Quality and Landfill Leachate

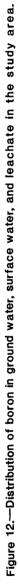
Two samples of leachate were obtained from each of the methane-venting wells, 152 and 154. These wells are finished in the waste material in order to prevent buildup and migration of methane gas. They also provide access to the leachate being produced by the waste material. Both leachates are highly mineralized (table 2), with concentrations of total dissolved solids (TDS, residue at 180 °C) of 3,170 mg/L for 152 and 26,300 mg/L for 154. Although most chemical constituents are typically elevated in a solidwaste leachate, data indicate that boron, iron, ammonia, and total dissolved solids are the most reliable indicators of ground-water pollution by landfill leachate (Clark and Piskin, 1977). Total hardness also is a reliable indicator in this instance because of high concentrations (2,500 mg/L and 12,000 mg/L) in the leachates as compared with native ground water.

The highest concentrations of boron in the study area (fig. 12) are seen in the leachate and in water from wells 107, 115, 111, and 112. All of these wells are located immediately downgradient from the landfill. The concentrations of boron range from 650 to 830 μ g/L in these wells, compared with a median value of 415 μ g/L for all wells in the dolomite aquifer in the study area.

Boron is a reliable indicator of ground-water contamination by leachate, perhaps because of its conservative nature. The more important dissolved-boron species are anionic or uncharged; therefore, they are not extensively adsorbed on other mineral surfaces (Hem, 1985). Sources of boron in refuse are the common cleaning agent borax and borosilicate glass.

Levels of iron in well 107 appear elevated compared to nearby wells 104 and 108. Concentrations of dissolved iron in well 112 (1,500 to 2,400 μ g/L) are highly elevated in





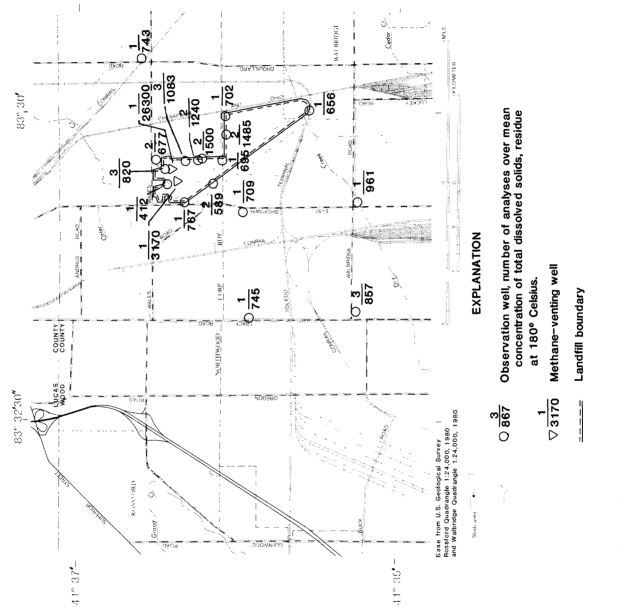
comparison to other carbonate wells (median concentration, 74 mg/L). The high iron concentration, coupled with near neutral pH, and sufficient carbon dioxide, may encourage the growth of iron-oxidizing bacteria, which may cause the precipitation of iron in 112. Iron bacteria are ubiquitous in the environment, but evidence of their metabolic activities is not normally encountered unless the proper conditions for their growth are present. Rust-red-colored precipitates (ferric hydroxide³) were present on the inner walls of the PVC well casing of 112. Evidence for existence of the precipitate was its presence on the cables of water-level measuring devices used on well 112.

The highest levels of ammonia found are as ammonium ion (NH_4^+) in the dolomite aquifer downgradient of the landfill. Ammonium ion is a useful indicator of leachate because large amounts of ammonia (NH_3) are produced from the microbial degradation of organic compounds in landfill refuse. The concentration of ammonium ion (NH_4^+) in the leachate ranges from 43 to 930 mg/L as NH_4 . Ammonium ion has a median concentration of 0.52 mg/L in the dolomite aquifer and is found in all wells in the dolomite aquifer. Concentrations in downgradient wells 107 and 112 are slightly elevated and range from 0.52 to 0.81 mg/L as NH_4 .

The relative age of a leachate may be inferred by its organic N and NH_4 load. In this case, well 152 is more mature because of the lower organic N and NH_4 load. Also, 152 is located in the oldest solid-waste cell in the landfill, and microbial degradation and subsequent migration of leachate has been occurring longer, which has resulted in decreased values of nitrogen compounds. On the other hand, 154 is located in the most recent solid-waste cell, therefore, it has the highest organic N and NH_4 load and is still actively undergoing decomposition. The temperature of the leachate is 5 °C higher for 154 in January 1985, perhaps indicating a higher rate of microbial activity.

All ground waters in the study area exceed the OEPA drinking-water standard (table 2) for TDS. The highest concentration of TDS in the dolomite aquifer is in wells 111 and 112 (fig. 13) located directly downgradient of the landfill. Concentrations also are somewhat elevated in well 107. The TDS in the leachate ranges from 3,170 to 26,300 mg/L. Leachates characteristically have high concentrations of TDS as compared with native ground water because of the degradation of refuse, which supplies a varied and concentrated source of chemical constituents. As the concentration of dissolved components in leachate increases (TDS), the density of that leachate becomes greater than that of the native ground water. The increased density of the leachate compared with native ground water (as measured by TDS) initiates movement of the leachate downgradient. It is not necessary to have a hydraulic-head difference (the ground-water mound) for movement of the leachate away from the landfill to occur.

³Not confirmed by X-ray diffraction analysis.





The significance of increased hardness may result from the displacement of calcium and magnesium from cation-exchange positions on clays exposed to landfill leachate (Griffen and others, 1976). Concentration of hardness ranges from 2,500 to 12,000 mg/L in the leachates, compared with a median value of 440 for wells in the dolomite aquifer. Consistently higher values of hardness (580 to 730 mg/L) are seen in well 112, and slightly elevated values of hardness (580 mg/L) are present in well 111, and, in one analysis, 550 mg/L for well 107.

In summary, water from wells 107, 111, 112, and 115 contains elevated concentrations of certain constituents that suggest mixing of leachate with native ground water. The data, although suggestive of leachate migration, are not conclusive because of the variability of ground-water quality in the study area. However, data on geology, groundwater flow, landfill construction, and surface-water flooding indicate sources and probable routes of leachate migration. In an attempt to qualify and further explain water quality and hydrogeologic data, analyses of oxygen and hydrogen-stable isotope ratios of ground water were used.

Oxygen and Hydrogen Stable Isotopes

Water samples for analysis of oxygen and hydrogen stable isotopes were collected from 2 till wells, 12 dolomite-aquifer wells, and 1 surface-water site. In conjunction with the isotope samples, water also was analyzed for major, minor, and trace constituents and nutrients. Samples were collected after a period of intense precipitation in spring 1985, which caused surface-water flooding and ground-water-level increases of more than 8 ft in the vicinity of the landfill.

Isotopes are atoms of the same element whose nuclei contain the same number of protons but a different number of neutrons. The heavier isotopes are almost always less abundant, and the notation "R" refers to the ratio O-18:O-16 or to the ratio D:H where deuterium (D) is a heavier, less common isotope of hydrogen (H). Isotopic compositions are reported as parts per thousand (permil) differences of rare to common relative to a standard known as V-SMOW or Vienna-Standard Mean Ocean Water. This relative difference is called the delta value (δ) and is defined for hydrogen and oxygen as:

 $\delta D_{\text{sample}} = [(R_{\text{sample}}/R_{\text{V-SMOW}})] - 1 \times 10^3$ $\delta O - 18_{\text{sample}} = [(R_{\text{sample}}/R_{\text{V-SMOW}})] - 1 \times 10^3$

V-SMOW is a water standard with an isotopic composition close to the average ocean water; therefore, it is used in hydrologic investigations. Stable isotopes of O and H are considered to be ideal water-source indicators or chemical tracers because concentrations neither decay with time nor are the isotopes known to be removed from water by exchange processes during movement through most aquifer materials (Winograd and

Friedman, 1972). Silicate hydrolysis reactions, which are not a major control of water chemistry in carbonate terrains, may operate to increase the deuterium content and decrease the oxygen-18 content of water (International Atomic Energy Agency, 1983).

 δ D and δ O-18 can be positive, negative, or zero. A positive value indicates enrichment of a sample in oxygen 18 and deuterium relative to V-SMOW; conversely, negative values of δ O-18 and δ D indicate depletion of oxygen 18 and deuterium isotopes relative to V-SMOW. Values of δ O-18 and δ D in the study area are negative and appear in table 2 for well water and in table 10 (at back of report) for surface water from Otter Creek.

