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Effects of Land Use on Quality of Water in Stratified-Drift Aquifers in Connecticut

U.S. GEOLOGICAL SURVEY Open-File Report 91-200

Prepared as part of the
U.S. Geological Survey
Toxic-Waste--Ground-Water Contamination Program



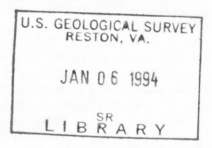


EFFECTS OF LAND USE ON QUALITY OF WATER IN STRATIFIED-DRIFT AQUIFERS IN CONNECTICUT

by Stephen J. Grady

U.S. GEOLOGICAL SURVEY

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U.S. DEPARTMENT OF THE INTERIOR

BRUCE BABBITT, Secretary

U.S. GEOLOGICAL SURVEY

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CONVERSION FACTORS AND VERTICAL DATUM

Multiply	Ву	To obtain
	Length	
inch (in.)	25.4 0.3048	millimeter
<pre>foot (ft) mile (mi)</pre>	1.609	meter kilometer
	Area	
acre square mile (mi²)	4,047 2.590	square meter square kilometer
	Flow	
inch per year (in/yr) foot per day (ft/d)	25.4 0.3048	millimeter per year meter per day
	Hydraulic Conductivit	Y
foot per day (ft/d)	0.3048	meter per day

Temperature in degrees Celsius (°C) can be converted to degrees Fahrenheit (°F) as follows:

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

Sea level--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.

IN CONNECTICUT

by Stephen J. Grady

ABSTRACT

Human activities associated with agricultural, residential, commercial and industrial land uses have affected the quality of water in the four stratified-drift aquifers examined in Connecticut. A study to evaluate quantitatively the effects of human activities, expressed as land use, on regional ground-water quality was initiated in 1984 as part of the U.S. Geological Survey's Toxic-Waste--Ground-Water Contamination Program. Water-quality data were collected from 116 shallow, stainless-steel wells installed beneath or immediately downgradient from seven types of land use within the Pootatuck, Pomperaug, Farmington, and Hockanum River valleys in Connecticut. Analysis of variance on the ranked concentrations of 21 largely uncensored or slightly censored constituents, and contingency-table analysis of the frequency of detection of 49 moderately to highly censored constituents indicate that 27 water-quality variables differ at the 0.05 level of significance for samples from at least one land-use area.

For most constituents, concentrations or detection frequencies are lowest in samples from the undeveloped areas, which characterize background water-quality conditions. The effect of agricultural land use on ground-water quality reflects tillage practices; tilled areas affect the water quality to a greater degree than do untilled areas. Twenty percent of the wells in the tilled agricultural areas yielded water with concentrations of nitrate plus nitrite nitrogen exceeding 10 milligrams per liter. Atrazine detections in one-third of the wells in areas of tilled agricultural land use were significantly more common than in the undeveloped areas.

Ground-water quality beneath sewered residential areas is more severely affected by inorganic and organic nonpoint-source contaminants than is water quality beneath unsewered residential areas. Median concentrations or detection frequencies of most physical properties and inorganic constituents of ground water are higher in sewered than in unsewered residential areas. Generally low concentrations (less than 1.0 microgram per liter) of one or more of 17 volatile organic compounds were detected in samples from 62 percent of the wells in the unsewered residential areas. Most of these compounds were detected in less than 10 percent of the ground-water samples from the unsewered residential areas, however, and consequently, their frequency of detections was not significantly different than in samples from other land-use areas. The detection of chloroform in ground-water samples from 47 percent of the wells in the sewered residential areas is significantly higher than the frequency of detection of chloroform in samples from the undeveloped, tilled agricultural, and unsewered residential areas.

The quality of ground water is adversely affected beneath commercial areas more so than beneath all other land-use areas. Median concentrations of sodium (22.5 milligrams per liter), chloride (36 milligrams per liter), and dissolved solids (286 milligrams per liter) are highest in ground-water samples in commercial areas. Detections of tetrachloroethylene, trichloroethylene, and 1,2-transdichloroethylene were significantly more common in ground-water samples from the commercial areas than in samples from one or more of the other land-use areas. Tetrachloroethylene was detected in water samples from 50 percent of the observation wells in the commercial areas at concentrations of up to 1,300 micrograms per liter. Trichloroethylene and 1,2-transdichloroethylene were found at concentrations of up to 20 and 55 micrograms per liter, respectively, in samples from more than 40 percent of the wells in the commercial areas.

Although industrial areas occupy only a small part of each of the study areas, they have a disproportionately large effect on ground-water quality. One or more of 12 volatile organic compounds were detected in water samples from 91 percent of the observation wells in the industrial areas. Chloroform, the most commonly detected compound, was found at concentrations of 0.2 to 1.3 micrograms per liter in samples from more than half of the industrial-area wells. Thirty-six percent of these wells yielded water with detectable concentrations of tetrachloroethylene (0.2 to 4.9 micrograms per liter) and 95 percent contained 1,1,1-trichloroethane (0.2 to 180 micrograms per liter). The frequency of 1,1,1-trichloroethane, 1,1-dichloroethane, and 1,1-dichloroethylene detections at concentrations equal to or exceeding 1.0 microgram per liter were significantly higher for samples from the industrial areas than for samples from one of more of the other land-use areas.

INTRODUCTION

Ground water is a vital resource for the Nation, supplying more than half of the population with drinking water, but its quality is being degraded by an increasing number and variety of contaminants. Incidents of ground-water contamination by toxic chemicals have been widely publicized, heightening concern for the protection of ground-water supplies. At present, only a very small part of the Nation's total ground-water resources is known to be contaminated. There are insufficient data, however, particularly about the occurrence of trace elements and synthetic organic chemicals, to assess adequately the full extent of ground-water contamination within the Nation's aguifers.

In 1984, the U.S. Geological Survey's Toxic-Waste--Ground-Water Contamination Program initiated studies of the effects of human activities on the quality of ground water in 14 regions of the Nation (Ragone, 1984). Each study has endeavored to relate present ground-water quality to the hydrologic, climatic, and land-use characteristics of that region and, by so doing, provides a basis for appraising ground-water quality in other areas of the Nation that have similar characteristics (Helsel and Ragone, 1984). The study in Connecticut has focused on unconsolidated glacial sand and gravel (stratified-drift) aquifers that are among the most productive sources of public water supply in the glaciated northeastern United States (fig. 1).

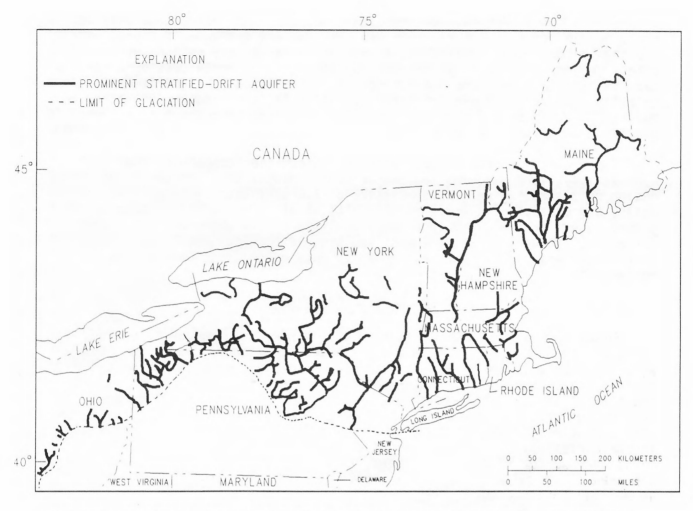


Figure 1.--Prominent stratified-drift aquifers in the northeastern United States. (Modified from McGuinness, 1964.)

Stratified-drift aquifers that occupy many of the principal river valleys of Connecticut and the region are highly susceptible to contamination from human activities at the land surface. The same hydrologic characteristics that enable these aquifers to yield large quantities of water also facilitate the entry and migration of contaminants. These characteristics include a generally shallow water table, high permeability of predominantly coarse-grained sediments, and hydraulic connection with surface-water bodies.

Purpose and Scope

This report presents the final results of the study to determine the effects of land-use activities on the quality of the water in four stratified-drift aquifers in Connecticut. The quality of ground water in the aquifers is described, with emphasis on trace elements and synthetic organic compounds that have anthropogenic sources. The water chemistry is explained in terms of natural hydrogeologic factors and human factors. The hypothesis that human activities, as defined by a variety of land uses, affect ground-water quality is tested by statistical comparisons of water quality beneath seven land-use categories. Statistically significant relations between land use (or other causative factors) and ground-water quality that may be transferable to similar settings throughout the glaciated Northeast are identified.

The report includes a compilation of information concerning the hydrogeology, land use, and sources of contaminants; description of the design of a sampling network consisting of 116 stainless-steel observation wells, and a sample-collection protocol to obtain representative samples of ground water for field and laboratory analyses of 179 water-quality variables, including trace elements and organic chemical compounds; and a statistical analysis of the water-quality data using distribution-free, nonparametric statistical methods.

Previous Investigations

Grady and Weaver (1988) presented preliminary findings of this study based on selected historical data and limited sampling in two of the four aquifers studied. Their report described the methods used to define the hydrogeology, land use, and water quality of stratified-drift aquifers and the statistical methods used to evaluate relations between these factors. Information presented by Grady and Weaver (1988) was included in a review by Cain and others (1989) of the preliminary results of the regional nonpoint-source contamination studies conducted in New York (Eckhardt and others, 1989), New Jersey (Barton and others, 1987), Florida (Rutledge, 1987), Nebraska (Chen and Druliner, 1987), and Colorado (Cain and Edelmann, 1986) as well as Connecticut.

Grady (1989) and Grady and Weaver (1989) presented interim results of the study in Connecticut based on data collected in all four aquifers through 1988. Most of the preliminary and interim findings presented in the above reports are confirmed in this report; however, the additional data available at this time allow the author to refine the land-use categories previously used.

<u>Acknowledgments</u>

The author wishes to acknowledge the cooperation, information, and assistance provided by public officials of the Towns of Avon, Ellington, Farmington, Granby, Manchester, Newtown, Plainville, Simsbury, Southbury, South Windsor, Vernon, and Woodbury that greatly facilitated the successful completion of this study. In addition, access was provided by the Connecticut Departments of Environmental Protection and Transportation to public lands for data collection. The author is also grateful to the numerous private citizens, corporations, and utilities that allowed access to their properties for data collection.

APPROACH

The regional nonpoint-source contamination studies followed a common five-step approach developed by Helsel and Ragone (1984) that included (1) reconnaissance, (2) experimental design, (3) data collection, (4) data analysis, and (5) verification of results. During the reconnaissance phase of the study, all pertinent hydrogeologic, land use, and water-quality information on the study areas was compiled and analyzed, and limited additional water-quality data were collected. Hypothetical relations among land use, hydrogeology, and water quality were proposed, and preliminary statistical comparisons of the water-quality data associated with different land uses were made. Grady and Weaver (1988) described the information obtained and the preliminary findings of the reconnaissance phase of the Connecticut study.

Information obtained during the reconnaissance phase directed the manner in which additional data were collected to test the hypothesis that human activities have a predictable, quantifiable effect on ground-water quality. Testing the hypothesis required (1) accurate characterization of hydrologic and land-use conditions, (2) comprehensive chemical analyses of representative ground-water samples, and (3) valid statistical analysis.

DESCRIPTION OF STUDY AREAS

Four stratified-drift aquifers were selected for this study because their hydrogeologic characteristics and state of development are typical of glacial-valley aquifers in Connecticut as well as much of the region. The location of the stratified-drift aquifers in the Pootatuck, Pomperaug, Farmington, and Hockanum River valleys is shown in figure 2.

Hydrogeologic Characteristics of Stratified-Drift Aquifers

A wealth of information is available on the hydrogeologic characteristics of stratified-drift aquifers in the northeastern United States because many of these aquifers have been investigated for potential water supplies. Typically, these studies have focused on detailed definition of boundaries and hydraulic properties of the stratified-drift aquifers. Hydrogeologic information was obtained through test drilling, seismic-refraction surveys, aquifer tests, and ground-water-flow models. Currently, the U.S. Geological Survey's Regional Aquifer-System Analysis Program is studying the hydrologic characteristics of glacial-valley aquifers in the northeastern United States (Lyford and others, 1984). Recent geological studies (Koteff and Pessl, 1981; Stone and Borns, 1986; Warren and Stone, 1986) have advanced the understanding of the deposition and sedimentology of glacial meltwater deposits that in turn control hydrologic characteristics.

Stratified-drift sediments--interbedded layers of generally well-sorted gravel, sand, silt and clay--were deposited in most of the principal valleys of the northeastern United States by meltwaters from continental glaciers that covered the region during the Pleistocene Epoch. The origin and depositional environment of these sediments strongly influence aquifer geometry, and thus, the aquifers are generally elongate, narrow, and relatively thin (fig. 3). Variations in the thickness and texture of stratified drift resulting from the pattern of deglaciation and topography of the land disrupt the continuity of the stratified-drift aquifers.

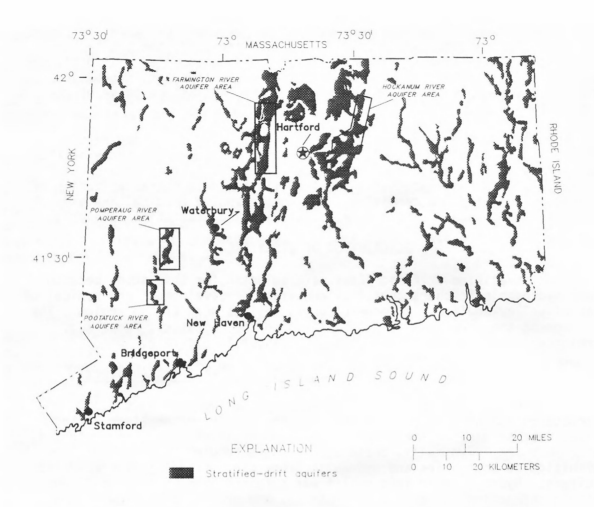


Figure 2.--Location of four stratified-drift aquifers studied in Connecticut.

Consequently, most major river valleys contain several individual aquifers that extend from a few hundred feet to several miles in length. The width of these aquifers is generally limited to a few thousand feet or less, and their maximum saturated thickness is generally less than 100 ft (feet). Stratified-drift aquifers underlie approximately 7.9 mi² (square miles) of the Pootatuck River valley, 12.7 mi² of the Pomperaug River valley, 21.5 mi² of the Hockanum River valley, and 87.6 mi² of the Farmington River valley.

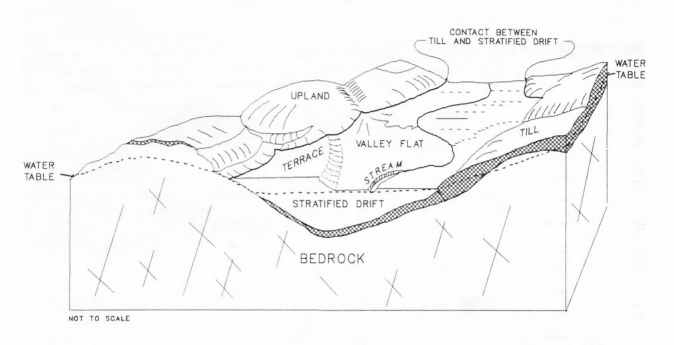


Figure 3.--Idealized three-dimensional geometry of a typical glacial-valley, stratified-drift aquifer. (Modified from Frimpter, 1981, fig. 1.)

Stratified-drift deposits commonly exhibit abrupt horizontal and vertical changes in texture that reflect their complex depositional environment. However, except for some glaciolacustrine deposits, the stratified drift contains relatively little silt and clay, and the bulk of the aquifers consist of sand and gravel. Hydraulic conductivity primarily reflects the grain size and degree of sorting of the individual layers, and ranges from less than 1 to greater than 10,000 ft/d (feet per day) for stratified drift in the northeastern United States (Lyford and others, 1984, p. 12). Average horizontal hydraulic conductivity values in the range of 20 to 170 ft/d are typical of the four aquifers studied in Connecticut (Haeni, 1978; Ryder and others, 1981: Handman and others, 1986; Mazzaferro, 1986).

Ground water in stratified-drift aquifers occurs mostly under unconfined conditions, and the water table is generally only a few feet to a few tens of feet below land surface. The mean depth to the water table measured in the 116 observation wells installed for this study was 11.2 ft. Natural recharge occurs directly through infiltration of precipitation upon the land surface above the aquifer, but more than half of the recharge to stratified-drift aquifers may be derived from upland runoff (Morrissey and others, 1988). The amount of recharge under natural (nonpumping) conditions averages about 25 in./yr (inches per year) for stratified-drift aquifers studied in Connecticut (Haeni, 1978; Mazzaferro, 1986). Seasonal variations in precipitation and evapotranspiration cause the depth to the water table to fluctuate from a few feet to more than 10 ft within the stratified-drift aquifers (fig. 4).

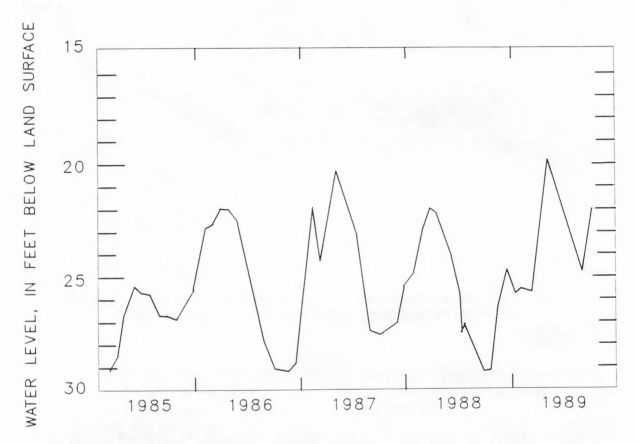


Figure 4.--Water levels in an observation well in the stratified-drift aguifer within the Pomperaug River valley, 1985-89.

Ground water is in constant movement from areas of high water-table altitudes which typically occur along the margins of the valleys, toward areas of lower water-table altitudes that generally coincide with major streams, lakes, and wetlands within the central portions of the valleys. The rate of flow depends on the hydraulic gradient, and the porosity and hydraulic conductivity of the aguifer. Grady and Weaver (1988, p. 13) have estimated the average linear velocity of ground-water flow from aguifer properties and hydraulic gradients in two stratified-drift aguifers in Connecticut. Values ranged from 0.01 to 10 ft/d; however, average linear velocities of 1 to 3 ft/d were considered to be most representative (Grady and Weaver, 1988, p. 13). Comparable velocity values of 0.3 to 1.3 ft/d were measured in natural-gradient tracer tests in glacial-outwash aguifers at Cape Cod, Massachusetts (LeBlanc, 1987, p. B-3) and Borden, Ontario. Canada (Mackay and others, 1986, p. 2027). Based upon these estimates of ground-water velocity, the residence time of water in stratified-drift aquifers must be brief, as the flowpaths are commonly only a few hundred to a few thousand feet in length. Consequently, the residence time for most ground water in stratified-drift aquifers is likely to be less than 1 year to 2 decades (Grady and Weaver, 1988, p. 13-14).

The short temporal and spatial scales of the ground-water-flow systems in most stratified-drift, glacial-valley aquifers strongly influenced the design of the water-quality sampling network. For those contaminants that move through the aquifer at the same rate as the water, the effects of only recent land-use practices should be observed. Consequently, evaluation of the current pattern and recent history of land-use development within the study areas was necessary. Land-use information also was compiled at a scale comparable to the hydrogeologic scale in order to avoid misrep-resenting possible associations between ground-water quality and land use.

Land Use

Information on the existing land use and recent changes in land use were compiled for each of the four aquifer areas. The 1970 statewide landuse inventory (Connecticut Office of Policy and Management, 1970) was the principal source used for this study. Identified and mapped from 1:12,000 scale aerial photography flown in March 1970, the statewide land-use inventory was compiled at a scale of 1:24,000 and therefore provides a level of resolution suitable for comparison with the hydrogeologic information. The 1970 land-use inventory, outdated in some areas of the State, was updated using more recent aerial photography, local land-use and zoning information, and field verification.

Sixty-nine land-use categories were mapped in the 1970 Connecticut land-use inventory that correspond to Level III and IV categories of Anderson and others (1976). Reconnaissance-phase and interim-study evaluations (Grady and Weaver, 1988; 1989; and Grady, 1989) used only four broadly defined land-use categories--undeveloped, agricultural, residential, and mixed industrial plus commercial--so that statistical comparisons of ground-water quality could be made from the small number of samples. Sufficient data were collected during the balance of the study to refine the land-use classifications to include seven categories: undeveloped, untilled agricultural, tilled agricultural, unsewered residential, sewered residential, commercial, and industrial.

The undeveloped land-use category consists primarily of mixed deciduous woodlands that are generally second-growth forest on fields that were once cleared for agriculture but have been abandoned for at least the past 50 years. This category also includes all inactive, unused, unproductive, or abandoned open lands and areas cleared for construction prior to development, plus wetlands and surface-water bodies. In Connecticut, the most extensive tracts of undeveloped land are frequently in the upland areas beyond the stratified-drift boundaries; however, in the study areas, some undeveloped land has been maintained as public reservoir watershed, parks, and wildlife preserves, or as private woodland recreational areas. Areas that are predominantly undeveloped may encompass some minor discordant land uses such as low-density residential development, secondary roads, and small agricultural plots, but the quality of ground water beneath such areas is assumed to be largely unaffected by human activities. Undeveloped areas. therefore, provide background water-quality conditions with which to compare and evaluate the effects from human activities within other land-use categories.

Untilled agricultural land use includes alfalfa and hay production, pasture lands, ornamental shrub nurseries, and turf farms. Also included in this category are recreational athletic fields and golf courses that, although not truly an agricultural use, receive applications of fertilizers, soil conditioners, and pesticides. The principal criteria for inclusion within this land-use category, therefore, was that the land was used for the active production of some type of cover (hay, grass, shrubs) without any significant mechanical disturbance of the natural soil profile, and the probable (though not documented) use of agricultural chemicals. The most common chemical applications to untilled agricultural lands include nitrogen and phosphorous fertilizers; potash, borax, and lime for soil treatments; and a variety of pesticides (predominantly herbicides).

Tilled agricultural lands are primarily cultivated for silage and sweet-corn production. Some 73 percent of the active agricultural land in Connecticut is in corn production; other row crops include strawberries, tobacco, and a variety of vegetables. Little of the cultivated farmland is irrigated, and, where irrigation is used, the source of water is most commonly a nearby surface-water body. Agricultural chemicals that are applied to tilled acreage include organic (manure) as well as inorganic fertilizers, soil treatments and micronutrients, and pesticides. The most common pesticides used on corn are the preemergent, broadleaf weed herbicides, atrazine and alachlor.

Residential land use encompasses any area of 5 or more acres where housing structures of any type predominate. The category includes a range of housing densities from high-density urban areas with more than 8 dwellings per acre to single, rural residences. Because population-density data are available only for census tracts that, in the study areas, include entire towns or large parts of the towns, it was not possible to subdivide the residential land-use category by this criterion. However, information on which tracts of residential development are served by sanitary-sewer lines and which tracts utilize individual septic systems was readily available. Consequently, the residential land-use category has been subdivided into unsewered residential and sewered residential categories. Although there is overlap in the housing densities included in each of the two categories, the unsewered residential areas tend to have larger lot

sizes (generally greater than 1 acre), are commonly underlain by coarse-grained stratified drift, and underlie areas where depth to the water table is greater. The sewered residential areas, conversely, tend to have higher housing densities, are more likely to include commercial structures within the area mapped as residential, and are more common where fine-grained materials or shallow depths to the water table predominate.

