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# **GROUNDWATER CHEMISTRY** AT THE NEVADA TEST SITE: RECEIVED **Data and Preliminary Interpretations** JUL 19 1993 OSTI

Prepared by

Jenny B. Chapman and Brad F. Lyles

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March 1993

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# **GROUNDWATER CHEMISTRY AT THE NEVADA TEST SITE: Data and Preliminary Interpretations**

by

Water Resources Center Desert Research Institute University and Community College System of Nevada

> prepared for DOE Nevada Operations Office U.S. Department of Energy Las Vegas, Nevada

> > Publication No. 45100

March 1993

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#### ABSTRACT

The interpretation of chemical analyses of groundwater collected at and near the Nevada Test Site (NTS) has been vital in developing conceptual models of groundwater flow in the area. These conceptual models are tested using recent chemical data generated by the Desert Research Institute, as well as historic analyses from the U.S. Geological Survey. A total of 81 wells are represented by analyses from 1957 to 1990, with generally excellent agreement between repeat samples from the same location. As identified by previous workers, three hydrochemical facies are represented by the samples: Ca-Mg-HCO3 water in carbonate rocks or alluvium derived from carbonates, Na-K-HCO3 water in volcanic rocks and alluvium derived from volcanic rocks, and a mixed facies found in many carbonate and alluvium water samples, and some volcanic waters. There is a general lack of lateral continuity in chemical characteristics along presumed flowpaths within each hydrologic unit (alluvium, carbonate, and volcanic). Though a lack of continuity between basins on the east side of the NTS was expected for water in alluvial and volcanic units due to the absence of interbasin flow, chemical differences observed within individual basins suggest a dominance of vertical over lateral flow. Groundwater in volcanic materials on the east side of Yucca and Frenchman Flats and on the west side of Pahute Mesa and Yucca Mountain has a nearly pure Na-K-HCO3 signature that reflects contact with primarily volcanic material. Groundwater in volcanic units in the middle of the NTS and on the east side of Pahute Mesa contains a higher proportion of Ca, Mg, Cl, and SO<sub>4</sub> than the other volcanic waters and indicates the contribution of water from the upper carbonate aquifer and/or hydrothermally altered regions. The lack of lateral chemical continuity in volcanic water on Pahute Mesa is attributed to stratigraphic and structural complexities in the volcanic units of the caldera complex. Areal variations in conservative ions in the carbonate aquifer indicate that though the lower carbonate aquifer may be considered a single hydraulic unit in gross flow terms, it may be considerably more complex when evaluating solute transport. Striking differences between groundwater in carbonate units at the south end of Yucca Flat and groundwater upgradient indicate the contribution of water from volcanic units, either by downward leakage or by lateral flow from structurally juxtaposed formations. A similar contribution of groundwater from volcanic formations is indicated by sodium and chloride concentrations in the carbonate aquifer near Fortymile Wash. The reduction in dissolved solutes in the carbonate aquifer at the southeast end of the NTS, as compared to water sampled at the south end of Yucca Flat, suggests a major contribution of groundwater flow coming from east of the NTS.

## INTRODUCTION

The dissolved ionic constituents of groundwater are a record of the minerals the water has contacted (either dissolving or precipitating) as it moves through soil and rock. As such, the chemistry of the water can be used to trace groundwater movement. The chemistry of groundwater in the Nevada Test Site (NTS) area has been instrumental in the development of the most basic flow hypotheses for the region. For example, the movement of water from volcanic aquifers into the regional carbonate aquifer in the vicinity of the NTS was deduced in large part by an increase in sodium concentration in the carbonate groundwater sampled at and downgradient from the NTS (Schoff and Moore, 1964). Low sodium concentrations in groundwater from the carbonate aquifer in the vicinity of Indian Springs were used as evidence that a groundwater divide directs flow from the NTS toward Ash Meadows rather than Las Vegas. Recognizing the importance of groundwater chemistry to hydrogeologic interpretations, the U.S. Geological Survey (USGS) performed extensive sampling in the NTS region that culminated with publications by Blankennagel and Weir in 1973 and Winograd and Thordarson in 1975. Three hydrochemical facies in and near the NTS were recognized: a Ca-Mg-HCO<sub>3</sub> facies from groundwater in carbonate rocks east and south of the NTS, a Na-K-HCO<sub>3</sub> facies from groundwater in volcanic rocks, and a Ca-Mg-Na-HCO<sub>3</sub> (mixed facies) in carbonate rocks beneath the NTS and in the east-central Amargosa Desert and Ash Meadows areas (Schoff and Moore, 1964).

The chemical data from which so many important conclusions were drawn are scattered through a number of USGS publications, and a few are referenced as personal communications or data on file in Denver USGS offices. Most of these data were obtained from samples collected in the late 1950s and early 1960s, though a few new wells were drilled and sampled in the late 1970s. The late 1980s and early 1990s have seen a resurgence of water sampling on the NTS, with samples collected from pre-existing wells and 28 new wells. These samples have been collected and analyzed by the Desert Research Institute (DRI).

This document compiles the USGS and DRI analyses in an effort to present a reasonably complete dataset for groundwater chemistry at the NTS. No exhaustive effort was made to identify all historic data; rather, the focus was on compiling the bulk of the analyses used to formulate important ideas about groundwater flow at the NTS. Some data, particularly recent samples from the Yucca Mountain area, have no doubt been overlooked. In cases where both old (1957–1977) and new (1983–1990) samples were available for the same well, the analyses have been compared using trilinear diagrams of ion percentages. In most cases, analyses compared well, so the datasets were combined and mean values for the constituents and pH for a given well were calculated. By dividing the wells into three groups based on the reported lithology of the sampling horizon (alluvium, volcanic, carbonate), the areal distribution of ions was mapped and compared to general groundwater flow directions.

## **Data Sources**

Chemical data for the period 1957 to 1977 were gathered from the following USGS reports: Schoff and Moore (1964), Blankennagel and Weir (1973), Young (1965), Thordarson *et al.* (1962), Robinson and Beetem (1965), and Dinwiddie and Weir (1979). These data are presented in Appendix A. Well names used in the appendices and throughout the text of this report are consistent with the names used in the 1991 edition of the Raytheon Services Nevada "NTS Drilling and Mining Summary." Wells are reported in the appendices in order of increasing Nevada state north coordinate.

All analyses but one reported from the period 1983 to 1990 were performed by the Water Analysis Laboratory of the Desert Research Institute. The one exception is the analysis from UE-25p #1, performed by the USGS and reported by Benson and McKinley (1985). Some of these DRI analyses have appeared previously in specific project reports prepared by DRI for the Department of Energy. These data are presented in Appendix B. Together, the USGS and DRI analyses represent 81 wells at and near the NTS (Figure 1). Average ion and pH values computed by combining the 1957-1977 and 1983-1990 datasets are given in Appendix C.

The chemical analyses reported in the appendices were performed on samples collected in several different ways. Samples were either collected during drilling and well development, with permanent pumps installed in supply wells, with temporary pumps used for aquifer testing and sampling, or with bailers. The degree to which a sample is representative of formation water varies with each method, but also varies with how a method was applied (e.g., number of pumping volumes before sampling) and sample collection and preservation procedures (e.g., filtering and acidifying cation samples). No attempt was made to assign quality ratings to the analyses assembled for this report because detailed sampling information was rarely available. However, it may aid the reader to keep in mind the following generalizations. Many of the analyses reported in Appendix A and the 1988 analyses from Pahute Mesa #3 in Appendix B were collected during drilling and well development and the resulting analyses could be influenced by residual drilling fluids. In some cases, this means that the producing horizon is actually different from sample to sample (e.g., samples were collected from multiple formations in UE-16d Eleana, USGS HTH #1, and U-19d #2). In Appendix B, samples from NTS supply wells were collected by pumping, and due to the lengthy pumping history of these wells, these samples are the most likely to be representative of formation water. These supply wells are those with "Water Well" in the name, plus UE-16d Eleana. Wells located around Yucca Mountain (wells with a "WT" for water table in the name) were sampled with a low volume piston pump and collected after pH, temperature, and electrical conductivity had stabilized (Matuska, 1989). The other samples in Appendix B were collected with a discrete bailer. The bailer was evacuated at the surface, lowered to a depth corresponding to a screened or open interval, then a valve was opened for sample collection. Because the water collected was present in the well bore, not the formation, there is the possibility that degassing of CO2 could have occurred and altered the pH and ion concentrations, and that reactions with casing materials could have occurred.

## COMPARISON BETWEEN 1957-1977 DATA AND 1984-1990 DATA

Eighteen wells have chemical analyses available from both the earlier and later sampling periods. Of these 18, 12 have no major change in salinity and relative ion percentages between the

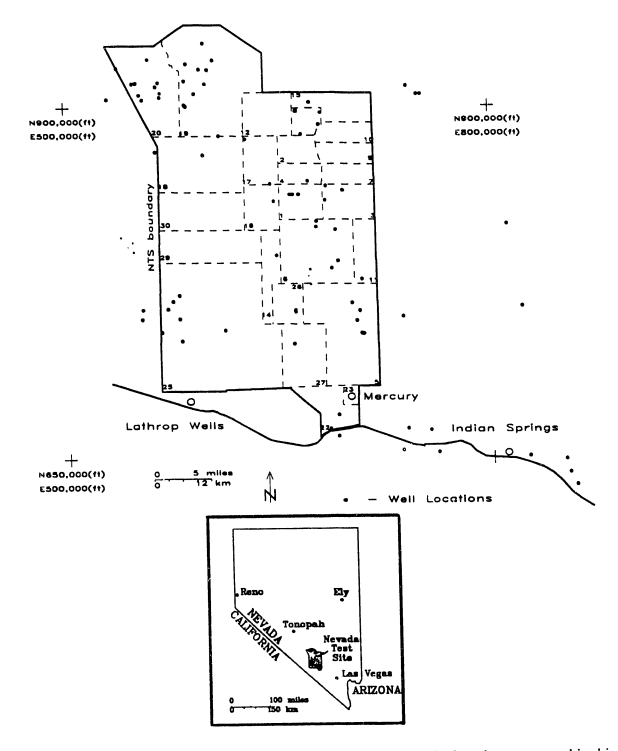


Figure 1. Location of the Nevada Test Site and wells with chemical analyses reported in this study. Well names can be found on detailed maps later in the report.

two groups. In a few cases (e.g., Water Well 2), there is no obvious break between the 1957–1977 and 1984–1990 data groups but there is a relatively large spread in the ion ratios of the earlier analyses. This may be caused by continuing well development with pumping, resulting in more representative formation water with time. The six wells with significant differences between the old and new analyses are discussed below in the following order: UE–15d Water Well, UE–16f Eleana, UE–16d Eleana, Army 6A, USGS Test Well D, and USGS HTH #1. Because there is little change in overall salinity of samples from a given well, trilinear plots are used to discuss temporal changes of the total milliequivalents per liter of cations (left triangle), percentages of the total milliequivalents per liter of anions (right triangle), and a combined representation of the anions and cations by projecting these points onto a central diamond.

A sample collected from UE-15d Water Well in 1989 is within 10% of the ion ratios of samples collected in 1961 and 1962 (Figure 2). However, two samples collected in 1990 have shifted to approximately 15 to 20% more calcium relative to other cations, as compared to the earlier samples. Total salinity is essentially unchanged. It seems unlikely that preservation or analytical errors could be the source of the difference because there are two independent samples defining each group, and anion percentages remain stable. Rather, the difference may be related to a reduction in pumping at the well that began in 1990 and culminated with the pump becoming inoperative in 1991. This well is completed in fractured quartzite with a dolomite unit at the bottom of the hole. The 1960's and 1989 samples probably reflect water produced primarily from the quartzite. The change in chemistry in 1990 could reflect mixing of water from the dolomite in the well bore in the long periods between pumping in 1990. The shift in chemical composition is important because there has been an increase in the tritium concentration in this well over the last 15 years (Lyles, 1990 and 1992). Identifying the formations contributing water to the well would aid in determining the source of the tritium. To this end, the inoperative pump in UE-15d Water Well should be pulled and hydrologic logging performed.

Analyses are available from UE-16f Eleana from samples collected at the time of drilling and testing (1977) and in 1988. The three samples show a progressive increase in the amount of HCO<sub>3</sub> relative to the other anions, with the 1988 sample having greater than 95% HCO<sub>3</sub>, less than 5% Cl, and essentially no SO<sub>4</sub> (Figure 3). This trend could be the result of well development, meaning that the most recent sample is the most representative of formation water.