The precision of the isotope analyses for this investigation is ± 0.1 permil for δ O-18 and ± 1.5 permil for δ D. These values represent the 95-percent confidence interval. Values of δ O-18 range from -7.20 permil to -12.2 permil, whereas values for δ D range from -40.0 permil to -84.5 permil.

Figure 14 provides a preliminary indication of the areal variability in isotopic content of ground water and of the potential for multiple sources of ground water in the study area. Waters with three distinctly different isotopic signatures are present in wells upgradient of the landfill. Waters from wells 117 and 119 have a heavier signature than waters from wells 118 and 105, which are significantly lighter. Water from well 120, also upgradient of the landfill, has an isotopic composition that is intermediate to the other two groups of upgradient wells.

The isotopic composition of ground waters within and near the landfill perimeter are even more variable. However, the waters from the glacial tills (wells 113 and 123) have similar compositions that are only slightly lighter than a surface water from Otter Creek. The waters from the tills are notably heavier than any upgradient water in the dolomite aquifer.

Oxygen and hydrogen isotope data for all waters sampled, including leachates, are plotted on a stable-isotope plot for water in figure 15. The values of δD and δO -18 for all waters except leachates plot in a linear trend. A least-squares regression line was fit through data collected from wells in the dolomite aquifer, glacial tills, and surface water. Selected wells in the dolomite aquifer were excluded from the regression on the basis of previous water-quality data indicating probable leachate effects and cement-grout effects. Because the isotope data may indicate differences between ambient and affected water chemistry, wells 107, 111, 115, and 112 were not included in the regression equation because of prior evidence of leachate. The leachates also were excluded from the regression. Waters from wells 106 and 110 also were excluded because of prior evidence of cement-grout contamination. The resulting regression equation, $\delta D = 7.8$ (δO -18) + 10.5, describes a local water line and is similar to Craig's (1961) global meteoric water line of $\delta D = 8$ (δO -18) + 10. This similar linear relation indicates that ground water in the study area can be considered to represent atmospheric precipitation or surface water that has not been isotopically altered as it has entered the ground-water system.

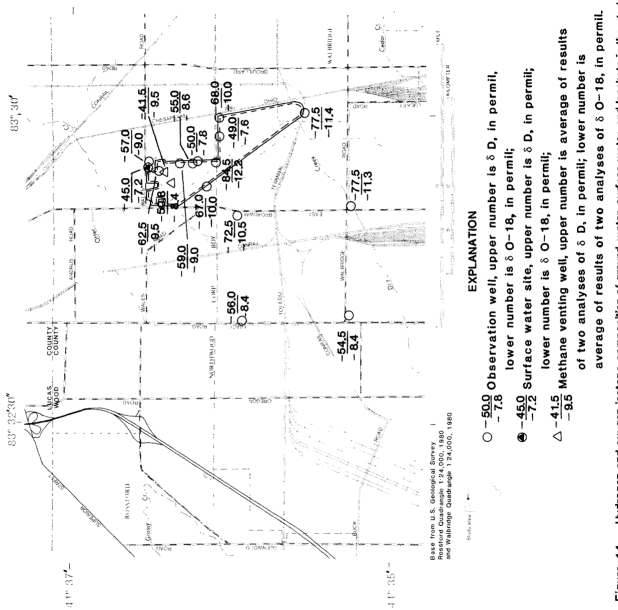
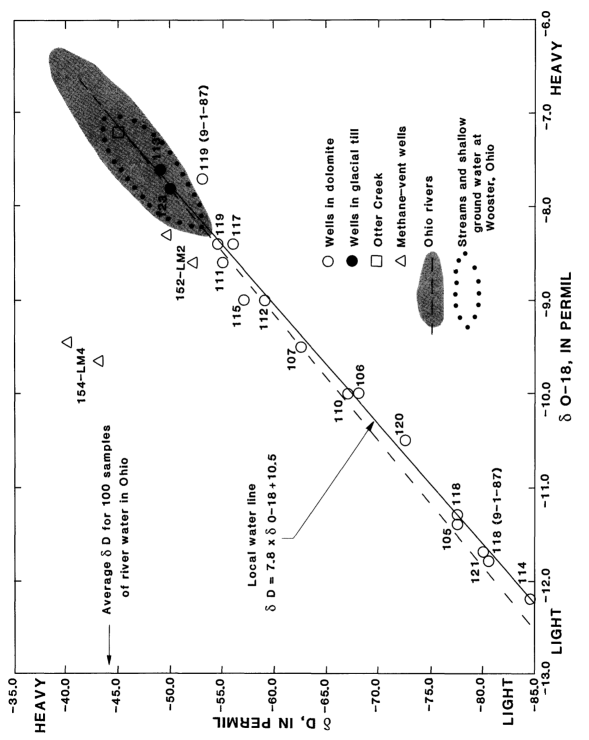
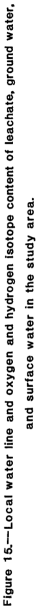


Figure 14.-- Hydrogen and oxygen isotope composition of ground water, surface water, and leachate in the study area.





In an attempt to better understand the time variations in isotopic content of ground water in the Northwood area, wells 118 and 119 were resampled for isotopic analysis on September 1, 1987. These additional data are plotted in figure 15 with date identifiers. These data indicate that variations of 3 permil in δ D and 0.5 permil in δ O-18 might be expected due to analytical precision of isotope analyses and natural variability in waters from a single well.

The recent seasonal variation in the isotopic compositions for river waters in Ohio as determined on a preliminary basis during the 1984-87 water years (Carol Kendall, U.S. Geological Survey, oral commun., 1988) is shown by the stippled elliptical area at the heavy (upper right) part of the figure. These data plot along an "Ohio water line" that is dashed into the part of the plot with lighter isotopic compositions. The average δ D for 100 samples of river water in Ohio is -44 permil. Additional data for seasonal variations in isotopic composition of ground water and surface water from a stream-aquifer system near Wooster in northeastern Ohio (J. E. Dysart, U.S. Geological Survey, oral commun., 1986) also are shown by a dotted ellipse to further illustrate the seasonal variability in recent meteoric water in Ohio. The waters from the till wells (113, 123) and Otter Creek have compositions consistent with a recent meteoric water source. Few waters from the dolomite aquifer (wells 111, 117, 119) approach the isotopic composition of recent meteoric waters.

The isotopic data provide information on generation of leachate in the landfill, which produces waters of anomalous isotopic content. Leachate samples for isotopic analysis were collected from two methane-venting wells, 152 and 154. Both leachates are characterized by elevated concentrations of many common and trace constituents as compared with other wells sampled. Well 154 yields the more concentrated leachate of the two, with a mean specific conductance of 27,900 μ S/cm. Leachate from well 152 has a mean specific conductance of 5,400 μ S/cm.

Baedecker and Fisher (1977), Baedecker (1979), and Baedecker and Back (1979) studied the chemical and isotopic signatures of landfill leachate and waters affected by leachate. These studies showed that one result of the chemical reactions that produce leachate is an isotopic shift to values 0 to 10 permil heavier and 10 permil heavier for δ O-18 and δ D, respectively, in leachate relative to native ground waters. All leachate samples are heavier in deuterium and oxygen-18 than native ground water from the dolomite aquifer.

Each of the leachate wells was sampled twice, so the data (table 2) also provide some information on time variations in isotopic content. The leachate wells were first sampled in January 1985, then resampled in April 1985 after a period of intense precipitation. Both wells indicate a depletion of deuterium and oxygen-18 in recharge water between January and April 1985. The lightest compositions of river water in Ohio often occur in March or April due to light precipitation (Carol Kendall, U.S. Geological Survey, oral commun., 1988). Lighter recharge water may explain the lighter leachate composition in April as compared with January. The most noticeable difference isotopically between leachate and ground water is the shift of the leachate composition to the left of the local water line. In comparing the two leachates, it appears that the more concentrated the leachate is, the greater the shift to the left of the local water line. The shift indicated on figure 16 may be a composite of two separate shifts, a δ D shift off the water line to heavier values and also a δ O-18 shift to lighter values. The shift to heavier δ D has been noted by Baedecker and coworkers and is considered to be characteristic of landfill leachate. A shift to lighter δ O-18, if present, is possibly related to interactions between refuse and water, but these effects are not well documented in the scientific literature pertaining to landfills.