A variety of potential sources of contaminants are associated with unsewered and sewered residential areas. A prominent source of inorganic and organic contaminants in unsewered areas are the individual septic systems that receive domestic wastes, wastewaters, and discarded household chemicals. Exfiltration from leaky sanitary sewer lines may provide a source for similar contaminants within the sewered residential areas. Other contaminant sources associated with residential areas include the infiltration of street runoff carrying deicing chemicals, metals, and hydrocarbons. Use of lawn and garden chemicals and wastes from domestic animals provide sources for additional nutrients and pesticides. Leaky household fuel tanks may contaminate ground water with fuel oil or gasoline.

Commercial land use in the study areas consists of a variety of trades and services including gas stations and automobile repair shops, dry cleaners, printers and photographic finishers, restaurants, retail outlets, and offices. Commercial areas have been mapped as such when they extend over an area of 5 acres or more. Generally, they fall within three characteristic patterns: (1) the central business section or downtown areas of a city or town; (2) shopping centers, frequently in outlying areas; and (3) commercial strip developments along major highways or streets. Much of the land surface in commercial land-use areas is impervious--either covered by the buildings themselves, parking lots, storage "yards", and the network of streets and roads that facilitate vehicular access. A variety of nonpoint-source contaminants may originate from commercial areas as a result of leaks from gasoline, fuel oil, or chemical storage tanks; inadvertent leaks and spills or improper disposal of chemicals and wastes; infiltration of street and parking lot runoff; disposal of wastes and wastewaters to onsite septic systems in unsewered commercial areas; and exfiltration from sanitary sewers in areas served by municipal wastewater-treatment systems.

The industrial land-use category includes a complex mixture of individual types of industrial facilities that can collectively be described as "light" manufacturing. The term light manufacturing is used in the context of industries devoted to designing, assembling, finishing, and packaging of products rather than "heavy" manufacturing that involves the extraction and processing of basic or raw materials or producing heavy machinery. In Connecticut, light manufacturing largely includes small, modern (often high-technology) facilities for the fabrication of electronic equipment and components (motors and generators; switchgear and controls; electric housewares, and communication equipment; lighting and wiring; semiconductors, capacitors, coils and transformers); metal products (cutlery, hand tools, and hardware; plumbing and heating equipment; springs, valves, and wire products); metal plating or polishing; textile and apparel; lumber and wood products; chemicals and allied products; and food processing, warehousing and distribution. Individual industrial facilities or, more commonly, industrial parks have been mapped where they extend over areas of 5 acres or more. These facilities include office, warehouse, plant and laboratory buildings, parking and storage areas, ancillary facilities, roadways, and open areas.

Ground-water contamination has frequently been associated with industrial point sources. Generally, however, the multiple sources of contaminants that arise from manufacturing operations within diffuse areas of industrial land use can be considered as a nonpoint source. These multiple sources include industrial pits, ponds, and lagoons that receive wastes or wastewater discharges; leaks, spills, and improper discharges of hydrocarbons and other chemicals; infiltration of runoff from parking lots, storage areas, and roadways; effluent from on-site septic systems or exfiltration from leaky sanitary sewers; and atmospheric deposition of industrial air pollutants. The most common contaminants associated with industrial sources include trace metals, volatile organic compounds from solvents and cleaners, and other synthetic organic chemicals.

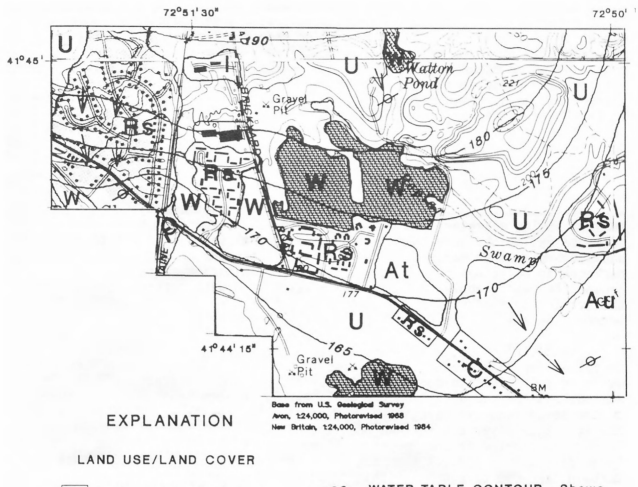
COLLECTION AND ANALYSIS OF WATER-QUALITY DATA

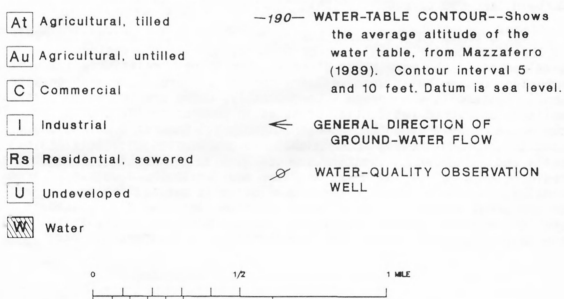
In addition to knowledge of the hydrogeologic and land-use conditions within the study areas, accurate water-quality data are needed to ascertain relations between land use and ground-water quality. Grady and Weaver (1988, p. 18-20) have shown how the historical ground-water-quality information available for the selected aquifers was too limited, largely outdated, and inappropriate for this purpose. Consequently, an observation-well network, sampling protocol, and statistical analysis program were designed to provide sufficient numbers of representative ground-water samples for comprehensive chemical analyses of nonpoint-source contaminants and valid statistical inference.

Observation-Well Network

The principal consideration in designing the observation-well network was to enable collection of ground-water samples that would provide an accurate representation of water-quality conditions within an aquifer that reflect a particular land use. To accomplish this, it was necessary to (1) locate wells within or immediately downgradient from well-defined and areally extensive land-use areas, (2) construct shallow wells with short screened intervals open to the uppermost part of the ground-water-flow system, (3) install the wells with the minimum amount of disturbance of the natural aquifer materials and without introducing any foreign materials, and (4) avoid the use of casing and screen materials that could alter the chemistry of the water samples.

Land-use maps and water-table-contour maps were overlain to locate potential observation wells at sites within or along the downgradient side of target land-use polygons. Because the extent of all land-use polygons was 5 or more acres, and generally was a few tens of acres, a single land use commonly would overlie all or most of the length of a typical ground-water-flow path. Consequently, it was possible to locate sites for observation wells within or immediately downgradient from polygons of target land-use areas such that the quality of ground water could be ascribed to a particular land use. An example of the positioning of the observation wells that demonstrates the compatibility of land use and hydrogeology is illustrated in figure 5. Appendix A (at the back of this report) identifies the land use associated with each of the 116 observation wells.





1 KILOMETER

Figure 5.--Location of selected observation wells within or immediately downgradient from polygons of target land-use areas.

CONTOUR INTERVAL 10 FEET ATIONAL GEODETIC VERTICAL DATUM OF 1929 When appropriate downgradient locations for observation wells were selected, other hydrogeologic siting criteria and requirements also needed to be satisfied. Wells were not installed where the depth to the water table exceeded 50 ft, because a thick unsaturated zone could impede downward migration of contaminants introduced at or near the land surface. If predominantly fine-grained sediments (clay and silt) that could adsorb contaminants or otherwise retard their movement were encountered in the aquifer zone to be sampled, a well was not installed at the site. Also, there could be no known major point sources of contaminants in the vicinity of the observation well so as not to complicate determination of the contaminant source. Finally, it was necessary to have access to the site for the drill rig and the permission of the landowner to install the well.

Well installation was accomplished with a truck-mounted hollow-stem auger that was used to drill a hole to depths of generally 5 to 10 ft below the water table. Type 304 stainless-steel, flush-joint casing (2-in. nominal inside diameter) and 2-ft long, wire-wound, continuous-slot, screened wellpoints were then driven an additional several feet into undisturbed aquifer materials. All casing and screens were precleaned and packaged in polyethylene tubing to prevent contamination. The annulus around the casing above the water table was filled with cuttings brought to the surface by the augers. No cements or grouts were used to avoid introducing any materials that could affect the water quality.

The distribution, by land-use category and aquifer, of the 116 wells installed for this study is identified in table 1. Although the number of wells is sufficient to compare water quality by land use statistically, it was not possible to apportion the wells evenly in each aquifer. Only three of the seven land-use categories were sampled by the six wells installed in the Pootatuck River aquifer and one land-use category, sewered residential, is not represented in the Pomperaug River aquifer (table 1). These inequities mainly reflect the fact that there is little or no occurrence of certain land-use categories within these aquifers.

Differences in the nature and intensity of land use are also reflected in the distribution of wells. For example, much of the residential development in the Farmington River aquifer is new, relatively low-density, suburban housing built upon well-drained soils where the use of on-site septic systems is widespread. Consequently, there are three times as many wells in unsewered residential areas as in sewered residential areas. In the Hockanum River aquifer, however, nearly all the residential land use consists of high-density housing that was constructed on poorly drained soils and is served by sanitary sewers. Only 1 of the 15 wells installed in residential areas of the Hockanum River aquifer is in an unsewered area. Finally, the differences in the number of wells per aquifer are in part due to the areal extent of the aquifers, with the least wells emplaced in the smallest aquifer, the Pootatuck, whereas the most wells were installed in the most extensive aquifer, the Farmington River aquifer.

Table 1.--Distribution of wells by land-use category and aquifer

	Aguifer								
Land-use category	Pootatuck River	Pomperaug River	Farmington River	Hockanum River	Total number of wells				
Undeveloped	0	3	7	5	15				
Agricultural, untilled	1	1	5	4	11				
Agricultural, tilled	0	7	6	2	15				
Residential, unsewered	2	7	21	1	31				
Residential, sewered	0	0	7	14	21				
Commercial	0	4	3	5	12				
Industrial	3	1	4	3	11				
Total number									
of wells	6	23	53	34	116				

Well depth, measured from the land surface to the bottom of the well screen, of the 116 observation wells ranged from 6.4 to 55.6 ft and averaged 20.9 ft. Depth to the water table, measured at each of the 116 wells on several occasions, ranged from 2.3 to 44.0 ft and averaged 11.2 ft. The depths of wells and the depths to the water table are listed in Appendix A and are compared in the seven land-use categories in figure 6. Wells in the agricultural areas are slightly shallower than wells in the other land-use areas. This reflects the fact that many agricultural fields occur within the valley floodplains where the depth to the water table is typically shallow. Other land uses encompass a larger variety of topographic settings where the depth to the water table commonly ranges from 10 to greater than 40 ft. However, no statistically significant differences in the depth of wells or the depth to the water table were observed among the land-use categories using analysis of variance (ANOVA) on the ranks of the well-depth and water-level data (Sokal and Rohlf, 1969; Conover and Iman, 1981; Helsel. 1983).

Sampling Protocol

Ground-water samples were collected using an all Teflon $^{\perp}$ / and stainless-steel, positive-displacement, gas-bladder pump pressurized with an oilless air compressor. Pumps of this design and construction have been found superior to other mechanisms for retrieving ground-water samples because they minimize (1) loss of volatile organic chemicals by degassing, (2) sorptive interaction of dissolved chemical species with solid surfaces, and (3) contamination of samples by leaching of pump materials (Barcelona and others, 1983; 1984; Devlin, 1986).

 $[\]frac{1}{\text{Use}}$ of trade names in this report is for identification purposes only and does not constitute endorsement by the U.S. Geological Survey.

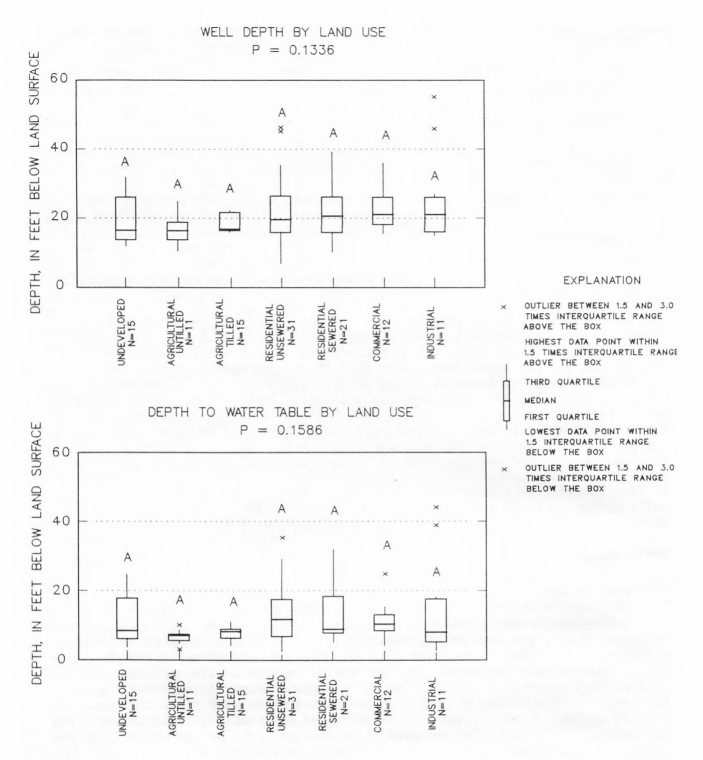


Figure 6.--Statistical comparisons by land use of depth of observation wells and depth to the water table in four stratified-drift aquifers in Connecticut. (P-value is the attained significance level for that statistical comparison; N is the number of wells sampled per land-use category; sample populations are designated by letter symbols (A, B, C, and D); sample populations that share the same letter are not significantly different at the 0.05 level of significance).

Prior to sample collection, the pump was cleaned with a detergent wash, deionized-water rinse, and then a 1:10 acetone (or methanol)-to-deionized-water solution was flushed through the pump followed by several additional volumes of deionized water. Three or more well volumes were removed from each observation well and specific conductance, pH, and water temperature were monitored until they stabilized before samples were collected. Sample collection and preservation were performed according to methods prescribed by Wood (1976), Classen (1982), Wershaw and others (1987), and Fishman and Friedman (1989).

Most ground-water samples were collected during four periods: April 1985, July through September 1986, June through September 1987, and June through July 1988. A few samples were collected during October-December 1988 to complete analytical coverage for all wells. Seven wells, however, were sampled monthly from June 1987 through June 1988 to determine the seasonal variability of selected water-quality constituents. Appendix A includes the date and time of all samples collected for this study.

Analytical determinations of 3 physical properties, 8 major inorganic chemical species, 3 nutrient species, 19 trace inorganic chemicals, 36 volatile organic compounds, 24 pesticides, and 4 other organic compounds were performed on water samples from at least 110 of the 116 observation wells (see table 2). Twenty-seven additional pesticide compounds were analyzed in a more limited set of samples from 82 to 97 wells, and analyses for 4 other pesticide compounds were performed on samples from 12 or fewer wells. Finally, samples from all 116 wells were analyzed by gas chromatography using a flame-ionization detector (GC/FID) as a tool for screening the samples for the presence of a wide variety of organic compounds. Where the GC/FID results indicated a substantial presence of one or more unknown organic compounds, gas chromatography/mass spectrometric (GC/MS) analysis was performed to identify and quantify individual baseneutral and acid-extractable (BNA) organic compounds. Samples from 24 wells were analyzed by GC/MS. All laboratory analytical determination were performed by the U.S. Geological Survey. All quantitative water-quality data are published in the annual hydrologic data report for Connecticut (Cervione and others, 1987; 1988; 1989a; 1989b; 1990). GC/FID results and other semiguantitative chromatographic data are on file at the Connecticut Office, U.S. Geological Survey, Hartford, Conn.

Statistical Analysis

Nonparametric statistical procedures have been used to test for relations between water-quality variables and land-use categories because these data are typically not normally distributed and sample size is small. The water-quality data are bounded at zero or some detection limit, commonly are highly skewed, and include infrequent extreme values or "outliers." When traditional methods of statistical inference that assume a normal distribution are applied to such data sets, the results are generally invalid (Helsel, 1987). Nonparametric statistical procedures are robust—that is, the power of the test to detect differences between groups of data is high regardless of the underlying population distribution. Rank transformation of the data, replacing each of the data points by its respective rank when all of the observations are ordered from smallest to largest, is a commonly used nonparametric procedure (Conover, 1980, p. 337; Helsel, 1987, p. 185).

Table 2.--Water-quality properties or constituents analyzed and detected in ground-water samples

[Detection limit is the lowest detectable constituent concentration or water-quality property unit of measurement that could routinely be achieved by the field or laboratory analytical methods. When analytical conditions resulted in multiple detection limits for any property or constituent, the detection limit reported below is the lowest detection limit common to the analytical methods used. Units: °C, degrees Celsius; μ S/cm, microsiemens per centimeter at 25 degrees Celsius; mJ/c, milligrams per liter; μ J/c, micrograms per liter. Number of detections is the number of wells in which the property or constituent was measured at or above the reported detection limit (when more than one analysis was available for a well the median value was used.) Values in parenthesis indicate the number of wells in which the constituent was reported at concentrations below the lowest common analytical detection limit.]

Water-quality property or constituent	Detection limit	Number of wells	Number of detec- tions	Water-quality property or constituent	Detection limit	Number of wells	Number of detec- tions
PHYSICAL PROPERTIES				VOLATILE ORGANIC COMPOUN	ns		
pH	0.1 units	115	115	Benzene	0.2 µg/L	113	3
Specific conductance	1.0 µS/cm		116	Bromoform	.2 µg/L	113	0
Temperature	.5 °C	115	115	Carbon tetrachloride	.2 µg/L	113	0
remperature		110	113	Chloromethane	.2 µg/L	113	0
				Chlorobenzene	.2 µg/L	113	0
MAJOR INORGANIC COMPOUNDS				Chlorodibromomethane	.2 µg/L	113	0
Calcium	.1 mg/L	115	115	Chloroethane	.2 µg/L	113	1
Magnesium	.1 mg/L	115	115	Chloroform	.2 µg/L	113	27
Hardness, as CaCO3	.1 mg/L	115	115	2-Chloroethylvinylether	.2 µg/L	113	0
Sodium	.1 mg/L	115	115	1,2-Dibromoethane	.2 µg/L	113	0
Chloride	.1 mg/L	116	116	1,2-Dichlorobenzene	.2 µg/L	114	3
Sulfate	.1 mg/L	116	116	1,3-Dichlorobenzene	.2 µg/L	113	0
Silica, as SiO2	.1 mg/L	115	115	1,4-Dichlorobenzene	.2 µg/L	113	0
Dissolved solids	.1 mg/L	116	116	Dichlorobromomethane	.2 µg/L	113	0
				Dichlorodifluoromethane	.2 µg/L	113	5
NUTRIENTS				1,1-Dichloroethane	.2 µg/L	113	. 7
Nitrate plus nitrite, as	N $.1 \text{ mg/L}$	116	100	1,2-Dichloroethane	.2 µg/L	113	1
Ammonia, as NH4	.01 mg/L	116	67	1,1-Dichloroethylene	.2 µg/L	113	5
Orthophosphorus, as PO4	.01 mg/L	116	47	1,2-transDichloro-			
				ethylene	.2 µg/L	113	18
TRACE ELEMENTS				1,2-Dichloropropane	.2 µg/L	113	6
Arsenic	1.0 µg/L	116	25	1,3-Dichloropropene	.2 µg/L	113	0
Barium	1.0 µg/L	115	115	1,3-cisDichloropropene	.2 µg/L	113	0
Beryllium	0.5 µg/L	115	17	1,3-transDichloro-			
Boron	20.0 µg/L	116	80	propene	.2 µg/L	113	0
Cadmium	1.0 µg/L	115	11	Ethylbenzene	.2 µg/L	113	1
Chromium	1.0 µg/L	116	26	Methylbromide	.2 µg/L	113	0
Cobalt	3.0 µg/L	115	9	Methylene chloride	3.0 µg/L	115	2
Copper	10.0 µg/L	115	6	Styrene	.2 µg/L	113	0
Iron	3.0 µg/L	115	110	1,1,2,2-Tetrachloro-			
Lead	10.0 µg/L	115	15	ethane	.2 µg/L	113	1

Table 2.--Water-quality properties or constituents analyzed and detected in ground-water samples--Continued

11-1		Maria	Number	Haban analik		No 1	Number
Water-quality		Number		Water-quality		Number	of
property or	Detection	of	detec-	property or	Detection	of	detec-
constituent	limit	wells	tions	constituent	limit	wells	tions
TRACE ELEMENTScontinued				VOLATILE ORGANIC COMPOUN	DScontinu	ied	
Lithium	4.0 µg/L	115	50	Tetrachloroethylene	0.2 µg/L	113	18
Manganese	1.0 µg/L	115	112	Toluene	.2 µg/L	113	15
Mercury	.1 µg/L	115	19	1,1,1-Trichloroethane	.2 µg/L	113	22
Molybdenum	10.0 µg/L	115	4	1,1,2-Trichloroethane	.2 µg/L	113	1
Nickel	1.0 µg/L	116	99	Trichloroethylene	.2 µg/L	113	11
Silver	1.0 µg/L	116	12	Trichlorofluoromethane	.2 µg/L	113	9
Strontium	1.0 µg/L	115	115	Vinyl chloride	.2 µg/L	113	2
Vanadium	6.0 µg/1	115	2	Xylene	.2 µg/L	112	3
Zinc	3.0 µg/L	115	108	Ny tene	731-		200
				DAGE WEUTDAL AND ACTO EV	TDACTABLE	SOMBOUND	
PESTICIDES	1	0.5	0	BASE-NEUTRAL AND ACID EX			
Alachlor	.1 µg/L	85	0	Acenaphthene	5.0 µg/L	24	0(1)
Aldicarb	.5 μg/L	82	0	Acenaphthylene	5.0 µg/L	24	0
Aldicarb sulfone	.5 μg/L	82	0	Anthracene	5.0 µg/L	24	0
Aldicarb sulfoxide	.5 µg/L	82	0	Benzo(b)fluoranthene	10.0 µg/L	24	0
Aldrin	$.01 \mu g/L$		0	Benzo(k)fluoranthene	10.0 µg/L	24	0
Ametryne	.1 µg/L	85	0	Benzo(a)pyrene	$10.0 \mu g/L$	24	0
Atrazine	.1 µg/L	85	7	Benzo(a)anthracene	$10.0 \mu g/L$		0
Carbaryl	$.5 \mu g/L$	82	0	Benzo(g,h,i)perylene	$10.0 \mu g/L$	24	0
Carbofuran	.5 µg/L	82	0	Bis(2-chloroethyl)-			
Chlordane	$.1 \mu g/L$	113	3	ether	5.0 µg/L	24	0
Chlorpyrifos	.01 µg/L	. 88	0	Bis(2-chloroethoxy)-			
Cyanazine	.1 µg/L	85	1	methane	$5.0 \mu g/L$	24	0
DDD	.01 µg/L	113	1	Bis(2-chloroisopropyl)	-		
DDE	.01 µg/L	. 113	0(2)	ether	5.0 µg/L	24	0
DDT	.01 µg/L	113	0	Bis(2-ethyhexyl)-			
2,4-D	.01 µg/L	. 97	5	phthalate	5.0 µg/L	24	7(2
2,4-DP	.01 µg/L	. 97	1	Butyl benzyl phthalate	5.0 µg/L	24	0
2,4,5-T	.01 µg/L		3	Chrysene	10.0 µg/L	24	0
Dicamba	.01 µg/L		0	4-Bromophenyl phenyl-			
Diazinon	.01 µg/L		2	ether	5.0 µg/L	24	0
Dieldrin	.01 µg/L		5(1)	2-Chloronaphthalene	5.0 µg/L	24	0
Disulfoton	.01 µg/L		0	2-Chlorophenol	5.0 µg/L	24	0
DEF	.01 µg/L		0	4-Chloro-3-methyl-			
Endosulfan	.01 µg/L		0	phenol	30.0 µg/L	24	0
Endrin	.01 μg/L		0	4-Chlorophenyl phenyl-			
Ethion	.01 µg/L		0	ether	5.0 µg/L	24	0
Fonofos	.01 µg/L		0	2,4-Dichlorophenol	5.0 µg/L		0
Heptachlor	.01 µg/l		0	Diethyl phthalate	5.0 µg/L		1
Heptachlor epoxide	.01 µg/L		0	Dimethyl phthalate	5.0 µg/L		0
3-Hydroxycarbofuran	.5 μg/L	82	0	Dibenz(a,h)anthracene	10.0 µg/L		0
Lindane	.01 μg/L		0	2,4-Dimethylphenol	5.0 μg/L		0
Malathion	.01 μg/l		0	2,4-Dinitrophenol	20.0 µg/L		0
		82	0	4,6-Dinitro-2-methyl-	20.0 pg/	2 1	J
Methomy l	.5 µg/L		0	phenol	30.0 µg/L	. 24	0
Methoxychlor	.01 µg/l			2,4-Dinitrotoluene	5.0 µg/l		0
Metolachlor	.1 µg/L	85	0	2,4-01111110101delle	J.0 µg/1	. 24	U

