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Completion Report: Pesticide and Nitrate Monitoring Results for Craighead, Mississippi, and Poinsett Counties, Arkansas: Phase II

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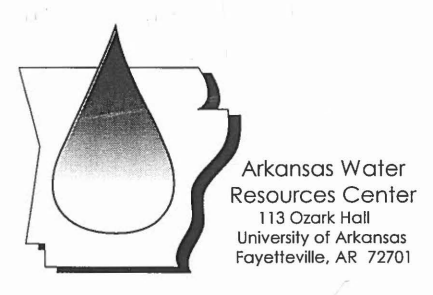


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COMPLETION REPORT: PESTICIDE AND NITRATE MONITORING RESULTS FOR CRAIGHEAD, MISSISSIPPI, AND POINSETT COUNTIES, ARKANSAS PHASE II

1993

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APPENDIX A

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PESTICIDE AND NITRATE MONITORING RESULTS FOR CRAIGHEAD, MISSISSIPPI, AND POINSETT COUNTIES, ARKANSAS

Kenneth F. Steele, William R. Clayton, Terry W. Nichols, Paul F. Vendrell

June 1996

ABSTRACT

Because of the concern for potential contamination of ground water by agricultural chemicals, 38 wells drilled in the Mississippi River Valley alluvial aquifer in Mississippi County and the eastern parts of Craighead and Poinsett Counties, Arkansas were analyzed for pesticides and nitrate. The pesticide, fluometuron, was detected in one sample at a concentration of 0.5 mg/L. Bentazon was detected in three samples at concentrations of 2.5, 0.3, and 0.3 mg/L. The occurrences of the pesticides appear to represent isolated incidents rather than a widespread aquifer contamination. All detections were below health and safety standards. Nitrate is present in several wells at concentrations above 0.15 mg/L, one of which exceeded the EPA established maximum contaminant level for drinking water of 10 milligrams per liter as nitrogen. Except for two wells nitrate and iron are not present together at concentrations above 0.15 mg/L. This is probably due to microbially mediated reactions. Nitrate concentrations above 0.15 mg/L is only present in wells that are less than 60 feet deep and near permeable soils. Iron is present in wells that are not near permeable soils or wells that are greater than 40 feet deep, and may exceed 1 mg/L in some cases.

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PESTICIDE AND NITRATE MONITORING RESULTS FOR CRAIGHEAD, MISSISSIPPI, AND POINSETT COUNTIES, ARKANSAS.

Kenneth F. Steele, William R. Clayton, Terry W. Nichols, Paul F. Vendrell

INTRODUCTION

Mississippi, Craighead and Poinsett counties in northeastern Arkansas are important agricultural areas where it is economically beneficial to use pesticides and fertilizers. In order to determine if agricultural practices are contaminating the ground water, water was collected from shallow wells and analyzed. This investigation is part of a monitoring program designed to determine pesticide contamination of shallow ground water in the most vulnerable areas of Arkansas. The study area includes all of Mississippi County and parts of Craighead and Poinsett counties east of Crowley's Ridge. The study area is bounded on the east by the Mississippi River, on the south by Crittenden and Cross counties, and on the north by Greene County and the state of Missouri (Figure 1).

PURPOSE AND SCOPE

This report describes and discusses the occurrence of pesticides, nitrate, iron and lead in water from the alluvial aquifer in the study area in northeastern Arkansas. The description and occurrence includes the discussion of parameter relationships to each other and to the soils near the wells that were sampled. Information on other major and minor water quality parameters is provided by Clayton (1995).

BACKGROUND

According to the U.S. Environmental Protection Agency (EPA) nationwide evidence has shown that agricultural pesticides can contaminate ground water (U.S. EPA, 1990). Because of this, the EPA has encouraged each state to develop a management plan for dealing with possible pesticide contamination of ground water.

The responsibility for developing a pesticide management plan for Arkansas has been given to the Arkansas State Plant Board (ASPB). The Arkansas Soil and Water Conservation Commission (ASWCC) coordinated development of a vulnerability model (Fugitt, 1992) to determine potential problem areas for pesticide contamination of ground water with ASPB. ASWCC also has been involved in some monitoring (especially of deeper wells). The plan developed by the Arkansas State Plant Board in collaboration with the Arkansas Water Resource Center (AWRC) includes an education program designed to prevent contamination, and a monitoring program to determine if contamination has occurred. The monitoring program uses the vulnerability model to determine potential problem areas (Arkansas State Plant Board, 1992). The areas chosen for monitoring were those considered to be most vulnerable to contamination by pesticides. The vulnerability assessment was based, in part, on the pesticide version of the

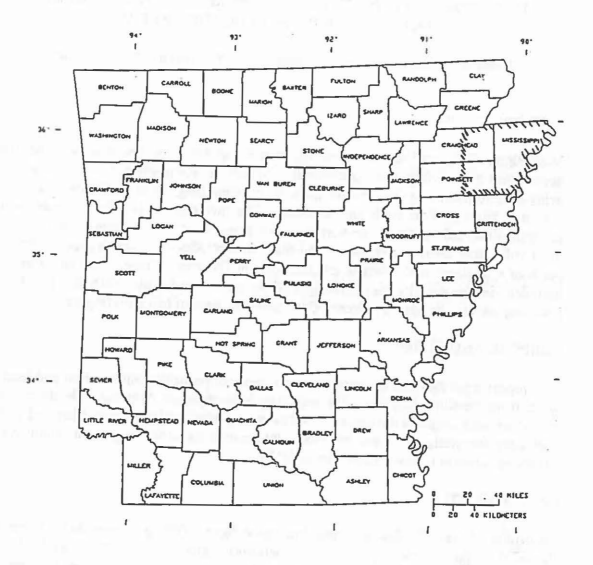


Figure 1. Location of the study area.

DRASTIC method for determining areas sensitive to ground water contamination (Aller et al., 1987).

DRASTIC determines sensitivity based on seven factors:

- Depth to the water table,
- net Recharge,
- · Aquifer media,
- Soil media,
- Topography,
- Impact of the vadose zone, and
- hydraulic Conductivity

In the study area, depth to the water table, aquifer media, topography, and hydraulic conductivity are nearly uniform (Ryling, 1960). The three remaining factors, net recharge, soil media, and impact of the vadose zone are all indicators of how easily water can move from the surface to the aquifer.

The vulnerability in a county also considers the amount of pesticides used in the county. Pesticide use was based on information about crop production by county, and an estimate of the type and amount of pesticides used for specific crops. The most vulnerable areas in the state were determined by considering pesticide use and DRASTIC sensitivity (Nichols and Wilkes, 1992).

Sample collection and analyses were performed by the Arkansas Water Resource Center (AWRC) Water Quality Laboratory. Monitoring in Ashley County, previously determined to be the most vulnerable area, has been completed (Nichols et al., 1993). Metolachlor was detected at 0.71 µg/L in only one of 23 wells. The well was later re-sampled for verification of the detection; the re-sample did not contain detectable levels of pesticide. This present study, involves the second most vulnerable area (Mississippi County and the eastern parts of Craighead and Poinsett counties).

LAND USE AND CLIMATE

The study area is predominantly farm land, but forest remains in some areas, especially in the floodway adjacent to the St. Francis River, and areas near the Mississippi River that are not protected from flooding. Soybeans, cotton, and rice are the main crops grown. Some areas are also used for wheat, corn, grain sorghum, and cattle grazing (Arkansas Agricultural Experiment Station, 1995). Crop statistics for each of the three counties in the study area are given in Table 1 (Arkansas Agricultural Experiment Station, 1995). Note that Table 1 gives totals for all of Craighead and Poinsett counties, i.e., it includes the parts of these counties not in the study area.

Table 1. Agricultural statistics for Craighead, Mississippi and Poinsett Counties.

Values given are in acres.

	Craighead County	Mississippi County	Poinsett County
Total land area	454,913	574,922	484,987
Land in farms	350,402	484,751	404,585
Soybean	104,000	223,000	143,000
Cotton	84,000	200,000	55,000
Rice	78,000	20,000	111,000
Wheat	14,500	58,000	12,000
Sorghum	6,500	16,500	7,000
Com	5,600	3,700	1,700

Source: Arkansas Agricultural Experiment Station, 1995.

The climate of the study area is characterized by typically hot, humid summers and cool winters. Cold winter storms occur but are of short duration. Average precipitation is nearly 50 inches per year. January is the wettest month, but the heaviest rainfall events occur during the spring (Ferguson and Gray, 1971; Ferguson, 1979; and Gray and Ferguson, 1977).

HYDROGEOLOGY AND GEOLOGY

The uppermost aquifer in the area is the Mississippi River Valley alluvial aquifer. It is composed of Mississippi River alluvial deposits. These deposits grade upward from coarse sand and gravel to silt and clay (Plafcan and Fugitt, 1985).

The lower half to two thirds of the aquifer is composed of braided-stream deposits laid down during glaciation. The upper part of the aquifer was deposited by the Mississippi River after it became a meandering stream. These deposits consist of point-bar sands and some gravel (Saucier, 1964; Saucier and Snead, 1989).

In most of the study area the depth to ground water is less than 20 feet. The base of the aquifer is 100-200 feet below the land surface (Ryling, 1960). Wells in the alluvial aquifer yield 1000-3000 gallons per minute (gpm). The direction of ground-water flow is generally to the southwest. The Mississippi River acts as a source of recharge, but this may change seasonally with changes in river level, resulting in changes in the direction of flow near the river (Plafcan and Fugitt, 1985). Ryling (1960) reports ground-water velocities of 0.2 to 0.3 feet per day (fpd) in northwestern Mississippi County, and velocities of 0.1 to 0.8 fpd near the Mississippi River. Velocities near the river are greatly affected by the level of the river. Hydraulic conductivity values for the aquifer range from 120-390 fpd (Mahon and Ludwig, 1990).

Overlying the alluvial aquifer are finer-grained sands, silts and clays that were deposited by the Mississippi River and smaller streams as backswamp, meander-belt and braided-stream deposits. These units form the overlying confining layer for the alluvial aquifer (Ackerman, 1989).

The Memphis aquifer underlies the alluvial aquifer. This aquifer is mainly sand with some interbedded clay and comprises the lower part of the Claiborne Group. The boundary between the two aquifers is an erosional surface, an angular unconformity. In the northeastern part of the study area the two aquifers are connected. In the southwestern part they are separated by the Jackson-Claiborne clay, which includes the upper part of the Claiborne Group and the Jackson Group (Broom and Lyford, 1982).

SOILS

Soils in Mississippi County are related to the past and present deposits of the Mississippi River in which they have developed. Near the Mississippi River, loamy to sandy soils are

developed on the natural levees of the river. Clayey soils occur throughout much of the central part of the county and are developed in backswamp deposits. Sandy or loamy soils occur in some areas of the central part of Mississippi County. Northwest of the Left Hand Chute of the Little River the soils become more loamy with some sandy soils. Locally, sandy soils exist as a result of sand blows that occurred during the 1811-1812 New Madrid earthquakes (Ferguson and Gray, 1971).

Soils in the part of Craighead County included in the study area are generally loamy to fine sandy loam; however, in the St. Francis River basin, which crosses the county from north to south, the soils are clayey. These soils were developed in clays deposited by the St. Francis River (Ferguson, 1979). In the selected portion of Poinsett County soils are generally clayey, although there are some sandy soils. These grade to loamy soils in the western part of the study area (Gray and Ferguson, 1977).

PARAMETERS AND METHODS

A total of 38 wells were sampled for this study. Samples for pesticide, nitrate, iron, lead, and other analyses were collected on seven trips to the study area from November 1993 to October 1994. See Clayton (1995) for details of collection and analysis of other cations and anions, and field parameters. Sample locations are shown in Figure 2. Locations and depths of wells are given in Table 2. The appropriate U.S. Geological Survey 7 1/2-minute quadrangles for each well, and well use are also listed in this table.

For all of these wells pH, temperature and specific conductance were measured at the time of sampling. Well depth was obtained from the owner. Wells that contained detectable levels of pesticides were resampled at a later date, except for one well, M4, that was no longer in use. Before water was collected, the wells were purged until temperature, conductance and pH became stable. This was done to ensure that the collected water represented ambient aquifer conditions.

The project involved monitoring for the following 13 herbicides.

metolachlor; norflurazon; fluometuron; alachlor; atrazine; cyanazine; bentazon; 2,4-D; metribuzin; acifluorfen; molinate; diuron;

These pesticides were chosen because of their extensive use in the area, their high leaching potential and their long half-life in soil. Solubility, half-life, adsorption coefficient (K_{oc}), and leaching potential are given in Table 3. Data are taken from Nichols and Wilkes (1992) which is based on data from the Cooperative Extension Service.

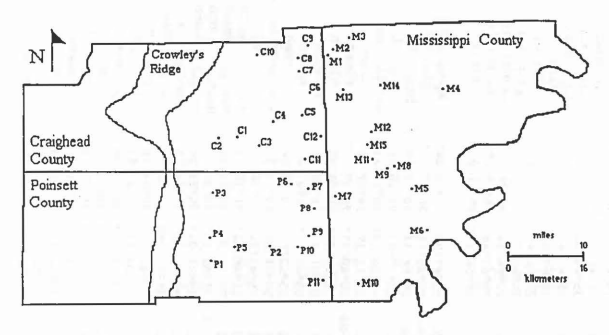


Figure 2. Locations of wells sampled.

Table 2. Location, depth and use of wells.

Well	Depth in feet	Latitude	Longitude	USGS quadrangle	Well use
61	25	250461225	000001071	I also Gime	D
C1 C2	25	35°46'22"	90°29'27"	Lake City	Domestic
	30	35°46'25"	90°33'09"	Needham	Domestic
C3	20	35°45'26"	90°27'42"	Lake City	Agriculture
C4	20	35°47'27"	90°25'30"	Lake City	Agriculture
C5	25	35°48'12"	90°21'01"	Caraway	Agriculture
C6	20	35°51'46°	90°18'53°	Caraway	Agriculture
C7	50	35°53'31'	90°20'30"	Leachville	Agriculture
C8	50	35°56'31"	90°20'32"	Leachville	Agriculture
C9	35	35°57'38"	90°19'08*	Leachville	Agriculture
C10	18	35°56'36"	90°27'02"	Dixie	Domestic and
					agriculture
C11	30	35°43 .55 .	90°20'24"	Rivervale	Agriculture
C12	50-60	35046.19.	90°17'58"	Caraway	Agriculture
M1	100+	35°56'34"	90017'00"	Leachville	Agriculture
M2	100+	35°57 '23 "	90°16'02"	Leachville	Agriculture
M3	100+	35°58'15"	90°13'33°	Manilla North	Agriculture
M4	50	35°52'19"	89°59'24"	Luxora	Agriculture
M5	30-50	35°40'33"	90°05'00"	Kieser	Agriculture
M6	125	35°34'29"	90°02'32"	Wilson	Pond
M7	25	35°38'43"	90°16'07"	Rivervale	Domestic
м8	50	35°43'12"	90°07'30"	Etowah	Agriculture
M9	45	35°42'54"	90°08'38"	Etowah	Agriculture
M10	55	35°28'09"	90°13'02"	Frenchmans Bayou	_
M11	65	35043'36"	90010'30"	Etowah	Agriculture
M12	27	35046'42"	90°10'39-	Manilla South	Domestic
M13	30	35°51'52"	90014.52	Manilla South	Agriculture
M14	20-40	35°52'19"	90°08'52"	Manilla South	Agriculture
M15	30	35045'24"	90°11'16"	Manilla South	Domestic
P1	100+	35°31'29"	90°34'14"	McCormick	Agriculture
P2	100	35°33'16"	90°25'48"	Marked Tree	Agriculture
P3	14	35°40'33"	90034118	Trumann	Domestic
P4	40	35°34'31"	90°34'53"	McCormick	Agriculture
P5	14	35°36'15*	90°31'09"	McCormick	Agriculture
P6	30	35°41'10"	90°22'34°	Hatchie Coon	Agriculture
P7	60	35°40'21"	90°20'29	Rivervale	Agriculture
P8	15	35°38'04"	90°19'23	Rivervale	Agriculture
P9	20	35°34'10"	90°20'01"	Lepanto	Hand pump
P10	50	35°32'54"	90°21'39	Lepanto	Domestic
P11	65	35°29'10*	90°18.01.	Tyronza	Domestic
	0.0	10	20 70 01	1,101124	PAMERCY

Table 3. Pesticide characteristics and uses.

Pesticide	Solubility mg/L	Half Life days	Soil Sorption K ∞	Leaching Potential	Trade Name	Crops Used on
2,4-D	890	10	20	medium	Amine	corn, rice, wheat
Acifluorfen	250,000	14	113	medium	Blazer	soybean
Alachlor	240	15	170	medium	Lasso	corn, cotton, soybean
Atrazine	33	60	100	large	Aatrex	com
Bentazon	230,000	20	34	medium	Basagram	soybean, corn, rice
Cyanazine	170	14	190	medium	Bladex	corn, cotton, wheat
Diuron	42	90	480	medium	Bladex	corn, cotton, wheat
Fluometuron	110	85	100	medium	Cotoran	cotton
Linuron	75	60	400	medium	Lorox	corn, cotton, winter wheat
Metolachlor	530	90	200	medium	Dual	corn, cotton, soybean
Metribuzin	1220	40	60	large	Sencore/Lexone	soybean, corn, wheat
Molinate	970	21	190	medium	Odram	rice
Norflurazon	28	90	600	medium	Zorial	cotton

From Wauchope and Goss, 1988

Three different methods were used for pesticide analysis. EPA method 507 (U.S. Environmental Protection Agency, 1991a), and EPA method 515.2 (U.S. Environmental Protection Agency, 1992) use gas chromatography for analysis, but use different detectors and different extraction processes. The third method is National Pesticide Survey method 4 which uses high performance liquid chromatography (HPLC) (Table 4). Iron and lead were analyzed using inductively coupled plasma emission spectrometry (EPA, 1991b). Nitrate-N was analyzed by ion chromatography (Plafcan et al, 1989). See Clayton (1995) for details of inorganic methods and analytical data.

Laboratory analyses were performed at the AWRC Water Quality Laboratory. All analyses were conducted in accordance with EPA accepted methodology. Estimated detection limits for all analytes for which a method is approved are published with the method. Method detection limits are specific to the laboratory conducting the analyses and are computed from results obtained in that laboratory. Both of these limits are reported in Table 4. A quality control report for the pesticide data by Nichols et al. (1994) is given in Appendix A. See Clayton (1995) for quality control information for the inorganic analyses.

Table 4. EPA analytical methods, and detection limits.

Group	Units	Parameter	Method Detection Limit	Estimated Detection Limit
EPA 200.7	mg/L	Fe	0.001	0.003
		Pb	0.003	0.003
EPA 300.0	mg/L	NO ₃ -N	0.005	0.01
EPA 507	µg/L	molinate	0.085	0.15
		atrazine	0.032	0.13
		metribuzin	0.09	0.15
		alachlor	0.103	0.38
		metolachlor	0.141	0.75
		norflurazon	0.122	0.5
NPS4	μg/L	cyanazine	0.236	0.15
		fluometuron	0.056	0.13
		diuron	0.083	0.15
		linuron	0.085	0.085
EPA 515.2	μg/L	2,4-D	0.069	0.28
	X. CERC	bentazon	0.676	0.63
		acifluorfen	0.208	0.25

RESULTS AND DISCUSSION

Pesticide Data

With the exception of well P8, all 38 wells were analyzed for the 13 pesticides. The extracts of the water from well P8 contained an oily residue and was not analyzed to

prevent contamination of the equipment. Pesticides were detected in samples taken from four wells. Bentazon was detected in wells M4, M5 and P1 with concentrations of 2.5, 0.3 and 0.2 micrograms per liter (mg/L), respectively. Well C4 had a 0.5 mg/L fluometuron detection (Table 5). The locations of these wells are shown in Figure 3.

Table 5. Wells With Pesticides Detections

Well	Pesticide	Pesticide ug/L	NO ₃ -N mg/L	Depth feet	Soil Association
M4	bentazon	2.5	0.05	50	Sharkey-Steele complex
M5	bentazaon	0.3	0.02	30-50	Sharkey-Steele clay
P1	bentazon	0.2	0.02	100+	Sharkey clay
C4	fluometuron	0.5	4.62	20	Sharkey clay

The detections for wells M4 and P1 were confirmed by the Arkansas State Plant Board using a gas chromatograph with a mass spectrometer detector. There was no additional sample from well M5 available for confirmation. The fluometuron detection from well C4 was confirmed by using an alternate column on the HPLC. Three of these wells were later resampled for verification of the contamination. Well M4 could not be resampled, even though two separate attempts were made, because the pump was no longer working. Of these samples collected for verification none contained detectable levels of pesticides.

