Geochemical characterization of groundwater evolution south of Grand Canyon, Arizona (USA)

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Abstract
Better characterization of the geochemical evolution of groundwater south of Grand Canyon, Arizona (USA), is needed to understand natural conditions and assess potential effects from breccia-pipe uranium mining in the region. Geochemical signatures of groundwater at 28 sampling locations were evaluated; baseline concentrations for select trace elements (As, B, Ba, Cr, Li, Mo, Rb, Se, Sr, Th, Ti, U, V) were established, and anomalous chemistry characteristics were identified. Concentrations at some groundwater sites exceeded the USEPA drinking water standard for As of 10 μg/L (Red Canyon, Miners, JT, Havasu, and Warm Springs) and U of 30 μg/L (Salt Creek Spring). Four springs from the study area (Blue, Havasu, Fern, and Warm Springs) had unique chemistry, which may indicate a deep flow path or potential contribution of fluids from lower in the crust. Other springs in the study area were distinguished by major anion water type: sulfate, bicarbonate, and a mixture of the two. Water type distinctions were somewhat spatially segregated, with sulfate type present on the western side of the study area, bicarbonate type on the eastern side, and a mixture of the two interspersed between the endmember sites. Sulfate-type water from this study area had low strontium isotopic ratio ($^{87}$Sr/$^{86}$Sr) values. The location of spring discharge within single drainages of the Grand Canyon may influence chemistry, as groundwater discharging from bedrock was altered after flowing through alluvial material. Geochemical analysis of groundwater in Grand Canyon indicates the importance of continued monitoring and better understanding of short-term chemical fluctuations.

Keywords Uranium · Strontium · Geochemistry · Springs · USA

Introduction

On the arid South Rim of the Grand Canyon in Arizona (USA), water is a limited resource. Local populations and ecosystems are dependent on groundwater. The geochemical evolution of the groundwater as it moves through the subsurface, affecting the suitability for consumption, is not well understood. Increased development, uranium mining, and climate change may introduce changes to the groundwater system and focused studies are needed to understand the timing and effects from these changes. In 2012, then US Secretary of the Interior, Ken Salazar, initiated a removal of over 1 million acres in three segregation areas of federal land (north, east, and south) in the Grand Canyon region from new uranium mining activities for the following 20 years, subject to valid existing rights (US Department of the Interior 2012). A key factor in the decision for the withdrawal was the limited amount of scientific data and resulting uncertainties on the
potential effects of uranium mining activities on cultural, biological, and water resources in the area. The US Geological Survey (USGS) has been conducting studies to address the uncertainties of potential uranium mining effects.

**Purpose and scope**

The purpose of this report is to document understanding of the geochemical evolution of groundwater discharging from the Redwall-Muav aquifer south of Grand Canyon. Establishing an understanding of the geochemical signature of the groundwater discharging south of Grand Canyon will provide baseline geochemical information, identify anomalous groundwater samples and their sources, and help elucidate what potential geochemical changes may occur due to breccia-pipe uranium mining. The scope of this report includes an assessment of 28 groundwater sites (25 springs and 3 wells) located south of the Grand Canyon spanning from Blue Spring in the eastern to Warm Spring in the western parts of the study area (Fig. 1).

Targeted samples were collected for this study from 2016 to 2018 and the data were combined with data from previous studies of South-Rim groundwater chemistry by Monroe et al. 2005, Bills et al. 2007, and USGS 2019. Major ion and trace element chemistry was used to characterize the groundwater and evaluate changes in groundwater chemistry as compared to historical sample collection. The dissolved ion chemistry was used to characterize water type and as input for statistical analysis that formally determined the similarities and differences between groundwater along the South Rim and helped develop a conceptual model of groundwater geochemical evolution. Groundwater recharge sources, proposed flow paths, and estimated mean ages (Solder et al. 2020; Solder and Beisner 2020; USGS 2019) were used in conjunction with dissolved ion and isotopic data in conceptual model development. Isotopic ratios of uranium and strontium provided additional evidence of flow paths through stratigraphic sections and the extent of water–rock interaction.

Fig. 1 Map showing the study area including groundwater sample locations and known breccia pipe deposits, some of which are mineralized. Shaded relief represents the topographic elevation changes where the color tan represents lower elevations and purple to white represent higher elevations. Squares represent samples collected between 2016 and 2018 and circles represent samples collected by other USGS studies between 2000 and 2015. Location of mineralized breccia pipes can be found in the following references: Alpine 2010; Brown et al. 1992; Chenoweth 1986; Wenrich et al. 1997; Sutphin and Wenrich 1989; Van Gosen and Wenrich 1989, 1991; Finch et al. 1990; Wenrich 1992; Wenrich and Aumente-Modreski 1994; Gardner 1998; Mazeika 2002.
Hydrogeologic setting

The study area is located in the Colorado Plateau physiographic province. The youngest rocks in the study area are Tertiary volcanic rocks primarily present to the south of Grand Canyon which are underlain by Mesozoic and Paleozoic sedimentary rocks. Groundwater samples in this study were from the Redwall-Muav aquifer (Mississippian to Cambrian in age), which is lower in the Paleozoic stratigraphic sequence. A less significant portion of groundwater is present in the perched Coconino aquifer (Permian in age) near the top of the Paleozoic sequence. Water is thought to recharge the system to the south of Grand Canyon near the San Francisco peaks (Crossey et al. 2009; Bills et al. 2007; Solder et al. 2020; Solder and Beisner 2020) and some additional smaller areas of localized recharge closer to the South Rim have been proposed (Knight and Pool 2016; Solder et al. 2020; Solder and Beisner 2020; Fig. 2). The extensive system of fractures, faults, and karst features in the area control general groundwater flow direction and speed, but the magnitude of influence and location of these features are not well constrained (Huntoon 1974; Jones et al. 2017).

