

Prepared in cooperation with the National Park Service

Groundwater Flow, Quality (2007–10), and Mixing in the Wind Cave National Park Area, South Dakota

Scientific Investigations Report 2011–5235

U.S. Department of the Interior
U.S. Geological Survey

Front cover. Fluorescein dye in Wind Cave in Windy City Lake, a lake that merged with Calcite Lake before the end of 1999. Dye was injected by National Park Service staff into What the Hell Lake in February 2008 and was detected in Windy City Lake 2 months later. This photograph was taken November 2008 by Peter Sprouse, National Park Service.

Back cover. Upper left: Measuring the physical properties of water from Stairway Spring.
Upper right: Leading us to the water in Wind Cave! Photograph by Jason Walz, National Park Service.
Lower left: Squeezing through a narrow place above What the Hell Lake in Wind Cave. Photograph by Jason Walz, National Park Service.
Lower right: Downloading temperature data collected from Highland Creek. Photograph by Jason Walz, National Park Service.

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By Andrew J. Long, Marc J. Ohms, and Jonathan D.R.G. McKaskey

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Conversion Factors

SI to Inch/Pound

Multiply	By	To obtain
Length		
meter (m)	3.281	foot (ft)
kilometer (km)	0.6214	mile (mi)
Volume		
liter (L)	33.82	ounce, fluid (fl. oz)
liter (L)	2.113	pint (pt)
liter (L)	1.057	quart (qt)
liter (L)	0.2642	gallon (gal)
liter (L)	61.02	cubic inch (in ³)
Flow rate		
cubic meter per second (m ³ /s)	35.31	cubic foot per second (ft ³ /s)
meter per day (m/d)	3.281	foot per day (ft/d)

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

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

Vertical coordinate information is referenced to the North American Vertical Datum of 1988 (NAVD 88).

Horizontal coordinate information is referenced to the North American Datum of 1983 (NAD 83).

Altitude, as used in this report, refers to distance above the North American Vertical Datum of 1988 (NAVD 88) unless otherwise specified.

Water year (WY) is the 12-month period, October 1 through September 30, and is designated by the calendar year in which it ends. Thus, the water year ending September 30, 2006, is called the “2006” water year.

Specific conductance is given in microsiemens per centimeter at 25 degrees Celsius (µS/cm at 25°C).

Concentrations of chemical constituents in water in this report are given in milligrams per liter (mg/L) or micrograms per liter (µg/L).

Abbreviations and Acronyms

δ ² H	stable isotope of hydrogen
δ ¹⁸ O	stable isotope of oxygen
CFC	chlorofluorocarbon
³ H	tritium
PCA	principal components analysis
RPD	relative percent difference
USEPA	U.S. Environmental Protection Agency

Groundwater Flow, Quality (2007–10), and Mixing in the Wind Cave National Park Area, South Dakota

By Andrew J. Long,¹ Marc J. Ohms,² and Jonathan D.R.G. McKaskey¹

Abstract

A study of groundwater flow, quality, and mixing in relation to Wind Cave National Park in western South Dakota was conducted during 2007–11 by the U.S. Geological Survey in cooperation with the National Park Service because of water-quality concerns and to determine possible sources of groundwater contamination in the Wind Cave National Park area. A large area surrounding Wind Cave National Park was included in this study because to understand groundwater in the park, a general understanding of groundwater in the surrounding southern Black Hills is necessary. Three aquifers are of particular importance for this purpose: the Minnelusa, Madison, and Precambrian aquifers. Multivariate methods applied to hydrochemical data, consisting of principal component analysis (PCA), cluster analysis, and an end-member mixing model, were applied to characterize groundwater flow and mixing. This provided a way to assess characteristics important for groundwater quality, including the differentiation of hydrogeologic domains within the study area, sources of groundwater to these domains, and groundwater mixing within these domains. Groundwater and surface-water samples collected for this study were analyzed for common ions (calcium, magnesium, sodium, bicarbonate, chloride, silica, and sulfate), arsenic, stable isotopes of oxygen and hydrogen, specific conductance, and pH. These 12 variables were used in all multivariate methods. A total of 100 samples were collected from 60 sites from 2007 to 2010 and included stream sinks, cave drip, cave water bodies, springs, and wells.

In previous approaches that combined PCA with end-member mixing, extreme-value samples identified by PCA typically were assumed to represent end members. In this study, end members were not assumed to have been sampled but rather were estimated and constrained by prior hydrologic knowledge. Also, the end-member mixing model was quantified in relation to hydrogeologic domains, which focuses model results on major hydrologic processes. Finally, conservative tracers were weighted preferentially in model calibration, which distributed model errors of optimized values, or

residuals, more appropriately than would otherwise be the case. The latter item also provides an estimate of the relative effect of geochemical evolution along flow paths in comparison to mixing. The end-member mixing model estimated that Wind Cave sites received 38 percent of their groundwater inflow from local surface recharge, 34 percent from the upgradient Precambrian aquifer, 26 percent from surface recharge to the west, and 2 percent from regional flow. Artesian springs primarily received water from end members assumed to represent regional groundwater flow.

Groundwater samples were collected and analyzed for chlorofluorocarbons, dissolved gasses (argon, carbon dioxide, methane, nitrogen, and oxygen), and tritium at selected sites and used to estimate groundwater age. Apparent ages, or model ages, for the Madison aquifer in the study area indicate that groundwater closest to surface recharge areas is youngest, with increasing age in a downgradient direction toward deeper parts of the aquifer. Arsenic concentrations in samples collected for this study ranged from 0.28 to 37.1 micrograms per liter ($\mu\text{g/L}$) with a median value of 6.4 $\mu\text{g/L}$, and 32 percent of these exceeded 10 $\mu\text{g/L}$. The highest arsenic concentrations in and near the study area are approximately coincident with the outcrop of the Minnelusa Formation and likely originated from arsenic in shale layers in this formation. Sample concentrations of nitrate plus nitrite were less than 2 milligrams per liter for 92 percent of samples collected, which is not a concern for drinking-water quality. Water samples were collected in the park and analyzed for five trace metals (chromium, copper, lithium, vanadium, and zinc), the concentrations of which did not correlate with arsenic. Dye tracing indicated hydraulic connection between three water bodies in Wind Cave.

Introduction

Established in 1903, Wind Cave and the surrounding land became the eighth national park in the United States and the first one created to protect a cave (Wind Cave National Park, 2008). Wind Cave is located in western South Dakota (fig. 1) and is the fifth longest cave in the world with 218 kilometers (km) of accessible passages (Gulden, 2011). Groundwater is an important resource for the park and, in particular, for

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Wind Cave. Groundwater drips from the ceiling of the cave at numerous locations and exists in ponds, lakes, and streams on the cave floor. The water table of the Madison aquifer is accessible at the deepest part of this cave, where subterranean lakes exist. These lakes are in hydraulic connection with the regionally extensive Madison aquifer, which exists in the northern Great Plains in several States, including South Dakota, North Dakota, Montana, Wyoming, Saskatchewan, and Manitoba. To understand the hydrology and hydrochemistry of Wind Cave, it is essential to understand the surrounding Madison aquifer within which the cave exists.

The National Park Service is concerned about water quality and possible sources of current (2011) or future groundwater contamination in Wind Cave National Park. To address these concerns, the U.S. Geological Survey in cooperation with the National Park Service conducted a study during 2007–11. The objective of this study was to characterize groundwater flow, quality, and mixing in the Wind Cave National Park area (fig. 1). This objective was carried out by assessing or estimating (1) groundwater gradients, flow directions, and transit times; (2) the occurrence and geospatial distributions of arsenic and nitrate and possible associations with trace metals; (3) groundwater sources for Wind Cave and sources of arsenic in the park, and (4) the relative proportions of different source waters contributing to Wind Cave and other areas of interest.

Purpose and Scope

The purpose of this report is to document results of a 4-year study of groundwater flow, quality, and mixing within and surrounding Wind Cave National Park in western South Dakota. Analytical results from a total of 100 samples collected from 2007 to 2010 from 60 selected sites including stream sinks, cave drip, cave water bodies, artesian and shallow springs, and wells are presented. Samples collected for this study were analyzed for common ions, arsenic, stable isotopes of oxygen and hydrogen, specific conductance, and pH. Data from these samples were used in principal component analysis (PCA), cluster analysis, and end-member mixing, which are described in this report. Samples from selected sites also were analyzed for nitrate plus nitrite to determine if concentrations were of concern and for trace metals to assess possible existence of unusually high concentrations or correlations of these metals with arsenic. Samples were analyzed for chlorofluorocarbons (CFCs), dissolved gasses, and tritium for the purpose of estimating groundwater age, or residence time. Dye tracing using a fluorescent dye was used to determine the hydraulic connectivity between water bodies in Wind Cave.

This report provides information that is useful for understanding sources of groundwater to the park and the hydrologic interactions between the park's groundwater and surrounding groundwater. This report will be useful for identifying possible sources of contamination if such events occur in the future. Water-quality concerns, primarily related to arsenic, are addressed, and possible sources are described.

Description of Study Area

The study area is located in the southern part of the Black Hills of western South Dakota, which is a dome-type structure with sedimentary layers of Paleozoic age dipping radially outward on the flanks (fig. 1, table 1). Detailed geologic and hydrogeologic descriptions can be found in Gries and Martin (1981), Strobel and others (1999), and Redden and DeWitt (2008). Hydrochemical groundwater and cave studies in the Black Hills area include Back and others (1983), Bakalowicz and others (1987), Alexander and Davis (1989), Palmer and Palmer (1989), Ford and others (1993), Naus and others (2001), and Heakin (2004). Back (2011) investigated the hydrochemistry of Wind Cave National Park and interpreted data collected during 2007; these 2007 data are included in the analyses described in this report.

Underlying Paleozoic sedimentary layers and exposed at the central core of the Black Hills are Precambrian-age fractured metamorphic and igneous rocks, within which the Precambrian aquifer is contained (unit pC; fig. 1, table 1). The Paleozoic Formations overlie the Precambrian rocks and are exposed in roughly concentric rings surrounding the central core of the Black Hills. The Cambrian- and Ordovician-age Deadwood Formation is the lowermost Paleozoic Formation and contains the Deadwood aquifer (unit O&D; fig. 1). The Devonian-age Englewood Limestone and Mississippian-age Madison Limestone overlie the Deadwood Formation. The Madison aquifer (unit MDme; fig. 1) is a mature karst aquifer contained within the regionally extensive Madison Limestone (locally called the Pahasapa Limestone) and the Englewood Limestone, which are referred to as the Madison Limestone for the remainder of this report. The Madison aquifer contains fractures and solution-enlarged caves, and at least 15 large Black Hills caves are contained within this formation (Greene and Rahn, 1995). Wind Cave National Park maintains a database of more than 250 caves in the Black Hills that are more than 8 meters (m) in length. The lower part of the Madison Limestone and the underlying Englewood Limestone generally have lower permeability than the upper part of the Madison aquifer in the Black Hills (Greene, 1993). Overlying the Madison Limestone is the Pennsylvanian- and Permian-age Minnelusa Formation (fig. 1), which contains the Minnelusa aquifer. The Minnelusa Formation is composed of interbedded sandstone, limestone, dolostone, and shale, with solution openings in carbonate layers including caves as much as 1,500 m in length (table 1). At the base of the Minnelusa Formation is a red clay shale that varies between 0 and 15 m thick in some areas of the Black Hills and is a residual weathered soil developed on the surface of the Madison Limestone (Cattermole, 1969; Gries, 1996). Overlying the Minnelusa Formation is the Permian-age Opeche Shale, which is a confining unit that underlies the Permian-age Minnekahta Limestone and the Permian- and Triassic-age Spearfish Formation (fig. 1). Overlying the Spearfish Formation are formations described in table 1, including the Tertiary-age White River Group, which contains the White River aquifer that overlies the Madison

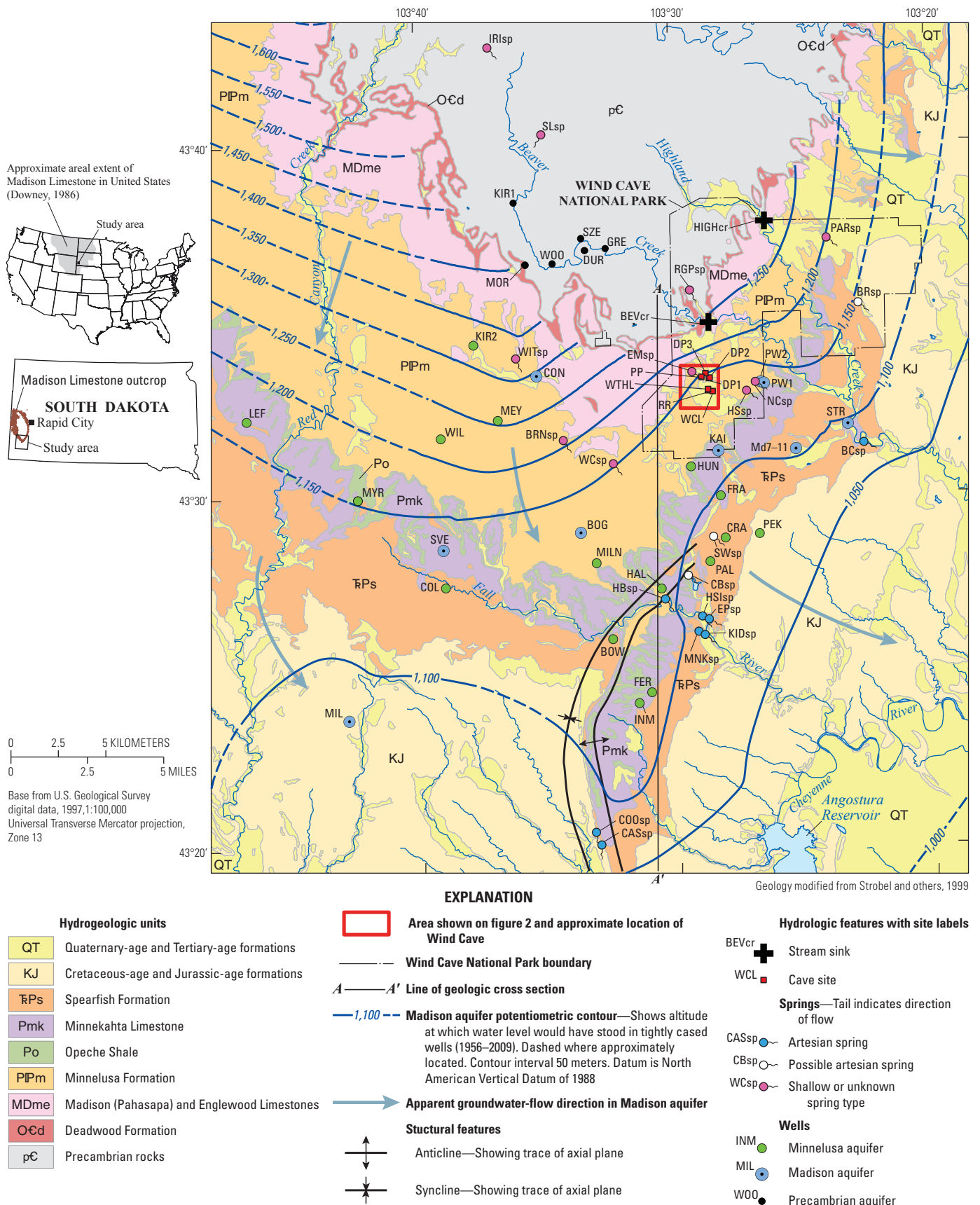


Figure 1. Study area showing hydrogeologic units and potentiometric surface of the Madison aquifer (geology modified from Strobel and others, 1999).

and Minnelusa aquifers in some areas and might contribute recharge to these aquifers.

Wind Cave is located in the southeastern Black Hills, where sampled cave sites consist of cave drip (sites DP1, DP2, DP3) and subterranean water bodies (sites WCL, WTHL, PP, and RR; figs. 1 and 2, table 2). Land-surface altitudes in the study area range from about 1,700 m above the North American Vertical Datum of 1988 (NAVD 88) in the northwest to about 1,000 m in the southeast, and streams generally flow to the south and southeast (fig. 1). Surface recharge to the Minnelusa and Madison aquifers (units PIPm and MDme; fig. 1) occurs on outcrop areas of the Minnelusa and Madison Formations. Additionally, streams flowing across these outcrop areas sink, fully or partially, into solution openings and fractures, primarily into the Madison aquifer and secondarily into the Minnelusa aquifer. Highland Creek flow rates at site HIGHcr (fig. 1) ranged from 0.02 to 0.05 cubic meters per second (m^3/s) from 2002 to 2010 [30 measurements during this time; Wind Cave National Park written commun., (2010)]. Daily streamflow measurements for Beaver Creek at station BEVcr from 1990 to 2009 ranged from 0 to 2.4 m^3/s with a mean of 0.06 m^3/s (U.S. Geological Survey, 2011). The

watersheds upstream from the Beaver Creek and Highland Creek (site HIGHcr) sites are almost entirely within Precambrian rocks.

Collectively, artesian springs are a large source of groundwater discharge in the study area. Eight artesian springs are located in the southeast part of the study area (fig. 1, table 2), including five along the Fall River (sites HBsp, HSIsp, KIDsp, MNKsp, EPsp), two near the southern boundary of the study area (sites CASsp and COOsp), and one near Beaver Creek (site BCsp) at the eastern boundary. Flow from Cascade Springs (site CASsp) ranged from 0.37 to 0.71 m^3/s with a mean of 0.55 m^3/s from 1976 to 1995 (station 06400497; U.S. Geological Survey, 2011).

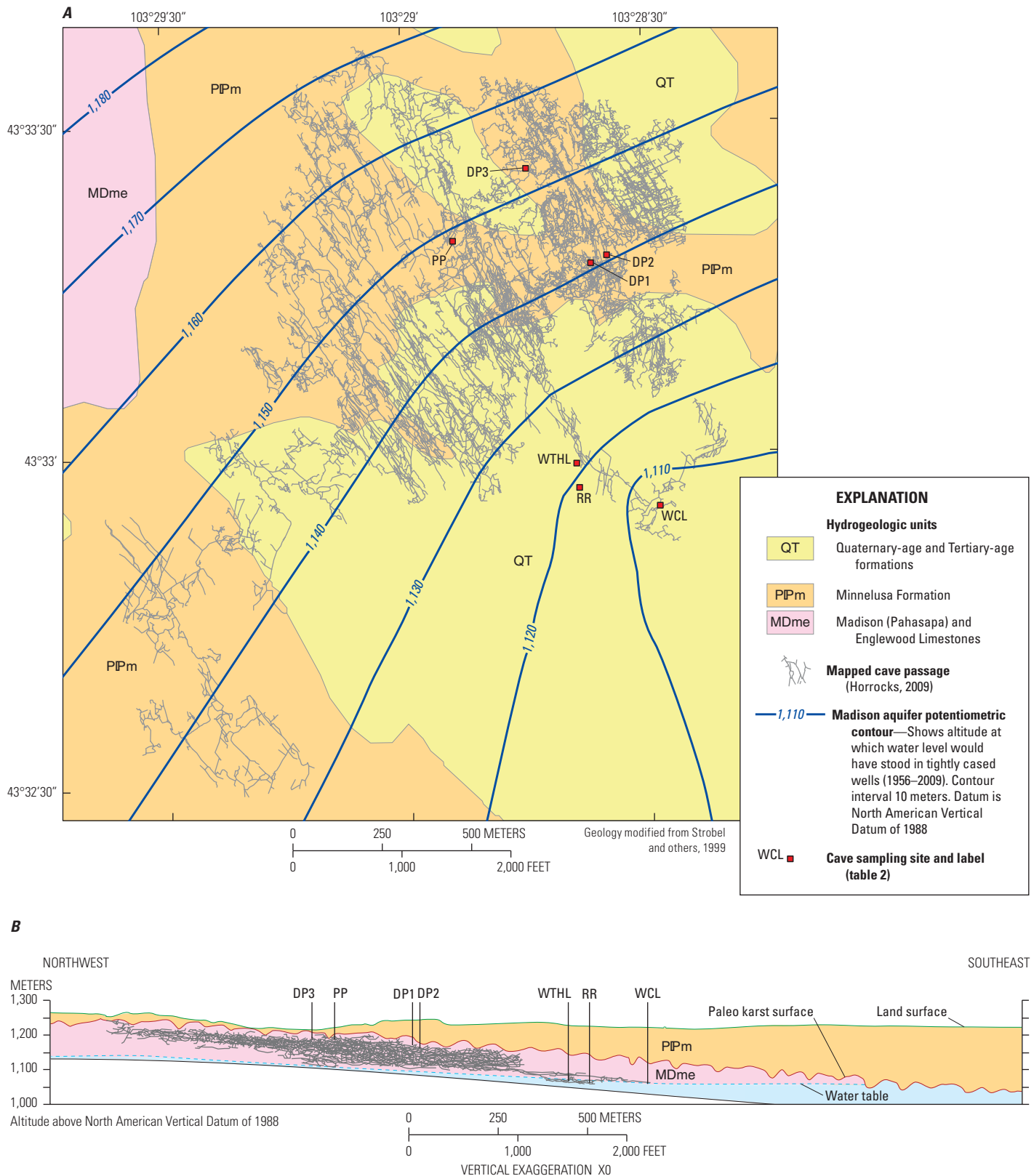
Artesian springs generally emerge from or near outcrop areas of the Spearfish Formation (fig. 1), which generally has low permeability because of high shale content but also has flowing groundwater in fractures, cavities created by dissolved gypsum, and vertical breccia pipes. According to Hayes (1999), the throats of artesian springs in the southern Black Hills probably are breccia pipes that allow groundwater from deep bedrock aquifers to emerge from overlying formations. Cascade Springs (site CASsp), Cool Spring (site

Table 1. Description of hydrogeologic units.

[Summarized from Strobel and others, 1999; Redden and DeWitt, 2008]

Unit label (fig. 1)	Formation and description	Aquifer name or hydrogeologic description	Pore water type	Thickness (meters)	Age
Qt	Alluvium, colluvium, gravel, claystone, poorly indurated sandstone, conglomerate	Unconsolidated aquifers	Intergranular	0–140	Quaternary and Tertiary.
KJ	Undifferentiated shale, limestone, sandstone, conglomerate, siltstone, claystone, gypsum	Aquifers and confining units	Intergranular, fractures	800–2,300	Cretaceous and Jurassic.
TRs	Spearfish Formation - shale with interbedded sandstone, siltstone, and gypsum	Low permeability in general ^a	Fractures, intergranular, dissolved gypsum	110–240	Triassic and Permian.
Pmk	Minnekahta Limestone	Minnekahta aquifer	Fractures, solution openings	10–20	Permian.
Po	Opeche Shale	Confining unit	Intergranular	10–50	Permian.
PIPm	Minnesula Formation - interbedded sandstone, limestone, dolostone, and shale	Minnelusa aquifer	Fractures, intergranular, solution openings	110–360	Permian and Pennsylvanian.
MDme	Madison (Pahasapa) and Englewood Limestones - also contains dolostone	Madison aquifer, mainly in upper part	Solution openings, fractures	90–320	Mississippian and Devonian.
OEd	Deadwood Formation - glauconitic sandstone, shale, siltstone, and conglomerate	Deadwood aquifer	Fractures, intergranular	0–150	Ordovician and Cambrian.
pC	Fractured metamorphosed quartzite and metagraywacke with isolated areas of granite and pegmatite	Precambrian aquifer	Fractures	Unknown	Precambrian.

^a Except for fractured areas and areas of dissolved gypsum cavities.



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Table 2. Sampled cave sites and springs.