The bentazon detections from wells M5 and P1 are below the method detection limit of $0.676 \mu g/L$. They are reported as detections because they produced an identifiable peak on the chromatogram. The quantitative concentrations of these detections may not be reproducible or accurate but the presence of a trace amount of pesticide is indicated.

The EPA has tentatively set a maximum contaminant level goal (MCLG) of 200 µg/L (0.02 mg/L) for bentazon. The maximum contaminant level goal is a non-enforceable concentration for drinking water set to protect against adverse human health effects. The EPA has not set an enforceable maximum contaminant level (MCL) for bentazon. An MCL is enforceable only for public water supply, not for domestic wells. The EPA has not set an MCLG or an MCL for fluometuron. None of the detections for bentazon are above the MCLG. The EPA has set health advisories for bentazon and fluometuron. Health advisories are concerned with only non-carcinogenic health effects. These advisories are levels of a contaminant in drinking water that are not expected to cause any non-carcinogenic health problems (EPA, 1994). None of these levels have been exceeded. Wells C4, M4 and M5 are reported to have depths of 20, 50, and 30-50 ft, respectively. These well depths are in distinct contrast to well P1, which is reported as being greater than 100 ft.

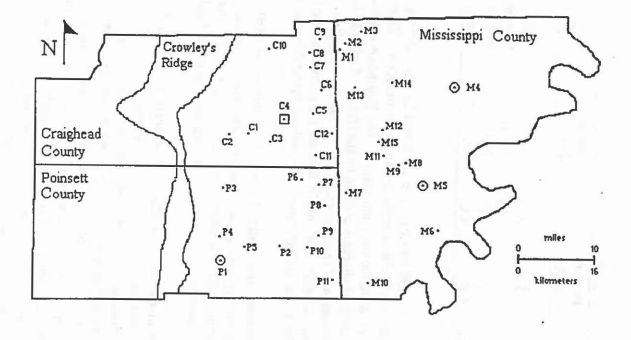


Figure 3. Locations of wells that contain pesticides. Bentazaon detections are circled and the floumetron detection is in a square.

Pesticide Contamination

A pesticide must travel through the soil and through the unsaturated or vadose zone in order to reach the water table. Then the pesticide must travel through the aquifer to a well if it is to be detected. There are several factors which can affect the movement of pesticides. Pesticides will move with water (Cheng and Koskinen, 1986); therefore, where there is more recharge to the aquifer it is more likely for a pesticide to be carried to the ground water. In most cases the pesticide will not move through the soil at the same rate as the water. This is due to the adsorption of some of the pesticide onto soil particles. The amount of the chemical that is adsorbed depends on the chemical properties of the pesticide as well as properties of the soil. Important soil properties are organic carbon content, clay content, field moisture capacity, pH and cation exchange capacity. Of these, the organic carbon content is the most important (Helling and Dragun, 1980). Unfortunately, there are only incomplete and indefinite data on organic carbon content available for the study area. Because a pesticide will only move appreciable distances through permeable materials with moving water, pesticides should only be able to contaminate the aquifer through soils that allow recharge to the aquifer.

Pesticides may also reach the ground water by movement along the annulus of an improperly cased well, or by direct contamination of the well. These are situations that are the result of negligence or mishandling of pesticides.

The permeability of soils in the study area is described in the soil surveys for the counties in the study area (Ferguson and Gray, 1971; Gray and Ferguson, 1977; Ferguson, 1979). In these reports, permeability is described by how rapidly water moves through the soil. The categories used are very slow, slow, moderately slow, moderate, moderately rapid and rapid.

One would expect a relationship between pesticide detections and the sensitivity of the site. An investigation of this relationship using geographical information systems was not possible because the necessary detail for soils and other factors determining sensitivity were not available in a magnetic useable format. Because soils are one of the most critical factors determining sensitivity and detailed soils maps were available for the area, the relationship between permeability of soils and pesticide detections was investigated. In order to limit the number of soils considered for each well, only those soils within one mile were used. It is reasonable to impose such a limit because at some point all soil types would be encountered. Maps showing the soils within one mile of each well are included in Appendix B. Soils through which water movement is moderately rapid or rapid will be considered permeable in this report. Wells C4 and M4 are within one mile of at least one permeable soil, but wells M5 and P1 are not. This indicates that bentazon and fluometuron detections may not be related to nearby soils. This could be because pesticides may travel more than one mile in the aquifer, or they could leach through less permeable soils, or the detections could be the result of well contamination.

The occurrence of bentazon in three wells does not represent widespread aquifer contamination. The bentazon detections were separated by at least 14 miles (23 kilometers). None of the wells in between, or closer to the contaminated wells contained detectable levels of pesticides. Therefore, it is thought that the detections are separate incidents.

All of the contaminated wells are used in agriculture or gardening. It is possible that the wells were contaminated by back-siphoning during the mixing or application of the pesticides. It is also possible that the pesticide could have been carried down along macropores, e.g., the zone between the well casing and the surrounding material. If either of these cases occurred, the pesticide detections may represent a contaminated well as opposed to a contaminated aquifer. It is not possible to determine which potential pathway of contamination was taken by the pesticides. In either case, re-sampling indicates that the pesticide detections were temporary, localized occurrences that do not represent a long-term or widespread aquifer contamination.

Inorganic Data

The results of the cation, anion, and other parameter analyses are given in Clayton (1995). Ryling (1960) reported that water from the alluvial aquifer is typically of calcium bicarbonate type water. The results of analyses of the wells in this study support this statement (Clayton, 1995).

Only three of the inorganic parameters will be discussed in this report. Nitrate is included because of its agricultural importance, associated use with pesticides, and its potential health hazard. Iron is included because of its staining properties and potential interaction with nitrate. Because of its potential health hazard lead is also included (Table 6).

High iron concentrations are a common problem with water from the alluvial aquifer. Ryling (1960) reported iron concentrations as high as 26 mg/L from the alluvial aquifer. Because he does not report whether or not his samples were filtered, it is assumed that his analyses were for raw water, i.e., unfiltered samples. Iron concentrations over 0.3 mg/L are generally considered undesirable and can stain plumbing fixtures and clothing. Iron concentrations from this study, based on filtered samples, are listed in Table 6. Values range from less than the detection limit of 0.003 to 15.3 mg/L, with an average value of 3.5 mg/L.

The EPA has set an MCLG for lead at zero (EPA, 1994). Eighteen of the 31 wells for which lead was determined contained lead at or above the detection limit of 0.003 mg/L. One sample, taken from well P6, contained a lead concentration of 0.026 mg/L. This is above the action level of 0.015 mg/L which has been set by the EPA for public supply wells. The wells sampled were purged before sample water was collected. If the wells or the pumping equipment contain lead, persons drinking from the wells without purging above the action level of 0.015 mg/L which has been set by the EPA for public supply

Table 6. Nitrate, lead, and iron concentrations for individual wells.

Well	NO ₃	Fe	Pb
C1	14.35	0.605	0.002
C2	0.04	NA	NA
C3	< 0.01	NA	NA
C4	4.62	<0.003	< 0.003
C5	2.20	0.004	0.007
C6	< 0.01	0.009	< 0.003
C7	7.70	0.027	< 0.003
C8	< 0.01	1.751	< 0.003
C9	7.46	0.096	< 0.003
C10	6.52	0.014	< 0.003
C11	0.77	0.908	< 0.003
C12	0.80	0.111	< 0.003
M1	0.04	0.008	< 0.003
M2	0.03	0.008	0.004
M3	0.05	1.943	< 0.003
M4	0.05	NA	NA
M5	0.02	10.360	< 0.003
M6	< 0.01	2.442	0.003
M7	< 0.01	NA	NA
M8	0.02	9.820	< 0.003
M9	< 0.01	4.774	< 0.003
M10	< 0.01	11.175	0.003
M11	0.01	9.945	0.003
M12	2.00	0.129	< 0.003
M13	6.49	< 0.003	< 0.003
M14	0.02	0.105	< 0.003
M15	8.04	< 0.003	0.003
P1	0.02	5.170	< 0.003
P2	0.11	1.578	0.004
P3	0.41	NA	NA
P4	< 0.01	NA	NA
P5	< 0.01	NA	NA
P6	0.01	3.083	0.026
P7	0.04	13.710	0.003
P8	0.02	6.140	0.005
P9	0.07	3.173	0.003
P10	0.07	5.740	0.003
P11	0.04	15.320	0.005

NA = not analyzed

them first could be ingesting significantly more lead than indicated by the analyses in this report.

Relationships with Depth

To investigate the relationship of well depth and nitrate concentrations, wells were separated into two groups--shallow wells and deep wells. Wells 40 feet deep or less were considered shallow, and wells deeper than 40 feet were considered deep. Forty feet was chosen as the depth for demarcation because it is approximately the median depth for wells in this study. Iron has higher concentrations in deep wells. Nitrate has higher concentrations in shallow wells. Statistical significance was determined by using the Student t-test for sample means (Wine, 1964). T-test values that exceed the critical value indicate that within a 95% confidence limit, the deep and shallow samples were taken from different populations. Averages and standard deviations for deep and shallow wells, as well as t-test and critical values, are given in Table 7. Other parameter and well depth relationships are discussed in Clayton (1995).

Table 7. Averages and standard deviations for 17 deep and 21 shallow wells.

			T-test	Critical t value
Parameter	Wells ≤ 40 ft	Wells > 40 ft	value	at $\alpha = 0.05$
Depth ft				0
Average	25.6	71.8		
Std. dev.	7.7	25.9		
Fe mg/L			2.22	2.06
Average	1.82	5.22		
Std. dev.	3.16	5.19		
Nitrate-N mg/L			2.07	2.04
Average	2.53	0.53		
Std. dev.	3.92	1.86		

Oxidation, Reduction and Denitrification

Nitrate contaminated ground water can cause the sometimes fatal condition methemoglobinemia in infants (Korom, 1992). Several wells contained detectable levels of nitrate. Concentrations below 0.40 mg/L as nitrogen are considered trace amounts in this study based on background concentrations for another aquifer (Steele and McCalister, 1996). Twelve wells contained nitrate above 0.40 mg/L. The locations of these 12 wells are shown in Figure 4. Well C1 exceeded the MCL of 10 mg/L (reported as N). This well contained 14.35 mg/L nitrate-N.

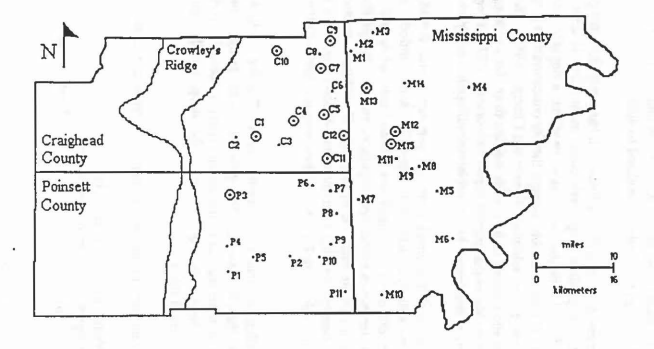


Figure 4. Locations of wells with nitrate. Wells with anomalously high nitrate concentrations are circled.

usually as ammonium nitrate (NH₄NO₃) or urea [CO(NH₂)₂], is applied to the land surface as fertilizer. The nitrogen in urea will be hydrolyzed by urease to ammonium (NH₄⁺). This can occur over a period of several days (Alexander, 1977). The ammonium nitrate will disassociate to ammonium and nitrate (NO₃⁻). The ammonium can then be converted to nitrate by microorganisms in a process known as nitrification. This process requires oxygen and typically occurs above the water table, in the soil zone. This also can occur over a period of several days (Alexander, 1977; Freeze and Cherry, 1979). Septic tanks are another source of nitrogen that could be oxidized to nitrate.

The mobility of the ammonium ion is limited because it has a positive charge, which allows it to be adsorbed onto soil particles, especially clays; its mobility is controlled by cation exchange capacity of the soil. Nitrate, on the other hand, has a negative charge, and is not significantly adsorbed. Nitrate is soluble enough that its concentration in ground water is limited by availability rather than solubility (Freeze and Cherry, 1979). The high solubility and lack of adsorption allow nitrate to be readily leached from the soil zone to the ground water, and to be very mobile in the ground water (Alexander, 1977; Freeze and Cherry, 1979). Nitrate mobility is also limited by microbial immobilization and plant uptake.

Because nitrate can be readily leached from the soil zone, one would expect nitrate contamination to be more likely in areas with permeable soils. Indeed, all twelve wells with nitrate concentrations above 0.15 mg/L are within 1 mile of at least one permeable soil, through which water moves moderately rapidly or rapidly. Maps showing the soils within one mile of each well are included in Appendix B. Nitrate occurrences are also related to depth. The deepest well that contains nitrate greater than 0.15 mg/L is 55 feet deep. This relationship is shown in Figure 5. There are nine wells in this study that are deeper than 55 feet.

Twenty-four wells within one mile of permeable soils. Twelve of these wells do not contain nitrate greater than 0.15 mg/L. The permeable soils around these wells could conceivably allow nitrate to be leached to the ground water if excessive fertilizer is applied at the surface. Five of these wells are 60 or more feet deep, which might protect them from nitrate contamination (Table 8).

Table 8. Nitrate and well depth means with respect to distance to permeable soils.

	Wells ≤ mile from permeable soil	Wells > 1 mile from permeable soil	T-test value	Critical t-value at $\alpha = 0.05$
Parameter				
Depth ft				
Average	47.8	45.4		
Std. dev.	27.3	31.0	0.24	2.03
Nitrate-N mg/L				
Average	0.03	2.56		
Std. dev.	0.03	3.9	3.21	2.07

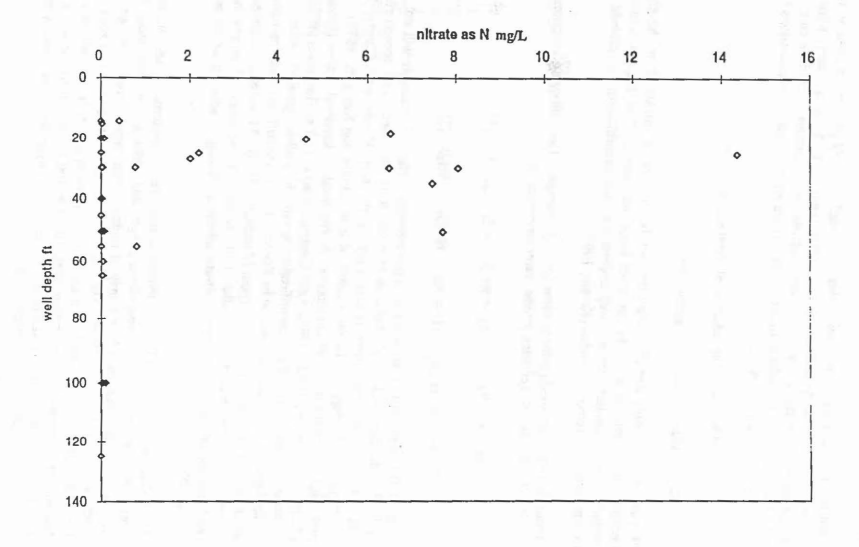


Figure 5. Nitrate versus well depth.

Because nitrate concentrations greater than 0.15 mg/L are only present in shallow wells with some of the nearby soils having moderately rapid or rapid permeability, nitrate is being removed from the water. If this were not the case the nitrate would be carried to wells near less permeable soils or to deep wells. There are two possible explanations for the removal of nitrate. They are:

- (1) denitrification -- the reduction of nitrate to N2O or N2, or
- (2) reduction of nitrate to ammonium.

Of these methods only denitrification permanently removes the nitrate from the aquifer because the ammonium could be converted back into nitrate. The factors controlling reduction to ammonium are not well understood, but denitrification is probably the dominant process removing nitrate (Korom, 1992).

Denitrification is an oxidation-reduction (redox) reaction. The following reactions can occur in a progressively reducing ground water environment.

$$5CH_2O + 4NO_3 = 2N_2(g) + 5HCO_3 + H^+ + 2H_2O$$
 (1)

$$CH_2O + Fe(OH)_{3(s)} + 7H + = Fe_2^+ + HCO_3^- + 10H_2O$$
 (2)

where CH₂O represents metabolizable organic matter. The first reaction will not occur until all of the dissolved oxygen has been removed from the water. The second reaction will not occur until the nitrate is consumed by the first (Freeze and Cherry, 1979). Fe(OH)₃₍₃₎ may be replaced by other forms of iron (Stumm and Morgan, 1970). These processes only occur in significant amounts as the result of bacterial action (Freeze and Cherry, 1979;Korom, 1992). Gillham and Cherry (1978) as well as Edmunds (1973) have described situations in which denitrification occurs in shallow ground water. Other denitrification studies are summarized by Korom (1992). Denitrifying bacteria use nitrate as an electron acceptor in place of oxygen (Alexander, 1977). Therefore, nitrate can give an indication of the oxidation potential of the water. The presence of nitrate indicates oxidizing conditions. The absence of nitrate indicates reducing conditions or the lack of a source of nitrate.

The reduction of iron from ferric to ferrous is mediated by bacteria. These bacteria use iron as an electron acceptor in areas where oxygen and nitrate are not available. For this process to occur there must be a supply of metabolizable organic matter to serve as an electron donor (Alexander, 1977). The iron could be leached from the soil zone or from the aquifer material. The supply of organic matter is much greater in the soil zone than in the aquifer. Therefore, it seems more likely that the iron is reduced in the soil zone and then leached to the aquifer. Chapelle and Lovely (1992) have described a case where iron is known to be reduced in a confined aquifer.

Iron was only considered important if it was above 0.15 mg/L (the mean value for all wells). Eighteen of the 31 wells that were analyzed for cations contained iron concentrations greater than 0.15 mg/L. Because soluble iron occurs under reducing conditions, iron occurrences should be related to local soils. Impermeable soils typically become saturated after rain. Saturated conditions prevent oxygen from being replenished to the soil so that oxygen used by microorganisms is not replaced. Certain types of bacteria are then able to use iron (or nitrate, if present) in place of oxygen. The iron is reduced to the soluble Fe²⁺ ion and can be leached to the ground water.

Eleven of the 18 wells with iron concentration exceeding 0.15 mg/L are not within one mile of a permeable soil. Four of these wells are 40 feet deep or less. The other seven wells, including two that also had nitrate, are associated with permeable soils; however, only one of these wells is less than 40 feet deep.

Stumm and Morgan (1970) state that nitrate is not compatible with iron in waters with neutral pH. It is commonly observed that ground water does not contain both iron and nitrate (Korom, 1992), as is the case for this study. With only two exceptions, no well has more than 0.15 mg/L of both iron (Fe) and nitrate (NO₃⁻, reported as N). This relationship is shown in Figure 6. One of the exceptions is well C1. As previously mentioned, this well contains 14.35 mg/L nitrate-N as well as 0.61 mg/L Fe. The other exception is well C11, which contains 0.77 mg/L nitrate and 0.91 mg/L Fe.

Although it is possible that the mutual exclusion of iron and nitrate could be due to the progressive reduction of ground water by reactions (1) and (2) stated above, this is not likely because these reactions require a sufficient supply of organic substrate. It is much more likely that the of iron reduction is occurring in the soil zone, due to the availability of organic carbon.

Denitrification can also proceed as the reaction between iron and nitrate. The reaction between NO₃ and Fe²⁺ is:

$$3Fe^{2+} + NO_3^- + 4H + = NO(g) + 2H_2O + 3Fe^{3+}$$
 $E^0 = -0.19$

Where E° is the standard potential of the reaction. A negative standard potential indicates that the reaction will occur spontaneously (Krauskopf, 1979). Any iron converted to Fe³+ would quickly be precipitated, most likely as Fe(OH)₃ (Stumm and Morgan, 1970). The fact that there are two wells that contain iron and nitrate above 0.15 mg/L, as well as other wells that contain smaller amounts, indicates that the reactions do not happen instantaneously. Supporting this conclusion, Freeze and Cherry (1979) state that many oxidation-reduction reactions occur slowly. Korom (1992) reports that this reaction does not happen at significant rates without microbial activity. Certain bacteria can use iron as an electron donor in place of organic carbon. The nitrate serves as the electron acceptor (Korom, 1992). This explains why iron and nitrate are not usually present together. It also helps to explain why there is more iron in deeper parts of the aquifer. Oxygen and

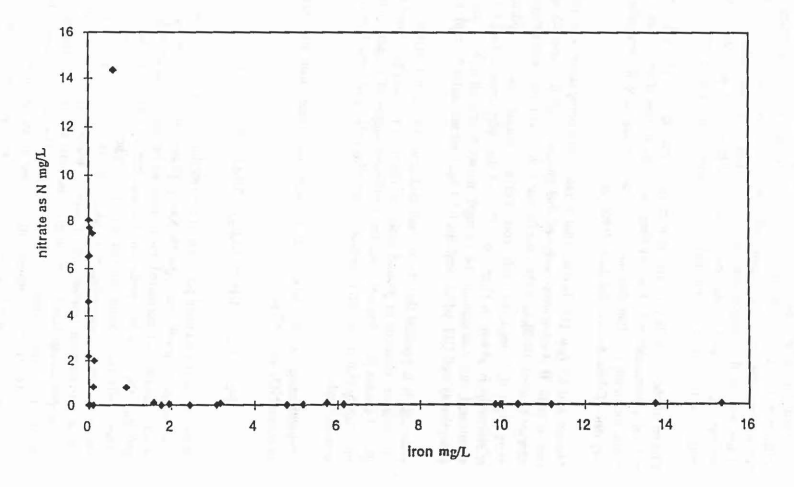


Figure 6. Nitrate versus iron.

nitrate which could oxidize the iron are likely consumed in shallow parts of the aquifer. In fact, denitrification may be occurring in the soil zone where carbon substrate is available.