The study of groundwater geochemistry south of Grand Canyon has a rich history. Metzger (1961) assessed the potential water supply in Grand Canyon National Park. Huntoon (1974, 1982) described surface water and groundwater flow of the North and South Rims of Grand Canyon and presented analysis of karst flow in the system. Foust and Hoppe (1985) and Goings (1985) established seasonal trends and baseline values for major and trace dissolved constituents in spring waters. Zukosky (1995) contributed stable isotopic ($\delta^2$H and $\delta^{18}$O) and rare earth element analysis. Fitzgerald (1996) contributed tritium and uranium isotope analysis to the spring data set. Ingraham et al. (2001) explored the possibility of using stable isotopes as a tracer for anthropogenic influence at Indian Garden Spring, and Liebe (2003) focused on uranium...
and uranium isotopes. Monroe et al. (2005) comprehensively explored the hydrogeochemistry of springs discharging along the South Rim of Grand Canyon, adding radiocarbon, strontium isotopes, and an extensive trace element suite. Crossey et al. (2006) utilized noble gas and radiocarbon analysis to understand contribution from fluids lower in the earth’s crust. Bills et al. (2007) built upon the Monroe et al. (2005) study by resampling springs over time. Tobin et al. (2018) provided a review of all hydrogeologic and geochemical studies conducted on groundwater in Grand Canyon.

Breccia pipe deposits

Solution-collapse features known as breccia pipes are found throughout northern Arizona, South Dakota, Wyoming in the USA (Epstein 2005), and internationally with examples from Canada (Ford 1997), Australia (Feltrin and Oliver 2014), Spain (Gutiérrez et al. 2001), China (Yaoru et al. 2002), and Lithuania and England (UK) (Friedman 1997). Breccia pipes in the Grand Canyon region are thought to form by dissolution and karst development in the underlying Redwall Limestone unit, with progressive collapse moving upwards through time into overlying rock units, forming a rubble (breccia) filled column that can be as much as 3,000 ft (~915 m) or more in height (Alpine and Brown 2010; Fig. 3). Breccia pipes are roughly circular in plan view, about 300 ft (~90 m) in diameter, and are often characterized by inward dipping beds along the margins (Otton and Van Gosen 2010). Other collapse features found throughout the Grand Canyon region include sinkholes and localized shallow collapse caused by dissolution of gypsum in the Kaibab and Toroweap formations (Billingsley et al. 2008). In the absence of breccia exposed at the surface, the most effective way to determine if a breccia pipe underlies a collapse feature is by obtaining a geologic core from drilling.

Some breccia pipes contain concentrated deposits of uranium, copper, silver, lead, zinc, cobalt, and nickel minerals (Wenrich 1985; Wenrich et al. 1989; Finch et al. 1992). Uranium mineralization in breccia pipes likely occurred in multiple stages as mineralizing fluids moved through the pipe. Initial reducing conditions within areas of the pipe led to reduction of oxidized uranium, U(VI), and formation of lower-solubility U(IV) minerals, such as uraninite. The dissolved uranium minerals were then deposited when a subsequent mineralizing fluid with differing characteristics moved through the pipe (Wenrich 1985; Huntoon 1996). High-grade uranium ore deposits are most commonly located in breccia adjacent to the Coconino Sandstone, Hermit Formation, and the upper members of the Supai Group (Otton and Van Gosen 2010).

Mining of breccia-pipe-ore zones for copper, lead, zinc, and silver in the Grand Canyon region began in the 1860s, with uranium mining beginning in the 1950s. One of the early mines in the region was the Orphan Lode Mine (located at the head of the Horn Creek drainage and represented by the red diamond south of site 10 on Fig. 1) which was originally developed as a copper mine in 1893. Uranium ore was discovered in the mine workings from the Orphan Lode Mine by the US Geological Survey (USGS) in 1951. The Orphan Mine produced 4.3 million pounds of uranium oxide from 1956 to 1969 and was the largest uranium producer in the region during the early 1950s and 1960s (Chenoweth 1986). The uranium deposits found at the Orphan Lode Mine led to the regional exploration for additional uranium mineralized breccia pipe deposits.

Methods

Data collection

Water samples collected for this study from two wells and 12 spring sites between September 2016 and May 2018, are described in detail in the following. The analytical results were combined with reported data from Red Canyon, JT, Miners, Monument, Lonetree, and Salt Creek Springs (published in Monroe et al. 2005), Fossil Canyon, Matkatamiba, Ruby, Sapphire, Serpentine, and Slate Springs (from Bills et al. 2007), and Warm Springs and Bar 4 Well (USGS 2019; Fig. 1). All samples were collected by the USGS using standard field methods (USGS 2006; 2018), and filtered samples were passed through a 0.45-μm capsule filter. Detailed field and analytical descriptions are published in Beisner et al. (2017).

Field parameters including pH, water temperature, specific conductance, dissolved oxygen, and barometric pressure were measured on site just before water samples were collected. Spring samples were collected as close to the point of issuance from the ground as possible using a peristaltic pump with flexible silicon tubing. Wells were purged of three casing volumes and parameters were within stability criteria prior to collection of water samples (USGS 2006).

Water samples were analyzed for major, trace, and rare-earth elements by the USGS National Research Program Laboratory in Boulder, Colorado (USGSMTCO) using methods from Garbarino and Taylor (1996), (2001), and Garbarino and Taylor (1979). The precision for all methods analyzed by the USGSMTCO was 4% or better depending on the element. Nitrate (NO₃) plus nitrite (NO₂) was analyzed by the USGS National Research Program Laboratory in Menlo Park, California, using methods described in Bullen et al. (1996) with precision of ±0.00002 or better at the 95% confidence level. ALS Environmental in Fort Collins, Colorado, analyzed water samples for uranium isotopes (²³⁴U, ²³⁵U, and ²³⁸U)
using alpha-counting methods (ASTM D 3972) as a USGS contract laboratory.

Quality assurance

One field blank and one replicate were analyzed along with the samples collected between 2016 and 2018. The field blank of certified inorganic-free blank water was collected at the Pipe Spring site in September 2016. A replicate sample was collected from Indian Garden Spring in September 2016. Multiple standard reference samples were analyzed by the USGSTMCO during every analytical run. A detailed analysis of the standard reference sample performance for the respective laboratories is presented in Beisner et al. (2017).

In the field blank from 2016 the following chemical analytes had a value above the reporting level concentration (in parentheses); aluminum (0.4 μg/L), nickel (0.06 μg/L), antimony (0.005 μg/L), tin (0.1 μg/L), and sulfate (0.02 mg/L). The measured concentrations in the blank for the aforementioned constituents were similar to concentrations in blanks collected by the same method reported in Beisner et al. (2017). For the majority of those analytes, the environmental sample values were more than 10 times greater than the blank concentrations and likely do not bias the environmental values. The aluminum value from the blank of 0.4 μg/L is similar to values from several samples and some environmental samples are below the threshold of 10 times the blank value for tin and antimony. The aforementioned blank was analyzed with a batch of nine other samples in the dataset. The reporting level from USGSTMCO changes between batch runs of samples.