[Δh , estimated height in meters (m) of the Madison aquifer hydraulic head above (positive values) that of the Minnelusa aquifer; NAVD 88, North American Vertical Datum of 1988; --, not applicable]

Site name	Descriptive name	Source of flow	Hydrogeologic domain	Δh (m)	Altitude of site, in meters above NAVD 88
Cave drip sites					
DP1	Cave drip 1 (C53)	Madison aquifer	East	--	--
DP2	Cave drip 2 (UD17)	Madison aquifer	East	--	--
DP3	Cave drip 3 (NP38A)	Madison aquifer	East	--	--
Cave water body sites					
WCL	Windy City Lake	Madison aquifer	East	--	1,105 ^a
WTHL	What the Hell Lake	Madison aquifer	East	--	1,127
PP	Petey's puddle	Madison aquifer	East	--	--
RR	Rebel River	Madison aquifer	East	--	1,118
Artesian spring sites					
HBsp	Hot Brook Spring	Minnelusa aquifer ^b	Artesian 1	-20	1,106
HSIs	Hot Springs Intake Spring	Madison aquifer ^c	Artesian 1	10	1,065
KIDsp	Kidney Spring	Madison aquifer ^c	Artesian 1	20	1,066
MNKsp	Minnekahta Spring	Madison aquifer ^c	Artesian 1	20	1,079
EPsp	Evans Plunge Spring ^d	Madison aquifer ^c	Artesian 1	10	1,063
BCsp	Beaver Creek Spring	Madison aquifer ^c	Artesian 2	50	1,260
CASp	Cascade Springs	Madison aquifer ^c	Artesian 2	10	1,049
COOsp	Cool Spring	Madison aquifer ^c	Artesian 2	20	1,052
BRsp	Boland Ridge Spring	Unknown ^e	Artesian 2	40	1,127
SWsp	Stairway Spring	Unknown ^e	Artesian 2	10	1,143
CBsp	Cold Brook Spring	Unknown ^e	Artesian 2	-20	1,127
Shallow spring sites					
PARsp	Parker Spring	White River aquifer	East	30	1,269
HSsp	Horse Shelter Spring	Minnelusa aquifer	East	-60	1,207
NCsp	Negro Canyon Spring	Minnelusa aquifer	East	-40	1,184
BRNsp	Brown Spring	Minnelusa aquifer	West	-170	1,378
EMsp	Elk Mountain Spring	Minnelusa aquifer	West	-110	1,330
WCsp	Woodcock Spring	Minnelusa aquifer	West	-140	1,293
WITsp	Witch Spring	Minnelusa aquifer	West	-90	1,409
IRIs	Irish Spring	Precambrian aquifer	PC	--	1,645
RGPsp	RG Pegmatite Spring	Precambrian aquifer	PC	--	1,351
SLsp	Southerland Spring	Precambrian aquifer	PC	--	1,621

^a Altitude at which staff gage equals zero (surveyed May 28, 2008).

^b Assumed to flow primarily from Minnelusa aquifer but could include water from other aquifers.

^c Assumed to flow primarily from Madison aquifer but could include water from other aquifers.

^d No sample available.

^e Possibly artesian flowing from Madison or Minnelusa aquifer or both.

COOsp), Hot Brook Spring (site HBsp), Evans Plunge Spring (site EPsp), and Beaver Creek Spring (site BCsp) are considered artesian and probably flow primarily from the Madison aquifer (Rahn and Gries, 1973; Back and others, 1983; Hayes, 1999; Naus and others, 2001; Driscoll and others, 2002). Consistent with these previous studies identifying the Madison aquifer as the primary source of artesian springs is an assessment of vertical hydraulic-head gradients described in this report, where the Madison aquifer was estimated to be higher than in the Minnelusa aquifer for all artesian springs except Hot Brook Spring (table 2).

In addition to the previously classified springs, Hot Springs Intake Spring (site HSIsp) and Kidney Spring (site KIDsp) were classified in this study as artesian (previously unclassified) because of similarity in temperature and hydrochemistry to those of the other artesian springs near the Fall River (fig. 1). Three springs (sites BRsp, SWsp, and CBsp) are not known to be artesian but are hydrochemically and geologically similar to known artesian springs, and thus the Madison aquifer and possibly the Minnelusa aquifer might be a source of flow for these springs (table 2). Also, breccia pipes that do not extend to the land surface probably exist in the study area and facilitate groundwater exchange between the Madison, Minnelusa, and possibly other overlying aquifers (Brobst and Epstein, 1963; Hayes, 1999).

Methods

Assessment of hydraulic gradients and flow direction were used in interpreting hydrochemical analyses. Plotting the geospatial distributions of arsenic concentrations, nitrate plus nitrite concentrations, specific conductance values, and groundwater age were useful for characterizing hydrochemical differences and gradients across the study area and provided supplemental information for assessing groundwater flow. Multivariate methods to estimate groundwater mixing consisted of principal component analysis (PCA), cluster analysis, and end-member mixing applied to hydrochemical data. Application of these multivariate methods presented in this report is a summary of results described by Long and Valder (2011).

Sample Collection and Hydrochemical Data

Sites that were sampled for this study include stream sinks; cave drip; cave water bodies; artesian springs primarily flowing from the Madison aquifer; shallow springs flowing from the White River, Minnelusa, and Precambrian aquifers; and wells open to the Minnelusa, Madison, and Precambrian aquifers (fig. 1, table 3). A total of 100 samples were collected from 60 sites from 2007 to 2010, and 93 percent of the sites were sampled at least once during 2009 and 2010. Multiple samples were collected during different months or years for

19 sites, with as many as 5 samples per site from 2007 to 2010, and were used to assess seasonal and annual variability. Five samples were collected from 5 of the 19 sites (four samples during 2007 and one during 2009), and 2 to 4 samples were collected from 2007 to 2009 for the other 14 sites.

Samples collected for this study were analyzed for common ions (calcium, magnesium, sodium, bicarbonate, chloride, silica, and sulfate), arsenic, stable isotopes of oxygen ($\delta^{18}\text{O}$) and hydrogen ($\delta^2\text{H}$), specific conductance, and pH. These 12 variables were used in principal component analysis (PCA), cluster analysis, and end-member mixing. Also, samples collected during 2007 were analyzed for nitrate plus nitrite (table 3). Samples analyzed for chlorofluorocarbons (CFCs), dissolved gasses (argon, carbon dioxide, nitrogen, and oxygen), and tritium (^3H) were collected at selected sites in the study area (table 3) for the purpose of estimating groundwater age, or residence time. Hydrochemical data for all sites described in this report as well as other sites in the study area, including historical data, are available from the National Water Information System (U.S. Geological Survey, 2011).

Samples were collected according to methods described in U.S. Geological Survey (variously dated). Samples were analyzed for common ions and arsenic by the U.S. Geological Survey National Water Quality Laboratory in Lakewood, Colorado, using methods described by Fishman and Friedman (1989), Fishman (1993), and Garbarino and others (2006). Values of $\delta^{18}\text{O}$ and $\delta^2\text{H}$ were determined by the U.S. Geological Survey Reston Isotope Laboratory in Reston, Virginia, using a gaseous hydrogen equilibration procedure described by Révész and Coplen (2008a, 2008b). Samples were analyzed for CFCs and dissolved gasses at the Reston Chlorofluorocarbon Laboratory in Reston, Virginia. CFCs were analyzed using a purge-and-trap gas chromatography procedure (Shimadzu GC-8AIE) with an electron capture detector, and dissolved gases (argon, carbon dioxide, nitrogen, and oxygen) were determined by gas chromatography (Hewlett Packard model 5890; <http://water.usgs.gov/lab/>). Samples were analyzed for tritium at the Tritium Laboratory in Menlo Park, California, by electrolytic enrichment and liquid scintillation (Thatcher and others, 1977).

Quality-control samples consisted of 12 pairs of replicate samples and 5 field-equipment blank samples. The relative percent difference (RPD) is defined as the difference between the replicate and sample values divided by the sample value times 100. The RPDs for each variable for each replicate pair was 6 percent or less for 96 percent of the data. Larger RPD values resulted from one sodium pair (14 percent RPD), one silica pair (9 percent RPD), and two arsenic pairs (11 and 23 percent RPDs). The largest RPD (23 percent) was for a replicate pair with low arsenic concentrations of 0.25 and 0.28 $\mu\text{g/L}$. The conservative tracers, chloride, $\delta^{18}\text{O}$, and $\delta^2\text{H}$, had RPDs of less than 3, 1, and 1 percent, respectively. Laboratory analyses of all field-equipment blank samples were below detection limits.

Table 3. Water-quality sampling sites and constituents analyzed in samples.

[SC, specific conductance; CFC, chlorofluorocarbons; --, not applicable or not analyzed]

Site name	Descriptive name	Station number	Station name	Source aquifer	Number of samples ^a	Constituents analyzed in samples						
						Common ions, arsenic, stable isotopes, SC, pH	Nitrate plus nitrite	Trace metals	Tritium	CFC	Dissolved gases	Fluorescein dye
Stream sink sites												
BevCr	Beaver Creek sink	06402430	Beaver Creek near Pringle, S. Dak.	--	3	Yes	Yes	Yes	Yes	--	--	--
HIGHcr	Highland Creek sink	433745103261900	Highland Creek above Madison outcrop near Pringle, S. Dak.	--	5	Yes	Yes	--	--	--	--	--
Cave drip sites												
DP1	Cave drip 1 (C53)	433302103281507	6S 5E12DBAB7	Madison	2	Yes	Yes	--	--	--	--	--
DP2	Cave drip 2 (UD17)	433302103281508	6S 5E12DBAB8	Madison	3	Yes	Yes	Yes	--	--	--	--
DP3	Cave drip 3 (NP38A)	433302103281509	6S 5E12DBAB9	Madison	3	Yes	Yes	Yes	Yes	--	--	--
Cave water body sites												
PP	Petey's puddle	433302103281504	6S 5E12DBAB4	Madison	1	Yes	Yes	--	--	--	--	--
RR	Rebel River	433302103281506	6S 5E12DBAB6	Madison	1	Yes	Yes	--	--	--	--	Yes
WCL	Windy City Lake	433302103281501	6S 5E12DBAB	Madison	2	Yes	Yes	--	--	--	--	Yes
WTHL	What the Hell Lake	433302103281502	6S 5E12DBAB2	Madison	2	Yes	Yes	--	Yes	Yes	--	--
Artesian spring sites												
HBsp	Hot Brook Spring	432703103302801	7S 5E10DCBA	Minnelusa ^b	5	Yes	Yes	Yes	Yes	Yes	Yes	--
BCsp	Beaver Creek Spring	433128103223401	6S 6E14CDB	Madison ^c	5	Yes	Yes	--	Yes	Yes	Yes	--
CASsp	Cascade Springs	432006103330501	8S 5E20CDAB	Madison ^c	3	Yes	--	--	Yes	Yes	Yes	--
COOsp	Cool Spring	432028103331601	8S 5E20BDCB	Madison ^c	1	Yes	--	--	Yes	Yes	Yes	--
HSIsP	Hot Springs Intake Spring	432632103285302	7S 5E14DDDA	Madison ^c	5	Yes	Yes	--	--	Yes	Yes	--
KIDsp	Kidney Spring	432605103285401	7S 5E14DDD	Madison ^c	1	Yes	--	--	Yes	Yes	Yes	--
MNKsp	Minnekahta Spring	432605103290901	7S 5E14DCD	Madison ^c	1	Yes	--	--	Yes	Yes	Yes	--
BRsp	Boland Ridge Spring	433525103224401	5S 6E26BBCD	Unknown ^d	1	Yes	--	Yes	--	--	--	--
CBsp	Cold Brook Spring	432744103293401	7S 5E11BAAB	Unknown ^d	1	Yes	--	--	--	--	--	--
SWsp	Stairway Spring	432849103283201	6S 5E36CDBB	Unknown ^d	1	Yes	--	--	--	--	--	--

Table 3. Water-quality sampling sites and constituents analyzed in samples.—Continued

[SC, specific conductance; CFC, chlorofluorocarbons; --, not applicable or not analyzed]

Site name	Descriptive name	Station number	Station name	Source aquifer	Number of samples ^a	Constituents analyzed in samples						
						Common ions, arsenic, stable isotopes, SC, pH	Nitrate plus nitrite	Trace metals	Tritium	CFC	Dissolved gases	Fluorescein dye
Shallow spring sites												
PARsp	Parker Spring	433717103235401	5S 6E15BBAC	White River	1	Yes	--	Yes	--	--	--	--
BRNsp	Brown Spring	433137103342101	6S 5E18CABD	Minnelusa	1	Yes	--	--	--	--	--	--
EMsp	Elk Mountain Spring	433332103291801	6S 5E 2ACBD	Minnelusa	1	Yes	--	Yes	--	--	--	--
HSsp	Horse Shelter Spring	433258103270801	6S 6E 7BAAC	Minnelusa	1	Yes	--	--	--	--	--	--
IRIs	Irish Spring	434249103370401	4S 4E11CCBB	Minnelusa	1	Yes	--	--	--	--	--	--
NCsp	Negro Canyon Spring	433312103264701	6S 6E 6DCAA	Minnelusa	1	Yes	--	Yes	--	--	--	--
WCsp	Woodcock Spring	433056103322201	6S 5E21BCBC	Minnelusa	1	Yes	--	--	--	--	--	--
WITsp	Witch Spring	433611103335801	5S 4E35DDBC	Minnelusa	1	Yes	--	--	--	--	--	--
RGPsp	RG Pegmatite Spring	433551103291901	5S 5E23DBCD	Precambrian	1	Yes	--	Yes	--	--	--	--
SLsp	Southerland Spring	434020103350101	4S 4E25DB	Precambrian	1	Yes	--	--	--	--	--	--
Well sites												
BOW	--	432555103323201	7S 5E20AACA	Minnelusa	1	Yes	--	--	--	--	--	--
COL	--	432727103390201	7S 4E 9BACA	Minnelusa	1	Yes	--	--	--	--	--	--
CRA	--	432846103280501	6S 5E36DCBC	Minnelusa	1	Yes	--	--	--	--	--	--
FER	--	432437103305701	7S 5E27CCCD	Minnelusa	1	Yes	--	--	--	--	--	--
FRA	--	432437103305701	7S 5E27CCCD	Minnelusa	1	Yes	--	--	--	--	--	--
HAL	--	432720103303701	7S 5E15BABBB	Minnelusa	1	Yes	--	--	--	--	--	--
HUN	--	433034103284701	6S 5E23DB	Minnelusa	4	Yes	Yes	--	Yes	Yes	Yes	--
INM	--	432332103314801	7S 5E33ABDD	Minnelusa	1	Yes	--	--	--	--	--	--
KIR2	--	433420103374901	5S 4E34BDDB	Minnelusa	1	Yes	--	--	--	--	--	--
LEF	--	433215103464401	6S 3E 5CCD	Minnelusa	1	Yes	--	--	--	--	--	--
MEY	--	433215103365801	6S 4E11CCDB	Minnelusa	1	Yes	--	--	--	--	--	--
MILN	--	432806103330801	8S 5E 5CADD	Minnelusa	1	Yes	--	--	--	--	--	--
MYR	--	433003103420701	7S 3E36CBCC	Minnelusa	1	Yes	--	--	--	--	--	--
PAL	--	432806103284101	7S 5E 1CBCC	Minnelusa	1	Yes	--	--	--	--	--	--
PEK	--	432852103264401	6S 6E31DBDA	Minnelusa	1	Yes	--	--	--	--	--	--

Table 3. Water-quality sampling sites and constituents analyzed in samples.—Continued

[SC, specific conductance; CFC, chlorofluorocarbons; --, not applicable or not analyzed]

Site name	Descriptive name	Station number	Station name	Source aquifer	Number of samples ^a	Constituents analyzed in samples							
						Common ions, arsenic, stable isotopes, SC, pH	Nitrate plus nitrite	Trace metals	Tritium	CFC	Dissolved gases	Fluorescein dye	
Well sites—Continued													
WIL	--	433141103390901	6S 4E16CABB	Minnelusa	1	Yes	--	--	--	--	--	--	--
BOG	--	432858103334201	6S 5E31DA	Madison	2	Yes	Yes	--	Yes	Yes	--	--	--
CON	--	433326103352001	6S 4E 1DB	Madison	2	Yes	Yes	--	Yes	Yes	Yes	--	--
KAI	--	433114103281601	6S 5E24BAAA	Madison	2	Yes	Yes	--	--	--	--	--	--
Md7–11	--	433115103251401	6S 6E21BBBB (CU91A)	Madison	1	Yes	Yes	--	Yes	Yes	Yes	--	--
MIL	--	432340103421501	7S 3E36CBDC	Madison	2	Yes	Yes	--	--	Yes	--	--	--
PW1	--	433311103263101	6S 6E 6DDA	Madison	2	Yes	Yes	--	Yes	Yes	Yes	Yes	Yes
PW2	--	433311103263102	6S 6E 6DDA2	Madison	2	Yes	Yes	--	Yes	Yes	Yes	Yes	Yes
STR	--	433150103230501	6S 6E15ABDD	Madison	5	Yes	Yes	--	Yes	Yes	Yes	--	--
SVE	--	432825103391201	7S 4E 4BAC	Madison	1	Yes	Yes	--	Yes	Yes	Yes	--	--
DUR	--	433658103332301	5S 5E17BCDB	Precambrian	1	Yes	--	--	--	--	--	--	--
GRE	--	433701103323401	5S 5E17ACDD	Precambrian	1	Yes	--	--	--	--	--	--	--
KIR1	--	433821103360901	5S 4E 2DDBD	Precambrian	1	Yes	--	--	--	--	--	--	--
MOR	--	433635103354301	5S 4E13CCAC	Precambrian	1	Yes	--	--	--	--	--	--	--
SZE	--	433718103333101	5S 5E17BBBA	Precambrian	1	Yes	--	--	--	--	--	--	--
WOO	--	433636103343901	5S 5E18CCAC	Precambrian	1	Yes	--	--	--	--	Yes	--	--

^a Not including quality-control replicate samples.^b Assumed to flow primarily from Minnelusa aquifer but could include water from other aquifers.^c Assumed to flow primarily from Madison aquifer but could include water from other aquifers.^d Possibly artesian flowing from Madison or Minnelusa aquifer or both.

Principal Component and Cluster Analyses

Principal component analysis (PCA) is a linear transformation of data in multidimensional space, where the transformed axes, or principal components, align with the greatest variances in the data (Davis, 2002). Each principal component is a newly created variable, which is a linear combination of all original variables. PCA is a method that is used to graphically plot complex multivariate datasets and elucidate data patterns that otherwise might not be noticed. The term “scores” refers to the values of the new variables in the transformed space, and by plotting the data points as scores in this space, sample relations and groupings may become evident. PCA commonly is used to identify extreme-value points, which might be considered as possible end members (Davis, 2002).

The assignment of data points to a specified number of groups, or clusters, based on similarity of data is referred to as cluster analysis. The method described in this report partitions the data points by iteratively assigning each data point to a cluster that minimizes the sum of Euclidian distances between data points and the nearest cluster centroid (Seber, 1984; Spath, 1985). The scores from the PCA were used as the data from which clusters were generated to reduce the clustering error caused by data error or multicollinearity (Suk and Lee, 1999).

End-Member Mixing

For an end-member mixing analysis, it is assumed that each water sample consists of water from one or more end members in varying proportions. An end member is defined as water having a characteristic geochemical signature that best represents a source of groundwater inflow to the system. An end member represents the hydrochemistry of a particular source of groundwater inflow, such as recharge within the study area or groundwater flowing into the study area. In some cases, an end member might be a point source of inflow, such as a sinking stream; in other cases, it might represent the integration, or characteristic hydrochemical signature, of a distributed source, such as areally distributed recharge or regional groundwater inflow. A two end-member model adapted from Fritz and others (1976) is described as

$$\hat{c} = f_1 E_1 + f_2 E_2, \quad (1)$$

where \hat{c} is the concentration of a mixed water sample, f_1 and f_2 are the fractions, or mixing proportions, of end-members 1 and 2, respectively, and E_1 and E_2 are the respective end-member concentrations.

A common approach in applying an end-member mixing model is to collect samples from assumed end-member waters and to then determine the mixing proportion, or contribution, of each end member in samples assumed to contain mixed water. In this study, it was not assumed that end members had been sampled, but rather these were estimated by inverse

modeling using a generalized form of equation 1 that allows for any number of end members and variables:

$$c_{i,j} = \sum_{k=1}^n f_{i,k} E_{j,k}, \quad (2)$$

where $c_{i,j}$ is the concentration of variable j for site i ; $f_{i,k}$ is the fraction, or mixing proportion, of end-member k that is associated with site i ; and $E_{j,k}$ is the end-member concentration for variable j and end-member k . Equation 2 was programmed in Fortran, and the mixing proportions $f_{i,k}$ and end-member hydrochemical values $E_{j,k}$ were estimated by inverse modeling using the parameter optimization software, PEST (Doherty, 2005), which uses optimization methods described by Levenberg (1944) and Marquardt (1963). This process began with user-specified initial estimates for the values of $f_{i,k}$ and $E_{j,k}$. Then, the calculated concentrations $\hat{c}_{i,j}$ were compared to observed values $c_{i,j}$, and the differences, or residuals, between calculated and observed values $\hat{c}_{i,j} - c_{i,j}$ were minimized by optimizing the values of $f_{i,k}$ and $E_{j,k}$ iteratively. New residuals were calculated for each iteration, and $f_{i,k}$ and $E_{j,k}$ were adjusted for the next iteration until no further reduction of residuals occurred.

The primary limitation of this model is that hydrochemical evolution of groundwater along a flow path is neglected. For the current study, this limitation was diminished by weighting the calibration data for conservative tracers more heavily than for other variables during inverse modeling.

Groundwater Age Dating

Samples were collected and analyzed for CFCs and tritium for the purpose of estimating groundwater age, or the amount of time that a sampled groundwater has been out of contact with the atmosphere. This also is known as groundwater residence time and is important because it provides estimates of groundwater transit times, if greater than about 5 to 10 years. The CFCs are anthropogenic tracers that are present in the atmosphere and have become assimilated with rainwater and surface water (Cook and others, 1996; Oster and others, 1996; Plummer and Busenberg, 2000). The CFCs then enter groundwater with infiltrating precipitation or sinking streams. Groundwater age can be estimated by comparing concentrations in groundwater samples to historical atmospheric concentrations for different CFC species, including CFC-11, CFC-12, and CFC-113. The CFCs in water exposed to the atmosphere become equilibrated with atmospheric concentrations, but in groundwater that is not in contact with the open atmosphere, CFCs generally remain at the concentration of recharge water, except in reducing environments. Atmospheric tritium concentrations in past decades were elevated when CFC concentrations were relatively low, and both of these were at low concentrations or not present in the atmosphere before about 1950 (fig. 3).

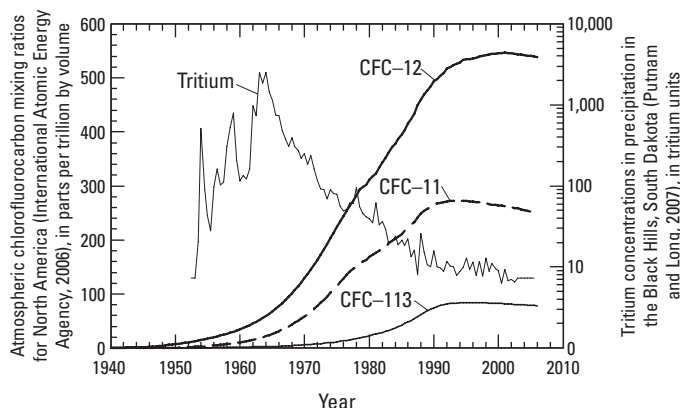


Figure 3. Chlorofluorocarbon (CFC) and tritium concentrations in the atmosphere.