A generalized conceptual model is shown in Figure 7. The general direction of ground-water flow is indicated by the arrow. Water in well A is likely to contain nitrate because the well is shallow and is surrounded by permeable soils. Water from wells B and C would not likely contain nitrate because well B is too deep and well C is surrounded by impermeable soils. Ground water that moves to deep parts of the aquifer or under impermeable soils will become reducing and denitrification will occur. Wells B and C would also contain iron. The iron could be reduced in the soils during saturated conditions and leached to the aquifer, or the iron could come from the aquifer material. Well A would not have iron because the ground water would be oxidizing and iron would exists in the insoluble ferric state. Wells B and C would have higher total dissolved solids than well A.

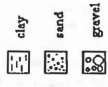
Nitrate and Pesticide Relationships

Pesticides and nitrate enter the ground water after being applied to the land surface. One would expect nitrate to be present in the wells containing pesticides. However, of the four wells that contained pesticide, only well C4 had significant amounts of nitrate. There are two reasons for the lack of a relationship between nitrate and pesticide contamination. One reason is that nitrate is more mobile. Pesticides are readily adsorbed to clay and organic matter. It is likely that the pesticides are held in the soil zone. In the soil zone, the pesticide can be degraded by microorganisms. Nitrate, on the other hand, is not significantly adsorbed to clay or organic matter and is extremely soluble. This explains the reason that nitrate is more common in ground water. The other difference between nitrate and pesticides is that nitrate can be removed by denitrification and would not be stable in reducing ground-water conditions. If a pesticides reaches the aquifer it can persist because it may not be readily degraded below the soil zone.

CONCLUSION

The data show that there is no major pesticide contamination in this area of northeastern Arkansas. Although the possibility of future contamination is not ruled out, current pesticide management practices appear to be adequately protecting the ground water from contamination.

Nitrate, on the other hand, is reaching the ground water. In one case the nitrate exceeded the maximum contaminant level for drinking water. Nitrate contamination was noted only in wells within one mile of permeable soils. This indicates that the nitrogen management practices for permeable soils should address this problem. Because nitrate is not present in deep wells or wells that are not near permeable soils, it is assumed that denitrification is occurring in the soil and perhaps the aquifer where ground water becomes reducing with



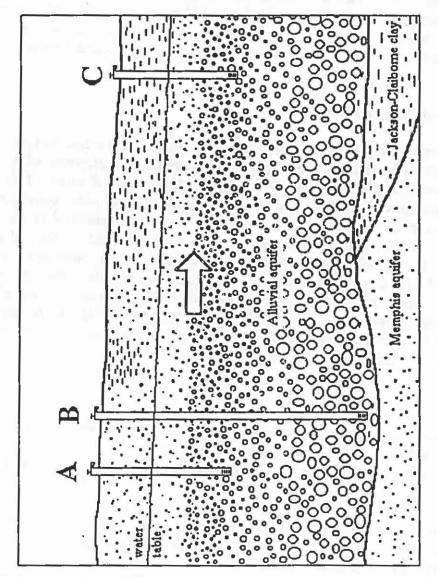


Figure 7. Conceptual model of the alluvial aquifer.

increased distance from the recharge areas. Microorganisms are using nitrate to oxidize ferrous iron or organic matter if it is available. The iron is probably being reduced in impermeable soils and then leached to the ground water or may be dissolved from the aquifer material.

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APPENDIX A

Quality Assurance Report

August 1, 1993 - October 19, 1994

Arkansas State Pesticides in Ground Water

Monitoring Project

PHASE II:

Mississippi, Craighead, and Pointsett Counties.

QUALITY ASSURANCE REPORT: ARKANSAS STATE PESTICIDES IN GROUND WATER MONITORING PROJECT PHASE II MISSISSIPPI, CRAIGHEAD AND POINSETT COUNTIES

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T. Nichols, P. Vendrell, K. Steele1

I. Introduction

Between November 1, 1993 and October 19, 1994, forty water samples were drawn from 37 wells in Mississippi, Craighead and Poinsett Counties. Ten liters of water were collected from each well, providing enough water to have a sample and a field fortified sample for each of the three methods, as well as extra water for duplicate analysis. Table 1 shows a list of the pesticides analyzed in these samples and the methods used.

Table 1. Phase II Analytes.

Metolachlor EPA/507.1 groundwater ug/L 0.75 Alachlor EPA/507.1 groundwater ug/L 0.38 Molinate EPA/507.1 groundwater ug/L 0.15 Atrazine EPA/507.1 groundwater ug/L 0.13 Metribuzin EPA/507.1 groundwater ug/L 0.15 Norflurazon EPA/507.1 groundwater ug/L 0.50 Linuron NPS/4 groundwater ug/L 0.25 Flumeturon NPS/4 groundwater ug/L 0.10 Cyanazine NPS/4 groundwater ug/L 0.58 Diuron NPS/4 groundwater ug/L 0.58 Diuron NPS/4 groundwater ug/L 0.58 Diuron NPS/4 groundwater ug/L 0.070 2,4-D EPA/515.2 groundwater ug/L 0.20 Bentazon EPA/515.2 groundwater ug/L 0.20 Acifluorfen EPA/515.2 groundwater ug/L 0.096	Compound	Source/Method	Matrix	Units	EDL
	Alachlor Molinate Atrazine Metribuzin Norflurazon Linuron Flumeturon Cyanazine Diuron 2,4-D Bentazon	EPA/507.1 EPA/507.1 EPA/507.1 EPA/507.1 EPA/507.1 NPS/4 NPS/4 NPS/4 NPS/4 EPA/515.2 EPA/515.2	groundwater groundwater groundwater groundwater groundwater groundwater groundwater groundwater groundwater groundwater	ug/L ug/L ug/L ug/L ug/L ug/L ug/L ug/L	0.38 0.15 0.13 0.15 0.50 0.25 0.10 0.58 0.070 0.20 0.20

Of the 37 wells tested, four showed trace levels of pesticides. Bentazon was found in three wells at 2.5, 0.3 and 0.2 ug/L. Fluometuron at 0.5 ug/L was found in one well. The well with 2.5 ug/L of Bentazon was unavailable for retesting despite two return trips to the well. The other three wells were resampled but no trace of pesticide was found again.

II. Interpretation of QC data.

During the project, seven trips were made to collect water. The samples collected on each trip were extracted and analyzed as a batch, with each batch being subdivided into the three methods of analysis indicated in Table 1. The tabulated quality control data follow this format. Thus, for each sampling trip the

¹Arkansas Water Resources Center, University of Arkansas.

reported analysis results are accompanied by three QC sheets, one for each method. The following paragraphs are intended as an aid in interpreting the QC data.

The major QA/QC concern of this study is to demonstrate an ongoing ability to detect small amounts of pesticides in various ground waters. Primary to this purpose are the three (one for each method) field fortified samples collected from each well and spiked with low levels of the appropriate pesticides. Table 2 shows the concentrations of these pesticides in the "field spikes." Extraction and analysis of these field spikes were done for every well and for every method, far exceeding EPA's recommendation that one in ten samples be field fortified. A consistent, high recovery of the pesticides spiked into the various ground waters is good indication that sample extraction and analysis are acceptable, that nothing in the ground waters is preventing the detection of pesticides in the non-fortified samples and that sample handling procedures are adequate to avoid pesticide degradation.

As a further check that small amounts of pesticide will not go unnoticed, 2X standards (containing pesticide concentrations at about two times the estimated detection limit for the pesticide) were analyzed with each batch. Concentrations for the 2X standards are also included in Table 2. For each batch and each method, peak areas for a 2X standard are reported to demonstrate instrument capability to detect very small amounts of pesticides. EPA holding times for samples and extracts were met without exception and samples and extracts were held at or below 4°C at all times.

Table 2. Spiking Levels.

METHOD	PESTICIDE	CONCENTRA	CONCENTRATION (ug/L)			
		FIELD SPIKE	2X STANDARD			
507	Molinate Atrazine Metribuzin Alachlor Metolachlor Norflurazon	2.00 2.06 2.10 4.08 13.72 5.90	0.40 0.41 0.42 0.81 2.74 1.18			
515.2	2,4-D Bentazon Aciflurofen	3.00 7.21 3.15	0.60 1.44 0.63			
NPS4	Cyanazine Fluometuron Diuron Linuron	6.42 1.10 0.99 3.03	1.28 0.22 0.20 0.60			

Recovery of a spiked pesticide from any field spike should be within the normal range of recovery for the laboratory doing the work. For EPA method 507 (EPA507) and National Pesticide Survey method 4 (NPS4), this laboratory has a history of successful analyses from which to determine a "normal" range of recovery for each analyte. Table 3 shows the mean recoveries and associated standard deviations for ten of the pesticides in this study. These were derived from 29 field spikes collected previously in a study of ground water in Ashley County.

Table 3. Summary of Recoveries for EPA Method 507 and National Pesticide Survey Method 4 - Mean, Standard Deviation and Range.

Chemical	N	Mean (M)	Std. Dev.(s)	Range (M±3s)
Molinate Atrazine Metribuzin Alachlor Metolachlor Norflurazon	29 29 29 29 29	82.2 97.0 103.3 97.3 95.1 99.0	7.8 13.5 17.5 14.7 13.0 20.5	58.8 - 105.6 56.4 - 137.6 50.8 - 155.8 53.3 - 141.2 56.0 - 134.2 37.4 - 160.5
EPA507 surrogate	84	100.0	17.0	49.0 - 151.0
Cyanazine Fluometuron Diuron Linuron NPS4 surrogate	29 29 29 29 80	88.9 83.8 87.3 84.0 79.5	13.3 16.8 16.1 13.9 7.7	49.0 - 128.8 33.3 - 134.3 39.1 - 135.5 42.1 - 125.8 56.5 - 102.4

The normal range of recovery is defined as the mean plus or minus 3 standard deviations. For example the mean recovery for molinate was 82.2% with a standard deviation of 7.8% yielding a range of 58.8 - 105.6% (the mean plus/minus 3 standard deviations). If the recovery of a particular analyte from a field spike is outside the normal range then the result for that analyte for that well is reported as suspect. In addition, surrogate recovery for the non-fortified samples must also fall in the normal range of surrogate recoveries which are defined in the same way. A surrogate is a pure compound not expected to be in the sample. A known amount of surrogate is added to the sample before extraction as a check on the sample preparation procedure. The normal range for surrogate recoveries for EPA507 and NPS4 are also given in Table 3.

EPA method 515.2(EPA515), used to analyze for bentazon, aciflurofen and 2,4-D, did not have a history in this laboratory prior to this study. Lacking such a history, EPA suggests 60% to 140% as the appropriate range for "start-up" work when analyzing

spiked reagent water. Experience suggests that recoveries from ground water will be more variable thus the range for this study was expanded to 55% to 145%. In future studies, the recoveries from this study will be used to develop an acceptable range of recoveries. Results for EPA515 are reported as suspect due to matrix effects if the spike recovery or the surrogate recovery was not in the specified range. In actuality, none of the recoveries in this study were so low as to cause suspicion of false negatives.

As specified in the Quality Assurance Project Plan (QAPP), QC data for nitrate consist of percent relative standard deviation (%RSD) between duplicate measurements of one sample and percent recovery (%REC) from a sample spiked with a known amount of nitrate-nitrogen. The maximum allowable %RSD is 10% and the allowable range for %REC is 80-120%. QC measurements are made on one sample from each sampling trip.

III. QA/QC Summary.

For the forty wells-including three resamples-there were a total of 520 data points (40 times 13 pesticides) of which 36, or 6.9%, have been reported as suspect. Thirty-three of these points are reported as suspect due to omission of the surrogate during analysis. Spike recoveries for all 33 points were within acceptable limits (as specified above), and there is no positive reason to doubt the validity of these results. But EPA requires that the data be reported as suspect because of the lack of surrogate information.

Spike recoveries for the three EPA515 analytes in Miss #10 were all below the minimum acceptable level, 55%. As the three points are from the same method and the same well, it is quite possible that the addition of the spike into the sample bottle was performed inaccurately. Alternatively, a problem in the extraction procedure may have caused these low recoveries. In this case, there is no positive proof of ability to recover the analytes from this particular matrix and the results are considered to be suspect due to matrix effects.

Suspect results have been highlighted with grey shading on the analysis reports and the three results associated with low spike recoveries are outlined as well as shaded. Being able to recover the minimum acceptable amount, or more, of the pesticides in the field spikes-except for the three results just mentioned-assures the researchers that no significant amounts of pesticide have gone undetected. The authors feel the pesticide QC data for these analysis results are adequate for the stated purposes of the study.

The nitrate data reported here are all acceptable. All the spike recoveries were very close to 100%. No %RSD was calculated for two of the trips as both of the duplicate measurements were at or below the detection limit. These duplicates are acceptable

even though the %RSD could not be computed. The %RSD associated with the second trip to Poinsett County is 24%. This was computed from duplicate measurements both of which were right at the detection limit (0.01 mg/L). The first measurement was 0.014 and the second .011. Realistically, these measurements are very close to each other and indicate that the measurement process is working properly. The %RSD is not a good measure when the reported concentrations are so close to the detection limit. Alternatives to using the %RSD to evaluate duplicate mesurements are now being considered.

ANALYSIS RESULTS AND QUALITY CONTROL DATA

RESULTS OF PESTICIDE MONITORING: TRIP #1 TO MISSISSIPPI COUNTY-NOVEMBER, 1993.

(unk = unknown, NC = not collected, ND = not detected)

(= suspect, see text)

	1	2	3	4	5	6
WELL ID:	MISS #1	MISS #2	MISS #3	MISS#4	MISS #5	MISS #6
DATE SAMPLED:	NOV 1,1993	NOV 1,1993	NOV 1,1993	NOV 2,1993	NOV 2,1993	NOV 2,1993
LATITUDE:	35° 56' 34"	35° 57' 23°	35° 58' 15"	35° 52' 19"	36° 40' 33"	35° 34' 29"
LONGITUDE:	90° 17' 00*	90° 16' 02"	90° 13' 33"	89° 59' 24°	90° 05' 00"	90° 02' 32"
DEPTH OF WELL, ft:	100+	100+	100+	50	30-50	125
pH, standard units:	7.6	7.4	7.5	6.7	6.7	7.1
CONDUCTIVITY AT 25° C , umhos/cm:	543	462	483	811	653	486
TEMPERATURE, ° C :	16	18	16	16	17	17
NITRATE, mg/L:	0.04	0.03	0.04	0.05	0.07	< 0.01
ACIFLUORFEN, ug/L	ND	ND	NO	ND	ND	ND
ALACHLOR, ug/L:	ND	NĐ	ND	ND	ND	ND
ATRAZINE,ug/L:	ND	ND	ND	ND	ND	ND
BENTAZON, ug/L	ND	ND	ND.	2.5*	0:3@	NÖ
CYANAZINE, ug/L:	ND	ND	ND	ND	ND	ND
DIURON, ug/L:	ND	ND	ND	ND	ND	ND
FLUOMETURON, ug/L:	ND	ND	ND	ND	ND	ND
LINURON, ug/L:	ND	ND	ND	ND	ND	ND
METOLACHLOR, ug/L:	ND .	ND	ND	ND	ND	ND
METRIBUZIN, ug/L:	ND	ND	ND	ND	ND	ND
MOLINATE, ug/L:	ND	ND	ND .	ND .	ND	ND
NORFLURAZON, ug/L	ND	ND	ND	, ND	ND	ND
2,4-D, ug/L	ND	ND	ND	ЙÓ	ND	ND

CONFORMATION POSITIVE, UNABLE TO COLLECT SAMPLE FOR VERIFICATION

EPA METHOD 507

		SURROGATE	MOLINATE	ATRAZINE	METRIBUZIN	ALACHLOR	METOLACHLOR	NORFLURAZON
FIELD FORTIFIED	SAMPLES							
	MISS #1	87	102	105	102	95	115	114
	MISS #2	91	99	110	107	99	121	123
	MISS #3	97	106	113	114	101	124	121
	MISS #4	103	105	107	104	97	119	116
	MISS #5	93	99	86	97	98	110	105
	MISS #6	83	110	114	120	105	129	127
NON CORTIFIED	CAREDI FO							
NON-FORTIFIED	MISS #1	96						
	MISS #2	70						
	MISS #3	77						
	MISS #4	71						
	MISS #5	69						
	MISS #B	86						
LAB BLANKS								
LAD BLANKS	P266	78						
	P267	76		*				
			CONCENTRA	ATIONS FOR I	LAB BLANKS			
	P266		0	0	o	0	0	0
	P287		o	o	ŏ	ō	o	o
			PEAK AREAS	S FOR A 2X	STANDARD			
	2X STANDARD		MOLINATE 13211	ATRAZINE 43630	METRIBUZIN 9039	ALACHLOR 12527	METOLACHLOR 33287	NORFLURAZON 27811
			DUPLICA"	TE ANALYSIS				
			SISI D DUIDLICA	TE CHODOCA	TE AREA COMPA	ADICON		
		P202	FIELD DUPLICA	P204	TE AREA COMP	%RSD		
		89889		81591		9.68		
			MACHINE NUE	LICATE CURO	DOATE ADEA OO	MADA DICON		
			MACHINE DUP		OGATE AREA CO			
		1ST RUN		2ND RUN		%RSD 3.48		
		113799		109910		3.48		

QUALITY CONTROL DATA FOR PESTICIDE MONITORING: TRIP #1 TO MISISSIPPI COUNTY - NOVEMBER, 1993

EPA METHOD 515

		SURROGATE	2,4-D	BENTAZON	ACIFLUROFEN
FIELD FORTIFIED SAMPLES					
MISS #1		122	104	104	113
MISS #2		102	87	97	96
MISS #3		88	74	83	82
MISS #4		98	85	129	94
MISS #5		113	91	116	110
MISS #6		81	74	87	85
NON-FORTIFIED SAMPLES					
MISS #1		129			
MISS #2		122			
MISS #3		OMITTED			
MISS #4		OMITTED			
MISS #5		OMITTED			4
MISS #6		OMITTED			
LAB BLANKS					
P284		95			
P265		OMITTED			
		CONCENTRAT	IONS FOR	LAB BLANKS	
			•	_	
P264 P265			0	0	0
F203			U	0	O
		PEAK AREAS	FOR A 2X	* STANDARD	
		SURROGATE	2,4-D	BENTAZON	ACIFLUROFEN
2X STANDARD		38266	35088	22657	106688
		DUPLICATE	ANALYSI	S	=
		EIELD DUDU ICATI	E SUPPOS	ATE AREA COMPA	DISON
	P214	TIEED DOI EIGHT	P215	ALE AILEA COMI A	%RSD
	48794		61104		8.81
	40/94		01104		0.01
		MACHINE DUPLIC	CATE - SURP	OGATE AREA CON	PARISON
	1ST RUN		2ND RUN		%RSD
	49909		49776		0.27

ANALYTE CONCENTRATIONS ARE ABOUT 2 TIMES THE EPA ESTIMATED DETECTION LIMIT

QUALITY CONTROL DATA FOR PESTICIDE MONITORING: TRIP #1 TO MISISSIPPI COUNTY - NOVEMBER, 1993

NPS METHOD 4

			CYANAZINE	FLUOMETURON	DIURON	LINURON	SURROGATE
	FIELD FORTIFIED SAMPLES						
	MISS #1		99	105	98	90	74
	MISS #2		94	103	90	80	73
	MISS #3		100	101	96	90	82
	MISS #4		102	103	97	90	80
	MISS #5		90	88	91	81	79
	MISS #6		92	94	87	81	81
	NON CORTIFIED CAMPLES						
	NON-FORTIFIED SAMPLES MISS #1						85
	MISS #2						78
	MISS #2						87
	MISS #4						77
	MISS #5						78
	MISS #6						87
	141133 #0						0,
>	LAB BLANKS						
_	P268						74
_	P270					31	76
			CONCENTRAT	TIONS FOR LAB	RIANKS		
			CONCENTIA	HONG FOIL EAD	DEANNO		
	P268		0	0	0	0	
	P270		0	0	0	0	
			PEAK AREAS	FOR A 2X* ST	ANDARD		
			CYANAZINE	FLUOMETURON	DIURON	LINURON	SURROGATE
	2X STANDARD		2988	922	2178	8545	210814
			DUBLICATE	ANALYSIS			
			DOFFICATE	ANALISIS			
			FIELD DUPLICAT	E - SURROGATE AF	REA COMPAR	ISON	
	P2			P220		%RSD	
	185	203		184481		11.03	

[&]quot;ANALYTE CONCENTRATIONS ARE ABOUT 2 TIMES THE EPA ESTIMATED DETECTION LIMIT

QUALITY CONTROL DATA FOR PESTICIDE MONITORING: TRIP #1 TO MISISSIPPI COUNTY - NOVEMBER, 1993 NITRATE

SPIKE RECOVERY

WELL NUMBER

% RECOVERY

MISS 6

102%

DUPLICATE ANALYSIS

1ST MEASUREMENT

2ND MEASUREMENT

% RSD

0.07 mg/L

0.07 mg/L

1.00%

RESULTS OF PESTICIDE MONITORING: TRIP #1 TO CRAIGHEAD COUNTY-NOVEMBER, 1993.