The replicate sample from 2016 was within 1% of the associated environmental sample for major ions and the average of trace element difference was 5%. The following elements had greater than a 20% difference between the replicate and...
environmental samples (percent difference in parentheses); beryllium (22%), cadmium (22%), phosphorus (29%), and antimony (57%). A detailed analysis of replicate samples collected from north of Grand Canyon is presented in Beisner et al. (2017).

Data analysis

As the majority of trace elements had one or more values below a laboratory reporting level, with several elements containing multiple reporting levels, specific statistical methods (Helsel 2012) for data with values below a censoring threshold (laboratory reporting level) were used to analyze the majority of analytes presented in this study. Detailed description of the statistical tests is given in Beisner et al. (2017). A $p$-value threshold of 0.05 (95% confidence level) was used to indicate statistical significance for all mentioned statistical tests.

Robust regression on order statistics (ROS) using the cenros function from the NADA software package in R (R Core Team 2018; Lee 2017) was applied to chemical analysis from spring samples for each element to calculate the median, mean, and standard deviation of the data. Groundwater in this study was compared to groundwater north of Grand Canyon (Beisner et al. 2017), using the cendiff function from the NADA package in R (Lee 2017; Helsel and Lee 2006).

Nonmetric multidimensional scaling (NMDS) was performed on the ranks of the uscores (Helsel 2012, 2016) using the metaMDS function from the vegan package in R (Oksanen et al. 2016) using Euclidean distance and two dimensions. Zero dissimilarities were handled by adding a small positive value and no autotransform since there are no community data (Helsel 2012). The following analytes were included in the NMDS: As, B, Ba, Li, Mo, Rb, Se, SO$_4$, Sr, U, V, and Zn; these were chosen based on interest for uranium mining in the region including groundwater north of Grand Canyon (from Beisner et al. 2017) and others that showed large variation across the study area (B, Ba, Rb, and SO$_4$). NMDS stress values ≤0.1 are considered fair with good ordination and no real prospect of misleading interpretation, values ≤0.05 indicate good fit, and values ≥ 0.2 are deemed suspect (Clarke et al. 2014).

The analytes used in the NMDS analysis were evaluated for correlation by calculating Kendall’s Tau using the cenken function from the NADA package in R (Lee 2017). A cluster analysis was used to identify similar groups in the spring samples by evaluating minimum differences within groups and maximum differences among groups using the hcistl function for the analytes used in the NMDS analysis. The Calinski-Harabasz criterion was used to determine the number of clusters that maximizes the difference between clusters, while minimizing the differences within clusters with the cascadeKM function of the vegan package in R (Oksanen et al. 2016).

PHREEQCI version 3.5.0 (Charlton and Parkhurst 2002; Parkhurst and Appelo 2013) was used to determine saturation indices of calcite, dolomite, and gypsum. The WATEQ4F database was used with the default redox potential (pe) value of 4, since none of the water samples indicate reducing conditions.

Results

Parameters and major ions

Groundwater temperature ranged from 9 to 26.5 °C with a median temperature of 18.3 °C. pH ranged from 6.4 to 8.8 with a median of 7.4. Specific conductance ranged from 293 to 4,890 μS/cm with a median of 604 μS/cm. Dissolved oxygen ranged from 3 to 11.4 mg/L in the data set, except for Bar 4 Well which had the lowest dissolved oxygen value (0.7 mg/L), with a median dissolved oxygen of 7.2 mg/L.

The majority of sites can be classified as having calcium-magnesium-bicarbonate type water. Except for Blue Spring, which was dominated by sodium type water, the major cations were calcium and magnesium. For the major anions, the majority of sites were dominated by bicarbonate. Some springs were a mixture of bicarbonate-sulfate (Lonetree, Horn, Salt Creek, Slate, Sapphire, Serpentine, Royal Arch, and 140 Mile Springs). Sulfate groundwaters were generally located on the west side of the study area and include Ruby, Fossil Canyon, Matkatamiba, Bar 4 Well, and National Canyon Springs. Blue Spring was dominantly chloride type water (Fig. 4).

Nitrate ranged from <0.002 to 1.4 mg/L as N, with a median value of 0.51 mg/L as N. All sample concentrations of nitrate were below the US Environmental Protection Agency (USEPA) drinking water maximum contaminant level (MCL) of 10 mg/L (USEPA 2019) except for one elevated value at Monument Spring. This outlier was noted in Monroe et al. 2005 as possibly influenced by wastewater effluent from the South Rim of Grand Canyon or by exposure to geologic features not encountered by water discharging at other springs.

Trace elements

Some groundwater sample concentrations in this study exceeded the USEPA drinking water standards for As and U (Table 1; USEPA 2019). Red Canyon, Miners, JT, Havasu, and Warm Springs waters exceeded the USEPA drinking water standard for As of 10 μg/L (Fig. 5a), and Salt Creek Spring water exceeded the USEPA drinking water standard for U of 30 μg/L (Fig. 5b).

Median and mean trace element concentrations for all 27 groundwater sites south of Grand Canyon in this study (25 springs and 2 wells; Supai Well No. 3 not included as it is primarily supplied by water from Havasu Spring) are listed in
Some elements were not statistically quantified due to the following reasons: the majority of the sample concentrations were below the censoring threshold from blanks in Beisner et al. 2017 (Al, Be, Cd, Cu, Mn, Pb, Sb, and Zn), more than 80% of the sample concentrations were below the highest laboratory reporting level (Co and Fe), or more than 60% of the sample concentrations were below the laboratory reporting level (Ni).

Eleven sites (Blue, JT, Miners, Grapevine, Pipe, Canyon Mine Well, Horn, Salt Creek, Hermit, East Boucher, and National Canyon Springs) had three or more samples over time reported in Monroe et al. 2005, Bills et al. 2007, and USGS 2019. Change of uranium concentration over time at these sites was small, varying between 10 and 30%. There are not enough data to statistically analyze trends over time, but clear monotonic trends are not obvious. Monroe et al. (2005) sampled the majority of the springs presented in this report during different times of the year and found that there was temporal consistency of chemical results for individual springs. Generally, springs discharging south of Grand Canyon presented in this report show less temporal variability.