The three 1977 samples from UE-16d Eleana were collected during drilling and show up to 10% shifts in ion percentages from one sample to another (Figure 4). Presumably, these variations are related to different contributing horizons at the time of sampling. The 1977 sample most different from the 1990 sample was collected on 6-19-77 and is recorded as being from the Eleana Formation (quartzite). The well was recompleted in 1981 with the Eleana Formation cemented off and perforations installed across the Tippipah Limestone, so it is not surprising that there are differences between the 6-19-77 sample and the one in 1990. The greatest similarity in cations exists between the 1990 sample and the one collected on 6-14-77. The 6-14 sample was the only earlier sample

Date (m/d/y)	Depth (m)	
A 11/12/61	591	438
B 03/27/62	1829	436
C 09/11/89	495	435
D 04/18/90	495	493
E 08/01/90	495	408

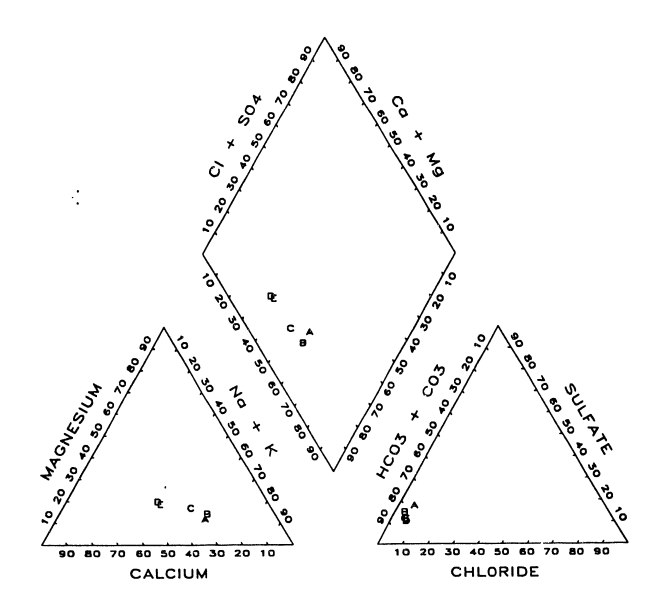


Figure 2. Trilinear diagram showing relative ion percentages for samples collected from UE-15d Water Well. The depth of the 1989 and 1990 samples is the location of the pump. The well was open to 1829 m.

Date (m/d/y)	Depth (m)	TDS (mg/i)
A 08/20/77	207256	1040
B 09/25/77	394-436	
C 09/20/88	378	975

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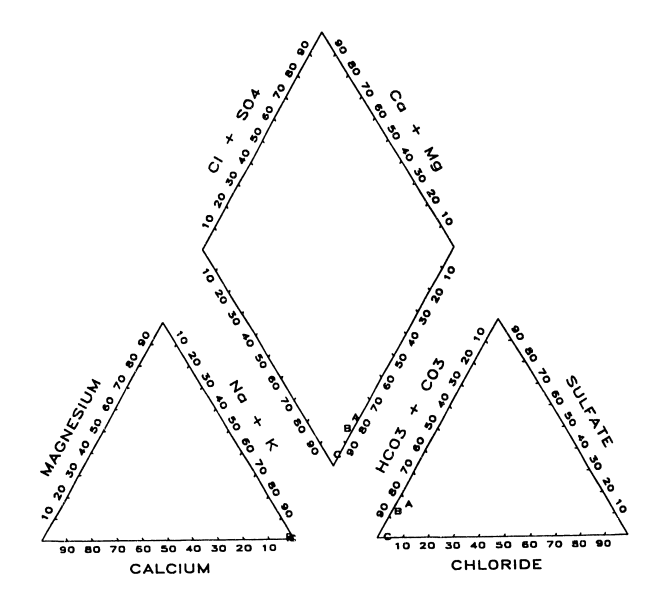


Figure 3. Trilinear diagram showing relative ion percentages for samples collected from well UE-16f Eleana.

Date (m/d/y)	Depth (m)	
A 05/23/77	25-253	309
B 06/14/77 C 06/19/77	25-646 468-527	386
D 06/15/90	335	418

2

3

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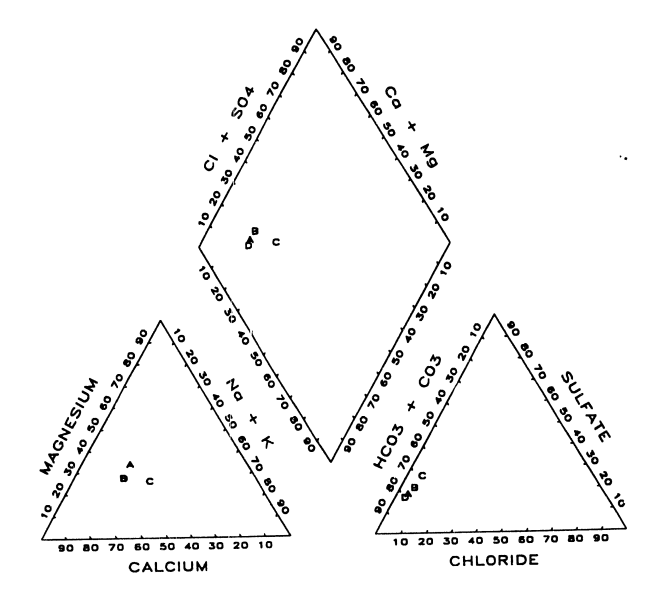


Figure 4. Trilinear diagram showing relative ion percentages for samples collected from well UE-16d Eleana.

that was collected by pumping, and the similarity in chemistries suggests that the Tippipah Limestone was probably the major contributing zone to the borehole during pumping in 1977.

The 1958 analysis of water from well Army 6A differs drastically from two analyses of samples collected in 1986 (Figure 5). The total dissolved solids (TDS) concentration in the more recent samples is about 7% lower than the earlier one, but the major difference is a large increase in the amount of magnesium at the expense of sodium, and an increase in SO<sub>4</sub> at the expense of HCO<sub>3</sub>. Schoff and Moore (1964) were suspicious of the 1958 analysis because it yielded a Na-K water in an area where other water was a Ca-Mg type. The later analyses do not resolve the problem, but instead make it one of a Mg-SO<sub>4</sub> type water in a Ca-Mg-HCO<sub>3</sub> area. Schoff and Moore concluded that the earlier analysis did not represent formation water and suggested that it could have been contaminated by cement used in the well during completion. However, the measured pH (8.0) was much lower than what would be expected from a cement-contaminated water (greater than 11). The later analyses also have reasonable pH values (around 8.6), and cation/anion charge balances are very good for both the 1958 analyses.

Though cement contamination seems unlikely, water introduced during drilling could have altered the *in situ* chemistry. Given that the well has a low production rate (1 to 2 gpm) and was abandoned immediately, water introduced during drilling in 1955 may not have been purged by the time the sample was collected in 1958. The difference between the 1958 and 1986 analyses may indicate that diffusive and/or advective flow have equilibrated the well with its surroundings. The log of well Army 6A includes quartzite and limestone, but only quartzite and siltstone are intersected by the perforated interval (Moore, 1962). Both the old and new analyses indicate equilibrium with dolomite, and some source of sulfate is needed, neither of which are consistent with a quartzite aquifer. Though anomalous, there are other wells with unusual chemistries in the area (USGS HTH "F" and Army 6) that have been attributed to hydrothermal activity and/or as yet undetected evaporite deposits in the subsurface. Pumping the well at a low flow rate, monitoring field parameters, then sampling after purging many well volumes could reduce the uncertainty about the chemistry of this well.

Samples collected in 1986 from USGS Test Well D are substantially lower in TDS than the one collected in 1961. This difference in TDS is not reflected in any change in cation ratios (Figure 6), but a shift to a higher proportion of HCO<sub>3</sub> in the 1986 samples is due to lower Cl and sulfate concentrations in those samples as compared to the 1961 sample. The 1960 sample was collected from a bailer after three hours of nearly continuous bailing had removed 12,000 liters (3100 gallons) from the well (Thordarson *et al.*, 1962). Though care was taken to minimize contamination of groundwater by using a cable-tool rig and not introducing bentonite or chemical additives in the zone of saturation, the aquifer was penetrated by at least 16 meters (53 ft) and water was injected in the hole for cuttings removal, before groundwater was recognized. The hole also experienced serious caving problems both from the overlying tuff units when the hole penetrated the carbonates (with blocks of tuff falling into the bailer), and from interbedded siltstone and argillite formations. Fine material from these horizons may have dissolved into the well water and added to the TDS concentration. Though no extensive well development or pumping is known to have occurred at

Date (m/d/y)	Dapth (m)	TDS (mg/i)
A 04/27/58	382	718
B 06/25/86	320	666
C 06/25/86	324	673

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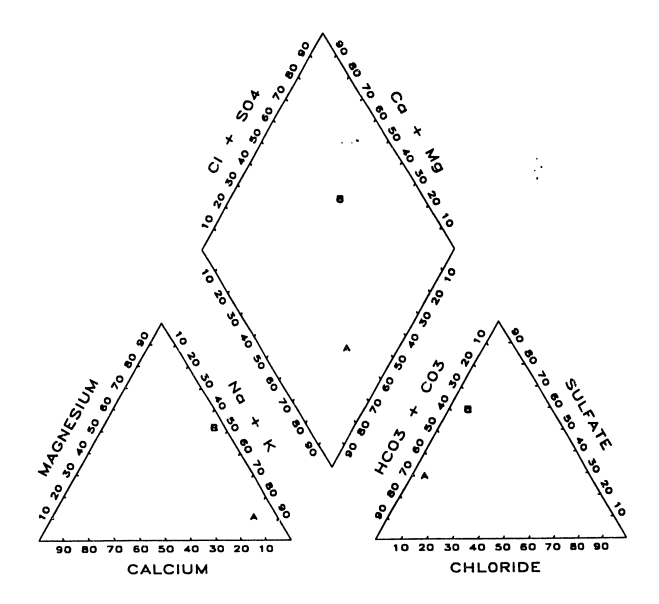


Figure 5. Trilinear diagram showing relative ion percentages for samples collected from well Army 6A.

	Date (m/d/y)	Depth (m)	TDS (mg/i)
A	01/09/60	594	398
	06/23/86	531	321
ē	06/23/86	561	321

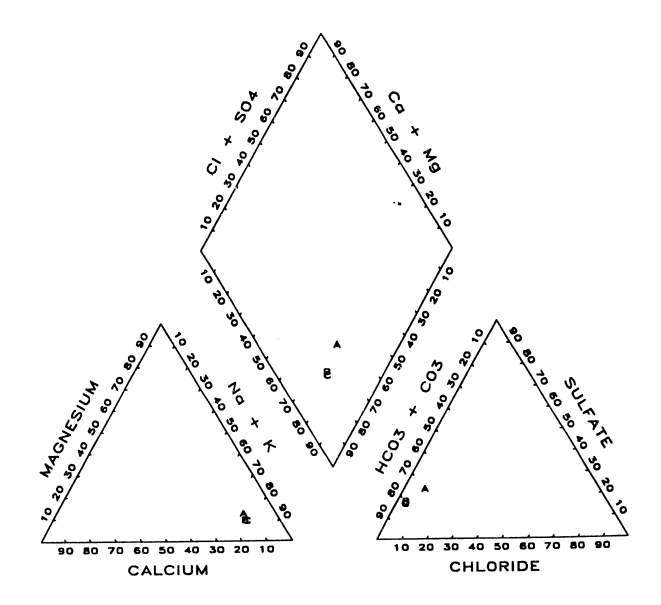


Figure 6. Trilinear diagram showing relative ion percentages for samples collected from USGS Test Well D.

USGS Test Well D between 1961 and 1986, equilibration of well water with the carbonate aquifer by advective or diffusive flow is reasonable over this time scale. Though the ionic composition of the 1986 samples may be more representative of formation water, the lower pH of the 1961 sample may be closer to *in situ* conditions. The 1986 samples were collected with a discrete bailer without previous purging of wellbore water, so some degassing of  $CO_2$  and elevation of pH could have occurred. As with Army 6A, new samples from this well should be collected after pumping and monitoring field parameters.

Some of the variation observed at well USGS HTH #1 (Figure 7) comes from sampling from two discrete aquifers. Samples from the underlying carbonate aquifer were only collected in 1961 and 1962. These are the samples with the highest calcium percentages. The other samples collected in 1960 and from 1986–1990 were collected from volcanic zones and show higher proportions of sodium and bicarbonate, with the 1986–1990 set clustering more in the Na and HCO<sub>3</sub> apexes. All of the samples together appear to form a mixing line between a higher salinity, carbonate–influenced water and a lower salinity, volcanic–influenced water. This suggests that differences between individual samples relate to their collection position in the borehole rather than any hydrochemical changes with time. Flow relationships between the five perforated intervals in USGS HTH #1 and how they relate to the 1986–1990 samples are discussed by Lyles *et al.* (1991).