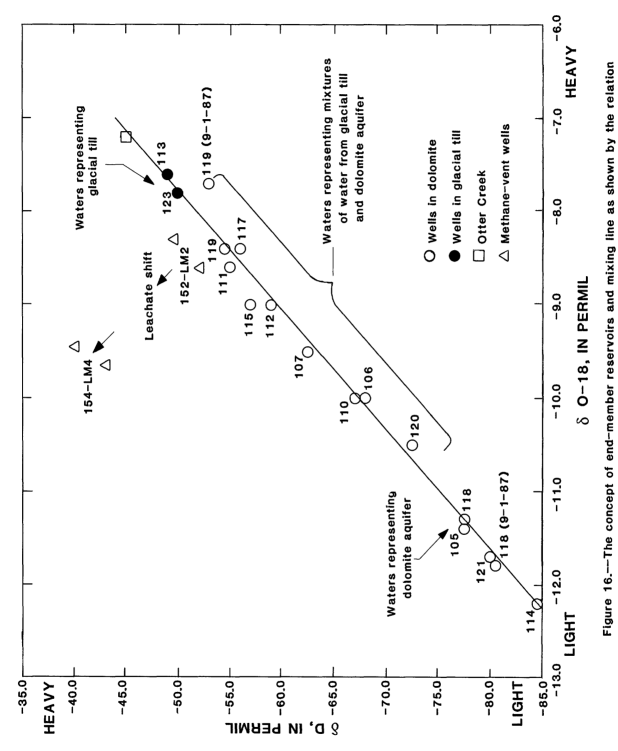
Figure 16 shows two isotopically distinct bodies of ground water along the local water line:

- 1. A reservoir of water in the dolomite aquifer, which is presumed to be relatively old, is removed from direct contact with recent precipitation, and is characterized by light isotopic content of oxygen and hydrogen.
- 2. A reservoir of ground water in the glacial tills which has been recharged relatively recently by precipitation and is characterized by a heavy isotopic signature for oxygen and hydrogen.

The concept of a leaky till reservoir of isotopically heavy water in the tills hydraulically above the dolomite aquifer may help explain some of the variability in isotopic content observed for waters in wells completed in the dolomite aquifer. A simple mixing of till waters with lighter waters from the dolomite aquifer would produce intermediate waters with compositions along the local water line. By applying this mixing concept, the group of waters from the dolomite aquifer that have the highest proportion of till water are from wells 119, 111, 117, 115, 112, and 107.

Wells 117 and 119, although upgradient of the landfill, show a large component of till water. The isotopic data support earlier conclusions regarding the construction of these wells. It appears that wells 117 and 119 both have fairly good access to the lower till-bedrock contact, which previous reports indicate may be a source of small domestic supplies. As a consequence of this mixing, the chemistry of wells 117 and 119 is not truly representative of the dolomite aquifer or the till, and shows characteristics of both.

Other wells in the dolomite aquifer having large proportions of till water are 107, 111, 112, and 115. These wells are directly downgradient of the northernmost landfill cells. Geologic and water-level data presented earlier suggest that these wells have the greatest potential for showing the effects of increased recharge or leakage from the landfill. Water-quality data from these wells showed elevated levels of constituents that are indicators of leachate-affected ground waters. The relatively heavy isotopic signature





in water from these wells supports the idea that the dolomite aquifer receives more recharge from overlying strata in the vicinity of the landfill than in other parts of the study area.

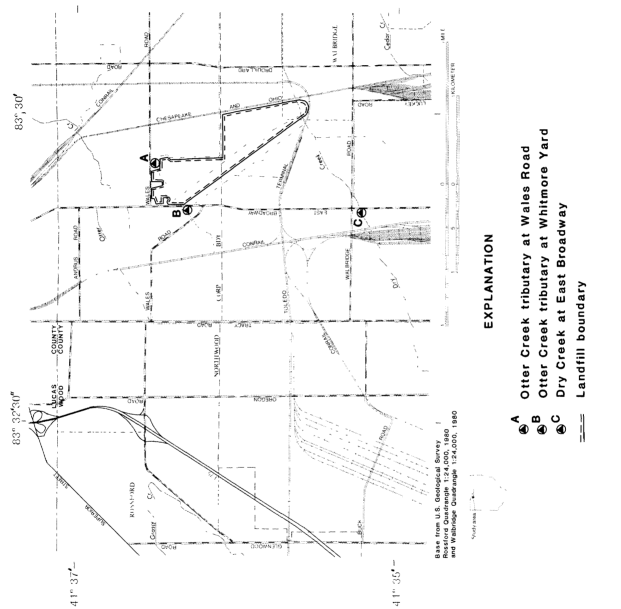
The local water line provides a snapshot of isotopic compositions in the groundwater systems of the study area and represents waters unaffected by leachate. Well 115 shows a shift off the line (fig. 15) in the direction of the leachates. Two interpretations are possible based on existing data: (1) the shift is characteristic scatter (3-permil range for δ D, 0.5-permil range for δ O-18) from natural variability in δ D and δ O-18, or (2) the shift is preliminary evidence of a hydrologic process involving mixing of landfill leachate with ground water beneath the landfill. In the second case, the process would involve three-component mixing of till water, ground water from the dolomite aquifer, and landfill leachate. The isotopic ratios are not shifted to the left of the local water line a significant amount in three of the four wells having other chemical irregularities that indicate leachate is present. The shift to heavier values of δ D (deuterium enrichment) is considered a preliminary indicator of the presence of landfill leachate in well 115, but more detailed study of seasonal variations in isotopic composition is required to resolve this question of leachate in the dolomite aquifer.

Of the constituents mentioned previously as leachate indicators, boron, because of its conservative chemical characteristics, is a particularly reliable indicator of leachate-affected ground water. Ground waters in four wells (107, 111, 112, and 115) in the dolomite aquifer all have boron concentrations above the level of 415 μ g/L, which is considered the background level of this study. Water from well 115 also plots to the left of the local isotopic mixing line. These observations are consistent in suggesting a leachate component exists in the dolomite aquifer.

Isotopic and chemical-quality data indicate that the water quality of the till is distinctly different from that of the dolomite aquifer. In addition, water-quality data indicate major chemical differences between well 113, which is finished in the upper/lower till, and well 123, which is finished in the lower till. Because of these differences, plus the limited amount of water-quality information available on the till, it is not possible to determine if water quality in the till has been affected by the landfill. However, the proximity of the landfill base to the fractures and sand zones near the upper till/lower till contact may allow preferential downward flow to the dolomite aquifer as opposed to lateral flow into the tills.

Surface Water

Samples were collected from three stream sites (fig. 17) during April 1984 and July 1985. Dry Creek at East Broadway just south of Walbridge Road was selected as a control site away from the influence of the landfill. Two sites were chosen on Otter Creek, at Whitmore Yard upstream of the landfill and at Wales Road downstream of the landfill. Results of water-quality analyses are presented in table 10 (at back of report).





Dry Creek is a perennial stream with a streambed composed of gravel, cobbles, and clay. Otter Creek, in this part of its reach, is an intermittent stream and has no flow during periods of sustained dry weather. The composition of the streambeds at both sites on Otter Creek is primarily clay. Observations made while sampling show both Dry Creek and Otter Creek at Whitmore Yard support aquatic plants and animals; no such aquatic life was observed at the Wales Road site.

During April 1984, when flow at all sites was estimated at 1.0 ft³/s (cubic feet per second) or less, water quality in Dry Creek and Otter Creek at Whitmore Yard was comparable. Concentrations of most constituents were slightly higher at the Whitmore Yard site, except for nitrogen, sodium, and chloride, which were elevated in Dry Creek. Otter Creek at Wales Road, downgradient of the landfill, had consistently higher values of most constituents.

Streambed sediments were sampled and analyzed for trace metals and organic compounds. Sediment from the Wales Road site contained the highest concentrations of arsenic and chromium, whereas sediment in Dry Creek contained the highest amount of mercury. All toxic metals detected were at trace levels, that is, less than $10 \,\mu$ g/g (micrograms per gram of bed material).

Organic compounds detected in sediment from the Whitmore Yard site on Otter Creek were: benzo-b-fluoranthene, 180 μ g/kg (micrograms per kilogram of bed material); benzo-a-anthracene, 170 μ g/kg; fluoranthene, 68 μ g/kg; and pyrene, 90 μ g/kg. Fluoranthene (100 μ g/kg) and pyrene (70 μ g/kg), as well as phenanthrene (110 μ g/kg), were detected in sediments at the Otter Creek site at Wales Road. None of these compounds were detected in sediments from Dry Creek. These compounds are polynuclear aromatic hydrocarbons (PAHs), which are ubiquitous in the environment and result from the burning of fossil-based fuels.

Phenol ranging in concentrations from 1 to 7 μ g/L was present in the streams. It is possible that the presence of phenol and PAHs is related to the railway transportation facilities, which since have been removed.

