Characterizing the hydrogeochemistry of two low-temperature thermal systems in Central Mexico

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ABSTRACT

Two low-temperature geothermal systems located at the Trans-Mexican Volcanic Belt with presence of fluoride and arsenic were studied with the aim to determine hydrogeochemical indicators of the toxic elements' presence, and to propose adequate geothermometers. The hydrogeological and geochemical study was carried out in Ixtapan de la Sal and Tonatico (IxS-T) and Santa Cruz de Juventino Rosas (JR), both located at the limits of the Trans-Mexican Volcanic Belt (TMVB). In these regions, low-temperature geothermal activity is present (T = 32–47 °C), and various fault and fracture systems have been identified. Several faults are active, enabling the upward flow of deep geothermal water. The geothermal waters of IxS-T manifest in the form of springs and have high Na+ and Cl − concentrations, whereas those of JR are captured in wells and mainly present high Na+ and HCO3 − concentrations. The hydrochemistry of water samples was analyzed to determine the dominant hydrogeochemical processes in both regions. These data were also useful for understanding the natural origin of the high levels of arsenic and fluoride reported in the water, which are likely due to mineral dissolution processes. The concentrations of these elements surpassed the permissible limits according to Mexican law (Asmax = 0.025 mg/L; F − = 1.5 mg/L) and represent a toxicity risk for the local populations. The groundwater at JR supplies all needs of the local population, while the water at IxS-T is mainly used for recreational and health spa purposes. Increasing trends of As and F − in the sedimentary aquifer of IxS-T are related with the increase in TDS, Cl −, HCO3 −, and SiO2, while silicate alteration releasing Na and HCO3 − are related with As and F − presence in the volcanic aquifer of JR. Reservoir temperature was adequately estimated with K2/Mg and Na-K-Ca (Mg corrected) geothermometers at IxS-T, and with chalcedony and quartz geothermometers at JR.

1. Introduction

In Mexico, high concentrations of Asmax and F − have been reported in the groundwater of several regions (Armienta and Segovia, 2008; Betancourt-Lineares et al., 2013; Esteller et al., 2015). The presence of these elements depends on the geological environment, pattern of groundwater flow, pH, temperature, and, in some cases, anthropogenic processes (Smedley and Kinniburgh, 2002; Peters and Blum, 2003). A hydrogeological and geochemical study was performed in two regions located at the limits of the Trans-Mexican Volcanic Belt (TMVB). Ixtapan de la Sal and Tonatico (IxS-T) and Santa Cruz de Juventino Rosas (JR) (distance between these regions is 220 kms) (Fig. 1a), the first one located in a sedimentary basin and the second in a volcanic environment. In both areas presence of Asmax and F − has been reported. Both regions have experienced recent volcanic activity (C. Culiacán; age 2.10 ± 0.24 Ma; Nevado de Toluca; age 2.6 ± 0.2 Ma), have a basement of sedimentary origin that presents low-grade metamorphism, and have various intricate and complex fault and fracture systems (Ban et al., 1992; García-Palomó et al., 2002). Some of these faults are active, enabling the upward flow of deep water. Therefore, these systems can be defined as low-temperature geothermal systems (Morales-Arredondo et al., 2016a, 2016b; Rodríguez et al., 2015; Martínez-Florentino, 2015). These characteristics are common along to the TMVB, and should be important to consider when classifying the geothermal water in the entire zone. In general, the geothermal waters of IxS-T rise to the ground surface as springs, containing travertine and high concentrations of Na+ and Cl −, and have a temperature of around 35 °C. The geothermal waters of JR are captured in wells, and high concentrations of mainly Na+ and HCO3 − have been detected.

In these thermal manifestations, the concentrations of arsenic and...
Fig. 1. a) Location of the studied sites and b) geology of the Ixtapan de la Sal-Tonatico region and c) geology of the Juventino Rosas region. The location of the sampling points is indicated.