(unk = unknown, NC = not collected, ND = not detected)

1	_	suspect.	200	tes

	1	2	3	4	5	6
WELL ID:	CH#1	CH#2	СН#3	CH#4	CH#5	CH#6
DATE SAMPLED:	NOV 22,1993					
LATITUDE:	35° 46' 22"	35° 46' 25°	35° 45' 26"	35° 47' 27"	35° 48' 12"	35° 51' 46"
LONGITUDE:	90° 29' 27"	90° 33' 09°	90° 27' 42°	90° 25' 30"	90° 21' 01°	90° 18' 53°
DEPTH OF WELL, ft:	25	30	20	20	25	20
pH, standard units:	6.5	7.5	7.9	6.9	7.4	7.6
CONDUCTIVITY AT 25° C , umhos/cm:	314	684	262	170	404	274
TEMPERATURE, ° C:	16	16	17	18	18	17
NITRATE, mg/L:	14.35	0.04	<0.01	5.5	2.2	< 0.01
ACIFLUORFEN, ug/L	ND	ND	ND	ND	ND	ND
ALACHLOR, ug/L:	ND	ND	ND	ND	ND	ND
ATRAZINE,ug/L:	ND	ND	ND	ND	ND	ND
BENTAZON, ug/L	ND	ND	ND	ND	ND	ND
CYANAZINE, ug/L:	ND	ND .	ND	ND	ND	ND
DIURON, ug/L:	ND	ND	ND	ND	ND	ND
FLUOMETURON, ug/L:	ND	ND	ND	0.5@	ND	ND
LINURON, ug/L:	ND	ND	ND	ND	ND	ND
METOLACHLOR, ug/L:	ND	ND	ND	ND	ND	ND
METRIBUZIN, ug/L:	ND	ND	ND	ND	ND	ND
MOLINATE, ug/L:	ND	ND	ND	ND	ND	ND
NORFLURAZON, ug/L	ND	ND	ND	ND	ND	ND
2,4-D, ug/L	ND	ND	ND	ND	ND	ND

QUALITY CONTROL DATA FOR PESTICIDE MONITORING: TRIP #1 TO CRAIGHEAD COUNTY - NOVEMBER, 1993.

EPA METHOD 507

		SURROGATE	MOLINATE	ATRAZINE	METRIBUZIN	ALACHLOR	METOLACHLOR	NORFLURAZON
FIELD FORTIFIED S	AMPLES							
	CH#1	90	81	96	90	98	97	99
	CH#2	94	80	92	84	93	94	94
	CH#3	97	82	91	86	96	95	94
	CH#4	103	84	93	86	99	96	93
(CH#5	121	91	97	91	101	101	99
!	CH#6	115	86	96	92	101	101	99
NON-FORTIFIED SA	AMPI ES							
	CH#1	112						
	CH#2	88						
	CH#2	100						
	CH#4	98						
	CH#5	91						
	CH#6	112						
LAB BLANKS								
	P366	80						
	P367	120						
			CONCENTRA	TIONS FOR	LAB BLANKS			
			CONCENTRA	TIONS FOR	LAB BLANKS			
1	P366		- 0	o	0	0	0	0
ı	P367		0	О	0	0	0	0
			PEAK AREAS	FOR A 2X	* STANDARD			
			MOUNTE	A TD A 7151E	METRIBUZIN	ALACHLOR	METOLACHLOR	NORFLURAZON
9	2X STANDARD		MOLINATE 12878	ATRAZINE 28087	7999	8830	30392	22191
			DI IDI ICA:	TE ANALYSIS				
			DUFLICA	IE ANALI SIG				
			FIELD DUPLICA		TE AREA COMPA			
		P303		P305		%RSD		
		122476		113663		7.46		
			MACHINE DUP	LICATE - SURR	OGATE AREA CON			
		1ST RUN	MACHINE DUP	LICATE - SURRI	OGATE AREA CON	MPARISON %RSD 2.77		

^{*}ANALYTE CONCENTRATIONS ARE ABOUT 2 TIMES THE EPA ESTIMATED DETECTION LIMIT

QUALITY CONTROL DATA FOR PESTICIDE MONITORING: TRIP #1 TO CRAIGHEAD COUNTY - NOVEMBER, 1993.

EPA METHOD 515

			SURROGATE	2,4-D	BENTAZON	ACIFLUROFEN
FIELD FORTIFIED S	SAMPLES					
	CH#1		85	85	90	82
	CH#2		115	101	104	104
	CH#3		93	65	74	72
	CH#4		112	95	98	95
	CH#5		115	97	104	100
	CH#8		123	104	108	104
NON-FORTIFIED SA	AMPLES					
	CH#1		113			
	CH#2		135			
	CH#3		122			
	CH#4		129			
	CH#5		138			
	CH#6		106			
LAB BLANKS						
	P361		128			
	P362		103			
			CONCENTRAT	IONS FOR	LAB BLANKS	
	P361			o	0	0
	P362			0	0	0
			PEAK AREAS F	OR A 2X	* STANDARD	
	2X STANDARD		SURROGATE 49469	2,4·D 12270	BENTAZON 36110	ACIFLUROFEN 136383
			DUPLICATE	ANALYSIS	S	
			FIELD DUPLICATE	- SURROGA	TE AREA COMPA	RISON
		P329		P330		%RSD
		39274		37819		3.77
			MACHINE DUPLIC	ATE - SURR	OGATE AREA CON	MPARISON
51		1ST RUN		2ND RUN		%RSD
170		36034		36311		0.77

^{*}ANALYTE CONCENTRATIONS ARE ABOUT 2 TIMES THE EPA ESTIMATED DETECTION LIMIT

QUALITY CONTROL DATA FOR PESTICIDE MONITORING: TRIP #1 TO CRAIGHEAD COUNTY - NOVEMBER, 1993 NPS METHOD 4

PERCENT RECOVERIES

		CYANAZINE	FLUOMETURON	DIURON	LINURON	SURROGATE
	FIELD FORTIFIED SAMPLES		-			
	CH#1	119	98	99	97	75
	CH#2	114	76	70	94	80
	CH#3	110	76	71	95	82
	CH#4	119	92	75	99	82
	CH#5	115	76	74	97	85
	CH#8	121	81	61	89	83
	NON-FORTIFIED SAMPLES					
	CH#1					75
	CH#2					78
	CH#3					72
	CH#4	6				65
	CH#5					66
	CH#6					73
D	LAB BLANKS	- 9.				
16	P369B					75
0.	P370B					81
		CONCENTRA	TIONS FOR LAB	BLANKS		
	P369B	0	o	0	0	
	P370B	0	0	0	0	

DUPLICATE ANALYSIS

FIELD DUPLICATE - SURROGATE AREA COMPARISON
P314 P315 %RSD
164729 149542 9.66

MACHINE DUPLICATE - SURROGATE AREA COMPARISON

none

^{*}ANALYTE CONCENTRATIONS ARE ABOUT 2 TIMES THE EPA ESTIMATED DETECTION LIMIT

QUALITY CONTROL DATA FOR PESTICIDE MONITORING: TRIP #1 TO CRAIGHEAD COUNTY - NOVEMBER, 1993 NITRATE

SPIKE RECOVERY

WELL NUMBER

% RECOVERY

CH#2

96%

DUPLICATE ANALYSIS

1ST MEASUREMENT

2ND MEASUREMENT

% RSD

14.35 mg/L

14.41mg/L

0.40%

RESULTS OF PESTICIDE MONITORING: TRIP #1 TO POINSETT COUNTY - DECEMBER, 1993.

(unk = unknown, NC = not collected, ND = not detected)

(= suspect, see text)

	1	2	3	4	5	6
WELL ID:	POIN #1	POIN #2	POIN #3	POIN #4	POIN #5	MISS #7
DATE SAMPLED:	DEC 6,1993	DEC 6,1993	DEC 6,1993	DEC 6,1993	DEC 6,1993	DEC 7,1993
LATITUDE:	35° 31' 29"	35° 33' 16"	35° 40' 33"	35° 34' 31 '	35° 36' 15"	35° 38' 43"
LONGITUDE:	90° 34' 14"	90° 25' 48"	90° 34' 18°	90° 34' 53"	90° 31' 09°	90° 16' 07"
DEPTH OF WELL, ft:	100+	100	14	40	14	25
pH, standard units:	7.4	7.3	7.3	7.3	7.6	7
CONDUCTIVITY AT 25° C, umhos/cm:	548	554	487	286	199	597
TEMPERATURE, ° C :	15	16	17	16	18	16
NITRATE, mg/L:	< 0.01	0.11	0.41	0.01	<0.01	< 0.01
ACIFLUORFEN, ug/L	ND	ND	ND	ND	ND	ND
ALACHLOR, ug/L:	ND	ND .	ND .	ND	ND	ND
ATRAZINE,ug/L:	ND	ND	ND	ND	ND	ND
BENTAZON, ug/L	0.2@	ND	ND	ND	ND	ND
CYANAZINE, ug/L:	ND	ND	ND	ND	ND	ND
DIURON, ug/L:	ND	ND	ND	ND	ND	ND
FLUOMETURON, ug/L:	ND	ND	ND	ND	ND	ND
LINURON, ug/L:	ND	ND	ND	ND	ND	ND -
METOLACHLOR, ug/L:	ND	ND	ND	ND	ND	ND
METRIBUZIN, ug/L:	ND	ND	ND	ND	ND	ND
MOLINATE, ug/L:	ND	ND	ND	ND	ND	ND
NORFLURAZON, ug/L	ND	ND	ND	ND	ND	ND
2,4-D, ug/L	ND	ND	ND	ND	ND	ND

QUALITY CONTROL DATA FOR PESTICIDE MONITORING: TRIP #1 TO POINSETT COUNTY - DECEMBER, 1993.

EPA METHOD 507

PERCENT RECOVERIES

		SUP	ROGATE	MOLINATE	ATRAZIN	E	METRIBUZIN	Α	LACHLOR	METO	DLACHLOR	NOR	FLURAZON
FIELD FORTIFIED SA	MPLES												
PC	OIN #1		117	104	121		112		118		120		127
PO	OIN #2		115	92	111		103		108		110		115
PC	OIN #3		120	95	110		101		108		111		114
PC	OIN #4		96	91	105		95		101		104		108
Po	OIN #5		108	87	107		96		103		105		107
M	IISS #7		96	78	83		69		85		95		95
NON-FORTIFIED SAN	MPLES												
PC	OIN #1		109										
PC	DIN #2		100										
PC	OIN #3		89										
PC	OIN #4		80										
PC	OIN #5		79										
М	IISS #7		82										
LAB BLANKS													
P	470		118										
P4	471		101										
P4	472		81										
				CONCENTRA	ATIONS FO	OR L	AB BLANKS				-2		
P4	470			0	0		0		0		0		0
P	471			0	0		0		0		0		0
P4	472			0	0		0		0		0		0

DUPLICATE ANALYSIS

MACHINE DUPLICATE - SURROGATE AREA COMPARISON

попе

^{*}ANALYTE CONCENTRATIONS ARE ABOUT 2 TIMES THE EPA ESTIMATED DETECTION LIMIT

PERCENT RECOVERIES

		SU	RROGATE	2,4-D	BENTAZON	ACIFLUROFEN
FIELD FORTIFIED SA	MPLES					
	POIN #1		78	77	80	70
	POIN #2		54	66	67	61
	POIN #3		71	68	74	71
	POIN #4	.4	63	72	70	69
	POIN #5		73	78	74	75
	MISS #7		58	60	62	58
NON-FORTIFIED SAM	APLES					
	POIN #1		87			
	POIN #2		90			
	POIN #3		85			
	POIN #4		110			
	POIN #5		84			
	MISS #7		93			
LAB BLANKS						
	P476		101			
	P477		116			
	P478		88			
		CON	ICENTRAT	TIONS FOR	LAB BLANKS	
	P478		0	0	0	0
	P477		0	0	0	0
	P478		0	0	0	o
		PEA	K AREAS	FOR A 2X	• STANDARD	
		SH	RROGATE	2,4-D	BENTAZON	ACIFLUROFEN
			44507	5283	20528	85859
			44007	0200	20020	

DUPLICATE ANALYSIS

FIELD DUPLICATE - SURROGATE AREA COMPARISON

P414 31409

%RSD 40898 26.25

MACHINE DUPLICATE - SURROGATE AREA COMPARISON

P415

21

QUALITY CONTROL DATA FOR PESTICIDE MONITORING: TRIP #1 TO POINSETT COUNTY - DECEMBER, 1993

NPS METHOD 4

PERCENT RECOVERIES

	CYANAZINE	FLUOMETURON	DIURON	LINURON	SURROGATE
FIELD FORTIFIED SAMPLES					
POIN #1	97	112	112	100	73
POIN #2	118	130	115	104	86
POIN #3	105	132	112	99	80
POIN #4	120	103	108	93	67
POIN #5	130	115	116	102	86
MISS #7	119	109	110	98	84
NON-FORTIFIED SAMPLES					
POIN #1					62
POIN #2					82
POIN #2					89
POIN #4					74
POIN #5					90
MISS #7					82
WISS #7					02
LAB BLANKS					
P474					87
P475					82
	CONCENTRA	TIONS FOR LAB E	BLANKS		
P474	0	0	0	0	
P475	0	o	ō	0	
	PEAK AREAS	FOR A 2X* STA	NDARD		
	CYANAZINE	FLUOMETURON	DIURON	LINURON	
	1278	1262	1766	6671	
	DUPLICAT	E ANALYSIS			
	FIELD DUPLICAT	re - SURROGATE AR	EA COMPARI	SON	

MACHINE DUPLICATE - SURROGATE AREA COMPARISON

%RSD

3.72

one

P429

182481

P420

189393

QUALITY CONTROL DATA FOR PESTICIDE MONITORING: TRIP #1 TO POINSETT COUNTY - DECEMBER, 1993 NITRATE

SPIKE RECOVERY

WELL NUMBER

% RECOVERY

POIN 1

100%

DUPLICATE ANALYSIS

1ST MEASUREMENT

2ND MEASUREMENT

% RSD

<0.01mg/L

0.01mg/L

NC

RESULTS OF PESTICIDE MONITORING: TRIP #2 TO MISSISSIPPI COUNTY - MARCH, 1994.

(unk = unknown, NC = not collected, ND = not detected)

(= suspect, see text)

	1	2	3	4	6
WELL ID:	MISS#5R	MISS#8	MISS#9	CH#4R	POIN #1R
DATE SAMPLED:	MAR 28,1994	MAR 28,1994	MAR 28,1994	MAR 29,1994	MAR 29,1994
LATITUDE:	35° 40' 33°	35° 43' 12"	35° 42' 54"	35° 47' 27°	35° 31' 29"
LONGITUDE:	90° 05' 00"	90° 07' 30"	90° 08' 38°	90° 26' 30°	90° 34' 14"
DEPTH OF WELL, ft:	30-50	50	45	20	100+
pH, standard units:	6.7	7.2	7.1	7.2	7.4
CONDUCTIVITY AT 25° C, umhos/cm:	476	766	515	159	476
TEMPERATURE, ° C:	17	17	16	17	17
NITRATE, mg/L:	0.02	0.02	<0.01	4.62	0.02
ACIFLUORFEN, ug/L	ND	ND	ND	ND'	ND
ALACHLOR, ug/L:	ND	ND	· ND	ND	ND
ATRAZINE,ug/L:	ND	ND	ND	ND	ND
BENTAZON, ug/L	מא	ND	ND	ND	ND
CYANAZINE, ug/L:	ND	ND	ND	ND	ND
DIURON, ug/L:	ND	ND	ND	ND .	ND
FLUOMETURON, ug/L:	ND	ND	ND	ND	ND
LINURON, ug/L;	ND	ND	ND	ND	ND
METOLACHLOR, ug/L:	NO	ND	ND	ND	ND
METRIBUZIN, ug/L:	ND	ND	ND	ND	ND
MOLINATE, ug/L:	ND	ND	ND	ND	ND
NORFLURAZON, ug/L	ND	ND	ND	ND	ND
2,4-D, ug/L	ND	ND	ND	ND	ND

EPA METHOD 507

PERCENT RECOVERIES

		SURROGATE	MOLINATE	ATRAZINE	METRIBUZIN	ALACHLOR	METOLACHLOR	NORFLURAZON
FIELD FORTIFIE	D SAMPLES							
	MISS #5R	77	81	98	93	92	96	95
	MISS #8	75	82	98	93	93	98	97
	MISS #9	83	90	102	93	96	101	98
	CH #4R	110	100	107	101	106	107	100
	POIN #1R	96	98	105	100	100	108	101
NON-FORTIFIED	SAMPLES							
	MISS #5R	omitted						
	MISS #8	68						
	MISS #9	72						
	CH #4R	99						
	POIN #1R	61						
LAB BLANKS								
EAD DEATH	P561	82						
	P583	71						
				. =:0::0 =00				
			CONCENTRA	A HONS FOR	LAB BLANKS			
	P561		O	0	0	0	0	0
	P563		0	0	0	0	0	0

DUPLICATE ANALYSIS

FIELD DUPLICATE - SURROGATE AREA COMPARISON

P410 P418 %RSD 84088 90408 7.24

MACHINE DUPLICATE - SURROGATE AREA COMPARISON

none

EPA METHOD 515

PERCENT RECOVERIES

			SURROGATE	2,4-D	BENTAZON	ACIFLUROFEN
FIELD FORTIFIED	SAMPLES					
	MISS #5R		50	57	83	59
	MISS #8		66	79	87	67
	MISS #9		75	92	79	82
	CH #4R		64	138	79	75
	POIN #1R		84	107	108	104
NON-FORTIFIED :	SAMPLES					
	MISS #5R		omitted			
	MISS #8		omitted			
	MISS #9		omitted		4	
	CH #4R		omitted			
	POIN #1R		omitted			
LAB BLANKS						
LAD DEAMING	P580		63			
	P569		74			
			7.			
			CONCENTRA	TIONS FOI	R LAB BLANK	S
	P580			0	0	0
	P569			0	ō	o
						_
			PEAK AREAS	FOR A 2	X* STANDAR	D
				2,4-D	BENTAZON	ACIFLUROFEN
	2xSTANDARD		7	47322	29491	152952
			DUPLICATE A	NALYSIS		
			FIELD DUPLICAT		SATE AREA CON	
		P539		P530		%RSD
		omitted		38834		none
			MACHINE DUPL			
		1ST RUN		2ND RUN		%RSD
		33156		30151		9.49