Chemical-quality analyses from the streams during low-flow conditions in July 1985 show that water from Dry Creek contained higher levels of most constituents than did water from the Otter Creek sites. As in the 1984 analyses, Otter Creek at Wales Road showed higher levels of most constituents than did the Whitmore Yard site. Although concentrations of individual constituents increased during low flow in Dry Creek, most constituents at both Otter Creek sites decreased. This suggests that water quality in Otter Creek is more influenced by runoff than by interactions of stream water with streambed materials or ground water.

To determine if compounds in water that runs off the landfill surface were entering Otter Creek, a series of water-quality samples were collected at the Wales Road site during a storm on March 28, 1985. In addition to the landfill, which is the closest drainage area that could contribute to runoff, a waste-hauling business is located just northwest of well 107 and also may affect stream quality. Heavy overland flow was observed entering Otter Creek during the storm, and the stage increased rapidly during the first hour (fig. 18). Values of specific conductance increased from 480 to 520 μ S/cm during the first 40 minutes, reached a peak of 555 μ S/cm 160 minutes after the storm began, and then declined.

Most chemical constituents reached their peak concentration 40 minutes after the storm, then declined as the stage in the stream rose further. The first sample was obtained after the stream had risen 0.6 ft in approximately 15 minutes; therefore, the concentrations are more indicative of an initial "flush" than of ambient conditions. After an initial "flush" of constituents into a stream, the water chemistry typically becomes more dilute because relatively unmineralized precipitation enters the channel directly and as runoff from adjacent areas. An increase after the initial flush and subsequent dilution may indicate a second source. The change of sodium and chloride concentrations seems to support this conclusion. Because of road salting in winter periods, levels of sodium and chloride generally are higher in streams. The probable reaction to a runoff event would be an increase in these levels during the initial flushing and then a decrease as dilution occurs. Sodium, chloride, silica, and fluoride concentrations indicate this effect and show a further increase, perhaps indicating an additional source.

Constituent concentration on March 28, 1985	Sample 1	Sample 2	Sample 3
Sodium (mg/L)	33	15	20
Chloride (mg/L)	34	26	38
Silica (mg/L)	8.8	2.5	2.8
Fluoride (mg/L)	.3	.2	.4
Specific conductance (µS/cm)	480	520	550
Calcium (mg/L)	46	53	52
Magnesium (mg/L)	15	17	15
Sulfate (mg/L)	120	130	120
Iron (m/L)	13	15	14
Alkalinity (mg/L as CaCO,)	77	81	80
Hardness (mg/L as CaCO,)	180	200	190
Potassium (mg/L)	2.7	3.2	3.3

Calcium, magnesium, sulfate, iron, alkalinity, and hardness remained relatively stable during the event. The only constituent that steadily increased during the storm was potassium, which increased from 2.7 to 3.3 mg/L. Potassium is normally a very conservative constituent and is not subject to great variation. The increasing concentrations are likely due to leaching of potassium from dead vegetation adjacent to the stream channel.

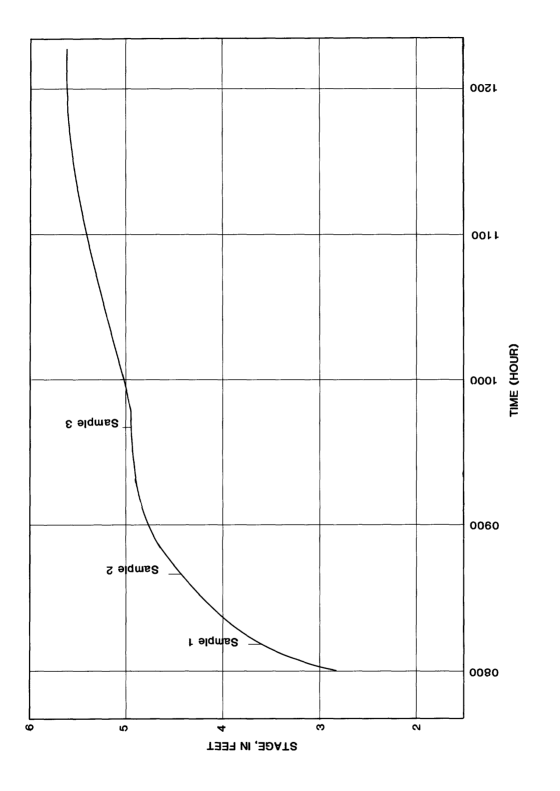


Figure 18.---Stage fluctuation and time of water-quality sampling for Otter Creek tributary at Wales Road, March 28, 1985. Potassium ions assimilated by plants become available for resolution when the plants mature and die (Hem, 1985). As part of the natural cycle, potassium is leached into the soil by rains during the dormant season, and some contribution to runoff would be expected.

The contribution of dissolved solids to Otter Creek during extreme runoff conditions is not very large, although results are not conclusive because of the limitations of one set of storm samples. In terms of concentration, the largest increase was 10 mg/L for sulfate. From the minimal increases seen in constituents during the storm, it does not appear that there is a significant decline in water quality of Otter Creek at high flow from sources upstream from Wales Road.

SUMMARY AND CONCLUSIONS

Hydrogeology and water quality near a landfill in northwestern Ohio were evaluated. The landfill, located near Northwood, Ohio, in Wood County, is used for disposal of solid and hazardous waste. Land use in the area is a mixture of heavy industry, transportation, housing, and agriculture.

A thick sequence of clays and tills of Wisconsinan age covers dolomites of Silurian age throughout the entire study area. The landfill cells are excavated into the upper till and extend to within 10 ft of the upper-till/lower-till contact, which is characterized by sand and gravel zones, mottling, and increased water content. Saturated seams of sand and gravel are found throughout both tills but are more prevalent in the lower till. There are no data to suggest the zones of sand or gravel are continuous; however, their prevalence and relatively higher permeability than the surrounding clay till may allow the zones to act as pathways for recharge to the underlying dolomite aquifer.

The primary aquifers are the Greenfield Dolomite and underlying Lockport Dolomite of Silurian age, which produce water by means of secondary openings such as fractures, solution channels, and voids. The dolomite aquifer is separated from the overlying tills by a thin veneer of glacial outwash, which locally produces small amounts of water.

Water-level measurements in wells completed in the northernmost landfill cells indicate that the refuse is saturated to near land surface. Water-level data indicate that parts of both the upper and lower tills are saturated. Vertical head relations between the landfill, upper till, lower till, and carbonate aquifer indicate that water is flowing downward from the landfill to the dolomite aquifer.

Potentiometric maps of the dolomite aquifer show a ground-water mound centered under the landfill and elongated in the direction of regional ground-water flow. Groundwater levels adjacent to the landfill near the center of the mound fluctuate as much as 14 ft per year, in contrast to fluctuations of less than 3 ft per year in upgradient wells. These factors may indicate that the landfill is a source of recharge to the underlying dolomite aquifer.

The chemical composition of ground water differs considerably throughout the study area. Native ground water is very hard, with a median pH of 7.55. The water type ranges from calcium-magnesium sulfate to calcium-magnesium-sodium sulfate. The composition of native ground water in the dolomite aquifer is controlled by chemical interaction between ground water and minerals occurring naturally in the aquifer. The effects of cement/grout contamination produced a distinctive water chemistry in selected monitoring wells characterized most notably by pH values of 8.9 or greater.

The chemistry of waters from wells completed in the calcareous glacial till was consistent with carbonate-mineral saturation. However, low levels of fluoride and strontium in till waters create conditions of undersaturation for fluorite, celestite, and strontianite that differ markedly from dolomite waters. Sulfate concentrations in till waters are nearly double the concentrations in water from the dolomite aquifer. Two possible sources for the elevated concentrations of sulfate in till waters are: (1) dissolution of gypsum in the till, and (2) oxidation of sulfide minerals or hydrogen sulfide gas by oxygen dissolved in water recharging the till.

All ground waters in the study area exceed the OEPA drinking-water standards for total dissolved solids. Elevated concentrations of boron, iron, ammonia, total dissolved solids, and hardness were detected in leachate and in several wells downgradient of the landfill. These constituents are considered to be the most reliable indicators of ground-water contamination by landfill leachate. Downgradient wells 107, 111, 112, and 115 contained concentrations of boron ranging from 650 to 830 mg/L, as compared with a median concentration of 415 μ g/L for all wells in the dolomite aquifer. Wells 107 and 111 exceed the OEPA water-quality standard for phenolic compounds. Well 112 exceeds OEPA standards for iron and manganese.

Analysis of stable-isotope data shows that waters from wells 107, 111, 112, and 115 have heavy isotopic signatures of hydrogen and oxygen. This indicates that they have recently received recharge from overlying strata. Wells completed in the dolomite aquifer directly downgradient of the landfill show the highest proportion of water from the till.

