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Research paper

Chemical and isotopic evaluation of sulfur sources and cycling in the Pecos River, New Mexico, USA

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1. Introduction

Sulfur is an essential element for life because nearly all organisms in terrestrial ecosystems require it as a macronutrient for the formation of amino acids, enzymes, vitamins, and other compounds (Solomon et al., 2005). Since sulfur is biologically active in many ecosystems, concentrations and isotopic compositions of dissolved sulfate in surface waters usually fluctuate in response to environmental changes (Van Steenwerck and Krouse, 1994). Over the last several decades, many studies have documented the influence of changes in atmospheric acid deposition on sulfur dynamics in temperate forest ecosystems of North America and Europe (e.g., Richter et al., 1983; Mayer et al., 1995b; Mitchell et al., 1998; Shanley et al., 2005). However, studies that evaluate the influence of land use change and climate variability on sulfur dynamics in the environment are scarce (e.g., Rock and Mayer, 2009). Such studies are extremely rare for the dryland ecosystems in the American Southwest, which are often impacted by high levels of total dissolved solids (TDS) from naturally-occurring ancient evaporites.

It is highly desirable to know how land use activities have influenced the sulfur dynamics in dryland ecosystems. Unlike temperate ecosystems, dryland ecosystems usually receive low inputs of mean annual precipitation and have high potential evapotranspiration. Dryland rivers are characterized by long periods of low flow, high hydrologic diversity, and extremely large flow variability (Davies et al., 1996). Agricultural practices introduce additional water and dissolved salts from surface water and groundwater sources onto cultivated farlands. Dissolved salts tend to build up on irrigated farmlands during dry climatic conditions and may be flushed out during floods or storms (Hernandez, 1978; Yuan et al., 2007). The great potential of land salinization and degradation in dryland environments
has concerned farmers and soil scientists for many decades (Scofield, 1940; El-Ashty et al., 1985).

The Pecos River is one of the saltiest river systems in the American Southwest. Over the last several decades, much of the research on the Pecos River has been driven by the need to improve the understanding of the mechanisms that control its water salinity and chemistry. Howard and White (1938) described marked increases in dissolved salt loading of the river near the Artesia and Malaga areas. Gibbs (1970) examined the relationship between the TDS and the Na/Na+Ca ratio and underscored the importance of evaporation-crystallization in controlling the water salinity and stream chemistry of the Pecos River. Yuan and Miyamoto (2005) developed a hydrochemical model to discriminate dominant processes controlling the water chemistry of the Pecos River. Yuan et al. (2007) reported that change in water chemistry of the Pecos River is driven by large-scale climatic variability such as the Pacific Decadal Oscillation (PDO). More recently, Yuan and Miyamoto (2008) reported $\delta^{18}O$ and $\delta^D$ values of surface waters in the Pecos River basin. These studies not only demonstrated that the water chemistry of the Pecos River is controlled by climatic, hydrologic, and lithologic variables but also suggested that land use activities may play an important role in regulating water chemistry of the Pecos River. Nevertheless, there is a lack of studies that evaluate the role of watershed biogeochemical processes on stream chemistry. Particularly, little is known about the influence of agricultural practices on sulfur cycling and transport in the dryland environment of the American Southwest.

Stable isotopic compositions of dissolved sulfate ($\delta^{18}O_{SO_4}$ and $\delta^{34}S_{SO_4}$ values) have been widely used to delineate sulfate sources and assess sulfur cycling in watersheds. Sulfates from different sources may be characterized by distinct isotopic signatures (e.g., Krouse and Gripenko, 1991). Dissolved sulfates in stream waters may retain their sulfur isotopic signatures of the source rocks due to minimal isotope fractionation during sulfur transformations under aerobic conditions (e.g., soil adsorption/desorption, mineral precipitation/dissolution, plant assimilation, and mineralization of organic sulfur). Substantial sulfur isotope fractionation occurs during bacterial (dis-similatory) sulfate reduction (Nakai and Jensen, 1964; Ingvorsen et al., 1981). In contrast, little sulfur isotope fractionation occurs during the oxidation of sulfide minerals and organic sulfur to form sulfate (e.g., Mayer et al., 1995a; Schiff et al., 2005; Shanley et al., 2005; Tuttle et al., 2009). On the other hand, $\delta^{18}O_{SO_4}$ values can change considerably because some of the oxygen in dissolved sulfate can be incorporated from ambient water (H$_2$O) and dissolved oxygen molecules (O$_2$) during biologically or chemically mediated sulfate oxidation. The oxygen isotope ratio of the newly formed sulfate depends on a number of factors, including the reaction pathways, the relative contributions to SO$_4^{2-}$ from H$_2$O and O$_2$, the $\delta^{18}O$ values of ambient H$_2$O and O$_2$, and the ambient environmental conditions (pH, temperature, and specific oxidizing agents) (Van Stempvoort and Krouse, 1994; Balci et al., 2007).