![Trilinear Piper plot representing the proportional major ion composition of groundwater samples. Squares represent samples collected between 2016 and 2018 and circles represent samples collected by other USGS studies between 2000 and 2015](image_url)
than springs north of Grand Canyon on the margins of the Kaibab Plateau that show rapid response from seasonal recharge pulses (Jones et al. 2017; Tobin et al. 2018).

Radioisotopes

Uranium isotopes

Natural uranium consists of three isotopes $^{238}$U, $^{235}$U, and $^{234}$U with relative abundances of approximately 99.27%, 0.72%, and 0.0057%, respectively. The ratio $^{234}$U/$^{238}$U can vary widely in natural waters, as a result of processes such as alpha recoil, which allows for preferential mobility of $^{234}$U relative to $^{238}$U during interactions between water and solid phases (Faure and Mensing 2005). Uranium in undisturbed rocks and minerals older than approximately 1 million years reaches a state of radioactive equilibrium where the rate of decay of the short-lived $^{234}$U is limited by the rate of decay of the long-lived $^{238}$U parent. As a result, the $^{234}$U/$^{238}$U activity ratio (UAR) is expected to equal unity (defined as secular

Table 1 Statistical values for trace elements from 27 groundwater sites (25 springs and 2 wells) discharging south of Grand Canyon. All values reported in units of µg/L, p-values for ‘cenros’ all less than 0.05 (highest 5.2 × 10^{-5} for Fe), SD standard deviation; Min minimum; Max maximum

| Element | Median | Mean | SD | Min | Max | Percent censored | High outlier sites | Notes of why not to include statistical values
|---------|--------|------|----|-----|-----|------------------|-------------------|-----------------------------------------------------|
| Al      | -      | -    | -  | <0.1| 19  | 19%              | -                 | > 90% of samples below censoring threshold of 4.5 |
| As      | 2.3    | 4.9  | 5.3| 0.46| 17  | 0%               | Red Canyon, Miners [Havasu, JT, Warm - nonoutlier exceedance] |
| B       | 108    | 148  | 186| 24  | 1,000| 0%              | Warm              |
| Ba      | 64     | 111  | 109| 9.6 | 358 | 0%               | Grapevine         |
| Be      | -      | -    | -  | <0.007| 0.04| 41%            | -                 | All samples below censoring threshold of 0.07 |
| Cd      | -      | -    | -  | <0.001| 0.09| 41%            | -                 | >90% of samples below censoring threshold of 0.05 |
| Co      | -      | -    | -  | <0.003| 0.4 | 48%            | -                 | All values less than highest reporting level of 2 |
| Cr      | 0.4    | 0.42 | 0.25| <0.08| 0.95| 26%            | -                 | >90% of samples below censoring threshold of 1.2 |
| Cu      | -      | -    | -  | <0.01| 1.5 | 15%            | -                 | >80% of values less than highest reporting level of 2 |
| Fe      | -      | -    | -  | <0.2 | 5,012| 52%            | -                 | >80% of samples below censoring threshold of 0.7 |
| Li      | 22     | 39   | 53 | 2.5 | 230 | 0%           | Havasu, Fern, Blue, Warm |
| Mn      | -      | -    | -  | <0.03| 83  | 11%            | -                 | >80% of samples below censoring threshold of 0.7 |
| Mo      | 1.9    | 2.5  | 2  | 0.23| 8.8 | 0%             | Bar 4 Well        |
| Ni      | -      | -    | -  | <0.01| 0.7 | 67%            | -                 | >60% of samples were less than the laboratory reporting level |
| Pb      | -      | -    | -  | <0.002| 0.28| 19%            | -                 | >58% of samples below censoring threshold of 0.05 |
| Rb      | 7.9    | 7.9  | 3.9| 2.2 | 17  | 0%             | -                 | >90% of samples below censoring threshold of 0.26 |
| Sb      | -      | -    | -  | 0.006| 0.27| 0%             | -                 | |
| Se      | 4.9    | 6.3  | 4.4| 1.1 | 17  | 0%             | -                 | |
| Sr      | 309    | 714  | 857| 76  | 2,805| 0%            | Matkatamiba, Fossil Canyon, Ruby, National Canyon, Bar 4 Well |
| Th      | 0.001 | 0.003| 0.005| <0.0002| 0.023| 15%            | Matkatamiba, Bar 4 Well |
| Tl      | 0.02  | 0.03 | 0.03| <0.006| 0.15 | 4%            | Blue, Monument, Matkatamiba, Bar 4 Well |
| U       | 4.6    | 6.1  | 6  | 1   | 31  | 0%             | Salt Creeka |
| V       | 1      | 1.5  | 1.6| <0.1| 6.7 | 11%            | Havasu, Fern, Warm |
| Zn      | -      | -    | -  | 0.4 | 43  | 0%             | -                 | >90% of samples below censoring threshold of 20 |

*a Indicates exceedance of USEPA drinking water standard

b Censoring threshold was determined from blanks presented in Beisner et al. (2017)
Fig. 5 Elemental concentration in groundwater samples for a arsenic and b uranium
equilibrium or UAR = 1.0). Water in contact with high-grade uranium ore having a recent history of oxidation and leaching is likely to represent a mixture of material with UARs both greater than and less than 1.0. Typical values for natural groundwaters fall in the 1–3 range but values in excess of 10 are possible (Kronfeld 1974; Osmond and Cowart 1976; Szabo 1982).

The UAR values of groundwater samples from this study generally decreased with increasing uranium concentrations (Fig. 6). The lowest UAR values were from Horn Creek and 140 Mile Springs (1.25 and 1.30 respectively), although the low uranium concentration (less than 10 μg/L) may reflect interaction with a nearby source of uranium that has undergone chemical change or mixing with other waters before discharging at the spring. Salt Creek Spring had the highest uranium value (31 μg/L) from this study and had a correspondingly low UAR of 1.55. The Canyon Mine Well is drilled into the Muav Limestone next to a mineralized breccia pipe deposit and had the second highest uranium value (13 μg/L) and a moderately low UAR (2.1). The mineralized zone of the Canyon Mine breccia pipe occurs in the Coconino Sandstone through the Supai Group, stratigraphically above the screened interval of the Canyon Mine Well.