Infiltration of water into the landfill and subsequent chemical reactions in the refuse produces a leachate having an anomalous isotopic content. Well 115 shows a shift off the local mixing line in the direction of the leachate composition. The shift to heavier values of δD may be an indication of the presence of leachate in this well.

A comparison of observations and stream-water quality at three sites indicates the absence of aquatic life in Otter Creek near Wales Road and increased levels of most constituents downstream from the landfill. The concentration of arsenic, chromium, iron, and manganese in streambed sediments also is higher downstream from the landfill. Analysis of water-quality samples taken during a period of high overland runoff suggests that surface runoff from the landfill does not significantly affect instream water quality.

Analysis of hydrogeologic, isotopic, and chemical-quality data suggests that the landfill is affecting ground-water flow and quality in the dolomite aquifer. Water-quality changes downgradient from the landfill are limited to increased levels of common ions, trace metals, and selected organic compounds. Additional or greater changes, if present, may not have been detected because of well placement or the nature of ground-water flow characteristics in the lower till and dolomite aquifer.

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WATER-QUALITY DATA

[µS/cm, microsiemens per centimeter at 25 degrees Celsius; mg/L, milligrams per liter; µg/L, micrograms per liter. A dash indicates "data not obtained" or "not applicable." ND indicates constituent not detected. (T), till wells; (M), methane-venting wells.]

Local number	Date of sample	Spe- cific con- duc- tance (µS/cm)	pH (stan- dard units)	Temper- ature, air, (deg. C)	Temper- ature, water (deg. C)	Baro- metric pres- sure (mm of Hg)	Oxygen, dis- solved (mg/L)	Oxygen demand, chem- ical (high level) (mg/L)	Hard- ness (mg/L as CaCO ₃)
WO-101	04-10-84	1,010	7.4	13.0	11.0		<0.1		410
WO-102	0 4-11-84 07-10-85	984 985	7.4 7.6	15.0 20.0	11.5 12.5	 743	<.1 .7	 12	400 420
WO-104	07-10-85	9 50	7.7	20.0	12.5	743	<.1	<10	440
WO-105	04-04-85	860	7.9	12.0	11.5	740	2.8	10	370
WO-106	04-04-85	950	8.9	10.0	11.0	740	2.4	14	340
WO-107	04-05-84 08-01-84 04-03-85 07-10-85	971 912 1,190 1,130	7.8 7.4 7.5 7.6	10.0 35.0 10.0 25.0	11.0 12.5 12.0 12.0	 735 743	1.7 1.2 1.2 1.5	 34 11	350 310 550 430
WO-108	08-01-84	1,260	7.4	35.0	12.0		3.9		440
WO-110	04-05-84 08-02-84 04-04-85	1,040 667 825	11.5 9.4 9.1	10.0 25.0 8.0	11.0 12.0 10.5	 740	1.9 2.4 3.6	 14	260 220 330
WO-111	04-03-85 07-11-85	1,400 1,500	7.9 7.9	12.0	11.0 11.5	740 749	<.1 2.2	10 15	580 580
WO-112	04-05-84 08-01-84 04-03-85 07-11-85	1,560 1,360 1,320 1,440	6.7 6.6 6.8 6.8	10.0 24.0 10.0 25.0	11.0 11.5 11.0 11.5	 735 743	<.1 <.1 <.1 <.1	 18 15	730 580 620 700
WO-113 (T)	04-04-85 07-11-85	1,620 1,800	7.3 7.1	10.0 27.0	6.5 16.5	740 743	10.4 .8	42 48	1,100 1,100
WO-114	04-04-85	925	7.8	8.0	11.0	740	1.0	19	440
wo-115	04-04-84 03-28-85 07-09-85	979 950 940	7.7 7.9 8.1	11.0 15.0 32.0	11.0 12.0 16.0	745 	<.1 <.1 .6	17 <10	230 270 280
WO-117	04-02-85	9 75	7.5	8.0	11.0	745	<.1	<10	490
WO-118	04-11-84 04-02-85	1,530 1,230	7.0 7.4	16.0 10.0	11.0 11.0	 745	<.1 <.1	 14	710 620
WO-119	04-10-84 08-02-84 04-02-85 07-10-85	1,100 1,060 1,050 1,080	7.0 7.0 7.4 7.4	12.0 25.0 8.0 25.0	11.0 12.0 11.0 13.0	 745 743	<.1 <.1 .4 .2	 15 <10	560 510 610 570
WO-120	03-27-85	975	7.7	11.0	11.0	740	.8	14	440
WO-123 (T)	04-04-85 07-11-85	1,850 1,980	7.7 7.7	14.0 22.0	15.0	740 		54 89	580 600
WO-152 (M)	01-29-85 04-03-85	4,950 5,850	6.8 6.7	-4.0 15.0	12.0 19.0	750 735	2.6	390	2,500
WO-154 (M)	01-29-85 04-03-85	28,000 27,800	5.8 5.8	-4.0 12.0	17.0 18.0	750 735	.8	41,000	12,000
Ohio EPA water s	drinking tandards		7.0-10.5						
Laborator detecti	y on limit						.1	10	

Local number	Date of sample	Hard- ness, noncar- bonate (mg/L as CaCO ₃)	Calcium, dis- solved (mg/L) as Ca)	Magne- sium, dis- solved (mg/L as Mg)	Sodium, dis- solved (mg/L as Na)	Potas- sium, dis- solved (mg/L as K)	Alka- linity field (mg/L as CaCO ₃)	Carbon dioxide, dis- solved (mg/L as CO ₂)	Sulfide, total (mg/L as S)
WO-101	04-10-84	290	99	39	56	2.1	119	9.2	ND
WO-102	04-11-84	290	89	37	57	2.2	108	8.3	ND
	07-10-85	320	96	39	61	2.4	105	5.1	.6
WO-104	07-10-85	340	98	41	59	2.1	89	3.4	<.5
WO-105	04-04-85	260	83	33	64	2.0	99	2.4	ND
WO-106	04-04-85	250	80	29	89	2.6	85	.2	ND
107	04-05-84	240	76	22	68	1 0	109	3.3	
WO-107	08-01-84	240	78	33 32		1.9	109	8.3	<.5
	04-03-85	370	140	44	81	3.1	181	11	ND
	07-10-85	270	98	38	81	2.5	157	7.6	ND
WO-108	08-01-84	250	110	40			192	15	
WO-110	04-05-84	35	100	2.0	27	3.6	223	.0	.5
	08-02-84	190	67	13			30	.0	ND
	04-04-85	290	83	25	58	3.4	38	.0	ND
WO-111	04-03-85	500	140	52	130	2.5	79	1.9	ND
	07-11-85	510	140	53	130	2.6	72	1.7	ND
WO-112	04-05-84	230	170	68	69	2.8	492	190	<.5
	08-01-84	170	140	56	~-	~	413	201	
	04-03-85	300	150	55	89	2.8	324	99	ND
	07-11-85	290	170	63	79	2.9	410	126	<.5
WO-113 (T)	04-04-85	800	300	76	14	4.2	266	26	ND
	07-11-85	760	310	79	34	4.5	340	52	.8
WO-114	04-04-85	350	100	41	46	2.1	96	2.9	ND
WO-115	04-04-84	130	56	23	100	2.6	107	4.1	<.5
	03-28-85	160	58	24	110	2.4	106	2.6	ND
	07-09-85	170	61	25	110	2.9	106	1.6	.5
WO-117	04-02-85	360	110	47	44	2.4	128	7.8	ND
WO-118	04-11-84	550	150	82	59	2.3	168	33	ND
WO-110	04-02-85	480	130	63	58	2.3	133	10	ND ND
WO-119	04-10-84	370	120	57	29	2.4	187	36	ND
	08-02-84	330	110	56			175	34	<.5
	04-02-85	440	130	64	32	2.6	174	13	ND
	07-10-85	380	120	60	30	2.6	186	14	<.5
WO-120	03-27-85	320	97	43	48	3.4	114	4.4	ND
NO 100 (T)	04 04 05	200	120	5.0	222	7 0	100	7.0	ND
WO-123 (T)	04-04-85	390 460	130 140	59 59	220 250	7.8 7.4	188 140	7.2 5.4	ND <.5
	07-11-05	400	140	59	250	/.4	140	5.4	<.J
WO-152 (M)	01-29-85 04-03-85	 650	 460	 310	400	 44	1,630	500 695	
	V4-VJ-0J	000	400	210	400	74	1,800	090	1.7
WO-154 (M)	01-29-85 04-03-85	6,100	3,100	 1,100	3,300	1,500	7,190 6,200	22,100 19,000	 130
Ohio EPA o water st	drinking tandards								
Laboratory	Y on limit								.5