The purpose of our study was to determine the chemical and isotopic compositions of water and dissolved sulfate of surface and groundwater samples from the Pecos River basin and examine the relationships among those chemical and isotopic parameters to gain novel insights about causes controlling salinity in this dryland ecosystem. The main objectives of this work were 1) to discriminate major sources of dissolved sulfate in this dryland river, 2) to examine the major processes controlling sulfate cycling and transport in the watershed, and 3) to evaluate the influence of land use activities on sulfur cycling and transport in the Pecos River basin. It was anticipated that the obtained information would be valuable for dryland ecosystem management in the Pecos River and that the study would add to the currently sparse knowledge about sources and cycling of sulfur in the arid and semiarid regions of North America (e.g., Rock and Mayer, 2009; Tuttle et al., 2009).

2. Study area

The Pecos River originates in the southern part of the Sangre de Cristo Mountains, flows south across eastern New Mexico, and enters Red Bluff Reservoir in western Texas (Fig. 1). It drains an area of 50,000 km$^2$ in eastern New Mexico (USGS, 2010), which can be divided into an upper basin of 6,000 km$^2$ above Santa Rosa Lake and a lower basin of 44,000 km$^2$ above Red Bluff Reservoir. The hydrology of the Pecos River is rather heterogeneous and strongly influenced by changes in topography, climate, and the lithologic properties of the underlying rocks (Harrington, 1957; Summers, 1972; Yuan and Miyamoto, 2004; Yuan and Miyamoto, 2005; Yuan et al., 2007). In the upper basin, the Pecos River receives snowmelt from the headwater region located in the high mountains of the southern Rockies (elevation above 2000 m), with a relatively low mineral content (TDS~250 mg/L) (SWQDB, 2003). These mountains consist of conglomerates, sandstones, and limestones of Paleozoic age, and granites, gneisses, and schists of Precambrian age (Sidwell, 1941; Harrington, 1957). In the lower basin, the Pecos River receives water from multiple sources, including natural spring water, agricultural drains, runoff from monsoonal rainfall, and stream flow upstream with a relatively high mineral content (TDS>1000 mg/L) (Yuan and Miyamoto, 2005; Yuan et al., 2007).

The abrupt increase in the mineral content of the Pecos River near Santa Rosa is largely due to the dissolution of ancient evaporites in the lower basin, as indicated by many “bottomless” lakes scattered in the area (Harrington, 1957; Yuan and Miyamoto, 2005). The geologic framework of this basin has been documented extensively (e.g., Summers, 1972; Risser, 1987; Havenor, 2003). Permian evaporites (halite, anhydrite, and gypsum) occur in several geological units in the area, such as the San Andres, Bernal, Seven Rivers, Yates, Tansill, Castile, Salado, and Rustler Formations. The stratigraphic sequence varies throughout the basin. The Bernal Formation containing seams of anhydrite and limestone prevails in the northern part of the basin (Risser, 1987), with a northwestern limit near the Bernal and Villanueva area (Lucas, 1991). The Artesia group (Seven Rivers, Yates, and Tansill Formations) occurs in the middle part of the basin (mainly in the Roswell Basin). The upper Permian group (Castile, Salado, and Rustler Formations) mainly occurs in the southern part of the basin. The lower Permian unit (San Andres Formation) containing gray dense limestone and anhydrite occurs in many parts of the basin.

The Pecos River traverses several different hydroclimatological zones with distinct vegetation communities (e.g., alpine tundra, evergreen needle-leaf forest, shrubland, and grassland). Annual precipitation is on average 300 mm although it can exceed 700 mm in the mountainous headwater areas (Thomas, 1963). The upper Pecos River is affected by the Pecos mine drainage (Fig. 1) but has otherwise experienced little land use change. There are three irrigation districts in the lower basin, namely Fort Sumner Irrigation District (FSID), Pecos Valley Artesian Conservancy District (PVACD), and Carlsbad Irrigation District (CID). FSID and CID divert water from the river while PVACD pumps groundwater from the Roswell Artesian basin.

3. Materials and methods

Surface and groundwater samples were collected from eighteen locations in the Pecos River basin for chemical and isotopic analyses. The sampling was carried out during the snowmelt season in March 2010. Consideration of sampling site selection included site accessibility, spatial coverage, the degree of increases in dissolved sulfate flux along the Pecos River, and the significance of aquifers for agricultural practices in this region. Surface water samples were collected by hand-dipping along the river shore at a water depth of 10–15 cm. Clean sample bottles were filled slowly to minimize post-sampling alteration in isotopic compositions. Groundwater samples were taken...
from farm wells near the Agricultural Science Center of New Mexico State University in Artesia. Two groundwater samples were taken from the shallow alluvial aquifer and the other two samples from the deep Artesian aquifer.

Water samples were shipped to Cleveland State University for subsampling, treatment, and chemical and isotopic analyses. About 30 mL of water was subsampled and filtered from each water sample for chemical analyses at Cleveland State University, using an ion chromatography unit (ICS-1500). The relative errors for Cl\(^{-}\) and SO\(_4^{2-}\) were 1.0% and 1.5%, respectively. About 1 mL of water was subsampled and filtered for analysis of δ\(^{18}\)O and δD of water at Dr. Karr’s laboratory of Duke University, using a Thermo Finnigan TC/EA with GC-PAL autosampler attached to a Thermo Finnigan Delta Plus XL continuous flow mass spectrometer via a Conflo III interface. The isotopic values are reported using the standard δ notation relative to Vienna-Standard Mean Ocean Water (V-SMOW). The analytical precisions for δ\(^{18}\)O and δD of water were ±0.1‰ and ±1.5‰, respectively.