Groundwater Flow, Quality, and Mixing Assessments

Results of this study are described first in relation to general groundwater flow, the understanding of which is necessary for interpreting the results of methods applied to hydrochemical data. This assessment consists of a description of groundwater hydraulic gradients and flow directions, sources and flow rates of springs, and dye tracing. Potentiometric maps of the Madison and Minnelusa aquifers (figs. 1 and 4, respectively) were constructed and used to estimate general groundwater-flow directions, horizontal hydraulic gradients in the Madison and Minnelusa aquifers, and vertical hydraulic gradients between these two aquifers. Next, the geospatial distributions and other relations of water-quality constituents in the study area are described. Geospatial distributions of hydrochemical data primarily indicated differences between surface recharge areas with young groundwater and deep aquifer areas with older groundwater. Finally, an assessment of groundwater mixing is described, where five hydrogeologic domains and groundwater flow between these domains were characterized on the basis of multivariate analyses of hydrochemical data.

Groundwater Flow

The mapped passages of Wind Cave are almost entirely within the Madison aquifer's unsaturated zone and below an unsaturated part of the Minnelusa aquifer (fig. 2). Infiltration of precipitation on the outcrop of the Minnelusa Formation seeps downward, enters unsaturated parts of the Madison aquifer, and drips from the ceiling of Wind Cave at numerous sites. Similarly, this type of recharge to the Madison aquifer from the Minnelusa aquifer is assumed also to occur elsewhere in the study area. Windy City Lake (site WCL; figs. 1 and 2) is a body of water at the deepest part of the cave and exists because of this hydrologic connection between the Madison

and Minnelusa aquifers. Windy City Lake rises and declines with the surrounding water table as indicated by a comparison of water levels in the lake and well Md7-11 (fig. 5). Before 1999, another similar lake existed in proximity to Windy City Lake called Calcite Lake. Sometime before the end of 2000, the Madison aquifer water table and the levels of both lakes increased in altitude, which resulted in the merging of the two lakes.

What the Hell Lake (site WTHL; figs. 1 and 2) was filled unexpectedly in 1996 after being dry for decades (Shouse, 2004). Wind Cave National Park staff observed this water body on July 18, 1996, and stated that a new pool had formed, which did not exist during a previous visit to the same site on June 12, 1996, and this site has contained water continuously from that time until the present (Wind Cave National Park, written commun., 2011). The filling of site WTHL might have been the result of perched groundwater in caverns above this site that emptied rapidly. Perched water frequently is observed in Black Hills caves and results from areas of impermeable limestone or the accumulation of clay sediments on cave floors (Long, 2009). Rapid outflow from a perched water body might have been the result of spillover through an outlet as water levels were rising or from the dislodgement of clay that had filled an outlet. Years 1995–2000 was a period in which the water level at site WCL was rising (fig. 5), which indicates relatively high recharge and infiltration rates. Groundwater in a karst aquifer unsaturated zone can be decades old in some cases (Even and others, 1986), and if perched groundwater above site WTHL had been in storage for decades and then released, this could explain why site WTHL was dry for decades before filling.

Groundwater Gradients and Flow Directions

A potentiometric-surface map of the Madison aquifer was constructed on the basis of water-level data available from the U.S. Geological Survey (2011) and helps to characterize groundwater flow (fig. 1). Apparent groundwater-flow directions are perpendicular to potentiometric contours, but these might not be the actual flow directions because of possible anisotropic permeability resulting from cave and fracture orientations (Greene and Rahn, 1995). Anisotropic permeability can cause flow to deviate from the apparent flow direction (Freeze and Cherry, 1979). Back and others (1983) described the general groundwater flow in the Madison aquifer for the Black Hills and surrounding area, indicating that groundwater flows from the surface recharge area in the northwestern part of study area to the southwest, then sweeps east and then northeast around the southern end of the Black Hills. This description generally is consistent with the apparent flow directions indicated by the potentiometric contours shown on figure 1. These apparent flow directions provide a generalization of groundwater flow in the study area and are not necessarily accurate. A low hydraulic gradient in the area of Wind Cave, as indicated by widely spaced contours (1,150-m to

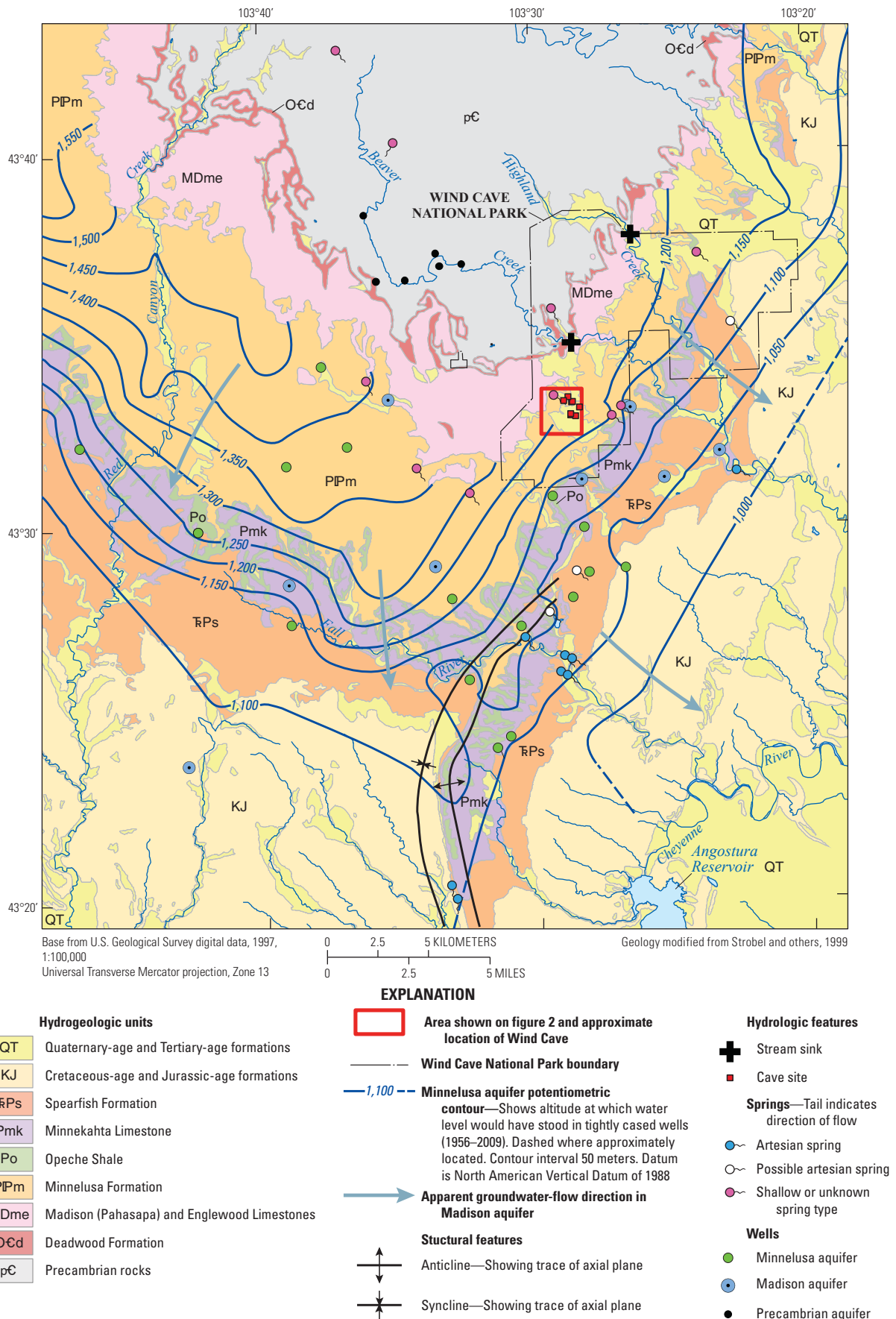


Figure 4. Potentiometric surface of the Minnelusa aquifer.

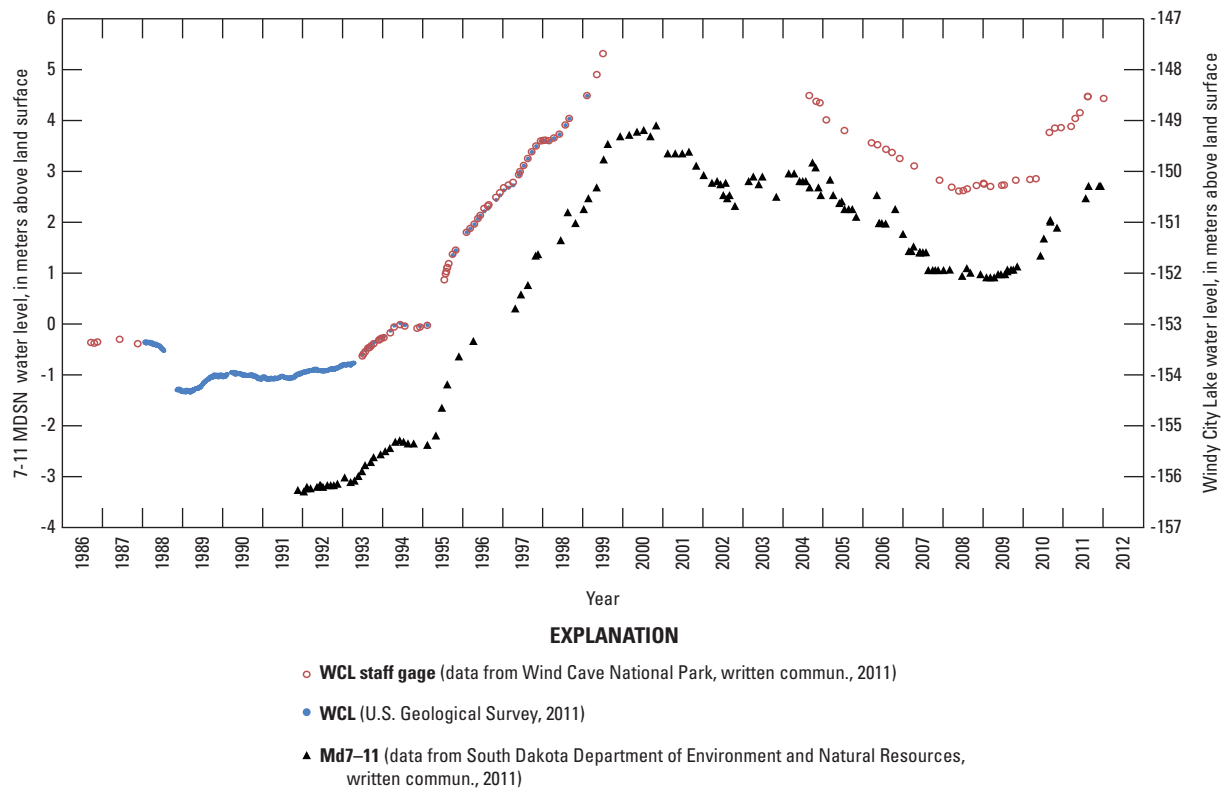


Figure 5. Water-level hydrographs for Windy City Lake (site WCL) and well Md7-11.

1,100-m contours; fig. 1), likely indicates anomalously high permeability resulting from extensive karst development.

A potentiometric-surface map of the Minnelusa aquifer also was constructed on the basis of water-level data available from the U.S. Geological Survey (2011). In general, the apparent groundwater-flow directions in the Minnelusa aquifer are similar to those in the Madison aquifer, except that in the western part of the study area the apparent flow direction is more westerly than in the Madison aquifer (figs. 1 and 4). The potentiometric surface of the Minnelusa aquifer (fig. 4) was subtracted from that of the Madison aquifer (fig. 1) to determine the direction and magnitude of the vertical hydraulic gradient (fig. 6). The resulting difference in hydraulic heads between the aquifers was used to estimate sources of spring flow. The Minnelusa aquifer has a potentiometric surface that is higher than that of the Madison aquifer within and near the outcrop of the Minnelusa Formation; farther from this outcrop, the Madison aquifer has the higher potentiometric surface (fig. 6). In places where the Madison and Minnelusa aquifers are connected hydraulically, groundwater flows from the aquifer with the higher hydraulic head into the aquifer with the lower hydraulic head. If breccia pipes are the throats of artesian springs as described by Hayes (1999), and if these throats hydraulically connect the Madison and Minnelusa aquifers, then the primary source of a spring would be the aquifer with

the highest hydraulic head of the two. Artesian springs generally are located in areas where the Madison aquifer has the higher potentiometric surface of the two aquifers, which indicates that the Madison aquifer probably is the primary source of these springs.

Land-surface altitudes where Precambrian rocks are exposed generally are higher than within the outcrop of the Madison Limestone in the study area (figs. 1 and 7). Groundwater levels in the Precambrian aquifer generally are near the land surface and higher than in the Madison aquifer (U.S. Geological Survey, 2011), and this groundwater gradient indicates a possibility that groundwater might flow from the Precambrian aquifer through the Deadwood aquifer and into the Madison aquifer. Evidence for this includes apparent groundwater ages estimated from the CFC data for CFC-12 for sites CON and WTHL (table 4). The apparent age is an estimate of age where the entire sample is assumed to contain water of a single age. Well CON is located near the southwestern edge of the Madison Limestone outcrop (fig. 1) and penetrates the aquifer by about 15 m. If not for an apparent groundwater age of 22–26 years (table 4), groundwater at this well might be assumed to represent almost entirely surface recharge because of its location. However, the apparent age indicates that a large fraction of this water did not result from recent precipitation but might have originated in the upgradient Precambrian

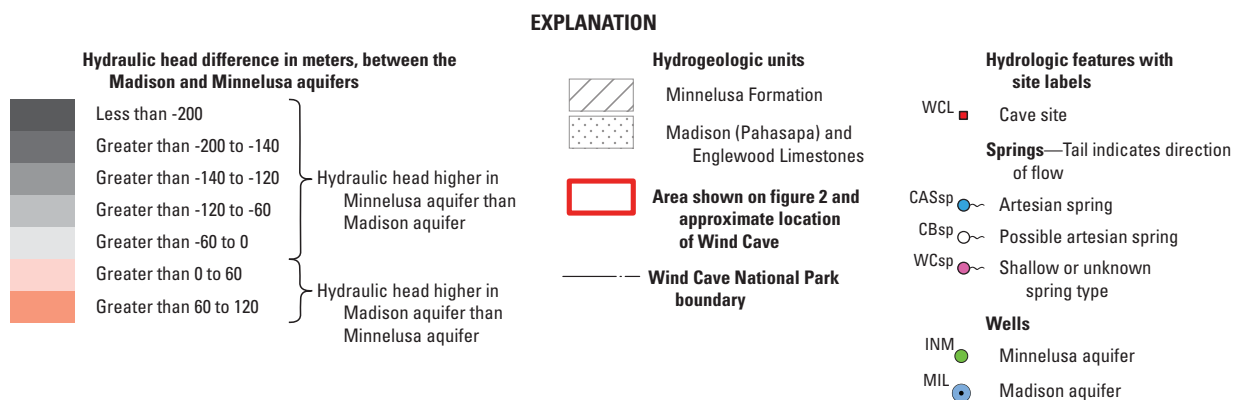
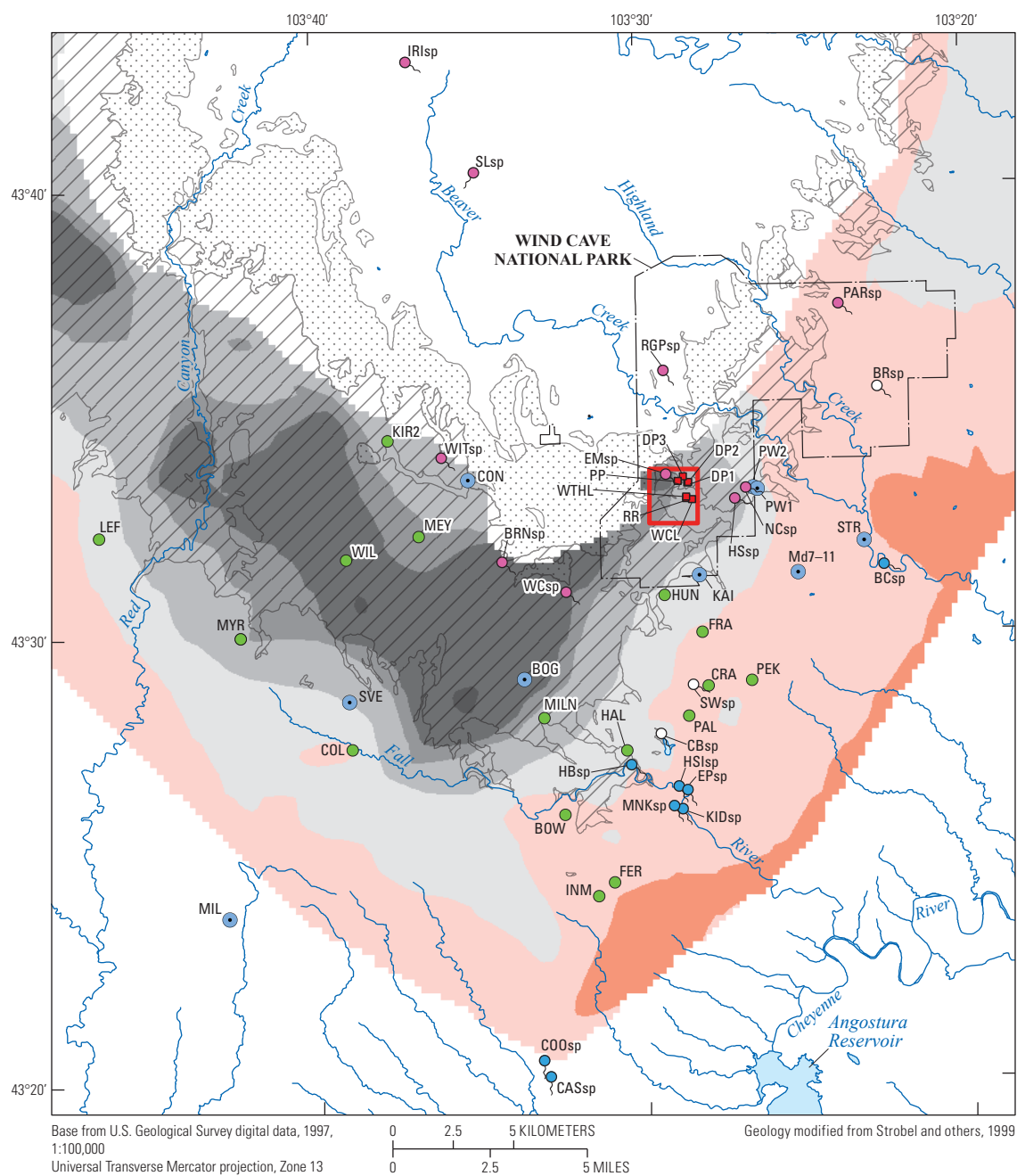


Figure 6. Difference in hydraulic head between the Madison and Minnelusa aquifers.

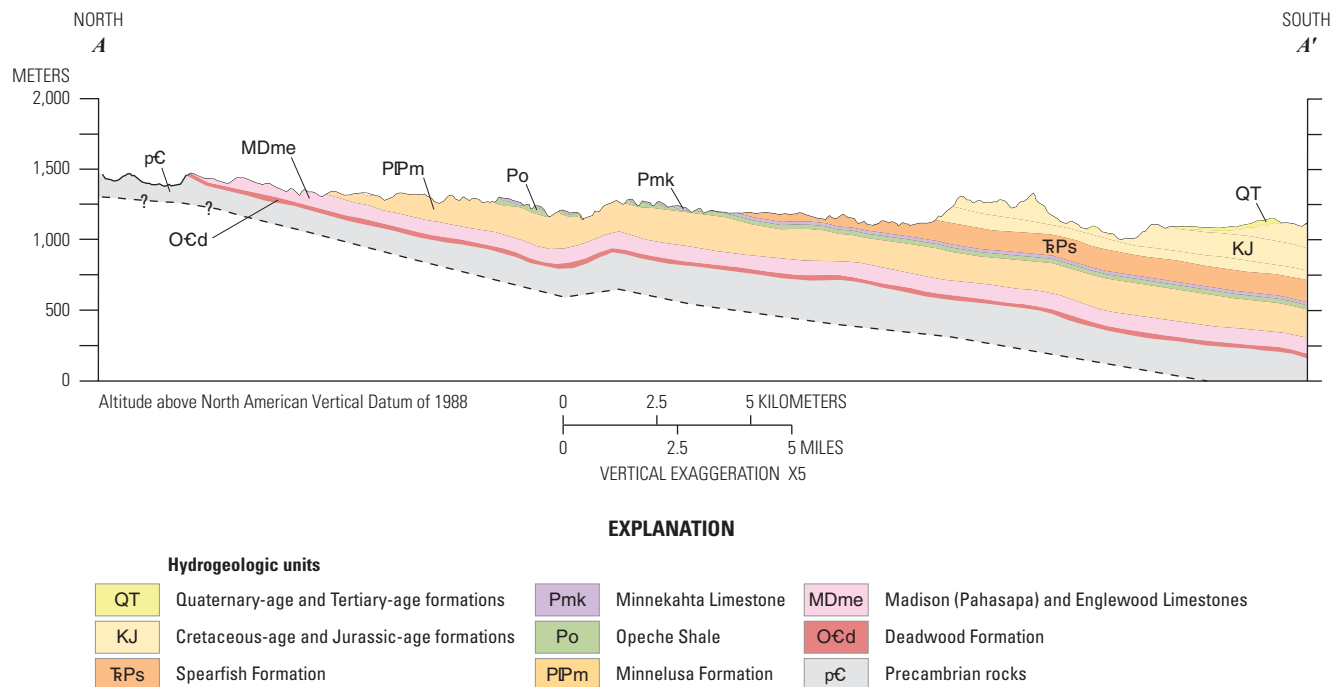


Figure 7. Geologic cross section A–A' shown on figure 1.

aquifer. Site WTHL, also located near the Madison Limestone outcrop, is a water body in Wind Cave (fig. 1) that is fed by groundwater emerging from the cave floor with an apparent age of 20–27 years (table 4), which also indicates the possible presence of water originating from the Precambrian aquifer.

Dye Tracing

Dye tracing was conducted to determine the connectivity and transit times between different groundwater sites in and near Wind Cave. Fluorescein dye was injected into What the Hell Lake (site WTHL), and charcoal dye receptors and water samples were collected and analyzed to determine the presence and concentration of dye in Rebel River (site RR; fig. 1), Windy City Lake (site WCL; fig. 1), and two wells in the park (wells PW1 and PW2; fig. 1), all of which are hydraulically downgradient from site WTHL (fig. 1). A dye receptor is a packet of charcoal, which adsorbs fluorescent dyes in water. These receptors were deployed at sites and submerged in water for at least one month. After receptors were retrieved, the dye was removed by an eluent, which was then analyzed for the presence of dye, the concentration of which is affected by the dye concentration in the water during deployment and the duration of deployment.

Wind Cave National Park staff deployed and collected the dye receptors and water samples and sent them to the Crawford Hydrology Laboratory at Western Kentucky University for analysis (table 5). The initial set of dye receptors were deployed on December 3, 2007. Dye receptors

were replaced each time they were retrieved. Analyses of dye receptors and water samples were conducted at Crawford Hydrology Laboratory with a Shimadzu Model RF-5301PC Synchronously Scanning Spectrofluorophotometer, which can detect dye concentrations as low as 5 parts per trillion (<http://www.dyetracing.com>). Four liters of liquid fluorescein dye (40-percent fluorescein by weight) were injected into What the Hell Lake (site WTHL) in Wind Cave on February 26, 2008. Visual dye was present in the water at WTHL and RR during a site visit on March 25, 2008. Visual dye was present in the water at WCL during a site visit on May 28, 2008.