^{*}ANALYTE CONCENTRATIONS ARE ABOUT 2 TIMES THE EPA ESTIMATED DETECTION LIMIT

A25

NPS METHOD 4

PERCENT RECOVERIES

			CYANAZINE	FLUOMETURON	DIURON	LINURON	SURROGATE
FIELD FORTIFIED S	SAMPLES						
	MISS #5R		87	86	91	81	OMITTED
	MISS #8		85	85	91	79	OMITTED
	MISS #9		89	86	92	85	OMITTED
	CH #4R		85	84	88	78	OMITTED
	POIN #1R		89	80	88	75	OMITTED
NON FORTIFIED O	*******						
NON-FORTIFIED SA	MISS #5R						102
	MISS #8						84
							84
	MISS #9 CH #4R						88
	POIN #1R						94
LAB BLANKS							
END DEMINIO	P566					ř	102
	P567						98
	1007						
			CONCENTRA	TIONS FOR LAB	BLANKS		
	P566		0	0	0	0	
	P567		0	О	0	0	
			PEAK AREAS	FOR A 2X* STA	NDARD		
			CYANAZINE	FLUOMETURON	DIURON	LINURON	
			5507	2899	7252	17983	
			9907	2699	1202	17903	1000
			DUPLICAT	E ANALYSIS			
			FIELD DUPLICAT	TE - SURROGATE AR	EA COMPARI	SON	
		P544		P546		%RSD	
		349302		357098		2.21	
		343302		007000		G. 1 G. T	

MACHINE DUPLICATE - SURROGATE AREA COMPARISON

enon

NITRATE

SPIKE RECOVERY

WELL NUMBER

% RECOVERY

MISS 5R

100%

DUPLICATE ANALYSIS

1ST MEASUREMENT

2ND MEASUREMENT

% RSD

<0.01 mg/L

<0.01 mg/L

0.00%

RESULTS OF PESTICIDE MONITORING: TRIP #3 TO MISSISSIPPI COUNTY - JUNE, 1994. (unk = unknown, NC = not collected, ND = not detected) (= suspect, see text)

	1	2	3	4	5	6
WELL ID:	MISS#10	MISS #11	MISS#12	MISS#13	MISS #14	MISS#15
DATE SAMPLED:	JUNE 13,1994	JUNE 13,1994	JUN 13,1994	14-Jun-94	JUN 14,1994	JUN 14,1994
LATITUDE:	35° 28' 09"	35° 43' 36"	35° 46' 42"	36° 51' 52"	35° 52' 19"	35° 45' 24"
LONGITUDE:	90° 13' 02"	90° 10' 30"	90° 10' 39"	90° 14' 52"	90° 08' 52"	90° 11' 16"
DEPTH OF WELL, ft:	55	85	277	30 -	20-40	30
pH, standard units:	7.2	7.1	6.7	7.2	6.7	6.1
CONDUCTIVITY AT 25° C , umhos/cm:	454	560	513	557	184	262
TEMPERATURE, ° C :	17.5	17.5	19	19	17	18
NITRATE, mg/L:	< 0.01	0.01	2	6.49	0.02	8.04
ACIFLUORFEN, ug/L	ND	ND	ND	ND	ND	ND
ALACHLOR, ug/L:	ND	ND	ND	ND	ND	ND
ATRAZINE,ug/L:	ND	ND	ND	ND	ND	ND
BENTAZON, ug/L	ND	ND	ND	ND	ND	ND
CYANAZINE, ug/L:	ND	ND	ND	ND	ND	ND
DIURON, ug/L:	ND	ND "	ND	ND	ND	ND
FLUOMETURON, ug/L:	ND	ND	ND	ND	, ND	ND
LINURON, ug/L:	ND	ND	ND	ND	ND	ND
METOLACHLOR, ug/L:	ND	ND	ND	ND	ND	ND
METRIBUZIN, ug/L:	ND	ND	ND	ND	ND	ND
MOLINATE, ug/L:	ND	ND	ND	ND	ND	ND
NORFLURAZON, ug/L	ND	ND	ND	ND .	ND	ND
2,4-D, ug/L	ND	ND	ND	ND	ND	ND

PERCENT RECOVERIES

		SURROGATE	MOLINATE	ATRAZINE	METRIBUZIN	ALACHLOR	METOLACHLOR	NORFLURAZON
FIELD FORTIFIED	SAMPLES							
	MISS#10	67	112	73	71	77	79	84
	MISS #11	73	111	80	74	80	81	87
	MISS#12	80	108	77	88	74	80	84
	MISS#13	70	88	98.	95	94	95	105
	MISS #14	63	107	79	73	78	79	85
	MISS#15	65	69	80	69	75	77	81
		*			91			
NON-FORTIFIED	SAMPLES							
	MISS#10	80						
	MISS #11	77						
	MISS#12	84						
	MISS#13	53						
	MISS #14	77						
	MISS#15	67						
LAB BLANKS								
	P881	99						
	P882	78						
			CONCENTRA	ATIONS FOR I	AB BLANKS			
	P881			0			•	
	P882		0	0	0	0	0	0
	F002		Ü			O	Ü	Ü
			PEAK AREAS	S FOR A 2X*	STANDARD			
	2X STANDARD		MOLINATE 98519	ATRAZINE 17004	METRIBUZIN 7646	ALACHLOR 8012	METOLACHLOR 25403	NORFLURAZON 23966
	ZX STANUANU		30013	17004	7040	8012	20403	23500
			DUPLICA	TE ANALYSIS				
		9	FIELD DUPLICA	TE - SURROGA	TE AREA COMPA	RISON		
		P833		P835		%RSD		
		205925		148891		32.28		
			MACHINE DUP	LICATE - SURRO	GATE AREA CO	MPARISON		

none

A29

^{*}ANALYTE CONCENTRATIONS ARE ABOUT 2 TIMES THE EPA ESTIMATED DETECTION LIMIT

PERCENT RECOVERIES

			SURROGATE	2,4-D	BENTAZON	ACIFLUROFEN
FIELD FORTIFIED S	AAADI EC					
FIELD FOR TIFIED 3	MISS#10		51	48	36	43
	MISS #11		94	75	75	85
	MISS#12		74	58	57	62
	MISS#13		85	69	75	80
	MISS #14		92	71	77	88
			101	69	79	81
	MISS#15		101	69	/9	81
NON-FORTIFIED SA	AMPLES		×.			
	MISS#10		omitted			
	MISS #11		105			
	MISS#12		84			16
	MISS#13		95			
	MISS #14		83			
	MISS#15		84		1	
						9.
LAB BLANKS			1922/12			
	P878		80			
	P879		81			
			CONCENTRAT	IONS FOR	LAB BLANKS	
	P878			0	0	o
	P879			o	ō	ō
			15			
			PEAK AREAS F	FOR A 2X	* STANDARD	
			SURROGATE	2.4-D	BENTAZON	ACIFLUROFEN
	2X STANDARD		64868	70727	26088	130838
					-	
			DUPLICATE	ΔΝΔΙ ΥΝ	S	
			DO. CIONIE	,		
			FIELD DUPLICATE		ATE AREA COMPA	
		P820		P829		%RSD
		63465		87714		6.48

MACHINE DUPLICATE - SURROGATE AREA COMPARISON

none

ANALYTE CONCENTRATIONS ARE ABOUT 2 TIMES THE EPA ESTIMATED DETECTION LIMIT

NPS METHOD 4

			CYANAZINE	FLUOMETURON	DIURON	LINURON	SURROGATE
FIELD FORTIFIED S	SAMPLES		1.0				
	MISS#10		- 75	100	83	72	91
	MISS #11		90	73	77	66	59
	MISS#12		117	75	93	82	77
	MISS#13		118	71	68	72	85
	MISS #14		108	79	92	81	76
	MISS#15		121	81	61	89	83
NON-FORTIFIED SA	AMPLES						
	MISS#10						91
	MISS #11						72
	MISS#12						80
	MISS#13						68
	MISS #14						77
	MISS#15						94
LAB BLANKS							
	P875						70
	P876						77
			CONCENTRA	TIONS FOR LAB	BLANKS		
			001102111111				
	P875		0	0	0	0	
	P876		0	0	0	0	
			PEAK AREAS	FOR A 2X* STA	NDARD		
			CYANAZINE	FLUOMETURON	DIURON	LINURON	
			4840	4504	5403	15995	
			4040	4004	0403	10880	
			DUPLICATE	E ANALYSIS			
			FIELD DUPLICAT	E - SURROGATE AR	EA COMPARI	SON	
		P814	and the	P815		%RSD	
		387340		389431		0.54	
			MACHINE DUPLI	CATE - SURROGATE	AREA COM	PARISON	
		1ST RUN		2ND RUN		%RSD	
		319460		325181		1.77	

^{*}ANALYTE CONCENTRATIONS ARE ABOUT 2 TIMES THE EPA ESTIMATED DETECTION LIMIT

NITRATE

SPIKE RECOVERY

WELL NUMBER

% RECOVERY

MISS 14

100%

DUPLICATE ANALYSIS

1ST MEASUREMENT

2ND MEASUREMENT

% RSD

6.49 mg/L

6.51mg/L

0.22%

RESULTS OF PESTICIDE MONITORING: TRIP #2 TO CRAIGHEAD COUNTY - SEPTEMBER, 1994

(unk = unknown, NC = not collected, ND = not detected)

(= suspect, see text)

	1	2	3	4	5	8
WELL ID:	CH #7	CH #8	CH #9	CH #10	CH #11	CH #12
DATE SAMPLED:	AUG. 31, 1994					
LATITUDE:	35° 53' 31"	35° 58' 31"	35° 57' 38"	35° 56' 36"	35° 43' 55°	35° 46' 19°
LONGITUDE:	90° 20' 30"	90° 20' 32°	90° 19' 08°	90° 27' 02°	90° 20' 24°	90° 17' 58°
DEPTH OF WELL, ft:	50	507	35	18	30	50-60
pH, standard units:	6.8	7.4	6.7	7.1	6.1	6.3
CONDUCTIVITY AT 25° C , umhos/cm:	385	476	492	534	202	445
TEMPERATURE, ° C:	21	17.5	16	18.5	17	17.5
NITRATE, mg/L:	7.70	<0.01	7.48	6.52	0.77	0.80
ACIFLUORFEN, ug/L	ND	ND	ND	ND	ND	ND
ALACHLOR, ug/L:	ND	ND	ND	ND	ND	ND
ATRAZINE,ug/L:	ND	ND	ND	ND	ND	ND
BENTAZON, ug/L	ND	ND	ND	ND	ND	ND
CYANAZINE, ug/L:	ND	ND	ND	ND	ND	ND
DIURON, ug/L:	ND	ND	ND	ND	ND	ND
FLUOMETURON, ug/L:	ND	ND	ND	ND	ND	ND
LINURON, ug/L:	ND -	ND	ND	ND	ND	ND
METOLACHLOR, ug/L:	ND	ND	ND	ND	ND	ND
METRIBUZIN, ug/L:	ND	ND	ND	ND	ND	ND
MOLINATE, ug/L:	ND	ND	ND	ND	ND	ND
NORFLURAZON, ug/L	ND	ND	ND	ND	ND	ND
2,4-D, ug/L	ND	ND	ND	ND	ND	ND

EPA METHOD 507

		SURROGATE	MOLINATE	ATRAZINE	METRIBUZIN	ALACHLOR	METOLACHLOR	NORFLURAZON	INT. STD.
FIELD FORTIFIED	SAMPLES								
	CH #7	110	87	68	74	88	93	65	75
	CH #8	104	78	69	70	78	81	80	во
	CH #9	131	126	102	108	111	109	104	78
	CH #10	127	115	113	113	120	120	112	89
	CH #11	106	112	108	107	113	117	103	112
	CH #12	111	111	111	110	115	116	107	94
NON-FORTIFIED	SAMPLES								
	CH #7	90							107
	CH #8	90							118
	CH #9	81							108
	CH #10	96							130
	CH #11	92							111
	CH #12	85							118
LAB BLANKS									
	P1170	122							103
	P1172	97							117
			CONCENTRA	ATIONS FOR	LAB BLANKS				
	P1170		0	0	0	0	0	0	
	P1172		0	0	0	0	0	0	
		PEAK AREAS FOR A 2X* STANDARD							
			MOLINATE	ATRAZINE	METRIBUZIN	ALACHLOR	METOLACHLOR	NORFLURAZON	
	2X STANDARD		3830	6331	3437	3694	10158	7800	
			DUPLICA	TE ANALYSIS	S				
			FIELD DUPLICA	ATE - SURROGA	TE AREA COMPA	RISON			
		P1203	VS.	P1205		%RSD			
		64523		70970		9.52			
			MACHINE DUP	LICATE - SURR	OGATE AREA COI	MPARISON			
		1ST RUN		2ND RUN	nacerous, t. A. att. Status att. att. A. de Tall	%RSD			
		89907		72962		20.81			
ANALYTE CON	CENTRATIONS AR		S THE EPA ESTI		TION LIMIT				
MINETIL	SELLINITIONS ON								

QUALITY CONTROL DATA FOR PESTICIDE MONITORING: TRIP #2 TO CRAIGHEAD COUNTY - SEPTEMBER, 1994.

EPA METHOD 515

PERCENT RECOVERIES

			SURROGATE	2,4-D	INT, STD.	BENTAZON	ACIFLUROFEN
FIELD FORTIFIED	SAMPLES						
	CH #7		87	81	93	89	76
	CH #8		78	73	82	87	66
	CH #9		86	87	87	100	78
	CH #10		70	75	78	75	65
	CH #11		84	93	84	95	86
	CH #12		92	81	86	77	68
NON-FORTIFIED	SAMPLES						
	CH #7		65		78		
	CH #8		78		77		
	CH #9		70		92		
	CH #10		71		76		
	CH #11	1.	60		79		
	CH #12		61		90		
FORTIFIED REAG	FNT WATER						
, otto near	P1180		92	68	79	69	62
	P1182		85	66	100	67	80
LAB BLANKS							
LAD BLANKS	P1179		74		67		
	P1181		68		69		
			CONCENTRAT	IONG FOR I	40 DI 41110	A.	
			CONCENTRAT	IONS FOR L	AR BLANKS		
	P1179			0		0	0
	P1181			0		0	o
							2

PEAK AREAS FOR A 2X* STANDARD

попв

DUPLICATE ANALYSIS

	FIELD DUPLICATE - SURROGATE AREA COM	MPARISON
P1220	P1229	%RSD
175961	151807	14.74

MACHINE DUPLICATE - SURROGATE AREA COMPARISON

1ST RUN 2ND RUN %RSD 177159 178699 0.87

^{*}ANALYTE CONCENTRATIONS ARE ABOUT 2 TIMES THE EPA ESTIMATED DETECTION LIMIT

QUALITY CONTROL DATA FOR PESTICIDE MONITORING: TRIP #2 TO CRAIGHEAD COUNTY - SEPTEMBER, 1994. QUALIT

NPS METHOD 4

PERCENT RECOVERIES

			CYANAZINE	FLUOMETURON	DIURON	LINURON	SURROGATE	INT. STD.
FIELD FORTIFIED	SAMPLES							
TILLD FORTIFIED	CH #7		69	74	80	65	77	99
	CH #8		76	70	84	70	82	95
	CH #9		70	67	82	70	78	107
	CH #10		77	74	90	78	93	89
	CH #11		75	79	84	70	79	99
	CH #12		75	59	87	65	71	97
NON-FORTIFIED S	SAMPLES							
	CH #7						71	98
	CH #8						71	104
	CH #9						71	104
	CH #10						91	103
	CH #11						68	93
	CH #12						79	105
FORTIFIED REAGI	ENT WATER							
	P1175		74	68	88	75	71	110
LAB BLANKS								
	P1174						71	97
	P1178						72	105
			CONCENTRA	TIONS FOR LAB E	BLANKS			
	01174		0	0	0	0		
	P1174 P1176		0	0	0	0		
	P1170		U	Ü	Ü	•		
			PEAK AREAS	FOR A 2X* STA	NDARD			
			CYANAZINE 667	FLUOMETURON 2732	DIURON 8126	LINURON 14123		
			DUPLICAT	E ANALYSIS				
			DOI LICKT	LAMALION				
			FIELD DUPLICA	TE - SURROGATE AR	EA COMPARI			
		P1214		P1215		%RSD		
		296357		290816		1.96		
			MACHINE DUD	ICATE PURDOCATE	ADEA COM	DADISON		
		1ST RUN	WIACHINE DUPL	ICATE - SURROGATE 2ND RUN	AREA COM	%RSD		
		408853		418761		2.93		
		,00000						

*ANALYTE CONCENTRATIONS ARE ABOUT 2 TIMES THE EPA ESTIMATED DETECTION LIMIT

CONTROL DATA FOR PESTICIDE MONITORING: TRIP #2 TO CRAIGHEAD COUNTY - SEPTEMBER, 1994.

NITRATE

SPIKE RECOVERY

WELL NUMBER

% RECOVERY

CH #8

106%

DUPLICATE ANALYSIS

1ST MEASUREMENT

2ND MEASUREMENT

% RSD

7.70mg/L

7.76 mg/L

0.84%

RESULTS OF PESTICIDE MONITORING: TRIP #2 TO POINSETT COUNTY - OCTOBER, 1994.

(unk = unknown, NC = not collected, ND = not detected)

= suspect, see tex

	1	2	3	4	5
WELL ID:	POIN #6	POIN #7	POIN #9	POIN #10	POIN #11
DATE SAMPLED:	Oct. 19, 1994	OCT 19, 1994	19-Oct-94	OCT 19, 1994	19-Oct-94
LATITUDE:	35° 41' 10°	35° 40' 21°	35° 34' 10"	35° 32' 54"	35° 29' 10°
LONGITUDE:	90° 22' 34"	90° 20' 29"	90° 20' 01"	90° 21' 39"	90° 18' 01"
DEPTH OF WELL, ft:	30	60	20	50	65
pH, standard units:	7.4	7.1	6.8	7.1	7
CONDUCTIVITY AT 25° C , umhos/cm:	725	453	604	830	534
TEMPERATURE, ° C :	18	18	18	18	17.5
NITRATE, mg/L:	0.01	0.04	0.13	0.07	0.04
ACIFLUORFEN, ug/L	ND	ND	ND	ND	ND
ALACHLOR, ug/L:	ND	ND	ND	ND	ND
ATRAZINE,ug/L:	ND	ND	ND	ND	ND
BENTAZON, ug/L	ND	ND	ND	ND	ND
CYANAZINE, ug/L:	ND	ND	ND	ND	ND
DIURON, ug/L:	ND	ND	ND	ND	ND
FLUOMETURON, ug/L:	ND	ND	ND	ND	ND
LINURON, ug/L:	ND	ND	ND	ND	ND
METOLACHLOR, ug/L:	ND	ND	ND	ND	ND
METRIBUZIN, ug/L:	ND	ND	ND	ND	ND
MOLINATE, ug/L:	ND	ND	ND	ND	ND
NORFLURAZON, ug/L	ND	ND	ND	ND	ND
2,4-D, ug/L	ND	ND	ND	ND	ND

QUALITY CONTROL DATA FOR PESTICIDE MONITORING: TRIP #2 TO POINSETT COUNTY - OCTOBER, 1994.

EPA METHOD 507

PERCENT RECOVERIES

		SURROGATE	MOLINATE	ATRAZINE	METRIBUZIN	ALACHLOR	METOLACHLOR	NORFLURAZON	INT. STD.
FIELD FORTIFIED	SAMPLES								
TIELD TOTTITIES	POIN #6	163	124	100	104	113	113	111	80
	POIN #7	112	92	86	71	82	92	95	85
	POIN #9	122	90	78	74	84	99	101	86
	POIN #10	116	91	96	90	84	94	98	95
	POIN #11	98	108	100	90	90	92	95	101
NON-FORTIFIED	SAMPLES								
	POIN #6	89							100
	POIN #7	56							111
	POIN #9	95							115
	POIN #10	87							116
	POIN #11	84							114
FORTIFIED REAG	SENT WATER P1468	72	73	77	61	68	86	92	95
	F1408	12	/3	//	01	08	80	92	85
LAB BLANKS									
	P1467	87							94
	P1469	79 -							125
			CONCENTRA	ATIONS FOR	LAB BLANKS				
	P1467		0	0	0	0	0	0	
	P1469		0	0	0	0	0	0	
	F1405		Ü	Ü	Ü	Ü	Ü	U	
			PEAK AREA	S FOR A 2X	* STANDARD				
			MOLINATE	ATRAZINE	METRIBUZIN	ALACHLOR	METOLACHLOR	NORFLURAZON	
	2X STANDARD		2899	5158	2689	2882	7870	6691	
			DUPLICA:	TE ANALYSIS	3				
			FIELD DUPLICA	TE - SURROGA	TE AREA COMPA	RISON			
		P1503		P1505		%RSD			
		65591		90162		31.55			
			MACHINE DUP	LICATE - SURR	OGATE AREA COM	MPARISON			
		1ST RUN		2ND RUN		%RSD			
		80659		72098		11.21			
*ANALYTE CON	CENTRATIONS ARI	E ABOUT 2 TIMES	S THE EPA ESTI	MATED DETECT	TION LIMIT				

^{*}ANALYTE CONCENTRATIONS ARE ABOUT 2 TIMES THE EPA ESTIMATED DETECTION LIMIT

QUALITY CONTROL DATA FOR PESTICIDE MONITORING: TRIP #2 TO POINSETT COUNTY - OCTOBER, 1994.