Table 2.-Water-quality analyses of ground water near Northwood, Ohio-Continued

Table 2.-Water-quality analyses of ground water near Northwood, Ohio-Continued

Local number	Date <i>o</i> f sample	Sulfate, dis- solved (mg/L as SO4)	Chlo- ride, dis- solved (mg/L as Cl)	Fluo- ride, dis- solved (mg/L as F)	Silica, dis- solved (mg/L as SiO ₂)	Solids, residue at 180 deg. C dis- solved (mg/L)	Solids, sum of constit- uents, dis- solved (mg/L)	Nitrogen, nitrite, dis- solved (mg/L as N)	Nitro- gen, NO ₂ +NO ₃ dis- solved (mg/L as N)
WO-101	04-10-84	430	7.9	1.7	8.1		720	.01	<0.10
WO-102	04-11-84 07-10-85	440 400	17 18	.3 2.0	8.2 8.6	 743	740 710	<.01 <.01	<.10 <.10
WO-104	07-10-85	450	10	1.9	7.4	767	740	<.01	<.10
WO-105	04-04-85	360	14	2.1	7.4	656	650	<.01	<.10
WO-106	04-04-85	400	9.9	1.9	8.7	70 2	690	<.01	<.10
WO-107	04-05-84 08-01-84 04-03-85 07-10-85	390 400 480 420	10 13 12	.2 1.4 1.9	6.7 10 9.0	716 940 806	670 900 780	<.01 <.01 <.01	<.10 16 <.10
WO-108	08-01-84	500				91 2			
WO-110	04-05-84 08-02-84 04-04-85	200 340 390	26 9.9	.2 1.4	3.8 4.8	546 632	500 610	<.01 <.01	<.10 <.10
WO-111	04-03-85 07-11-85	710 740	10 9.2	1.4 1.4	9.9 9.2	1,230 1,250	1,100 1,100	<.01 <.01	<.10 <.10
WO-112	04-05-84 08-01-84 04-03-85 07-11-85	450 330 440 430	17 13 13	.8 1.1 1.0	20 18 20	1,130 1,010 1,110	1,100 980 1,000	<.01 <.01 <.01	<.10 <.10 <.10
WO-113 (T)	04-04-85 07-11-85	740 840	26 35	.2 .3	9.9 16	1,420 1,550	1,300 1,500	<.01 <.01	<.10 <.10
WO-114	04-04-85	390	15	2.0	8.0	695	690	<.05	<.10
WO-115	04-04-84 03-28-85 07-09-85	420 380 380	8.6 8.7 8.1	.2 1.9 1.9	7.5 7.7 8.3	653 702	680 680 680	<.01 .05 <.01	<.10 <.10 <.10
WO-117	04-02-85	410	6.3	1.5	9.0	745	730	<.01	<.10
WO-118	04-11-84 04-02-85	750 570	9.2 9.6	1.7 1.8	8.5 8.3	961	1,200 950	<.01 <.01	<.10 <.10
WO-119	04-10-84 08-02-84 04-02-85 07-10-85	480 460 440 430	3.3 4.4 4.3	1.6 1.6 1.6	11 12 11	892 846 834	840 810 790	<.01 <.01 <.01	<.10 <.10 <.10
WO-120	03-27-85	380	11	1.8	7.4	709	680	.04	<.10
WO-123 (T)	04-04-85 07-11-85	910 890	23 19	.6 .6	10 10	1,550 1,590		.01 <.01	.97 <.10
WO-152 (M)	01-29-85 04-03-85	 17	1,100	.2	28	 3,170	 3,500	<.01	<.10
WO-154 (M)	01-29-85 04-03-85			.4	22	 26,300		.01	.36
Ohio EPA water s	drinking tandards	250	250			500	500	10	
Laborator detecti	y on limit							.01	.1

Table 2.-Water-quality analyses of ground water near Northwood, Ohio-Continued

Local number	Date of sample	Nitrogen, ammonia, dis- solved (mg/L as N)	Nitrogen, ammonia, dis- solved (mg/L as NH4)	Nitrogen, organic, dis- solved (mg/L as N, calc.)	Nitrogen, ammonia + organic dis- solved (mg/L as N)	Alumi- num, dis- solved (µg/L as Al)	Arsenic, dis- solved (µg/L as As)	Boron, dis- solved (µg/L as B)	Cad- mium dis- solved (µg/L as Cd)
WO-101	04-10-84	.38	.49	.72	1.1	<100			
WO-102	04-11-84 07-10-85	.33 .34	.43	2.8 .16	3.1 .50	10	1		1
WO-104	07-10-85	.38	.49	.02	.40				
WO-105	04-04-85	.47	,61	.53	1.0	<10	<1	380	<1
WO-106	04-04-85	.49	.63	.31	.80	10	<1	480	<1
₩O-107	04-05-84 08-01-84 04-03-85 07-10-85	.40 .50 .44	.52 .64 .57	.50 .40 .16	.90 .90 .60	30 10	1 <1	650	<1 16
WO-108	08-01-84	.11							
WO-110	04-05-84	.28	.36	.02	.30	300			
	08-02-84 04-04-85	.31	.40	.39	.70	<10	<1	240	<1
WO-111	04-03-85 07-11-85	- 44 - 40	.57 .52	.26 .10	.70 .50	20	<1	830	<1
WO-112	04-05-84 08-01-84	. 63 . 47	.81 .61	.17	.80	30 10	1 <1	790	<1
	04-03-85 07-11-85	.51	.66	.33 .19	.80 .70				<1
WO-113 (T)	04-04-85 07-11-85	.13 .19	.17 .24	.57 .51	.70 .70	20	<1 	120	<1
WO-114	04-04-85	.38	.49	.42	.80	20	<1	250	<1
WO-115	04-04-84 03-28-85 07-09-85	.40 .30 .49	.52 .39 .63	.00 .30 .11	.40 .60 .60	300 40	1	720	2
WO-117	04-02-85	.41	.53	.39	.80	10	<1	310	<1
WO-118	04-11-84 04-02-85	.42 .39	.54 .50	.08 .61	.50 1.0	100 10	<1	430	<1
WO-119	04-10-84 08-02-84	.45	.58	5.8	6.2	<10	1		<1
	04-02-85 07-10-85	.46 .45	.59 .58	.14 .15	.60 .60	20 	<1 	400	1
WO-120	03-27-85	.30	.39	.30	.60	30	1	310	<1
WO-123 (T)	04-04-85 07-11-85	.30 .63	.39 .81	.60 .37	.90 1.0	80	1 	400	<1
WO-152 (M)	01-29-85 04-03-85	33.0	43	27	 58.0	 50	3	1,900	5
WO-154 (M)	01-29-85 04-03-85	720	 930	230	- - 950	1,500	2	480,000	5
Ohio EPA water s	drinking tandards						50		10
Laboratory detection	y on limit	.10	.10		.10	10	1		1

Local number	Date of sample	Chro- mium, dis- solved (ug/L as Cr)	Iron, dis- solved (ug/L as Fe)	Lead, dis- solved (ug/L as Pb)	Lithium, dis- solved (ug/L as Li)	Manga- nese, dis- solved (ug/L as Mn)	Sele- nium, dis- solved (ug/L as Se)	Stron- tium, dis- solved (ug/L as Sr)	Carbon, organic total (mg/L as C)
WO-101	04-10-84		42			53			1.7
WO-102	04-11-84 07-10-85	10	23 17	6 		1 <1	<1 	20,000 20,000	1.7 .8
WO-104	07-10-85		11			3		19,000	.7
WO-105	04-04-85	<10	71	<1	27	5	<1	20,000	1.1
WO-106	04-04-85	10	10	<1	30		<1	20,000	1.0
WO-107	04-05-84 08-01-84	10	110 80	1		9 7	<1 	20,000	1.3 1.0
	04-03-85 07-10-85	<10 	190 230	3	36	16 11	<1 	21,000 22,000	1.0 .8
WO-108	08-01-84		64			54			1.3
WO-110	04-05-84		12			<1			2.4
	08-02-84 04-04-85	<10	<3 12	 <1	29	<1 4	<1	15,000	1.5 .9
WO-111	04-03-85 07-11-85	<10	120 76	<1 	27 	13 11	<1 	14,000 14,000	1.1 1.1
WO-112	04-05-84 08-01-84	10	2,300 2,100	1		550 320	<1 	18,000	8.7 3.5
	04-03-85 07-11-85	<10 	1,500 2,400	<1 	32	260 330	<1 	18,000 18,000	2.0 2.7
WO-113 (T)	04-04-85 07-11-85	<10	15 1,400	<1	28 	45 260	<1 	650 2,200	12 14
WO-114	04-04-85	<10	51	<1	30	5	<1	23,000	1.1
WO-115	04-04-84		25			2			1.9
	03-28-85 07-09-85	10	91 8	<1	16	2	<1 	19,000 19,000	1.7 1.0
WO-117	04-02-85	<10	170	<1	32	4	<1	23,000	1.3
WO-118	04-11-84 04-02-85	<10	150 110	 <1	 34	5 4	 <1	 28,000	1.7 .9
WO-119	04-10-84	10	64	5		1	<1	20,000	2.7
	08-02-84 04-02-85 07-10-85	<10	68 71 90	<1 	 44 	2 5 2	<1 	21,000 20,000	1.6 .8 .8
WO-120	03-27-85	10	460	<1	21		<1	18,000	1.9
WO-123 (T)	04-04-85 07-11-85	<10	150 60	<1	70	250 390	1	8,800 8,600	7.1 9.8
₩0-152 (M)	01-29-85 04-03-85	 10	13,000	 5	 60	 560	 <1	 25,000	 120
WO-154 (M)	01-29-85 04-03-85	150	 82,000	 65	 1,900	 290	 <1	 38,000	 >790
Ohio EPA water si	drinking tandards	50	300	50		50			
Laborat <i>or</i> detectio	y on limit	10	3	1		1	1		.1