Table 2.--Water-quality properties or constituents analyzed and detected in ground-water samples--Continued

			Number				Number
Water-quality		Number	of	Water-quality		Number	of
property or	Detection	of	detec-	property or	Detection	of	detec-
constituent	limit	wells	tions	constituent	limit	wells	tions
PESTICIDEScontinued				BASE-NEUTRAL AND ACID EXT	RACTABLE CO	MPOUNDS-	-continue
Metribuzin	$0.1 \mu g/L$	85	0	2,6-Dinitrotoluene	5.0 µg/L	24	0
Methyl parathion	$.01 \mu g/L$	113	0	Di-n-octyl phthalate	10.0 µg/L	24	0(1)
Methyl trithion	$.01 \mu g/L$	113	0	Di-n-butyl phthalate	5.0 µg/L	24	0(2)
Mirex	.01 µg/L	112	0	Fluoranthene	5.0 µg/L	24	0(1)
1-Naphthal	.5 µg/L	82	0	Fluorene	5.0 µg/L	24	0(1)
0xamy1	.5 µg/L	82	0	Hexachlorobenzene	5.0 µg/L	24	0
Parathion	.01 µg/L	113	0	Hexachlorobutadiene	5.0 µg/L	24	0
Perthane	$.1 \mu g/L$	113	0	Hexachlorocyclo-			
Phorate	.01 µg/L	88	1	pentadiene	5.0 µg/L	24	0
Picloram	.01 µg/L	10	0	Hexachloroethane	5.0 µg/L	24	0
Prometone	.1 µg/L	85	2	Ideno(1,2,3-cd)pyrene	10.0 μ g/L	24	0
Prometryn	.1 µg/L	85	0	Isophorone	$5.0 \mu g/L$	24	0
Propazine	.1 µg/L	85	1	Napthalene	5.0 µg/L	24	0(1)
Propham	.5 µg/L	82	0	Nitrobenzene	$5.0 \mu g/L$	24	0
Silvex	.01 µg/L	110	2	2-Nitrophenol	5.0 µg/L	24	0
Simazine	.1 µg/L	85	2	4-Nitrophenol	30.0 µg/L	24	0
Simetryn	.1 µg/L	85	0	n-Nitrosodimethylamine	5.0 µg/L	24	0
Toxaphene	$1.0 \mu g/L$	113	0	n-Nitrosodiphenylamine	5.0 µg/L	24	0
Trifluralin	.1 µg/L	85	0	n-Nitrosodipropylamine	5.0 µg/L	24	0
Trithion	.01 µg/L	113	0	Phenanthrene	5.0 µg/L	24	0
				Phenol	5.0 µg/L	24	0(2)
OTHER ORGANIC CONSTITUENT	S			Pentachlorophenol	30.0 µg/L	24	0
Total organic carbon	.1 mg/L	116	112	Pyrene	5.0 µg/L	24	0
Methylene-blue-active	3.			1,2,4-Trichlorobenzene	5.0 µg/L	24	0
substances	.01 mg/L	116	115	2,4,6-Trichlorophenol	20.0 µg/L	24	0
Gross polychlorinated	,						
biphenyls	.1 µg/L	113	0				
Gross polychlorinated							
naphthalenes	.1 µg/L	113	0				

Four statistical procedures have been used to describe the data and to test for differences between water-quality variables that were segregated by land use. First, descriptive statistics have been used to describe the central tendency (median) and variability of the data (interquartile range) within each group. The median or 50th percentile of the data's distribution is the preferred measure of central tendency for nonnormally distributed data because, unlike the mean, it is not influenced by the magnitude of extreme values. Similarly, the interquartile range—the difference between the 75th percentile (or 3rd quartile) and the 25th percentile (1st quartile)—provides a measure of sample variability that is less influenced by outliers than is the standard deviation for nonnormal data. For censored data distributions—that is, if some of the data are reported as less than a specified detection level—descriptive statistics were sometimes estimated by the method of Helsel and Gilliom (1986) and Gilliom and Helsel (1986).

If more than 25 percent of the data for a variable are below the analytical detection limit, the interquartile range was estimated; if more than 50 percent of the data are below the detection limit, the median was also estimated, if possible. For some variables there were too few detections, or all detections were at the same concentration, and estimates of median and interquartile range could not be made. Boxplots are used to display the population distribution of water-quality variables that have 75 percent or more of the data at or above the analytical detection limit.

Analysis of variance of rank-transformed data for 20 uncensored or slightly censored water-quality variables was used to test for significant differences in water quality among land uses (one-way ANOVA) and among land uses and underlying aquifers (two-way ANOVA) (Sokal and Rohlf, 1969: Conover and Iman, 1981; Helsel, 1983). The uncensored or slightly censored variables -- those with 75 percent or more of the values equal to or greater than the analytical detection level--generally include the physical properties, major inorganics, and some of the trace elements. An ANOVA of the ranked boron concentrations was also performed, although only 69 percent of the boron observations were at or above the detection limit. Nearly half of the censored boron observations were for samples from two land-use categories. The null hypothesis tested by ANOVA states that there is no significant difference in the mean rank--a measure of the central tendency of the sample population of each water-quality variable -- for any land use. Rejection of the null hypothesis at the 0.05 level of significance is considered evidence of a relation between land use and ground-water quality, aquifer and ground-water quality, or some interaction of land use and aquifer with ground-water quality. An unbalanced ANOVA design is required because the number of wells in each of the land-use categories is not the same. The SAS Systems General Linear Models (GLM) procedure was used (SAS Institute Inc., 1985).

Where the GLM procedure indicated that land use had a significant effect on a water-quality variable, a multiple-comparison procedure, Tukey's honest significant-difference test (Sokal and Rohlf, 1969; Stoline, 1981) can be used to discriminate the land uses that differ. The Tukey test uses the within-group variance to calculate, for all possible pairwise comparisons, the minimum difference in mean rank between groups of data that is required to consider the groups significantly different (Helsel and Ragone, 1984). As with the ANOVA, a 0.05 level of significance (alpha) or the probability of falsely detecting differences that are the result of chance rather than the independent variable (land use), was chosen for the Tukey tests. Tukey's honest significant-difference test was used because it controls the overall error rate at alpha = 0.05, and it is applicable to the uneven sample sizes.

Many of the water-quality variables, especially trace elements and organic compounds, are moderately censored; fewer than 75 percent of the samples have detectable concentrations. For many of these variables, the level of censoring is severe; fewer than 25 percent of the samples have concentrations above the detection level. For these variables, the only statistical test that was applicable was contingency-table analysis (Conover, 1980; Helsel and Ragone, 1984). Contingency-table analysis compares the frequency of detection, or the frequency of exceeding some designated concentration, for the variable by land-use category. The procedure is to tabulate the observations by row, according to land-use

category, and by column according to some measure of the water quality, such as whether the concentration of a variable is above or below the detection limit or some other concentration threshold, or whether the concentration falls within a specified range of values. The null hypothesis tested by the contingency table is that the probability of an observation being classified into one of the column categories is independent of the classification (row) it belongs to. Rejection of the null hypothesis at the 0.05 level of significance is evidence that the frequency of detection of a water-quality variable is associated with land use.

Multiway (7x2) contingency tables were first computed using an exact probability determination on the basis of a multinomial distribution (Sokal and Rohlf, 1969, p. 590; Conover, 1980, p. 160). The exact probability computed by the contingency-table analysis is the probability of obtaining the observed cell counts or more extreme cell counts (greater deviations from expected) given the marginal totals; this probability is used in preference to the chi-square approximation because of the low to zero cell frequencies for some water-quality variables. If a significant difference in the frequency of detection was noted by this procedure, separate 2x2 contingency tables of all possible two-row combinations of land uses were computed using Fisher's exact test (Sokal and Rohlf, 1969, p. 595) to determine which pairs of populations differ (Iman and Conover, 1983, p. 309).

EFFECTS OF LAND USE ON GROUND-WATER QUALITY

The chemistry of ground water is affected by natural factors and human activities. As water moves through the hydrologic cycle, its composition is modified by a number of chemical, physical, and biological processes. Particles of dust and atmospheric gases dissolve to produce low concentrations of most major inorganic constituents in the precipitation that falls on the land surface. Near saltwater sources, precipitation commonly contains increased concentrations of sodium and chloride. Industrial and automobile emissions produce elevated levels of nitrate and sulfate and the low pH characteristic of acid rain.

The composition of water moving through the soil and the unsaturated zone is further modified by evapotranspiration and reactions with the porous media, interstitial water and gases, and organic matter. The duration of contact between water molecules and earth materials, as well as the solubility, sorptive, and ion-exchange properties of those materials, controls the chemistry of natural waters. Although the texture of stratified-drift sediments can be highly variable, the mineral composition is predominantly siliceous and only minor calcite, sodic plagioclase and potassium feldspars, and accessory minerals are reactants (Rogers, 1987, p. 1531). The short flow paths and brief residence time of ground water in stratified-drift aquifers also influence the extent of natural geochemical evolution such that the dissolved solids concentrations rarely exceed 300 mg/L (milligrams per liter) (Rogers, 1987; p. 1531; Melvin and others, 1988, p. 192).

The quality of ground water can be impaired by human activities, such as those commonly associated with agriculture, industrialization, and urbanization. The common causes of ground-water contamination in Connecticut are applied pesticides, improperly handled and disposed of solvents, leachate from solid-waste disposal sites, leakage from petroleum storage tanks, and improperly stored road salt (Melvin and others, 1988).

During 1979-86, these causes accounted for 882 of the 928 incidents of public- and private-well contamination reported by the Connecticut Department of Environmental Protection (1986, p. 35). Generally, contamination is restricted to an area within a few hundred to a few thousand feet of a readily identifiable point source, such as a landfill, surface impoundment, underground storage tank, known chemical spill, or injection well. However, degradation of water quality from nonpoint sources that introduce contaminants over an area of a few acres to tens or even hundreds of acres may be a more pervasive problem and is the focus of this report. Nonpoint sources of contaminants include application of agricultural pesticides and fertilizers, discharge of wastewater to septic tanks and cesspools, leakage from sewage pipelines, road salt applications, and infiltration of runoff from paved areas, as well as the aggregate effect of multiple, small point sources (for example, leaks, spills, and discharges of hydrocarbons or other chemicals within an industrial park).

Land use is a convenient framework for describing the prevailing activities within a relatively uniform demographic area, although it may have little relation to the severity and magnitude of ground-water degradation within or generated from that area. A host of other causative factors, such as population density; septic-system density; percentage of impervious area; traffic patterns; street-cleaning practices; nature and density of storm sewers; age and density of underground storage tanks; deicing-chemical application rates; design, construction characteristics and condition of sanitary (or combined sanitary and storm) sewers; vegetation cover; tillage practices; agricultural-chemical application rates; and irrigation practices affect the amount of contaminants generated from areal sources. Because many of these factors cannot be quantified and are highly variable in the study areas, land use is used as a collective surrogate for nonpoint-contaminant sources of differing type and intensity.

Ground-Water Quality Beneath Land-Use Areas

Table 2 shows that 80 of the 179 water-quality variables analyzed for in ground water from the four stratified-drift aquifers have been detected or measured in at least one sample. These include all 33 of the inorganic constituents and physical properties but only 47 of the 146 organic constituents tested (including 8 constituents reported at concentrations below the lowest common analytical detection limit). The statistical procedures previously described were performed for all water-quality variables included in table 2 that were detected in ground-water samples, except the BNA organic compounds. Only 24 samples were analyzed by the GC/MS method for the target BNA organic compounds listed in table 2. Other nontarget BNA compounds were identified and quantified if mass spectra could be matched to National Bureau of Standards library reference spectra—/. Those analyses were performed to identify and quantify both target and nontarget BNA compounds subsequent to detection of unknown compounds by GC/FID scans.

 $[\]frac{2}{D}$ Data for tentatively identified organic compounds (TIOCs) in this report are based on comparison of sample spectra with library spectra followed by visual examination by GC/MS analysts. TIOC data have not been confirmed by direct comparison with reference standards. Therefore, TIOC identification is tentative, and reported concentrations are semiquantitative.

Attained significance levels (p-values), or the probability that observed differences are due to chance rather than the tested effect, are shown in table 3 for the 21 water-quality variables compared by ANOVA, and are listed in table 4 for the 49 variables compared by contingency-table analysis. Only 23 of the 70 tested water-quality variables have significantly different populations for one or more types of land use. While inorganic chemical constituents and a few organic constituents have both natural and manmade sources, statistically significant differences in sample populations for data segregated by land use are due to human sources rather than natural factors. Table 5 (at the end of the report) is a statistical summary by land-use category of all detected water-quality variables, except the BNA-extractable organic compounds, showing the sample size, number of detections, median and interquartile range, and letter symbols that identify similarities and differences in sample populations.

Table 3.--Attained significance levels (p-values) for two-way analysis of variance on the ranks of water-quality variables compared by land use and aguifer

[The p-value is the probability that observed differences are due to chance rather than the tested effect. Asterisk indicates effect is significant at the 0.05 level]

Water-quality variable	e p-value for the tested effect				
	Land use	Aquifer	Interaction		
Temperature	0.0864	0.7078	0.8194		
Specific conductance	.0001*	.0035*	.0504		
pH	.0489*	.2542	.2914		
Calcium	.0002*	.0429*	.3170		
Magnesium	.0001*	.0001*	.0386*		
Hardness, as CaCO3	.0001*	.0167*	.1704		
Sodium	.0001*	.0001*	.0857		
Chloride	.0003*	.0264*	.7146		
Sulfate	.0050*	.0007*	.2064		
Silica, as SiO2	.0166*	.4045	.0424*		
Dissolved solids	.0001*	.0061*	.0664		
Nitrate plus nitrite, as N	.0001*	.0924	.1704		
Barium	.0180*	.0001*	.4859		
Boron	.0001*	.0632	.5346		
Iron	.0327*	.9311	.2199		
Manganese	.0619	.7568	.1124		
Nickel	.0477*	.6557	.0394*		
Strontium	.0003*	.4255	.1874		
Zinc	.8599	.8040	.2295		
Total organic carbon	.4023	.1862	.5694		
Methylene-blue-active substances	.0006*	.1578	.0014*		

Table 4.--Attained significance levels (p-values) for contingency-table tests of water-quality variables compared by land use

[The p-value is the probability that observed differences are due to chance rather than to the tested effect. Asterisk indicates effect is significant at 0.05 level]

Water-quality variable	p-value	Water-quality variable	p-value
Ammonia, as NH4	0.0460*	Ethylbenzene	0.9999
Orthophosphorus, as PO4	.2807	Methylene chloride	.7666
Arsenic	.6513	Tetrachloroethylene	.0013*
Beryllium	.0700	1,1,2,2-Tetrachloroethane	.9999
Cadmium	.4997	1,1,1-Trichloroethane	.0570
Chromium	.4821	1,1,2-Trichloroethane	.1947
Cobalt	.2563	Trichloroethylene	.0073*
Copper	.2940	Trichlorofluoromethane	.3239
Lead	.6378	Toluene	. 4495
Lithium	.0614	Vinyl chloride	.7067
Mercury	.5891	Xylene	.0510
Molybdenum	.3471	Atrazine	.0131*
Silver	.1296	Cyanazine	.5176
Vanadium	1.0000	Prometone	.0616
		Propazine	.6588
Benzene	.5926	Simazine	.8908
Chloroethane	.1875	Chlordane	1.0000
Chloroform	.0018*	Diazinon	.8972
Dichlorodifluoromethane	.6543	Dieldrin	.9739
1,2-Dichlorobenzene	.7970	DDD	.9999
1,1-Dichloroethane	.1494	2,4-D	.4640
1,2-Dichloroethane	.9999	2,4-DP	.7423
1,1-Dichloroethylene	.1450	2,4,5-T	.8826
1,2-Dichloropropane	.2040	Silvex	.9999
1,2-transDichloroethylene	.0382*	Phorate	1.0000

Undeveloped Land

The quality of ground water in undeveloped areas is largely unaffected by human activities and is considered to represent background conditions. The effects of human activities on ground-water quality in other land-use areas are generally quantified by comparison to values measured in the sample population from the undeveloped areas. The only probable nonpoint sources of contaminants in undeveloped areas are minor tracts of discordant land uses situated within the more extensive undeveloped areas, and atmospheric deposition of contaminants.

Median concentrations and (or) frequencies of detection of 19 inorganic constituents and physical properties are lowest (or equally as low as for another type of land use) for ground-water samples from the undeveloped land-use area (table 5). The mean rank of the sample population or the frequency of detection for 13 of these variables is significantly lower for

the undeveloped area than 1 or more of the other 6 land-use areas. Also, within-sample-population variability, as measured by the interquartile range, is smaller (or equally as small as for another land use) for samples from the undeveloped area for 12 of the 21 inorganic constituents and physical properties for which this statistic can be compared. The lowest pH (5.2) and temperature $(6.1\ ^\circ\text{C}\ (\text{degrees Celsius}))$ were recorded for undeveloped-area samples, as were the lowest concentrations of barium $(6\ \mu\text{g/L}\ (\text{micrograms per liter}))$, chloride $(1.4\ \text{mg/L})$, and dissolved solids $(30\ \text{mg/L})$. The frequency of detecting arsenic, lithium, molybdenum, and silver was lowest (or equally as low as for another land-use area) for ground-water samples from the undeveloped areas. These findings are indicative of the dilute, slightly acidic, soft water that is characteristic of a short residence time within the dominantly silicate-mineral matrix of the stratified-drift aguifers.

Median iron concentration and interquartile range, however, are higher in the undeveloped area than the other land-use areas. This may reflect the depletion of dissolved oxygen during infiltration through soils with high organic-matter content, which are more prevalent in undeveloped areas. Where dissolved oxygen is depleted by the oxidation of organic material in wetland and forest soils, dissolution of iron oxides in the soil and aquifer materials is facilitated. The mean rank of iron concentrations in samples from the undeveloped land-use category, however, is not significantly different from that of any of the other land-use areas. The inability of the ANOVA to discern a significant difference between iron concentrations in the undeveloped areas and the other land-use areas is due to the large variability of iron concentrations in the undeveloped area that range from less than 3.0 to 18,000 μ g/L. Low concentrations of cobalt (3.0 to 6.0 $\mu q/L$) and lead (10 to 20 $\mu q/L$) were detected most often in the undeveloped areas, however their detection frequencies are not significantly different from other land-use areas. The infrequent detections of cobalt and lead may also be related to natural geochemical factors.

Median total organic carbon (TOC) and methylene-blue-active substances (MBAS) concentrations are also lowest (or equally as low as another land use) in the undeveloped areas. There were no detections of 26 of the 37 organic constituents listed in table 5 in ground-water samples from the undeveloped areas. Seven volatile organic compounds were detected, generally at trace-level concentrations, in some undeveloped-area wells. The fluorocarbon trichlorofluoromethane and 1,2-transdichloroethylene, a break-down product of widely used solvents, were each detected in two wells at concentrations of less than 1.0 $\mu g/L$. Toluene and benzene were present in the water from one undeveloped-area well at 2.2 and 0.7 μ g/L respectively. Neither compound was detected in a subsequent sample from this well, suggesting the possibility of a transitory source such as a small hydrocarbon spill in the vicinity of the well. GC/FID analysis of a sample from this well reported an unidentified BNA compound, but the estimated concentration was considered to be too low for positive identification and quantification using GC/MS analysis. In addition, chloroform, dichlorodifluoromethane, and 1,2-dichloropropane have each been detected at concentrations of 0.8, 0.6, and 1.8 μ q/L respectively in the undeveloped area.

The herbicides simazine (0.2 μ g/L) and 2,4-D (0.06 μ g/L) were detected in water samples from two wells in undeveloped areas. Their source may have been applications to control weeds along powerline corridors that transect the areas. An alternative source for all or some of the trace-level organic compounds found in the undeveloped samples may be atmospheric deposition. Eisenreich and others (1981) and Shulters and others (1987) report that volatile, BNA, and pesticide compounds were found at nanogram to microgram per liter concentrations in precipitation samples collected downwind of both urban and agricultural areas.

Agricultural Land

Agricultural land use has been subdivided into untilled and tilled categories as previously described in the section on land use. principal difference between the two land-use categories with respect to nonpoint-source contaminants is related to the intensity of agricultural practices. Wall and Magner (1988) found lower nitrate and chloride concentrations in ground water beneath areas of limited agricultural intensity with little row-crop production as compared to areas of moderate to intense agricultural activity with predominant row-crop cultivation. Hallberg (1986, p. 5) cites that a 3- to 60-fold increase in nitrate concentration between forested-pasture-grassland areas and intensely cultivated and fertilized areas has been documented by numerous investigators. Although data are sparse on the actual application rates, anecdotal information suggests that untilled areas included in this study receive lesser amounts of fertilizers and pesticides than tilled areas. Numerous macropores in the undisturbed soil profiles of untilled areas, however. may facilitate movement of contaminants to the water table (White, 1985; Dick and others, 1986; Hallberg, 1986, p. 38).

Untilled areas

Ground-water quality beneath untilled agricultural areas is minimally affected by human activities. Although median concentrations of most inorganic constituents are somewhat elevated with respect to background levels (see table 5), only magnesium and hardness are significantly higher than in the undeveloped areas. Median ammonia-nitrogen and nitrate plus nitrite-nitrogen concentrations are about 10 times greater in ground water beneath the untilled agricultural land use than the undeveloped; however, the differences are not statistically significant. The highest reported concentration of sulfate (120 mg/L) was for a sample from the untilled agricultural area. Applications of fertilizers and inorganic chemical soil treatments, including lime, are the most likely cause for elevated concentrations of these constituents.

Median iron, manganese, silica, and MBAS concentrations are lowest (or equally as low as for another land use) in ground-water samples from the untilled agricultural areas. The interquartile range of the concentrations of several inorganic constituents (chloride, silica, orthophosphorus, and strontium) is smallest for the untilled agricultural areas. Ground-water quality in the untilled agricultural areas resembles background conditions with respect to the generally low median concentrations and interquartile ranges for most inorganic ground-water constituents.