EPA METHOD 515

PERCENT RECOVERIES

		SURROGATE	2,4-D	INT. STD.	BENTAZON	ACIFLUROFEN
FIELD FORTIFIED	SAMPLES					
	POIN #6	94	111	73	91	118
	POIN #7	112	100	79	112	93
	POIN #9	123	108	81	61	103
	POIN #10	89	99	77	96	86
	POIN #11	91	133	78	77	135
NON-FORTIFIED S	SAMPLES					
	POIN #6	90		71		
	POIN #7	67		72		
	POIN #9	76		68		
	POIN #10	98		73		
	POIN #11	66		73		
FORTIFIED REAG	ENT WATER					
	P1463	78	72	88	62	61
LAB BLANKS						
	P1466	65		74		
		CONCENTRAT	TIONS FOR	LAB BLANKS		
	P1466		o		o	0

PEAK AREAS FOR A 2X* STANDARD

none

DUPLICATE ANALYSIS

	FIELD DUPLICATE - SURROGATE AREA C	OMPARISON
P1530	P1539	%RSD
271382	145341	60.49
	MACHINE DUPLICATE - SURROGATE ARE	A COMPARISON
1ST RUN	2ND RUN	%RSD
192766	200758	4.06

^{*}ANALYTE CONCENTRATIONS ARE ABOUT 2 TIMES THE EPA ESTIMATED DETECTION LIMIT

NPS METHOD 4

PERCENT RECOVERIES

			CYANAZINE	FLUOMETURON	D!URON	LINURON	SUP	ROGATE	INT. STD.
FIELD FORTIFIED	SAMPLES								
	POIN #6		80	87	85	90		88	124
	POIN #7		79	87	84	91		84	125
	POIN #9		88	97	96	101		106	117
	POIN #10		84	92	90	93		88	119
	POIN #11		82	92	89	93		99	119
NON-FORTIFIED S	CAMADI ES								
MON-FORTIFIED S	POIN #6							96	113
	POIN #7							92	109
	POIN #9							80	141
	POIN #10							105	104
	POIN #11							73	157
	FOR4 # 11							,,	107
FORTIFIED REAGI				72 -	22				
	P1470		82	87	90	94		86	128
LAB BLANKS									
	P1471							91	123
	*								
			CONCENTRA	TIONS FOR LAB E	BLANKS				
	P1471		0	o	o	o			
			PEAK AREAS	FOR A 2X* STA	NDARD				
			CYANAZINE	FLUOMETURON	DIURON	LINURON			
	2X STANDARD		5617	3239	7137	15782			
			DUPLICAT	E ANALYSIS					
			FIELD DUPLICA	TE - SURROGATE AR	EA COMPARI				
		P1514		P1515		%RSD			
		122510		141898		14,67	in-		
			MACHINE DUPL	ICATE - SURROGATE	AREA COM				
		1ST RUN		2ND RUN		%RSD			
		459833		459551		0.08			

^{*}ANALYTE CONCENTRATIONS ARE ABOUT 2 TIMES THE EPA ESTIMATED DETECTION LIMIT

CONTROL DATA FOR PESTICIDE MONITORING: TRIP #2 TO POINSETT COUNTY - OCTOBER, 1994.

NITRATE

SPIKE RECOVERY

WELL NUMBER

% RECOVERY

POIN 7

98%

DUPLICATE ANALYSIS

1ST MEASUREMENT

2ND MEASUREMENT

% RSD

0.014 mg/L

0.011mg/L

24.00%

APPENDIX B

Tables B1 - B3 explain the symbols used to identify each soil. Table B4 lists the family, subgroup and order for each of the soil series. Table B5 lists the soil surrounding each well. Figures B1-B38 show the soils within one mile of each well as presented in the soil surveys for each county (Ferguson and Gray, 1971; Gray and Ferguson, 1977; Ferguson, 1979). On these maps, cross lines have been drawn to indicate permeable soils, those through which water movement is rapid or moderately rapid.

Table B1. Key to soil symbols used in Craighead County.

Symbol	Name
1	Amagon fine sandy loam
2	Amagon silt loam
3	Beulah fine sandy loam, 0 to 1 percent slopes
4	Beulah fine sandy loam, gently undulating
7	Bruno loarny sand
12	Commerce very fine sandy loam
14	Convent fine sandy loam
15	Dubbs fine sandy loam, 0 to 1 percent slopes
16	Dubbs fine sandy loam, gently undulating
18	Dundee fine sandy loam
19	Dundee silt loam
20	Dundee-Bruno-Commerce complex
22	Foley silt loam
23	Fountain silt loam
34	Mhoon fine sandy loam
35	Mhoon soils, frequently flooded
36	Roellen silty clay loam
37	Sharkey clay

Table B2. Key to soil symbols used in Mississippi County.

Symbol	Name
Aa	Alligator clay
An	Amagon sandy loam
Вр	Borrow pits
Br	Bowdre silty clay loam
Bv	Bruno-Crevasse complex
Cm	Commerce silt loam
Cn	Convent fine sandy loam
Cr	Crevasse loarny sand
Du	Dundee silt loam
Dv	Dundee-Dubbs-Crevasse complex
Ec	Earle-clay
Fe	Forestdale silt loam
Fo	Forestdale silty clay loam
Fr	Forestdale-Routon complex
Ha	Hayti fine sandy loam
lb	Iberia clay
Je	Jeanerette silt loam
Mo	Marganfield fine sandy loam
Rd	Routon-Dundee-Crevasse complex
Sc	Sharkey silty clay loam
Sh	Sharkey silty clay
Sk	Sharkey-Crevasse complex
Sm	Sharkey-Steele complex
Sn	Sharkey and Steele soils
So	Steele loamy sand
Sr	Steele silty clay loam
Ss	Steele and Crevasse soils
St	Steele and Tunica soils
Td	Tiptonville and Dubbs silt loams
Tu	Tunica silty clay

Table B3. Key to soil symbols used in Poinsett County.

Symbol	Name
Aa	Alligator clay
An	Arnagon silt loam
BeU	Beulah fine sandy loam, undulating
BoU	Bowdre silty clay loam, undulating
Cu	Convent sift loam
DbU	Dubbs silt loam, undulating
DdA	Dundee silt loam, 0 to 2 percent slopes
Ec	Earle sitty clay loam
Fo	Foley-Calhoun complex
Ha	Hayti solls
Mo	Mhoon silt loam
Sc	Sharkey clay
Sm	Sharkey-Steele complex
SN	Sharkey soils, frequently flooded
TnA	Tunica clay, 0 to 1 percent slopes
TnU	Tunica clay, undulating

Table B4. Classifications of soil series.

Series	Family	Subgroup	Order
Alliantor	Very-fine, montmortilonitic, add, thermic	New Joseph Landson, and a	l-aNasla
Alligator	Very-fine, montmorillonitic, add, thermic	Vertic Haplaquepts	Inceptisols Alfisols
Amagon Beulah	EXTENSION FOR THE PART OF THE PROPERTY OF THE PART OF	Typic Ochraqualis	
Bowdre	Coarse-loamy, mixed, thermic	Typic Dystrochrepts	Inceptisols Moliisols
	Clayey over loamy, mixed, thermic	Fluvaquentic Hapludolls	
Bruno	Sandy, mixed, thermic	Typic Udifluvents	Entisols
Calhoun	Fine-silty, mixed, thermic	Typic Glossudalfs	Alfisols
Commerce	Fine-silty, mixed, nonacid, thermic	Aeric Fluvaquents	Entisols
Convent	Coarse-silty, mixed, nonacid, thermic	Aeric Fluvaquents	Entisols
Crevasse	Mixed, thermic	Typic Udipsamments	Entisols
Dubbs	Fine-silty, mixed, thermic	Typic Haptudalfs	Alfisols
Dundee	Fine-silty, mixed, thermic	Aeric Ochraqualis	Alfisois
Earle	Clayey over loamy, montmorttlonitic, add, thermic	Vertic Haplaquepts	Inceptisois
Foley	Fine-sitty, mixed, thermic	Albic Glossic Natraqualis	Alfisols
Forestdale	Fine, montmorillonitic, thermic	Typic Ochraquatis	Alfisols
Fountain	Fine-sitty, mixed, thermic	Typic Glossaqualfs	Alfisols
Hayti	Fine-sitty, mixed, nonacid, thermic	Typic Fluvaquents	Entisols
Iberia	Fine, montmorillonitic, noncalcareous, thermic	Vertic Haplaquolls	Mollisols
Jeanerette	Fine-silty, mixed, noncalcareous, thermic	Typic Argiaquolis	Mollisols
Mhoon	Fine-silty, mixed, nonacid, thermic	Typic Fluvaquents	Entisols
Morganfleld	Coarse-silty, mixed, nonacld, thermic	Typic Udifluvents	Entisols
Roellen	Fine, montmorillonitic, thermic	Vertic Haplaquolls	Mollisols
Routon	Fine-sitty, mixed, thermic	Typic Ochraqualis	Alfisols
Sharkey	Very-fine, montmortilonitic, nonacid, thermic	Vertic Haplaquepts	Inceptisols
Steele	Sandy over clayey, mixed, nonacid, thermic	Aquic Udifluvents	Entisols
Tiptonville	Fine-sitty, mixed, thermic	Typic Argludolis	Mollisols
Tunica	Clayey over loamy, montmorillonitic, nonacid, thermic	Vertic Haplaquepts	Inceptisols

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Table B5. Soil that immediately surrounds each well.

Well	Soil
C1	Dubbs fine sandy loam, gently undulating
C2	Fountain silt loam
C3	Mhoon fine sandy loam
C4	Sharkey clay
C5	Commerce very fine sandy loam
C6	Dundee fine sandy loam
C7	Dubbs fine sandy loam, 0 to 1 percent slopes
C8	Dundee fine sandy loam
C9	Dundee fine sandy loam
C10 C11	Dubbs fine sandy loam, gently undulating Convent fine sandy loam
C12	Commerce very fine sandy loam
M1	Dundee silt loam
M2	Routon-Dundee-Crevasse complex
МЗ	Routon-Dundee-Crevasse complex
M4	Sharkey-Steele complex
M5	Sharkey silty clay
M6	Bowdre silty clay loam
M7	Dundee silt loam
M8	Tunica silty clay
M9	Tunica sitty clay
M10	Alligator clay
M11	Steele and Crevasse soils
M12	Routon-Dundee-Crevasse complex
M13	Routon-Dundee-Crevasse complex
M14	Routon-Dundee-Crevasse complex
M15	Amagon sandy loam
P1	Sharkey clay
P2	Sharkey clay
P3	Dundee silt loam, 0 to 2 percent slopes
P4	Dundee silt loam, 0 to 2 percent slopes
P5	Mhoon silt loam
P6	Hayti soils
P7	Hayti soils
P8	Hayti soils
P9	Sharkey-Steele complex
P10	Sharkey-Steele complex
P11	Sharkey clay

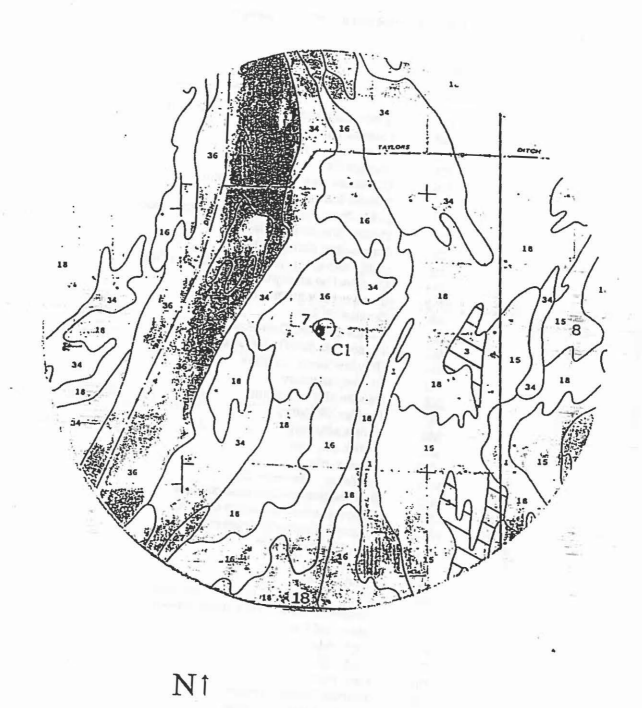
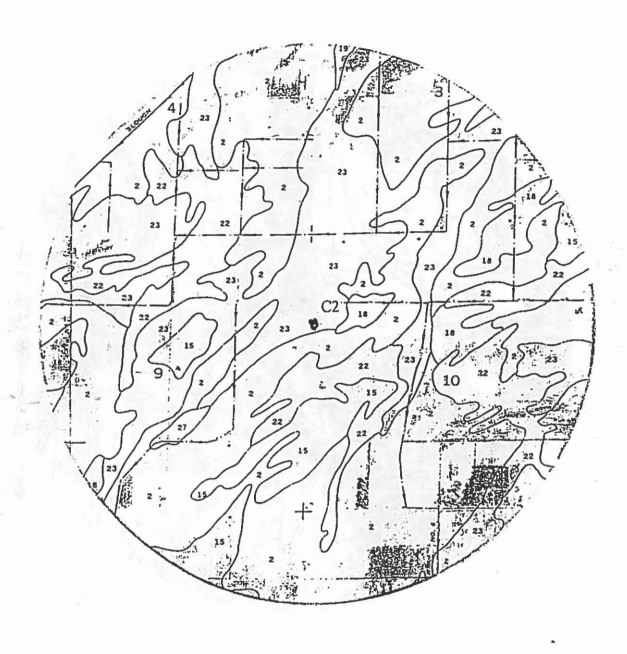


Figure B1. Soils within one mile of well C1. Permeable soils are indicated by cross lines.



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Figure B2. Soils within one mile of well C2. Permeable soils are indicated by cross lines.

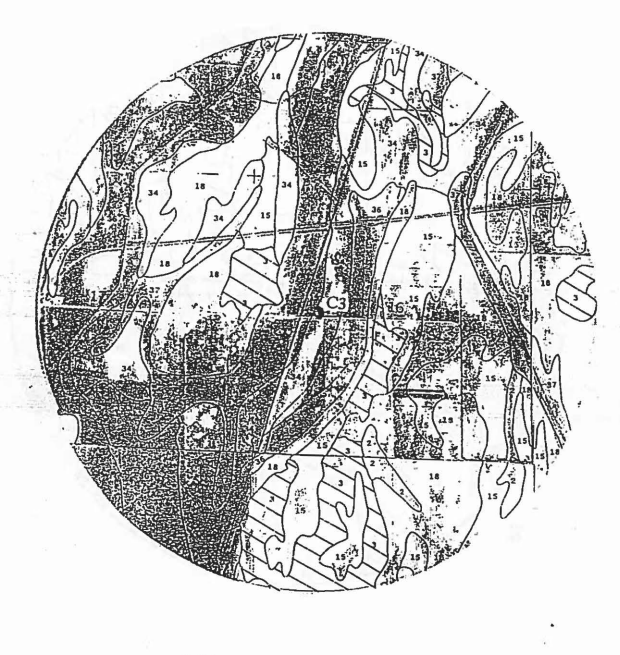


Figure B3. Soils within one mile of well C3. Permeable soils are indicated by cross lines.

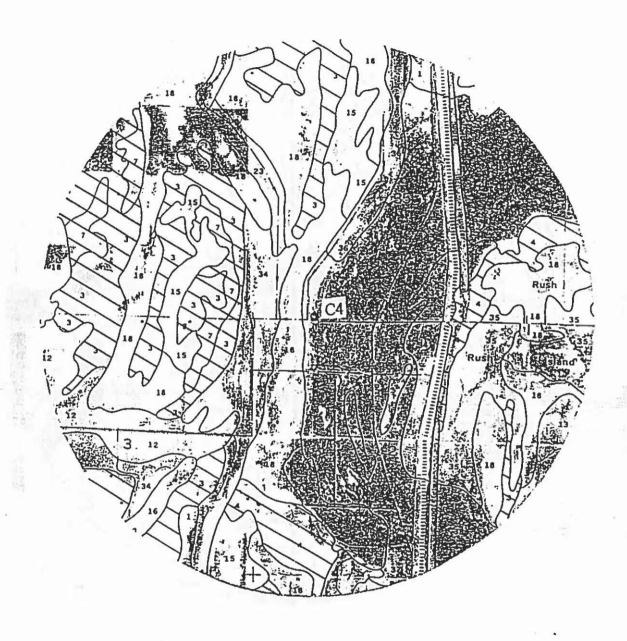
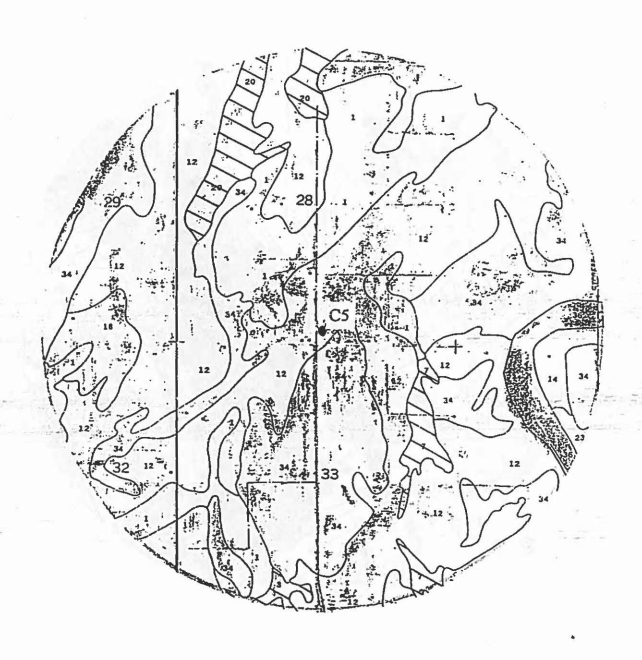


Figure B4. Soils within one mile of well C4. Permeable soils are indicated by cross lines.



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Figure B5. Soils within one mile of well C5. Permeable soils are indicated by cross lines.

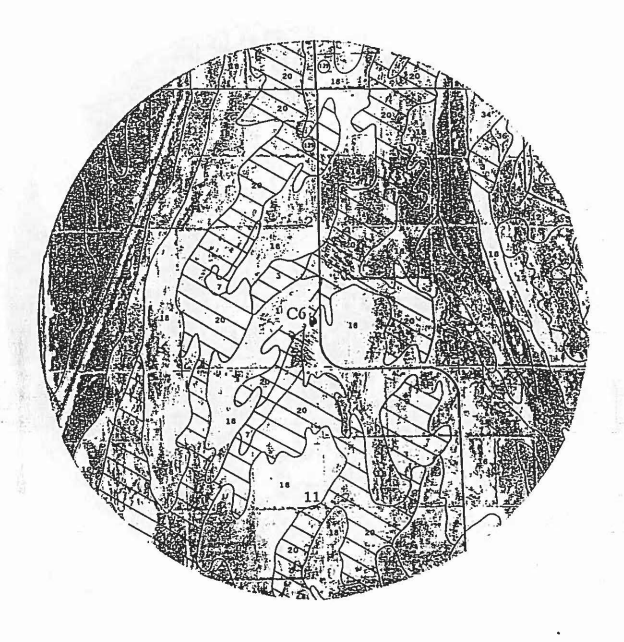


Figure B6. Soils within one mile of well C6. Permeable soils are indicated by cross lines.

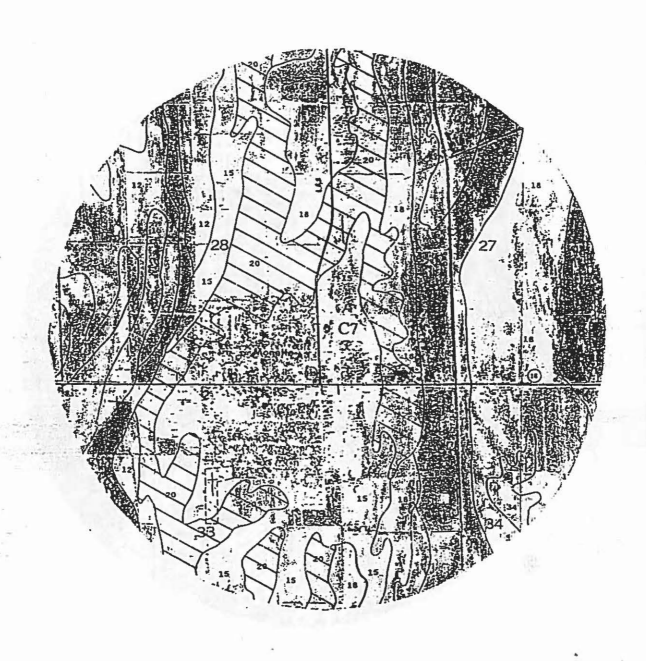


Figure B7. Soils within one mile of well C7. Permeable soils are indicated by cross lines.