Table 2.-Water-quality analyses of ground water near Northwood, Ohio-Continued

				Methy-	Oil and grease,		
Local number	Date of sample	Cyanide, total (mg/L as Cn)	Phenols, total (µg/L)	lene blue, active sub- stance (mg/L)	total recov- erable gravi- metric (mg/L)	δ D stable isotope ratio (permil)	δ O-18 stable isotope ratio (permil)
wo-101	04-10-84		<1				
WO-102	04-11-84	<.01	<1				
	07-10-85	<.01	<1	.01	1		
wo-104	07-10-85	<.01	5	.02	<1		
wo-105	04-04-85			.04		-77.5	-11.4
WO-106	04-04-85			.04		-68.0	-10.0
vo-107	04-05-84	<.01	<1				
	08-01-84				2		
	04-03-85			.05		-62.5	-9.5
	07-10-85	<.01	4	.02	<1		
WO-108	08-01-84				<1		
wo-110	04-05-84		<1				
	08-02-84				2		
	04-04-85			.05		-67.0	-10.0
10-111	04-03-85			.02		-55.0	-8.6
0-111	07-11-85	<.01	4	.01	<1		
10-112	04-05-84	<.01	<1				
	08-01-84				2		
	04-03-85			.03		-59.0	-9.0
	07-11-85	<.01	<1	.03	<1		
ю -113 (т)	04-04-85			.16		-49.0	-7.6
	07-11-85	<.01	5	.12	2		
10-114	04-04-85			.02		-84.5	-12.2
NO-115	04-04-84		<1				
	03-28-85			.01		-57.0	-9.0
	07-09-85	<.01	1	.07	<1		
10-117	04-02-85			.02		-56.0	-8.4
10-118	04 11 04		-1				
10-118	04-11-84 04-02-85		<1	.04		-77.5	-11.3
vo-119	04-10-84	<.01	<1				
	08-02-84				1		
	04-02-85			.04		-54.5	-8.4
	07-10-85	<.01	<1	.02	<1		
10-120	03-27-85			.02		-72.5	-10.5
۲ ∩−123 / ۳۱	04-04-85			.06		-50.0	-7.8
(0-123 (1)	07-11-85	<.01	2	.19	3		-/.8
	00 0						
10-152 (M)	01-29-85 04-03-85			.45		-49.5 -52.0	-8.3 -8.6
10-154 (M)	01-29-85 04-03-85			2.6		-40.0 -43.0	-9.4 -9.6
hio EPA (water st	drinking Landards	.01	1	.5			
aboratory	/ on limit	.01	1	.01	1		

Table 2.-Water-quality analyses of ground water near Northwood, Ohio-Continued

<pre>[deg., degrees; mg/L, milligrams per liter; pg/L, micrograms per liter; ps/cm, microsiemens per centimeter at 25 degrees Celsius; recov., recoverable; bot. mat., bottom material; ac-ft, acre-foot; nd, not detected. Dash indicates no data available.]</pre>	µg/L, micro bottom mate	<pre>µg/L, micrograms per liter; µS/cm, microsiemens per bottom material; ac-ft, acre-foot; nd, not detected.</pre>	ter; ps/cm acre-foot	, micros ; nd, nc	tiemens per t detected	centimet . Dash j	sntimeter at 25 degrees Dash indicates no data	degrees no data
Station name	Site desig- nation	Date	Temper ature, water (deg. C		Temper- ature, air (deg. C)	Baro- metric pres- sure (mm of Hg)	ft ir fg	Stream- flow, instan- taneous (ft ³ /s)
Otter Creek tributary at Wales Road near Northwood, Ohio	æ	04-09-84 03-28-85 03-28-85 03-28-85 03-28-85 03-28-85	8.0 12.0 12.0 12.0 27.0		5.0 15.0 15.0 15.0 30.5	 745 745 	•	5
Otter Creek tributary at Whitmore Yard near Northwood, Ohio	ф	04-09-84 07-09-85	10.5 29.5		17.0 29.0		·	 .05
Dry Creek at E. Broadway near Walbridge, Ohio	υ	04-10-84 07-09-85	4.5 24.5		14.0 29.0		·	- 2
			Spe- cific con- duc- tance, field (µS/cm)	Oxygen, dis- solved (mg/L)	Oxygen demand, chem- ical, high (mg/L)	pH, field (stan- dard units)	pH, lab (stan- dard units)	Carbon dioxide, dis- solved (mg/L as CO2)
Otter Creek tributary at Wales Road near Northwood, Ohio	A	04-09-84 03-28-85 03-28-85 03-28-85 03-28-85 07-09-85	797 480 550 499	11.9 	45	8.0 7.9 7.9 8.0	7.9 7.3 7.5 8.0	4.7 2.0 2.6
Otter Creek tributary at Whitmore Yard near Northwood, Ohio	£	04-09-84 07-09-85	649 365	19.4 7.3		8.0 9.3	7.9 9.2	4.2
Dry Creek at E. Broadway near Walbridge, Ohio	υ	04-10-84 07-09-85	685 744	15.5 6.1	 19	8.10 7.90	7.9 8.0	3.1 4.4

Table 10.-Analyses of surface-water quality and sediment near Northwood, Ohio

Table 10Analyses of surface-water quality and sediment near Northwood, Ohio-Continued	f surface-w	ater quality	r and sedir	ment near	Northwood	, Ohio-Cor	ntinued		
Station name	Site desig- nation	Date	Alka- linity, total, field (mg/L as CaCO3)	Oil and grease, total recov., gravi- metric (mg/L)	Nitro- gen dis- solved N)	Nitro- gen, organic, dis- solved (mg/L as N)	Nitro- gen, ammonia, dis- solved (mg/L as N)	Nitro- gen, , nitrite, dis- solved (mg/L as N)	i î T
Otter Creek tributary at Wales Road near Northwood, Ohio	R	04-09-84 03-28-85 03-28-85 03-28-85 03-28-85 03-28-85 03-29-85	246 77 81 80 134	I I I I ⁴	1.4 1.1	0.19 .72 	0.21 .28 .30	0.01 .06 .03	
Otter Creek tributary at Whitmore Yard near Northwood, Ohio	щ	04-09-84 07-09-85	216 112	↓	1.5 .99	.19 .64	.21	.01.03	-1 -0
Dry Creek at E. Broadway near Walbridge, Ohio	υ	04-10-84 07-09-85	203 181	↓	3.1 1.2	.33	.07	.03	m 01
			Nitro- gen, nitrate, dis- solved d (mg/L as N)	Nitro- gen, am- monia + 1 organic, dissolved (mg/L as N)	Nitro- gen, NO2+NO3, dis- solved (mg/L as N) a	Phos- phate, ortho, p dis- solved (mg/L as PO4)	Phos- phorus, dis- solved (mg/L as P)	Phos- phorus, Ortho, dis- solved (mg/L as P)	Carbon, organic, total (mg/L as C)
Otter Creek tributary at Wales Road near Northwood, Ohio	Ł	04-09-84 03-28-85 03-28-85 03-28-85 03-28-85 03-28-85	0.99 1.14 	0.4 	1.00 1.20 -1.20		0.0	<0.01 08 	7.1 5.9 10
Otter Creek tributary at Whitmire Yard near Northwood, Ohio	щ	04-09-84 07-09-85		. 4	1.10 .29		.01	- 10 - 1	 8.9
Dry Creek at E. Broadway near Walbridge, Ohio	υ	04-10-84 07-09-85	2.67 .76	4.	2.70 .78		.02	10. ×	4.5 4.2