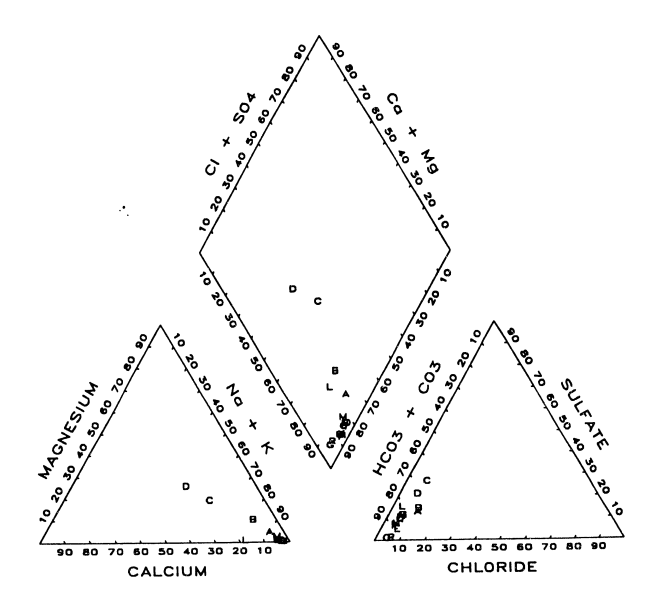
## **AREAL DISTRIBUTION OF DISSOLVED IONS IN GROUNDWATER**

The three hydrochemical facies identified by previous workers are also represented in the averaged dataset created by combining the 1957–1977 data with the 1984–1990 data. Groundwater ranges from a Ca-Mg-HCO<sub>3</sub> water to a Na-K-HCO<sub>3</sub> water, with a rough mixing line between the two on the cation triangle (Figure 8). Water from volcanic rocks predominates in the Na-K area, while water from the carbonate rocks has generally higher percentages of calcium and magnesium. Most waters from carbonate rocks have a mixed chemical character, as do some volcanic aquifer waters. Water from alluvial deposits spans the chemical spectrum, depending on the source rock for the alluvial material. With the exception of a few outliers, there is less variation in anion percentages, with most waters dominated by bicarbonate.

The chemistry of the combined dataset continues to strongly support the hypothesis of Schoff and Moore (1964) and others that two distinct types of water develop depending on whether groundwater moves through a volcanic or carbonate aquifer. Significantly, the mixed nature of many samples from the NTS area indicates a water that has moved through both types of material.

Given the importance of aquifer material on groundwater chemistry, the combined 1957–1990 dataset was subdivided into three groups: wells completed in alluvium (14 wells), wells completed in carbonate rocks (17 wells), and wells completed in volcanic rocks (50 wells). This simplistic designation, while easy to make in many cases, was more problematic in others. An example is UE–16f Eleana, which is completed in the Eleana Formation, comprised of siltstone, sandstone, and quartzite with minor limestone. This well was grouped with the carbonates because the lithology seemed to have more in common with that group than with the volcanics or alluvium. However, it

Date (m/d/y)	Depth (m)	TDS (mg/l)	Date (m/d/y)	Depth (m)	TDS (mg/i)
• • • • • •	171	151	J 04/26/89	651	124
A 10/01/60 B 10/20/60	337	150	K 04/26/89	686	125
C 05/09/61c	1137	278	L 09/24/90	472	153
$D \ 08/11/62c$	1282	208	M 09/24/90	588	127
E 07/24/86	461	141	N 09/24/90	622	126
F 07/28/87	725	129	0 09/25/90	649	124
G 07/28/87	1119	115	P 09/25/90	686	128
H 04/25/89	588	125	0 09/25/90	732	124
1 04/25/89	622	128	R 09/26/90	1122	107



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Figure 7. Trilinear diagram showing relative ion percentages for samples collected from well USGS HTH #1. Samples C and D (with a "c" after the date) were collected from carbonate rocks, while all of the others were from volcanic units.

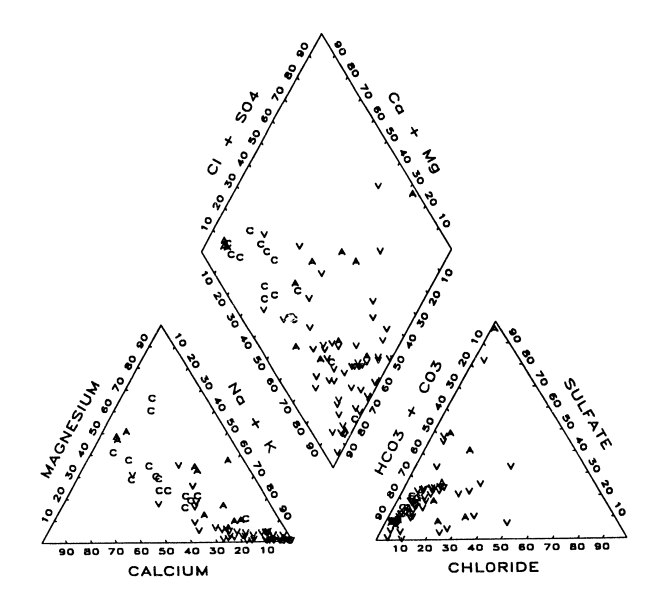


Figure 8. Trilinear diagram showing relative ion percentages for samples collected from all wells reported in this study, differentiated based on aquifer rock type. Aquifer designations are as follows: A=alluvium, C=carbonate, and V=volcanic. Averaged values used to construct this diagram are reported in Appendix C.

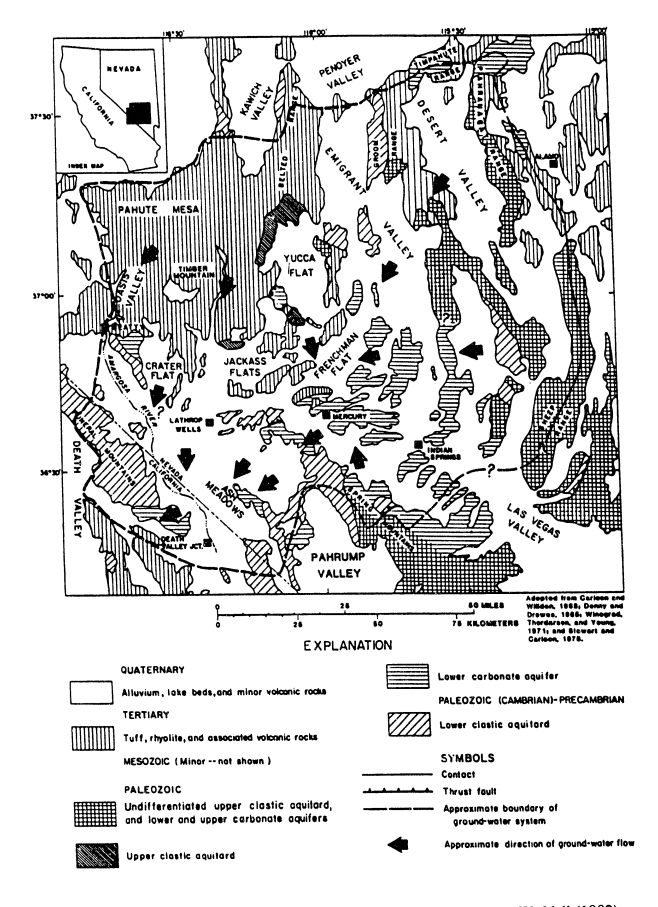
has essentially 100% Na+K in terms of cations, making it chemically unlike the other carbonate wells and more similar to the volcanics. Despite such problems, aquifer designations were made based on rock type rather than by second guessing the chemical interpretations to follow.

The chemical data were examined in the context of our present knowledge of the physical flow system. Groundwater flow directions in the NTS area have been suggested based on geologic and geophysical information, water-level data, precipitation data, spring discharge measurements, aquifer test data, as well as water chemistry. Though considerable uncertainty in flow boundaries and inter-unit flow exists, workers have generally agreed on the gross regional groundwater flow directions at the NTS. Flow is believed to move primarily in a north-to-south direction across the NTS with an additional component of northeast to southwest flow (Figure 9). Flow southward through Yucca Flat into Frenchman Flat is joined by a component of westerly flow from Emigrant Valley and other areas east of the NTS. After passing through Mercury Valley, there is a southwestward turn as groundwater moves toward the discharge area in Ash Meadows. Components of groundwater flow beneath Pahute Mesa move southward into the Fortymile Wash structural feature and on to discharge at Alkali Flat and in Death Valley. There is also southwestward flow from Pahute Mesa to Oasis Valley. Flow relationships between the east and west sides of the NTS are less certain, with postulated flow from Pahute/Rainier Mesas into Yucca Flat in the north and possibly from western Yucca and Frenchman Flats to Jackass Flats in the south.

The following sections compare the chemical data for each type of aquifer material with the postulated flow directions discussed above. Well water chemistry is represented on maps using Stiff diagrams. The Stiff diagrams show the geochemical similarities and differences among wells; the cation concentrations are plotted on the left side of each Stiff figure in units of milliequivalents per liter, while the anions are similarly plotted on the right side. The graphical similarities/differences between Stiff diagrams represent the geochemical similarities/differences of the water samples collected from each well (Hem, 1985).

## **Alluvium Wells**

All wells providing water samples from alluvial material in this study are located either in the eastern part of the NTS or east of the NTS (Figure 10). Regional (interbasin) groundwater flow beneath the eastern part of the NTS is believed to occur only through carbonate rocks (Winograd and Thordarson, 1975). Groundwater in alluvial deposits is isolated in each basin, so regional trends in ion concentrations would not be expected in wells completed in alluvium. For example, USGS Water Well A and UE–6d are in Yucca Flat, while Water Well 5B and Water Well 5C are in Frenchman Flat. Alluvial materials are absent in some locations and unsaturated in others between Yucca and Frenchman Flats so that water cannot flow from UE–6d to Water Well 5B without entering volcanic and carbonate units. This segmentation of alluvial water is reflected in the areal distribution of ions shown on Figure 11. The difference in ion concentrations within an individual basin (*e.g.*, sodium at Water Well 5B is 97 mg/L, while at Water Well 5C it is 129 mg/L) is also marked and suggests a lack of well–developed lateral flow systems that would homogenize chemical character within each basin. The logical extension of this observation is that vertical flow dominates



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Figure 9. General groundwater flow directions in the NTS area. From Waddell (1982).

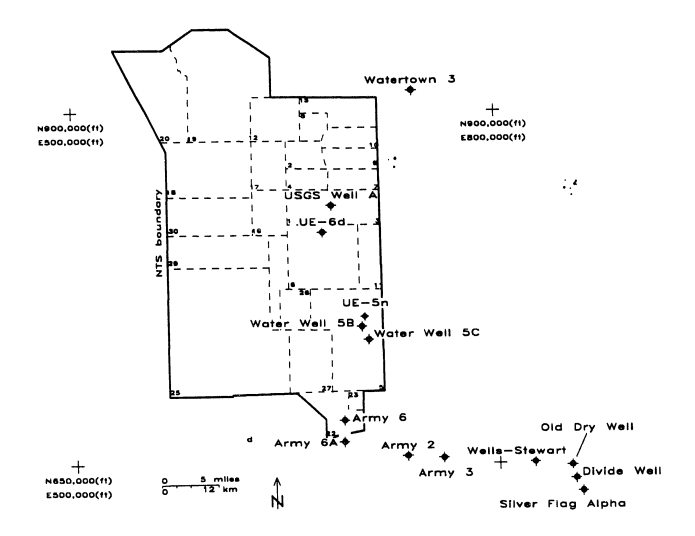


Figure 10. Wells completed in alluvial materials at and near the NTS for which water chemistry data are available. Nevada state coordinates are shown for reference.

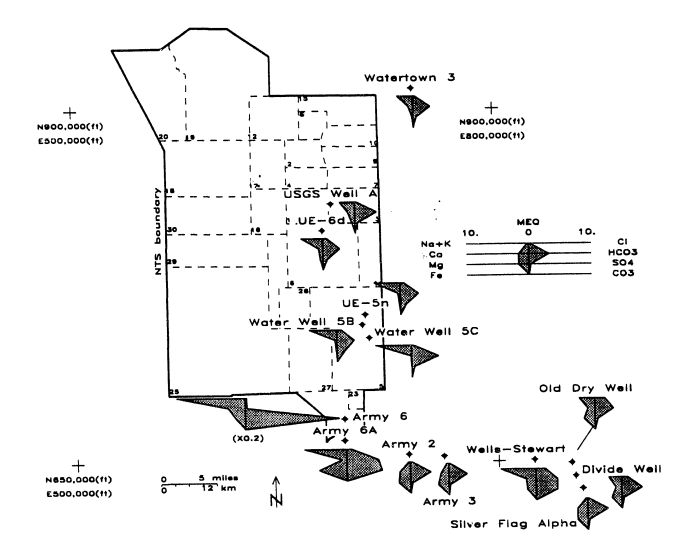


Figure 11. Areal variation in water chemistry for wells completed in alluvial materials, as represented by Stiff diagrams.

in the alluvium. The one exception appears to be the saturated alluvium in the vicinity of wells Army #2 and Army #3, where there is good consistency in ion concentrations.

## **Carbonate Wells**

As with the alluvium wells, all but one of the carbonate wells for which chemistry data were available are located on the eastern part of the NTS, or east of the NTS (Figure 12). Flow is believed to move through carbonate rocks from the north end of Yucca Flat, through Frenchman Flat and Mercury Valley. A westward flowing component joins this southerly component from the area sampled by USGS HTH #10 and USGS HTH #4, and points east, and this combined system moves in a southwestwardly direction toward Ash Meadows. Quartz and Sand Springs, also east of the NTS, occur in the Pintwater Range and represent perched water that does not contribute to the regional flow system. The one western carbonate well, UE-25p #1, is located in the Fortymile Wash drainage, where flow is believed to be more directly southward to Ash Meadows.