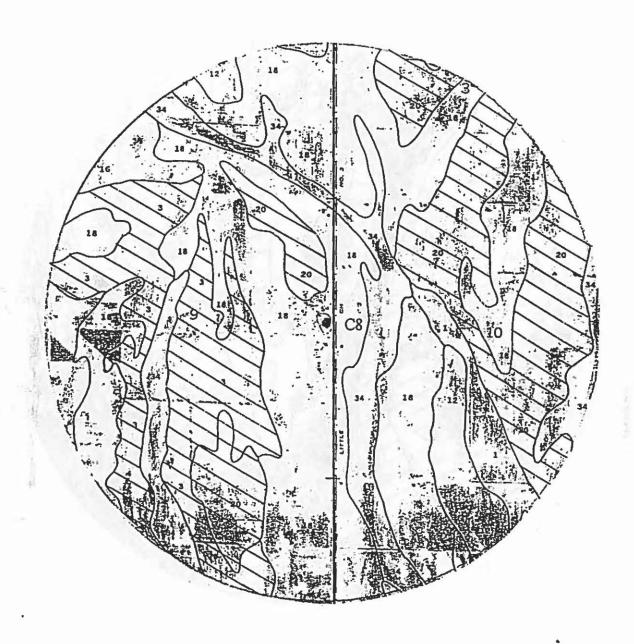


Figure B8. Soils within one mile of well C8. Permeable soils are indicated by cross lines.

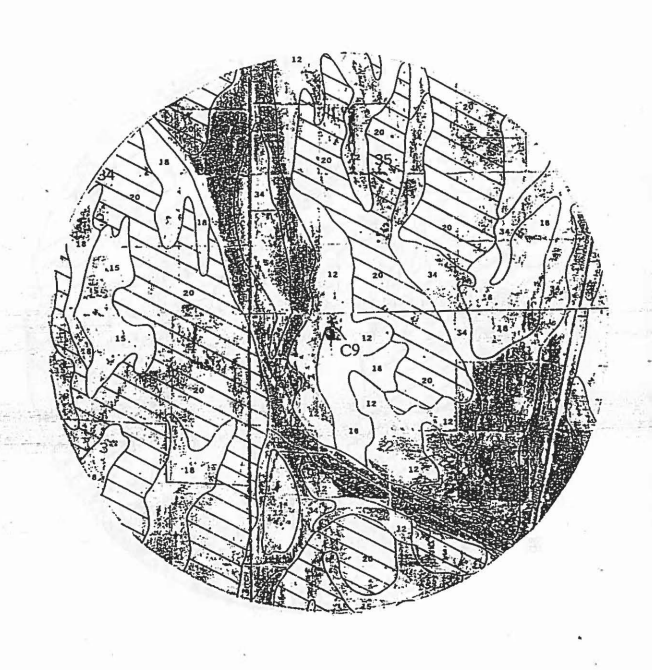


Figure B9. Soils within one mile of well C9. Permeable soils are indicated by cross lines.

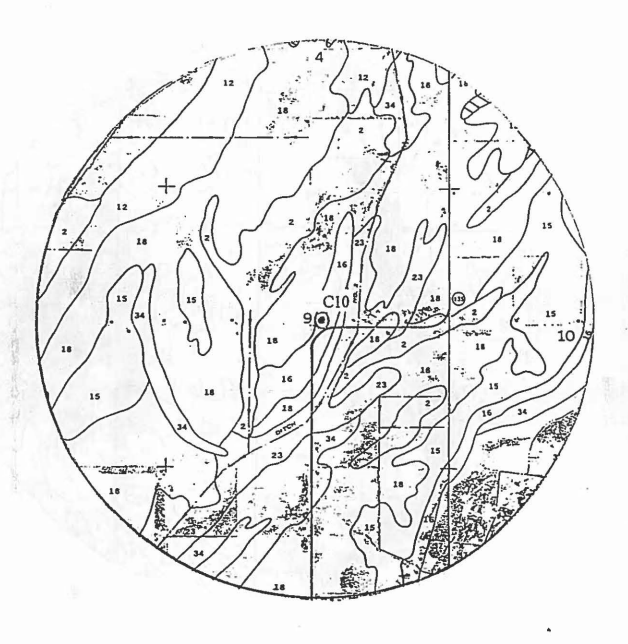


Figure B10. Soils within one mile of well C10. Permeable soils are indicated by cross lines.

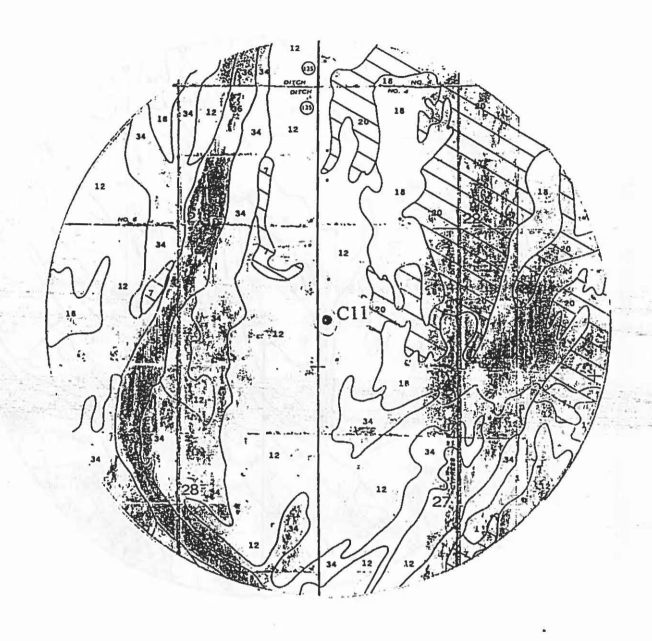


Figure B11. Soils within one mile of well C11. Permeable soils are indicated by cross lines.

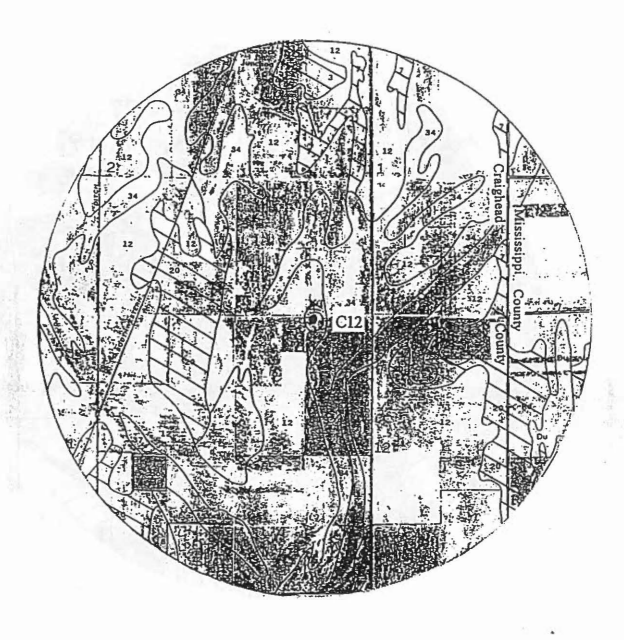


Figure B12. Soils within one mile of well C12. Permeable soils are indicated by cross lines.

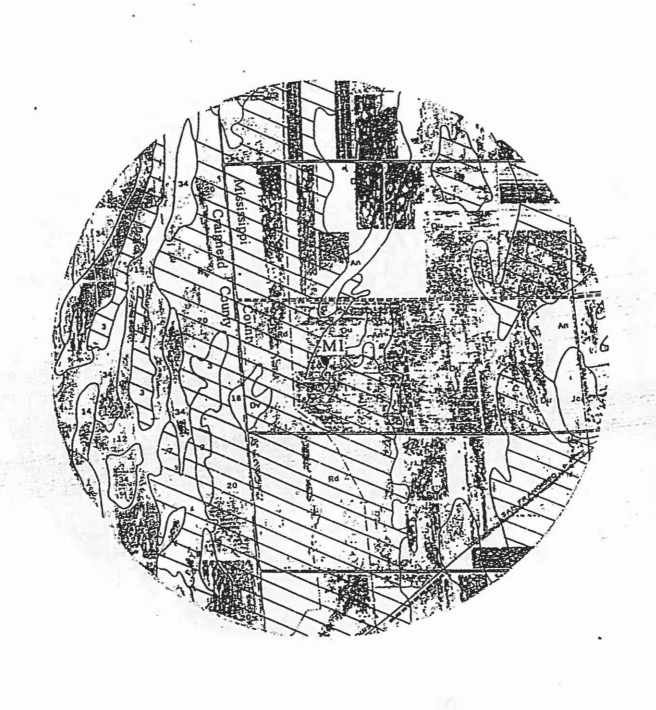
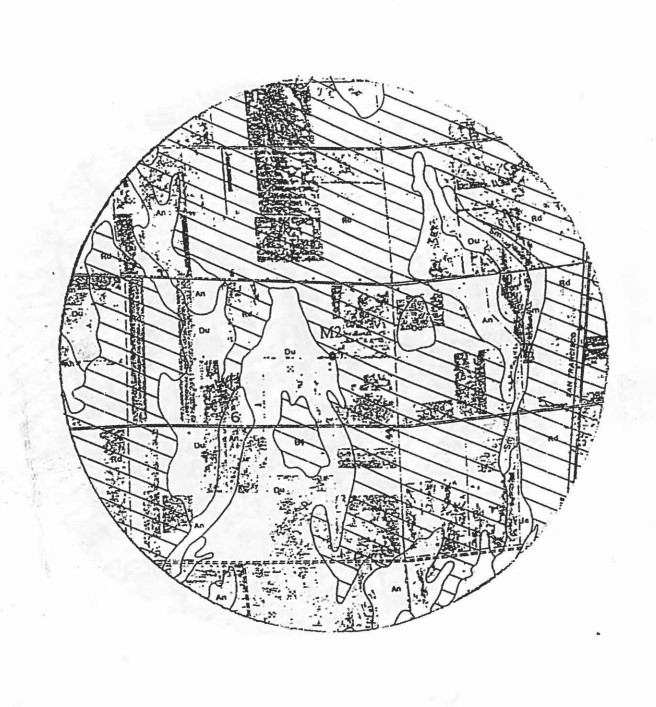
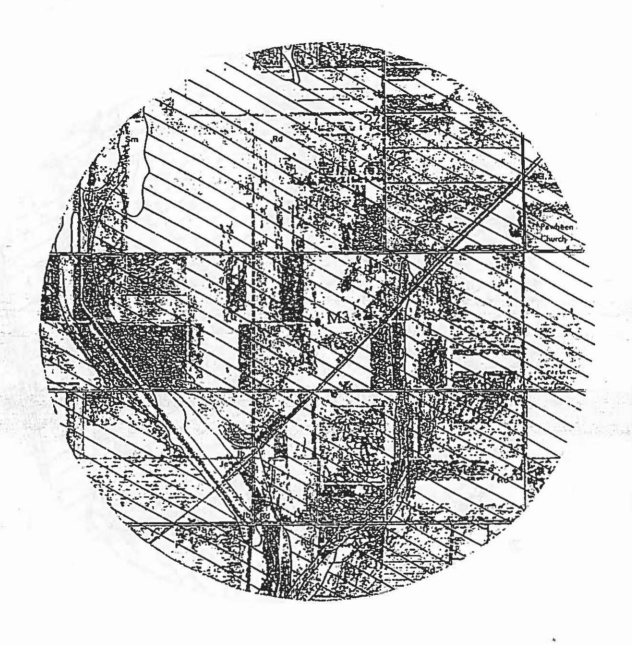


Figure B13. Soils within one mile of well M1. Permeable soils are indicated by cross lines.



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Figure B14. Soils within one mile of well M2. Permeable soils are indicated by cross lines.



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Figure B15. Soils within one mile of well M3. Permeable soils are indicated by cross lines.

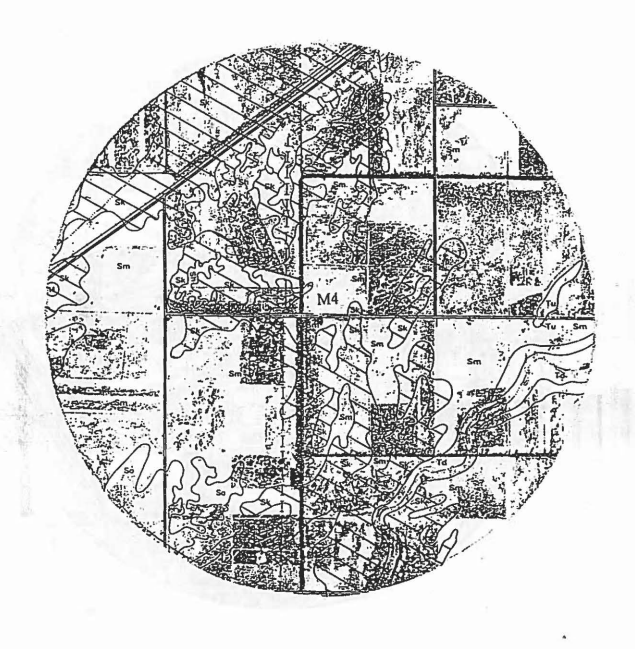


Figure B16. Soils within one mile of well M4. Permeable soils are indicated by cross lines.

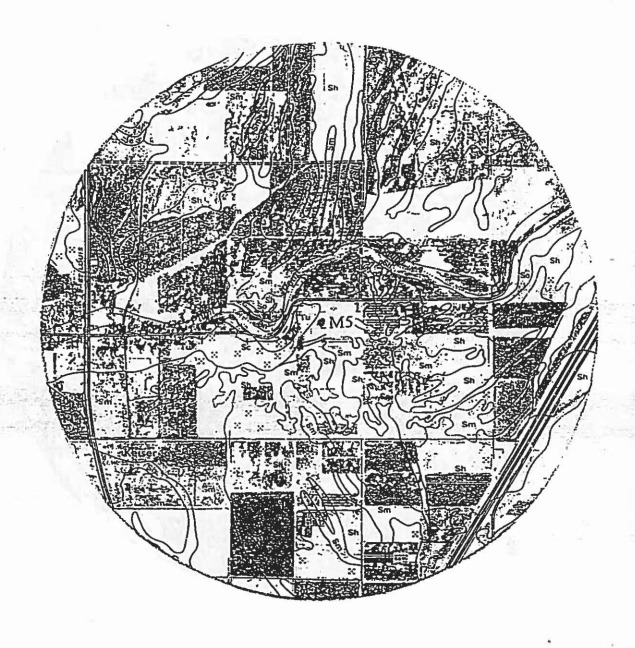


Figure B17. Soils within one mile of well M5. Permeable soils are indicated by cross lines.

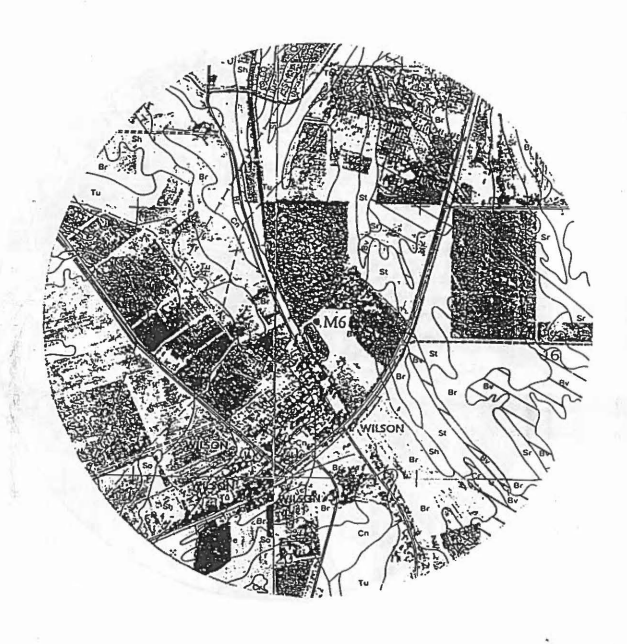


Figure B18. Soils within one mile of well M6. Permeable soils are indicated by cross lines.

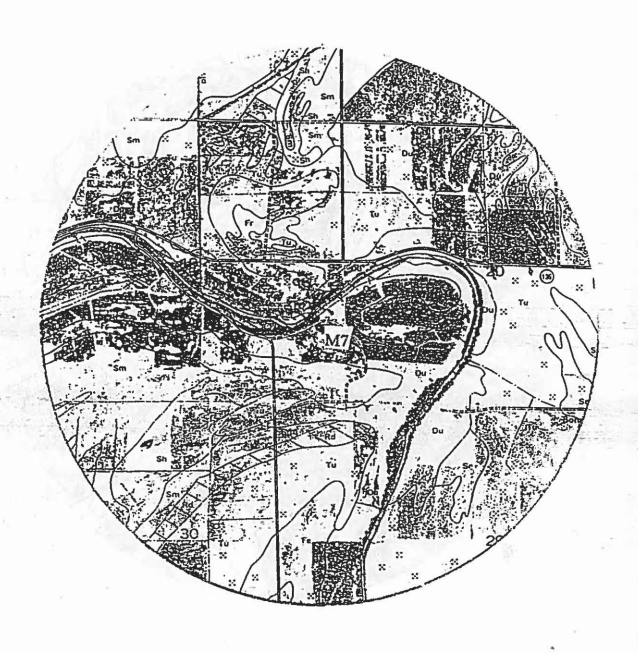
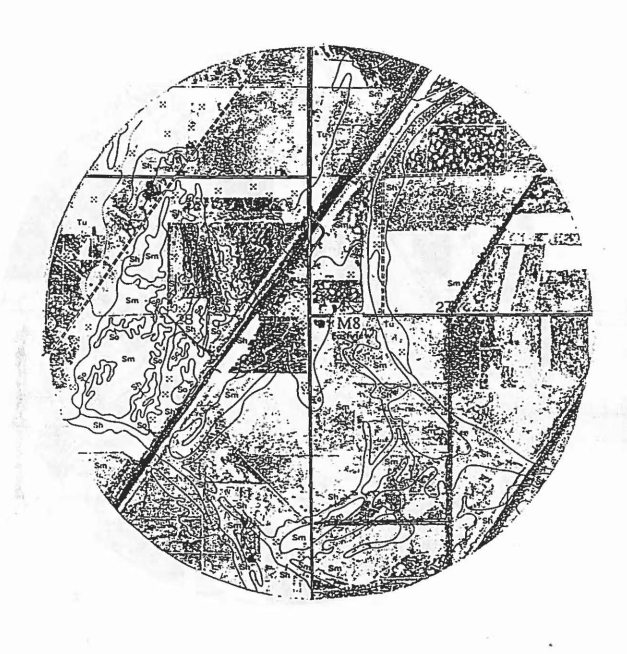


Figure B19. Soils within one mile of well M7. Permeable soils are indicated by cross lines.



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Figure B20. Soils within one mile of well M8. Permeable soils are indicated by cross lines.

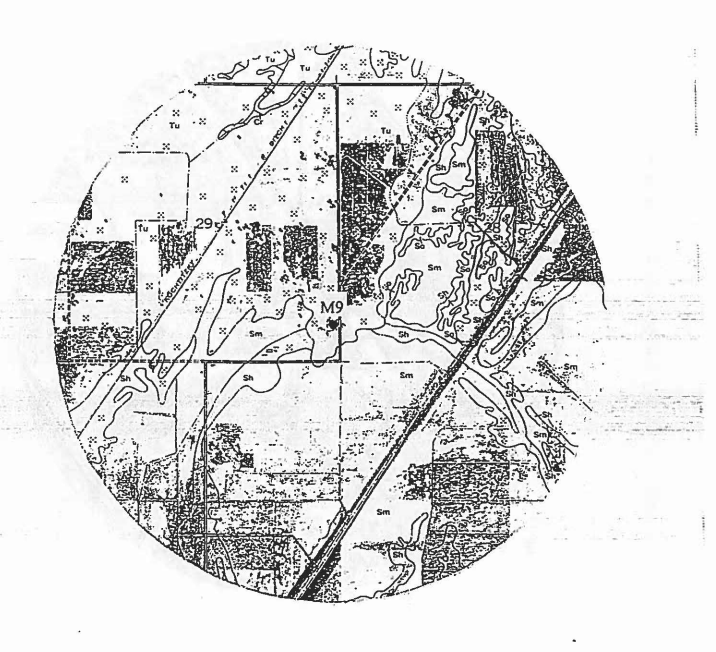


Figure B21. Soils within one mile of well M9. Permeable soils are indicated by cross lines.