Station name	Site desig- nation	Date	Carbon, organic, dis- solved (mg/L as C)	Carbon, organic, sus- pended, total (mg/L as C)	Cyanide, total (mg/L as CN)	Cyanide, total, in bot- tom ma- terial (ug/g as CN)	Sulfide, total (mg/L as S)	Hard- ness, (mg/L as CaCO3)	Hard- ness, noncar- bonate, field (mg/L as CaCO3)
Otter Creek tributary at Wales Road near Northwood, Ohio	A	04-09-84 03-28-85 03-28-85 03-28-85 03-28-85 03-28-85	7.1	0.5	· · · · · · · · · · · · · · · · · · ·	<0.5 	nd 	380 180 200 190 210	130 100 120 110 80
Otter Creek tributary at Whitmore Yard near Northwood, Ohio	В	04-09-84 07-09-85	5.9	۳.	01	<. 5 -	nd < .5	310 170	94 56
Dry Creek at E. Broadway near Walbridge, Ohio	υ	04-10-84 07-09-85	4.5	· 5	<	ہ ۲.5	nd 5	290 340	87 160
			Calcium, dis- solved (mg/L as Ca)	Magne- sium, dis- solved (mg/L as Mg)	Sodium, dis- solved (mg/L as Na)	Sodium ad- sorp- tion ratio	Percent sodium	Potas- sium, dis- solved (mg/L as K)	Chlo- ride, dis- solved (mg/L as Cl)
Otter Creek tributary at Wales Road near Northwood, Ohio	R	04-09-84 03-28-85 03-28-85 03-28-85 03-28-85 03-28-85	97 53 52 57	33 15 17 17	21 33 20 20 20	1.5 .5 .6	11 29 14 17	3.6 3.7 4.1	37 34 38 33 37
Otter Creek tributary at Whitmore Yard near Northwood, Ohio	В	04-09-84 07-09-85	83 47	25 12	15 13	4. S.	1 9 1	2.8 3.5	32 26
Dry Creek at E. Broadway near Walbridge, Ohio	υ	04-10-84 07-09-85	80 67	22 39	28 21	.5	17 12	1.9 7.6	57 45

Table 10.-Analyses of surface-water quality and sediment near Northwood, Ohio-Continued

		•							
Station name	Site desig- nation	Date	Sulfate, dis- solved (mg/L as SO4)	Fluo- ride, dis- solved (mg/L as F)	Silica, dis- solved (mg/L as SiO2)	Arsenic, dís- solved (µg/L as As)	Arsenic, total, in bot- tom ma- terial (µg/g as As)	Beryl- lium, recov. from bot- tom ma- terial (µg/g as Be)	- Boron, dis- solved (µg/L as B)
Otter Creek tributary at Wales Road near Northwood, Ohio	A	04-09-84 03-28-85 03-28-85 03-28-85 03-28-85 07-09-85	150 120 120 120 73	0.3 .3 .4	5.4 8.8 6.8 6.6	~		↓ 10 10	110
Otter Creek tributary at Whitmore Yard near Northwood, Ohio	щ	04-09-84 07-09-85	110 47		5.7 7.6		FI 	<10 	
Dry Creek at E. Broadway near Walbridge, Ohio	υ	04-10-84 07-09-85	84 150	. 2	3.6 9.4		-	<10	} }
			Cadmium, dis- solved (µg/L as Cd)	Cadmium, recov. from bot- tom ma- terial (µg/g as Cd)	Chro- mium, - recov. from bot- terial (µg/g)	Chro- mium, dis- solved (µg/L as Cr)	Copper, recov. from bot- tom ma- terial (µg/g as Cu)	Iron, dis- solved (µg/L as Fe)	Lead, dis- solved (ug/L as Pb)
Otter Creek tributary at Wales Road near Northwood, Ohio	Ą	04-09-84 03-28-85 03-28-85 03-28-85 03-28-85 07-09-85	4	4	∾	10	⊷	14 14 6	^
Otter Creek tributary at Whitmore Yard near Northwood, Ohio	ш	04-09-84 07-09-85		4	1 1	11	H	30 55	
Dry Creek at E. Broadway near Walbridge, Ohio	υ	04-10-84 07-09-85		↓	4	; ;	4	37 22	

Table 10.-Analyses of surface-water quality and sediment near Northwood, Ohio-Continued

Station name	Site desig- nation	Date	Lead, recov. from bot- tom ma- terial (ug/g as Pb)	Manga- - nese, dis- solved (µg/L as Mn)	Nickel, recov. from bot tom ma- terial (µg/g as Ni)	Stron- t- tium, dis- solved (µg/L as Sr)	Zinc, recov. from bot- tom ma- terial (µg/g) as Zn)	Alum- t- inum, dis- solved (µg/L as Al)	Lithium, dis- solved (µg/L as Li)
Otter Creek tributary at Wales Road near Northwood, Ohio	A	04-09-84 03-28-85 03-28-85 03-28-85 03-28-85 07-09-85	10 1	150 41 44 170	0 1 1 1 1 1 1	 1900 1800	√	<pre><100 </pre>	
Otter Creek tributary at Whitmore Yard near Northwood, Ohio	щ	04-09-84 07-09-85	<10	49 12	<10	1100	↓	100	
Dry Creek at E. Broadway near Walbridge, Ohio	υ	04-10-84 07-09-85	<pre>10</pre>	45 58	<10	 10000	↓	100	
			Sele- nium, dís- solved (µg/L as Se)	Sele- nium, total in bot- tom ma- I terial (ug/g)	Phenols, total b (ug/L)	Benzo b fluor- an- 1 thene bot. mat 1 (ug/kg)	Fluor- anthene, bot. mat. (ug/kg)	Phenan- threne, bot. mat. (µg/kg)	Pyrene, bot. mat. (µg/kg)
Otter Creek tributary at Wales Road near Northwood, Ohio	Ą	04-09-84 03-28-85 03-28-85 03-28-85 03-28-85 07-09-85	4	4	Ų 4	<10.0	100	110 	70.0
Otter Creek tributary at Whitmore Yard near Northwood, Ohio	В	04-09-84 07-09-85		4 ∣	1 ۲	180 	68.0 	<10.0	0.06
Dry Creek at E. Broadway near Walbridge, Ohio	υ	04-10-84 07-09-85		₽	4	<10.0	<10.0	<10.0	<10.0

τ 2 2 --

Station name	Site desig- nation	Date	Benzo a anthra- cene 1,2- benzan- thracene, bot. mat. (µg/kg)	4	Phenol (C6H- 5OH) total (µg/L)	Methy- lene blue active sub- stances (mg/L)
Otter Creek tributary at Wales Road near Northwood, Ohio	A	04-09-84 03-28-85 03-28-85 03-28-85 03-28-85 03-28-85	<10.0 		1.0	
Otter Creek tributary at Whitmore Yard near Northwood, Ohio	В	04-09-84 07-09-85	170		<1.0 	
Dry Creek at E. Broadway near Walbridge, Ohio	υ	0 4 -10-84 07-09-85	<10.0		<1.0 	
			Solids, Sc residue su at 180 cc deg. C, tu dis- solved so (mg/L) (r	Solids, sum of consti- tuents, dis- solved (mg/L)	Solids, dis- solved (tons per day)	Solids, dis- solved (tons per ac-ft)
Otter Creek tributary at Wales Road near Northwood, Ohio	đ	04-09-84 03-28-85 03-28-85 03-28-85 03-28-85 07-09-85	 331 315	500 310 300 300	0.06	0.67 .42 .45 .41
Otter Creek tributary at Whitmore Yard near Northwood, Ohio	В	04-09-84 07-09-85	 242	400 230	 0.03	.55
Dry Creek at E. Broadway near Walbridge, Ohio	υ	04-10-84 07-09-85	 508	400 460	0.27	. 69

Table 10.-Analyses of surface-water quality and sediment near Northwood, Ohio-Continued

			Mercury,		
			recov.	8 D	δ 0-18
			from bot-	stable	stable
			tom ma-	isotope	isotope
	Site		terial	ratio	ratio
	desig-		6/6n)	(per-	(per-
Station, name	nation	Date	as Hg)	(lim	(lim
Otter Creek tributary at Wales Road near	A	04-09-84	1.1	ł	ł
Northwood, Ohio		03-28-85	}		ł
		03-28-85	1	-45.0	-7.2
		03-28-85	ł		ł
		07-09-85	1	ł	
Otter Creek tributary at Whitmore Yard	В	04-09-84	.84	ł	ł
near Northwood, Ohio		07-09-85	}	!	ł
Dry Creek at E. Broadway near Walbridge,	U	04-10-84	2.4	1	ł
Chio		07-09-85	1	ļ	ł

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