JT and Miners Springs had low UAR values (1.78 and 1.49, respectively) but low uranium concentration (4.1 and 3.1 μg/L, respectively). The abandoned Grandview Mine (Fig. 1), which was a mineralized breccia pipe mine, is located near Miners Spring and more investigation of water upgradient of the spring emergence would provide additional context for these low UAR values. These two springs have tritium concentrations indicating a possible mixture of modern and premodern waters (Solder et al. 2020).

The ratio of $^{87}$Sr to $^{86}$Sr ($^{87}$Sr/$^{86}$Sr) can provide an indication of water–rock interaction as groundwater moves through the subsurface. Precambrian basement rocks of the region have the most radiogenic (highest) strontium isotopic ratios (greater than 0.714; Crossey et al. 2006, Patchett and Spencer 2001, Johnson et al. 2011, Bills et al. 2007), Tertiary volcanic rocks have the lowest strontium isotopic ratios (0.70463–0.70535; Bills et al. 2007), and Paleozoic sedimentary rocks have intermediate strontium isotopic ratios (0.70756–0.71216; Monroe et al. 2005 and Bills et al. 2007) as illustrated in Fig. 7.

Generally, groundwater samples with a higher strontium concentration had a lower strontium isotopic ratio (Fig. 7). Samples from Matkatamiba, National Canyon Springs, and Bar 4 Well had the highest strontium concentrations (2,400–2,805 μg/L) and the lowest strontium isotopic ratios (0.707853–0.70819) near the average values of the Kaibab and Toroweap Formations. The highest strontium isotopic ratio values were associated with some of the lowest strontium

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![Fig. 6](image-url)  
**Fig. 6** Uranium activity ratio (UAR) versus uranium concentration for groundwater samples.
concentrations in this dataset, some of which (JT, Lonetree, Pipe, Horn, Salt Creek, and Serpentine Springs) are greater than the highest strontium isotopic ratio values from sedimentary rocks from the Grand Canyon (0.71216; Monroe et al. 2005 and Bills et al. 2007). Strontium isotopic values (87Sr/86Sr) are not available for Fossil Canyon, Ruby, Sapphire, Slate, and Warm Springs.

The springs with low strontium concentrations may indicate less water rock interaction and may be influenced by strontium isotopic ratios of dissolved minerals different than the bulk sedimentary rock values from Monroe et al. 2005 and Bills et al. 2007. Of note, the only spring north of the Grand Canyon with a high strontium isotopic ratio and strontium concentration (Fence Spring; 0.71417, 1060 µg/L, respectively) had a distinct chemical signature of geothermal fluids (Crossey et al. 2006; Beisner et al. 2017). Strontium concentrations generally increase with sulfate concentration and may suggest that water in contact with gypsum along its flow path has elevated strontium. Groundwaters north of Grand Canyon include groundwater discharging from stratigraphic units including and above the Redwall Limestone, which show geochemical evidence of interaction with gypsum units in the Toroweap Formation to the Redwall Limestone (Beisner et al. 2017), were compared to 27 groundwater samples collected south of Grand Canyon from this study, to assess regional similarities for select elements (As, B, Li, Se, Sr, Tl, U, V).

As and Sr were significantly different between the north and south groups (p <0.05). As values were higher and Sr values were lower in the group of springs south of Grand Canyon compared with the group of springs north of Grand Canyon. Elevated As concentrations may be related to the volcanic rocks present near the recharge source for water on the South Rim of Grand Canyon (Fig. 2), that are not a present in the presumed recharge area for springs north of Grand Canyon (Beisner et al. 2017). Strontium concentrations generally increase with sulfate concentration and may suggest that water in contact with gypsum along its flow path has elevated strontium. Groundwaters north of Grand Canyon include groundwater discharging from stratigraphic units including and above the Redwall Limestone, which show geochemical evidence of interaction with gypsum units in the Toroweap Formation (Beisner et al. 2017). The other elements (B, Li, Se, Tl, U, and V) were not statistically different between the north and south groups.

Five springs in this study (Pipe, Horn, Indian Gardens, National Canyon, and 140 Mile) discharge from alluvial material rather than directly from bedrock. Changes in trace element chemistry in Horn Creek as water discharges from bedrock and moves through alluvial material have been observed.
Water at the head of the eastern fork of the Horn Creek drainage emerges from bedrock with elevated uranium concentration (257 μg/L), low UAR (1.12), calcium-magnesium-sulfate type water, and low tritium (0.9 pCi/L), then infiltrates into the alluvium and emerges lower in the drainage with lower uranium concentration (23 μg/L), similar UAR (0.95), calcium-magnesium-bicarbonate type water, and elevated tritium (2.6 pCi/L; Beisner and Tillman 2019). Monroe et al. 2005 similarly discussed changes in chemical constituent concentration as groundwater discharging at a higher elevation in Cottonwood and Monument Creek drainages moves through alluvial material, noting increasing concentration of several major ions and decreases of several trace elements and nutrients. Unfortunately, blank and replicate samples were not assessed in the Monroe et al. 2005 study, which limits quantitative assessment of some of the elements from the chemical comparison along a drainage, but similar trends as those observed in Horn Creek by Beisner and Tillman (2019) suggest that additional chemical changes within the drainages are taking place. The mechanisms for the chemical changes within the drainages are not well understood and could vary depending on the local condition and possibly be influenced by mixing with another source of groundwater or geochemical changes as the water moves through unconsolidated material. However, the overall significance is that springs discharging from alluvial material may not have the same chemical signature of the bedrock groundwater, and characterizing secondary processes is critical for understanding Grand Canyon spring discharge chemistry.

**Multivariate trace element analysis**

Multivariate statistical analysis of groundwater south of Grand Canyon shows a good ordination distinction between sites based on their elemental assemblage (Fig. 8). Analytes associated near uranium in the NMDS were Mo, SO₄, Se, and Sr (Fig. 8). These same analytes were related to U with a significant positive Kendall’s Tau value (Fig. 9). Uranium had a significantly negative Kendall’s Tau relationship with Ba and As (indicating that higher values of uranium are associated with lower values of Ba and As), which are located across the NMDS plot from uranium (Fig. 8).