Table 1

<table>
<thead>
<tr>
<th>Geological units JR</th>
<th>Hydraulic properties</th>
<th>Fault system</th>
<th>Geological units IxS-T</th>
<th>Hydraulic properties</th>
<th>Fault system</th>
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</thead>
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<tr>
<td>Alluvial deposit</td>
<td>Unconfined aquifer</td>
<td>E-W</td>
<td>Alluvial deposit</td>
<td>5.0 L/s/m</td>
<td>E-W</td>
</tr>
<tr>
<td>Volcano shield and monogenetic volcano</td>
<td>Recharge zone in some places are unconfined aquifer</td>
<td>E-W</td>
<td>Travertine</td>
<td>5.0 L/s/m</td>
<td>E-W</td>
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<tr>
<td>Andesites from tertiary</td>
<td>Have secondary porosity</td>
<td>E-W</td>
<td>Volcanic breccia, tuffs, lahars, pyroclastic and lava flows</td>
<td>Between 1.2 and 0.05 L/s/m</td>
<td>E-W and NE-SW</td>
</tr>
<tr>
<td>Rhyolitic domes</td>
<td>In some place act as recharge zone, low permeability</td>
<td>E-W</td>
<td>Conglomerate</td>
<td>0.05 to 1.0 L/s/m</td>
<td>E-W</td>
</tr>
<tr>
<td>Basaltic plateau tertiary</td>
<td>Recharge zone</td>
<td>E-W</td>
<td>Igneous rocks</td>
<td>5.0 L/s/m</td>
<td>E-W and NE-SW</td>
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<td></td>
<td></td>
<td></td>
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<td>1.0 to 2.0 L/s/m</td>
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<td>0.05 to 1.0 L/s/m</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>&lt; 0.05 L/s/m</td>
<td></td>
</tr>
<tr>
<td>ALLENDE andesite</td>
<td>1.6 × 10^{-4} m²/s</td>
<td>NE-SW and NWSE</td>
<td>Shale-sandstone</td>
<td>&lt; 0.05 L/s/m</td>
<td>NW-SE</td>
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<tr>
<td>CHICHINDARO rhyolite</td>
<td>Unknown properties, secondary porosity</td>
<td>NE-SW and NWSE</td>
<td>Limestone-shale</td>
<td>≈ 0.05 L/s/m</td>
<td>NW-SE</td>
</tr>
<tr>
<td>Ignimbrite sequence from Oligocene-Miocene</td>
<td>Low permeability, aquitard, in some circumstances act as basement</td>
<td>NE-SW and NWSE</td>
<td>Limestone-dolomite</td>
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<td>NW-SE</td>
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<td>SOYATA Formation</td>
<td>1.2 × 10^{-4} m²/s</td>
<td>Limestone</td>
<td>Metamorphic rocks</td>
<td>≈ 0.05 L/s/m</td>
<td>NW-SE</td>
</tr>
</tbody>
</table>
fluoride surpass the permissible limits for water for human use and consumption established by Mexican regulations (NOM-127-SSA-1994; modified in 2000; As = 0.025 mg/L; F⁻ = 1.5 mg/L), thereby representing a toxicity risk and possibly causing health problems for the local populations. Groundwater at JR is used to supply all water needs of the local population, while the water at Ixs-T is mainly used for recreational and balneological purposes.

This investigation had the objective of assessing the hydrogeochemical properties of two low-temperature geothermal systems located along the TMVB, the presence, origin, behavior and hydrogeochemical indicators of As and F⁻ in each area, and to identify the most representative geothermometer for each geological setting.

2. Location and geology

The municipalities of Ixtapan de la Sal and Tonatique (Ixs-T) are located in the State of Mexico. Physiographically, this region is located between the lower limit of the Trans-Mexican Volcanic Belt and the Sierra Madre del Sur (Fig. 1b). The climate is semi-warm subhumid with an average annual temperature between 18 and 24 °C and a summer and winter rainfall pattern. This area forms part of the Morelos-Guerrero Watershed and presents a high structural complexity resulting from three juxtaposed tectonic domains, of which the Telolapa-Ixtapan Complex dating to the Mesozoic Era is the most representative (Table 1). This complex is formed of Mesozoic sediments. platform rocks and volcanic-sedimentary rocks of marginal-sea volcanic island arc type (Campa, 1978). In addition, this volcanic-sedimentary complex presents low-grade metamorphism and green shales facies (Martínez-Florentino, 2015). A deformed and folded series of Cretaceous Flysch-type carbonate and clastic rocks is overlain on the prior complex in a discordant manner, forming the high sierras of the zone. The third element overlapping the others is a volcanic complex composed of rhyolite and pyroclastic rocks and lahars (Table 1). The Nevado de Toluca volcano is located to the north, and basaltic plateaus ranging from 2700 to 2900 masl in elevation are positioned to the south. In some places from Ixs-T existing salt deposits before prehispanic age, these deposits are not exploited (Reyes Reyes, 1998).