There are no significant differences in the mean rank of the concentrations or frequency of detection of any organic compound between the untilled agricultural and the undeveloped land uses (see table 5). Only six volatile organic compounds and one pesticide compound were detected in samples from the untilled areas. Trace-level concentrations (0.2 to 0.3 $\mu q/L$) of chloroform and 1,2-transdichloroethylene were each detected three times. Two detections of trichlorofluoromethane (0.3 and 0.7 μ g/L), and one detection each of chloroethane (0.3 μ g/L), 1,1,1-trichloroethane (0.2 μ g/L), and toluene (0.2 µg/L) were reported in ground-water samples from this land use. Because 9 of the 11 detections of volatile organic compounds were in samples from 3 wells, and there is no known agricultural use for these compounds, their presence may reflect unknown anthropogenic factors in the vicinity of the anomalous wells, or perhaps atmospheric deposition. Similarly, detections of unidentified BNA compounds in three GC/FID scans cannot be related to known human activities in this land-use area. GC/MS analyses performed on two of these samples were inconclusive. No BNA compounds were detected in one sample, and a trace-level concentration (0.7 uq/L) of one compound, methylbenzene (toluene), was reported in the other. Resampling the third well failed to confirm the GC/FID peaks reported in the initial sampling. The only pesticide detected in the untilled agricultural areas was cyanazine $(0.1 \, \mu \text{g/L})$, a herbicide commonly used on fallow agricultural lands.

Tilled areas

Ground-water quality beneath tilled agricultural lands is more severely affected than is water quality beneath untilled areas because of the more intense agricultural practices associated with this land use. The mean rank of the sample populations or the frequency of detection for specific conductance, calcium, magnesium, hardness, sulfate, dissolved solids, nitrate plus nitrite nitrogen, ammonia nitrogen, boron, and strontium is significantly higher for the tilled agricultural areas than the undeveloped areas (see table 5). Increased nitrate plus nitrite-nitrogen concentrations and ammonia-nitrogen detections can be attributed to widespread applications of inorganic and organic fertilizers on cultivated cropland. Applications of a variety of inorganic chemical compounds as soil treatments and micronutrients, including lime, dolomite, anhydrite, gypsum, and borax, increase the concentrations of calcium, magnesium, hardness, sulfate, boron, and strontium. Increased specific conductance and dissolved solids reflect the aggregate effects of fertilizer and inorganic chemical applications.

The median nitrate plus nitrite-nitrogen concentration of 4.7 mg/L is higher for ground-water samples from the tilled agricultural areas than for all other land-use areas (fig. 7). Twenty percent of the wells in the tilled agricultural areas yielded water that exceeded the 10 mg/L limit specified by the National primary drinking-water regulations (U.S. Environmental Protection Agency, 1986). In 11 samples collected from one well installed beneath a corn field that receives heavy applications of organic fertilizer (manure), nitrate plus nitrite-nitrogen concentrations ranged from 5.5 to 60 mg/L. Although the mean rank of the nitrate plus nitrite-nitrogen concentrations in water samples from the tilled agricultural areas is significantly higher than in those from the undeveloped areas, it is not significantly different from the untilled agricultural areas or any of the developed land-use areas. The

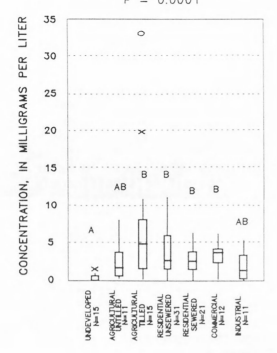
interquartile range of nitrate plus nitrite-nitrogen and sulfate concentrations is greater for samples from this land-use area than for samples from all other land-use areas. Large variability in the concentrations of these constituents may reflect differences in the quantity and composition of fertilizer and soil-treatment chemicals applied to different tilled crops by different farmers.

Although the use of phosphate fertilizers has increased the phosphorous content of agricultural drainage and contributed to algal growth problems in surface waters (Hem, 1985, p. 126-129), the low solubility of phosphorous and its use by biota as a nutrient generally result in low concentrations in ground waters. Both the estimated median concentration (0.002 mg/L) and the frequency of detection (20 percent) of orthophosphorus were lowest in water samples from the tilled agricultural areas but did not differ significantly from those of the other land-use areas. Several trace elements (arsenic, cadmium, cobalt, copper, chromium, lead, molybdenum, silver, and vanadium) were also detected least frequently (or equally as frequent as for another land use) in the tilled agricultural areas.

Although the mean rank of the MBAS concentrations for ground-water samples from the tilled agricultural areas is not statistically different from that of the other land-use areas, the median MBAS concentration, 0.05 mg/L, is elevated relative to five other land-use areas and equally as large as the median MBAS concentration for samples from the commercial areas. Samples from wells in the tilled agricultural areas had both the lowest (<0.01 mg/L) and the highest (0.38 mg/L) MBAS concentrations. Elevated MBAS concentrations may relate to agricultural pesticide use, because surfactants are used to aid dispersal of some pesticide compounds.

The herbicide atrazine, widely used for weed control in corn, was detected at low concentrations (0.1 to 0.2 μ g/L) in one-third of the wells in the tilled agricultural areas (fig. 7). The frequency of atrazine detections was significantly higher in this land use than in the undeveloped areas, where it was undetected. Although no atrazine was detected in ground water beneath the untilled agricultural, unsewered residential, and sewered residential land-use areas as well, the number of samples from these areas was too few to establish a valid statistical difference in the frequency of detection.

One additional pesticide and nine volatile organic compounds were detected in ground-water samples from the tilled agricultural areas. The insecticide dieldrin, commonly used to control corn pests prior to its prohibition in 1974, was detected at a concentration of 0.13 $\mu g/L$ in water from one well. The extremely low volatility and low solubility of dieldrin, as well as its high affinity for organic matter, may result in its residual occurrence and may account for the persistence of this compound in ground water more than a decade after all use presumably ceased. The most frequently detected volatile organic compound, 1,2-dichloropropane, is a soil fumigant and nematocide used on tobacco and strawberries in the study areas. It has been detected in water samples from three wells in the tilled areas at concentrations of 0.9 to 41 $\mu g/L$. The frequency of detection of 1,2-dichloropropane is highest in ground water from the tilled agricultural areas; however, it is not significantly higher than in the other land-use areas where it has been detected rarely or not at all (see table 5).



ATRAZINE DETECTIONS BY LAND USE (detection limit 0.1 microgram per liter) P = 0.0131

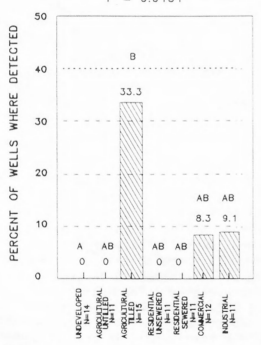


Figure 7.--Statistical comparisons by land use of nitrate plus nitritenitrogen concentrations and frequency of atrazine detections in ground-water samples from four stratified-drift aquifers in Connecticut. (P-value is attained significance level for that statistical comparison; N is number of wells sampled per land-use category; sample populations are designated by letter symbols (A, B, C, and D); sample populations that share the same letter are not significantly different at the 0.05 level of significance).

EXPLANATION

- OUTLIER GREATER THAN 3.0 TIMES INTERQUARTILE RANGE ABOVE THE BOX
- OUTLIER BETWEEN 15 AND 3.0 TIMES INTERQUARTILE RANGE ABOVE THE BOX

HIGHEST DATA POINT WITHIN
1.5 TIMES INTERQUARTILE RANGE
ABOVE THE BOX

THIRD QUARTILE

MEDIAN

FIRST QUARTILE

LOWEST DATA POINT WITHIN 1.5 TIMES INTERQUARTILE RANGE BELOW THE BOX The occurrence of three volatile organic compounds, 1,1-dichloroethane, 1,1-dichloroethylene, and 1,1,1-trichloroethane at concentrations of 0.3, 1.4, and 25 $\mu \rm g/L$ respectively, in the water from one tilled-area well is enigmatic. The lack of any known agricultural source for these compounds and the elevated concentration of the 1,1,1-trichloroethane suggests contamination from a nonagricultural source in the vicinity of the well. Trace-level concentrations (0.2 to 0.5 $\mu \rm g/L)$ of other volatile organic compounds in one or two water samples, 1,2-dichlorobenzene, dichlorodifluoromethane, 1,2-transdichloroethylene, trichlorofluoromethane, and xylene may also originate from nonagricultural sources.

GC/FID analyses of water samples from four wells completed in the tilled agricultural land use indicated that each sample contained one or more unidentified BNA organic compounds. Subsequent resampling and GC/FID analysis of water from three of the wells failed to detect any compounds in two of the samples, while the presence of organic compounds was confirmed in the third sample. GC/MS analysis was performed on the latter sample, as well as on the sample from the fourth well. One sample contained nine nontarget BNA organic compounds, including five aliphatic hydrocarbons and four unidentified alkane compounds, at concentrations of less than 1.0 to 20 $\mu \rm g/L$. The other sample contained 38 $\mu \rm g/L$ of an unidentified carboxylic acid ester and 5.8 $\mu \rm g/L$ of 1,3,5-cycloheptatriene. No agricultural source is apparent for any of the compounds identified in these samples.

Residential Land

Although several nonpoint sources may contribute to the degradation of ground-water quality in residential areas, the most important is perhaps on-site disposal of wastes through septic tanks and cesspools. Septic tanks represent the highest total volume of wastewater discharged directly to ground water and the most commonly recorded source of ground-water contamination (U.S. Environmental Protection Agency, 1977). Consequently, residential land use was differentiated into unsewered and sewered areas to evaluate if differences in ground-water quality exist between the two areas that may reflect the manner of sewage disposal.

Eckhardt and others (1989) observed statistically significant differences in the concentration or frequency of detection of several inorganic constituents and volatile organic compounds in ground water beneath sewered and unsewered areas of Long Island, New York. Ground water beneath sewered areas of Long Island was found to have higher total dissolved solids concentrations and greater frequency of volatile organic compound detections than unsewered areas. Nitrate and chloride concentrations were not significantly different between the two areas, although the implementation of sewering was anticipated to reduce levels of these constituents. Eckhardt and others (1989, p. 22-25) concluded that the higher population density of the sewered areas together with other sources of ground-water contamination complicated direct comparisons of ground-water quality between sewered and unsewered areas.

Unsewered areas

The quality of ground water beneath unsewered residential lands is affected to about the same degree with respect to inorganic constituents as is the water quality in the tilled agricultural areas. The mean rank of the

concentrations or the frequency of detection of specific conductance, calcium, magnesium, hardness, sodium, chloride, dissolved solids, nitrate plus nitrite nitrogen, boron, and strontium is significantly higher in samples from the unsewered land use than in those from the undeveloped areas (table 5). The mean rank of the sodium concentrations is also significantly higher in the unsewered residential areas than in both the untilled and tilled agricultural land uses, reflecting the absence of any substantial septic-tank or road-salt sources in those areas. Median concentrations of most inorganic constituents are intermediate; only strontium has a higher median concentration in water samples from the unsewered residential areas than in samples from all other land-use areas (fig. 8). The most likely source of strontium is from septic-tank effluents, inasmuch as the only documented domestic use of strontium known to the author is in toothpaste (Wennig and Kirsch, 1988, p. 632). Elevated levels of all 10 of the above water-quality variables could result from septic-tank effluents; however, other nonpoint sources, including applications of deicing chemicals and lawn and garden fertilizers, likely are sources as well.

Large variations in some water-quality constituents were found in unsewered residential areas, as evidenced by the fact that three inorganic constituents have their lowest and highest concentrations in water from this land use. Calcium concentrations ranged from 1.5 to 110 mg/L, strontium concentrations ranged from 9 to 940 $\mu g/L$, and hardness concentrations ranged from 5 to 270 mg/L (as calcium carbonate) in samples from unsewered residential-area wells. Lowest magnesium (0.3 mg/L) and silica (3.4 mg/L) concentrations also were detected in this area, whereas the highest temperature (20 °C) and concentrations of barium (400 $\mu g/L$), cadmium (8 $\mu g/L$), iron (20,000 $\mu g/L$), lithium (66 $\mu g/L$), manganese (7,400 $\mu g/L$), mercury (3.7 $\mu g/L$), and silver (1.8 $\mu g/L$) were reported in ground-water samples from the unsewered residential land use. This large variability may be attributed, in part, to the wide range in housing densities encountered in unsewered areas.

Evidence of widespread presence of organic contaminants from human activities in the unsewered residential land use is provided by the detection of one or more volatile organic compounds in samples from 62 percent of the wells in this category. Although 17 volatile organic compounds have been detected in ground-water samples from the unsewered residential areas, including 5 compounds (1,2-dichlorobenzene, 1,2dichloroethane, ethylbenzene, methylene chloride, and 1,1,2,2tetrachloroethane) found solely or most commonly in this area, the frequency of detection was not significantly higher for any of the compounds (table 5). The most commonly detected volatile organic compounds were toluene (0.2 to 0.3 μ g/L) and 1,1,1-trichloroethane (0.2 to 2.1 μ g/L), both of which were found in six wells. Chloroform was detected in samples from five wells at concentrations ranging from 0.2 to 1.3 μ g/L, tetrachloroethylene in samples from four wells at trace-level concentrations of 0.2 to 0.3 μ g/L, and trichloroethylene in water from three wells at concentrations ranging from 0.2 to 4.4 μ q/L. Other volatile organic compounds were detected just once or twice and, except for methylene chloride (8.8 to 11.0 μ g/L) and 1,2dichlorobenzene (3.0 μ g/L), all were present at concentrations of less than 1.0 μ g/L. This group of synthetic chemicals are low-molecular-weight. chlorinated hydrocarbons widely used as solvents, degreasers, refrigerants, fumigants, and insecticides, and are found in numerous household products.

Disposal of these products through septic systems is a likely source for some of the detections, but spills, leaks, and improper disposal of gasoline and motor oils, paint, varnish, or finish removers, disinfectants, and cleaning agents may also be sources.

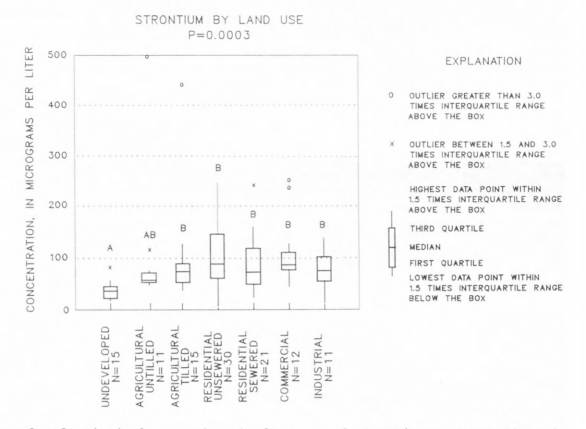


Figure 8.--Statistical comparison by land use of strontium concentrations in ground-water samples from four stratified-drift aquifers in Connecticut. (P-value is the attained significance level for that statistical comparison; N is the number of wells sampled per land-use category; sample populations are designated by letter symbols (A, B, C, and D); sample populations that share the same letter are not significantly different at the 0.05 level of significance).

Nearly half (48 percent) of the samples from wells in the unsewered residential areas submitted for GC/FID analyses contained one or more unidentified BNA organic compounds. GC/MS analyses were performed on 10 samples from wells with positive GC/FID results, and target BNA compounds (see table 2) were identified and quantified in 6 of these samples. These compounds included bis(2-ethylhexyl)phthalate at concentrations of 2.9 to 8.5 μ g/L in five of the samples, diethyl phthalate (1.9 to 5.5 μ g/L) and phenol (2.4 to 2.6 μ g/L) in two of the samples, and singular detections of di-n-butylphthalate (2.6 μ q/L) and napthalene (2.3 μ q/L). The above compounds are most commonly used as plasticizers, solvents, lubricants, and preservatives, and are found in numerous household products such as perfumes, pharmaceuticals, and moth and mosquito repellents. Trace levels of the polynuclear aromatic hydrocarbons (PAH) acenapthene, fluoranthene, and fluorene, common combustion byproducts, were also reported in one sample. In addition, most of the samples contained one or more nontarget BNA compounds, principally high-molecular weight, aliphatic hydrocarbons. Water from one well contained as many as 49 BNA compounds at concentrations ranging from less than 1.0 to 500 $\mu g/L$. The association of such compounds as fenchone (40 μ g/L), benzenepropanoic acid (500 μ g/L), trans-geraniol (10 μ g/L), para-menthan-8-ol (400 μ g/L), alpha-terpineol (200 μ g/L), and benzeneacetic acid (150 µg/L) with various phenols and alkanes suggests contamination from an adjacent septic source that receives household products, such as cleaners, toiletries, pharmaceuticals, food additives, preservatives, and flavoring agents.

Seven pesticides have also been detected in samples from the unsewered residential areas, including the insecticides chlordane, diazinon, dieldrin, and DDD, and the herbicides 2,4-D, 2,4,5-T, and silvex (table 5). Chlordane (0.1 to 0.3 $\mu g/L$), diazinon (0.01 to 0.07 $\mu g/L$), and dieldrin (0.03 to 0.063 $\mu g/L$) were each detected twice, and DDD (0.01 $\mu g/L$) once. These insecticides have commonly been used in residential areas to control pests such as termites, ants, cockroaches, yellowjackets, and garden insects. The herbicides 2,4-D (0.03 $\mu g/L$), 2,4,5-T (0.01 $\mu g/L$), and silvex (0.01 $\mu g/L$) were all detected in water from one well, where they may have been used to control weeds in residential lawns and gardens. None of the above pesticides were detected in enough samples from unsewered-residential area wells to be statistically different from other land-use areas.

Sewered areas

Ground-water quality beneath sewered residential areas is affected to a greater degree by human activities than is ground-water quality beneath the unsewered areas. The mean rank of the concentrations or the frequency of detection of specific conductance, calcium, magnesium, hardness, sodium, chloride, sulfate, dissolved solids, nitrate plus nitrite nitrogen, ammonia nitrogen, barium, boron, and strontium is significantly higher for samples from the sewered residential areas than for those from the undeveloped areas (table 5). The mean rank of the specific conductance of ground water, and the mean rank of the dissolved solids, sodium, barium, and boron concentrations are also significantly higher for the sewered areas than for the untilled agricultural areas. Mean ranks of sodium, chloride, barium, and iron concentrations are significantly higher for ground-water samples from the sewered residential areas than for those from the tilled agricultural areas. The mean rank of the barium concentrations is significantly higher for ground-water samples from the sewered residential areas than for samples from the the unsewered residential areas.

Median concentrations or frequency of detection of most physical properties and inorganic constituents are higher in water samples from the sewered areas than in those from the unsewered areas. Figure 9 shows that median concentrations of calcium (29 mg/L), hardness (93 mg/L), and sulfate (20 mg/L) are higher in ground water beneath the sewered residential land use than in the ground water beneath any other land-use area. Median concentrations of orthophosphorus (0.01 mg/L) and manganese (149 μ g/L), and detection frequencies of copper, chromium, and molybdenum are also highest in the sewered residential areas. Median nitrate plus nitrite-nitrogen concentrations are nearly identical for ground-water samples from sewered and unsewered areas. Although sewering may prevent nitrogen-rich septic effluents from reaching ground water, exfiltration from sewer lines is likely to occur in many places within the Northeast where soils are sandy and the water table may be less than 10 ft below sewer lines (Engineering Science, written commun., 1989). Effluents from leaking sewer lines could be contributing to the measured nitrate plus nitrite-nitrogen concentrations. Other sources of nitrogen, such as heavy use of lawn fertilizer in densely populated sewered areas, are masking any improvement in water quality that might result from use of sewers.

Highest reported concentrations of a number of inorganic constituents were measured in samples from the sewered residential areas. Extremely high specific conductance (3,530 $\mu\text{S/cm}$ (microsiemens per centimeter at 25 degrees Celsius)) and concentrations of sodium (480 mg/L), chloride (1,100 mg/L), and dissolved solids (2,050 mg/L) were measured in one ground-water sample from a sewered residential area. These analytical results indicate contamination of the shallow aquifer at this site from a local, concentrated salt source. A pile of sand containing deicing chemicals was left uncovered near this well for use by motorists during the previous winter. The general similarity of sodium and chloride concentrations in samples from unsewered and sewered residential areas indicates that use of deicing chemicals may be more important source of these constituents than septic-tank effluents.

Ground-water samples from the sewered residential areas also had the highest reported concentrations of several other inorganic constituents that may be derived from nonpoint sources. The highest concentrations of five trace elements—arsenic (9 μ g/L), chromium (6 μ g/L), copper (37 μ g/L), lead (210 μ g/L), and molybdenum (40 μ g/L)—were measured in samples from sewered residential areas. Concentrations of hardness (270 mg/L) and barium (400 μ g/L) were equally as high as concentrations observed in samples from unsewered areas. The broad interquartile ranges for calcium, lithium, manganese, and orthophosphorus concentrations and pH measurements indicate great variability in the sources of these constituents within the sewered residential areas.

The frequency of detection of only one volatile organic compound, chloroform, in ground-water samples from the sewered residential areas differed significantly from the frequency of detection observed in other land-use areas. Detection of low concentrations (0.2 to 1.0 $\mu g/L$) of chloroform in nearly 48 percent of the wells in the sewered residential land use was significantly higher than detections in the undeveloped, tilled agricultural, and unsewered residential areas (table 5). The widespread detection of chloroform may originate from domestic use of solvents and cleaning agents, exfiltration from water mains, or lawn irrigation with chlorinated municipal drinking water. Eight other volatile organic

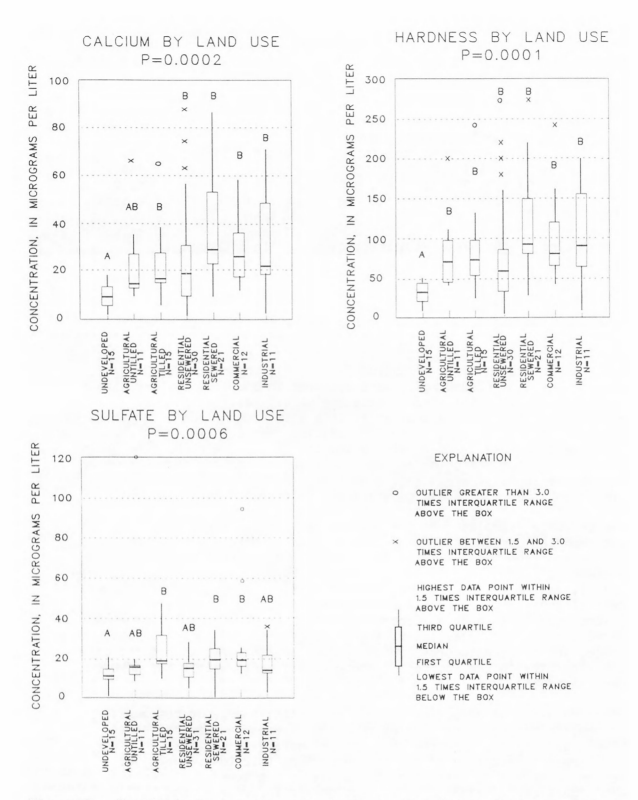


Figure 9.--Statistical comparisons by land use of calcium, hardness, and sulfate concentrations in ground-water samples from four stratified-drift aquifers in Connecticut. (P-value is the attained significance level for that statistical comparison; N is the number of wells sampled per land-use category; sample populations are designated by letter symbols (A, B, C, and D); sample populations that share the same letter are not significantly different at the 0.05 level of significance).

compounds were detected infrequently. Three compounds, including 1,1-dichloroethane (4.8 μ g/L), tetrachloroethylene (2.9 μ g/L), and trichloroethylene (19 μ g/L), were found at elevated concentrations. Trace concentrations of 1,1-dichloroethylene (0.6 μ g/L), 1,2-transdichloroethylene (0.2 and 0.3 μ g/L), 1,1-trichloroethane (0.2 to 0.4 μ g/L), trichlorofluoromethane (0.9 μ g/L), and toluene (0.2 to 0.3 μ g/L) have also been measured in samples from some sewered residential-area wells.