A cluster analysis on the data used in the NMDS analysis using the Calinski-Harabasz criterion indicates that there are three distinct groups of groundwater sites (Fig. 10, and dashed outlines on Fig. 8). Blue, Havasu, Fern, and Warm Springs are in one group and have higher values of B, Li, Rb, and V and lower values of Mo and Se compared to other sites. Lonetree, Horn, Salt Creek, Slate, Sapphire, Ruby, Serpentine, Royal Arch, Fossil Canyon, 140 Mile, Matkatamiba, Bar 4 Well, and National Canyon Spring form another group and are distinguished by higher values of Mo, Se, SO₄, and Sr and lower values of Ba. Red Canyon, JT, Miners, Grapevine, Pipe, Indian Gardens, Monument, Hermit, East Boucher Springs and Canyon Mine Well form another group and are distinguished by lower values of B, Li, SO₄, and Sr. The groundwater site groupings appear to overlap spatially from east to west with clustering of sites locally (Fig. 10).

**Groundwater geochemical evolution**

Multiple geochemical tracers were used to understand recharge source, groundwater residence time, and water–rock interaction of groundwaters analyzed in this study. A companion paper (Solder et al. 2020) and USGS (2019) present environmental-age tracer analysis for groundwaters presented in this report. Groundwater-age tracer analysis is combined with geochemical information and discussed in the following paragraphs. Detailed geochemical modeling was not performed for this study, as there were not enough samples along a flow path from recharge source to discharge available for the study area.

Using the categorical age classification for tritium of <1.3 pCi/L for premodern water, >12.8 pCi/L for modern water, and values between for a mixture of premodern and modern water from Beisner et al. (2017), no samples contained primarily modern water (water recharged after 1952) and the majority of the sites had a mixture of modern and premodern water and some sites were predominantly premodern water (Blue, Havasu, Supai Well No. 3, Fern, Boucher East, Canyon Mine Well, Hermit, Pipe, Indian Gardens Spring, and Bar 4 Well; Bills et al. 2007, Monroe et al. 2005, Solder et al. 2020). Tritium values were not available for Fossil Canyon, Ruby, Sapphire, Slate and Warm Springs (Bills et al. 2007). Spring sites with sulfate or sulfate/bicarbonate water had tritium in the range of a mixture of modern and premodern water and may indicate a component of water that potentially moved through the subsurface gypsum deposits along a shorter flow path. Bar 4 Well did have sulfate water type but no measurable tritium (Bills et al. 2007; USGS 2019).

Solder et al. (2020) calculated mean age for the springs sampled between 2016 and 2018 for this study as well as for groundwater data published in Monroe et al. 2005 and Bills et al. 2007. Sites classified as premodern water had mean ages ranging from approximately 1,050 to 19,200 years. Of the premodern samples, Pipe Spring had the lowest mean age of 1,050 years and the next lowest age was Indian Gardens Spring with 2,750 years. Tritium content was 3.5 pCi/L in Pipe Spring samples collected in 2000 (Monroe et al. 2005), which would indicate a mixture of modern and premodern water and may explain the low mean age compared with other springs with low tritium values. Waters with tritium concentrations indicating a mixture of modern and premodern water had lower mean age ranging from 90 to 2,900 years. Monument Spring had the highest mean age for the mixed
waters of 2,900 years (Monument Spring tritium value was 1.3 pCi/L, which is right at the threshold for premodern water) and the next highest age was Royal Arch Spring with a mean age of 2,400 years.

Dedolomitization may be occurring in some groundwater south of Grand Canyon, and may affect the radiocarbon age calculation (Plummer et al. 1990; Fitzgerald 1996). The calculated saturation index of gypsum was negative for all samples in this study, but values show a logarithmic increase to near saturation with increasing sulfate concentration (Fig. 11). Samples range from Grapevine Spring with 6 mg/L sulfate and SI of gypsum of −3.03 to National Canyon Spring with 991 mg/L sulfate and SI of gypsum of −0.42 (Fig. 11). Five sites (National Canyon, Matkatamiba, Ruby, Fossil Springs, and Bar 4 Well) had high sulfate concentration (>600 mg/L), saturation indices (SI) of gypsum less than 0.7, calcite SI plus or minus 0.25, and dolomite SI plus or minus 0.5, which may indicate dedolomitization

Four springs (Blue, Havasu, Fern, and Warm Spring) had unique chemistry and may represent a contribution of groundwater from deeper in the earth’s crust (Crossey et al. 2006) or long groundwater residence times. These sites are located at discrete locations in the study area near major structural geologic features that may allow for focused groundwater discharge and may also provide a pathway for interaction with deep crustal materials. Characteristics of these waters are premodern tritium values, radiocarbon ages on the order of 4,000–16,000 years (Solder et al. 2020), elevated As, B, Li, Rb, and V concentration, calcium-magnesium-bicarbonate type water (except for Blue Spring with sodium-chloride type water), and intermediate UAR (between 2

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**Fig. 8 Nonmetric multidimensional scaling (NMDS) plot for elemental analysis from groundwater sites**

| Site                  | Explanation                   |
|-----------------------|-------------------------------|
| Blue Spring           | 1                              |
| Canyon Mine Well      | 8                              |
| Red Canyon Spring     | 9                              |
| JT Spring             | 3                              |
| Miners Spring         | 4                              |
| Grapevine Spring      | 5                              |
| Lonetree Spring       | 6                              |
| Pipe Spring           | 7                              |
| Monument Spring       | 16                             |
| Salt Creek Spring     | 11                             |
| Horn Spring           | 10                             |
| Ruby Spring           | 17                             |
| Serpentine Spring     | 19                             |
| Royal Arch Spring     | 19                             |
| Hermil Spring         | 13                             |
| Fossil Canyon Spring  | 20                             |
| Phoenix Creek Spring  | 21                             |
| 140 Mile Spring       | 22                             |

Stress = 0.096

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Strontium isotopic ratio was intermediate for these springs (Table S1 of the ESM), suggesting groundwater chemistry is largely controlled by interaction with Paleozoic carbonate rocks rather than Precambrian basement rocks with more radiogenic strontium isotopic ratios (Crossey et al. 2006).