Dye tracing indicated that groundwater reached sites WCL and RR from the injection site (WTHL) in 28 days or less (table 5). Dye was detected in every dye receptor or sample from site WCL from March 2008 to August 2011, or 42 months. Assuming a straight-line flow path from site WTHL to site WCL (265 m), which is nearly perpendicular to potentiometric contours (fig. 2), the groundwater velocity was at least 9 meters per day (m/d), but might have been faster because the first sample collected from site WCL after injection was 28 days later. Residence time for dye at site WCL and this flow path combined was at least 42 months. Residence time at site WCL alone might be shorter than 42 months because of possible groundwater retention along the flow path, which could have been a source of ongoing input to this site. The last dye receptor retrieved from site RR was 92 days after injection and had an eluent concentration that was more than 200 times higher than in eluent from any receptors retrieved from site WCL (table 5). Dye was not detected with certainty

for any of the five samples collected from wells PW1 and PW2 (table 5).

Groundwater Quality

Hydrochemical data for samples collected from 60 sites are presented in table 12 in the “Supplemental Tables” section. Hydrochemical data presented in table 12 include pH, specific conductance, common ions, nitrate plus nitrite, arsenic, and stable isotopes of oxygen and hydrogen. Water samples from selected sites were analyzed for trace metals, and the analytical results are presented in table 6. The occurrence and geospatial distributions of specific conductance, nitrate plus nitrite, arsenic, and trace metals were assessed. Arsenic is a particular concern because of concentrations within the park that were higher than desirable for drinking-water quality.

Specific Conductance

Specific conductance is a measure of the electrical conductivity of water and indicates relative dissolved-solid content (Hem, 1985). The geospatial distribution of specific conductance in groundwater samples provides a general overview of water quality and hydrochemical differences in the study area. Specific conductance was interpolated between sampled sites and is shown as ranges of values in the study area (fig. 8). Specific conductance generally is lowest in recharge areas for the Madison, Minnelusa, and Precambrian aquifers and highest in the southern part of the study area, distant from recharge areas (fig. 8). The highest specific conductance values might indicate long residence times because time is required to dissolve minerals in aquifer media. A correlation between apparent groundwater age and specific conductance in the Madison aquifer exists in the Rapid City area (Long and others, 2008; South Dakota inset map on fig. 1). The southern part of the study area probably represents a mixture of groundwater that originated as surface recharge within the study area and as regional flow from the west and northwest.

Arsenic

Arsenic concentrations in samples collected for this study ranged from 0.28 to 37.1 mg/L with a median value of 6.4 mg/L (table 12), and 32 percent of these exceeded 10 mg/L, which is the Maximum Contaminant Level (MCL) for drinking water in the United States (U.S. Environmental Protection Agency, 2010). The geospatial distribution of arsenic in groundwater samples for the Madison, Minnelusa, and Precambrian aquifers indicates that the highest arsenic concentrations in and near the study area are approximately coincident with the outcrop of the Minnelusa Formation as shown in figure 9, which indicates an arsenic anomaly in this area. Figure 9 was based on samples collected for this study (table 12) plus data for all other water samples available in the National Water Information System (U.S. Geological Survey, 2011). The median arsenic concentration is about equal for

the Madison and Minnelusa aquifers (fig. 10). Overall, the Minnelusa aquifer has slightly larger arsenic concentrations than the Madison aquifer (fig. 10). Where sampling sites for the Madison and Minnelusa aquifers are proximal, arsenic concentrations generally are similar (fig. 9), so the concentration differences indicated by figure 10 might result partly from geospatial differences in sample locations between the two aquifers.

The exceptions to the association of arsenic with the outcrop of the Minnelusa Formation are samples from two wells open to the Precambrian aquifer (well MOR and WOO; fig. 9), which had arsenic concentrations of 35.2 and 12.7 mg/L, respectively. These concentrations, which are high relative to other Precambrian aquifer samples, might be the result of previous industrial activities in the town of Pringle (located at well MOR), which were the focus of U.S. Environmental Protection Agency (USEPA) investigations from 1991 to 2001 (Hayhurst, 2002), and included wood preservative treatment with chromated copper arsenate. Soil in the Pringle area was contaminated with arsenic, chromium, copper, zinc, pentachlorophenol, polycyclic aromatic hydrocarbons, dioxins, and furans. Dioxins and furans were detected in Beaver Creek and its sediments downstream from the former industrial site, with decreasing concentrations in a downstream direction (Hayhurst, 2002). Heakin (2004) documented the presence of phenol in Beaver Creek in 2002 and 2003 in the park and about 2 km upstream from the western park boundary.

The arsenic anomaly coincident with the outcrop of the Minnelusa Formation also exists when all groundwater and surface-water samples available from the National Water Information System (U.S. Geological Survey, 2011) in and near the study area are plotted (fig. 11). The area shown in figure 11 extends beyond the study area to show a larger representation of the arsenic anomaly and indicates that the Minnelusa and Madison aquifers have the highest arsenic concentrations when compared with other groundwater and surface-water sites in and near the southern Black Hills.

The geographic association of the arsenic anomaly with the Minnelusa Formation outcrop (excluding the Pringle area) indicates that arsenic in groundwater probably results from naturally occurring minerals in the marine shales of this formation. Marine shales commonly contain high concentrations of arsenic, which is released into the aqueous phase in aquifers as a result of oxidative weathering of iron-sulfide minerals, mainly pyrite (FeS_2) and arsenopyrite (FeAsS ; Muloin and Dudas, 2005). During sulfide mineral formation, arsenic precipitates with iron and sulfide and is incorporated into the mineral structure (Muloin and Dudas, 2005). Zhu and others (2008) concluded that microbial sulfide production enhances arsenic mobilization in pyrite-bearing black shale formations. Solid-phase arsenic concentration was analyzed in clay, sand and gravel, shales, and limestones in aquifers of the Lower Illinois River Basin (not shown in report), where arsenic concentrations in groundwater were as high as 110 mg/L, and the highest solid-phase concentrations were in shales (Warner, 2001). This author concluded that dark shales in the bedrock

Table 4. Chlorofluorocarbon (CFC) and tritium data.

[n, number of CFC samples; pptv, parts per trillion by volume; pCi/L, picocuries per liter; --, could not be estimated by this method]

Site name	Station number	Station name	Date	n	Calculated atmospheric mixing ratio, in pptv ^a			Apparent groundwater age, in years ^a					Results of binary mixing model ^b							Tritium, in pCi/L	Tritium, 2-sigma, in pCi/L		
					CFC– 11	CFC–12	CFC– 113	CFC–11	CFC–12	CFC–113	CFC–11/CFC–12 ratio age, in years ^a	Percent young water in mixture ^{a,c}	Number of samples used	CFC–113/ CFC–12 ratio age, in years ^a	Percent young water in mixture ^{a,c}	Number of samples used	CFC–11/ CFC–11 ratio age, in years ^a	Percent young water in mixture ^{a,c}	Number of samples used				
Stream sink site					Stream sink site																		
BevCr	06402430	Beaver Creek near Pringle, S. Dak.	04/16/2007	0	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	50.2	3.0		
Cave drip site					Cave drip site																		
DP3	433302103281509	6S 5E12DBAB9	04/25/2007	0	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	29.4	1.9		
Cave water body sites					Cave water body sites																		
WTHL	433302103281502	6S 5E12DBAB2	04/18/2007	2	196.8	439.8	55.0	24	20	20	--	--	--	--	--	--	18	72	2	15.7	1.3		
WTHL	433302103281502	6S 5E12DBAB2	07/16/2009	2	190.3	445.2	53.0	27	23	23	--	--	--	--	--	--	20	75	2	--	--		
Artesian spring sites					Artesian spring sites																		
HBsp	432703103302801	7S 5E10DCBA	04/20/2007	2	44.3	107.3	12.3	39	39	32	--	--	--	22	26	2	15	68	2	3.8	1.0		
HBsp	432703103302801	7S 5E10DCBA	05/28/2009	2	35.5	104.9	11.1	43	41	35	--	--	--	25	28	2	15	13	2	5.7	--		
BCsp	433128103223401	6S 6E14CDB	04/17/2007	3	31.9	82.7	10.3	41	41	33	--	--	--	21	20	3	19	63	2	4.5	1.0		
BCsp	433128103223401	6S 6E14CDB	05/28/2009	2	30.2	91.7	8.6	44	42	37	--	--	--	26	25	2	20	12	2	6.0	--		
CASsp	432006103330501	8S 5E20CDAB	06/19/2009	3	24.6	60.6	6.9	45	46	38	38	36	2	24	15	3	20	10	3	4.0	--		
COOsp	432028103331601	8S 5E20BDCB	06/19/2009	3	38.4	103.9	13.0	42	41	34	41	89	1	22	25	3	12	15	1	3.9	--		
HSIsp	432632103285302	7S 5E14DDDA	05/21/2009	2	42.2	157.5	11.8	42	38	34	--	--	--	29	52	2	20	16	2	--	--		
KIDsp	432605103285401	7S 5E14DDD	06/19/2009	3	28.8	169.9	13.0	44	37	34	--	--	--	29	55	3	--	--	--	.9	--		
MIL	432340103421501	7S 3E36CBDC	04/08/2010	3	21.0	3.1	.0	47	64	57	--	--	--	--	--	--	--	--	--	--	--		
MNKsp	432605103290901	7S 5E14DCD	05/28/2009	2	25.6	171.3	3.4	45	37	44	--	--	--	--	--	--	30	16	2	.5	--		
Well sites					Well sites																		
HUN	433034103284701	6S 5E23DB	04/17/2007	3	129.4	283.2	35.6	31	29	24	--	--	--	20	67	3	18	71	3	26.9	1.9		
BOG	432858103334201	6S 5E31DA	04/26/2007	3	96.5	170.2	23.9	34	35	27	30	66	1	19	37	3	18	63	3	1.6	1.0		
BOG	432858103334201	6S 5E31DA	07/15/2009	3	28.7	99.3	9.4	44	42	36	--	--	--	26	28	3	13	11	2	.6	--		
CON	433326103352001	6S 4E 1DB	07/15/2009	3	220.7	388.0	61.7	24	26	22	--	--	--	18	76	2	20	86	3	29.3	--		
Md7–11 ^d	433115103251401	6S 6E21BBBB (CU91A)	04/20/2007	2	4.6	3.0	.0	52	61	54	--	--	--	--	--	--	--	--	--	.6	.6		
PW1	433311103263101	6S 6E 6DDA	07/29/2009	2	147.3	316.1	38.3	32	29	26	--	--	--	23	77	2	21	--	2	21.4	--		
PW2	433311103263102	6S 6E 6DDA2	07/29/2009	3	182.3	115.8	13.7	27	41	34	--	--	--	24	29	3	21	20	1	9.2	--		
STR	433150103230501	6S 6E15ABDD	04/17/2007	3	70.6	152.5	17.9	36	36	29	36	97	1	21	38	3	19	62	3	16.6	1.3		
STR	433150103230501	6S 6E15ABDD	05/22/2009	2	66.0	162.2	19.3	39	38	31	--	--	--	23	40	2	19	25	2	17.4	--		
SVE	432825103391201	7S 4E 4BAC	09/02/2009	3	191.6	80.4	8.2	27	44	38	--	--	--	26	21	3	--	--	--	3.4	--		

^a Average of multiple samples. Age is the apparent age of the young fraction.

^b Plummer and Busenberg (2000).

^c Percentage younger than about 60 years.

^d Methane present. Apparent ages represents the minimum apparent age.

aquifer were the likely source of arsenic in the overlying and adjacent glacial drift aquifer. In groundwater of the Piedmont Province of Pennsylvania (not shown in report), arsenic concentrations higher than 10 mg/L were the result of arsenic mobilization from shales, whereas the arsenic concentrations in the pore water of igneous, crystalline, and calcareous rocks typically were lower (Serfes and others, 2006; Peters and Burkert, 2008; Rhine and others, 2008).

It is most likely that arsenic in the Minnelusa and Madison aquifers originates in shale layers of the Minnelusa aquifer and is transported downward into the Madison aquifer by advective flow. This is possible in areas where the hydraulic head in the Minnelusa aquifer is higher than in the Madison aquifer (fig. 6), which also is approximately coincident with the arsenic anomaly (fig. 9). This downward seepage is known to occur in the area of Wind Cave, where groundwater in the unsaturated zone of the Minnelusa aquifer seeps downward into the Madison aquifer and drips from the ceiling of the cave at numerous locations. Arsenic concentrations in sampled drip water (table 12) ranged from 24.3 to 34.2 mg/L (sites DP1, DP2, and DP3) and from 12.6 to 16.4 mg/L for water bodies in the cave (sites WCL, WTHL, PP, and RR). This indicates that arsenic concentrations in Wind Cave were highest for water that is known to originate in the Minnelusa aquifer. Cave drip probably contributes to cave water bodies, which also may receive contributions from the Precambrian aquifer

as previously described in the “Groundwater Gradients and Flow Directions” section. If so, this would result in a mixture in the cave water bodies of groundwater from the Minnelusa aquifer (cave drip) and the Precambrian aquifer. The sampling site closest to the cave for the Precambrian aquifer was a spring (RGPsp; fig. 1), which had an arsenic concentration of 2.0 mg/L (table 12). Therefore, mixing could have affected the observed arsenic concentrations in the cave water bodies, which had concentrations between those of cave drip and the Precambrian aquifer.

It is uncertain why the arsenic anomaly is coincident with the Minnelusa Formation outcrop rather than confined aquifer areas, but might be related to weathering of shales in the outcrop. As Muloin and Dudas (2005) stated, weathering of marine shales releases arsenic into groundwater. Further, as described by Tuttle and others (2009), during oxidative weathering of iron-sulfide minerals in marine shales, arsenic and other trace elements are dispersed or redistributed in the environment by aqueous and mechanical transport. The decrease in arsenic concentration downgradient from the outcrop areas of the Minnelusa and Madison Formations indicates that mobilized arsenic is not transported large distances in the Minnelusa and Madison aquifers. The reason for this is unclear, and arsenic concentrations did not correlate with either pH or dissolved oxygen. It is possible that dissolved arsenic originates from particular shale layers in the Minnelusa Formation,

Table 5. Results of laboratory analyses of the presence of fluorescein dye in charcoal dye receptors that were placed in Wind Cave water bodies and two wells. Dye was injected on February 26, 2008.

[ppb, parts per billion; B, background concentration; --, no sample; ND, nondetection]

Date collected	Days since injection	Years since injection	Windy City Lake (WCL)		Rebel River (RR), in ppb	Well PW1	Well PW2
			Dye receptor eluent, in ppb	Water sample, in ppb			
02/26/2008	0	0.0	B	--	ND	--	--
03/20/2008	23	.1	--	--	--	ND	ND
03/25/2008	28	.1	0.85	--	249,000	--	--
05/28/2008	92	.3	59.4	--	93,000	--	--
07/09/2008	134	.4	168	--	--	--	--
08/04/2008	160	.4	--	--	--	B	ND
09/11/2008	198	.5	--	--	--	--	ND
11/13/2008	261	.7	376	--	--	--	--
01/07/2009	316	.9	96.3	--	--	--	--
06/30/2009	490	1.3	--	--	--	--	B
05/03/2010	797	2.2	302	--	--	--	--
07/16/2010	871	2.4	261	--	--	--	--
04/25/2011	1,154	3.2	--	.52	--	--	--
06/08/2011	1,198	3.3	21.4	--	--	--	--
08/18/2011	1,269	3.5	101.5	--	--	--	--

Table 6. Trace-metal data for water samples from selected sites within Wind Cave National Park.[µg/L, micrograms per liter; µS/cm, microsiemens per centimeter at 25 degrees Celsius; <, less than; E, estimated value^a; --, not applicable]

Site name	Station number	Station name	Date	Arsenic, dissolved (µg/L)	Chromium, dissolved (µg/L)	Copper, dissolved (µg/L)	Lithium, dissolved (µg/L)	Vanadium, dissolved (µg/L)	Zinc, dissolved (µg/L)	Specific conductance, unfiltered (µS/ cm)
Stream sink site										
BevCr	06402430	Beaver Creek near Pringle, S. Dak.	06/25/2008	4.4	<0.12	<1	34	2.7	<1.8	563
Cave drip sites										
DP2	433302103281508	6S 5E12DBAB8	06/26/2008	24.4	0.12	E 0.99	6	6.6	<1.8	360
DP3	433302103281509	6S 5E12DBAB9	06/26/2008	23.8	<.12	E .54	2	6.5	E 1.1	263
Artesian spring site										
BRsp	433525103224401	5S 6E26BBCD	06/25/2008	5.6	0.38	2.4	422	19.6	<3.6	2,830
Shallow spring sites										
PARsp	433717103235401	5S 6E15BBAC	06/25/2008	12.2	0.12	1.5	28	6.4	2.4	472
Emsp	433332103291801	6S 5E 2ACBD	06/25/2008	12	E .08	1.1	10	16.6	16.6	605
HSsp	433258103270801	6S 6E 7BAAC	06/26/2008	25.9	E .07	1.6	7	4.4	<1.8	390
NCsp	433312103264701	6S 6E 6DCAA	06/26/2008	16.7	<.12	E .92	7	6	E 1.2	408
NCsp	433312103264701	6S 6E 6DCAA	06/26/2008	16.7	<.12	E .57	7	6	1.8	408
RGPsp	433551103291901	5S 5E23DBCD	06/25/2008	1.7	<.12	<1	37	.72	4	363
Maximum Contaminant Level (MCL), in µg/L ^b				10	100	1,300	--	--	--	--

^a Value less than the lowest calibration standard (Childress and others, 1999).^b From U.S. Environmental Protection Agency (2010).

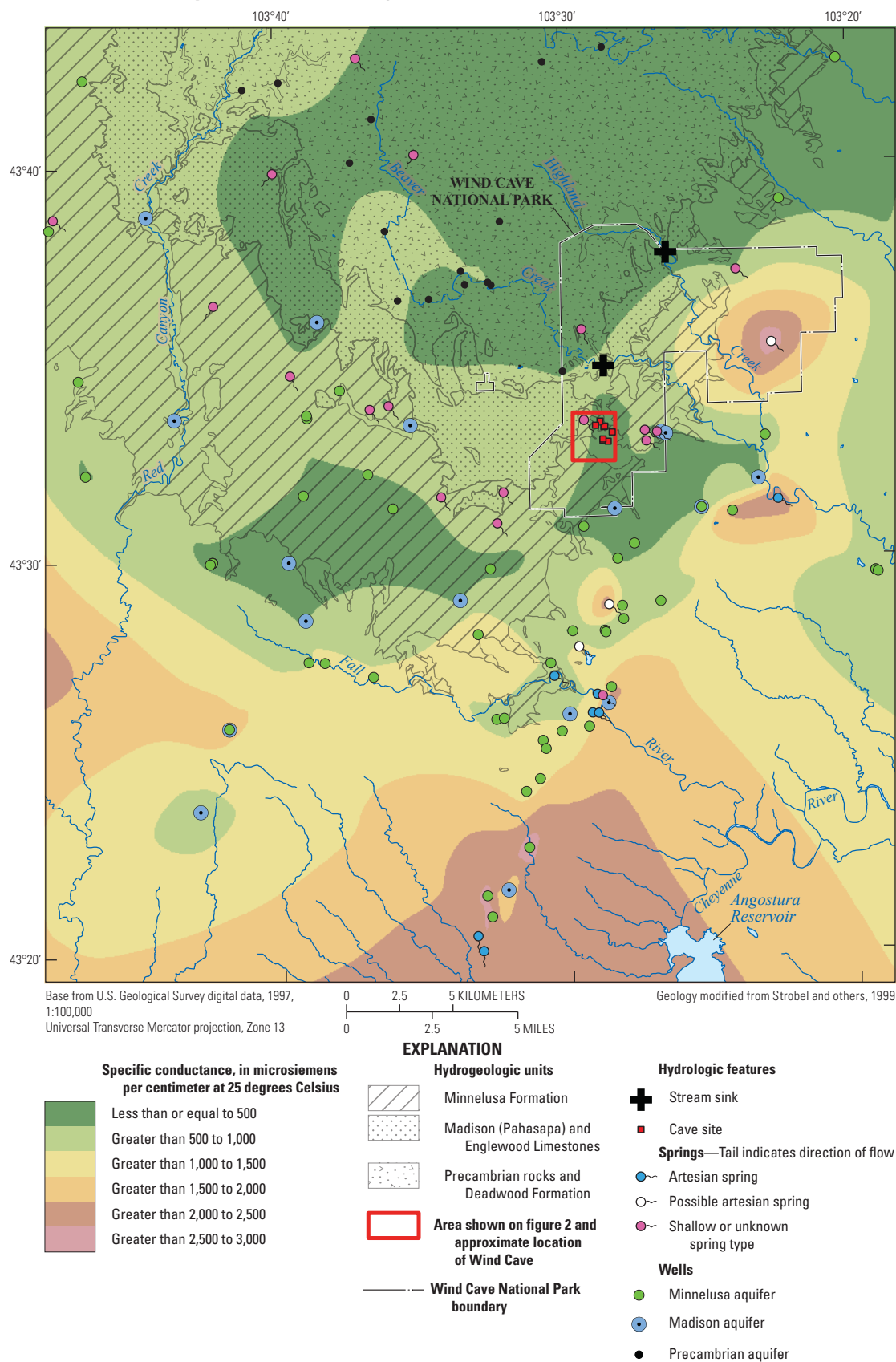


Figure 8. Geospatial distribution of specific conductance for the Minnelusa, Madison, and Precambrian aquifers in the study area. (All relevant sites with specific conductance data available from the National Water Information System [U.S. Geological Survey, 2011] are included in map.)

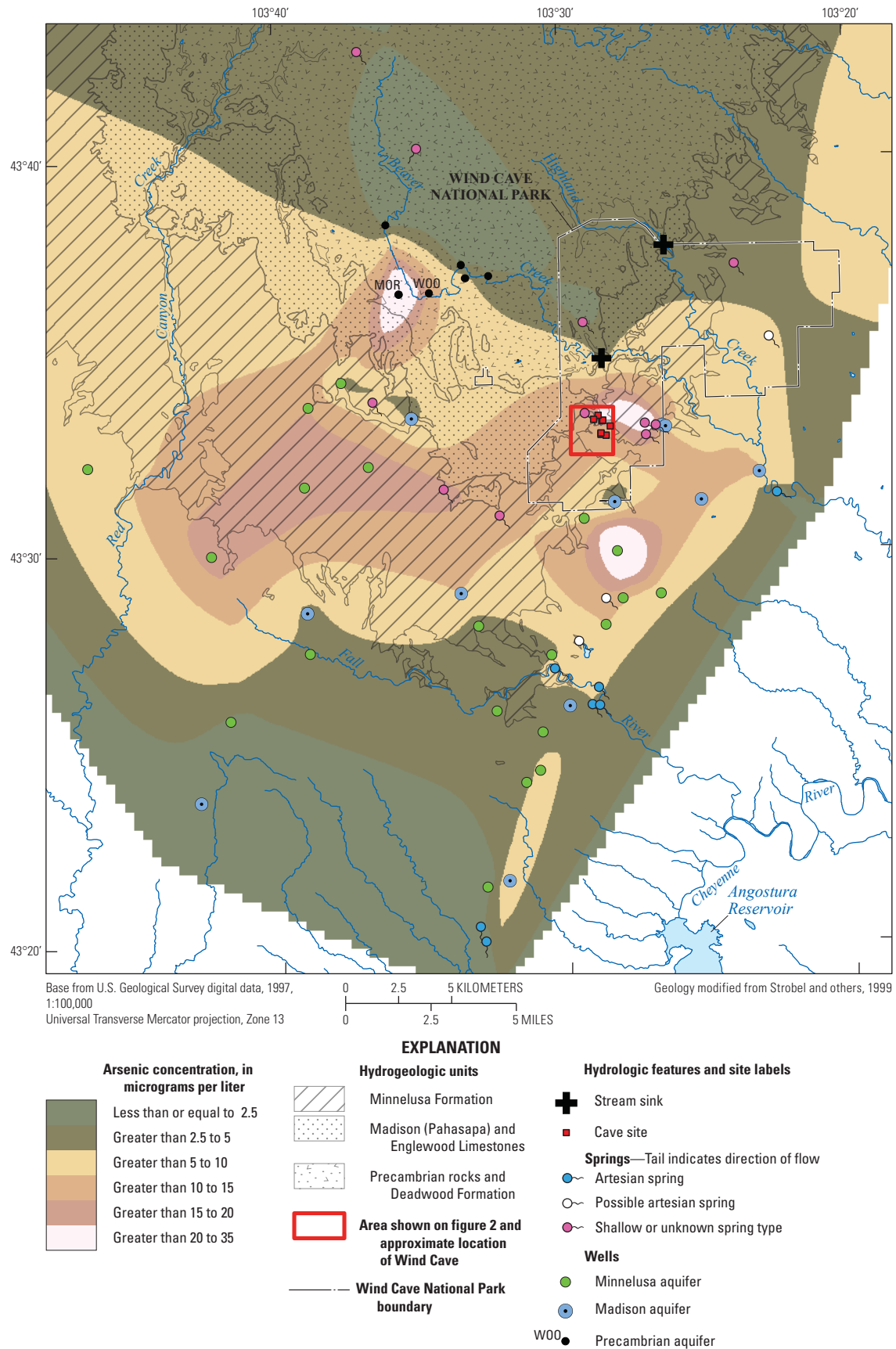


Figure 9. Geospatial distribution of arsenic concentrations for the Minnelusa, Madison, and Precambrian aquifers in the study area. (All relevant sites with arsenic data available from the National Water Information System [U.S. Geological Survey, 2011] are included in map.)