Though the carbonate aquifer is considered a regional aquifer system, the groundwater chemistry reveals that the units intersected by the carbonate wells are not chemically homogeneous. Though some of the variation in ion composition seen on the carbonate map (Figure 13) can be attributed to the lumping of all pre-Tertiary units for mapping purposes (in particular, UE-15d Water Well, UE-16d Eleana, and UE-16f Eleana are not sampling the lower carbonate aquifer), unaccounted shifts in ion concentrations occur. For example, USGS Test Well D and UE-1c sample carbonate units about 3000 m (10,000 ft) apart in mid-Yucca Flat. Dissolved chloride is over 1.5 times higher at USGS Test Well D and dissolved sodium is almost three times greater than at UE-1c. UE-1c is located farther to the south (downgradient) and thus would be expected to either have similar or higher dissolved ion concentrations than USGS Test Well D. Moving farther southward along the presumed flow direction, chloride and sodium values increase markedly at USGS Water Well C and Water Well C-1, indicating a contribution of water from volcanic units. Water could be moving into the carbonates by enhanced downward leakage through volcanics and alluvium beneath Yucca Lake or laterally where volcanic units are juxtaposed with carbonates along faults. USGS Water Well C and Water Well C-1 are located in the upper plate of a low-angle thrust fault (Winograd and Thordarson, 1975), which probably abuts volcanic units to the north of the wells. The very old radiocarbon age of water in Water Well C-1 (greater than 30,000 years before present; Boughton, 1986) indicates that the water has had a long residence time, whether derived from lateral or vertical flow. Relatively high chloride and sodium concentrations at UE-25p #1 similarly indicate a contribution of volcanic water to the carbonate aquifer in the Fortymile Wash area. Saturation calculations show that all of the carbonate waters are saturated with respect to calcite and quartz, whether they have mixed with volcanic water or not.

The wide concentration variations in conservative ions such as chloride and sodium suggest a complex carbonate system possibly containing isolated flow units such as the fault block containing the C wells. Transmissivity in the lower carbonate aquifer is structurally controlled and known to have a wide range in values. However, hydraulic testing indicates that water-bearing fractures are reasonably well connected, creating a grossly homogeneous aquifer (Winograd and Thordarson, 1975). In addition, Winograd and Thordarson report nearly identical chemical analyses

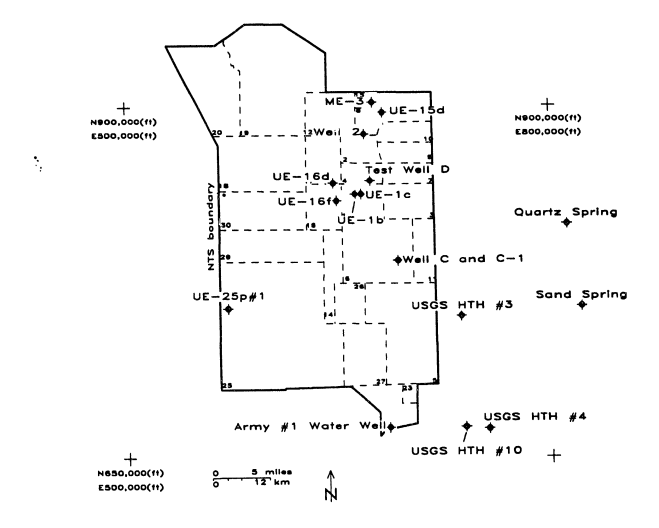
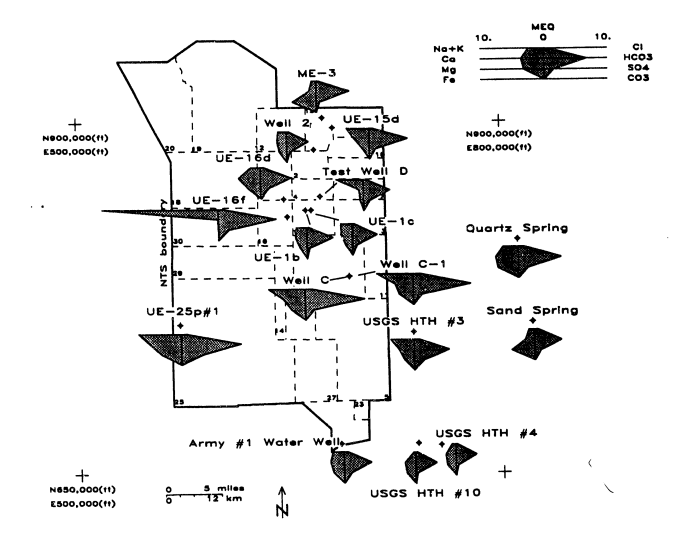
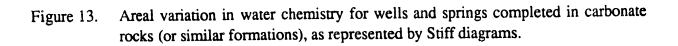


Figure 12. Wells and springs completed in carbonate rocks (or similar formations) at and near the NTS for which water chemistry data are available. Nevada state coordinates are shown for reference.







for water from different formations comprising the lower carbonate aquifer, sampled within a single well. Despite this earlier work, the range in groundwater chemistry in the carbonate aquifer across the eastern portion of the NTS indicates that though the lower carbonate aquifer may be considered a single hydraulic unit in gross flow terms, it may be considerably more complex when considering solute transport.

One feature of the carbonate hydrochemical map that can be explained is the significant decrease in dissolved ion concentrations that occurs between the south end of Yucca Flat and Mercury Valley, as typified by the halving of chloride and reduction by two-thirds in sodium concentrations from USGS Water Well C to Army #1 Water Well. Such a reduction in conservative ions could only occur with the addition of a lower salinity water to the system. Leakage from the alluvium and volcanic aquifers in Frenchman Flat cannot be a source of dilution because sodium concentrations are much higher in those units than those at Army #1 Water Well. A more likely source of dilution, suggested by most flow models, is the movement of water from points east and south in the carbonate units. Groundwater in the USGS HTH #4 and #10 area probably originates as recharge in the northern Spring Mountains (Byer, 1991; and Sadler et al., 1991). The low salinity and relatively low calcite saturation index at these wells supports the relatively young ages (approximately 5000 years) calculated for groundwater in the area using a discrete-state compartment model (Sadler et al., 1991). Using ion concentrations averaged between those of USGS HTH #10 and USGS HTH #4 to represent underflow from the east, the relative contributions from Yucca Flat (as represented by USGS Water Well C and Water Well C-1) and the eastern area required to produce the water sampled from Army #1 Water Well are approximately 30% water from Yucca Fiat and 70% water from the east. These approximate percentages are upheld by all of the major ion concentrations with the exception of sulfate (Table 1). Several other wells in the vicinity of Army #1 Water Well have high sulfate concentrations (Army 6 and Army 6A), suggesting a local source for dissolved sulfate, either from hydrothermally altered locks or gypsum deposits.

Ion	Concentration in Well C	Concentration in HTH 4 & 10	Concentration in Army #1	Percent from Yucca Flat	Percent from east of NTS
Ca	65	36	44	35	65
Mg	28	18	21	30	70
Na	125	10	38	25	75
K	14	2	5	30	70
Cl	32	6	16	40	60
SO <sub>4</sub>	65	16	53	75	25
HCO <sub>3</sub>	555	197	257	15	85

Table 1. Dissolved ion concentrations in mg/L for wells in three areas of the carbonate aquifer and calculated mass percentages needed to create water at Army #1 Water Well from the other two types.

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Another possible explanation for the difference between the "C" wells and Army #1 Water Well is that water from USGS Water Well C and Water Well C-1 is not representative of the carbonate water composition leaving Yucca Flat. This would assume that the well C structural block is isolated from the rest of the flow system and that flow from Yucca Flat is typified by the composition of water from Water Well 2, UE-1b, and UE-1c. This scenario would allow little or no contribution of water from the east, and is not compatible with the conceptual groundwater flow models.

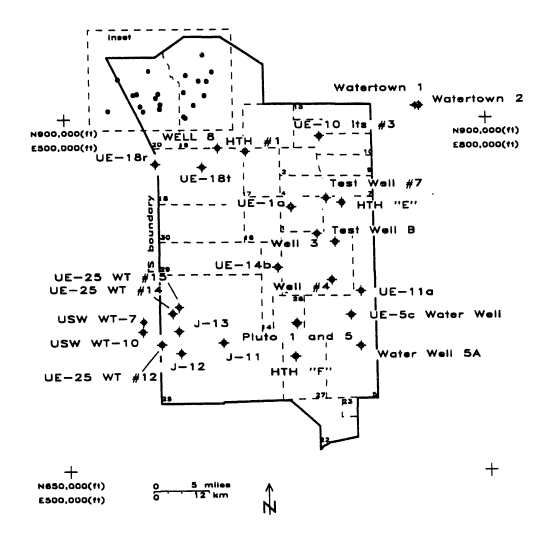
## **Volcanic Wells**

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Wells that penetrate volcanic units are found throughout the NTS (Figures 14 and 15). In the eastern area, coincident with the areas where there are alluvium and carbonate wells, the volcanic aquifers are not believed to form continuous interbasin flow units. As with groundwater in the alluvium, it is believed that water in the volcanics must discharge to the carbonate aquifer to flow from Yucca Flat to Frenchman Flat and then to Mercury Valley. Spatial variations in ion concentrations from these wells suggest the absence of a well-integrated lateral groundwater flow system (Figure 16). One general pattern is suggested by the data, however. Wells on the west side of Yucca and Frenchman Flats tend to have higher calcium and magnesium concentrations and lower pH values than wells to the east. This is balanced by lower sodium concentrations and/or higher chloride and sulfate concentrations. The wells showing the higher calcium and magnesium values are USGS HTH "F", Pluto 1, Pluto 5, UE-1a, and to a lesser extent Well 3 and Water Well #4. An exception to this trend is well UE-14b, located in Mid Valley, which does not have notably high calcium and magnesium concentrations, but does have higher sulfate than the eastern wells. The higher calcium and magnesium concentrations in the western Yucca and Frenchman wells (UE-1a, Well 3, Water Well #4) may reflect the movement of groundwater into volcanic units from the upper carbonate aquifer in the structurally complex area along the Eleana Formation subcrop bounding western Yucca Flat. The impact of hydrothermally altered rocks is believed to account for the higher sulfate concentration in J-11, USGS HTH "F" and possibly UE-14b. The Pluto wells sample perched water that probably receives calcium via surface recharge processes.

A lack of intrabasin chemical continuity is particularly noteworthy in Frenchman Flat where both sodium and bicarbonate are twice as high in Water Well 5A than in UE-5c Water Well, less than 10 km (six miles) to the north. In this case, and perform others, structural features may have isolated the volcanic units. Water Well 5A is on the opping of the Rock Valley Fault System from UE-5c Water Well and a 4 m (13 ft) difference in head values attests to some hydraulic discontinuity between the two areas.

Though chemical variations in water from volcanic wells on Pahute Mesa are generally less than variations found elsewhere on the NTS, ion concentrations in well water on the Mesa do not vary in any pattern consistent with postulated flow directions (Figure 17). This may indicate that the complex sequence of volcanic units beneath Pahute Mesa does not constitute a single, unified flow system. Within the saturated zone, the vertical and horizontal distribution of permeable ash-fall and ash-flow tuffs and lava flows that comprise the aquifers of Pahute Mesa is highly variable, even over short distances. This stratigraphic variability, coupled with structural control of groundwater flow through fractures in a complex caldera environment, results in a hydrologic system that does



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Figure 14. Wells completed in volcanic rocks at and near the NTS for which water chemistry data are available. Well names for the Pahute Mesa area can be found on Figure 15. Nevada state coordinates are shown for reference.

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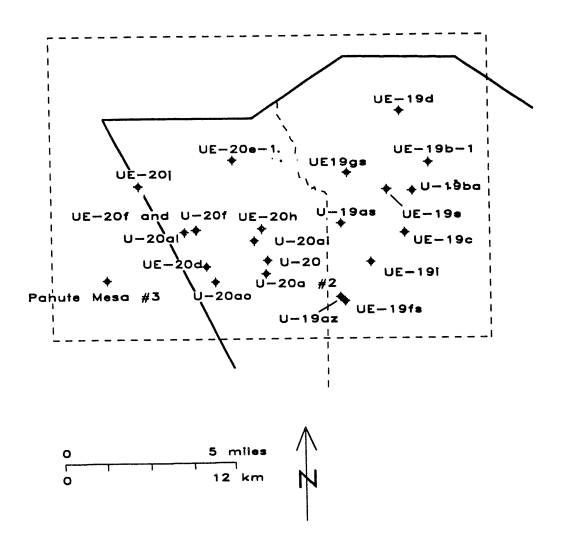


Figure 15. Wells completed in volcanic rocks on Pahute Mesa for which water chemistry data are available. For location relative to the NTS, see Figure 14.