More than half (52 percent) of the wells in the sewered residential areas contained one or more unidentified BNA organic compounds in samples analyzed by the GC/FID method. Of five samples selected for GC/MS analysis, no BNA compounds could be identified in two samples, while the plasticizer bis(2-ethyhexyl)phthalate was found in three samples at concentrations of 10 to 420 $\mu \rm g/L$. No other target BNA compounds and few nontarget BNA compounds were reported in these samples.

Seven pesticides were also detected in ground-water samples from the sewered residential land use. The insecticides chlordane (0.1 $\mu g/L$), dieldrin (0.02 $\mu g/L$), and phorate (0.01 $\mu g/L$) were each detected in one of three wells, while the herbicides 2,4-D (0.03 $\mu g/L$), 2,4-DP (0.09 $\mu g/L$), 2,4,5-T (0.06 $\mu g/L$), and silvex (0.02 $\mu g/L$) were all detected in one well. These herbicide compounds are often combined in mixtures used to control brush and weeds, and such a combination may have been applied in the vicinity of this well.

A large number of nonpoint-contamination sources and a high-population density may explain the higher median concentrations and greater number of inorganic water-quality variables that are statistically significant in the sewered compared to the unsewered residential areas. Eckhardt and others (1989) found that specific conductance and concentrations of dissolved solids and chloride were highest in ground water on Long Island, New York beneath high-density (more than five dwellings per acre) residential areas. Population density affects water quality because contaminant loadings are determined by factors such as percentage of impervious area, traffic patterns, street-cleaning practices, type of storm-water drainage system, and deicing-chemical application rates, which differ in urban and suburban settings.

Sewered areas in this study tend to resemble an urban rather than suburban setting, in that they are more populous and have more impervious area than unsewered areas. Pervious areas of the sewered, residential land are more likely to be cultivated lawns and shrubbery than vacant fields or wooded lots, and sewered residential areas are more likely than unsewered areas to encompass small tracts of industrial and commercial land use. primary source of contaminants in these areas is chemicals leached from street refuse--the detritus, litter, and dust that accumulates on the impervious areas--that infiltrate to the water table in areas of natural (no storm or combined storm-sanitary sewers) storm drainage (Novotny and Chesters, 1981, p. 407). Street refuse commonly contains inorganic and organic matter derived from motor-vehicle emissions and road wear, decaying vegetation and animal wastes, discarded litter and garbage, deicing chemicals, leaks and spills from vehicles transporting hydrocarbons and other chemicals through urban areas, and atmospheric deposition. Other nonpoint sources include exfiltration from sanitary sewers; application of lawn and garden chemicals; improperly disposed of hydrocarbons and other

chemicals on pervious areas, in seepage pits, in dry wells, or in old septic tanks not replaced during sewering; and leaks or spills from underground storage tanks or from commercial tanks and pipelines within or traversing residential areas.

Commercial Land

The quality of ground water beneath commercial areas is adversely affected more so than it is in all other land-use areas. A variety of nonpoint sources contribute contaminants in commercial areas. Infiltration of runoff from streets, parking lots, and other impervious areas carries inorganic and organic contaminants, including deicing chemicals, metals, grease, oil and hydrocarbon fuels. Leaks and spills, particularly from underground storage tanks, and improper discharges of solvents, fuels, and other chemicals into drains, sumps, and dry wells may introduce contaminants directly into the subsurface, thereby facilitating their movement to the water table. Like the sewered residential areas, commercial areas within the stratified-drift aquifers studied are, with few exceptions, served by sanitary sewers. Removal of wastes and wastewater from commercial facilities by sanitary sewers tends to eliminate a major potential source of nonpoint contaminants. However, exfiltration from sanitary sewer lines can occur, contributing inorganic and organic contaminants to the ground water.

The mean rank of specific conductance measurements, the mean rank of concentrations of calcium, magnesium, hardness, sodium, chloride, sulfate, dissolved solids, nitrate plus nitrite nitrogen, boron, and strontium; and the frequency of detection of ammonia nitrogen is significantly higher for water samples from the commercial areas than for those from the undeveloped areas (table 5). Five of the above variables (specific conductance, sodium, chloride, dissolved solids, and boron) are also significantly higher in the commercial areas than in the untilled agricultural areas, and the mean rank of sodium, chloride, and iron concentrations are significantly higher than in the tilled agricultural areas. Figure 10 shows that median concentrations of sodium (22.5 mg/L), chloride (36 mg/L), and dissolved solids (286 mg/L) are highest for the commercial areas. The use of highway deicing chemicals, a likely source for these constituents, is generally greater in commercial areas than in other land-use areas because the major transportation routes often transect or are bounded by commercial areas. Fifty percent of the wells in the commercial areas yielded water with sodium concentrations that exceeded the State's drinking-water standard of 20 mg/L (Connecticut Department of Health Services, 1985).

Median water temperature (14 °C), and ammonia-nitrogen (0.035 mg/L) and zinc (8.2 $\mu \rm g/L)$ concentrations were also highest, but not significantly higher, in ground water from the commercial areas compared to other land-use areas. Although the median nickel concentration (8.5 $\mu \rm g/L)$ was highest in the commercial areas, and the ANOVA indicated that nickel was significantly different for one or more land uses (see table 3), the Tukey's multiple-comparison test could not distinguish a difference at the 0.05 level of significance. The large variability of nickel concentrations in ground water beneath commercial areas appears to obscure the difference between the generally high concentrations in the commercial areas and low concentrations present in the undeveloped and untilled agricultural areas. Ammonianitrogen detections, however, were significantly more common in the

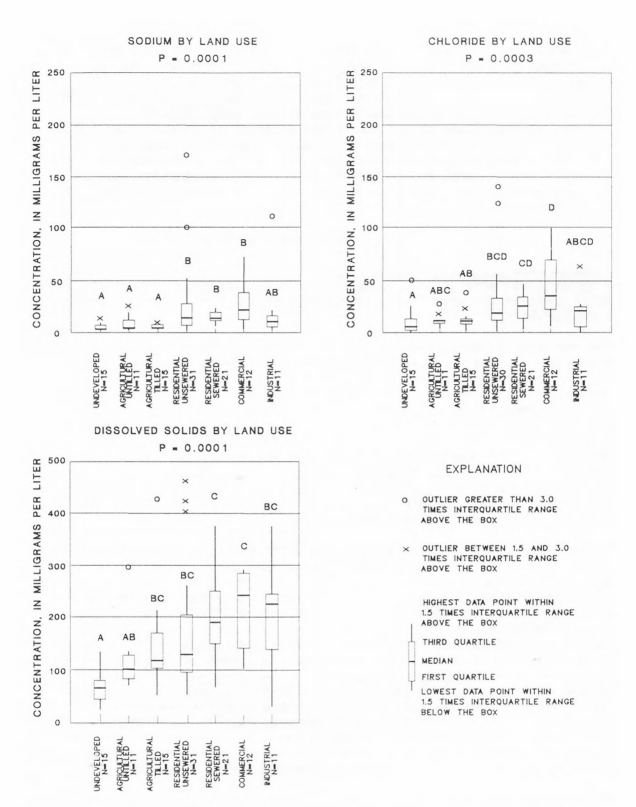


Figure 10.--Statistical comparisons by land use of sodium, chloride and dissolved solids concentrations in ground-water samples from four stratified-drift aquifers in Connecticut. (P-value is the attained significance level for that statistical comparison; N is the number of wells sampled per land-use category; sample populations are designated by letter symbols (A, B, C, and D); sample populations that share the same letter are not significantly different at the 0.05 level of significance).

commercial areas than the undeveloped areas (fig. 11). The highest reported ammonia-nitrogen concentration (1.7 mg/L) was in the commercial area, as were highest reported magnesium (23 mg/L) and beryllium (1.8 μ g/L) concentrations. Although low concentrations (less than 1.0 μ g/L) of beryllium were detected in ground-water samples from most land-use areas, the frequency of detections at concentrations equal to or greater than 1.0 μ g/L was significantly higher in samples from the commercial areas than in samples from the sewered residential areas. Variability in the specific conductance of ground water and in the concentrations of magnesium, sodium, chloride, dissolved solids, and nickel is greatest for samples from the commercial land-use areas.

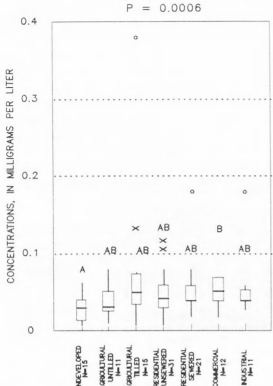
Mean rank of the MBAS concentrations in water samples for the commercial area is significantly higher than in those for the undeveloped area. Median MBAS (0.05 mg/L) concentration is equally as high as for the tilled agricultural area (fig. 11). Use of anionic-surfactant detergents in commercial areas is the most likely source of elevated MBAS concentrations in the ground water.

The frequency of detection for 3 of 11 volatile organic compounds found in ground-water samples from wells in the commercial areas is significantly higher than for samples from one or more of the other land-use areas (fig. 12). Tetrachloroethylene detections in water samples from 50 percent of the wells in the commercial area are significantly higher than tetrachloroethylene detections in samples from the undeveloped, untilled and tilled agricultural, and unsewered residential areas. An extreme tetrachloroethylene concentration of 1,300 $\mu \rm g/L$ was reported for a sample from one well in a commercial area that apparently was contaminated by drycleaning fluid. Although not indicative of the degree of contamination most commonly associated with nonpoint sources, this incident demonstrates the magnitude of possible effects on ground-water quality from human activities.

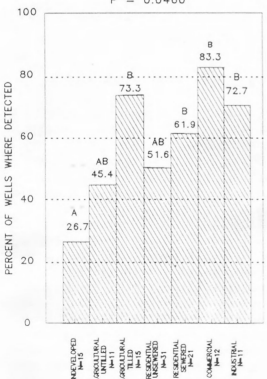
Tetrachloroethylene concentrations of 0.3 to 26 $\mu g/L$ were observed in other samples from the commercial areas. Detections of trichloroethylene, at concentrations ranging from 0.2 to 20 $\mu g/L$, were significantly more frequent in water samples from wells in the commercial areas than in samples from the undeveloped, untilled and tilled agricultural, and unsewered and sewered residential areas (fig. 12). Because trace-level concentrations (0.2 to 0.4 $\mu g/L$) of 1,2-transdichloroethylene were detected in all land-use areas, the frequency of detection in the commercial areas was significantly higher than only the frequency of detection in the unsewered residential areas. However, comparison of detections of this compound at concentrations equal to or greater than 1.0 $\mu g/L$ (fig. 12), shows that the frequency of detection of 1,2-transdichloroethylene at these concentrations in the commercial area, where four wells yielded water with concentrations of 1.8 to 55 $\mu g/L$, is significantly higher than that in the undeveloped, tilled agricultural, and unsewered and sewered residential areas (fig. 12).

Other volatile organic compounds found in ground-water samples from commercial land-use areas at concentrations exceeding 1.0 μ g/L include 1,1,1-trichloroethane (4.1 μ g/L), xylene (3.9 μ g/L), dichlorofluoromethane (2.5 μ g/L), and toluene (1.6 μ g/L). Trace concentrations of chloroform (0.2 to 0.8 μ g/L), 1,1-dichloroethane (0.5 μ g/L), 1,1-dichloroethylene (0.2 μ g/L), and vinyl chloride (0.3 μ g/L) have also been detected in ground-water samples from the commercial areas. Volatile organic compounds are used widely as solvents and degreasing agents, or are components of fuels used and stored in commercial areas.

METHYLENE-BLUE-ACTIVE SUBSTANCES BY LAND USE



AMMONIA, AS NITROGEN (detection limit 0.01 milligram per liter) P = 0.0460



EXPLANATION

- O OUTLIER GREATER THAN 3.0 TIMES INTERQUARTILE RANGE ABOVE THE BOX
- OUTLIER BETWEEN 1.5 AND 3.0 TIMES INTERQUARTILE RANGE ABOVE THE BOX

HIGHEST DATA POINT WITHIN
1.5 TIMES INTERQUARTILE RANGE
ABOVE THE BOX

THIRD QUARTILE

MEDIAN

FIRST QUARTILE

LOWEST DATA POINT WITHIN
1.5 TIMES INTERQUARTILE RANGE
BELOW THE BOX

Figure 11. -- Statistical comparisons by land use of the concentration of methylene-blue-active substances and the frequency of ammonia-nitrogen detections in ground-water samples from four stratified-drift aquifers in Connecticut. (P-value is the attained significance level for that statistical comparison; N is the number of wells sampled per land-use category; sample populations are designated by letter symbols (A, B, C, and D); sample populations that share the same letter are not significantly different at the 0.05 level of significance).

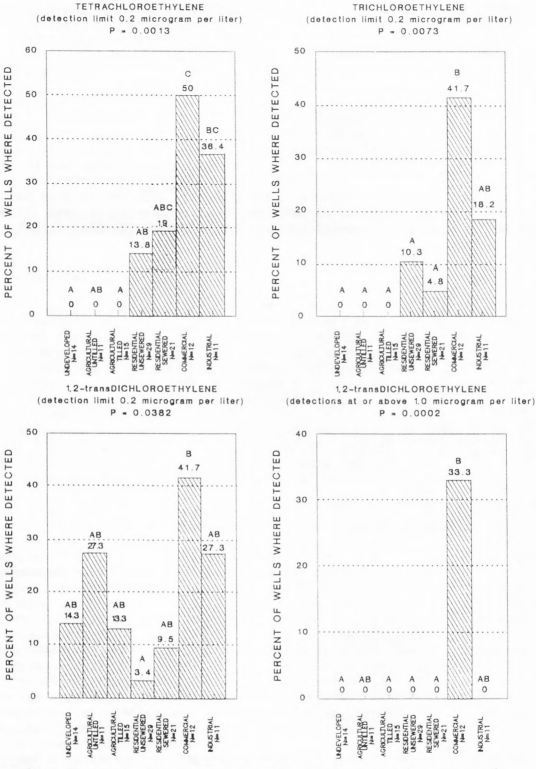


Figure 12.--Statistical comparisons by land use of the frequency of tetrachloroethylene, trichloroethylene, and 1,2-transdichloroethylene detections in ground-water samples from four stratified-drift aquifers in Connecticut. (P-value is the attained significance level for that statistical comparison; N is the number of wells sampled per land-use category; sample populations are designated by letter symbols (A, B, C, and D); sample populations that share the same letter are not significantly different at the 0.05 level of significance).

GC/FID analyses indicated that water samples from 5 of the 12 wells (42 percent) in the commercial land-use areas contained 1 or more unidentified BNA organic compounds. GC/MS analyses were performed on three samples that appeared to have sufficiently high concentrations to enable identification of BNA compounds. Target or nontarget BNA compounds were identified in two of the three samples. The plasticizer bis(2-ethylhexyl)phthalate was reported in each sample at a concentration of 21 $\mu g/L$ in one and 14.000 $\mu g/L$ in the second. The extremely high concentration observed in the latter sample, however, is not likely related to use of this compound as a plasticizer, in which the phthalate ester is encased in a polymer matrix. The occurrence of two other phthalate esters, di-n-octylphthalate (6.7 μ g/L) and di-n-butylphthalate (2.4 μ g/L), and low concentrations of several nontarget BNA compounds (nonanoic acid, nonyl alcohol, 1.2-benzenedicarboxylic acid. and an unidentified alkane) in this sample may be related to a commercial source where the compounds have nonplasticizer uses such as pesticide carriers or insect repellents, or are used in cosmetics, fragrances, or oils (Peakall, 1975).

Four herbicide compounds, atrazine (2.0 $\mu g/L$), propazine (0.1 $\mu g/L$), simazine (0.4 $\mu g/L$), and 2,4,5-T (0.4 $\mu g/L$), were each detected once in ground-water samples from the commercial areas. The first three compounds, all triazine herbicides, were detected in one well adjacent to an interstate highway. It is possible that these compounds were used to control weed growth in turf areas along the highway. The observed atrazine concentration in this well exceeds concentrations measured in the tilled agricultural areas where this compound is used extensively on corn.

Industrial Land

Industrial areas occupy a smaller part of the land surface above the stratified-drift aguifers than do any of the other land uses, but they have a disproportionate effect on ground-water quality. Several nonpoint sources of contaminants are associated with the typically high-technology, lightindustry facilities prevalent in the study areas. These industries are substantial users or producers of industrial chemicals and byproducts. Commonly, such industries are clustered within industrial parks that are isolated from residential developments, and they are not served by sanitary sewers in some areas. Improper or inadvertent discharge of chemicals or wastes through on-site septic systems and (or) industrial pits, ponds and lagoons have contaminated public-supply, domestic, and industrial wells (Handman and others, 1979, p. 46; Melvin and others, 1988, p. 196). Additional nonpoint sources of contaminants in industrial areas include leaks and spills of chemicals and hydrocarbons, and infiltration of runoff from paved areas (roadways, parking lots and storage yards), and the use of fertilizers and pesticides on lawns and ornamental shrubs within industrial properties.

The mean rank of the specific conductance measurements; the mean rank of the concentrations of calcium, magnesium, hardness, dissolved solids, boron, and strontium; and the frequency of detection of ammonia nitrogen is significantly higher for ground-water samples from the industrial areas than for samples from the undeveloped areas (see table 5). In addition, the mean rank of the boron and silica concentrations in samples from industrial-area wells is significantly higher than that for ground-water samples from the untilled agricultural areas. The mean rank of pH measurements is significantly higher for ground-water samples from the industrial areas than for samples from the unsewered residential areas. Median pH (6.9) and specific conductance (364 μ S/cm) of ground water are highest in the

industrial areas (fig. 13). Median concentrations of magnesium (5.2 mg/L), silica (18 mg/L), and boron (50 μ g/L) are also highest for water samples from the industrial land-use areas (fig. 14).

The highest measured pH (9.2); the largest concentrations of silica (35 mg/L), boron (360 $\mu g/L$), orthophosphorus (6.6 mg/L), cobalt (25 $\mu g/L$), and nickel (630 $\mu g/L$); and the largest variability in concentrations of hardness, silica, barium, and boron concentrations were observed in groundwater samples from wells in the industrial land-use areas. The frequency of detection of orthophosphorus, arsenic, beryllium, and silver is highest in ground-water samples from the industrial areas but is not significantly higher than for other land-use areas. However, comparison of beryllium detections at concentrations equal to or greater than 1.0 $\mu g/L$ (fig. 15) shows a significantly higher frequency of elevated beryllium concentrations in water samples from the industrial areas than for samples from the sewered residential areas.

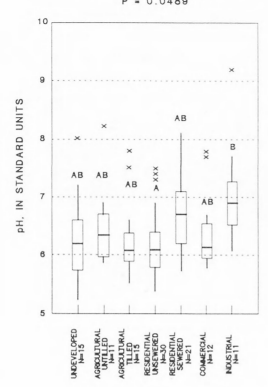
Twelve volatile organic compounds have been detected in ground-water samples from 10 of the 11 wells (91 percent) in the industrial areas, and the frequency of detection of 5 of these compounds is significantly higher for this land use than for other land uses. Detections of chloroform at concentrations of 0.2 to 1.3 μ g/L in water samples from more than 50 percent of the wells in the industrial area (fig. 15), and of tetrachloroethylene (0.2 to 4.9 μ g/L) in 36 percent of the wells (see fig. 12), are significantly more frequent than in both the undeveloped and tilled agricultural areas. Chloroform detections were also significantly more frequent in samples from the industrial-area wells than from those in the unsewered residential areas.

Comparison of the frequency of detection of three other compounds at concentrations equal to or greater than 1.0 $\mu g/L$ also demonstrates statistically significant differences between the industrial area and one or more of the other land-use areas (fig. 16). Elevated concentrations of 1,1-dichloroethane (2.4 to 4.8 $\mu g/L)$ and 1,1-dichloroethylene (1.4 to 1.8 $\mu g/L)$ occur significantly more often in ground water beneath industrial areas than beneath unsewered residential land. Concentrations of 1,1,1-trichloroethane ranging from 1.0 to 180 $\mu g/L$ were observed in samples from 36 percent of the wells in the industrial area—a frequency that was significantly higher than that in the undeveloped and the sewered residential areas. Use of these compounds as solvents and degreasing agents in industrial processes is the most probable source of their presence in ground water in this area.

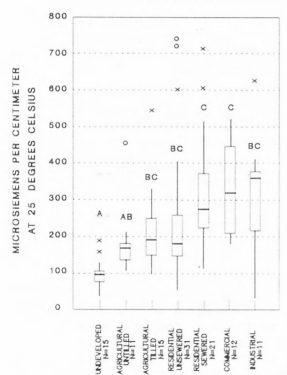
Trichloroethylene was found in water samples from two wells in the industrial land use at concentrations of 3.3 and 14 $\mu g/L$ respectively. Two additional compounds, dichlorodifluoromethane and 1,1,2-trichloroethane were detected at concentrations of 1 $\mu g/L$ or more. Trace-level concentrations (0.2 to 0.6 $\mu g/L$) of benzene, 1,2-transdichloroethylene, trichlorofluoromethane, and toluene have also been detected in ground-water samples from the industrial areas. GC/FID analyses indicated that three samples contained one or more unidentified BNA organic compounds, but only one sample was judged to contain sufficient concentrations to warrant GC/MS analysis. Subsequent analysis indicated the presence of three nontarget compounds at concentrations less than 5 $\mu g/L$ each.

Four pesticide compounds were detected in ground-water samples from the industrial land use. The herbicides prometone (0.1 to 2.0 μ g/L) and 2,4-D (0.2 μ g/L) were each detected in two samples, and atrazine (0.01 μ g/L) in one. A trace concentration of the insecticide dieldrin (0.01 μ g/L) was also detected in one sample.

FIELD pH BY LAND USE P = 0.0489



SPECIFIC CONDUCTANCE BY LAND USE P = 0.0001



EXPLANATION

- OUTLIER GREATER THAN 3.0 TIMES INTERQUARTILE RANGE ABOVE THE BOX
- OUTLIER BETWEEN 1.5 AND 3.0 TIMES INTERQUARTILE RANGE ABOVE THE BOX

HIGHEST DATA POINT WITHIN
1.5 TIMES INTERQUARTILE RANGE
ABOVE THE BOX

THIRD QUARTILE

MEDIAN

FIRST QUARTILE

LOWEST DATA POINT WITHIN
1.5 TIMES INTERQUARTILE RANGE
BELOW THE BOX

Figure 13.--Statistical comparisons by land use of pH and specific conductance of ground-water samples from four stratifieddrift aquifers in Connecticut. (P-value is the attained significance level for that statistical comparison; N is the number of wells sampled per land-use category; sample populations are designated by letter symbols (A, B, C, and D); sample populations that share the same letter are not significantly different at the 0.05 level of significance).