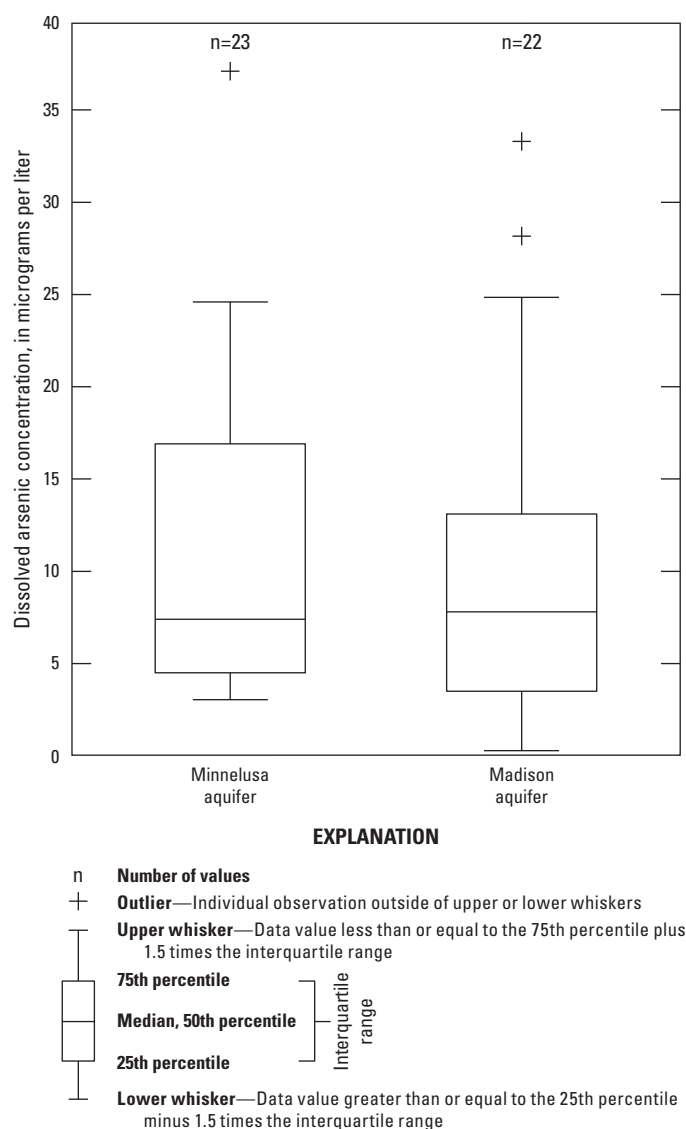


Figure 10. Boxplots of arsenic concentrations in the Minnelusa and Madison aquifers.

which was assessed by considering a possible correlation between arsenic concentrations and the sampling depth from the top of the Minnelusa aquifer; however, no such correlation is apparent on a scatter plot (fig. 12).

Nitrate Plus Nitrite

Potential sources of nitrate in groundwater include nitrogen fertilizer, mining, landfills, atmospheric deposition, and human or animal waste (Meyer, 1987; Taylor, 2003; Katz, 2004; Rahn, 2006). Concentrations of nitrate plus nitrite in samples collected for this study are presented in table 12 and were less than 2 milligrams per liter (mg/L) for 92 percent of these samples and ranged from 0.13 to 2.38 mg/L. Concentrations in this range are not a concern for drinking-water quality

because the MCL for nitrate drinking water is 10 mg/L (U.S. Environmental Protection Agency, 2010). The area with the highest nitrate plus nitrite concentrations is on the western side of the outcrop areas of the Madison and Minnelusa aquifers with concentrations decreasing to the south and east (fig. 13). The reason for this geospatial distribution is unclear and unaccounted for by land-use differences. Grazing of domestic stock occurs in many areas outside of Wind Cave National Park, and wildlife grazing occurs inside the park. Wind Cave National Park monitors nitrate concentrations in precipitation for the park and reports a 10-year average of 1.4 mg/L (Wind Cave National Park, written commun., 2011).

Site WCL in Wind Cave had a concentration of nitrate plus nitrite of 2.38 mg/L, which is not visible in the interpolated shaded areas on figure 13 because other proximal cave sites had lower concentrations ranging from 0.5 to 1.6 mg/L. Cave sites had a median concentration of 1.3 mg/L, and other sampled sites in the park had a median concentration of 0.2 mg/L. The two wells within the park (sites PW1 and PW2) had concentrations of 0.6 and 0.2 mg/L, respectively.

Trace Metals

Exploratory sampling of trace metals (chromium, copper, lithium, vanadium, and zinc) for selected sites in the park was conducted to assess possible existence of unusually high concentrations or correlations of these metals with arsenic. The USEPA regulates arsenic, chromium, and copper with MCLs of 10, 100, and 1,300 $\mu\text{g/L}$, respectively (U.S. Environmental Protection Agency, 2010). The highest concentrations for chromium and copper were 0.38 and 2.4 $\mu\text{g/L}$, respectively, or less than 1 percent of the MCL (table 6). The concentration of lithium for spring BRsp was 422 $\mu\text{g/L}$, which was much higher than the other sites, and specific conductance for this site also was much higher than the other sites (table 6). Although the USEPA has not set an MCL for drinking water for lithium, it can be toxic to plants. For example, injury to some crops from lithium in irrigation water occurred at concentrations as low as 50 $\mu\text{g/L}$, but in other cases, injury was not observed at concentrations lower than 5,000 $\mu\text{g/L}$ (Bingham and others, 1964). Lithium concentrations of 600 micrograms per gram ($\mu\text{g/g}$) in the leaves of Bush bean plants resulted in severe toxicity, and lithium accumulates in plant tissue (Wallace and others, 1977). Based on all samples collected in the park, arsenic indicated no apparent correlation with any of the five other trace metals when observed on scatter plots. Because of the generally low concentrations of trace metals and the lack of correlation with arsenic, additional sampling was not conducted.

Groundwater Mixing

All variables used in PCA were log transformed, which resulted in datasets that approximated normal distributions, and all variables were then standardized to a mean of zero and standard deviation of one. Stable-isotope values (table 12), all

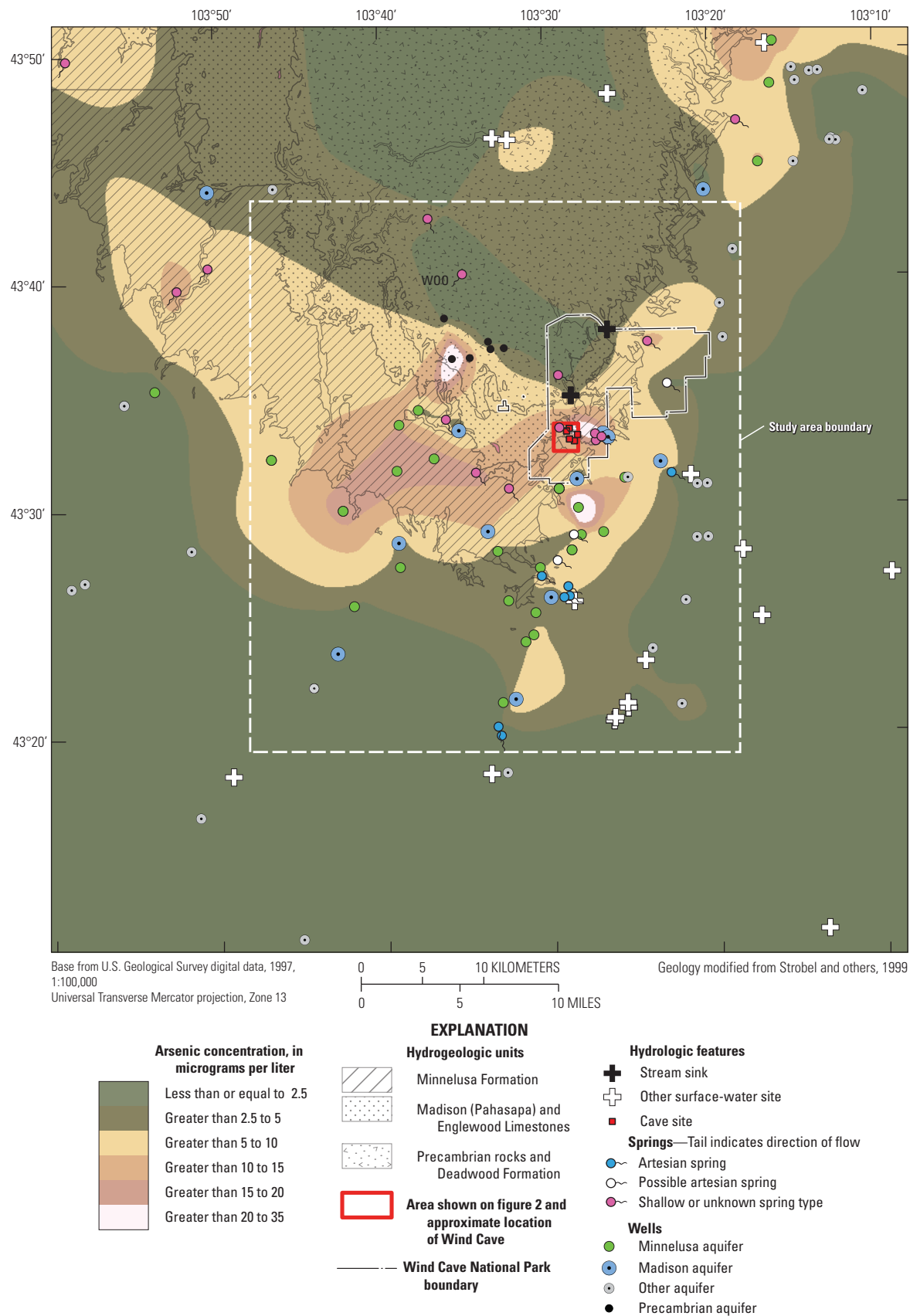


Figure 11. Geospatial distribution of arsenic concentrations for all groundwater and surface-water samples in and near the study area. (All relevant sites with arsenic data available from the National Water Information System [U.S. Geological Survey, 2011] are included in map.)

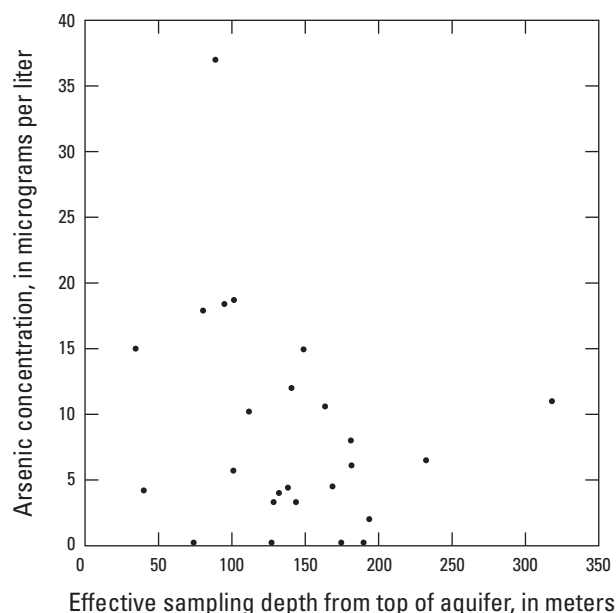


Figure 12. Scatter plot of arsenic concentrations in the Minnelusa aquifer and sampling depth from the top of the aquifer.

of which were negative, were multiplied by -1 for log transformation, then multiplied by -1 after transformation for consistency with the original sign. Values of pH were converted to hydrogen ion activity prior to normalization and standardization. To avoid giving disproportionate weight to any site, data for each site with multiple samples (excluding quality-control replicate samples) were averaged for PCA and end-member mixing so that each site had only one value per variable. Ten sites had missing data for one variable, and one site had missing data for four variables (1.2 percent of the dataset). However, because multiple samples were available for all of these sites, no values were missing in the site-averaged dataset. For end-member mixing, data were standardized to equalize variable weights and were not log transformed. The following results of PCA, cluster analysis, and end-member mixing are summarized from Long and Valder (2011).

Principal Component and Cluster Analyses

Results of PCA are shown as sampling sites plotted on the first two principal components, which explain 65 percent of the total variance of the data (fig. 14). Principal components 1, 2, and 3 represented 48, 17, and 11 percent, respectively, of the total variance, with higher order principal components representing fractions of the total variance that ranged from 0.03 to 7.9 percent. For cluster analysis, the number of clusters is specified by the user. Different numbers of clusters were tested with a maximum of six clusters. Five clusters were chosen because the five clusters that formed were separated by divisions that could be explained hydrogeologically and thus also

would work well as a basis for an end-member mixing model (fig. 15). The PC cluster consists mainly of Precambrian aquifer sites. The West cluster consists of Minnelusa and Madison aquifer sites mainly located near or within the western surface recharge area of these aquifers. The East cluster is similar to the West cluster, except it includes all cave sites and the two stream sinks. The Artesian 1 cluster consists of all artesian springs near the Fall River and wells open to the Madison and Minnelusa aquifers in that general area, and the Artesian 2 cluster consists of the remaining artesian springs, which are all located within or near the outcrop of the Spearfish Formation. Because these cluster areas could be interpreted and differentiated hydrogeologically, these geospatial areas of the groundwater-flow system are referred to as “hydrogeologic domains.” Hydrochemical data for these domains are summarized in table 13.

The relative influences of principal components 1 and 2 on the five clusters are shown in figure 14. Principal component 1 separates the Artesian 1 and 2 clusters from the three clusters to the north and thus distinguishes groundwater affected by regional flow from that affected by local recharge. Hydrochemical differences between the Artesian 1 and 2 clusters account for the greatest variability in the dataset, which is indicated by the distances between clusters on figure 14. The PC, East, and West clusters all are separated from each other on principal component 2, which therefore distinguishes groundwater affected by different local recharge sources. Also, differentiation of the East cluster from the PC and West clusters is associated with principal component 1. The Artesian 1 and 2 clusters are separated on principal component 2 and account for the largest variability of this principal component (fig. 14). Therefore, in addition to distinguishing local recharge, principal component 2 describes differences between artesian springs.

Similarly to the way scores determine the plotting positions for sites (fig. 14), the term “loadings” refers to the plotting positions of the original variables (Davis, 2002). Sites and variables are plotted together and used to assess relations of variables and clusters (fig. 16). Because of the different ranges of the values of scores and loadings, these values were scaled to range from -1 to 1 so that sites and variables could be plotted together (fig. 16). The vector lengths of the variables indicate their respective relative influences, and all variables except arsenic, pH, and bicarbonate have similar influence. Calcium, magnesium, sulfate, and specific conductance plot in the Artesian 2 cluster indicating a strong influence of these variables with this cluster, whereas the Artesian 1 cluster is influenced by sodium and chloride. Stable isotopes $\delta^{18}\text{O}$ and $\delta^2\text{H}$ plot near the East and West clusters and thus might help distinguish between these clusters, which is consistent with a gradient in these isotope values from east to west in the study area (Naus and others, 2001). The variables silica and magnesium are heavily loaded on principal component 2 and therefore are important in distinguishing clusters that are separated on that principal component. Most of the remaining variables mainly are loaded on principal component 1 and

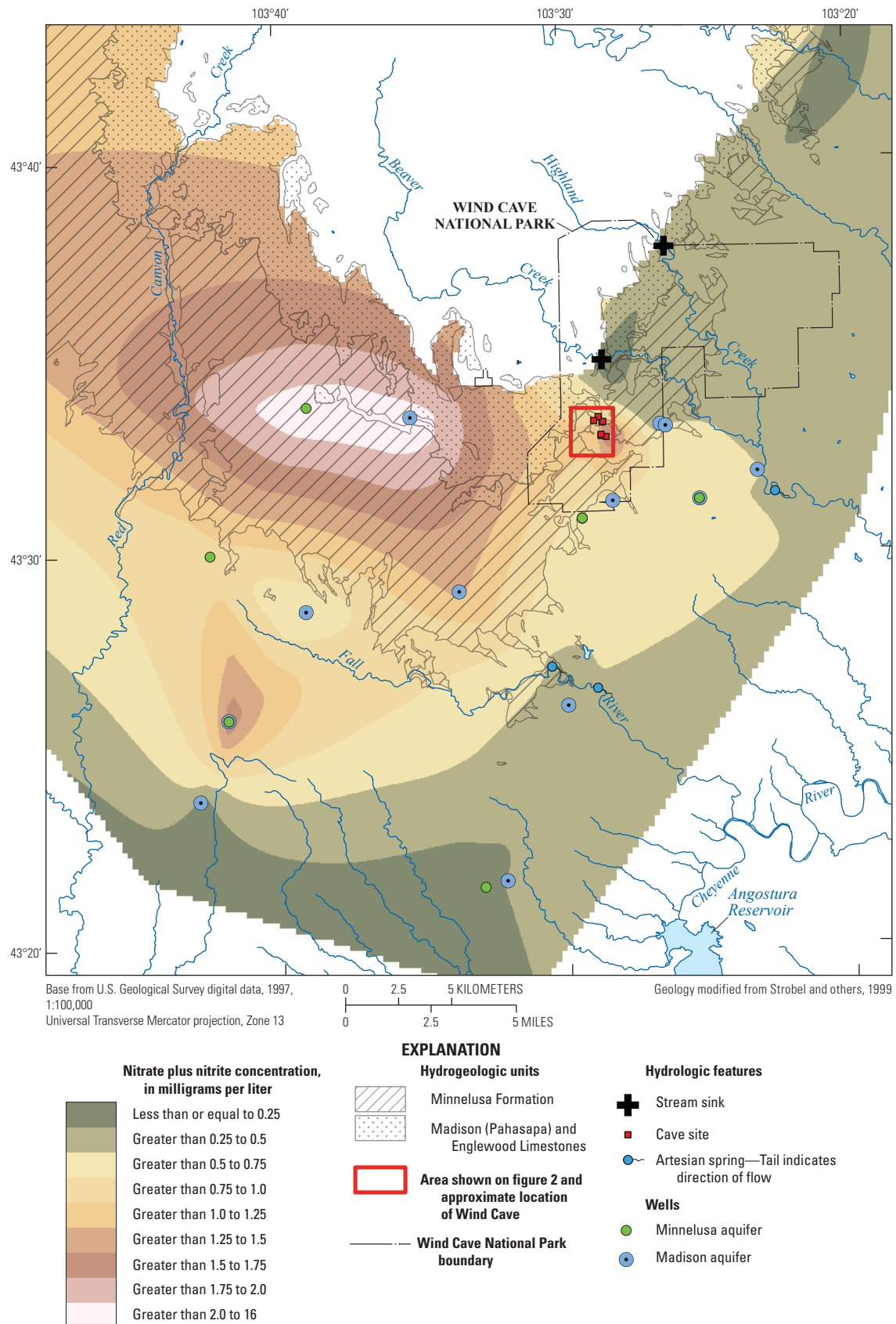


Figure 13. Geospatial distribution of nitrate plus nitrite for the Minnelusa and Madison aquifers in the study area. [All relevant sites with nitrate plus nitrite data available from the National Water Information System (U.S. Geological Survey, 2011) are included in the map].

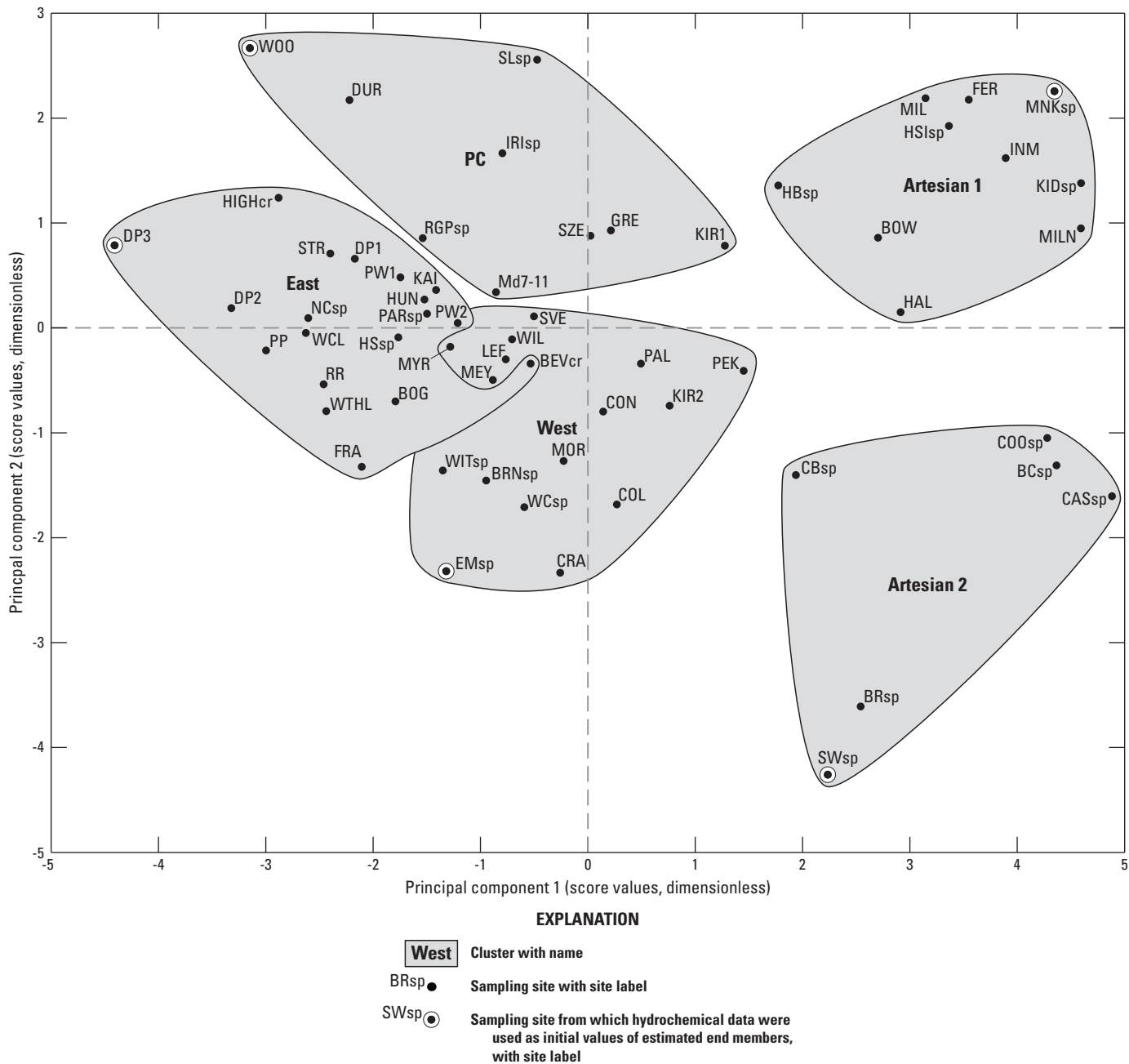


Figure 14. Results of principal component analysis (PCA) and cluster analysis for site-averaged data plotted on the first two principal component axes. The scores are the plotting positions for sites sampled.

thus primarily distinguish the Artesian 1 and 2 clusters from the three clusters to the north and secondarily distinguish clusters that are separated on principal component 2. Variables that plot proximally on figure 16 indicate correlation (Davis, 2002). For example, the conservative tracers $\delta^{18}\text{O}$ and $\delta^2\text{H}$ are correlated with each other and inversely correlated with sodium and chloride. The variables calcium, sulfate, and specific conductance also are correlated.