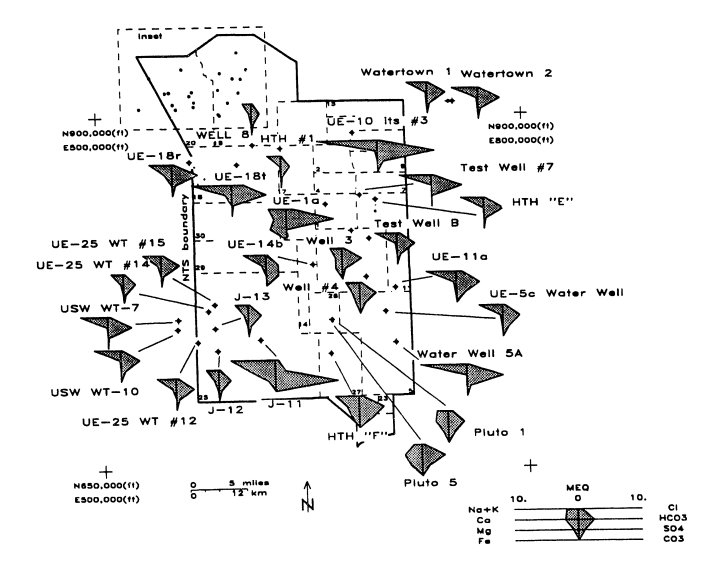


Figure 16. Areal variation in water chemistry for wells completed in volcanic rocks, as represented by Stiff diagrams. The Pahute Mesa area is shown separately on Figure 17 for clarity.

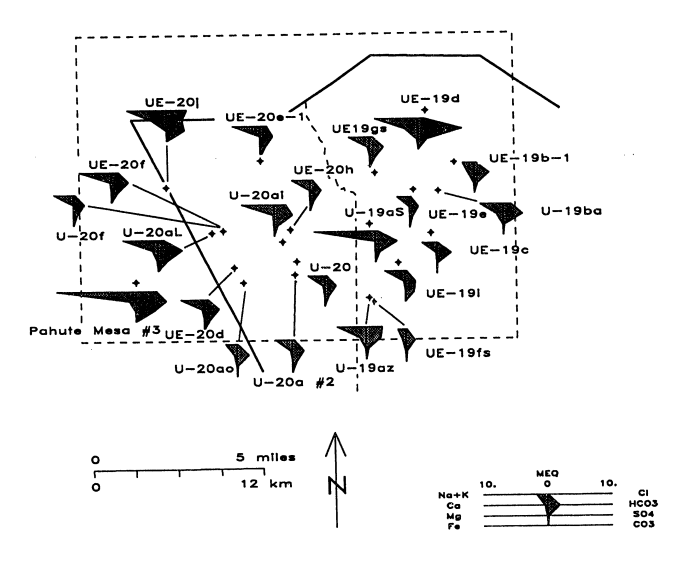


Figure 17. Areal variation in water chemistry for wells completed in volcanic rocks on Pahute Mesa, as represented by Stiff diagrams.

not have predictable continuity in lateral directions. The apparent lack of consistent chemical changes in downgradient directions may actually result from an incomplete understanding of flow path directions. In addition, some variability results from sampling perched horizons that may have been contaminated by drilling fluid (at U-19ba and U-19az).

Blankennagel and Weir (1973) observed that hydraulic heads decreased with depth beneath the eastern part of Pahute Mesa (indicating a potential for downward flow) and that heads increased with depth beneath western Pahute Mesa. In addition, they also observed that while sodium and bicarbonate were the dominant ions in all Pahute Mesa groundwater, there was proportionally more calcium in the area of downward flow and more sulfate and chloride in the zone of upward flow. The new data blur the distinction for cations, but the difference in anions is similar (Figure 18). Sulfate is often associated with hydrothermal alteration and is consistent with an area where deeper, older water is moving upward in the groundwater system.

Another cluster of volcanic aquifer samples is located in the southwest corner of the NTS (Figure 16). A major hydrologic feature in this area is Fortymile Wash, which is believed to mark a zone of higher hydraulic conductivity and greater recharge (Byer, 1991; Sadler *et al.*, 1991). With the newer samples, the chemical similarity between J–12 Water Well and J–13 Water Well can now be extended further north along the wash to UE–25 WT #14, which has an ionic composition very similar to the downgradient wells. Even farther to the north, UE–25 WT #15 is also similar to the other Fortymile Wash samples but has a higher percentage of sodium and bicarbonate. Well UE–18t is located much farther north, near the head of the wash and has a greater dissolved ion content than the southern Fortymile Wash wells. The lower salinity of the downgradient wells may reflect the diluting effect of infiltrating recharge along the length of the wash.

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The southern Fortymile Wash wells have about five times the amount of calcium and over 10 times the amount of magnesium as the two wells (USW WT-7 and USW WT-10) in the drainage basin west of Yucca Mountain. The western wells have a dominantly Na-HCO<sub>3</sub> signature characteristic of water from volcanic aquifers. The higher concentration of calcium and magnesium in the Fortymile Wash area may indicate a contribution of water from underlying carbonate rocks, infiltration of rainfall that dissolves near-surface calcium deposits, or mixing with water from areas east of Fortymile Wash, where higher dissolved ion concentrations are attributed to interaction with hydrothermally altered rocks. Groundwater from J-11 represents this higher Ca-SO<sub>4</sub> water, which is also found at USGS HTH "F." The chemistry of the carbonate water in Fortymile Wash, as sampled by UE-25p#1, indicates mixing of volcanic water into the carbonate unit rather than movement of water from the carbonates to volcanics. Either this mixing occurs outside the area sampled by the volcanic wells (e.g., where the carbonates abut the nearby caldera complexes), or the calcium and magnesium in the volcanic wells must be from one of the sources other than the underlying carbonate. Based on geochemical and isotopic data, Matuska (1989) concluded that flow from the carbonates into the volcanics in the Fortymile area was unlikely.

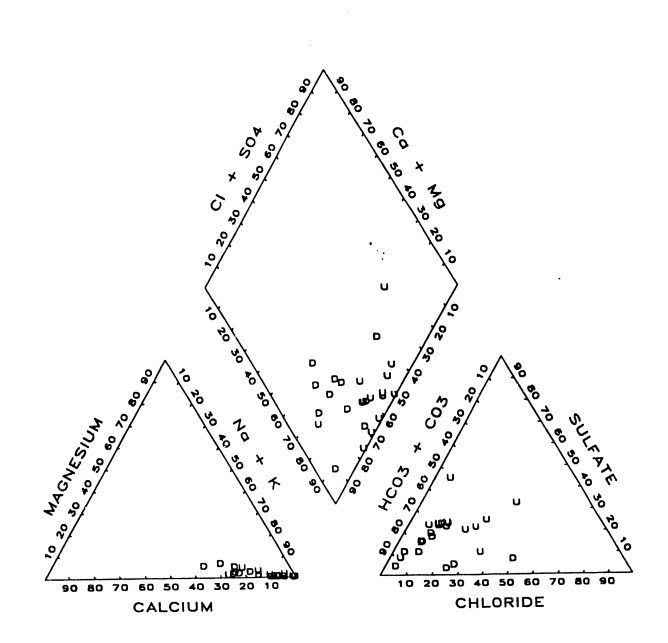


Figure 18. Trilinear diagram showing relative ion percentages for samples collected from volcanic aquifers on Pahute Mesa, differentiated based on their location relative to the areas of downward ("D") and upward ("U") flow defined by Blankennagel and Weir (1973).

### CONCLUSIONS

Groundwater samples from 81 locations at and around the NTS have been analyzed for major dissolved constituents. Twenty-seven of these locations are new additions sampled by DRI after publication of major interpretive reports by the USGS. Eighteen locations were sampled by both DRI and the USGS with generally excellent agreement between results. Most cases with discrepancies can be explained by continued well development or changes in well completion over the years.

The new data support the occurrence of three hydrochemical facies in the NTS area, identified by previous workers. The distribution of these facies, in particular the presence of mixed cation water in the carbonates, has led to significant conclusions regarding gross groundwater flow by suggesting the movement of water from volcanic aquifers into the carbonate aquifer. However, when examined in detail, the hydrochemistry of the samples in this study is not fully compatible with concepts of regional lateral flow in the carbonate aquifer. There are wide variations in concentrations in generally conservative ions that suggest that though the carbonate aquifer may be considered a single hydrologic unit in gross flow terms, it may be considerably more complex when considering solute transport. The chemistry in wells at the south end of Yucca Flat and near Fortymile Wash indicates a greater contribution of water from volcanic units to the carbonate aquifer in these areas than found elsewhere at the NTS. Ionic concentrations indicate that at the southern end of the NTS, the majority of water in the carbonate system originated from areas east and south of the NTS, with only about 30% of the water coming from southern Yucca Flat.

While differences in chemistry in alluvium and volcanic wells between basins in the eastern NTS were expected because of a lack of interbasin flow, the absence of homogeneous chemical character within single basins suggests a lack of well-developed lateral flow systems and dominance of vertical groundwater flow in these units within each basin. Relatively higher calcium and magnesium in volcanic wells in the south-central part of the NTS (west side of Yucca and Frenchman Flats and east side of Jackass Flats) may reflect movement of water from the upper carbonate aquifer into the volcanics and contributions from hydrothermally altered areas. The low-calcium water found in eastern Yucca and Frenchman Flats and west of Yucca Mountain is typical of water that has encountered primarily volcanic material along its flowpath. Consistent chemical patterns could not be identified in the volcanic aquifers of Pahute Mesa, presumably because of complex stratigraphic and structural controls. In the southwestern part of the NTS, chemical homogeneity supports groundwater flow in volcanic units along Fortymile Wash, while the calcium and magnesium concentrations suggest a contribution of water from hydrothermally altered rocks in the area east of Fortymile Wash or from infiltration of rainfall that dissolved near-surface calcium deposits.

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## **APPENDIX A**

Water chemistry data from samples collected between 1957 and 1977 from wells at and near the NTS. Data are from the USGS, with data sources listed in the text. Well names are consistent with those used in the 1991 edition of the Raytheon Services Nevada "NTS Drilling and Mining Summary." Wells are in order of increasing Nevada state north coordinate.

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Well Name	northing easting	date (m/d/y)	Hq	Ca	Mg	Na	×	Ð	SO4	НСО3	SiO <sub>2</sub>
								66.0	164	210	19.0
Wells-Stewart	651000 825000	05/01/64	7.6	33.0	18.0	0.621	12.0		5	196	66
Armu #3	654163 760615	01/28/59	7.9	27.0	21.0	6.8	1.4	0.0	11		
		05/03/61	7.8	35.0	23.0	5.6	1.2	5.0	15	507	14.0
	MICCI 700000	01010	80	24.0	16.0	222.0	11.0	23.0	169	486	9.3
Army 6A		00117140	20	340	17.0	13.0	2.5	6.10	17	197	20.0
USGS HTH #4		70/01/60	. r		210	37.0	52	16.0	53	256	21.0
Army #1 Water Well		70/01//0	1.1	0.14	010	38.0	49	17.0	58	254	20.0
Army #1 Water Well		79/01//0	Ċ, ţ	46.U	0.12	38.0	5.2	16.0	52	256	19.0
Army #1 Water Well	-	10/20/04	, r U r		10.01	0.00	1 C	67	15	<u>19</u>	14.0
<b>USGS HTTH #10</b>		02/26/63		0.16	10.0	1.1 A 0 A	100	50	14	201	15.0
<b>USGS HTH #10</b>		07/04/04	0./	0.10	0.01	1001 0	0.0	35.0	3599	98	22.6
Army 6		UNKUOWI	c	0.107	0.04	0.00	0.0 A A	32.0	4	199	32.0
USGS HTH "F"		19/02/20	ن. د و	0.61	0.1	0.44	50	110	181	278	31.0
USGS HTH "F"		00/11/07	5.1 0.0	09.0	0.00	0.00		0.11	74	121	26.0
J-12 Water Well	733508 581012	04/25/58	8.2	9.6	1.9	40.U	<b>7.</b> C	0.0	5 2	118	40.0
J-12 Water Well	733508 581012	02/19/59	7.4	14.0	1.5	42.0	4.4	0.0	\$ \$	011	
I-12 Water Well	733508 581012	03/31/62	8.0	14.0	2.5	39.0	0.0	0.0 0	5	(11 (4)	0.00
I 17 Water Well	733508 581012	05/26/64	7.4	17.0	0.9	40.0	4.6	7.4	17		0.00
Wheter Well 5A		04/04/57	8.6	3.2	0.0	160.0	6.2	12.0	25	346	0.80
		09/18/57	8.8	1.8	0.5	158.0	5.8	0.0	25	341	57.0
		NAMA/58	0.0	1.6	1.0	165.0	6.6	9.0	30	336	67.0
Water Well DA		05/190/00	<b>X</b> 0	<b>P</b> 6	00	158.0	6.0	0.6	24	344	58.0
Water Well 5A		60140170	0.0	7 04 7 <del>-</del>	0.0	163.0	5.6	11.0	27	345	49.0
Water Well 5A		70/77/20	000	5.1	104	163.0	6.4	11.0	27	384	50.0
Water Well SA		101000	) 0 0 F	020	14.0	157.0	16.0	20.0	484	102	67.0
J-11		10/01/60	0.7	0.00	13.0	1540	16.0	18.0	479	104	67.0
1-11					12.0	143.0	150	18.0	449	102	68.0
J-11		80/01/71	0.	0.70	0.01		2.1 2 2	10.0	36	266	56.0
Water Well 5C	742860 705888	04/01/57	8.1	3.6	1.0	0.021		0.01		763	55.0
Water Well 5C	742860 705888	09/18/57	8.8	1.8	0.7	128.0	0.0	0.2	\$ 2		0.02
Water Well SC	742860 705888	03/24/58	8.8	2.4	0.5	130.0	6.8	8.0	<b>q</b> ;	707	0.00
Water Well SC	742860 705888	12/09/58	8.5	1.6	1.0	126.0	5.8	10.0	24	<b>C07</b>	0.90
		-	8.7	3.1	0.5	128.0	5.6	8.9	25	258	53.0
			0	00	0.6	130.0	5.6	8.5	24	309	50.0
Water Well 5C	000CU/ 000274/		5	2	}	) ) 8	,				

ar the NTS, from USGS reports cited in the text. All concentrations are in mg/L. -. 110 ç • ,