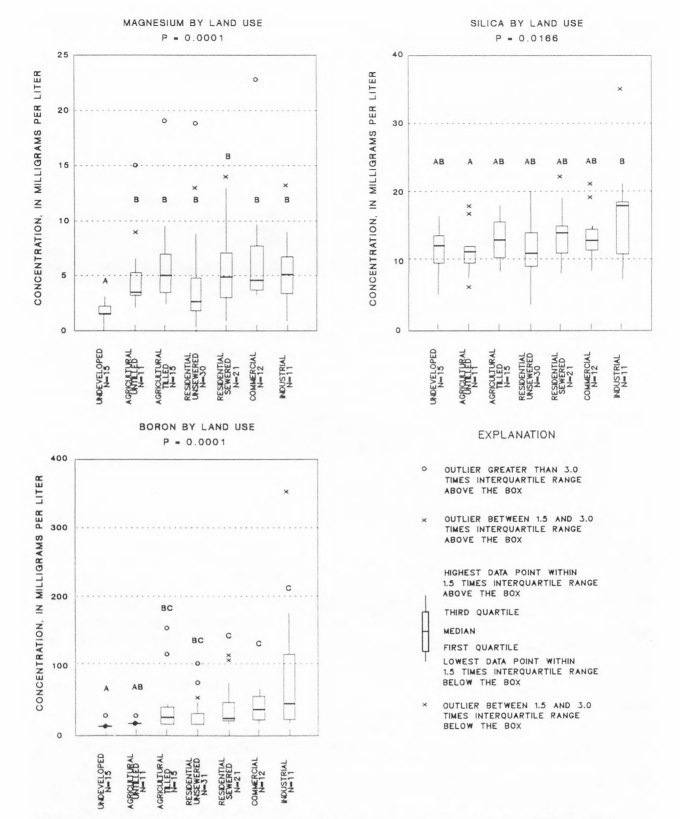


Figure 14.--Statistical comparisons by land use of magnesium, silica, and boron concentrations in ground-water samples from four stratified-drift aquifers in Connecticut. (P-value is the attained significance level for that statistical comparison; N is the number of wells sampled per land-use category; sample populations are designated by letter symbols (A, B, C, and D); sample populations that share the same letter are not significantly different at the 0.05 level of significance).

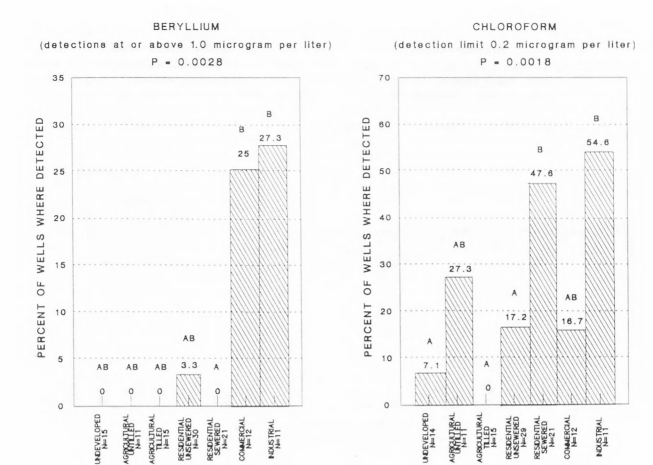
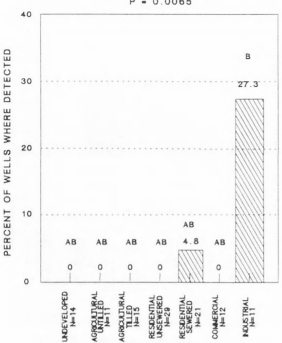
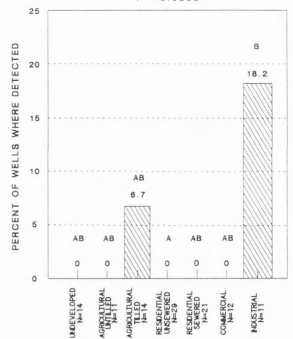


Figure 15.--Statistical comparisons by land use of the frequency of detection of beryllium at concentrations equal to or exceeding 1.0 microgram per liter and chloroform in ground-water samples from four stratified-drift aquifers in Connecticut. (P-value is the attained significance level for that statistical comparison; N is the number of wells sampled per land-use category; sample populations are designated by letter symbols (A, B, C, and D); sample populations that share the same letter are not significantly different at the 0.05 level of significance).

1,1-DICHLOROETHANE (detections at or above 1.0 microgram per liter) P = 0.0065



1,1-DICHLOROETHYLENE (detections at or above 1.0 microgram per liter) P = 0.0365



1, 1, 1-TRICHLOROETHANE (detections at or above 1.0 microgram per liter) P = 0.0195

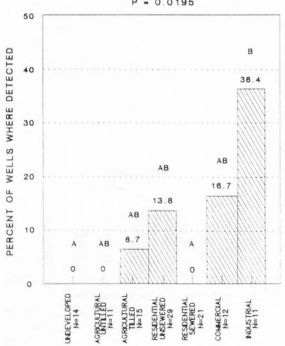


Figure 16.--Statistical comparisons by land use of the frequency of detection of 1,1-dichloroethane, 1,1-dichloroethylene and 1,1,1- trichloroethane at concentrations equal to exceeding 1.0 microgram per liter in ground-water samples from four stratifieddrift aquifers in Connecticut. (P-value is the attained significance level for that statistical comparison; N is the number of wells sampled per landuse category; sample populations are designated by letter symbols (A, B, C, and D); sample populations that share the same letter are not significantly different at the 0.05 level

of significance).

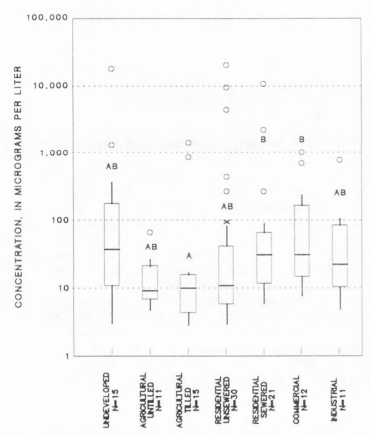
Comparisons of water-quality variables among aquifers were made to evaluate whether the effects of land use on ground-water quality are consistent for each of the stratified-drift aquifers studied. Differences in water quality among aquifers are caused by natural geochemical and biochemical factors and by differences in the type and the intensity of human activities. Statistically significant differences in concentrations of naturally occurring constituents in ground water beneath undeveloped areas of the respective aquifers could be considered evidence of the effects of natural factors. In the absence of such differences in the water quality of undeveloped areas, inconsistent relations between land use and ground-water quality among the aquifers can be considered a consequence of variations in effects of human activities.

Statistical comparison of water quality between aguifers was impeded in this study by the uneven distribution of land-use categories among the four aquifers. As a result, the number of wells per land-use category in each of the aguifers ranging from 0 to 21 (see table 1). Consequently, interaguifer comparisons were only done for the 21 uncensored or slightly censored waterquality variables which were analyzed by ANOVA. Table 3 includes p-values for the two-way ANOVA on the rank-transformed data for the effects of land use and aquifer on ground-water quality. The type III (partial sum of the squares) factorial model used by the ANOVA for these comparisons tests the significance of each factor in the model (including the "main effects" of land use and aquifer, and the interaction between these effects) in the presence of all other factors. Interaction refers to the dependence of the effect of one factor on the level of another factor and can contribute a positive (synergism) or negative (interference) increment to the level of the variable (Sokal and Rohlf, 1969, p. 306). By evaluating all factors simultaneously, the effect of each can be measured while compensating for the presence of the others.

Although 17 water-quality variables appear to be significantly related to land use, only 9 (specific conductance, calcium, magnesium, hardness, sodium, chloride, sulfate, dissolved solids, and barium) appear to be significantly related to aquifer. An interaction between land use and aquifer appears to have a significant effect on the concentrations of four variables--magnesium, silica, nickel and MBAS.

Based on the two-way ANOVA results identified in table 3, land use appears to be the only significant factor affecting the sample populations for five variables (pH, nitrate plus nitrite nitrogen, boron, iron, and strontium). For nitrogen, boron, and strontium, the magnitude and variability of concentrations under natural conditions are so small that any elevated levels are clearly the result of human contributions. The conclusion that land use is the sole influencing factor is misleading, however, for pH and iron, whose concentrations are naturally highly variable. Iron concentrations, for example, which differ by more than four orders of magnitude in the study areas, clearly reflect natural geochemical and biochemical conditions that control the solubility of iron in ground water (Hem, 1985, p. 75-84) but also are apparently influenced by land use (fig. 17).

P = 0.0189



EXPLANATION

- OUTLIER GREATER THAN 3.0 TIMES INTERQUARTILE RANGE ABOVE THE BOX
- OUTLIER BETWEEN 1.5 AND 3.0 TIMES INTERQUARTILE RANGE ABOVE THE BOX

HIGHEST DATA POINT WITHIN
1.5 TIMES INTERQUARTILE RANGE
ABOVE THE BOX

THIRD QUARTILE

MEDIAN

FIRST QUARTILE

LOWEST DATA POINT WITHIN
1.5 TIMES INTERQUARTILE RANGE
BELOW THE BOX

Figure 17.--Statistical comparison by land use of iron concentrations in ground-water samples from four stratified-drift aquifers in Connecticut. (P-value is the attained significance level for that statistical comparison; N is the number of wells sampled per land-use category; sample populations are designated by letter symbols (A, B, C, and D); sample populations that share the same letter are not significantly different at the 0.05 level of significance level).

It is likely that the statistically significant differences observed between the "elevated" iron concentrations associated with the sewered residential and commercial land uses, and the low concentrations measured in the tilled agricultural areas (see fig. 17) are the result of natural and human factors. First, neither the low iron concentrations in ground-water samples from the tilled agricultural areas nor the elevated iron concentrations in samples from the sewered residential and commercial areas are significantly different from iron concentrations in ground water from

the undeveloped area. Thus, natural variability in iron concentrations could explain the observed differences among these land-use areas if the conditions controlling iron solubility differed. It is possible that ground water in the tilled agricultural areas have higher dissolved oxygen concentrations unfavorable to iron solubility because the depth to the water table is generally shallower and less variable than it is in other land-use areas (see fig. 6b). Iron, however, is also a nonpoint-source contaminant associated with wastewater, solid wastes, and stormwater runoff.

The results of interaquifer comparisons for the one water-quality variable, magnesium, that demonstrates significant differences for all three effects--land use (p = 0.0001), aquifer (p = 0.0001), and interaction between land use and aquifer (p = 0.0386), are summarized in table 6. The interaquifer comparisons described in table 6 were obtained by segregating the data by a classification scheme that reflected both the land-use and aquifer categories, and by conducting a one-way ANOVA on the ranks of the magnesium concentrations in each composite land-use/aquifer category. Table 6 includes the median and interquartile range for each of the 17 land-use/aquifer categories for which at least 3 wells were sampled. Populations were compared by Tukey's honest significant-difference test; sample populations are designated by letter symbols (A, B, C, and D), and sample populations that share the same letter are not significantly different at a 0.05 level of significance.

The Tukey-test results indicate that the significant land-use effect is manifest in the difference between magnesium concentrations in the undeveloped land-use areas and in one or more of the developed land-use areas in the Farmington and Hockanum aquifers. Specifically, magnesium concentrations in the tilled agricultural areas of the Farmington aquifer are significantly higher than those in the undeveloped areas of that aquifer. In the Hockanum aquifer, magnesium concentrations in the untilled agricultural, sewered residential, and commercial areas are significantly higher than in the undeveloped areas. No land-use difference was observed for the Pomperaug aquifer, and no comparison can be made for the Pootatuck aquifer.

The results of the Tukey multiple comparison test for the same land uses in different aguifers show that the only significant difference is between unsewered residential areas in the Pomperaug and Farmington aguifer. Magnesium concentrations are significantly higher in the Pomperaug than in the Farmington aguifer, but the source of the higher concentrations in the Pomperaug aguifer is unclear. Interaquifer comparisons for the other three land-use areas that were sampled in both aquifers show no other significant or even consistent differences in magnesium concentrations. It is possible that the apparent significant relation of magnesium concentration to aquifer for the unsewered residential categories is the result of some characteristic inherent to that land use in the Pomperaug aquifer--for example, the density of housing with on-site septic systems may be somewhat higher in the Pomperaug than the Farmington aguifer. It is also possible, however, given the small sample size, that the difference is an artifact of the limited data. It is unlikely that the apparent significant relation of the concentrations of magnesium and the other eight variables to aquifer are the result of natural geochemical differences between these aguifers, inasmuch as no differences have been described for comparisons of groundwater quality in undeveloped lands.

Table 6.--Statistical summary of magnesium concentrations by land use and aquifer

[Concentrations in milligrams per liter; N = number of wells per land-use/aquifer category; dashes indicate insufficient number of samples to calculate statistic; IQR = interquartile range; Tukey = results of Tukey's honest significant-difference test; sample populations are designated by letter symbols (A, B, C, and D); sample populations that share the same letter are not different at the 0.05 level of significance]

			Land-us	e category			
	Undevel-			Residential			
Aquifer	oped	untilled	tilled			Commercial	Industrial
Pootatuck							
N	0	1	0	2	0	0	3
Median							8.8
IQR							5.2
Tukey							С
Pomperaug							
N	3	1	7	7	0	4	1
Median	2.5		4.4	5.4		4.85	
IQR	. 5		4.0	4.85		15.3	
Tukey	ABC		BC	C		BC	
Farmingto	n _						
N	7	5	6	20	7	3	4
Median	1.4	3.2	4.8	2.05	2.5	4.4	2.7
IQR	. 5	1.3	2.18	2.02	4.9	1.3	3.58
Tukey	А	ABC	BC	AB	ABC	ABC	ABC
Hockanum							
N	5	4	2	1	14	5	3
Median	.9	7.7			5.5	6.6	4.7
IQR	1.4	9.35			3.62	5.25	1.8
Tukey	A	C			C	C	ABC

The effect of the interaction between land use and aguifer is demonstrated in table 6 by the inconsistency of the relation of land use to magnesium concentrations for different aquifers. For example, although magnesium concentrations in the tilled agricultural areas of the Farmington aquifer are significantly higher than they are in the undeveloped areas of that aguifer, there is no difference in magnesium concentrations in groundwater samples from these land uses in the Pomperaug aguifer. Therefore, the effect of tilled agricultural land use on the magnesium concentration of ground water depends on the aguifer sampled, and the results may not be extrapolated to other aguifers. Similarly, the relation between concentrations of magnesium and land use are not consistent among the aguifers for other land-use categories as well--for the untilled agricultural and sewered residential land-use areas with respect to undeveloped land-use areas in the Farmington and Hockanum aguifers, and for the commercial areas with respect to undeveloped areas of all three aguifers sampled. However, the apparent interaction between land use and aguifer may be attributable to the small sample size and the inherent large variability of the magnesium concentrations demonstrated by the wide interquartile ranges for the very limited data available for many of the land-use/aguifer categories.

Because only four variables appear to be significantly related to the interaction of land use and aquifer, and three of these did not show significant differences attributable to aquifer alone, it is the author's conclusion that the effects of land use on ground-water quality are largely independent of aquifer. However, the small number of wells sampled in most land-use/aquifer categories and the limited number of variables compared constrains the extrapolation of these results to other stratified-drift aquifers. Additional data are needed in order to establish if the relations between land use and water quality demonstrated by this study of four stratified-drift aquifers in Connecticut can be extrapolated to similar aquifers throughout the northeastern United States.

The data needed to establish the validity of relations between land use and ground-water quality on a regional scale could best be obtained in two ways. First, augmenting the number of wells per land use in each aquifer to approximately 20 to 30 wells per land-use category per aquifer would improve confidence in the statistical inference. Secondly, inclusion of several additional stratified-drift aquifers in areas of differing geologic terrane, hydrologic setting, and population density would ensure that the statistical comparisons included conditions representative of the entire region.

Although stratified-drift aquifers are the most productive aquifers in the region and supply some of the most populated areas, they have limited distribution. In most of the glaciated Northeast, where stratified-drift aquifers are absent, shallow till and bedrock aquifers are the only source of domestic and commercial self-supplied water. A complete appraisal of the effects of land use on ground-water quality in the region needs to include information on the effect of nonpoint-source contaminants on water quality in these aguifers as well.

SUMMARY AND CONCLUSIONS

The quality of water in four stratified-drift aguifers in Connecticut has been affected by nonpoint-source, inorganic and organic contaminants that can be related to human activities within specific land-use areas. Differences observed in the concentration, the frequency of detection, or the frequency of observations equal to or exceeding 1 μ g/L of 27 waterquality variables for sample populations from seven land-use categories are statistically significant at the 0.05 level of significance. Water-quality variables that differ significantly for one or more land-use areas include the physical properties pH and specific conductance; the major inorganic constituents calcium, magnesium, hardness, sodium, chloride, sulfate, silica, and dissolved solids; the nutrients nitrate plus nitrite nitrogen and ammonia nitrogen; the trace elements barium, beryllium, boron, iron, nickel, and strontium; methylene-blue-active substances; the volatile organic compounds chloroform, 1,1-dichloroethane, 1,1-dichloroethylene, 1,2transdichloroethylene, tetrachloroethylene, 1,1,1-trichloroethane, and trichloroethylene; and the pesticide atrazine. Other organic contaminants were detected in water samples from one or more wells in this study, but their frequencies of detection within any land-use area were not statistically different. These variables include 14 volatile organic compounds, 13 pesticides, and 9 base-neutral and acid-extractable compounds.

The following relations between ground-water quality and land use were observed:

- (1) The quality of ground water in stratified-drift aquifers in undeveloped areas is assumed to be unaffected by human activities and is considered to represent background conditions. Median concentrations and detection frequencies of 19 physical properties or inorganic constituents in ground water are lowest for samples from the undeveloped areas. The mean rank of the concentrations or the frequency of detection of 13 of these properties or constituents is significantly lower for ground-water samples from the undeveloped areas than it is for those from 1 or more of the other 6 land-use areas. Fewer organic compounds were detected in ground water from the undeveloped areas than in ground water from all other land-use areas.
- (2) The quality of ground water in agricultural areas reflects tillage practices, with greater effect occurring beneath tilled areas than untilled areas because the former endure more intense agricultural activities, including heavier fertilizer and pesticide applications, than do untilled areas. Although median concentrations of most inorganic constituents are elevated in the untilled areas with respect to the undeveloped areas, only concentrations of magnesium and hardness are significantly higher. Organic compounds were detected in few ground-water samples from the untilled agricultural areas and the source of those compounds is ambiguous. No organic compound was detected significantly more often in samples from the untilled agricultural areas than in samples from the undeveloped areas. Only one pesticide was detected in ground-water samples from the untilled agricultural areas $(0.1~\mu\text{g/L})$ of cyanazine in one well), probably reflecting the minimal use of such agricultural chemicals in these areas.

Conversely, the mean rank of the specific conductance measurements; the mean rank of the concentrations of calcium, magnesium, hardness, sulfate, dissolved solids, nitrate plus nitrite nitrogen, boron, and strontium; and the frequency of detection of ammonia nitrogen in ground water beneath tilled agricultural areas is significantly higher than it is in the undeveloped areas. Median nitrate plus nitrite-nitrogen concentration is highest for the tilled agricultural areas where 20 percent of the wells yielded water with nitrate plus nitrite-nitrogen concentrations higher than 10 mg/L. The frequency of detecting the herbicide atrazine—at low concentrations (0.1 to 0.2 $\mu \text{g/L}$) in 33 percent of the wells in the tilled agricultural areas—was significantly higher than in the undeveloped areas, where it was undetected. The insecticides dieldrin and 1,2-dichloropropane have also been detected infrequently in ground-water samples from the tilled agricultural areas.

(3) The quality of ground water beneath residential areas is widely affected by inorganic and organic nonpoint-source contaminants. Sewered residential areas have a greater effect on water quality than unsewered areas. The greater housing density, more heterogeneity of land use, and higher number of potential nonpoint-contaminant sources in the sewered residential areas compared to the unsewered residential areas may obscure any improvements in water quality attributable to removal of wastewater by municipal sewer systems. The mean rank of the specific conductance measurements and the mean rank of the concentrations of calcium, magnesium, hardness, sodium, chloride, dissolved solids, nitrate plus nitrite nitrogen, boron, and strontium is significantly higher for samples from the unsewered and sewered residential areas than for those from the undeveloped areas. Median concentrations or detection frequencies for most of the physical properties and inorganic constituents of ground water are higher in the sewered than for the unsewered areas. The similarity of nitrate plus nitrite-nitrogen concentrations for sewered and unsewered residential areas may indicate exfiltration from sanitary sewers. Alternatively, some additional source of nitrogen, possibly increased lawn and garden fertilizer use in the more densely populated sewered areas, is replacing the release of nitrogen from on-site domestic septic systems as the principal source of this constituent in ground water. The similarity of sodium and chloride concentrations for sewered and unsewered areas suggests that use of deicing chemicals may be a more important source than septic-tank effluents for these constituents.

Although 62 percent of the wells in the unsewered residential areas yielded ground-water samples containing one or more volatile organic compound, the frequency of detections was not statistically significant for any specific compound. Chloroform detections in 47 percent of the wells in the sewered residential areas, however, were significantly more common than in the undeveloped, tilled agricultural, and unsewered residential areas.

(4) The quality of ground water is most adversely affected by human activities in commercial areas. Although most commercial areas are served by sanitary sewers that remove wastes and wastewater from the immediate area, exfiltration of wastewater from leaky sewer lines may be contributing inorganic and organic contaminants to the ground water. Numerous other nonpoint sources may also contribute contaminants to the soil and, ultimately to the aquifer beneath commercial land-use areas. The mean rank of specific conductance measurements; the mean rank of the concentrations of

calcium, magnesium, hardness, sodium, chloride, sulfate, dissolved solids, nitrate plus nitrite nitrogen, boron, and strontium; and the frequency of detection of ammonia nitrogen is significantly higher in ground-water samples from the commercial areas than in samples from the undeveloped areas. Measurements and concentrations of five of the above water-quality variables--specific conductance, sodium, chloride, boron, and dissolved solids--were significantly higher in ground-water samples from the commercial areas than in samples from the untilled agricultural areas. The mean rank of the concentrations of sodium, chloride, and iron in water samples from the commercial-area wells is significantly higher than in samples from the tilled agricultural areas. Median concentrations of sodium (22.5 mg/L), chloride (36 mg/L), and dissolved solids (286 mg/L) were highest in ground water from the commercial areas. Beryllium detections at concentrations equal to or exceeding 1.0 μ g/L were significantly more frequent in the commercial areas than in the sewered residential areas.

The mean rank of MBAS concentrations is significantly higher for ground-water samples from the commercial areas than it is for samples from the undeveloped areas. Detections of tetrachloroethylene in samples from 50 percent of the commercial-area wells were significantly more frequent than they were in samples from wells in the undeveloped, untilled and tilled agricultural, and unsewered residential areas. Trichloroethylene detections were significantly more frequent in ground water beneath the commercial areas than they were in the undeveloped, untilled and tilled agricultural, and unsewered and sewered residential areas. A third volatile organic compound, 1,2-transdichloroethylene, was widely detected at concentrations lower than 1.0 µg/L in ground-water samples from all land-use areas, so that the frequency of detection in the commercial areas was different only with respect to the unsewered residential areas. Comparing detections of 1,2transdichloroethylene at concentrations equal to or exceeding 1.0 μ g/L, however, shows that elevated concentrations of this compound were present only in samples from the commercial areas. A significant difference was evident between the high frequency of elevated 1,2-transdichloroethylene detections in samples from wells in the commercial areas (33 percent) and those from the undeveloped, tilled agricultural, and unsewered and sewered residential areas.