Water samples were filtered and acidified with 3 M HCl to remove dissolved carbonates and bicarbonates in water. BaSO\(_4\) was precipitated using 0.5 M BaCl\(_2\), collected by filtration, rinsed with deionized water, and oven-dried at 60 °C. Analyses of δ\(^{18}\)O and δ\(^{34}\)S values of sulfate were conducted at the University of Calgary via continuous flow isotope ratio mass spectrometry. δ\(^{18}\)OSO\(_4\) and δ\(^{34}\)SSO\(_4\) values are reported with respect to the international standards V-SMOW and Vienna-Canyon Diablo Troilite (V-CDT). The analytical precisions for δ\(^{18}\)OSO\(_4\) and δ\(^{34}\)SSO\(_4\) values were ±0.5‰ and ±0.3‰, respectively.

Fig. 1. Google map showing locations of fourteen sampling sites along the Pecos River, New Mexico. The small red rectangle denotes approximate location of the Pecos mine drainage in the upper Pecos valley (Berger et al., 2000), the gray area highlights the extent of the Roswell Artesian Basin (Land and Huff, 2010), and the green areas represent the distribution of irrigated farmlands in this basin (Longworth and Carron, 2003). The solid blue line refers to the drainage boundary of the Pecos River.
4. Results

The analytical results of the chemical and isotopic measurements of surface and groundwater from the Pecos River are listed in Table 1. There are wide variations in water chemistry and stable isotope geochemistry parameters of river water. The concentrations of Cl\(^-\) and SO\(_4\)\(^2-\) of river water increased rapidly from 3 mg/L and 12 mg/L near Pecos in the mountainous headwater region to 79 mg/L and 1362 mg/L near Santa Rosa and to 2298 mg/L and 1800 mg/L near Pierce Canyon Crossing. \(^{18}\)O and \(\delta D\) values of river water increased progressively from -12.7% and -89.6% near Pecos to -5.0 and 11.5% near Malaga. The deuterium excess (\(d\)), defined as \(d = \delta D - 8\times^{18}\)O (Dansgaard, 1964), decreased considerably (from 12% to -2.5%) from the upper basin downward.

Sulfur and oxygen isotope ratios of dissolved sulfates were different in the upper and lower basins of the Pecos River (Fig. 2a and b). \(^{34}\)S\(_{SO_4}\) and \(^{18}\)O\(_{SO_4}\) values were +1.0 and -6.2‰ at the initial sampling site near Pecos but increased concurrently with flow distance in the upper Pecos River. In the lower basin, however, \(^{34}\)S\(_{SO_4}\) values were constant with 11.8±0.3‰ while \(^{18}\)O\(_{SO_4}\) values decreased considerably from 14.1‰ near Santa Rosa to 9.0‰ near Pierce Canyon Crossing. \(^{18}\)O is the difference between \(^{18}\)O of sulfate and \(^{18}\)O of water (i.e., \(^{18}\)O\(_{SO_4}\) = \(^{18}\)O\(_{water}\) - \(^{18}\)O\(_{water}\)). \(^{18}\)O\(_{SO_4}\) values were relatively low in the upper basin, increasing from 13.7‰ near Pecos to 16.3‰ near Dilia, reaching their maximum of 24‰ near Santa Rosa, and subsequently decreasing progressively to 14‰ in the Malaga and Pierce Canyon Crossing area (Fig. 2c).

Groundwater from the Roswell Basin is brackish with sulfate being increased progressively from 3 mg/L and 3 mg/L near Pecos to 2298 mg/L and 1800 mg/L near Santa Rosa. Flow to the river mouth near the New Mexico/Texas state line is estimated using Google Earth.

Flow measurements from USGS gaging stations over the last decade (2001–2010), the annual average discharge of the river increased slightly from 2.5 m\(^3\)/s near Pecos to 2.8 m\(^3\)/s near Antion Chico, and then decreased to 2.4 m\(^3\)/s at the gaging station above Santa Rosa Lake even though the river receives additional water from the Gallinas River at an annual average flow rate of 0.36 m\(^3\)/s. As documented previously (Harrington, 1957; Dinwiddie and Clebsch, 1973; Risser, 1987), a substantial amount of the river water infiltrates underground before reaching Santa Rosa Lake and comes back to surface in the reach between Santa Rosa and Puerto De Luna. The annual average discharge of the river peaks at Puerto De Luna (4.3 m\(^3\)/s) due to