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ppendix A. Chemical analysis for wells at and near the NTS, from USGS reports cited in the text. All concentrations are in mg/L (contin-	ued).
Append	

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Well Name	northing easting	date (m/d/y)	μd	Ca	Mg	Na	ĸ	D	SO4	HCO <sub>3</sub>	SiO <sub>2</sub>
	247360 T04762	DAIMIST	84	8.0	15	96.0	11.0	25.0	55	168	64.0
Waler well JB		12/12/100	00	5.6	1.0	102.0	10.0	24.0	59	155	16.0
		03/24/58	84	8.0	1.9	96.0	11.0	23.0	99	166	61.0
Water well 3B		02/22/60	- C	0.6	2.2	95.0	10.0	22.0	57	163	62.0
Water Well JD		06/13/64	4.8	7.8	1.8	94.0	11.0	22.0	57	178	54.0
Walch WCH JD		01/01/63	7.0	14.0	2.4	46.0	6.6	8.4	25	124	57.0
1 12 Water Well		05/25/64	7.1	14.0	1.8	48.0	5.0	7.4	23	136	58.0
1100-00 HTTH #2		05/10/62	7.3	51.0	21.0	83.0	7.6	23.0	25	328	24.0
		10/10/62	8.3	17.0	28.0	126.0	15.0	35.0	61	386	27.0
		06/14/64	6.9	74.0	29.0	123.0	14.0	34.0	8	576	29.0
		19/11/01	7.0	74.0	27.0	142.0	15.0	34.0	11	577	30.0
USUS Water Well C		01/19/62	7.2	70.0	19.0	130.0	12.0	23.0	75	560	30.0
USUS Water Well C		04/25/62	7.2	80.0	27.0	128.0	13.0	34.0	63	576	33.0
USUS Water Well C		10/16/64	6.7	47.0	26.0	119.0	14.0	35.0	2	472	30.0
Well 3		04/04/57	7.8	22.0	12.0	40.0	7.4	8.0	ជ	192	74.0
Well 3	-	09/18/57	8.0	19.0	13.0	39.0	7.4	6.0	21	<u>19</u>	74.0
Well 3		03/24/58	8.1	21.0	12.0	40.0	7.4	5.0	ជ	<b>1</b> 36	66.0
Well 3		12/02/58	7.8	19.0	14.0	37.0	7.4	5.0	21	190	72.0
WCII 3		12/19/61	7.5	20.0	11.0	39.0	7.6	6.5	23	189	76.0
Well 3		04/25/62	7.8	20.0	12.0	38.0	7.2	6.4	22	195	26.0
Well 3		05/27/64	7.9	22.0	11.0	38.0	8.0	5.6	19	196	64.0
I IF_16f Fleana		08/20/77	8.6	4.2	0.6	420.0	1.4	33.0	130	860	2.2
ULTIME Eleana		09/25/77	8.2	5.2	2.0	430.0	3.0	18.0	110	1000	7.2
11SGS Water Well A		09/21/60	7.6	28.0	4.6	51.0	8.7	6.0	18	212	62.0
11SGS Water Well A		04/25/61	8.0	22.0	7.4	53.0	8.8	6.5	21	206	81.0
11SGS Water Well A		12/19/61	7.6	22.0	6.5	50.0	8.4	6.0	23	207	82.0
11SGS Water Well A		06/13/64	7.8	23.0	6.6	50.0	8.4	5.6	18	212	72.0
HTTH "F"		03/31/60	9.0	1.6	0.0	81.0	2.6	6.0	16	187	61.0
Test Well #7		02/04/58	7.3	1.0	0.2	113.0	4.6	8.0	7	286	1.4
I IE-16d Eleana		05/23/77	7.8	68.0	30.0	28.0	7.8	9.2	49	270	18.0
I IF-16d Fleana		06/14/77	7.2	79.0	25.0	30.0	6.6	13.0	62	280	31.0
UE-16d Eleana		11/61/90	7.4	58.0	21.0	44.0	5.7	14.0	61	260	29.0

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Well Name	northing easting	date (m/d/y)	Hd	ß	Mg	Na	У	บ	SO4	HCO <sub>3</sub>	SiO <sub>2</sub>
			i i		00,	0 50 5	011	000	11	PLC.	18.0
C. Ilest Well	846600 672600	01/09/61	7.8	17.0	10.0	10/.01	0. <b>+</b> 1	0.07 1			45.0
1.TE1.8r	868100 564700	01/29/68	8.0	26.0	1.0	81.0	3.1	0.1	\$ :	707	0.05
118 HTTH 20211	876855 629310	10/01/60	7.6	2.0	0.0	36.0	1.0	7.0	11		0.46
		10/20/60	7.5	4.0	0.0	36.0	6.5	<b>6.</b> 3	12	78	38.0
14 UTU COCO		06/09/61	<u> </u>	21.0	11.0	62.0	1.7	13.0	59	180	20.0
1# HTH SOSO		00/00/01 08/11/67	77	20.0	11.0	35.0	3.2	8.7	35	150	19.0
<b>USGS HTH #1</b>		70/11/00	4 0 ¥	0.0 <b>7</b>	00	30.0	3.3	7.8	16	82	45.0
Water Well 8	-	10/CT/01	0.0	0.0 0 CC	19.0	24.0	5.6	L.L	23	186	53.0
Wair Well 2		70/07/100	7.6	30.05	10.0	28.0	6.6	5.6	21	190	46.0
Water Well 2		13/07/00	, r	0.00	011	080	16.0	17.0	65	357	8.5
UE-15d Water Well		10/71/11		0.04 10.0	12.0	0.00	15.0	11.0	¥	384	17.0
UE-15d Water Well		70/17/20	0.7	0.14	71	0.00	30	63	0	<b>9</b> 8	56.0
UE-19(S	900900 587084	08/18/65	8.1	0.11	1.0	0.72			Č	310	74.0
Marble #3	903096 674866	07/11/59	7.9	25.0	47.0	0.61	5.0 0.1	0.0	\$ 8		41.0
11-20a #2 Water Well	907448 568671	10/14/64	6.9	5.9	0.1	58.0	1.9	0.11	8	90 j	0.14
11_20ta #2 Water Well		03/10/66	T.T	6.1	0.1	55.0	0.2	11.0	12	9	46.0
Watertown 1		09/18/57	8.0	5.6	1.5	67.0	8.6	6.0	18	175	84.0
Watertown 1		04/25/58	8.2	4.8	2.4	68.0	8.8	6.0	53	171	86.0
		<b>DR/18/57</b>	8	5.6	12	86.0	9.2	6.0	19	215	90.0
		04/25/58	8.5	4.8	1.9	86.0	9.0	7.0	77	204	81.0
		03/08// 20	8.2	1.4	0.1	81.0	0.2	23.0	<del>4</del>	12	37.0
UE-201		D3/D8/66	83	1.4	0.1	83.0	0.1	24.0	42	120	46.0
UE-200	-		7.8	43	0.1	88.0	1.7	23.0	4	137	47.0
UE-204			8.4	21.0	0.1	68.0	0.5	8.8 8.9	53	143	52.0
			8.5	8.5	0.1	107.0	2.6	24.0	<del>4</del>	192	45.0
				5.0	0.1	75.0	0.5	7.0	20	8	39.0
		11/75/50	7.8	16.0	2.9	56.0	5.6	8.0	25	170	78.0
		VS/DU/SU	76	10	0.1	29.0	0.1	2.6	S	8	41.0
UE-19C Water Well			0.4	13.0	0.1	141.0	0.2	L.L	0	400	30.0
UE-190 Walci Well			0 0	0.8	0.1	. 69.0	0.6	6.6	30	103	36.0
U-201			01	04	0.1	0.69	0.8	7.0	23	8	C 55
107-O					0.1	113.0	2.0	40.0	48	164	47.0
UE-ZUI				0.6	10	640	1.8	15.0	30	107	49.0
UE-20h	14/100 010816	c0/07/90	1.0	2	•	) ;; )	}	1			

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Chemical analysis for wells at and near the NTS, from USGS reports cited in the text. All concentrations are in mg/L (co	
Appendix A.	

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Well Name	northing easting	date (m/d/y)	Hd	Ca	Mg	Na	К	ច	SO4	НСО3	SiO <sub>2</sub>
			6.0	00	01	2000	5.5	19.0	56	174	66.0
U-19aS	075095 347616	C0/10/00		) ( 1		0001	0 4	11.0	<i>LL</i>	205	66.0
11-10aS	919248 586326	06/07/65	8.4	1.2	0.1	189.0	0.0	0.14	1		010
0-01 II	010748 586376	06407/65	9.3	1.6	0.1	263.0	7.1	12.0	51	107	0.40
		UMD7165	74	0.5	0.1	50.0	12	11.0	ຊ	8	50.0
UE-19e	66606C 006176		6 0	77	0.1	43.0	0.8	3.7	16	8	56.0
UE-19e	66606C ME/76	10/10/00	4 C	AK D	1.7	138.0	64	115.0	135	150	44.0
UE-20j	928306 538531	+0/17/01	0.1		7.1	0.001	0.6	00	36	146	46.0
UE-19gS	931338 587843	03/27/65	8.2	17.0	0.1	0.00	, . ,		3 5	172	50.0
1 IE_190S	931338 587843	02/08/66	8.2	2.8	0.1	84.0	<b>C.I</b>	0.77	<del>.</del>		
	022700 606835	06/01/64	6.7	20.0	1.7	43.0	2.9	13.0	24	130	4/.0
UE-190#1		10/13/64	A L	24.0	24	42.0	3.0	6.8	21	150	41.0
UE-196#1	933/UU 00000	-DICT INT			ic	112.0	20	57.0	43	130	44.0
UE-20e #1	934466 560958	<b>40/CI/00</b>	1.1	4.0	1.0	0.211		0.00	Ę	110	36.0
1 IF-20e #1	934466 560958	03/08/66	8.5	0.2	0.1	0.68	<b>N</b> .7	20.02	;		
	045001 FMMM	03/24/64	7.5	29.0	2.9	173.0	3.6	39.0	63	474	0.07
7# 061-0		NAITCAN A	70	44.0	5,0	150.0	3.7	33.0	8	380	58.0
7# p61-0			- r		<b>V</b> 1	148.0	40	30.0	99	320	49.0
U-19d #2	945991 600202	00/17/00	0.1	0.67			, t r d		5	481	55.0
C# 101-11	945991 600202	03/09/66	7.9	58.0	2.8	153.0	3.7	19.0	8	104	
11_104 #7		03/09/66	7.9	57.0	2.8	153.0	4.3	20.0	57	489	0.66
1= 2/1-0											

\*value from Blankennagel and Weir (1973), reported as 7.1 by Robinson and Beetern (1965).

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# **APPENDIX B**

Water chemistry data from samples collected between 1983 and 1990 from wells at and near the NTS. Samples were collected and analyzed by DRI. Well names are consistent with those used in the 1991 edition of the Raytheon Services Nevada "NTS Drilling and Mining Summary." Wells are in order of increasing Nevada state north coordinate.

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Chemical analysis for wells and springs sampled at and near the NTS by DRI between 1983 and 1990. All concentrations are in mg/L. Appendix B.