(5) Industrial land occupies less of the study areas than do the other land-use areas, but it has a disproportionately large effect on ground-water quality. The mean rank of the specific-conductance measurements; the mean rank of the concentrations of calcium, magnesium, hardness, dissolved solids, boron, strontium; and the frequency of detection of ammonia nitrogen is significantly higher in ground-water samples from the industrial areas than it is in samples from the undeveloped areas. The mean rank of boron and silica concentrations is significantly higher for ground-water samples from wells in the industrial areas than in the untilled agricultural areas, and the mean rank of pH is significantly higher for ground water in the industrial areas than in the unsewered residential areas. Beryllium detections at concentrations equal to or exceeding 1.0 $\mu g/L$ are significantly more frequent in the industrial areas than in the sewered residential areas.

In the industrial areas, water samples from 91 percent of the wells contained one or more of 12 volatile organic compounds. Chloroform detections in samples from more than half of the industrial-area wells were significantly more frequent than in samples from the undeveloped, tilled

agricultural, and unsewered residential areas. The frequency of tetrachloroethylene detections, found in 36 percent of the industrial-area wells, was significantly higher than in samples from wells in the undeveloped and tilled agricultural areas. The frequency of 1,1,1-trichloroethane detections at concentrations equal to or greater than 1.0 $\mu g/L$ was significantly higher in industrial areas than in undeveloped and sewered residential areas, and 1,1-dichloroethane and 1,1-dichloroethylene were found significantly more often at similarly elevated concentrations in ground water beneath the industrial areas than the unsewered residential areas.

Interaguifer comparisons for those water-quality variables using twoway ANOVA indicate that effects of aguifer on ground-water guality were significant for nine variables: specific conductance, calcium, magnesium, hardness, sodium, chloride, sulfate, dissolved solids, and barium. magnesium, however, appears to be significantly related to the interaction between land use and aquifer. Three additional variables, silica, nickel, and MBAS, appear to be significantly related to the interaction between land use and aguifer, but not for the aguifer alone. The significant differences among stratified-drift aquifers for some variables may mean that the effect of human activities associated with specified land uses is not the same in each aguifer. Consequently, relations among land use and some water-quality variables may not be extrapolated beyond aguifers in the study areas. However, the general lack of significant interaction between land use and aquifer, and the uncertainty generated by the small sample size for data on interaquifer differences, indicate that effects of land use on ground-water quality are largely independent of aquifer. Where differences in groundwater quality do exist among aquifers, they may reflect the intensity of human activities or may be artifacts of the limited data. Additional study is needed to establish the transfer value of relations between land use and water quality that were demonstrated for four stratified-drift aquifers in Connecticut to similar aguifers in the northeastern United States.

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SUPPLEMENTAL TABLES 65

Table 5.--Statistical summary by land use of water-quality variables detected in 1 or more of 85 to 116 observation wells in 4 stratified-drift aquifers in Connecticut

[All constituents are dissolved except as indicated. When multiple analyses are available from one well, the median value is used to calculate the statistic. Sample populations are designated by letter symbols (A, B, C, and D); sample populations that share the same letter symbol are not significantly different at alpha = 0.05. Asterisk indicates some constituent concentrations below analytical detection limit and therefore statistic estimated using method of Helsel and Gilliom (1986). Dashes indicate insufficient data to calculate statistic. Units: °C, degrees Celsius; μ S/cm, microsiemens per centimeter at 25°C; mg/L, milligrams per liter; μ g/L, micrograms per liter]

Water-quality variable,			L	and-use catego	ory		
units, and	Undevel-	Agricul	tural	Reside	ntial	Commercial	Industrial
statistic	oped	untilled	tilled	unsewered	sewered		
		РНУ	SICAL PROP	ERTIES			
Temperature (°C)							
Number of wells	15	10	15	31	21	12	11
Number of detections	15	10	15	31	21	12	11
Median	12	13	12.5	13	13	14	14
Interquartile range	2.5	1.6	1.4	2.6	1.6	1.7	1.7
Population	А	A	A	A	А	A	A
Specific conductance (µ5	S/cm at 25 °C						
Number of wells	15	11	15	31	21	12	11
Number of detections	15	11	15	31	21	12	11
Median	97	167	191	182	280	319.5	364
Interquartile range	40	62	105.5	136.5	189.5	267.2	181
Population	А	AB	ВС	ВС	С	С	ВС
pH (standard units)							
Number of wells	15	11	15	30	21	12	11
Number of detections	15	11	15	30	21	12	11
Median	6.2	6.4	6.1	6.1	6.7	6.2	6.9
Interquartile range	.9	.8	.5	.7	1.0	7	.8
Population	AB	AB	AB	А	AB	AB	В
		MAJOR IN	ORGANIC CO	NSTITUENTS			
Calcium (mg/L)							
Number of wells	15	11	15	30	21	12	11
Number of detections	15	11	15	30	21	12	11
Median	9.2	15	17	19	29	25.5	22
Interquartile range	8.9	19	14	21.6	40	22.3	31
Population	А	AB	В	В	В	В	В
Magnesium (mg/L)							
Number of wells	15	11	15	31	21	12	11
Number of detections	15	11	15	30	21	12	11
Median	1.5	3.5	5.1	2.8	4.9	4.6	5.2
Interquartile range	1.0	3.3	3.7	3.2	4.3	4.7	4.4
Population	Α	В	В	В	В	В	В

Table 5.--Statistical summary by land use of water-quality variables detected in 1 or more of 85 to 116 observation wells in 4 stratified-drift aquifers in Connecticut--Continued

Water-quality variable,	Land-use category									
units, and	Undevel-	Agricult	tural	Reside	ntial	Commercial	Industria			
statistic	oped	untilled	tilled	unsewered	sewered					
	М	AJOR INORGANI	IC CONSTIT	UENTScontin	ued					
Hardness, as CaCO3 (mg/L))									
Number of wells	15	11	15	30	21	12	11			
Number of detections	15	11	15	30	21	12	11			
Median	32	69	72.5	61	93	82	91			
Interquartile range	26	56	49.5	57.8	90	66.2	108			
Population	A	В	В	В	В	В	В			
Sodium (mg/L)										
Number of wells	15	11	15	30	21	12	11			
Number of detections	15	11	15	30	21	12	11			
Median	4.6	5.9	5.6	15	14	22.5	11			
Interquartile range	4.8	9.9	2.4	21.4	8.0	28.3	13.7			
Population	A	А	А	В	В	В	AB			
Chloride (mg/L)										
Number of wells	15	11	15	31	21	12	11			
Number of detections	15	11	15	31	21	12	11			
Median	7.1	12	11.5	22	27	36	22			
Interquartile range	15.4	4.0	5.5	21	20.5	50.5	21.8			
Population	А	ABC	AB	BCD	CD	D	ABCD			
Sulfate (mg/L)										
Number of wells	15	11	15	31	21	12	11			
Number of detections	15	11	15	31	21	12	11			
Median	11	15	19	15	20	19.5	15			
Interquartile range	5.9	6.0	17	9.0	11	8.5	14			
Population	A	AB	В	AB	В	В	AB			
Silica, as SiO2 (mg/L)										
Number of wells	15	11	15	30	21	12	11			
Number of detections	15	11	15	30	21	12	11			
Median	12	11	13	11	14	13	18			
Interquartile range	4.5	2.3	6.4	5.1	4.5	3.5	8.0			
Population	AB	A	AB	AB	AB	AB	В			
Dissolved solids, residue	e at 180 °C	(mg/L)								
Number of wells	15	11	15	31	21	12	11			
Number of detections	15	11	15	31	21	12	11			
Median	64	104	118	134	189	286	253			
Interquartile range	38	52	74	119.5	115.5	147.4	120			
Population	A	AB	ВС	ВС	С	С	ВС			
. opu rac ron										

Table 5.--Statistical summary by land use of water-quality variables detected in 1 or more of 85 to 116 observation wells in 4 stratified-drift aguifers in Connecticut--Continued

Water-quality			Lan	d-use categor	У		
variable, units, and	Undevel-	Agricultu	ıral	Resident	ial	Commercial	Industrial
statistic	oped	untilled	tilled	unsewered	sewered	oommer or a r	1110030110
			NUTRIENTS				
Nitrate plus nitrite, as	s N (mg/L)						
Number of wells	15	11	15	31	21	12	11
Number of detections	8	11	14	29	19	10	9
Median	.11	1.5	4.7	2.4	2.3	3.4	1.3
Interquartile range	.56	3.0	6.5	4.4	2.3	1.9	3.0
Population	А	AB	В	В	В	В	AB
Ammonia, as NH4 (mg/L)							
Number of wells	15	11	15	31	21	12	11
Number of detections	4	5	11	16	13	10	8
Median	.001*	.009*	.010	.010	.018	.035	.030
Interquartile range	.012*	.043*	.038	.115*	.086*	.049	.041
Population	Α	AB	В	AB	В	В	В
Orthophosphorous, as PO-	4 (mg/L)						
Number of wells	15	11	15	31	21	12	11
Number of detections	7	4	3	10	11	6	6
Median	.009*	.008*	.002	.007*	.010	.007*	.005
Interquartile range	.004*	.004*	.006		.024*		.019
Population	А	А	Α	А	А	Α	Α
		TRA	ACE ELEMENT	rs			
Arsenic (μg/L)							
Number of wells	15	11	15	31	21	12	11
Number of detections	2	2	2	9	3	3	4
Median				.28*			.55*
Interquartile range				.89*			.66*
Population	А	А	А	А	А	A	А
Barium (µg/L)							
Number of wells	15	11	15	30	21	12	11
Number of detections	15	11	15	30	21	12	11
Median	25	27	37	41.5	120	140	170
Interquartile range	40	73	69	56.6	143	104.9	161
Population	А	А	А	А	В	AB	AB
Seryllium (µg/L)							
Number of wells	15	11	15	30	21	12	11
Number of detections	1	0	2	5	1	4	4
Median				.10*		.25*	.55*
Interquartile range				.21*		.60*	.65*
Population	А	A	А	A	А	A	Α

Table 5.--Statistical summary by land use of water-quality variables detected in 1 or more of 85 to 116 observation wells in 4 stratified-drift aquifers in Connecticut--Continued

Water-quality variable,			Li	and-use catego	ry		
units, and	Undevel-	Agricul	tural	Residen	tial	Commercial	Industria
statistic	oped	untilled	tilled	unsewered	sewered		
		TRACE	ELEMENTS	continued			
Boron (µg/L)							
Number of wells	15	11	15	31	21	12	11
Number of detections	3	6	11	22	19	11	8
Median	6.6*	20	30	20	30	40	50
Interquartile range	5.7*	10*	39.4*	29.1*	27.5*	37.5	137.3
Population	А	AB	ВС	BC	C	С	C
Cadmium (µg/L)							
Number of wells	15	11	15	30	21	12	11
Number of detections	1	2	0	5	2	0	1
Median				.06*			
Interquartile range				.28*			
Population	А	A	А	Α	А	A	А
r opu rac ron	^	٨	^	^	^	^	^
Chromium $(\mu g/L)$							
Number of wells	15	11	15	31	21	12	11
Number of detections	4	2	1	6	8	3	2
Median	.27*			.15*	.62*	.75*	
Interquartile range	.90*			.43*	1.73*	1.54*	
Population	А	А	А	A	А	А	А
Cobalt (µg/L)							
Number of wells	15	11	15	30	21	12	11
Number of detections	3	0	0	2	2	0	2
Median	1.04*						
Interquartile range	1.63*						
Population	Α	Α	A	Α	A	A	A
ropulation	A	A	A	A	A	^	^
Copper (µg/L)							
Number of wells	15	11	15	30	21	12	11
Number of detections	2	0	0	1	3	0	0
Median					.61*		
Interquartile range					2.72*		
Population	А	А	А	А	А	А	А
Iron (µg/L)							
Number of wells	15	11	15	30	21	12	11
Number of detections	14	11	14	27	21	12	11
Median	37	9.0	10	11	32	31	23
Interquartile range	169	14	11.5	36	54	155	76
Population	AB	AB	А	AB	В	В	AB
Load (wall)							
Lead (µg/L)	1.5	1.1	1.5	20	21	12	11
Number of wells	15	11	15	30	21	12	11 2
Number of detections	3	1	0	3	4	2	
Median	2.53*			1.18*	.14*		
Interquartile range	4.43*			2.58*	2.15*		
Population	А	A	А	А	А	А	А

Table 5.--Statistical summary by land use of water-quality variables detected in 1 or more of 85 to 116 observation wells in 4 stratified-drift aquifers in Connecticut--Continued

Water-quality		Land-use category									
variable, units, and	Undevel-	Agricult	ural	Resident	rial	Commercial	Industria				
statistic	oped	untilled	tilled	unsewered	sewered	Commercial	industria				
		TRACE E	LEMENTSC	continued							
Lithium $(\mu g/L)$	4.5		4.5	20	0.1	1.0					
Number of wells	15	11	15	30	21	12	11				
Number of detections	2	6	7	10	12	8	5				
Median		6.0	3.8*	2.2*	4.0	5.5	3.7*				
Interquartile range		3.8*	3.2*	4.7*	5.6*	3.6*	2.5*				
Population	А	А	А	А	А	А	А				
Manganese (µg/L)											
Number of wells	15	11	15	30	21	12	11				
Number of detections	15	9	15	29	21	12	11				
Median	45	4.0	22	61.5	149	93.2	45				
Interquartile range	215.5	116.5	28	190	488	125.5	161.5				
Population	Α	A	A	A	A	A	A				
Mercury (µg/L)			1.5	20	0.1	10					
Number of wells	15	11	15	30	21	12	11				
Number of detections	3	4	2	6	2	1	1				
Median		.018*		.001*							
Interquartile range		.097*		.002*							
Population	А	А	Α	А	А	А	А				
Molybdenum (μ g/L)											
Number of wells	15	11	15	30	21	12	11				
Number of detections	0	0	0	1	3	0	0				
Median					.50*						
Interquartile range					2.49*						
Population	А	A	А	А	А	А	А				
Nichal ()											
Nickel (µg/L)	15	1.1	1.0	21	21	12	11				
Number of wells	15	11	15	31							
Number of detections	14	8	14	27	19	10	7				
Median	2.0	2.0	3.0	3.0	4.0	8.5	1.0				
Interquartile range	2.0	2.6	6.0	9.5	5.0	13	3.7*				
Population	А	А	А	А	А	А	А				
Silver (µg/L)											
Number of wells	15	11	15	31	21	12	11				
Number of detections	0	1	0	5	3	0	3				
Median				.43*	~ ~						
Interquartile range				.39*							
Population	А	А	А	А	А	А	А				
Stronting (mg/l)											
Strontium (µg/L)	15	1.1	16	20	21	12	11				
Number of wells	15	11	15	30	21						
Number of detections	15	11	15	30	21	12	11				
Median	37	56	76	91	76	89	79				
Interquartile range	25	23	45	90.8	80.5	42	57				
Population	Α	AB	В	В	В	В	В				

Table 5.--<u>Statistical summary by land use of water-quality variables detected in 1 or more of 85 to 116 observation wells in 4 stratified-drift aquifers in Connecticut--Continued</u>

Water-quality variable,			La	nd-use categor	ry		
units, and	Undevel-	Agricult	ural	Resident	tial	Commercial	Industria
statistic	oped	untilled	tilled	unsewered	sewered		
		TRACE E	LEMENTSC	ontinued			
Vanadium (µg/L)							
Number of wells	15	11	15	30	21	12	11
Number of detections	0	0	0	1	1	0	0
Median							
Interquartile range							
Population	Α	А	Α	Α	А	А	А
Zinc (µg/L)							
Number of wells	15	11	15	30	21	12	11
Number of detections	13	10	14	28	21	11	11
Median	6.0	7.0	7.0	8.0	7.0	8.2	7.0
Interquartile range	8.5	5.0	4.8	7.0	5.0	4.5	4.5
Population	А	A	А	А	А	A	А
		ORG	SANIC COMPO	LINDS			
		3113					
Total organic carbon (me							
Number of wells	15	10	15	31	21	12	11
Number of detections	15	10	15	31	21	12	11
Median	.60	1.20	.80	1.00	1.00	1.10	1.40
Interquartile range	.55	1.20	.70	.92	1.20	.75	1.30
Population	А	А	А	А	А	А	А
Methylene-blue-active s	ubstances (mg	/L)					
Number of wells	15	11	15	31	21	12	11
Number of detections	15	11	14	31	21	12	11
Median	.030	.030	.050		.040	.050	.040
Interquartile range	.025	.025	.040		.020	.030	.015
Population	A	AB	AB	AB	AB	В	AB
		VOLATILE	ORGANIC C	OMPOUNDS			
Panzana (ug/l)							
Benzene (µg/L) Number of wells	14	11	15	29	21	12	11
Number of detections	14	11	0	1	0	0	1
Median							1
Interquartile range Population	 A	Α	Δ	Α	 A	Α	Α
ropulation	A	A	А	A	А	А	A
Chloroethane (µg/L)							
Number of wells	14	10	15	29	21	12	11
Number of detections	0	1	0	0	0	0	0
Median							
Interquartile range							
Population	А	А	А	А	А	А	A

Table 5.--Statistical summary by land use of water-quality variables detected in 1 or more of 85 to 116 observation wells in 4 stratified-drift aguifers in Connecticut--Continued

Water-quality variable,			L	and-use categ	ory		
units, and statistic	Undevel- oped	Agricul untilled	tural tilled	Reside	ntial sewered	Commercial	Industria
		VOLATILE ORGA	ANIC COMPO	UNDScontinu	ed		
Chloroform $(\mu g/L)$							
Number of wells	14	11	15	29	21	12	11
Number of detections	1	3	0	5	10	2	6
Median		.11*		.02*			.20
Interquartile range		.13*		.09*			.34*
Population	А	AB	А	A	В	AB	В
Dichlorodifluoromethane	(µg/L)						
Number of wells	14	11	15	29	21	12	11
Number of detections	1	0	1	1	0	1	1
Median							
Interquartile range							
Population	Α	Α	Α	А	Α	Α	Α
1,2-Dichlorobenzene (µg,	/L)						
Number of wells	14	11	14	28	21	12	11
Number of detections	0	0	1	2	0	0	0
Median							
Interquartile range			'				
Population	Α	Α	А	А	А	А	Α
1,1-Dichloroethane (μ g/	1.)						
Number of wells	14	11	15	29	21	12	11
Number of detections	0	0	1	1	1	1	3
Median							.03*
Interquartile range							2.40*
Population	А	А	А	А	А	Α	А
1 2 0:-13							
1,2-Dichloroethane (μ g/		1.1	1.5	29	21	12	11
Number of wells	14 0	11 0	15 0	1	0	0	11
Number of detections Median			U	1	U	0	U
Interquartile range							
Population	Α	A	A	A	A	A	A
1,1-Dichloroethylene (μ							
Number of wells	14	11	15	29	21	12	11
Number of detections	0	0	1	0	1	1	2
Median							
Interquartile range							
Population	А	А	А	А	А	А	А
1,2-Dichloropropane (µg	/L)						
Number of wells	14	11	15	29	21	12	11
Number of detections	1	0	3	2	0	0	0
Median							
Interquartile range							
Population	Α	A	A	A	A	Α	Α

Table 5.--Statistical summary by land use of water-quality variables detected in 1 or more of 85 to 116 observation wells in 4 stratified-drift aquifers in Connecticut--Continued

Water-quality variable,			L	and-use catego	ory		
units, and statistic	Undevel- oped	Agricult untilled	tilled	Resider unsewered	ntial sewered	Commercial	Industrial
		VOLATILE ORGA	ANIC COMPO	UNDScontinue	ed		
1,2-transDichloroethyler							
Number of wells	14	11	15	29	21	12	11
Number of detections	2	3	2	1	2	5	3
Median		.11*				.09*	.14*
Interquartile range		.13*				2.84*	.11*
Population	AB	AB	AB	А	AB	В	AB
Ethylbenzene (µg/L)							
Number of wells	14	11	15	29	21	12	11
Number of detections	0	0	0	1	0	0	0
Median							
Interquartile range							
Population	А	Α	А	А	А	A	А
· opa · ac · on	,,	~	.,				
Methylene chloride (µg/	L)						
Number of wells	15	11	15	30	21	12	11
Number of detections	0	0	0	2	0	0	0
Median							
Interquartile range							
Population	Α	Α	Α	Α	Α	Α	Α
Tetrachloroethylene (µg,	/1.)						
Number of wells	14	11	15	29	21	12	11
Number of detections	0	0	0	4	4	6	4
Median				.09*		.16*	.03*
Interquartile range				.13*		9.40*	.60*
Population	Α	AB	А	AB	ABC	C	ВС
				,,_			
1,1,2,2-Tetrachloroetha	ne $(\mu g/L)$						
Number of wells	14	11	15	29	21	12	11
Number of detections	0	0	0	1	0	0	. 0
Median							
Interquartile range							
Population	А	А	А	А	А	А	Α
1,1,1-Trichloroethane (ua /1)						
Number of wells	14	11	15	29	21	12	11
Number of detections	0	1	1	6	6	3	5
Median		1		.03*		7	.03*
Interquartile range				.16*			2.80*
Population	Α	A	Α	Α	Α Α	A	Α
Τοραταστοπ	^	A	7	7	,	,,	,,,
1,1,2-Trichloroethane (μg/L)						
Number of wells	14	11	15	29	21	12	11
Number of detections	0	0	0	0	0	0	1
Median							
Interquartile range							
Population	Α	Α	А	Α	А	А	А

Table 5.--Statistical summary by land use of water-quality variables detected in 1 or more of 85 to 116 observation wells in 4 stratified-drift aquifers in Connecticut--Continued

Land-use category									
Undevel-	Agricult	ural	Residential		Commercial	Industria			
oped	untilled	tilled	unsewered	sewered					
	VOLATILE ORGA	NIC COMPOL	JNDScontinued						
14	11	15	29	21	12	11			
0	0	0	3	1	5	2			
					.04*				
					.65*				
А	Α	Α	А	Α	В	AB			
μg/L)									
14	11	15	29	21	12	11			
2	2		1	1		2			
А	Α	Α	А	А	Α	Α			
14	11	15	29	21	12	11			
			6			1			
			.12*	_					
А	Α	А	А	Α	Α	Α			
14	11	15	29	21	12	11			
			1	0	1	0			
А	Α	А	Α	Α	Α	Α			
					**				
14	11	15	29	21	12	10			
	0	1	0	0	2	0			
А	А	Α	А	А	Α	А			
		PESTICIDE	S						
14	11	15	11	11	12	11			
0	0	5	0	0	1	1			
A	AB	В	AB	AB	AB	AB			
	14 0 -A 14 2 A 14 0 A	oped Untilled VOLATILE ORGA 14	VOLATILE ORGANIC COMPOL 14 11 15 0 0 0 0 A A A A μg/L) 14 11 15 2 2 1 A A A A 14 11 15 1 0 A A A A 14 11 15 0 0 0 0 A A A A 14 11 15 0 0 0 0 A A A A PESTICIDE	oped untilled tilled unsewered VOLATILE ORGANIC COMPOUNDScontinued 14 11 15 29 0 0 0 3 A A A A A 14 11 15 29 1 1 1 0 6 6 12* .08* A A A A A 14 11 15 29 0 0 0 1 A A A A PESTICIDES	Oped Untilled Unsewered Sewered	Oped untilled tilled unsewered sewered VOLATILE ORGANIC COMPOUNDScontinued 14 11 15 29 21 12 0 0 0 3 1 5 0.04* 0.04* 0.65* A A A A A B μg/L) 14 11 15 29 21 12 2 2 1 1 1 0 14 11 15 29 21 12 0 0 0			