Table 1

<table>
<thead>
<tr>
<th>Sample ID</th>
<th>Site description</th>
<th>Latitude (°N)</th>
<th>Longitude (°W)</th>
<th>Cl(^-) (mg/L)</th>
<th>SO(_4)(^2-) (mg/L)</th>
<th>(\delta D) (‰, SMOW)</th>
<th>(\delta^{18})O (‰, SMOW)</th>
<th>(\delta^{18})O(_{SO_4}) (‰, CDT)</th>
<th>(\delta^{18})O(_{SO_4}) (‰, SMOW)</th>
<th>(d) (‰, SMOW)</th>
<th>(\Delta^{18})O (‰)</th>
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<td>PR01</td>
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<td>105.670</td>
<td>3</td>
<td>12</td>
<td>-89.6</td>
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<td>1.0</td>
<td>12.1</td>
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<td>-86.4</td>
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<td>11.7</td>
<td>14.7</td>
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<td>Dilia</td>
<td>35.171</td>
<td>105.049</td>
<td>3</td>
<td>14</td>
<td>-86.2</td>
<td>-12.0</td>
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<td>4.4</td>
<td>9.6</td>
<td>16.3</td>
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<td>Santa Rosa</td>
<td>34.939</td>
<td>104.691</td>
<td>79</td>
<td>1362</td>
<td>-71.9</td>
<td>-9.8</td>
<td>11.5</td>
<td>14.1</td>
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<td>104.626</td>
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<td>-72.8</td>
<td>-10.1</td>
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<td>13.4</td>
<td>8.0</td>
<td>23.5</td>
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<td>104.679</td>
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<td>1430</td>
<td>-75.0</td>
<td>-10.3</td>
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<td>-65.1</td>
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<td>104.398</td>
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<td>PCC</td>
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\(^a\) \(d = \delta D - 8\times^{18}\)O.
\(^b\) \(\Delta^{18}\)O = \(^{34}\)S\(_{SO_4}\) - \(^{18}\)O.
\(^c\) Pierce Canyon Crossing.
the additions of groundwater (Harrington, 1957) and decreases significantly (by 27%) in the reach between Puerto De Luna and Acme. The observed flow decrease in this reach is caused primarily by evaporative water loss in Sumner Lake and the water diversion for irrigation purposes in the FSID (Longworth and Carron, 2003). In the reach along the eastern margin of the Roswell Basin, the Pecos River gains significantly (23%) from various spring flows. Earlier studies documented a substantial amount (5 m³/s) of spring water flowing into the river near Roswell (Fiedler and Nye, 1933; Theis, 1965). Today the amount of spring water contributing to the river is considerably reduced due to extensive groundwater withdrawal for irrigation practices in the PVACD. The average discharge of the river decreased from 4.1 m³/s near Artesia to 2.3 m³/s near the Malaga Bend and Pierce Canyon Crossing area due to evaporative water loss in Brantley Lake and Lake Avalon, and the water diversion by the CID. River water in the Malaga and Pierce Canyon Crossing area consists of reservoir water from Lake Avalon, agricultural returnflows, and saline groundwater seepages (Hale et al., 1954; Havens and Wilkins, 1980; Yuan and Miyamoto, 2005; Yuan et al., 2007).

There exists a significant log-linear correlation of stream discharge and sulfate concentration at most gaging stations of the Pecos River (Yuan et al., 2007). The average concentrations of dissolved sulfate at various gaging stations on the Pecos River can be calculated using the empirical log-linear equations tabulated in Yuan et al. (2007, Table 3) and the average discharge over the last decade (Fig. 3a and b). The new measurements of dissolved sulfate reported in this study are consistent with those derived from the empirical equations (cf. Fig. 3b and d). Concentrations of dissolved sulfate in river water increase abruptly between the upper basin and the lower basin, most prominently in the reach between Santa Rosa and Puerto De Luna. But there are some discrepancies between the two curves, e.g., the significant increases in sulfate concentration in the reach adjacent to the Roswell Basin (Fig. 3d). This may be explained by a pronounced effect of spring water during non-irrigation low flow conditions when water samples were taken in March 2010.

Accordingly, the sulfate flux of the Pecos River fluctuates widely from reach to reach. Most of dissolved sulfate additions occur in the two areas, i.e., the Santa Rosa basin and the Roswell basin, as highlighted in Fig. 3c. In contrast, there are some decreases in sulfate loading in the reaches associated with the two major reservoirs (Sumner Lake and Lake Brantley), and in the Malaga Bend area. Considering the changes in sulfate fluxes (Fig. 3c) and drainage area from USGS (2010), the sulfate export rates of the river change considerably from 5.4 kg S/ha/yr in the upper basin to 74 kg S/ha/yr in the Santa Rosa basin to 3.9 kg S/ha/yr in the Roswell basin to −37 kg S/ha/yr in the Carlsbad basin. The relatively high export rate from the Santa Rosa basin is due to the dissolution of Permian evaporites (Yuan and Miyamoto, 2005) while the relatively low or even negative export rates of the Roswell and Carlsbad basins may be associated with land use activities.
5.2. Chemical and isotopic characteristics of river water