The municipality of Juventino Rosas (JR) is located in the state of Guanajuato and belongs to the hydrogeological province of Valle de Celaya (Fig. 1c). This study area is found between the physiographic regions of the Trans-Mexican Volcanic Belt and the Central Plateau. Lithologically, the area is formed by a sedimentary basin that overlies volcanic emissions of basic composition from the Pliocene-Pleistocene discordant manner, forming the high sierras of the zone. The third element overlapping the others is a volcanic complex composed of rhyolite and pyroclastic rocks and lahars (Table 1). The Nevado de Toluca volcano is located to the north, and basaltic plateaus ranging from 2700 to 2900 masl in elevation are positioned to the south. In some places from Ixs-T existing salt deposits before prehispanic age, these deposits are not exploited (Reyes Reyes, 1998).

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3. Methodology

Groundwater was sampled in JR in 2013 and 2014 from thermal springs, non-thermal springs, and wells in Ixs-T in 2014 and 2015. The most representative results are shown in Tables 2 and 3. The water sampling and chemical analyses were performed according to international standards (APHA-AWWA-WFPC, 2005). In the wells in JR, temperature, pH, and electric conductivity were measured in situ using a Conductronic PC 18 portable meter calibrated at each site. In Ixs-T, an OrionStar A329 multiparametric meter, also calibrated at each site, was used to determine temperature, pH, and electric conductivity.

Alkalinity was also analyzed in situ. The chemical determinations of JR samples were performed in the Laboratorio de Química Analítica of the Instituto de Geofísica of the Universidad Nacional Autónoma de México (UNAM). In this analysis, the major ions were determined. Chromium was measured by flame atomic absorption spectrophotometry, arsenic by hydride generation, boron by colorimetry and its reactions with carminic acid, fluorides by potentiometry with a selective electrode, lithium by nitrous oxide-acetylene flame emission spectrometry, and silicon by nitrous oxide-acetylene flame atomic absorption spectrometry, using a Perkin Elmer AAAnalyst 100 spectrometer. UV-visible light spectroscopy, according to the molybdisilic acid method, was used for samples with lower concentrations (up to 10 mg/L of Si). The concentration of silica was determined by calculating its stoichiometric ratio to silicon.

The chemical analysis of the samples from Ixs-T was carried out in the Laboratorio de Geoquímica Ambiental, Campus Juriquilla, UNAM. In this analysis, the cations were determined by inductively coupled plasma optical emission spectrometry (ICP-OES), and the concentrations As, B, Li, Si, Mg, Ca, Na, and K. The determination of anions was performed in the Laboratorio de Calidad del Agua of the Centro Interamericano de Recursos de Agua (CIRA) of the Universidad Autónoma del Estado de México (UAEMex). Sulfate was determined using the turbidimetric method and chloride through silver nitrate titration (APHA-AWWA-WFPC, 2005); fluoride was quantified with a selective electrode potentiometer. The concentration of silica was determined by calculating its stoichiometric ratio to silicon.

4. Results and discussion

The water samples from both JR and Ixs-T had a nearly neutral pH. In Ixs-T, the temperature of thermal water was around 35 °C, while that of non-thermal springs and wells ranged from 20 to 26 °C. The most abundant ions by mass were Cl⁻ and Na⁺, with the exception of well water, in which the dominant ions were Ca²⁺ and HCO₃⁻ (Table 2). In JR, the temperature of water captured in wells ranged from 28 to 48 °C, and the most abundant ions were Na⁺, Ca²⁺, and HCO₃⁻ (Table 3).

5. Piper diagram

In Ixs-T, based on the results of the Piper diagram, the samples of well water (Ixs-T W) can be categorized as Ca-Mg-HCO₃ type. The samples of non-thermal springs (Ixs-T S) can be characterized as Ca-Na-HCO₃-CI type and those of thermal springs (Ixs-T HS) as Na-Ca-Cl type (Fig. 2). In JR, the samples from thermal wells can be characterized as Na-HCO₃ and Na-Ca-HCO₃ type (JR) and those from non-thermal wells (JR Low T) as Ca-Mg-Na-HCO₃ and Ca-Mg-HCO₃-SO₄ type (Fig. 2). These results confirm that the water sampled in Ixs-T, considering thermal spring, non-thermal spring, and well water samples, are originated from different geochemical processes in the same geological environment.

Also, the hydrogeochemical evolution of the water sampled in Ixs-T from non-thermal springs evidences mixing processes, as specific chemical species of water proceeding from thermal springs and well water could be identified (Fig. 2) (Larsen et al., 2001), as explained below. In the JR region, a mixing process also appears to occur, likely indicating that deep geothermal water flows upward through the local fault system and then mixes with shallow water.