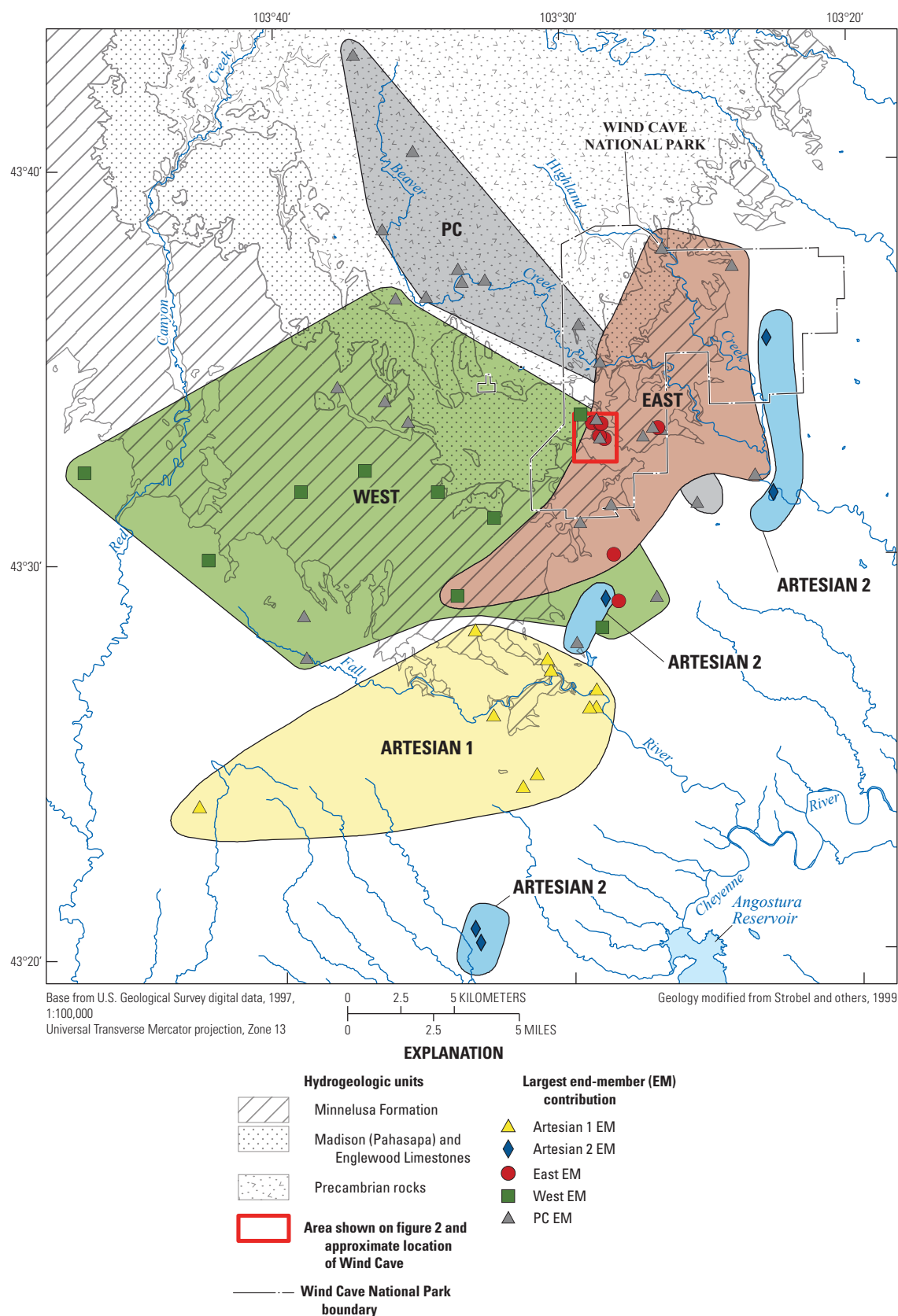


Figure 15. Results of cluster analysis showing the geographic locations of cluster areas. Results of end-member mixing are shown as the largest end-member contribution for each site in relation to the clusters.

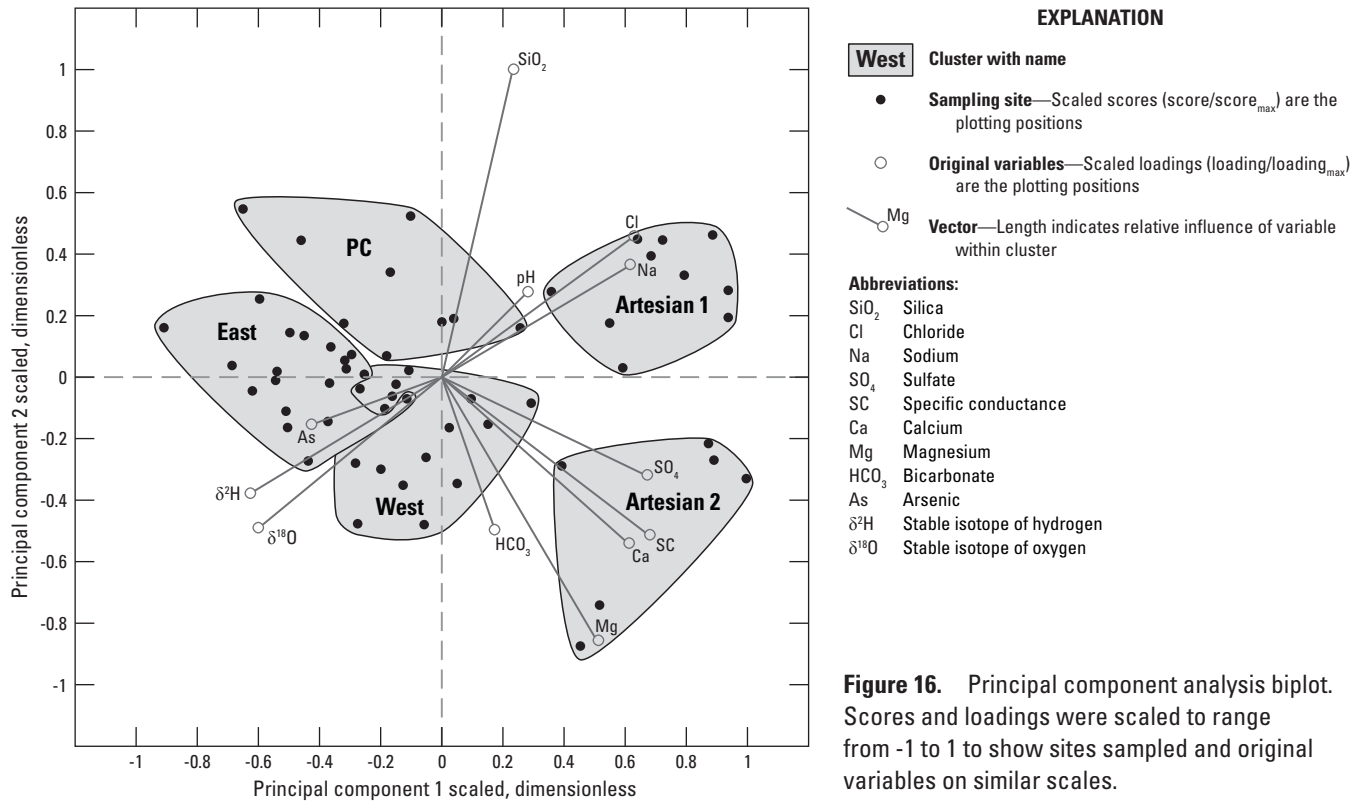


Figure 16. Principal component analysis biplot. Scores and loadings were scaled to range from -1 to 1 to show sites sampled and original variables on similar scales.

End-Member Mixing

For inverse modeling, initial values of $f_{i,k}$ and $E_{j,k}$ (equation 2) that are proximal to optimized values are desirable; otherwise these parameter values might be optimized to satisfy local minima of residuals, rather than the true minimum (Doherty, 2005). Initial values for estimated end-member concentrations $E_{j,k}$ were determined by PCA and cluster analysis by using the hydrochemistry of the site within each cluster that is farthest from the origin on figure 14. Such extreme-value points commonly are assumed to be end members (Christophersen and Hooper, 1992; Laaksoharju and others, 1999). During inverse modeling, end-member values were constrained to within 10 percent (plus or minus) of the total range of observation data for all sites.

As stated previously in the “Sample Collection and Hydrochemical Data” section, 12 variables were used in the PCA and end-member mixing: calcium, magnesium, sodium, bicarbonate, chloride, silica, sulfate, arsenic, $\delta^{18}\text{O}$, $\delta^2\text{H}$, specific conductance, and pH. Model error in end-member mixing is introduced by neglecting geochemical evolution along flow paths from the interaction of rocks and water. If only the conservative tracers ($\delta^{18}\text{O}$, $\delta^2\text{H}$, and chloride) are included in end-member mixing, then geochemical evolution need not be considered, but the model would be poorly constrained with only three variables for observation data. The model would be better constrained with 12 variables but would neglect geochemical evolution for 9 of the 12 variables. As a compromise,

all 12 variables were used, but larger residuals were allowed for nonconservative variables than for conservative tracers. This was accomplished by preferentially weighting the calibration data (observed values) of conservative tracers by a factor of 2.0 for inverse modeling.

Water contributions from the Precambrian aquifer to the Madison and Minnelusa aquifers result from streams that drain Precambrian rocks, receive base flow from the Precambrian aquifer, and then sink into the Madison and Minnelusa aquifers. Precambrian aquifer contributions also might occur by direct groundwater underflow. However, flow in the opposite direction is not possible. Therefore, the end-member mixing model was constrained so that Precambrian aquifer sites could have contributions only from the PC end member, but that this end member could contribute to all sites. Mixing proportions for Precambrian aquifer sites were not estimated by inverse modeling but were set as fixed values, where the PC end-member proportion was set to 100 percent and other end-member contributions to zero. Also, it was assumed that the unsaturated part of the Minnelusa aquifer at this location could receive water only from proximal sources; that is, the East, West, or PC end members. Therefore, constraints were placed on mixing proportions for cave drip sites to satisfy this assumption.

Model constraints, such as this, based on hydrologic information are known as “prior information” in inverse modeling. This reduced the number of parameter values to optimize from 300 to 255 and resulted in a well constrained model

with 2.8 times as many observed values (720) as parameter values to be optimized. This effectively forced the PC end member to have final estimated values that are approximately the average of observed values for Precambrian aquifer samples. Also, this forced all simulated Precambrian aquifer sites to have exactly the same values as this end member, which was not desirable but was considered the best of all options.

After inverse modeling, the coefficient of determination (R^2) from a linear regression model (Davis, 2002) of observed and calculated values was 0.94 for conservative tracers and 0.74 for the other nine variables. Differences, or residuals, from the regression lines are interpreted as resulting from four sources of errors: data errors, model errors, temporal variability, or processes other than pure mixing (that is, geochemical evolution along flow paths). Conservative tracers were assumed to be unaffected by geochemical evolution, and thus regression residuals for these tracers represent the former three sources of error only, whereas residuals for nonconservative variables represent all four sources of error. Therefore, the reduction in the R^2 value for nonconservative variables (21 percent smaller than for conservative tracers) is a model estimate of the effect of geochemical evolution relative to that of mixing. Grouping the sites by the largest end-member contribution resulted in 65 percent similarity to the cluster-analysis groupings (fig. 15), which indicates that 35 percent of the sites received their largest contribution from end members outside of their corresponding clusters.

End-Member Mixing Evaluation

Each end member was assumed to represent a different hydrogeologic domain, and thus, PCA was used to determine how each estimated end member paired with each cluster (table 7). The proximity of end members to clusters in principal-component space was the criterion. For all hydrogeologic domains except East, the largest end-member

contribution was from its own end member. The largest contribution to the East domain was from the PC end member, and the West domain received equal amounts from the West and PC end members (table 8). The East and West end members are interpreted to represent surface recharge to the Madison and Minnelusa aquifers (table 7), and thus end-member mixing estimated that the Precambrian aquifer contribution is larger than that from local surface recharge for the East domain and about equal for the West domain. The Artesian 1 and 2 end members combined (or regional groundwater flow) contributed 60 and 67 percent to the Artesian 1 and 2 domains, respectively (table 8).

The Artesian 1 and 2 end members are interpreted to represent regional groundwater flow, primarily from the Madison aquifer, because they primarily contribute to artesian springs that have relatively large discharge rates (tables 7 and 8). The Artesian 1 and 2 end members contributed from 1 to 7 percent to the East and West domains (table 8), which is consistent with the interpretation that artesian springs discharge regional groundwater flow that probably has little effect on the East and West domains. The West end member contributed about 1.5 times as much as the East end member to the Artesian 1 and 2 domains (table 8), which is consistent with general southward and eastward groundwater flow in the study area (fig. 1).

Mean end-member contributions for Wind Cave sites (table 9) are similar to those of the entire East domain (table 8), except that the largest contribution for the cave sites was from the East end member, or local surface recharge (38 percent), and the second largest was from the PC end member (34 percent). This analysis indicates that contamination to the Precambrian aquifer has potential to affect groundwater in Wind Cave. The West end member, or surface recharge from the west, contributed 26 percent. The Artesian 1 and 2 end-member contributions were 0 and 4 percent, respectively, to cave water bodies and 0 percent to cave drip sites

Table 7. Physical interpretation of end-member waters.

End member	Description of end member
PC	Precambrian aquifer.
East	Recharge to exposed areas of Madison and Minnelusa aquifers on the eastern side of the Black Hills.
West	Recharge to exposed areas of Madison and Minnelusa aquifers on the western side of the Black Hills.
Artesian 1	Regional Madison aquifer groundwater primarily contributing to the Artesian 1 domain.
Artesian 2	Regional Madison aquifer groundwater primarily contributing to the Artesian 2 domain.

Table 8. End-member contributions as an average to each cluster area.

[Shaded areas indicate the largest end-member contribution to each domain. PC, Precambrian aquifer]

Hydro-geologic domain	End-member (EM) contributions, in percent				
	PC EM	East EM	West EM	Artesian 1 EM	Artesian 2 EM
PC	95	3	2	0	1
East	39	33	25	1	3
West	34	21	34	4	7
Artesian 1	13	9	19	56	4
Artesian 2	10	9	15	14	53

Table 9. End-member contributions for Wind Cave sites.

[PC, Precambrian aquifer]

Site name	Site type	End-member (EM) contributions, in percent				
		PC EM	East EM	West EM	Artesian 1 EM	Artesian 2 EM
DP1	Cave drip	27	41	30	0	0
DP2	Cave drip	32	44	23	0	0
DP3	Cave drip	45	39	17	0	0
Drip mean		35	41	24	0	0
PP	Cave water body	33	34	30	0	3
RR	Cave water body	34	33	28	0	5
WCL	Cave water body	32	35	29	1	3
WTHL	Cave water body	30	34	31	0	5
Water-body mean		32	34	29	0	4
Overall mean		34	38	26	0	2

because of model constraints (table 9). This small contribution from the Artesian 2 end member to cave water bodies is possible because it represents regional flow from the west, which might contribute a small amount to the cave depending on anisotropic permeability.

Artesian springs were separated into two clusters because of differences in hydrochemistry, but the hydrogeologic distinction between these two groups of springs is not obvious. Values of temperature, specific conductance, and calcium plus sulfate for artesian springs were distinctly different between the two hydrogeologic domains (table 10). The reason for the differences in temperature is uncertain but might result from a geothermal anomaly in deep rocks near the Artesian 1 springs that affected upwelling water discharging from these springs. Geographically, the Artesian 1 domain is distinct from the Artesian 2 domain because it contains all springs near the Fall River in a small area, which combined with hydrochemical and thermal data, provides a basis for assuming that different hydrogeologic processes are occurring in these two areas. The Artesian 1 and 2 clusters are separated on principal component 2 and account for the largest variability of this principal component (fig. 14).

Back (2011) estimated that Beaver Creek Spring (site BCsp) had a contribution of 55–58 percent regional flow by using geochemical modeling and assuming that the spring was a mixture of regional flow, represented by well MIL, and local recharge, represented by Wind Cave water bodies. The end-member mixing model in the current study described in this report resulted in an estimate of 74 percent regional flow as a combination of the two regional-flow end members (Artesian 1 and 2) and 26 percent local recharge as a combination of the other three end members. The estimates from these two methods did not differ greatly, especially considering differences in the two methods, both of which have strengths

and limitations. The approach used by Back (2011) simulated geochemical evolution by physical processes along two flow paths originating from two end members that were assumed to be represented by the samples selected. In contrast, the approach described in this report allowed for as many as five end members, which were estimated and not assumed to have been sampled, but did not simulate geochemical evolution by physical processes.

The separation on figure 14 of the Artesian 1 and 2 clusters from the other three clusters indicates distinct hydrochemical differences; this does not, however, indicate hydraulic separation between the different hydrogeologic domains. Although the primary sources of groundwater and the groundwater flow paths to the different hydrogeologic domains may differ, the hydraulic head within any of these domains could affect the hydraulic head in the other domains. Results of the end-member mixing model indicate that all hydrogeologic domains (excluding Precambrian aquifer sites) received water from all upgradient end members (table 8), which indicates that the five domains all are hydraulically connected to some extent. For example, the Artesian 2 domain received 34 percent of its contribution from a combination of the PC, East, and West end members (table 8). This is consistent with the conclusions of Back (2011), which indicate that Beaver Creek Spring (site BCsp) is a mixture of local recharge and regional groundwater flow. Beaver Creek Spring and well STR have a spatial separation of only 1.2 km (fig. 1) but are within different hydrogeologic domains (Artesian 2 and East, respectively) and are hydrochemically dissimilar, as indicated on figure 14 by their distant plotting positions. Despite these hydrochemical differences, some degree of hydraulic connection between these two sites probably exists because of similar contributing end members indicated by the end-member mixing model: Beaver Creek Spring received 24 percent of its contribution

Table 10. Temperature, specific conductance, and calcium plus sulfate in water samples from artesian springs.[°C, degrees Celsius; $\mu\text{S}/\text{cm}$, microsiemens per centimeter at 25°C; mg/L, milligrams per liter]

Constituent	Mean	Minimum	Maximum
Artesian 1 hydrogeologic domain ^a			
Temperature, °C	26	24	32
Specific conductance, $\mu\text{S}/\text{cm}$	961	672	1,580
Calcium plus sulfate, mg/L	254	131	660
Artesian 2 hydrogeologic domain ^b			
Temperature, °C	19	18	19
Specific conductance, $\mu\text{S}/\text{cm}$	2,315	1,960	2,560
Calcium plus sulfate, mg/L	1,738	1,302	2,108

^a 4 springs, 12 samples.^b 3 springs, 7 samples.

from the PC and West end members combined, and well STR received 74 percent from these same end members.

Groundwater Age Dating

Age dating was useful for additional evaluation of end-member mixing results. Sample concentrations of the dissolved gasses argon, carbon dioxide, methane, nitrogen, and oxygen (table 11) were used to estimate the temperature of recharge water to improve the estimates of CFC-based groundwater age (table 4) and to assess possible CFC degradation by applying methods described by Plummer and Busenberg (2000). Groundwater samples commonly contain a mixture of water of different ages. The apparent age is an estimate of age that is based on the assumption that the sample contains water of a single age. Because any single sample from the Madison aquifer generally contains a mixture of ages ranging from 1 to more than 60 years (Long and Putnam, 2006), the apparent age is not useful for describing groundwater flow to a single well or spring. However, the apparent age is useful because it is assumed to be proportional to the average age of a groundwater sample, and thus, the apparent age is useful for assessing the general geospatial distribution of groundwater age.

Apparent ages for the Madison aquifer in the study area indicate that groundwater closest to surface recharge areas has the youngest apparent ages, with increasing apparent ages in deeper parts of the Madison and Minnelusa aquifers to the south (fig. 17). The geospatial distribution of apparent ages is consistent with that of specific conductance (fig. 8), where specific conductance generally is proportional to apparent age. Specific conductance increases with residence time in

carbonate aquifers because of the dissolution of aquifer rocks with time. Apparent ages generally are consistent with results of end-member mixing. Sites within the East and West hydrogeologic domains have the youngest apparent ages, and these domains were estimated to receive 90–100 percent of their contributions from local surface recharge or from the shallow Precambrian aquifer (table 8). Sites within the Artesian 1 and 2 hydrogeologic domains have the oldest apparent ages, and these domains were estimated to receive 60–67 percent of their water from end members representing regional flow (table 8). Apparent groundwater ages for Wind Cave are more than 20 years, which might be a result of partial groundwater inflow from the Precambrian aquifer and is consistent with results of the end-member mixing model.

Water from well MIL had the oldest apparent age of all samples, which indicates a regional source for this well and is consistent with end-member mixing results. Based on flow directions indicated by Back and others (1983), the regional source for this well probably is groundwater recharge in the northwestern part of the study area and also areas outside of the study area to the northwest. Also indicated by Back and others (1983), this regional flow sweeps around the southern end of the Black Hills and flows to the east, which would indicate that Cascade and Cool Springs (sites CASp and COOp) probably receive water from this regional source. The younger groundwater age for these springs relative to well MIL indicates that the springs also have contributions from water recharged at a nearer source, from faster flow paths, or both. A possible explanation for this difference in water sources might be karst conduits in the Madison aquifer that connect springs to local recharge areas. The Artesian 1 and 2 hydrogeologic domains each were estimated to receive 41 and 34 percent, respectively, of their water from the PC, East, and West clusters combined (table 8).

The apparent age for water from well Md7–11 was 61 years for CFC–12, which was much older than that of other proximal sites (table 4). An apparent age of 30 to 40 years might be expected for well Md7–11 based on apparent ages for proximal sites and its location near the Madison Limestone outcrop. This difference might be because this well is an observation well and not used for water production. Wells used for water production commonly are developed for maximum yield during installation. In a karst aquifer, this might result in the ability of the well to reach faster groundwater flow paths, such as conduits, than otherwise would be the case. Such well development is not necessary for an observation well that is used only to measure water levels. Therefore, water sampled from an observation well might represent water moving in a slower flow path with an older apparent age than for a production well. All wells other than Md7–11 were pumped regularly. Therefore, the apparent age for well Md7–11 was not plotted on figure 17 because it would result in a potentially misleading anomaly.

Methane was present in the samples from well Md7–11, which could cause degradation of CFCs resulting in a lowering of the concentration, particularly for CFC–11 (Plummer

Table 11. Dissolved gas data for water samples.