The analysis of chemical and isotopic compositions of river water has proven useful for the identification of water sources and flow pathways in the Pecos River (Yuan and Miyamoto, 2008). \( \delta^{18}O \) and \( \delta D \) values of river water in the Pecos River increase progressively from the upper basin downward (Fig. 4a and b). There are two positive isotopic excursions in the reaches between the Santa Rosa basin and the Roswell basin, indicating a pronounced isotopic enrichment of deuterium and \( ^{18}O \) induced by enhanced evaporation from open-water surfaces in lakes and wetlands and agricultural farmlands in the two basins. Moreover, \( \delta^{18}O \) and \( \delta D \) values of river water from the lower Pecos River fall slightly below the global meteoric water line (GMWL, \( \delta D = 8\delta^{18}O + 10 \)) (Craig, 1961) and form a local river water line (LRWL, \( \delta D = 6.0\delta^{18}O - 13.4 \)). The observed deviation from the GMWL usually reflects a differential isotopic enrichment of \( ^{18}O/^{16}O \) relative to \( D/H \) in the remaining water during evaporation (e.g., Cappa et al., 2003; Yuan and Miyamoto, 2008).

The d-excess (\( d \)) is defined as \( d = \delta D - 8\delta^{18}O \), which is the intercept of the GMWL (Dansgaard, 1964). As pointed out in Yuan and Miyamoto (2008), \( d \) is a better indicator of evaporation-induced heavy isotopic enrichment than either \( \delta^{18}O \) or \( \delta D \). This is not only because \( d \) is controlled by both \( \delta^{18}O \) and \( \delta D \) but also because the average \( d \) values of precipitation are more or less constant (~10‰) in many areas around the world. In the Pecos River, the \( d \) values of river water decrease from 12.1‰ near Pecos to −2.5‰ at Pierce Canyon Crossing (Fig. 4c), indicating a cumulative effect of evaporation with increasing flow distance downstream. There are, however, two positive excursions occurring in the Santa Rosa basin and the Roswell basin, indicating some groundwater discharge into the river water.

Assuming there is an insignificant amount of water inputs from local precipitation and water loss through evaporation, the relative contribution of groundwater additions over the total river flow may be estimated by a simple d-excess based binary model (Yuan and Miyamoto, 2008).

\[
f = \frac{d_d - d_u}{d_g - d_u}
\]

where

- \( d_u \) is the d-excess of river water upstream
- \( d_d \) is the d-excess of river water downstream
- \( d_g \) is the d-excess of local groundwater additions
- \( f \) is the fraction of local groundwater additions

To determine the fraction of local groundwater additions to the Pecos River near the Roswell basin, we assigned \( d_u = -2.6‰ \), the average \( d \) value of river water at the reach between Acme and Roswell (upstream), \( d_d = -0.3‰ \), the \( d \) value of the river water at Artesia (Table 1). The average \( d \) value of groundwater samples from the alluvial aquifer and the Artesian aquifer near the Agricultural Science Center of NMSU at Artesia (excluding sample PR11 from a water

Fig. 4. Changes in chemical and isotopic composition of surface water along the Pecos River. (a) \( \delta^{18}O \). (b) \( \delta D \). (c) d-excess. (d) Molar ratio of chloride to sulfate. Vertical gray bars highlight chemical and isotopic changes in the Santa Rosa-Sumner basin and the Roswell basin. Red dashed lines highlight positive excursions of \( \delta^{18}O \) or \( \delta D \) values in the reaches between the Santa Rosa basin and the Roswell basin.
storage pond affected by evaporation) is 8%, slightly lower than the average d values (9%) of groundwater samples from the Trans-Pecos and the Roswell basins (Hoy and Gross, 1982; Uliana et al., 2007). The average d value of spring waters near Roswell is 10.7‰ (Land and Huff, 2010). Thus, it is tempting to assign d = 9‰, the average d value of groundwater samples from the Roswell basin, as the d value for local groundwater additions. Using this deduced d value, we calculated the fraction of local groundwater additions over the total stream flow in the river to be about 20%. This 8% value (i.e., the calculated fraction) is slightly lower than that of the average annual flow gain (23%) in the reach between Acme and Artesia over the last decade, as estimated above. Similarly, the fraction of groundwater discharge into the reach between the Santa Rosa and Puerto De Luna is estimated around 39%, if assuming d = 10‰ as the d value for local groundwater. The average flow gain in the Santa Rosa basin over the last decade is 56% (Fig. 3a). The discrepancy suggests that the groundwater additions in this reach contain a significant amount of surface water from upstream reaches. Although uncertainties remain, the simple d-based binary modeling may serve as an independent approach for the separation, quantification and/or validation of the amount of water gains or losses in a specific river reach.