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Well Name	northing easting	date (m/d/y)	hq	Ca	Mg	Na	X	อ	S04	HCO3	้งหวั
											15.0
	K20574 858034	12/18/87	7.99	32.40	20.50	5.8	1.50	3.3	13.4	0.761	0.01
Suver riag Aupua		19/2010	8 37	17,10	18.70	71.3	9.20	65.9	25.4	193.0	21.0
Divide Well		10/07/10	010	27.20	20.60	17.7	3.42	3.4	11.3	202.0	26.0
Divide Well		19/61/71	01.0	27.50	20.50	030	8.96	107.0	36.1	168.0	22.0
Old Dry Well		19/17/10	0.0			15.0	2.19	5.3	13.0	195.0	26.0
Old Dry Well		18/61/71	0.10	M.77	07 77	0111	623	30.6	319.0	224.0	3.0
Army 6A	665641 690214	06/25/86	8.64	10.30	00.21	0.411	70.0	21.9	310.0	228.0	3.0
Armv 6A	665641 690214	06/25/86	8.60	10.50	73.60	110.0		0.10	113	750.0	203
Army #1 Water Well	670902 684772	04/17/90	8.03	37.60	21.60	39.4	5.68	10.0	1.10	0.702	
A amount of the second s	670900 684772	06/06/20	7.88	44.10	20.40	37.9	5.70	15.7	C.2C	0.002	0.02
Allity #1 Walst Well		06/10/00	8.03	44.10	20.50	38.3	5.82	15.6	52.3	260.0	20.3
Army #1 water well			S M	15 50	2.23	39.1	5.20	6.8	21.6	118.0	60.3
J-12 Water Well		04101100	0000		0.75	67.2	2.15	14.7	14.1	168.0	33.0
UE-25 WT #12		00/C7/HO	0.0	14 10		65.7	233	7.3	26.1	163.0	47.0
UE-25 WT #12		04/28/88	<b>8.02</b>	14.10		135.0	212 212	60	32.7	272.0	54.8
Water Well 5C	742860 705888	06/01/60	8.81	CC7	0.79	0.001		, r i o	33.6	183.0	47.0
<b>USW WT-10</b>	748771 553302	05/16/88	8.42	2.72	0.07	93.8	1.14		0.00	121.0	673
I_13 Water Well	749209 579651	06/10/90	8.02	13.50	1.91	42.2	10.0	0.0	11.0	0.121	
1100 mm CI-f		06/16/90	8.27	48.80	19.70	79.6	9.18	23.2	<b>66.8</b>	343.0	1/.4
		09/26/88	7.94	55.00	21.60	26.4	4.28	11.5	54.2	218.0	0.80
		02/10/86	8 77	8.14	2.39	88.4	8.27	18.7	42.1	173.0	48.0
UE-5n		11/07/24	107	40.50	9.83	36.2	7.68	23.7	46.9	150.0	54.0
Pluto I			8 17	2.63	018	96.6	2.07	12.5	7.2	228.0	20.0
I-IM MSD		00/04/00	27.0		30.00	150.00	12.00	28.0	160.0	569.0	41.0
UE-25p #1*			2010	41.00	44 RU	18.4	3.66	13.6	63.7	287.0	16.0
Sand Spring		10/60/00	10.1	10.00	A2 50	18.8	457	16.2	24.1	220.0	14.0
Sand Spring		01/03/88	0.10	10.00		22.5	6.26	12.1	44.0	163.0	60.2
<b>UE-5c Water Well</b>		06/01/60	10.8	00.4	90.7		5 5 4	75	18.6	116.0	58.0
UE-25 WT #14	761651 575210	03/26/88	7.31	9.23	6/.0	47.4		ر. ر ۲۰ ۲	16.0	168.0	53.0
$11F_{-25}$ WT #15	766117 579806	04/23/88	7.62	11.60	1.71	61.8	00.4		1.01	100.0	0.02
	777130 768280	08/26/88	8.08	9.05	2.72	106.0	4.66	24.6	6.60	0.112	0.00
		06/18/20	8.12	26.40	8.21	48.9	5.26	11.8	40.9	159.0	60.2
Water Well #4	784999 687900	06/11/60	8.13	24.70	8.18	48.1	5.34	11.4	41.2	158.0	61.1
AV GUCA VALUE # 1											

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Chemical analysis for wells and springs sampled at and near the NTS by DRI between 1983 and 1990. All concentrations are in mg/L (continued). Appendix B.

Well Name	northing easting	date (m/d/y)	ЪН	Ca	Mg	Na	×	ū	SO4	HCO3	SiO <sub>2</sub>
											0
Water Well C_1	790011 692132	09/11/89	7.85	66.50	28.20	123.0	14.70	34.2	64.8	589.0	30.0
Weter Wall C 1		04/16/90	7.69	76.20	29.50	123.0	13.80	33.8	63.8	587.0	30.1
		07/31/90	7.78	46.50	28.30	124.0	12.90	33.3	60.9	576.0	29.5
		06/11/60	7.85	74.50	28.20	126.0	13.20	33.2	65.0	584.0	30.6
		00/11/80	1.61	74.10	30.00	124.0	15.00	34.0	65.6	585.0	30.0
USUS Water Well C		04/16/90	7.65	75.70	29.80	123.0	13.90	34.0	64.5	589.0	29.7
		06/12/00	7.64	74.30	29.20	124.0	15.70	33.8	63.9	589.0	31.5
USUS Water Well		06/11/00	7.79	73.90	28.20	127.0	13.20	33.2	65.0	581.0	29.8
USUS WALLI WCIL C		09/01/88	8.37	13.40	0.26	80.4	1.12	7.4	92.5	119.0	44.0
UE-140 TICCE Test Wall B		08/29/88	8.28	6.73	0.57	75.5	3.64	14.2	29.0	172.0	18.0
U NOUS ICSU WEIL D		04/16/90	8.32	6.00	0.58	1.11	3.37	18.6	23.4	167.0	18.3
		06/12/86	8.30	21.20	9.65	58.2	11.60	8.2	18.5	245.0	53.0
		06/12/86	9.73	2.09	1.59	87.4	11.20	34.7	50.8	85.3	2.1
OE-00		05/09/87	000	76.20	39.10	53.8	3.66	25.1	66.6	425.0	52.0
Quartz Spring		01/02/88	8.21	66.90	38.80	54.5	2.17	30.6	79.8	383.0	43.0
Vuant opning TE_166 Eleans		09/20/88	9.41	1.52	0.40	412.0	1.41	19.7	1.7	0.667	6.0
		08/24/88	8.01	41.10	32.40	59.1	12.30	30.7	1.0	448.0	19.0
		08/31/88	8.08	37.90	13.50	31.6	11.50	ĽL	20.3	248.0	85.0
		08/31/88	7.98	34.10	13.20	33.7	12.70	6.7	34.1	238.0	98.0
ULTIC I IE_16d Fleans		06/12/90	7.95	78.00	24.60	30.2	6.98	11.0	58.2	363.0	31.3
11SGS Test Well D		06/23/86	8.38	13.80	5.76	84.8	8.02	6.6	40.1	239.0	45.0
11SGS Test Well D		06/23/86	8.36	13.20	5.53	84.4	9.26	7.0	37.9	242.0	45.0
1 IE_18t		09/23/88	8.63	22.20	1.00	141.0	8.16	64.4	10.8	331.0	7.0
11E-18r		09/23/88	8.15	20.60	0.95	75.0	3.32	6.9	23.4	225.0	52.0
IISGS HTTH #1		07/24/86	8.29	1.51	0.16	52.0	1.29	5.8	5.7	127.0	13.0
1# HTTH SOCO		07/28/87	00.6	1.46	0.20	48.2	0.30	3.2	5.6	96.1	23.0
1# HLH 3000		07/28/87	8.68	2.03	0.20	48.0	0.50	3.2	1.0	117.0	3.0
1# HLH SUSII		04/25/89	9.18	1.02	0.07	49.4	0.25	3.1	6.2	91.4	19.6
118GS HTH #1		04/25/89	9.14	1.02	0.07	49.3	0.25	3.1	6.2	96.9	19.7
<b>USGS HTTH #1</b>		04/26/89	9.28	1.37	0.07	50.3	0.22	3.1	6.0	6.18	21.6

Chemical analysis for wells and springs sampled at and near the NTS by DRI between 1983 and 1990. All concentrations are in mg/L (continued). Appendix B.

Well Name	northing easting	date (m/d/y)	Hd	Ca	Mg	Na	К	ธ	SO4	нсоз	SiO <sub>2</sub>
14 חידט אוז	876855 670310	04/26/89	9.27	1.02	0.07	50.6	0.25	3.1	6.2	88.1	19.8
		09/24/90	8.76	9.83	0.32	53.1	1.13	3.4	17.4	116.0	10.7
1# HIH \$9\$0		00/74/00	9.15	1.59	0.24	51.6	0.38	3.3	8.5	82.7	20.1
USGS HIH #1		00/17/00	915	1.91	0.78	52.1	0.79	3.3	8.8	80.7	19.8
USGS HIH #1		00/56/00	9.74	1.49	0.16	51.8	0.38	3.2	8.4	85.9	21.5
USGS HTH #1		06/54/00	100	1.38	0.16	51.4	0.41	3.3	8.6	78.5	20.0
USGS HTH #1	010670 000010	0612760	9.05	1 70	0.18	51.6	0.34	3.2	7.6	83.7	22.0
USGS HTH #1		06167160	918	2.12	0.22	51.4	0.77	3.6	0.9	93.5	2.5
USGS HIH #1		04/16/00	7 38	8.74	1.22	29.9	3.48	<i>L.</i> L	15.2	81.3	47.6
		0/01/00	092	8.05	1.21	30.4	3.71	7.6	15.0	76.6	49.0
Water Well 8		08/03/00	7 80	44.8	1.23	28.8	3.36	7.5	15.0	75.8	46.7
Water Well 8		00/11/00	7.81	8 44	125	28.5	3.42	7.2	14.7	76.0	47.4
Water Well 8		06/11/00	00 8	36.90	16.00	30.1	7.18	7.6	25.2	226.0	52.0
Water Well 2	00000000000000000000000000000000000000	00/11/60	7.88	36.20	15.40	30.0	6.80	ĽL	23.7	226.0	51.1
Water Well 2		06,01,000	8 07	35.80	14.40	28.7	7.27	7.5	24.3	225.0	51.8
Water Well 2			20.0 2 2 3	00.CC	0.38	209.0	21.50	10.2	42.8	543.0	62.0
UE-10115 #3			7C 00	47 80	15.10	78.8	14.60	15.0	43.0	394.0	27.0
UE-15d Water Well		00/11/60	7 83	58.40	15.70	45.1	16.80	14.7	32.5	327.0	74.8
UE-15d Water Well	400700 60/060	06/01/20	101	56.70	14.60	47.9	15.10	14.6	36.2	332.0	60.3
UE-15d Water Well	-	10/08/00	101	10.00	1.80	102.0	5.78	94.4	18.7	145.0	49.4
U-19az	010090 169106	10/06/50	8 1 A	882	1 24	38.0	1.90	3.2	8.1	114.0	47.0
U-20ao	000003 000200		8.31	19.60	0.72	190.0	15.90	114.0	131.0	209.0	55.0
Panute Mesa #3			00 0	1.14	1.53	1200.0	10.80	318.0	222.0	1005.0	13.0
Papule Mesa #2			8 34	19.20	0.74	190.0	15.60	114.0	137.0	209.0	55.0
Panule Mesa #3			8.29	22.70	0.60	148.0	10.40	95.7	126.0	157.0	46.0
Panute Mesa #2			9.83	1.43	1.55	1150.0	8.51	313.0	283.0	1086.0	15.0
Fallule Mesa #3			6612	33.20	1.46	135.0	10.50	98.1	129.0	152.0	61.0
Panute Mesa #3			8.05	34.00	1.51	134.0	10.90	98.9	130.0	149.0	61.0
Pehrute Mesa #3		-	8.12	34.40	1.45	134.0	10.90	99.3	131.0	149.0	61.0
Fahute Mesa #3			9.09	3.48	1.03	78.6	3.84	24.8	23.7	132.0	42.0

Appendix B. Ch in 1	Chemical analysis for wells and springs sampled at and near the NTS by DRI between 1983 and 1990. All concentrations are in mg/L (continued).	vells and spri	ngs samp	iled at an	d near the	NTS by I	)RI betwe	en 1983 an	d 1990. All c	oncentration	s are
Well Name	northing easting	date (m/d/y)	Hq	Ca	Mg	Na	К	อ	SO4	нсо <sub>3</sub>	SiO <sub>2</sub>
The Man #2	006077 530008	10/05/88	9.18	3.55	1.03	77.3	2.85	23.9	24.5	123.0	42.0
Pallute Mesa #3	800055 CUMU	10/05/88	8.78	15.80	1.36	138.0	10.20	88.4	127.0	102.0	4.0
Paliuic Meco #2		10/05/88	8.31	12.20	0.57	110.0	6.13	63.6	88.9	128.0	5.0
Pallulic Mesa #3 Debuta Mesa #3		10/20/88	9.17	3.30	0.30	43.2	0.80	4.3	10.3	65.5	38.2
Pahute Mesa #3		10/20/88	9.57	3.20	5.69	994.0	10.80	232.0	273.0	1220.0	5.8
Pahinte Mesa #3		10/27/88	8.04	34.20	1.39	138.0	11.40	97.0	123.0	155.0	61.5
Dahuta Mesa #3		10/27/88	8.14	34.90	1.42	141.0	11.60	95.0	122.0	165.0	60.2
Paluute Massa #2		10/28/88	8.17	34.00	1.39	137.0	11.40	97.5	124.0	153.0	60.8
Pallule Mesa #3		05/17/89	7.94	30.20	0.82	141.0	10.90	94.0	130.0	159.0	61.2
Pallute Mesa #3 Debute Mesa #3		05/17/89	7.95	28.40	0.79	140.0	12.60	93.5	129.0	159.0	61.1
Pallule Mesa #2		05/17/89	7.98	30.10	0.60	137.0	16.60	92.9	125.0	164.0	60.0
r 700 #7 Wotow Wall	007448	04/01/88	8.03	6.18	0.23	59.4	1.73	11.8	32.6	111.0	51.0
U-20d #2 Water Well	007448	04/10/88	8.27	6.34	0.24	62.6	2.27	11.2	38.4	112.0	52.0
U-20a #2 walls well	010582	05/23/87	8.12	6.38	0.28	57.3	1.67	12.0	31.4	111.0	48.0
11 20 Water Well		04/16/90	8.06	5.65	0.22	58.4	1.83	11.9	28.9	113.0	48.9
17 20 Water Well		08/02/90	8.25	5.40	0.67	58.3	1.78	11.4	30.7	111.0	48.2
U-20 Walci Well		06/11/60	8.33	6.15	0.44	57.0	1.74	11.4	30.7	107.0	49.2
U-20 Mater Well		03/12/85	8.43	4.29	1.05	115.0	7.17	63.5	26.0	175.0	58.0
1 IF_19c Water Well		04/16/90	8.52	1.29	0.10	35.0	0.38	2.6	6.6	70.2	45.2
1 IF-19c Water Well		08/07/90	8.51	1.42	0.22	30.2	0.79	2.4	5.8	67.1	44.2
1 IE-10c Water Well		06/11/60	8.59	1.54	0.25	32.9	0.76	2.4	6.1	64.9	45.4
11_20al		11/08/84	8.30	23.70	2.68	122.0	11.10	32.8	77.6	250.0	58.0
		11/08/84	8.43	13.10	2.05	117.0	11.00	30.5	68.0	213.0	59.0
11_10ha		11/06/90	7.81	20.70	1.13	78.3	6.06	39.0	9.7	181.0	49.4
11_10ha		11/06/90	8.06	20.50	1.09	77.8	6.08	43.9	9.7	181.0	49.1
U-19ba	926950 602900	11/27/90	8.07	20.80	1.16	0.67	5.47	40.9	10.2	189.0	47.2