Groundwaters with a dominant sulfate type water (Slate, Sapphire, Ruby, Fossil Canyon, Matkatamiba, Bar 4 Well, and National Canyon Spring) are located on the western side of the study area (orange group on multivariate analysis on Fig. 10). Gypsum deposits grade thicker from east to west in the study area (Billingsley et al. 2008; P. Huntoon, University of Wyoming, personal communication, 2019) and may explain the predominance of sulfate type water of samples in the west part of the study area (Fig. 4). Three of these sites have strontium isotopic data (National Canyon, Matkatamiba Springs, and Bar 4 Well), which indicate high strontium concentration and low strontium isotopic ratio that may indicate interaction with gypsum (CaSO\(_4\cdot2\)H\(_2\)O) deposits in the Kaibab and or Toroweap formations (Fig. 7). Ruby and Fossil Canyon Springs have elevated strontium concentrations (1,900 and 2,300 μg/L, respectively; Table S1 of the ESM), and may also have low strontium isotopic ratio which warrants follow up sampling. Matkatamiba and National Canyon Springs have tritium indicating a mixture of modern and premodern water (Table S1 of the ESM; Solder et al. 2020). National Canyon had a young mean age of 87 years and Matkatamiba had an intermediate age of 1,200 years. Bar 4 Well had no measurable tritium and an old radiocarbon age of 19,100 years, although the age may be less if dedolomitization is occurring along the flow path before the Bar 4 Well (Solder et al. 2020).

Other sites with a mixture of bicarbonate and sulfate type waters (Lonetree, Horn, Salt Creek, Serpentine, Royal Arch, and 140 Mile Springs; Fig. 4) are also part of the orange multivariate group (Fig. 10) and have tritium values indicating a mixture of modern and premodern water, low UAR (no data for Lonetree and Serpentine), and high strontium isotopic ratio.

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**Fig. 9** Correlation matrix for elemental analysis used in NMDS showing Kendall’s Tau values where blue shading represents positive and red negative correlation. Values with an X across the number represent samples with a p-value > 0.05.

|     | Ca  | Mg  | K   | Na  | Cl  | SO\(_4\) | Si  | Fe  | Al  | Mn  | Li  | Rb  | Sr  | Ba  | As  | V   |
|-----|-----|-----|-----|-----|-----|---------|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| Ca  | 1.00| 0.56| 0.34| 0.48| 0.62| 0.45    | 0.64| 0.62| 0.55| 0.53| 0.68| 0.77| 0.79| 0.75| 0.82| 0.68|
| Mg  | 0.56| 1.00| 0.35| 0.48| 0.53| 0.62    | 0.59| 0.58| 0.53| 0.57| 0.73| 0.79| 0.79| 0.78| 0.81| 0.74|
| K   | 0.34| 0.35| 1.00| 0.32| 0.45| 0.46    | 0.33| 0.33| 0.28| 0.30| 0.43| 0.51| 0.51| 0.51| 0.62| 0.54|
| Na  | 0.48| 0.48| 0.32| 1.00| 0.53| 0.55    | 0.49| 0.43| 0.37| 0.40| 0.46| 0.62| 0.59| 0.52| 0.62| 0.49|
| Cl  | 0.62| 0.53| 0.45| 0.53| 1.00| 0.50    | 0.65| 0.64| 0.60| 0.62| 0.79| 0.77| 0.76| 0.75| 0.81| 0.75|
| SO\(_4\) | 0.45| 0.62| 0.46| 0.55| 0.50| 1.00    | 0.44| 0.49| 0.46| 0.48| 0.62| 0.74| 0.73| 0.74| 0.77| 0.74|
| Si  | 0.59| 0.58| 0.33| 0.43| 0.65| 0.44    | 1.00| 0.58| 0.54| 0.55| 0.68| 0.72| 0.71| 0.71| 0.77| 0.71|
| Fe  | 0.62| 0.58| 0.33| 0.43| 0.64| 0.49    | 0.58| 1.00| 0.54| 0.54| 0.68| 0.72| 0.71| 0.72| 0.77| 0.72|
| Al  | 0.55| 0.53| 0.28| 0.37| 0.60| 0.46    | 0.54| 0.54| 1.00| 0.46| 0.68| 0.72| 0.71| 0.72| 0.77| 0.72|
| Mn  | 0.53| 0.57| 0.30| 0.40| 0.62| 0.48    | 0.55| 0.55| 0.46| 1.00| 0.71| 0.72| 0.71| 0.72| 0.77| 0.72|
| Li  | 0.68| 0.68| 0.46| 0.52| 0.79| 0.74    | 0.79| 0.79| 0.71| 0.71| 1.00| 0.81| 0.81| 0.81| 0.85| 0.81|
| Rb  | 0.77| 0.77| 0.51| 0.59| 0.77| 0.73    | 0.77| 0.77| 0.71| 0.71| 0.81| 1.00| 0.81| 0.81| 0.85| 0.81|
| Sr  | 0.79| 0.79| 0.51| 0.59| 0.79| 0.73    | 0.79| 0.79| 0.71| 0.71| 0.81| 0.81| 1.00| 0.81| 0.85| 0.81|
| Ba  | 0.75| 0.75| 0.51| 0.52| 0.75| 0.74    | 0.75| 0.75| 0.71| 0.71| 0.81| 0.81| 0.81| 1.00| 0.85| 0.81|
| As  | 0.82| 0.82| 0.62| 0.62| 0.82| 0.81    | 0.82| 0.82| 0.77| 0.77| 0.85| 0.85| 0.85| 0.85| 1.00| 0.85|
| V   | 0.68| 0.68| 0.54| 0.54| 0.68| 0.67    | 0.68| 0.68| 0.63| 0.63| 0.77| 0.77| 0.77| 0.77| 0.85| 1.00|
ratios (except Royal Arch Spring; Figs. 6 and 7). Lonetree, Horn, Salt Creek, and 140 Mile Spring water samples had low radiocarbon ages ranging from 175 to 1,000 years. Serpentine and Royal Arch Springs waters had older radiocarbon ages ranging from 2,000 to 2,400 years (Solder et al. 2020). Horn and 140 Mile Springs waters had the lowest UAR values from the study (Fig. 6) and had low radiocarbon ages of 360 and 175 years respectively (Solder et al. 2020).