5.3. Sulfur sources and cycling in the upper basin

The isotopic composition of dissolved sulfate can serve as an indicator of sulfate sources and transformations in a watershed (e.g., Caron et al., 1986; Berner et al., 2002; Shanley et al., 2005; Rock and Mayer, 2009). On global scales, $\delta^{34}$SO$_4$ and $\delta^{18}$O$_{SO_4}$ values of atmospheric deposition vary greatly, ranging from $-3$ to $0$‰ and from 7 to 17‰ in many industrialized countries (Krouse and Mayer, 2000). The average $\delta^{34}$SO$_4$ and $\delta^{18}$O$_{SO_4}$ values of atmospheric deposition in southern California and southern Nevada are 5±1.4% and 11.2±1.9‰ (Bao and Reheis, 2003). In northern New Mexico, $\delta^{34}$SO$_4$ values of meteoric waters (rain, snow, and lake water) range from 3.0 to 4.2‰. (Lynch et al., 1988; Mast et al., 2001). $\delta^{18}$O$_{SO_4}$ values of Colorado Rockies precipitation sulfate range from 10 to 15‰ (Kester et al., 2003; Scanlon et al., 2009). Atmospheric sulfate undergoes a series of processes in the soil zone, including assimilatory sulfate reduction by plants and microbes and oxidation and mineralization of organic sulfur. The latter processes cause a decrease of over 5‰ compared to the $\delta^{18}$O$_{SO_4}$ of atmospheric sulfate, since new oxygen atoms are introduced into the newly formed sulfate while there is little change in the $\delta^{34}$SO$_4$ (Mayer et al., 1995a; Mayer et al., 1995b; Shanley et al., 2005). Thus, the average $\delta^{34}$S and $\delta^{18}$O values of atmospheric-derived sulfate that has been recycled in the soil zone in this watershed is estimated to be 3.5‰ and 7.5‰ (Fig. 5d), significantly higher than the measured values of sulfate in river water from the upper Pecos River. This indicates that atmospheric deposition is not the only source of sulfate present in river water.

In fact, oxidation of sulfide minerals in the Pecos mine drainage near Tererro, New Mexico, has contributed to an elevated concentration (> 100 mg/L) of dissolved sulfate in Willow Creek, a tributary to the upper Pecos River (Berger et al., 2000). $\delta^{34}$S values of sulfide deposits from the Questa mine in northern New Mexico vary greatly from $-13.5$ to 0.5‰ (Campbell and Lueth, 2008). $\delta^{34}$SO$_4$ and $\delta^{18}$O$_{SO_4}$ values of alteration scars from the Questa mine range from $-10.6$ to 0‰ and from $-3$ to 3‰, respectively (Campbell and Lueth, 2008). Although not derived from the Pecos mine drainage, these data place an important range of isotopic variability for sulfates in this region. It is particularly useful to estimate the average $\delta^{18}$O$_{SO_4}$ value of the Pecos mine drainage as it depends largely on local climatic and drainage conditions (Taylor and Wheeler, 1993). As a first approximation of the relative contributions from the Pecos mine drainage we assigned 0‰ as the average $\delta^{18}$O$_{SO_4}$ value and deduced $\delta^{34}$SO$_4$ = -7.8‰ for sulfate oxidation-derived sulfate in this watershed by linear extrapolation (Fig. 5d).

Once $\delta^{34}$SO$_4$ and $\delta^{18}$O$_{SO_4}$ values of the two end-members of atmospheric deposition and acid mine drainage are assigned, their relative contributions to the pool of stream sulfate in the watershed can be quantified using a simple binary mixing model. Our calculations indicate that sulfate from the Pecos mine drainage accounts for 86% and 67% of the total dissolved sulfate in river water near Pecos and San Jose, respectively. This indicates that the majority of the stream sulfate at the two sites is from oxidation of sulfide minerals in the upper Pecos valley. However, stream sulfate in river water near Dilia is influenced by the occurrence of Permian evaporites because the isotopic data point corresponding to this site lies significantly above the mixing line between sulfate oxidation and soil sulfate end-members (Fig. 5d). The gypsum-bearing Bernal Formation prevailing in the Santa Rosa basin has a northwestern limit near the Bernal and Villanueva area (Lucas, 1991).

5.4. Sulfur sources and cycling in the lower basin

The lower Pecos River flows across the Permian basin in eastern New Mexico and receives dissolved salts from the dissolution of ancient evaporites (halite, anhydrite, and gypsum) (Van Denburgh and Feth, 1965; Hiss et al., 1969; Johnson, 1981; Yuan and Miyamoto, 2005). $\delta^{34}$SO$_4$ values of river water from the lower basin are quite constant with an average value of 11.8‰, almost identical to those of shallow groundwater from the alluvial aquifer but slightly lower than those of deep groundwater from the Artesian aquifer (Fig. 5d). All the $\delta^{34}$SO$_4$ values of river and groundwater samples are within the range of $\delta^{34}$SO$_4$ values (10.5 to 13.8‰) of Permian evaporites from Saladó, Castile, and San Andres Formations in this region (Claypool et al., 1980). Moreover, the average $\delta^{34}$SO$_4$ value (11.8‰) of river water from the lower Pecos River is identical to that of groundwater from the San Andres aquifer in the middle Rio Grande basin (Plummer et al., 2004). $\delta^{34}$SO$_4$ values of the deep groundwater are slightly elevated due to bacterial (dissimilatory) sulfate reduction as indicated by the presence of the distinctive smell of hydrogen sulfide (H$_2$S).