6. Behavior of the main constituents

The behavior of the major constituents of water samples from Ixs-T and JR, according to the relations between ions, was determined in order to assess the influence of the geological environment on the hydrogeochemical variability of sampled water (Fig. 3). In Ixs-T samples, the Na⁺ vs Cl⁻ relationship presented a linear tendency similar to that of the halite dissolution line, as observed in Fig. 3a (Hanor, 1994;
Carpenter, 1978). In Fig. 3b, almost all samples are located below the gypsum dissolution line, indicating that the concentration of Ca$^{2+}$ is slightly lower than that of SO$_4^{2-}$, which may be due to the dissolution of calcite. In Fig. 3c, the concentrations of HCO$_3^-$ are higher than those of SiO$_2$ (>-10.1), which could result from the alteration of carbonates (Appelo and Postma, 2005; Capaccioni et al., 2005), specifically carbonate or dolomite rocks, as deduced from Fig. 3d. Based on the above, all water samples from the IxS-T region present chemical evidence suggesting that the water has passed through carbonate or sedimentary rocks, thereby promoting the dissolution of ionic species. As presented in the lithological profile of this region in Fig. 1a, a large part of the basement is composed of dolomitic carbonate rocks.

With respect to sampled well water in IxS-T, the concentrations of alkaline earth metals (Ca$^{2+}$ + Mg$^{2+}$) were higher in comparison to those of alkaline metals (Na$^+$ + K$^+$), as shown in Fig. 3f. In addition, the concentrations of bicarbonates in solution (CO$_3^{2-}$ + HCO$_3^-$) exceeded those of sulfates and chlorides in solution (SO$_4^{2-}$ + Cl$^-$). These are common characteristics of recently infiltrated water and correspond with those of a local flow (Tóth, 1999; Chadha, 1999). In the samples from thermal springs, the concentrations of alkaline metals (Na$^+$ + K$^+$) exceeded those of alkaline earth metals (Ca$^{2+}$ + Mg$^{2+}$), while the concentrations of sulfates and chlorides in solution (SO$_4^{2-}$ + Cl$^-$) surpassed those of bicarbonates in solution (CO$_3^{2-}$ + HCO$_3^-$). These are common characteristics of deep water and of a system with intermediate or regional flow. The samples from non-thermal springs presented characteristics in between those of well water and thermal springs (Chadha, 1999), as observed in the Piper diagram of Fig. 2.

With respect to the water samples from JR, the concentration of Na$^+$ was greater than that of Cl$^-$, as shown in Fig. 3a, suggesting that the presence of Na$^+$ could be due to plagioclase weathering (Appelo and Postma, 2005). The concentration of Ca$^{2+}$ was greater than that of SO$_4^{2-}$. In this case, Hounslow (1995) indicates that the source of Ca$^{2+}$ could be the dissolution of calcite and/or silicates (Fig. 3b), but other processes as ionic exchange, pH or redox conditions may regulate the water chemistry (Russak and Sivan, 2010), in JR and IxS-T pH is neutral to slightly alkaline and the environment dominantly is reducing (Morales-Arredondo et al., 2016b; Martínez-Florentino, 2015). The concentrations of HCO$_3^-$ were superior to those of SiO$_2$ and found in a relationship of >3:1 to <10:1; this phenomenon is likely due to the alteration of the silicates present in the environment (Fig. 3c) (Hounslow, 1995; D'Amore et al., 1983; Drever and Hurcomb, 1986). The hydrogeochemical evolution of water in JR could be explained by the weathering of volcanic rocks of mostly acidic composition (Fig. 3d) (Ravenscroft and McArthur, 2004), although the water could have also had contact with carbonate rocks (Figs. 2 and 3b). Furthermore, in the environment, the mixing of deep geothermal water and shallow water is evident (Figs. 2 and 3d). Another process that could be occurring is ion exchange (Fig. 3e), as the presence of Ca$^{2+}$ and Mg$^{2+}$ decreases while the concentrations of Na$^+$ and K$^+$ increase in the direction of relative groundwater flow. This phenomenon has been observed during the weathering of silicates and causes the subsequent formation of clays that favor ion exchange processes (Ravenscroft and McArthur, 2004; Jalali, 2005), as previously reported in JR by Morales-Arredondo et al.
Based on the assumption that the concentrations of Mg²⁺ and SO₄²⁻ present a significant decrease in high-temperature geothermal environments (Stueber and Walter, 1991), which was not observed in the water samples of IxS-T or JR, a geothermometry study was proposed to determine the temperatures of these low-temperature reservoirs. Silica (SiO₂) geothermometers were used to calculate the minimum temperature of the water reservoirs because these are ideal for low temperatures systems with equilibrium between silica concentrations in fluid and rock (Karingithi, 2009; Fournier, 1977; Arnorsson, 1975). The concentration of silica in the water samples of JR, for instance, is governed by the chemical equilibrium with minerals such as chalcedony and quartz (Fig. 4), enabling the application of silica geothermometers. However, the water samples of IxS-T do not present equilibrium with silica minerals. This could be attributed to the distinct processes that occur in this aquifer, wherein whichever process is causing the high level of salinity is perhaps the most important. Giggenbach et al. (1983) and Giggenbach (1986) suggested that in high salinity environments, the use of SiO₂ geothermometers is not ideal. Therefore, for the IxS-T region, the temperature prediction for its brackish water was performed with a K₂/Mg geothermometer (Giggenbach, 1986), which shows rapid reequilibration in the water of superficial aquifers (Giggenbach et al., 1983). In comparison, the silica geothermometer shows that reequilibrium is established at a lesser velocity. In addition, a Na-K-Ca geothermometer with Mg correction was used to estimate the temperature and equilibrium conditions of deep aquifers (Fournier and Potter, 1979). The results suggest that IxS-T is a low-temperature geothermal system with a minimum reservoir temperature of 80 °C and a maximum of 111 °C but it is necessary to consider that the calculated temperatures for samples from IxS-T (NT1, NT2, NT3), with those geothermometers may not be representative due to mixing processes. On the other hand, JR is a low temperature system with a minimum reservoir temperature ranging from 40 to 99 °C (Morales-Arredondo et al., 2016a, 2016b) (Fig. 4, Table 4).