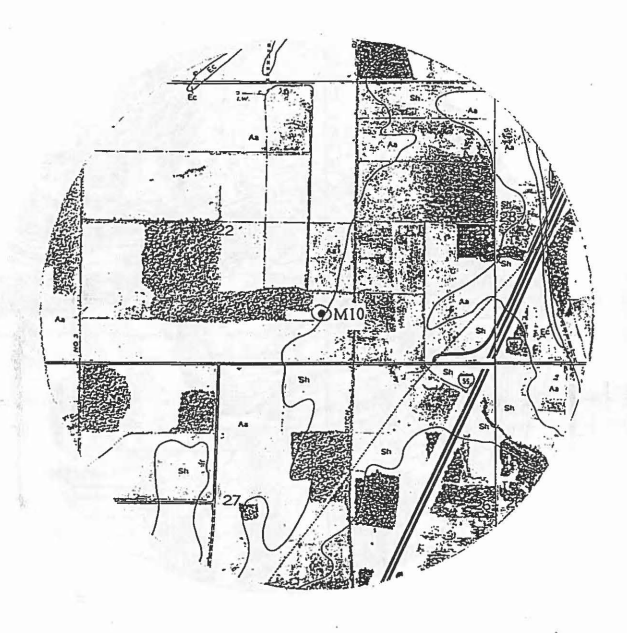


Figure B22. Soils within one mile of well M10. Permeable soils are indicated by cross lines.

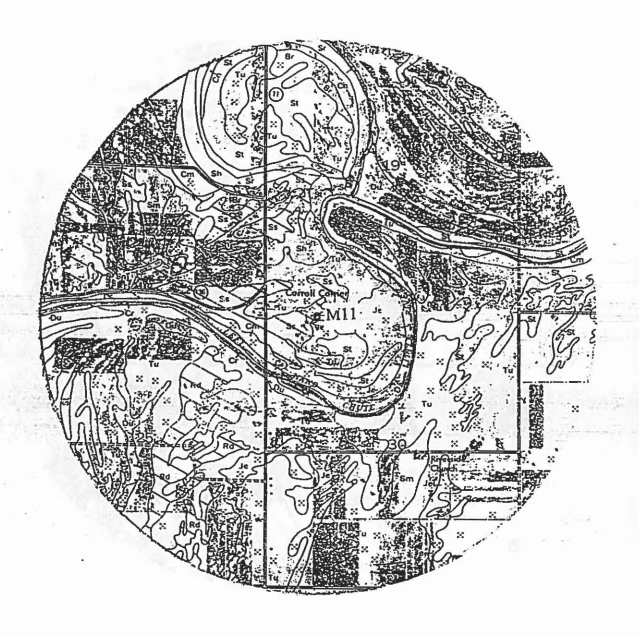


Figure B23. Soils within one mile of well M11. Permeable soils are indicated by cross lines.

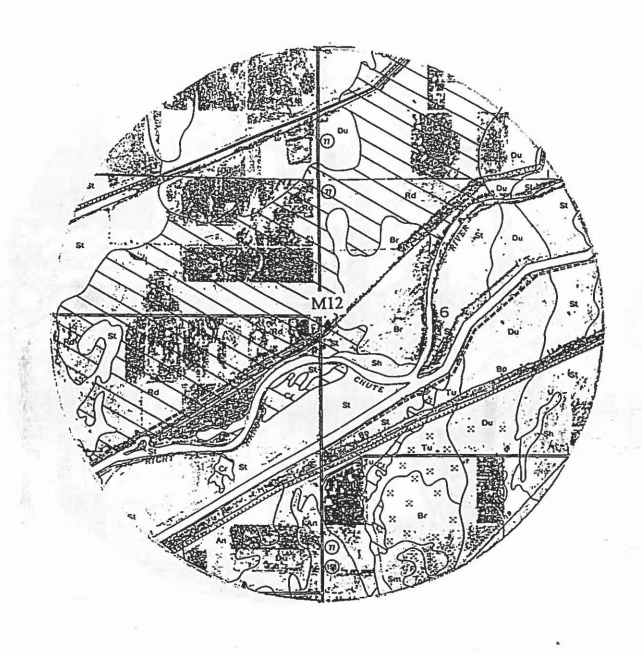


Figure B24. Soils within one mile of well M12. Permeable soils are indicated by cross lines.

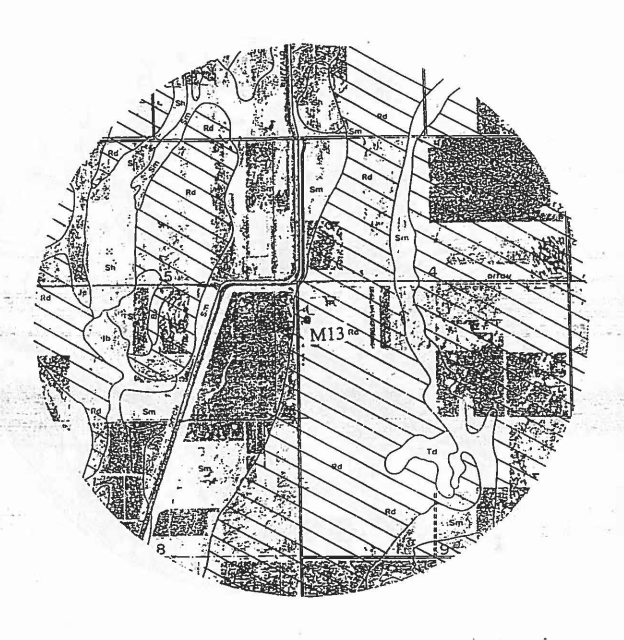


Figure B25. Soils within one mile of well M13. Permeable soils are indicated by cross lines.

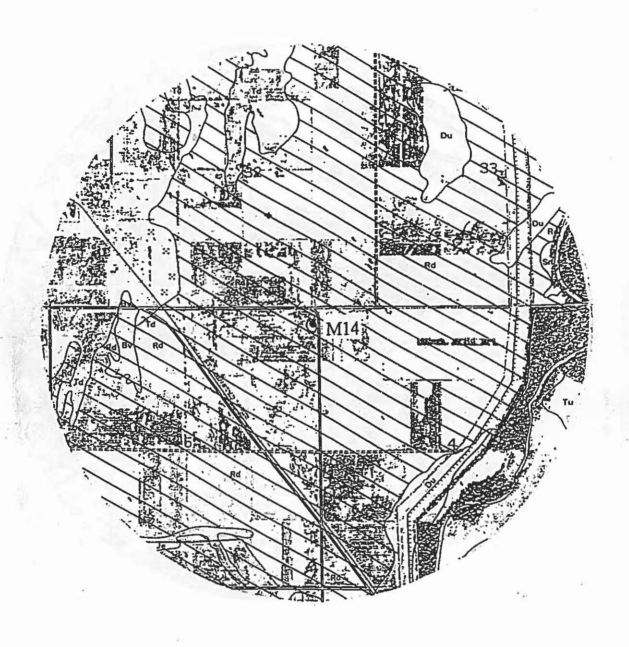


Figure B26. Soils within one mile of well M14. Permeable soils are indicated by cross lines.

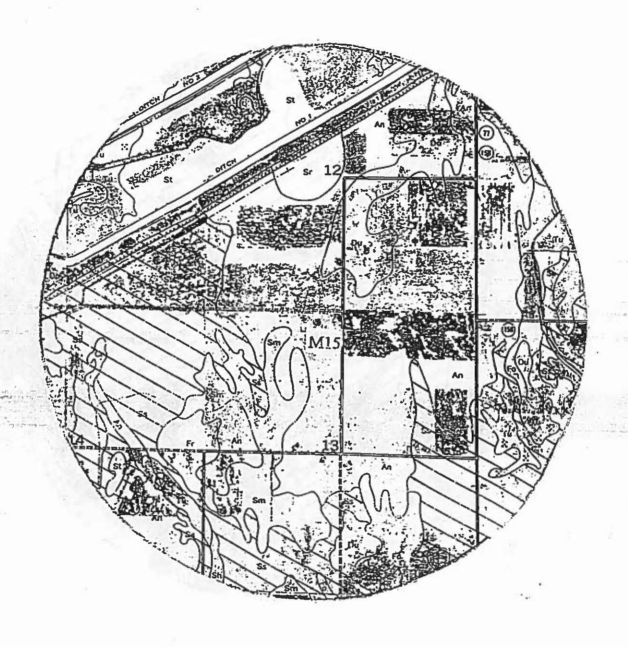


Figure B27. Soils within one mile of well M15. Permeable soils are indicated by cross lines.

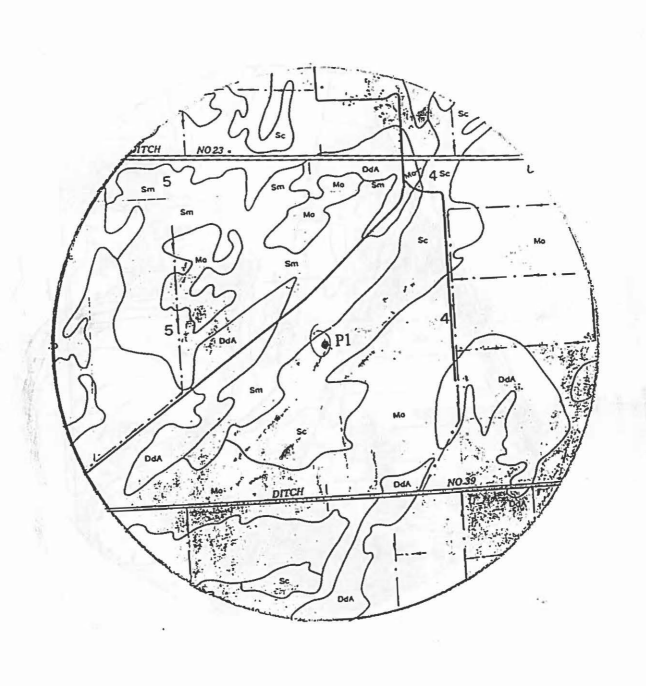


Figure B28. Soils within one mile of well P1. Permeable soils are indicated by cross lines.



Figure B29. Soils within one mile of well P2. Permeable soils are indicated by cross lines.

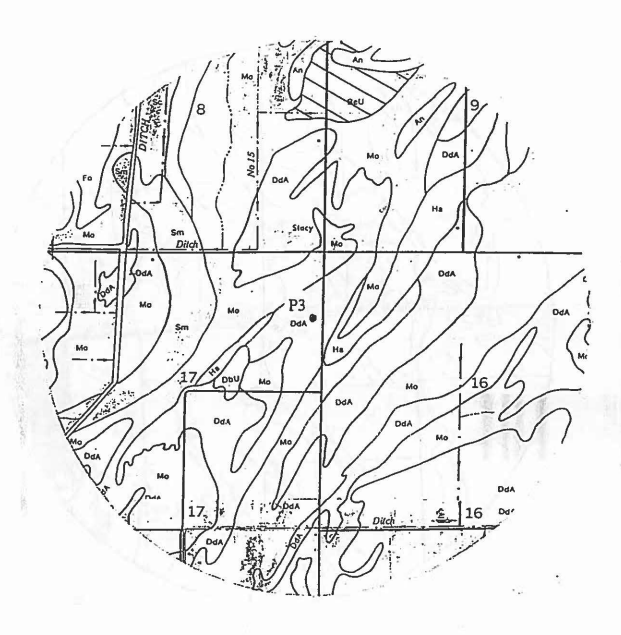


Figure B30. Soils within one mile of well P3. Permeable soils are indicated by cross lines.

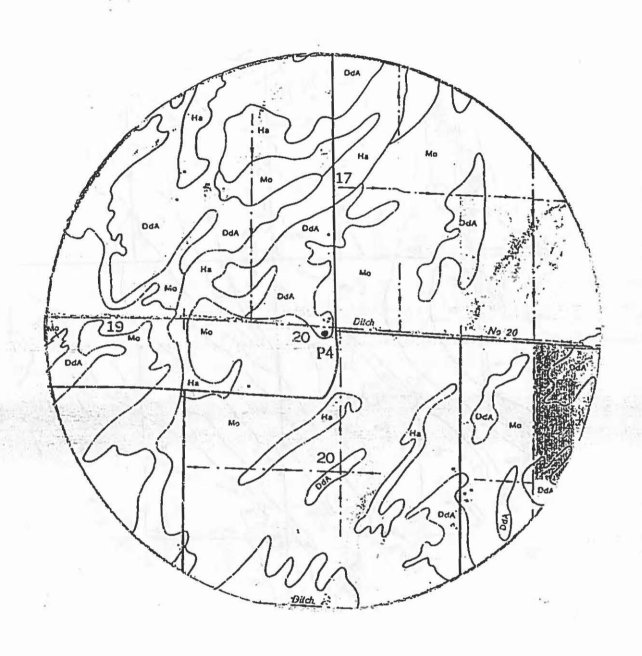


Figure B31. Soils within one mile of well P4. Permeable soils are indicated by cross lines.

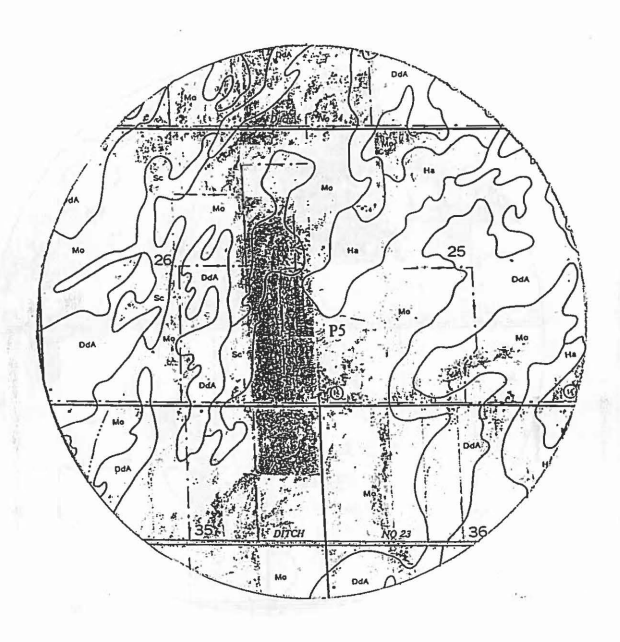
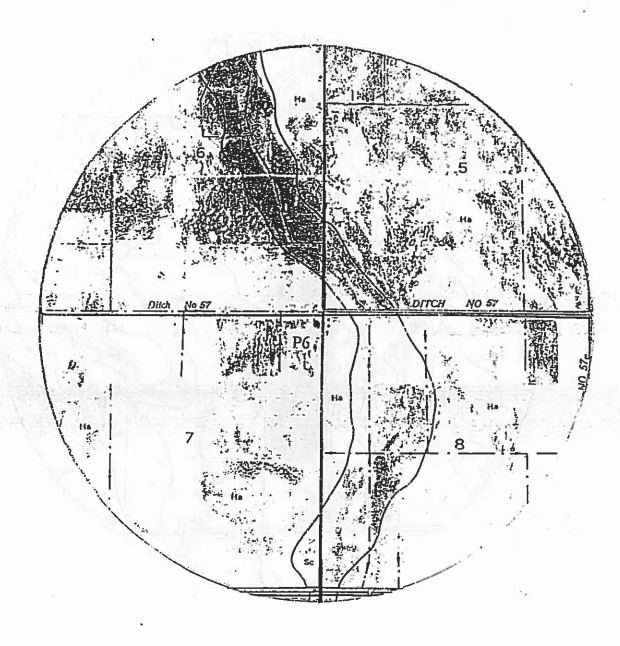


Figure B32. Soils within one mile of well P5. Permeable soils are indicated by cross lines.



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Figure B33. Soils within one mile of well P6. Permeable soils are indicated by cross lines.

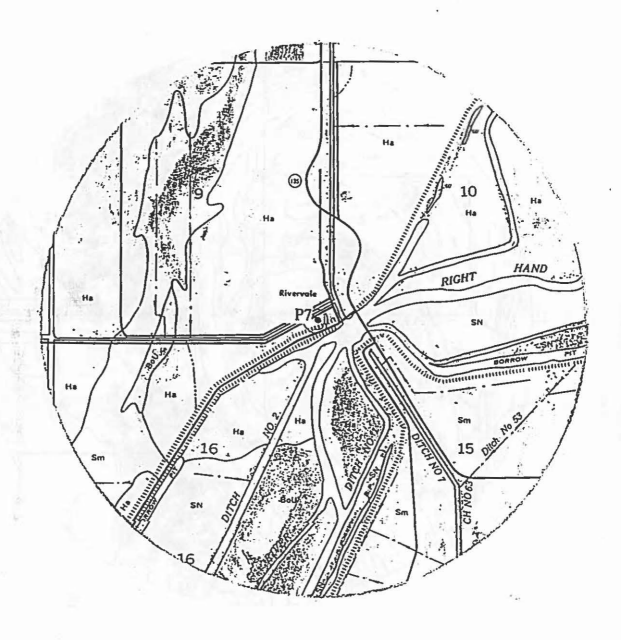


Figure B34. Soils within one mile of well P7. Permeable soils are indicated by cross lines.



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Figure B35. Soils within one mile of well P8. Permeable soils are indicated by cross lines.

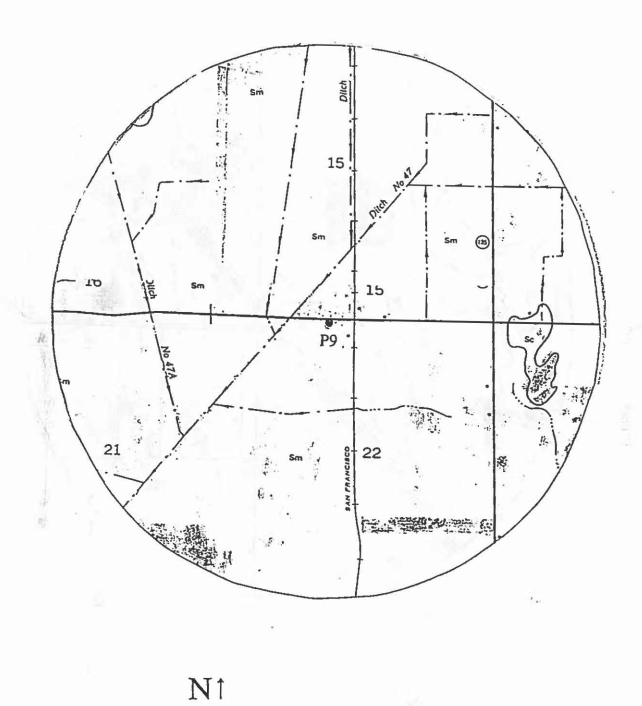
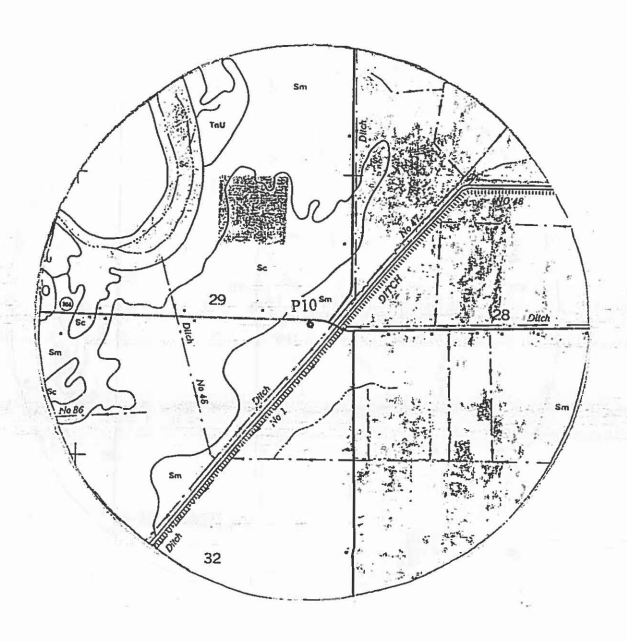


Figure B36. Soils within one mile of well P9. Permeable soils are indicated by cross lines.



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Figure B37. Soils within one mile of well P10. Permeable soils are indicated by cross lines.



Figure B38. Soils within one mile of well P11. Permeable soils are indicated by cross lines.