In contrast, $\delta^{18}$O$_{SO_4}$ values of river water decrease considerably from the Santa Rosa basin to the Roswell basin (Figs. 2b and 5d). Stream sulfate from the Santa Rosa basin has the highest values of $\delta^{18}$O$_{SO_4}$ (14‰) found in river water, exceeding the maximal $\delta^{18}$O$_{SO_4}$ value (12.2‰) of Permian evaporites previously documented in this region (Claypool et al., 1980). Although two major sulfate-bearing units (Bernal Formation and San Andres Formation) are documented in the Santa Rosa basin (Risser, 1987), the higher $\delta^{18}$O$_{SO_4}$ value of river water suggests that the stream sulfate is largely derived from the Bernal Formation. This is because groundwater samples from the Artesian aquifer (i.e., San Andres Formation) have lower $\delta^{18}$O$_{SO_4}$ values (-11‰). Like atmospheric sulfate deposition, stream sulfate derived from Permian evaporites undergoes sulfur cycling and transformation processes. For example, there is a 3.5‰ decrease in $\delta^{18}$O$_{SO_4}$ of river water in the section between Santa Rosa and Acme. Assuming the average $\delta^{18}$O value of recycled sulfate is 6‰, the average $\delta^{18}$O$_{SO_4}$ value of soil extracts from the southern High Plains (Scanlon et al., 2009), we estimated that 43% of dissolved sulfate has been recycled in the Santa Rosa and Sumner basin. In the Roswell basin, however, the Pecos River receives additional dissolved sulfate from spring waters as discussed above. The positive excursion, as highlighted in Fig. 2b, indicates that the $\delta^{18}$O$_{SO_4}$ values of spring inputs are higher than those of river water near Acme. The average $\delta^{18}$O$_{SO_4}$ value of spring water was calculated to be 12.9‰ by a binary mixing model, using values of sulfate flux and $\delta^{18}$O$_{SO_4}$ of river water. The estimated $\delta^{18}$O$_{SO_4}$ value is slightly higher than that of river and groundwater samples collected and measured in this area. The chemical and isotopic compositions of spring water near Roswell are significantly different from those of the groundwater near Artesia. Besides, there is a minimal (0.7‰) decrease in $\delta^{18}$O$_{SO_4}$ of river water in the reach.
between Roswell and Brantley Lake. This may indicate that about 13% of dissolved sulfate has been recycled. Lastly, there is a 1.7% decrease in δ18O of river water in the reach between Brantley Lake and Pierce Canyon Crossing, indicating 37% of dissolved sulfate has been recycled. In summary, a total of 63% of dissolved sulfate in river water mostly from the two major aquifers at a rate of 11.7 m³/s while PVACD pumps water mostly from the northern New Mexico (Campbell and Lueth, 2008); 2) New Mexico precipitation (Mast et al., 2001); 3) Permian evaporites (Claypool et al., 1980); 4) Soil extracts from the Southern High Plains (Scanlon et al., 2009). Δ18O = δ18O SO4 – δ18O H2O.

5.5. Influence of land use on sulfur cycling and transport

It has been shown that the rate of dissolved salt export from the Pecos River fluctuates over time due to changes in land use and climatic conditions (Yuan et al., 2007). Groundwater withdrawal, reservoir operation, and agricultural returnflows affect the rate of dissolved salt accumulation on farmlands in the region. There are three major irrigation districts (i.e., FSID, PVACD, and CID) in the Pecos River basin (Fig. 1). FSID and CID divert water from the Pecos River at average rates of 1.5 and 2.9 m³/s while PVACD pumps water mostly from the two major aquifers at a rate of 11.7 m³/s (Longworth and Carron, 2003; Wilson et al., 2003; Longworth et al., 2008). Irrigation water withdrawal by FSID has little effect on sulfate flux in the river while flow diversion by CID leads to a considerable reduction in sulfate flux in the river. Because of the proximity to the river, sulfates accumulated on the FSID irrigated farmlands are readily washed out into the river. We estimated that about 37% of riverine dissolved sulfate is transferred onto farmlands during irrigation seasons. More than half of dissolved sulfate accumulated is flushed back to the river without soil processing because only 43% of dissolved sulfate has been recycled from the observed changes in δ18O SO4 of river water. On the contrary, about 82% of riverine sulfate is transferred by CID onto irrigated farmlands and only 76% of the accumulated sulfate returns to the river by agricultural drains (Fig. 3c). As a result, irrigation farmlands in CID serve as a sink of dissolved river sulfate. In the Roswell basin, groundwater withdrawal by PVACD members introduces 17.6 kg/s (or 555 10⁶ kg/yr) of dissolved sulfate onto 40,000 ha farmlands. The sulfate flux from groundwater pumping is five times that observed in the Pecos River near Acme. Only 10% of the accumulated sulfate is exported to the river by spring flows and agricultural drains (Fig. 3c). The relatively low rate of
sulfate export from irrigated farmlands may be due in part to the relatively dry conditions prevailing in the region over the last decade.