emical analysis for wells and springs sampled at and near the NTS by DRI between 1983 and 1990. All concentrations are Ē q 11-

\*analysis reported by Benson and McKinley (1985)

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# **APPENDIX C**

Combined chemical dataset of samples collected from 1957 to 1990, with averages calculated for locations where multiple samples have been collected. The data in this table were used to construct the maps in Figures 11, 13, 16, and 17. Well names are consistent with those used in the 1991 edition of the Raytheon Services Nevada "NTS Drilling and Mining Summary." Wells are grouped according to producing formation and ordered within each group according to increasing north coordinate.

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Well Name	# of Samples	northing	easting	Hq	Ca	Mg	Na	м	ប	SO4	НСО3	SiO <sub>2</sub>
Formation: Alluvium	4											
rui IIIIIIIII. Alaha Cile Elee Alaha	-	630574	858034	8.0	32.4	20.5	5.8	1.5	3.3	13	197	15.0
ouver riag Aupua	- c	642680		8.2	19.7	19.7	44.5	6.3	34.7	18	198	23.5
	4 C	100000		84	22.3	21.3	54.0	5.6	56.2	25	182	24.0
UID LITY Well	1.	101010			22.0	18.0	123.0	12.0	55.0	164	210	19.0
Wells-Stewart		000100		0.0	0.00	210.0	0.044	1 4	60	=	186	6.6
Army #3	-	654163		6./ C -	0.12	0.12	0.0			15	20	14.0
Army #2	1	655582		7.8	35.0	23.0	0.0	7.1	0.0			1 Y
Army 6A	e	665641	690214	8.4	15.0	54.1	150.7	8.0	28.8	10	داد 22	1.0
	-	680800	690500	8.1	281.0	90.0	1291.0	0.0	35.0	3599	86	22.6
Water Well SC		742860	705888	8.7	2.4	0.6	129.0	6.0	9.1	8	271	55.1
Water Well 5R	. v	747359	704263	8.5	T.T	1.7	9.96	10.6	23.2	58	166	51.4
rate for	·	754460	706415	8.8	8.1	2.4	88.4	8.3	18.7	42	173	48.0
ULTON 1180-S Water Wall A	• •	833000		7.8	23.8	6.3	51.0	8.6	6.0	20	209	74.3
USUS WALLI WULL A	, c	814000		0.6	14.1	5.6	72.8	11.4	21.5	35	165	27.6
Uztertour 3 Watertour 3	۰ <del>-</del>	914990	742272	7.8	16.0	2.9	-56.0	5.6	8.0	22	170	78.0
Formation: Carbonate	-	20022		75	34.0	17.0	13.0	2.5	6.1	17	197	20.0
USGS HIH #4	\	0/9600		. r	0.40	000	38.1	54	16.1	53	257	20.1
Army #1 water well	0	7060/0			0.75	12.5	01	14	59	15	198	14.5
<b>USGS HTH #10</b>	- (	1011/0	C10461		0.10	101 101	0.7 213	84	23.1	75	336	20.7
<b>USGS HTH #3</b>	7	6810C/			C.C+	1.07	150.0	10.01	28.0	160	5695	41.0
UE-25p #1	-	1/196/		0.0	0.001	0.40	2 01	11	14.0	WV	754	15.0
Sand Spring	6	757105		8.3	6.67	7 <b>1</b> 6	10.0	<b>1</b> . <b>1</b>		5	222	N OC
Water Well C-1	9	110062		7.7	59.1	28.5	124.2	13.9	4.66 2.00	8 (		1.02
USGS Water Well C	œ	790082		7.4	71.1	27.0	127.1	14.0	32.6	67	8	C.05
Ouartz Spring	7	815654		8.2	71.6	39.0	54.2	2.9	27.9	73	<del>4</del>	47.5
TE-16f Fleans	<b>(</b> 1	832355		8.7	3.6	1.0	420.7	1.9	23.6	81	553	5.1
	) <b></b>	837000		8.1	37.9	13.5	31.6	11.5	T.T	20	248	85.0
05 10 11E_16	l <del>y</del>	837000		8.0	34.1	13.2	33.7	12.7	6.7	\$	238	98.0
UL-IC I IF-16d Fleana	4	844878		7.6	70.8	25.2	33.1	6.8	11.8	62	293	27.3
LINGS Test Well D	·	846600	_	8.2	14.7	7.1	92.1	10.4	11.2	50	251	36.0
Water Well 7	. v	88000		7.9	32.2	15.0	28.2	6.7	7.2	23	210	50.8
ITE 154 Woter Well		895709	2	10	40.8	130	73.2	15.5	14.5	<b></b>	359	37.5
						101		5		!		

,

Well Name	# of Samples	northing	easting	Hq	Ca	Mg	Na	×	Ū	SO4	НСО3	SiO <sub>2</sub>
Formation: Volcanic				ť	3.04	15 5	81 U	80	215	108	239	31.5
USGS HTTH "F"	7	731033	601100	U		0 5	0.10	2.0 1 2	76	3	110	\$0.3
J-12 Water Well	Ś	733508	581012	7.8	14.0	9.1	41.4	1.0		3 2		2 2 2
Water Well 5A	9	738361	707514	8.8	2.3	0.5	161.2	6.1	10.2	<u>୧</u> :	£ ;	
TTE_25 WT #12	2	739726	567011	8.1	12.9	0.1	66.5	2.2	11.0	20	100	40.0 1
		740890	611821	7.8	83.7	13.3	151.3	15.7	18.7	471	103	67.3
-11	⊲ -	748771	263302	48	2.7	0.1	<b>9</b> 3.8	1.1	7.8	\$	183	47.0
		000072	570651	7.3	13.9	2.1	45.6	5.8	L.L	23	126	58.6
J-15 Water Well	• •	152607	667558	7.9	55.0	21.6	26.4	4.3	11.5	2	218	58.0
		200001	667481	80	40.5	9.8	36.2	ĽL	23.7	47	150	54.0
	-	ULSSSL	552801	<b>8</b>	2.6	0.2	96.6	2.1	12.5	7	228	20.0
USW W1-/	4	010001	LOOUL	58	01	2.1	83.5	6.3	12.1	4	163	0
UE-JC Walet Well	-	133132	575710	0:0 6 L	6.0	0.8	42.4	5.5	7.5	19	116	58.0
UE25 W I #14		10010/	STORAS	76	116	1.7	61.8	4.6	11.5	16	168	53.0
UE23 W I #13		111001	708280	81	9.1	2.7	106.0	4.7	24.6	99	217	58.0
UD-118 Motor Wall #A	- 7	000782	687900	<b>8</b> .1	25.6	8.2	48.5	5.3	11.6	41	159	0
Mauch Would HT	• •-	794100	650111	8.4	13.4	0.3	80.4	1.1	7.4	93	119	44.0
ULTITO Tact Wall R	• •	812044	690713	8.3	6.4	0.6	76.3	3.5	16.4	8	170	0
USUS ICSU WCH D		817795	677762	7.8	20.4	12.1	38.7	7.5	6.1	21	193	64.6
WCII 3		UUULE8	000099	8.0	41.1	32.4	59.1	12.3	30.7		<b>448</b>	19.0
UE-18 UT-18 %E"		830080	696001	0.6	1.6	0.0	1.0	2.6	6.0	16	187	61.0
		601202	684654	7.3	1.0	0.2	113.0	4.6	8.0	7	286	1.4
LESU WELL # /	4	865800	598400	8.6	22.2	1.0	141.0	8.2	64.4	11	331	7.0
UL-101	• ~	868100		8.1	23.3	1.0	78.0	3.2	7.4	24	239	48.5
118028 HTTH #1	<b>,</b> [2	876855	629310	9.1	4.9	1.5	60.09	1.3	4.8	12	105	20.4
	- -	870468	600009	7.4	8.5	1.1	29.6	3.4	7.6	15	78	46.8
Walet well 0		887201	680700	8.3	7.8	0.4	209.0	21.5	10.2	43	543	62.0
	•		587084	81	11.0	1.6	29.0	3.0	6.3	6	88	56.0
UD-1910	• ~	001894	586010	8.0	19.9	1.8	102.0	5.8	94.4	19	145	49.4
U-1342		005600	556450	8.1	8.8	1.2	38.0	1.9	3.2	œ	114	47.0
0-20a0 Dahiita Meca #3	- v	906022	530998	8.6	20.0	1.3	277.8	10.	112.9	129	292	43.4
r autite Mesa #3 1170a #7 Water Well	; (	907448	568671	1.1	6.1	0.2	58.8	1.5	11.3	32	109	48.0
U-200 #2 Want Want	1	0							4			1

Appendix C. Combined data set of 1957 to 1990 data, with averages calculated for wells with multiple samples (continued).

	# of		·		ć	77-	No	ž	כ	0	ЧСОН	SiQ
Well Name	Samples	northing	easung	нd	C.	INI	PVI	4	3	400	60011	4
Formation: Volcanic (continued)	(continued)											
	~	9000K)	752226	8.4	5.2	1.6	86.0	9.1	6.5	21	210	85.5
Walciuwu 2	1 V		554300	87	73	0.1	85.4	1.0	20.6	4	143	45.4
UE-200	<del>-</del> د	010000	502107	7.0 L L	<b>6 5</b>	0.1	75.0	0.5	7.0	20	<u>8</u>	39.0
UE-191		010507		8.7 2	0 4	104	57.8	1.8	11.7	30	111	0
U-ZU Water Well		700016	202020	7.0	- <b>7</b>		115.0	72	63.5	26	175	58.0
U-20ai	- 1	007516	106000		n r r c		526	20	35	~	134	41.2
UE-19c Water Well	7	000/16	00102/	8.2	3.1	7.0	<b>0.</b>			) <b>;</b>		2 02
(J-20aL	(1	917335	549200	8.4	18.4	2.4	119.5	11.1	31./	51	767	U.00
11_20F		917825	551857	9.0	0.6	0.1	69.0	0.7	6.8	21	101	37.5
		017825	552007	7.2	4.8	0.1	113.0	2.0	40.0	<b>4</b> 8	164	47.0
	4 -	012015	LALTAS	81	06	0.1	64.0	1.8	15.0	30	107	49.0
	-	C10010			1.6	10	2173	61	14.0	35	213	72.0
U-19aS	<b>r</b>	947616		7.0	1.0	1.0					101	2 01
[1-19ha	6	926950		8.0	20.7	1.1	78.4	5.9	41.3	10	184	48.0
UTE 10.	· c	927300		7.8	2.1	0.1	46.5	1.0	7.4	18	83	53.0
	•	028306		7.0	46.0	1.2	138.0	6.4	115.0	135	150	44.0
	• (	021238		8.7	74	0.1	76.0	1.1	15.5	4	135	48.0
UE-1983	4 C		20100	11	0.00	10	2.04	3.0	6.6	23	140	44.0
UE-190#1	7	M/CCK	CC0000		0.77				200	54	175	U UV
UE-20e #1	6	934466	560958	8.1	0.3	0.1	C.16	7.7	0.00	f	3	??? ?
	ч	0.45001	<b>CUCUNA</b>	7.8	43 A	30	155.4	3.9	28.2	65	419	49.0

Appendix C. Combined data set of 1957 to 1990 data, with averages calculated for wells with multiple samples (continued).

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