The other springs in the study area (Red Canyon, JT, Miners, Grapevine, Pipe, Canyon Mine Well, Indian Gardens, Monument, Hermit, and East Boucher) had calcium-magnesium-bicarbonate type waters (green group from multivariate analysis in Fig. 10). The spring waters from this group on the east side of the study area (Red Canyon, JT, Miners and Grapevine Springs) have tritium indicating a mixture of modern and premodern waters and radiocarbon ages ranging from 345 to 1,500 years (Solder et al. 2020), high strontium isotopic ratio, high As and low U concentration (Table S1 of the ESM). Other lines of evidence, including groundwater flow modeling (Knight and Pool 2016) and isotopes of hydrogen and oxygen in water (Solder and Beisner 2020), suggest water present at the aforementioned springs has a short flow path and may have a more local source of recharge compared to the regional system. The groundwater from sites located to the west of the aforementioned springs generally have tritium indicating primarily premodern water and radiocarbon ages ranging from 1,050 to 12,000 years (except for Pipe and Monument spring as discussed at the beginning of this section; Solder et al. 2020).

Generally, the majority of water in this study area recharges the subsurface near the San Francisco Peaks and migrates down through permeable structural features to the Redwall-Muav aquifer (Fig. 2), consistent with previous conceptual models of groundwater flow (e.g., Hart et al. 2002; Monroe et al. 2005, Bills et al. 2007). Multiple independent geochemical constituents, as reported in this study and two companion papers (Solder et al. 2020; Solder and Beisner 2020), show additional sources of local recharge to the subsurface located north of the San Francisco Peaks along the Coconino Plateau. Locally, groundwater migrates from upper stratigraphic units containing gypsum to the Redwall-Muav aquifer through permeable structural features. In some parts of the study area, large-scale structural features such as faults, facilitate transport of older groundwater from the deep subsurface. The exchange between the shallow meteoric and deep subsurface groundwater flow systems along large structural features is apparent in the tracer chemistry (Solder et al. 2020; Solder and Beisner 2020), trace element chemistry (Fig. 11), and abundant dissolved carbon dioxide as reported by Crossey et al. (2006) and observed as over saturation and deposition of carbonate mineral species. Conversely, small-scale groundwater interaction with mineralized breccia pipes in the region is not well understood. The observed chemical anomalies that occur around known mineralized breccia pipes are the subject of further research.

Conclusions

The objectives of this study were to characterize the geochemical signatures of groundwater south of Grand Canyon, establish baseline concentrations for select trace elements, and
identify anomalous groundwater chemistries. Better understanding of the geochemical evolution of groundwater south of Grand Canyon is needed to understand potential effects from breccia-pipe uranium mining in the region.

Groundwater from 28 sampling locations south of Grand Canyon were analyzed for major ion, trace element, and radioisotopic constituents. Groundwater south of Grand Canyon was not statistically different from water north of Grand Canyon (from Beisner et al. 2017) for the following elements B, Li, Se, Tl, U, and V; however, As was higher in the southern groundwater and Sr was lower. Five groundwater sites (Red Canyon, Miners, JT, Havasu, and Warm Springs)
exceeded the USEPA drinking water standard for As of 10 μg/L. Salt Creek Spring exceeded the USEPA drinking water standard for U of 30 μg/L.

Four springs from the study area (Blue, Havasu, Fern, and Warm Springs) had unique chemistry, which may indicate a deep flow path or potential contribution of fluids from lower in the crust. The other springs in the study area were distinguished by major anion water type, sulfate, bicarbonate and a mixture of sulfate and bicarbonate. Water type distinctions were somewhat spatially segregated with sulfate type groundwater present on the western side of the study area where there are thicker gypsum deposits (Billingsley et al. 2008; P. Huntoon, University of Wyoming, personal communication, 2019), bicarbonate type groundwater on the eastern side of the study area, and a mixture of the two interspersed between the endmember sites. Sulfate type groundwater from this study has low strontium isotopic ratio (87Sr/86Sr) values in the range of rock values from the gypsum containing Kaibab and Toroweap formations. The water with high sulfate had higher tritium values indicating a component of younger water that may have moved from high in the stratigraphic section down to the Redwall-Muav aquifer along a focused preferential pathway (potential structurally enhanced conduit).

The location of spring groundwater sample collection within drainages of the Grand Canyon is important, and changes in chemistry may occur as groundwater discharges from bedrock and flows through alluvial material. The eastern fork of the Horn Creek drainage represents an area with groundwater discharging from bedrock high in the drainage that infiltrates into the alluvium and resurfaces lower in the drainage. The UAR is similar for the water with high uranium concentration at the head of the drainage, as it is for the groundwater emerging from the alluvium included in this study (Liebe 2003; Beisner and Tillman 2019). However, the tritium concentration is low for the water emerging from the Redwall Limestone at the head of the drainage (0.9 pCi/L) and increases to 3.4 pCi/L at the site in the alluvium described in this study, which may indicate mixing with a component of modern water in the alluvium (Beisner and Tillman 2019; USGS 2019). The UAR is generally insensitive to mixing with water that has low uranium concentration, such as precipitation recharging near the alluvium, whereas mixing with other regional groundwater with dissolved uranium and higher UAR would tend to drive the UAR value up in the mixed water.

Repeat sampling at shorter timescales would also be beneficial, especially at sites with a potential component of modern water, as changes in elemental concentration and tritium have been observed over monthly timeframes in groundwater at Pigeon Spring north of Grand Canyon (Beisner and Tillman 2018). Monument Spring has not been sampled since 2001 (Monroe et al. 2005) and resampling would lead to a better understanding of elevated nitrate concentrations and geochemical changes over time. Resampling at Slate, Sapphire, Ruby, Fossil Canyon, and Warm Spring would provide a comprehensive geochemical suite.

A conceptual understanding of the system indicates that the majority of water in this study area recharges the subsurface near the San Francisco Peaks and migrates down through permeable structural features to the Redwall-Muav aquifer. Additional sources of recharge from lower-elevation locations closer to Grand Canyon migrate from upper stratigraphic units containing gypsum to the Redwall-Muav aquifer through permeable structural features. Large structural features are present where older groundwater with no component of modern water and potential interaction with deeply sourced fluids discharges from the subsurface. Chemical anomalies occur around known mineralized breccia pipes and are the subject of further research.

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