The influence of land use activities on sulfur cycling and delivery is significant but varies from one irrigation district to another in the Pecos River. To further evaluate the influence of land use on sulfur cycling, we examined variations in δ18OSO4 values in relation to changes in the molar ratio of chloride to sulfate and the δ18O value of river water. δ18OSO4 is positively correlated with δ18O in the upper Pecos River and negatively correlated with δ18O in the lower basin (Fig. 5b). While the positive correlation is probably induced by varying degrees of mixing among isotopically distinct end-members of sulfide-derived sulfate, atmospheric sulfate, and evapotranspirated sulfate in the upper Pecos River, the negative correlation may be caused by the cumulative additions of recycled sulfate (with significantly lower δ18OSO4 values) in the downstream portions of the basin. The oxygen isotope exchange between sulfate and water in natural environments is very slow, at timescales of at least 10^5 yrs (Lloyd, 1968), and usually considered insignificant due to a relatively short life span of dissolved sulfate in most near-surface environments (Van Stempvoort and Krouse, 1994). A plot of Δ18O vs. [Cl⁻/J]SO4^2⁻ shows a separation of waters with different origins (Fig. 5c). Samples of groundwater and river water from the upper basin deviate significantly from the regression line derived from river waters in the lower basin. Δ18O values of river water from the Santa Rosa area are around 24‰, significantly higher than those of groundwater from the Roswell basin. This may signify variations in δ18O of dissolved sulfate from Permian evaporates in the two basins. The negative correlation between Δ18O and the molar ratio of chloride to sulfate in river water is interpreted to indicate the influence of land use on sulfur cycling and delivery. The molar ratio of chloride to sulfate of river water increases progressively due to a cumulative effect of land use, coinciding with decreases in δ18OSO4 due to sulfur cycling and increases in δ18O due to evaporation from the upper basin downstream. There is a total of 10‰ decrease in Δ18O in the lower Pecos River. Nearly half of that decrease is due to increases in δ18O of river water that resulted from evaporative enrichment. The other half is induced by decreases in δ18OSO4 of river water, most of which may be induced by sulfur cycling. It is evident that both sulfur cycling and evaporation can be enhanced by intensive land use activities. Thus, Δ18O may serve as an indicator of the extent of land use activities as it is related to evaporation, sulfur cycling, and relative accumulation of chloride to sulfate.

Interestingly, most of the 5‰ decrease in δ18OSO4 values occurs in river reaches associated with the two small irrigation districts (CID and FSID). There is a minimal (0.7‰) decrease in δ18OSO4 of river reaches adjacent to the largest irrigation district (PVACD) in the Roswell basin. We speculate that some of the isotopic variations may be attributed to differences in agricultural practices. PVACD farmers depend exclusively on groundwater for irrigation while CID and FSID clients use surface water from the river for irrigation. Additionally, sulfate inputs from spring water near Roswell have a more positive δ18OSO4 value which may offset some of the decrease in δ18OSO4 caused by sulfur cycling.

6. Conclusions

Chemical and isotopic measurements of surface and groundwater samples were carried out to identify and quantify sulfur sources and cycling in different parts of the Pecos River. The results of this study show that surface waters from the river are characterized by a wide range of variations in water chemical and isotopic compositions, with Cl⁻ ranging from 3 to 2298 mg/L, SO4^2⁻ from 12 to 1885 mg/L, δD from −89.6 to −42.0‰, δ18O from −12.7 to −4.9‰, δ18OSO4 from −6.2 to 12.2‰, and δ18OSO4 from 1.0 to 14.2‰. Detailed analysis of the chemical and isotopic data allowed us to divide the Pecos River into an upper basin above Santa Rosa Lake and a lower basin above Red Bluff Reservoir. River water from the upper basin is characterized by low concentrations of Cl⁻ and SO4^2⁻, low values of δ18O and δD, and low values of δ18OSO4 and δ34SSO4. Dissolved sulfate in the upper basin is from at least three different sources, namely the oxidation of sulfide minerals, the soil processing of atmospheric sulfate, and the dissolution of ancient evaporites. The dissolved sulfate from the oxidation of sulfides in the upper valley above Pecos has distinctively low values of δ18OSO4 and δ34SSO4 while the dissolved sulfate from the dissolution of evaporites in the lower part of the basin has high values of δ18OSO4 and δ34SSO4. The dissolved sulfate from the soil processing of atmospheric sulfate has intermediate δ18OSO4 and δ34SSO4 values. The relative contributions of these end-members to the pool of stream sulfate in river water can be evaluated, using the δ18OSO4 and δ34SSO4 values.

In the lower Pecos River, dissolved sulfate of river water is mainly from the dissolution of ancient evaporites. Despite lithologic variations, δ34SSO4 values of river water are rather constant throughout the lower basin, with an average value of 11.8‰, typical of δ34SSO4 values of Permian evaporites found in this region (Claypool et al., 1980; Plummer et al., 2004). On the other hand, δ18OSO4 values of river water decrease considerably due to sulfur cycling in the watershed. A 5‰ decrease in δ18OSO4 of river water in the lower Pecos River may indicate that 63% of the dissolved sulfate has been recycled in the watershed. Surprisingly, most of the sulfur cycling observed occurs in the two small irrigation districts (CID and FSID). There is little contribution to the decrease in δ18OSO4 of river water from the largest irrigation district (PVACD). The findings of this work imply that the influence of land use activities on sulfur cycling and transport may be more profound than previously thought.

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