Table 5.--<u>Statistical summary by land use of water-quality variables detected in 1 or more of 85 to 116 observation wells in 4 stratified-drift aquifers in Connecticut--Continued</u>

Water-quality			L	and-use categ	ory		
variable, units, and	Undevel-	Agricul	tural	Reside	ntial	Commercial	Industria
statistic	oped	untilled	tilled	unsewered	sewered		
		PEST	ICIDESco	ntinued			
Cyanazine (µg/L)							
Number of wells	14	11	15	11	11	12	11
Number of detections	0	1	0	0	0	0	0
Median							
Interquartile range							
Population	Α	Α	А	Α	Α	Α	Α
Prometone $(\mu g/L)$							
Number of wells	14	11	15	11	11	12	11
Number of detections	0	0	0	0	0	0	2
Median							
Interquartile range							
Population	Α	Α	А	А	Α	Α	А
Propazine (µg/L)							
Number of wells	14	11	15	11	11	12	11
Number of detections	0	0	0	0	0	1	0
Median							
Interquartile range							
Population	А	Α	А	А	А	А	Α
Simazine (µg/L)							
Number of wells	14	11	15	11	11	12	11
Number of detections	1	0	0	0	0	1	0
Median							
Interquartile range							
Population	Α	Α	Α	А	А	А	А
Chlordane (µg/L)							
Number of wells	13	10	15	31	21	12	11
Number of detections	0	0	0	2	1	0	0
Median							
Interquartile range							
Population	А	А	А	А	А	А	А
Diazinon (µg/L)							
Number of wells	13	10	15	31	21	12	11
Number of detections	0	0	0	2	0	0	0
Median							
Interquartile range							
Population	А	А	А	А	А	А	А
Dieldrin (µg/L)							
Number of wells	13	10	15	31	21	12	11
Number of detections	0	0	1	2	1	0	1
Median							
Interquartile range							
Population	A	A	А	А	А	А	А

Table 5.--Statistical summary by land use of water-quality variables detected in 1 or more of 85 to 116 observation wells in 4 stratified-drift aquifers in Connecticut--Continued

Water-quality variable,	Land-use category									
units, and	Undevel-	Agricult	tural	Residential		Commercial	Industria			
statistic	oped	untilled	tilled	unsewered	sewered					
		PEST	ICIDESco	ntinued						
DDD (µg/L)				,						
Number of wells	13	10	15	31	21	12	11			
Number of detections	0	0	0	1	0	0	0			
Median										
Interquartile range										
Population	Α	Α	А	Α	А	А	Α			
2 4 D (ug/L)										
2,4-D (µg/L)	1.2	10	12	25	16	11	10			
Number of wells	13 1	10 0	12 0	25 1	16 1	11	10			
Number of detections						0	2			
Median										
Interquartile range										
Population	А	А	Α	Α	А	А	А			
2,4-DP (µg/L)										
Number of wells	13	10	12	25	16	11	10			
Number of detections	0	0	0	0	1	0	0			
Median										
Interquartile range										
Population	А	А	Α	А	А	Α	Α			
2,4,5-T (μg/L)										
Number of wells	13	10	14	31	20	12	11			
Number of detections	0	0	0	1	1	1	0			
Median										
Interquartile range										
Population	А	А	А	А	А	Α	А			
Silvex (µg/L)										
Number of wells	13	10	14	31	20	11	11			
Number of detections	0	0	0	1	1	0	0			
Median		O	0	1	1					
Interquartile range										
Population	Α	A	A	A	A	А	A			
Phorate $(\mu g/L)$										
Number of wells	13	5	11	18	19	11	11			
Number of detections	0	0	0	0	1	0	0			
Median										
Interquartile range										
Population	А	A	Α	А	Α	Α	А			

Appendix A.--Summary of site information, sample date and time, and water level for wells sampled during this study

[Chemical analyses of water samples from wells listed below are published in a series of U.S. Geological Survey reports cotaining water-resources data for Connecticut: U.S. Geological Survey Water-Data Reports CT-85-1 to CT-89-1 (published annually), are maintained in the U.S. Geological Survey's Water Data Storage and Retrieval System (WATSTORE), or are on file in the U.S. Geological Survey's Hartford, Connecticut office. Depth of well is the distance below land surface to the bottom of the well screen. Land use: AT, tilled agricultural; AU, untilled agricultural; C, commercial; I, industrial; RS, sewered residential; RU, unsewered residential; and U, undeveloped. Depth to water is the distance below land surface to the static water level measured just prior to sampling the well; --, no data available]

Local well number	Site identification number	Depth of well (feet)	Land use	Date	Time	Depth to water (feet)
		POOTATU	CK AQUIF	ER		
NТ98	412312073160501	15.6	RU	04-08-85 08-26-86 08-19-87	1145 1320 1412	2.60 2.78 3.00
NT99	412320073165201	26.5	I	04-10-85 08-05-85	1400 1245	17.40 8.10
NT100	410 4200 721 65601	11. 4		08-27-86 08-19-87 11-09-88	1512 1108 1108	17.54 18.50 16.18
NT100	412438073165601	11.4	AU	04-08-85 08-26-86 06-24-87	1425 1122 1155	6.30 5.13 6.90
				07-27-87 08-25-87 09-23-87	1410 1000 1005	7.60 7.80 5.90
				10-20-87 11-18-87 02-16-88	0854 1017 1045	6.70 6.60 5.40
				03-17-88 04-26-88 05-23-88	1000 1000 1040	4.95 6.40 6.79
NT101	412453073170201	16.4	I	06-23-88 04-09-85 08-26-86	1130 1115 1324	7.58 8.10 7.97
				08-18-87 11-14-88 12-01-88	1407 1304 1420	9.40 7.87 7.55
NT102	412526073161802	16.3	RU	04-11-85 05-08-85	1145 1112	7.50 7.60
NT103	412308073164301	21.0	I	04-10-85 08-05-85 08-28-86 08-19-87	1114 1040 1145 1215	5.80 6.20 6.42 7.50
				11-09-88	1243	5.72

Appendix A.--Summary of site information, sample date and time, and water level for wells sampled during this study--Continued

Local well number	Site identification number	Depth of well (feet)	Land use	Date	Time	Depth to water (feet)
			UG AQUIF			
SB34	412939073131901	16.8	AT	04-05-85 09-02-86	1440 1440	7.00 8.20
SB35	412918073132501	26.1	С	08-03-87 04-11-85 08-05-85 09-02-86 08-18-87 08-10-88	1330 1410 1435 1325 1027 1100	7.80 9.60 9.90 10.50 10.80
SB36	412819073133101	11.6	RU	06-28-89 04-12-85 08-06-85 09-02-86 08-18-87	1252 0955 1110 1130 1202	8.75 3.10 3.50 3.62 3.50
SB37	412947073131801	21.5	AT	08-22-86 06-24-87 08-03-87 08-25-87 09-23-87 10-20-87 11-18-87 02-16-88 03-17-88 04-26-88 05-23-88	1335 1335 1205 1114 1120 1105 1128 1235 1128 1105 1153	9.50 8.30 9.30 9.70 8.20 8.50 7.60 6.92 7.89 7.97
SB38	412916073130001	25.3	I	06-28-88 08-25-87 06-28-88 11-15-88	0843 1312 0948 1115	9.12 12.60 12.35 12.15
WY 44	413128073121901	16.4	С	12-01-88 04-12-85 08-20-86 08-13-87 11-14-88	1512 1330 1155 1325 1433	11.29 11.90 12.30 12.70 12.63
WY 45	413317073120001	15.3	AT	04-04-85 09-03-86 08-07-87	1250 1326 1326	6.50 7.60
WY 46	413413073133101	11.6	U	04-01-85 04-12-85 08-22-86 08-11-87 08-22-88	1400 1150 0940 1530 1205	5.30 6.69 6.70 7.00
WY 47	413322073124501	15.6	RU	08-22-88 04-05-85 08-21-86 08-12-87	1240 1210 1316	5.10 5.79 5.70
WY 48	413311073125601	15.9	AT	08-12-87 04-02-85 08-21-86 08-04-87	1310 1310 1310 1450	7.60 5.23 8.10
WY 49	413223073133701	15.1	AU	04-05-85 08-20-86 08-03-87	1015 1335 1440	7.00 6.90 6.90

Appendix A.--Summary of site information, sample date and time, and water level for wells sampled during this study--Continued

Site identification number	Depth of well (feet)	Land use	Date	Time	Depth to water (feet)
PC	MPERAUG AQ	UIFERc	ontinued		
	·			1030	5.30
415520075114201	10.4	\wedge 1			5.70
413309073122901	45.6	RU			33.80
			05-08-85		33.90
			09-04-86	1215	34.10
			08-12-87	1438	32.30
		U			7.70
413130073122201	20.5	С			7.10
					8.00
.1.000.00701.00501	20 5				7.74
413326073123501	30.5	RU			12.10
					12.58
41222227212221	11 5	DII			12.30
					4.40 7.30
4133000/3125901	16.0	AI			7.20
					4.63
412220072125201	16 2	ΛΤ			4.03
4132280/3125201	10.2	AI			4.20
					4.40
413301073122101	20 8	RII			9.60
4133010/3122101	20.0	110			11.36
					12.72
413258073123001	44.7	RU			29.00
113230073123001		110			26.30
					28.69
			11-21-88	1032	29.67
			06-28-89	0952	25.89
413232073124001	25.8	C	08-19-86	1200	10.90
			08-26-87	1052	9.00
			07-26-88	0943	9.70
			11-15-88	1417	10.91
			11-21-88	1147	9.58
413053073141801	11.2	U			4.51
					4.80
			11-15-88	1230	4.11
	FARMING	TON AQUI	[FER		
414126072512101	20 0	DC	08 20 86	1225	8.14
414120072312101	20.9	1/2			8.50
					8.26
					7.66
414055072522301	20.9	RII			14.60
					6.80
					11.70
					6.80
1141120/200/01	20.0	,,,,			6.40
	identification number P0 413326073114201 413309073122901 413347073112601 413130073122201 413326073123501 413322073122301 413328073125901 413228073125201 413258073122101 413258073123001	dentification number number (feet)	Identification Of well Land Use	Identification number Nomperature Ceet Nomperature Nomperatu	Time PomPERAUG AQUIFERcontinued PomPERAUG AQUIFER PomPERAUG AQUIFER PomPERAUG AQUIFER PomPERAUG AQUIFER PomPERAUG AU PomPERAUG AU

Appendix A.--Summary of site information, sample date and time, and water level for wells sampled during this study--Continued

Local well number	Site identification number	Depth of well (feet)	Land use	Date	Time	Depth to water (feet)
Tidilibet					Time	(1660)
	FAR	MINGTON AQ	UIFERc	ontinued		
F323	414323072502501	21.1	AT	07-31-86 07-24-87	1330 1114	5.60 6.30
F324	414312072504301	21.1	AT	07-19-88 07-25-86 06-24-87 07-24-87 08-27-87 09-23-87 10-20-87 11-18-87 02-16-88 03-17-88 04-26-88 05-23-88 06-24-88	1320 0930 0905 1016 1320 1255 1307 1418 1320 1247 1323 0930	6.50 7.90 7.70 8.30 8.20 7.50 7.50 7.20 7.33 7.80 6.76 7.92
F325	414454072504801	25.5	U	07-19-88 07-30-86 08-14-87 07-29-88	1050 1215 1250 1220	8.00 17.30 17.20 17.80
F326	414521072510801	15.5	U	07-23-86 07-31-86 08-17-87 07-29-88	1020 1045	3.00 3.10 2.80
F327	414207072514801	16.4	I	07-29-88 07-24-86 08-20-87 10-25-88	1545 1036 1428	7.73 8.40 6.78
F328	414439072515401	25.7	RS	07-22-86 08-10-87 07-01-88	1530 1330	18.70 19.50 19.79
F329	414324072495601	38.5	RS	08-01-86 07-28-87 08-15-88 06-23-89 08-25-89	1625 0950 1645 1345 0955 1000	32.00 32.10 32.20 29.74 31.33
F330	414338072512601	16.1	U	08-23-69 07-24-86 07-28-87 07-29-88	1335 1340 0925	6.80 6.90 4.80
F331	414436072514401	25.9	RS	08-01-86 07-14-87 08-14-87 07-01-88	1145 1030 1030 1515	19.20 19.30 19.30 19.40 20.28
F332	414157072520801	46.2	I	10-25-88 03-30-88 11-23-88	1300 1000 1500	40.00 37.61
F333	414208072510601	55.6	I	08-01-86 08-20-87 11-08-88	1335 1155 1353	44.00 44.20 43.55

Appendix A.--Summary of site information, sample date and time, and water level for wells sampled during this study--Continued

Local well number	Site identification number	Depth of well (feet)	Land use	Date	Time	Depth to water (feet)
		MINGTON AQ				
A304	414905072491301	16.2	AT	08-07-86	1530	8.90
,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	11 130007 2 13 1001	1012	, , ,	08-06-87	1315	8.80
				07-25-88	1134	8.90
				11-18-88	1020	8.90
A305	414601072493901	20.6	AT	07-29-86	0945	8.90
				08-05-87	1145	8.70
1306	414637072493501	21.0	AU	07-27-88 07-23-86	1000 1340	8.83 7.90
1300	41403/0/2493301	21.0	٨٥	08-05-87	1305	7.90
				08-18-88	1110	8.20
				06-23-89	1235	7.13
				08-25-89	1130	7.64
1307	414714072494701	16.0	AU	07-29-86	1205	10.30
				08-06-87	1525	9.80
4308	414541072495101	25.3	U	07-28-86	1032	21.90
				08-17-87	1300	21.50
				07-27-88 08-19-88	1135 1345	22.90 22.20
4309	414828072492701	35.8	С	09-05-86	1400	24.10
1209	414020072492701	33.0	C	09-01-87	1355	24.40
				08-18-88	1430	24.60
1310	414626072495601	20.6	U	07-23-86	1115	15.00
				08-31-87	1427	15.30
				07-27-88	1300	15.20
				08-18-88	1250	15.40
4311	414839072505101	16.1	C	09-05-86	1030	8.50
				08-27-87	1215	8.40
1312	414747072500801	25.6	I	08-19-88 08-04-86	1200 1515	8.70 17.50
1312	414/4/0/2500601	25.0	1	09-01-87	1139	17.00
				07-25-88	1410	17.30
51445	415246072512001	16.0	AT	08-14-86	1010	3.82
				08-06-87	1124	3.90
				08-26-88	1055	3.80
51446	414922072490401	21.1	AT	08-07-86	1400	8.00
				08-05-87	1440	8.00
27.4.4.7	415051070402001	15.0	DII	07-25-88	0942	8.00
51447	415251072483001	15.8	RU	08-12-86 07-09-87	1525 0945	8.30 6.60
				08-17-88	1100	8.70
				10-31-88	1147	8.82
				06-26-89	0935	5.04
SI 448	415351072500501	25.8	U	08-13-86	1500	17.40
				07-09-87	1240	14.20
				10-31-88	1019	21.60
	41 51 1 00 70 51 0001	25 0	U	08-13-86	1107	18.80
SI 449	415118072512301	25.9	U			
SI 449	4151180/2512301	25.9	U	07-09-87 08-01-88	1405 1053	19.50 18.90

Appendix A.--Summary of site information, sample date and time, and water level for wells sampled during this study--Continued

Local well number	Site identification number	Depth of well (feet)	Land use	Date	Time	Depth to water (feet)
	FAR	MINGTON AQ	UIFERc	continued		
SI450	415234072480301	21.1	С	08-12-86 08-28-87 10-31-88	0945 1112 1310	14.80 13.70 15.30
SI451	415300072480301	45.7	RU	08-12-86 07-08-87 08-17-88	1400 1515 0930	34.30 32.30 34.50
				06-26-89 08-25-89 09-06-89	1100 1425 1345	32.75 32.53
SI 452	415340072472501	31.2	RU	08-11-86 07-07-87 08-16-88	1125 1330 1400	18.10 18.20 22.05
\$1453	415441072472601	20.3	RU	06-20-89 08-11-86 07-07-87 08-16-88 11-07-88 06-22-89	0940 1430 1107 1130 1524 1310	16.12 15.90 14.50 16.40 16.70
SI461 SI462 SI463 SI464 SI465 SI466 SI466	415037072485501 415045072512301 415011072513401 415035072512201 415308072480101 415328072490901 415316072483601	6.4 16.0 10.7 21.1 25.5 11.1 15.6	RU RU RU RU RU RU RS	08-25-89 07-18-88 07-14-88 07-14-88 07-14-88 07-07-88 07-07-88 07-13-88	1305 1300 1035 1400 1220 1020 1130 1420	14.88 5.70 6.00 7.90 12.70 22.20 2.30 7.90
SI468 SI469 SI470 SI471	415425072505801 415441072505201 415450072510401 415500072481801	14.4 15.3 16.3 19.5	RU RU RU RU	07-12-88 07-12-88 07-13-88 07-06-88	1405 1200 1220 0845	6.40 6.20 6.90 11.90
GR334 GR335 GR336	415642072474601 415706072475601 415747072475901	15.9 16.0 21.0	AU RU RU	06-30-88 07-01-88 07-01-88 12-02-88	1120 1217 1005 0932	5.60 6.60 14.60 13.82
GR337	415722072475701	26.2	RU	07-11-88 11-07-88	1045 1400	22.50
GR338	415643072501901	11.2	AU	06-28-88 07-18-88	1055 1025	4.70 4.80
GR339 GR340 GR341	415534072473201 415516072473301 415858072481001	26.0 30.9 34.9	RU RU RU	06-30-88 07-11-88 06-28-88	1330 1330 1340	13.60 24.70 23.20

Appendix A.--Summary of site information, sample date and time, and water level for wells sampled during this study--Continued

Local well	Site identification number	Depth of well (feet)	Land	Date	Time	Depth to water
number	number	(reet)	use	Date	Time	(feet)
		HOCKAN	UM AQUIF	ER		
M157	414853072303601	15.2	С	07-14-87 08-23-88	1135 1430	3.80 4.10
M158	414547072304801	11.5	RS	07-16-87 06-29-88	1400 1110	7.30 7.69
M159	414629072290601	15.2	U	06-25-87 07-28-88	1230 1152	6.10 7.60
1160	414624072283301	31.0	U	0 6 -25-87 07-15-87	1555 1410	14.80 15.00
M161	414857072302701	19.9	С	07-28-88 07-14-87 08-23-88	1017 1535 1211	20.20 3.60 3.80
M162	414741072284501	16.0	U	06-26-87 06-29-88	1200 1252	3.20 2.81
M163	414740072301801	15.0	RS	08-23-88 03-18-88 03-29-88 12-02-88	0930 1035 1330 1315	6.30 13.41 12.90 12.68
1164	414814072301901	15.0	I	07-17-87 06-29-88 10-26-88	0935 1404 1248	2.10 2.85 3.36
M165	414637072314401	35.2	RS	06-29-87 08-11-88 10-27-88	1315 1037 1200	22.60 23.30 23.52 22.67
4166	414709072330601	15.3	I	06-27-89 07-17-87 06-29-88 10-26-88	1145 1017 1109	3.00 3.08 3.13
M167	414709072332601	15.8	I	07-15-87 08-26-88	1630 0830	4.20
1168	414631072332301	16.3	RS	07-20-88	1425	6.11
M169	414548072330501	16.2	RU	07-21-88 07-20-88	1200 1330	6.11
M170	414618072331101	21.6	RS	07-21-88 07-21-88 07-21-88 07-22-88	1230 1200 1500 1209	11.50 17.50 17.50 17.50
M171	414528072305101	36.0	RS	07-27-88 06-29-88 12-01-88	0815 1510 0847	8.60 18.40 21.94
M172	414509072304301	26.0	RS	06-24-88 07-20-88	1200 1300	8.10
M173 M175	414901072305001 414454072311401	29.9 9.8	RS AU	06-23-88 08-15-88	1220 1415 1303	18.20 2.70 1.19
SW134 SW135	414914072310101 414913072304701	20.6 35.8	RS RS	08-18-89 06-23-88 06-29-88	1432 1145	7.50 12.70

Appendix A.--Summary of site information, sample date and time, and water level for wells sampled during this study--Continued

Local	Site	Depth	1			Depth
well number	identification number	of well (feet)	Land use	Date	Time	to water (feet)
	но	CKANUM AQU	IFERco	ntinued		
V89	415134072292601	20.6	RS	0 6- 23-87	1000	4.80
				07-22-87	1443	5.23
				08-24-87	1512	5.50
				09-22-87	1425	5.00
				10-19-87	1505	5.20
				11-19-87	1014	5.20
				02-17-88	1256	3.70
				03-18-88	1245	3.91
				04-26-88	1718	4.13
				05-24-88	1103	4.36
				06-24-88	1351	5.02
				12-02-88	1133	4.33
V9 0	415123072292601	14.0	RS	06-30-87	1015	11.90
				06-30-88	1058	12.18
V91	415102072291801	9.5	RS	06-30-87	1240	5.40
				06-30-88	1152	5.59
				10-27-88	1414	5.74
V92	415155072284601	25.6	C	06-23-87	1445	12.50
				06-30-88	1250	12.76
				11-07-88	1008	12.57
V93	415201072284601	15.7	RS	06-23-87	1230	6.00
		0.2		06-30-88	1340	6.52
V94	415013072263901	11.1	U	06-26-87	0930	3.60
				07-28-88	1330	4.20
V9 5	414936072300201	20.7	C	06-30-87	1445	8.70
				06-30-88	0958	9.20
				10-27-88	1318	9.17
10.6	41 400607000001	05.0	•	11-17-88	1350	8.50
V96	414926072300801	25.8	C	06-22-87	1522	13.00
				07-22-87	1220	13.44
				08-24-87	1620	13.70
				09-22-87	1525	12.90
				10-20-87	1430	13.40
				11-19-87	1112	12.90
				02-17-88	1412	12.00
				03-18-88	1140	12.47
				04-26-88	1813	12.80
				05-24-88	1202	12.60
				06-23-88	1402	13.33
				07-28-88	1509	12.70
TI 01	415616070202021	07.4		10-27-88	1218	10.44
EL81	415616072303801	27.4	U	06-22-87	1150	13.10
				07-31-87	1555	26.20
				08-24-87	1245	27.40
				02-17-88	1000	27.30
				03-18-88	1530	22.71
				04-26-88	1517	23.43
				05-24-88	0845	23.67
				06-24-88	1110	25.20

Appendix A.--Summary of site information, sample date and time, and water level for wells sampled during this study--Continued

Local well number	Site identification number	Depth of well (feet)	Land use	Date	Time	Depth to water (feet)
	НО	CKANUM AQU	IFERco	ntinued		
EL82	415458072291901	24.5	AU	07-31-87	1445	6.50
EL83	415450072272801	16.0	AU	07-01-88 07-21-87 07-05-88	0946 0930 1130	6.47 4.40 5.50
EL84	415453072273801	16.1	AT	11-18-88 07-21-87	1315 1255	3.35 4.01
EL85	415511072284201	21.2	AT	07-05-88 06-22-87	1300 1400	3.80 9.70
				07-23-87 08-24-87	1055 1402	10.52 11.10
				09-22-87 10-19-87	1310 1310	11.00
				11-19-87 02-17-88 03-18-88	0914 1115 1405	11.00 8.80 8.84
				04-26-88 05-24-88	1613 1002	9.95 10.13
				06-24-88 11-07-88	1226 1116	10.70
EL86	415441072274101	16.4	AU	07-21-87 08-15-88	1500 1045	6.20 6.20
				06-29-89	1043	5.70



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