[°C, degrees Celsius; mg/L, milligrams per liter; m, meters]

Site name	Station number	Station name	Date	Time	Sample temperature (°C)	Argon (mg/L)	Carbon dioxide (mg/L)	Methane (mg/L)	Nitrogen (mg/L)	Oxygen (mg/L)	Approximate recharge elevation (m)	Temperature of recharge water ^a (°C)
CASp	432006103330501	8S 5E20CDAB	06/19/2009	1135	19.31	0.58	38.53	0.00	16.12	3.38	1,830	9.7
CASp	432006103330501	8S 5E20CDAB	06/19/2009	1135	19.31	.57	38.28	.00	15.92	3.28	1,830	10.1
COOsp	432028103331601	8S 5E20BDCB	06/19/2009	1438	17.78	.56	31.78	.00	15.28	4.51	1,830	9.8
COOsp	432028103331601	8S 5E20BDCB	06/19/2009	1438	17.78	.56	31.71	.00	15.20	4.60	1,830	10.1
KIDsp	432605103285401	7S 5E14DDD	06/19/2009	1645	27.77	.47	30.47	.00	12.31	3.90	1,830	16.3
KIDsp	432605103285401	7S 5E14DDD	06/19/2009	1645	27.77	.47	29.85	.00	11.99	3.81	1,830	16.2
MNKsp	432605103290901	7S 5E14D	05/28/2009	1510	32.43	.56	27.92	.00	15.15	3.55	1,830	10.8
MNKsp	432605103290901	7S 5E14D	05/28/2009	1510	32.43	.55	27.88	.00	15.07	3.49	1,830	10.9
HSIsp	432632103285302	7S 5E14DDDA	05/21/2009	1602	27.38	.51	21.87	.00	13.48	4.25	1,830	12.9
HSIsp	432632103285302	7S 5E14DDDA	05/21/2009	1602	27.38	.51	21.30	.00	13.50	4.56	1,830	13.1
HBsp	432703103302801	7S 5E10DCBA	04/20/2007	1726	23.90	.55	18.14	.00	14.25	6.32	1,830	9.7
HBsp	432703103302801	7S 5E10DCBA	04/20/2007	1726	23.90	.54	18.17	.00	14.11	6.31	1,830	9.9
HBsp	432703103302801	7S 5E10DCBA	05/28/2009	1339	23.62	.57	18.60	.00	15.48	5.71	1,830	9.2
HBsp	432703103302801	7S 5E10DCBA	05/28/2009	1339	23.62	.57	18.57	.00	15.50	5.66	1,830	9.3
SVE	432825103391201	7S 4E 4BACA	09/02/2009	1040	18.12	.59	17.69	.00	16.03	7.39	1,830	8.1
SVE	432825103391201	7S 4E 4BACA	09/02/2009	1040	18.12	.59	17.91	.00	16.06	7.10	1,830	8.3
HUN	433034103284701	6S 5E23DBBB	04/17/2007	940	14.98	.62	10.20	.00	17.27	7.04	1,830	7.3
HUN	433034103284701	6S 5E23DBBB	04/17/2007	940	14.98	.61	10.11	.00	17.19	6.67	1,830	7.6
Md7–11	433115103251401	6S 6E21BBBB (CU91A)	04/20/2007	1350	13.40	.61	9.05	.01	17.10	1.93	1,830	8.4
Md7–11	433115103251401	6S 6E21BBBB (CU91A)	04/20/2007	1350	13.40	.60	8.80	.01	16.93	1.85	1,830	8.5
BCsp	433128103223401	6S 6E14CDB	04/17/2007	1230	19.12	.58	31.49	.00	16.04	3.07	1,830	9.9
BCsp	433128103223401	6S 6E14CDB	04/17/2007	1230	19.12	.57	31.81	.00	15.97	3.01	1,830	10.1
BCsp	433128103223401	6S 6E14CDB	05/28/2009	1020	18.48	.58	30.49	.00	16.05	3.04	1,830	10.2
BCsp	433128103223401	6S 6E14CDB	05/28/2009	1020	18.48	.58	30.72	.00	16.14	2.98	1,830	9.7
STR	433150103230501	6S 6E15ABDD	04/17/2007	1430	20.02	.61	6.91	.00	16.99	7.16	1,830	8.0
STR	433150103230501	6S 6E15ABDD	04/17/2007	1430	20.02	.61	6.62	.00	16.94	7.30	1,830	8.2
STR	433150103230501	6S 6E15ABDD	05/22/2009	1117	19.24	.61	8.50	.00	17.32	5.18	1,830	8.0
STR	433150103230501	6S 6E15ABDD	05/22/2009	1117	19.24	.61	8.56	.00	17.34	5.29	1,830	8.3
PW1	433311103263101	6S 6E 6DDAB	07/29/2009	1325	14.66	.61	9.14	.00	17.23	6.13	1,520	9.7
PW1	433311103263101	6S 6E 6DDAB	07/29/2009	1330	14.66	.61	9.15	.00	17.22	6.21	1,520	9.6

Table 11. Dissolved gas data for water samples.—Continued

[°C, degrees Celsius; mg/L, milligrams per liter; m, meters]

Site name	Station number	Station name	Date	Time	Sample temperature (°C)	Argon (mg/L)	Carbon dioxide (mg/L)	Methane (mg/L)	Nitrogen (mg/L)	Oxygen (mg/L)	Approximate recharge elevation (m)	Temperature of recharge water ^a (°C)
PW2	433311103263102	6S 6E 6DDAA	07/29/2009	1010	14.49	0.82	7.46	0.00	28.29	3.85	1,520	7.4
PW2	433311103263102	6S 6E 6DDAA	07/29/2009	1015	14.49	.82	7.45	.00	28.30	3.89	1,520	7.0
CON	433326103352001	6S 4E 1DBBB	07/15/2009	942	12.49	.63	41.14	.00	18.09	4.44	1,400	9.2
CON	433326103352001	6S 4E 1DBBB	07/15/2009	942	12.46	.63	41.95	.00	18.14	4.17	1,400	9.2
WOO	433636103343901	5S 5E18CCAC	04/26/2007	1730	15.22	.58	8.73	.00	15.86	8.76	1,830	9.3
WOO	433636103343901	5S 5E18CCAC	04/26/2007	1730	15.22	.58	8.71	.00	15.85	8.69	1,830	9.1
WOO	433636103343901	5S 5E18CCAC	07/15/2009	1422	15.90	.59	11.79	.00	16.26	9.06	1,830	9.0
WOO	433636103343901	5S 5E18CCAC	07/15/2009	1422	15.90	.59	11.68	.00	16.12	9.14	1,830	9.0

^a Determined by methods described by Plummer and Busenberg (2000).

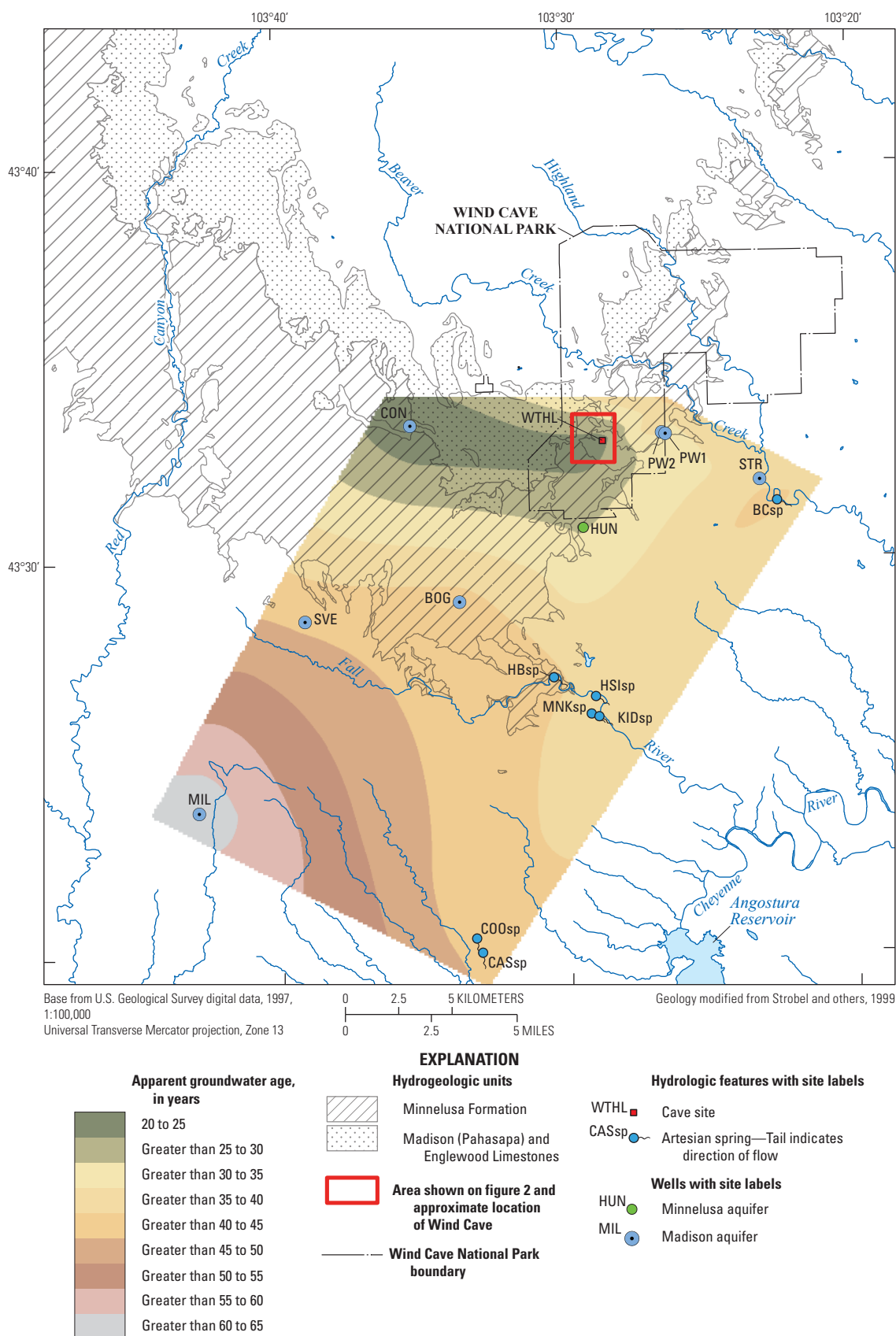


Figure 17. Geospatial distribution of apparent groundwater ages for the Madison aquifer estimated from CFC-12.

and Busenberg, 2000). This would result in an apparent age that is older than for the case of no CFC degradation, and the apparent ages for Md7–11 are, therefore, considered to be the minimum apparent ages. Well Md7–11 is under hydraulic pressure at the land surface and sealed at the wellhead. To prevent freezing of the wellhead during winter, an antifreeze solution that is lighter than water is kept inside the top several feet of the well casing. Although the antifreeze solution was removed, and the well was allowed to flow overnight to clear the well before sampling, it is possible that the methane was a result of this antifreeze solution.

Summary and Conclusions

A study of groundwater flow, quality, and mixing in relation to Wind Cave National Park in western South Dakota was conducted during 2007–10 by the U.S. Geological Survey in cooperation with the National Park Service because of water-quality concerns and to determine possible sources of groundwater contamination in the Wind Cave National Park area. Wind Cave is a primary natural resource for the park, and groundwater in the cave exists as cave drip at numerous locations and in underground streams, pools, and lakes. A large area surrounding the park was included in this study because to understand groundwater in the park, a general understanding of groundwater hydrochemistry and flow in the surrounding area of the southern Black Hills is necessary. Three primary aquifers interact with groundwater in Wind Cave: the Minnelusa, Madison and Precambrian aquifers. The Minnelusa aquifer is contained in the Minnelusa Formation, which is composed of interbedded limestone, dolostone, sandstone, and shale. The Madison Limestone and underlying Englewood Limestone contain Wind Cave and the Madison aquifer, which is a karstic aquifer. The Precambrian aquifer is contained in the underlying metamorphic and igneous fractured rocks of Precambrian age.

Groundwater and surface-water samples collected for this study were analyzed for common ions (calcium, magnesium, sodium, bicarbonate, chloride, silica, and sulfate), arsenic, stable isotopes of oxygen and hydrogen, specific conductance, and pH. These 12 variables were used in all multivariate methods. A total of 100 samples were collected from 60 sites from 2007 to 2010 and included stream sinks, cave drip, cave water bodies, springs, and wells. Groundwater samples were collected and analyzed for chlorofluorocarbons (CFCs), dissolved gasses (argon, carbon dioxide, methane, nitrogen, and oxygen), and tritium at selected sites and used to estimate groundwater age.

Multivariate methods were applied to hydrochemical data to characterize groundwater flow and mixing for these three aquifers. The first of these methods is principal component analysis (PCA), which is used to graphically plot complex multivariate datasets and elucidate data patterns that otherwise might not be noticed. The second method is cluster analysis,

which consists of the assignment of data points to a specified number of groups, or clusters, based on similarity of data. Finally, an end-member mixing model was used to estimate groundwater mixing and flow. This model is based on the assumption that each water sample consists of water originating from one or more end members, or sources of input, in varying proportions. Application of these methods provided a basis to assess characteristics important for groundwater quality, including the differentiation of hydrogeologic domains within the study area, sources of groundwater to these domains, and groundwater mixing within these domains.

In general, groundwater flows from the surface recharge area in the northwestern part of study area to the southwest, then sweeps east and then northeast around the southern end of the Black Hills. A low area in the potentiometric surface in the area of Wind Cave likely indicates anomalously high permeability resulting from extensive karst development. Vertical hydraulic gradients between the Madison and Minnelusa aquifers indicates that the primary source of artesian spring flow is the Madison aquifer. Dye tracing indicated hydraulic connectivity of three water bodies in Wind Cave. Fluorescent dye injected into What the Hell Lake reached Rebel River and Windy City Lake with minimum velocities of 9 meters per day.

Specific conductance generally is lowest in recharge areas for the Minnelusa, Madison, and Precambrian aquifers and highest in the southern part of the study area, distant from aquifer recharge areas. Arsenic concentrations in samples collected for this study ranged from 0.28 to 37.1 micrograms per liter (mg/L) with a median value of 6.4 mg/L, and 32 percent of these concentrations exceeded 10 mg/L, which is the Maximum Contaminant Level (MCL) established by the U.S. Environmental Protection Agency. The highest arsenic concentrations in and near the study area are approximately coincident with the outcrop of the Minnelusa Formation and likely originated from arsenic in shale layers in the Minnelusa aquifer. Sample concentrations of nitrate plus nitrite were less than 2 milligrams per liter for 92 percent of samples collected, which is not a concern for drinking-water quality. Water samples were collected in the park and analyzed for five trace metals (chromium, copper, lithium, vanadium, and zinc), the concentrations of which did not correlate with arsenic concentrations. Sample concentrations for chromium and copper were less than 1 percent of their respective MCLs established by the U.S. Environmental Protection Agency.

For end-member mixing, end members were not assumed to have been sampled but rather were estimated and constrained by prior hydrologic knowledge in the end-member mixing model. This model was quantified in relation to the five hydrogeologic domains that were determined by PCA and cluster analysis, which focused model results on major hydrologic processes. Conservative tracers (chloride and stable isotopes of hydrogen and oxygen) were weighted preferentially in model calibration, which distributed model errors of optimized values, or residuals, more appropriately than would otherwise be the case. The smallest residuals were for conservative

tracers, which is appropriate because geochemical evolution along groundwater flow paths for nonconservative variables was assumed to result in larger model residuals. The difference in residuals between conservative and nonconservative variables provided an estimate of the effect of geochemical evolution along flow paths relative to that of mixing, which was 21 percent.

Five hydrogeologic domains were distinguishable as a result of PCA, cluster analysis, and prior hydrogeologic knowledge. These are described as (1) the PC domain consisting mainly of Precambrian aquifer sites; (2) the West domain consisting of Madison and Minnelusa aquifer sites mainly located near or within the western surface recharge area of these aquifers; (3) the East domain, which is similar to the West domain, except it includes all cave sites and the two stream sinks; (4) the Artesian 1 domain consisting of all artesian springs near the Fall River and wells open to the Madison and Minnelusa aquifers in that general area; and (5) the Artesian 2 domain consisting of the remaining artesian springs, all of which are located within or near the outcrop of the Spearfish Formation. Each of the five end members estimated by the model were determined by PCA to be hydrochemically similar to one of the five domains. Thus, each domain included one estimated end member, and the end members were interpreted accordingly. The PC end member was interpreted to represent groundwater in the Precambrian aquifer; the East and West end members were interpreted to represent surface recharge in these domains; and the Artesian 1 and 2 end members were interpreted to represent regional groundwater flow contributing primarily to these domains. The end-member mixing model estimated that Wind Cave sites received 38 percent of their groundwater inflow from local surface recharge (East end member), 34 percent from the upgradient Precambrian aquifer, 26 percent originating from surface recharge to the west (West end member), and 2 percent from regional flow. The Artesian 1 and 2 end members combined contributed 60 and 67 percent to the Artesian 1 and 2 domains, respectively. This analysis indicates that contamination to the Precambrian aquifer has potential to affect groundwater in Wind Cave. Flow from the Precambrian aquifer into Wind Cave might occur from sinking streams that drain Precambrian rocks and also from direct groundwater transfer from the Precambrian aquifer, through the Deadwood aquifer, and into the Madison aquifer.

Apparent ages, or model ages, for the Madison aquifer in the study area indicate that groundwater closest to surface recharge areas is youngest, with increasing age in a downgradient direction toward deeper parts of the aquifer. Apparent groundwater ages for Wind Cave are more than 20 years, which might be a result of partial groundwater inflow from the Precambrian aquifer and is consistent with results of the end-member mixing model.

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Supplemental Tables

Table 12. Hydrochemical data for water samples.

[mg/L, milligrams per liter; µS/cm, microsiemens per centimeter at 25 degrees Celsius (°C); µg/L, micrograms per liter; ‰, per thousand; m, meters; --, no data or not applicable]

Site name	Station number	Station name	Source aquifer	Date	Time	Dissolved oxygen (mg/L)	pH	Specific conductance, unfiltered (µS/cm)	Temperature (°C)	Calcium, dissolved (mg/L)	Magnesium, dissolved (mg/L)	Sodium, dissolved (mg/L)	Nitrate + nitrite as nitrogen, dissolved (mg/L)	Acid neutralizing capacity ^a (mg/L as CaCO ₃)	Alkalinity ^b (mg/L as CaCO ₃)	Bicarbonate ^c (mg/L)	Chloride, dissolved (mg/L)	Silica, dissolved (mg/L)	Sulfate, dissolved (mg/L)	Arsenic, dissolved (µg/L)	δ ² H (‰)	δ ¹⁸ O (‰)	Well depth below land surface (m)
Stream sink sites												Stream sink sites											
BEVcr	06402430	Beaver Creek sink ^d	--	04/16/2007	1800	--	8.1	538	--	68.2	21.7	13.7	0.013	196	--	236	14.3	14.8	39.7	3.7	-90.10	-11.92	--
BEVcr	06402430	Beaver Creek sink ^d	--	06/25/2008	1120	--	7.8	563	19.3	72.7	22.9	13.6	--	--	--	278	25.6	16.5	65.6	4.4	--	--	--
BEVcr	06402430	Beaver Creek sink ^d	--	05/22/2009	910	--	7.9	559	11.1	77.6	26.6	13.6	--	--	263	318	22.6	17.0	34.6	4.2	-89.41	-11.70	--
HIGHcr	433745103261900	Highland Creek sink ^e	--	03/15/2007	1230	10.4	8.8	286	11.4	38.9	7.25	7.93	.270	--	--	149	3.06	19.2	14.5	3.3	-87.60	-11.95	--
HIGHcr	433745103261900	Highland Creek sink ^e	--	04/26/2007	1015	10.2	9.2	288	14.8	41.3	7.46	8.24	.161	131	--	146	3.01	17.4	14.6	3.0	-87.70	-11.90	--
HIGHcr	433745103261900	Highland Creek sink ^e	--	07/09/2007	1600	2.8	8.0	315	27.2	43.7	7.80	8.97	.114	138	--	163	3.04	22.4	13.7	3.9	-87.10	-11.78	--
HIGHcr	433745103261900	Highland Creek sink ^e	--	09/26/2007	1450	7.4	8.1	321	15.8	42.0	7.89	8.42	.111	138	--	166	2.90	19.4	15.0	2.4	-88.30	-11.88	--
HIGHcr	433745103261900	Highland Creek sink ^e	--	05/14/2009	1345	9.5	9.2	280	20.4	40.1	7.63	8.39	--	--	112	119	4.33	13.1	14.2	3.5	-87.30	-11.72	--
Cave drip sites												Cave drip sites											
DP1	433302103281507	6S 5E12DBAB7	Madison	04/27/2007	1400	9.5	8.3	385	--	35.4	16.9	12.6	1.36	139	--	167	13.2	17.6	21.3	34.2	-90.90	-12.42	--
DP1	433302103281507	6S 5E12DBAB7	Madison	07/29/2009	1130	12.9	7.8	348	--	33.8	19.0	8.27	--	--	146	177	7.25	16.0	22.6	32.4	-90.40	-12.15	--
DP2	433302103281508	6S 5E12DBAB8	Madison	04/25/2007	1200	9.6	8.4	339	--	45.7	13.0	4.29	1.55	157	--	188	3.88	17.5	10.7	29.3	-85.10	-11.69	--
DP2	433302103281508	6S 5E12DBAB8	Madison	05/22/2009	1515	--	--	340	--	47.3	13.1	4.28	--	--	162	195	3.60	18.2	10.6	29.3	-83.90	-11.54	--
DP2	433302103281508	6S 5E12DBAB8	Madison	07/29/2009	1200	12.1	7.9	255	--	50.1	13.3	4.40	--	--	169	205	3.71	17.2	10.7	25.9	-82.20	-11.30	--
DP3	433302103281509	6S 5E12DBAB9	Madison	04/25/2007	1400	9.7	8.4	252	--	37.2	8.19	.97	.455	120	--	144	1.27	19.3	7.44	25.2	-84.40	-11.61	--
DP3	433302103281509	6S 5E12DBAB9	Madison	05/22/2009	1500	--	7.3	255	--	37.7	8.06	.96	--	--	122	149	1.10	20.0	7.08	25.0	-84.00	-11.57	--
DP3	433302103281509	6S 5E12DBAB9	Madison	07/29/2009	1230	9.7	8.0	349	--	38.2	8.04	.95	--	--	125	151	1.17	19.0	7.79	24.3	-83.70	-11.54	--
Cave water body sites												Cave water body sites											
PP	433302103281504	6S 5E12DBAB4	Madison	04/27/2007	1200	9.4	8.1	331	--	34.4	19.8	4.47	0.959	157	--	189	3.07	12.6	6.62	13.1	-91.50	-12.25	--
RR	433302103281506	6S 5E12DBAB6	Madison	04/18/2007	1600	--	7.8	435	13.8	43.2	25.1	4.86	1.29	182	--	221	2.96	13.4	6.76	12.6	-90.70	-12.12	--
WCL	433302103281501	6S 5E12DBAB	Madison	04/18/2007	1430	--	8.0	335	14.8	35.7	19.4	7.67	2.38	149	--	180	4.46	13.9	9.17	16.4	-90.50	-12.15	--
WCL	433302103281501	6S 5E12DBAB	Madison	07/16/2009	1140	--	--	362	14.0	35.8	19.6	7.21	--	--	162	196	4.22	12.2	9.57	12.7	-91.60	-12.09	--
WTHL	433302103281502	6S 5E12DBAB2	Madison	04/18/2007	1230	--	7.8	427	13.8	47.4	27.1	5.08	1.32	211	--	255	3.02	13.2	6.91	12.8	-90.10	-12.11	--
WTHL	433302103281502	6S 5E12DBAB2	Madison	07/16/2009	1245	--	--	411	12.5	43.3	24.1	4.63	--	--	199	241	2.95	12.0	7.57	12.6	-92.50	-12.11	--
Artesian spring sites												Artesian spring sites											
HBsp	432703103302801	7S 5E10DCBA	Minnelusa ^f	03/15/2007	1000	6.8	7.3	683	23.9	63.9	24.7	36.0	0.431	--	--	256	45.6	18.2	69.3	6.6	-113	-15.05	--
HBsp	432703103302801	7S 5E10DCBA	Minnelusa ^f	04/20/2007	1600	6.9	7.0	700	24.0	64.5	25.3	36.9	.417	208	--	253	45.5	18.1	67.5	7.2	-113	-15.07	--
HBsp	432703103302801	7S 5E10DCBA	Minnelusa ^f	07/09/2007	1215	5.2	7.1	718	23.9	64.6	25.5	35.6	.426	206	--	251	45.8	18.9	68.7	4.7	-114	-15.08	--
HBsp	432703103302801	7S 5E10DCBA	Minnelusa ^f	09/27/2007	1100	7.3	7.1	672	23.9	65.4	25.4	36.8	.430	211	--	256	45.0	16.8	67.8	6.3	-115	-15.07	--
HBsp	432703103302801	7S 5E10DCBA	Minnelusa ^f	05/28/2009	1330	--	7.2	710	23.6	69.2	25.7	37.1	--	--	217	264	44.7	18.7	68.9	7.0	-115	-15.03	--
BCsp	433128103223401	6S 6E14CDB	Madison ^g	03/14/2007	1500	4.8	7.3	2,260	18.0	460	76.8	29.2	.409	--	--	219	32.5	16.0	1,280	2.1	-110	-14.42	--
BCsp	433128103223401	6S 6E14CDB	Madison ^g	04/17/2007	1230	6.2	7.1	2,390	19.1	443	73.3	27.4	.390	179	--	217	32.9	14.9	1,290	2.1	-110	-14.40	--
BCsp	433128103223401	6S 6E14CDB	Madison ^g	07/10/2007	1145	--	7.0	--	19.4	446	77.3	30.5	.490	177	--	215	33.8	16.5	1,290	1.7	-110	-14.41	--
BCsp	433128103223401	6S 6E14CDB	Madison ^g	09/26/2007	1320	5.2	6.9	2,350	18.9	478	79.4	30.8	.391	177	--	215	33.1	14.3	1,280	2.0	-109	-14.42	--
BCsp	433128103223401	6S 6E14CDB	Madison ^g	05/28/2009	945	--	6.7	2,350	18.5	477	78.2	29.7	--	--	186	227	34.5	16.6	1,310	1.0	-107	-14.00	--
CASsp	432006103330501	8S 5E20CDAB	Madison ^g	06/19/2009	1100	--	7.0	2,560	19.3	558	83.6	27.8	--	--	190	231	29.3	13.7	1,550	1.2	-117	-15.09	--
COOsp	432028103331601	8S 5E20BCDB	Madison ^g	06/19/2009	1425	--	6.9	1,960	17.8	327	75.3	30.3	--	--	193	235	33.1	14.5	975	2.3	-116	-15.01	--
HSIsp	432632103285302	7S 5E14DDDA	Madison ^g	03/15/2007	1100	5.0	7.3	1,020	27.2	87.5	28.0	70.0	.450	--	--	245	97.5	23.8	167	5.8	-120	-16.02	--
HSIsp	432632103285302	7S 5E14DDDA	Madison ^g	04/17/2007	1630	5.4	7.2	1,030	27.3	89.2	29.1	71.9	.456	198	--	240	99.1	22.7	168	5.7	-120	-16.13	--

Table 12. Hydrochemical data for water samples.—Continued

[mg/L, milligrams per liter; µS/cm, microsiemens per centimeter at 25 degrees Celsius (°C); µg/L, micrograms per liter; ‰, per thousand; m, meters; --, no data or not applicable]

Site name	Station number	Station name	Source aquifer	Date	Time	Dissolved oxygen (mg/L)	pH	Specific conductance, unfiltered (µS/cm)	Temperature (°C)	Calcium, dissolved (mg/L)	Magnesium, dissolved (mg/L)	Sodium, dissolved (mg/L)	Nitrate + nitrite as nitrogen, dissolved (mg/L)	Acid neutralizing capacity ^a (mg/L as CaCO ₃)	Alkalinity ^b (mg/L as CaCO ₃)	Bicarbonate ^c (mg/L)	Chloride, dissolved (mg/L)	Silica, dissolved (mg/L)	Sulfate, dissolved (mg/L)	Arsenic, dissolved (µg/L)	δ ² H (‰)	δ ¹⁸ O (‰)	Well depth below land surface (m)
Artesian spring sites—Continued																							
HSIsp	432632103285302	7S 5E14DDDA	Madison ^g	07/09/2007	1350	5.7	7.0	1,040	27.6	92.2	29.3	71.2	0.487	198	--	241	99.3	24.4	170	3.5	-120	-15.97	--
HSIsp	432632103285302	7S 5E14DDDA	Madison ^g	09/27/2007	1320	5.5	6.8	1,050	27.6	94.1	30.1	74.4	.489	204	--	248	94.0	21.9	174	5.8	-122	-15.91	--
HSIsp	432632103285302	7S 5E14DDDA	Madison ^g	05/21/2009	1600	7.1	7.2	1,030	27.4	96.2	30.1	70.2	--	--	206	251	96.4	25.0	187	6.3	-123	-15.88	--
KIDsp	432605103285401	7S 5E14DDD	Madison ^g	06/19/2009	1630	--	7.0	1,580	27.8	182	38.9	81.8	--	--	180	219	117	22.4	478	3.5	-125	-16.31	--
MNKsp	432605103290901	7S 5E14D	Madison ^g	05/28/2009	1450	--	6.7	1,300	32.4	123	32.9	90.8	--	--	195	238	125	27.4	275	5.2	-127	-16.44	--
BRsp	433525103224401	5S 6E26BBCD	Unknown ^h	08/13/2009	1545	4.5	7.2	2,900	29.6	594	149	16.9	--	--	134	163	3.21	13.8	1,940	2.2	-86.60	-11.11	--
CBsp	432744103293401	7S 5E 11BAAB	Unknown ^h	08/21/2009	1445	9.1	7.0	1,130	14.6	168	49.2	15.1	--	--	235	286	19.6	12.8	388	5.6	-101	-13.28	--
SWsp	432849103283201	6S 5E36CDBB	Unknown ^h	06/17/2010	1600	11.8	7.4	2,500	15.7	502	107	6.78	--	--	172	209	5.09	8.49	1,540	1.1	-90.80	-11.83	--
Shallow spring sites																							
BRNsp	433137103342101	6S 5E18CABD	Minnelusa ^f	06/11/2010	1020	13.8	7.1	568	10.4	59.0	42.2	5.88	--	--	295	359	3.64	11.4	20.5	16.9	-97.70	-12.73	--
EMsp	433332103291801	6S 5E 2ACBD	Minnelusa ^f	08/13/2009	915	6.8	7.4	656	12.7	47.9	61.8	3.10	--	--	358	436	2.04	11.2	20.4	15.3	-95.60	-12.48	--
HSsp	433258103270801	6S 6E 7BAAC	Minnelusa ^f	08/13/2009	1100	4.9	7.1	575	14.8	72.8	11.8	3.98	--	--	221	270	3.59	14.1	9.80	19.5	-95.90	-12.63	--
NCsp	433312103264701	6S 6E 6DCAA	Minnelusa ^f	08/13/2009	1015	7.7	7.2	409	12.8	75.0	6.50	2.88	--	--	210	256	2.40	12.3	7.12	16.9	-95.60	-12.55	--
WCsp	433056103322201	6S 5E21BCBC	Minnelusa ^f	08/21/2009	945	1.5	7.1	689	15.0	67.8	53.9	5.27	--	--	385	469	3.84	13.9	19.5	12.0	-95.20	-12.40	--
WITsp	433611103335801	5S 4E35DDBC	Minnelusa ^f	08/22/2009	1230	10.4	7.2	528	11.6	55.4	23.4	10.2	--	--	223	272	5.30	8.71	24.4	4.5	-90.90	-11.62	--
IRIsp	434249103370401	4S 4E11CCBB	Precambrian	06/17/2010	1420	10.9	6.7	508	14.7	46.9	12.6	13.8	--	--	156	190	10.5	21.2	19.2	2.7	-90.70	-12.06	--
RGPsp	433551103291901	5S 5E23DBCD	Precambrian	08/13/2009	1300	7.0	6.5	372	9.1	55.3	11.9	6.67	--	--	182	222	1.93	17.1	19.7	2.0	-89.50	-12.16	--
SLsp	434020103350101	4S 4E25DB	Precambrian	03/17/2010	1135	7.6	6.5	319	2.1	44.8	8.47	5.62	--	--	123	150	3.14	17.3	26.7	.82	-112	-14.91	--
PARsp	433717103235401	5S 6E15BBAC	White River	08/13/2009	1415	.8	7.2	607	32.4	61.8	18.4	11.5	--	--	258	314	5.15	21.0	8.38	11.1	-87.80	-11.10	--
Well sites																							
BOW	432555103323201	7S 5E20AACA	Minnelusa	05/15/2009	1500	4.6	7.3	890	22.7	99.8	33.9	39.1	--	--	208	253	48.6	18.1	198	3.3	-117	-15.28	152
COL	432727103390201	7S 4E 9BACA	Minnelusa	05/07/2009	1200	6.1	7.2	760	7.9	99.2	41.5	6.49	--	--	224	272	2.83	11.1	201	4.5	-101	-13.08	177
CRA	432846103280501	6S 5E36DCBC	Minnelusa	05/14/2009	1615	8.6	7.4	809	12.8	101	46.9	9.76	--	--	172	210	2.67	9.02	276	10.1	-93.20	-12.27	113
FER	432437103305701	7S 5E27CCCD	Minnelusa	05/08/2009	1325	9.2	7.7	1,100	32.9	97.5	29.4	86.6	--	--	185	224	123	27.5	207	5.6	-126	-16.26	107
FRA	432958103281401	6S 5E24CAAB	Minnelusa	05/08/2009	1535	6.3	7.9	434	10.7	47.2	21.3	15.1	--	--	200	242	3.32	9.00	32.9	37.1	-91.80	-12.08	128
HAL	432720103303701	7S 5E15BABB	Minnelusa	05/08/2009	925	5.6	7.5	985	14.1	112	38.5	46.1	--	--	243	295	67.3	14.4	210	4.0	-118	-15.40	43
HUN	433034103284701	6S 5E23DBBB	Minnelusa	03/14/2007	1100	7.3	7.5	400	15.1	48.2	17.1	9.57	0.589	--	--	235	5.88	15.5	17.3	7.9	-90.40	-12.17	250
HUN	433034103284701	6S 5E23DBBB	Minnelusa	04/17/2007	940	--	7.3	419	15.3	50.5	18.1	10.1	.608	190	--	230	5.97	15.4	17.2	7.7	-92.40	-12.23	250
HUN	433034103284701	6S 5E23DBBB	Minnelusa	09/27/2007	930	8.0	7.2	409	15.5	51.0	17.6	10.0	.596	192	--	233	5.75	15.0	17.0	7.4	-91.10	-12.21	250
HUN	433034103284701	6S 5E23DBBB	Minnelusa	05/21/2009	1510	7.6	7.3	414	13.9	51.1	17.9	9.32	--	--	198	242	5.77	15.2	17.9	6.6	-92.20	-12.18	250
INM	432332103314801	7S 5E33ABDD	Minnelusa	05/07/2009	1500	6.3	7.4	1,190	20.3	110	37.9	83.1	--	--	209	255	119	23.9	247	4.4	-125	-16.22	147
KIR2	433420103374901	5S 4E34BDDB	Minnelusa	06/11/2010	1340	9.3	7.2	771	8.3	107	30.9	13.4	--	--	318	387	41.1	12.9	24.1	3.0	-97.00	-12.83	5
LEF	433215103464401	6S 3E 5CCD	Minnelusa	05/14/2009	1110	6.5	7.5	467	13.2	48.0	26.1	9.70	--	--	224	272	3.31	11.9	21.5	6.5	-106	-13.81	235
MEY	433215103365801	6S 4E11CCDB	Minnelusa	06/11/2010	1510	39.0	7.4	495	10.5	55.8	30.9	6.69	--	--	223	271	9.31	13.4	29.0	18.5	-99.30	-12.88	104
MILN	432806103330801	8S 5E 5CADD	Minnelusa	05/07/2009	1650	4.8	7.0	1,490	23.1	179	52.6	75.1	--	--	242	295	99.2	24.0	433	3.2	-122	-15.91	152
MYR	433003103420701	7S 3E36CBCC	Minnelusa	05/08/2009	1110	12.0	7.5	444	11.2	59.0	20.2	5.27	--	--	225	274	2.79	12.7	13.5	24.6	-109	-14.14	85
PAL	432806103284101	7S 5E 1CBCC	Minnelusa	05/15/2009	1600	7.6	7.0	582	13.1	62.4	29.6	13.1	--	--	222	270	15.9	10.3	61.4	10.6	-109	-14.30	168
PEK	432852103264401	6S 6E31DBDA	Minnelusa	05/21/2009	1410	10.1	7.0	786	13.8	90.6	37.8	19.5	--	--	203	248	16.0	12.5	198	6.1	-107	-14.16	187
WIL	433141103390901	6S 4E16CABB	Minnelusa	06/25/2010	1030	10.5	7.2	479	11.8	53.0	29.8	8.07	--	--	220	268	7.14	14.0	22.7	18.7	-104	-13.32	129

Table 12. Hydrochemical data for water samples.—Continued

[mg/L, milligrams per liter; µS/cm, microsiemens per centimeter at 25 degrees Celsius (°C); µg/L, micrograms per liter; ‰, per thousand; m, meters; --, no data or not applicable]

Site name	Station number	Station name	Source aquifer	Date	Time	Dissolved oxygen (mg/L)	pH	Specific conductance, unfiltered (µS/cm)	Temperature (°C)	Calcium, dissolved (mg/L)	Magnesium, dissolved (mg/L)	Sodium, dissolved (mg/L)	Nitrate + nitrite as nitrogen, dissolved (mg/L)	Acid neutralizing capacity ^a (mg/L as CaCO ₃)	Alkalinity ^b (mg/L as CaCO ₃)	Bicarbonate ^c (mg/L)	Chloride, dissolved (mg/L)	Silica, dissolved (mg/L)	Sulfate, dissolved (mg/L)	Arsenic, dissolved (µg/L)	δ ² H (‰)	δ ¹⁸ O (‰)	Well depth below land surface (m)
Well sites—Continued												Well sites—Continued											
BOG	432858103334201	6S 5E31DAAC	Madison	04/26/2007	1730	8.4	7.8	466	15.2	44.7	29.9	6.29	1.29	217	--	262	2.63	12.7	10.5	15.8	-99.00	-12.90	201
BOG	432858103334201	6S 5E31DAAC	Madison	07/15/2009	1415	--	7.5	444	16.0	41.3	29.1	6.39	--	--	216	263	2.24	11.8	10.0	10.2	-97.20	-12.69	201
CON	433326103352001	6S 4E 1DBBB	Madison	04/23/2007	1200	8.6	7.7	579	8.7	65.4	29.9	10.8	2.24	240	--	292	8.10	12.6	28.9	6.4	-98.60	-12.86	30
CON	433326103352001	6S 4E 1DBBB	Madison	07/15/2009	940	--	7.2	753	12.5	86.3	37.0	14.3	--	--	316	385	17.5	13.0	34.3	3.6	-100	-13.02	30
KAI	433114103281601	6S 5E24BAAA	Madison	07/09/2007	1000	6.8	7.1	435	15.2	51.4	18.5	9.74	.616	188	--	229	5.74	16.3	16.4	6.8	-90.10	-12.24	238
KAI	433114103281601	6S 5E24BAAA	Madison	09/26/2007	1000	8.4	7.1	413	15.1	50.8	18.4	9.86	.626	188	--	228	5.52	14.9	16.4	7.1	-91.60	-12.24	238
Md7-11	433115103251401	6S 6E21BBBB (CU91A)	Madison	04/20/2007	1150	4.0	7.3	403	13.4	40.4	19.6	16.4	.310	170	--	206	5.95	12.8	23.5	.28	-89.70	-11.98	355
MIL	432340103421501	7S 3E36CBDC	Madison	04/23/2007	1430	3.9	7.3	743	21.5	77.8	26.6	37.5	.112	212	--	257	48.0	22.1	110	.99	-125	-16.64	651
MIL	432340103421501	7S 3E36CBDC	Madison	03/17/2010	1500	8.3	7.1	730	--	81.0	26.6	36.7	--	--	213	259	48.4	24.3	98.3	.47	-128	-16.54	651
PW1	433311103263101	6S 6E 6DDAB	Madison	05/24/2007	1230	7.5	7.3	385	13.8	46.2	15.9	12.8	.569	178	--	217	5.10	16.1	17.9	8.5	-89.90	-12.01	242
PW1	433311103263101	6S 6E 6DDAB	Madison	07/29/2009	1245	13.7	7.5	384	14.7	46.1	15.2	12.2	--	--	185	225	5.33	16.0	18.3	8.8	-90.00	-11.94	242
PW2	433311103263102	6S 6E 6DDAA	Madison	05/24/2007	1330	3.9	7.7	586	14.9	24.3	13.5	91.8	.210	223	--	270	20.2	11.1	43.8	26.7	-87.10	-11.88	209
PW2	433311103263102	6S 6E 6DDAA	Madison	07/29/2009	920	7.4	7.8	543	14.5	27.3	13.7	78.5	--	--	223	271	17.4	11.5	38.4	23.0	-88.30	-11.83	209
STR	433150103230501	6S 6E15ABDD	Madison	03/14/2007	1400	6.7	7.7	327	12.8	39.0	13.8	9.08	.511	--	--	189	2.48	16.3	11.2	8.9	-89.30	-11.86	286
STR	433150103230501	6S 6E15ABDD	Madison	04/17/2007	1430	7.4	7.5	343	20.0	39.8	14.1	9.32	.520	157	--	190	2.55	16.4	11.0	8.5	-88.20	-11.87	286
STR	433150103230501	6S 6E15ABDD	Madison	07/10/2007	925	--	7.3	356	23.1	40.5	14.4	9.31	.516	142	--	172	2.56	17.2	10.9	7.6	-88.50	-11.84	286
STR	433150103230501	6S 6E15ABDD	Madison	09/26/2007	1130	.5	6.8	342	16.7	40.0	14.1	9.38	.513	158	--	192	2.47	15.8	11.3	8.9	-87.10	-11.82	286
STR	433150103230501	6S 6E15ABDD	Madison	05/22/2009	1045	--	7.6	330	19.2	40.3	14.3	9.01	--	--	166	201	2.57	17.0	11.4	9.5	-88.30	-11.82	286
SVE	432825103391201	7S 4E 4BACA	Madison	04/23/2007	1800	8.4	7.7	455	12.1	53.0	26.8	4.67	.451	231	--	281	1.86	12.6	8.61	3.9	-114	-15.14	269
SVE	432825103391201	7S 4E 4BACA	Madison	06/26/2008	1305	--	6.5	452	16.6	50.9	25.6	4.33	--	--	--	288	1.82	11.6	8.81	--	--	--	269
SVE	432825103391201	7S 4E 4BACA	Madison	09/02/2009	1000	8.7	7.2	453	18.1	51.9	26.2	4.43	--	--	241	294	1.93	12.0	9.11	3.9	-116	-15.16	269
DUR	433658103332301	5S 5E17BCDB	Precambrian	06/17/2010	1145	8.0	6.7	299	14.5	43.1	7.63	6.66	--	--	130	158	3.82	22.9	14.4	5.7	-87.80	-12.00	85
GRE	433701103323401	5S 5E17ACDD	Precambrian	06/17/2010	1325	9.5	7.0	564	11.3	75.9	18.2	17.7	--	--	208	253	28.8	19.7	36.7	2.6	-92.70	-12.38	32
KIR1	433821103360901	5S 4E 2DDBD	Precambrian	06/17/2010	1020	7.8	6.9	718	7.1	88.3	24.0	24.8	--	--	252	307	53.8	19.3	40.6	1.8	-97.80	-12.84	8
MOR	433635103354301	5S 4E13CCAC	Precambrian	06/25/2010	1200	8.8	7.0	774	11.3	47.3	78.2	14.1	--	--	424	517	8.07	18.8	22.1	35.2	-91.80	-11.95	82
SZE	433718103333101	5S 5E17BBBA	Precambrian	06/03/2010	1455	9.3	7.0	523	12.2	77.0	15.9	11.7	--	--	217	265	15.1	19.4	40.5	1.9	-94.20	-12.63	38
WOO	433636103343901	5S 5E18CCAC	Precambrian	06/11/2010	1215	10.2	6.5	196	9.1	24.8	7.14	5.00	--	--	62	76	4.57	17.3	27.1	12.7	-87.40	-11.99	17

^a Water, unfiltered, inflection-point titration method (incremental titration method), field, milligrams per liter as calcium carbonate.

^b Water, filtered, fixed endpoint (pH 4.5) titration, laboratory, milligrams per liter as calcium carbonate.

^c Calculated from alkalinity by method of Eaton and others (1995).

^d Beaver Creek near Pringle, South Dakota, station 06402430.

^e Highland Creek above Madison outcrop, station 434249103370401.

^f Assumed to flow primarily from Minnelusa aquifer but could include water from other aquifers.

^g Assumed to flow primarily from Madison aquifer but could include water from other aquifers.

^h Possibly artesian flowing from Madison or Minnelusa aquifer or both.

Table 13. Summary of hydrochemical data for the five hydrogeologic domains.

[PC, Precambrian aquifer; SD, standard deviation; mg/L, milligrams per liter; $\mu\text{S}/\text{cm}$, microsiemens per centimeter at 25 degrees Celsius ($^{\circ}\text{C}$); $\mu\text{g}/\text{L}$, micrograms per liter; ‰, per thousand; m, meters; --, no data or not applicable]

Constituent	Units	Artesian 1		Artesian 2		East		West		PC	
		Mean	SD	Mean	SD	Mean	SD	Mean	SD	Mean	SD
Dissolved oxygen	mg/L	6.1	1.4	6.9	2.9	7.8	3.1	10.4	7.8	8.3	2.1
pH		7.2	.2	7.1	.2	7.8	.5	7.2	.3	6.8	.3
Specific conductance	$\mu\text{S}/\text{cm}$	981	271	2,268	495	395	94	605	137	434	160
Temperature	$^{\circ}\text{C}$	25.1	4.4	19.1	4.0	16.1	4.5	12.2	2.6	10.4	4.1
Calcium, dissolved	mg/L	97.3	34.1	445.2	120.7	46.0	12.1	66.4	19.9	55.2	20.8
Magnesium, dissolved	mg/L	31.1	6.9	84.9	26.4	16.2	6.2	36.8	14.7	13.9	5.9
Sodium, dissolved	mg/L	58.8	21.1	24.4	8.4	11.6	16.9	8.9	4.4	12.0	6.8
Bicarbonate ^a	mg/L	253	19	222	30	213	47	319	83	203	70
Chloride, dissolved	mg/L	79.4	31.5	25.7	12.2	5.8	5.8	8.2	9.5	14.2	17.1
Silica, dissolved	mg/L	21.7	3.6	14.2	2.4	15.6	2.9	12.3	2.1	18.6	2.9
Sulfate, dissolved	mg/L	182.3	116.1	1,282.1	402.6	16.5	12.0	55.0	78.0	27.6	9.6
Arsenic, dissolved	$\mu\text{g}/\text{L}$	4.7	1.9	2.1	1.3	13.9	9.7	11.4	8.8	3.4	3.8
$\delta^2\text{H}$	‰	-120.3	4.8	-105.8	10.0	-89.5	3.5	-101.4	7.5	-93.5	7.6
$\delta^{18}\text{O}$	‰	-15.80	.56	-13.80	1.34	-11.99	.36	-13.23	1.02	-12.55	.94
Well depth below land surface	m	120	47	--	--	120	107	128	81	36	30

^a Calculated from alkalinity method of Eaton and others (1995).

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