Different geothermometers were applied in JR and IxS-T to calculate reservoir temperatures (Fig. 4, Table 4), however possible mixing processes may affect the estimated temperature, for this reason the multicomponent fluid-mineral equilibria at a particular temperature was determined, to compare and evaluate the results obtained with the geothermometers. Reed and Spycher (1984) consider that if a set of minerals approaches equilibrium at a specific temperature, this has a close similarity to the reservoir temperature or to the source aquifer temperature of the particular water evaluated.

To estimate the reservoir temperature a geothermometric modeling was carried out. This technique simulates the theoretical temperature increase of the sampled waters to determine the one corresponding to the saturation of the minerals present in the reservoir, or to determine an equilibrium convergence point. In this case, as a condition to make the calculations we assumed that the water chemical composition did not change with increasing temperature. All simulations were made with PHREEQC©. Quartz and chalcedony were selected to model JR since both minerals have been reported in this zone and also reported to be in equilibrium with the fluid (Morales et al., 2015). Quartz, anhydrite and celestine being minerals reported in the zone (Martínez-Florentino, 2015) were selected to model IxS-T.

Results obtained for JR indicated that water is close to equilibrium with quartz and chalcedony for all wells (Fig. 5a, b and c), and boiling leads to sharp supersaturation with respect to both minerals in a temperature range between 65 and 105 °C for chalcedony and between 85 and 115 °C for quartz. These results are similar to some of the calculated values with geothermometers (Fig. 4, Table 4). The chalcedony geothermometer and the chalcedony equilibrium modeling, both
predict a lower temperature with respect to silica contents in groundwater. According to the equilibrium calculations obtained in IXS-T, anhydrite and celestine generally show over-saturation trends with increasing temperature, however, due to their different precipitation kinetics, both display a distinct temperature range to the saturation level, starting from a temperature of 93 °C for anhydrite and 145 °C for celestine; quartz shows under-saturation trends at lower temperature 51 °C (Fig. 5d and e). Fig. 5e shows two temperature ranges in which the saturation states of selected minerals converge towards equilibrium, the first at 78 °C (quartz-anhydrite) and the second at 110 °C (quartz-celestine); the second is the most representative and matches with the temperature values estimated with the applied geothermometers included in Table 4.

It should be made clear that mixing, degassing/boiling and other common processes such as precipitation may affect the calculation of equilibrium temperature in reservoir of both sites and it is necessary to estimate mixing processes to calculate the reservoir temperature.

9. Cl-B-Li ternary diagram

To understand processes occurring in geothermal systems, Li is one trace element that is often included because it is not significantly affected by the secondary processes that occur during rock dissolution. Determining the proportion of Li also aids to evaluate the origin of Cl and B, which are conservative elements in geothermal water. B can be an indicator of the degree of maturity of geothermal systems: its concentration decreases with respect to that of Cl and Li in old geothermal systems versus recent geothermal systems (Spivack et al., 1987; Mnjokava, 2007). Meanwhile, in deep geothermal systems, the concentration of Cl increases with respect to SO4^{2-}. This process is
mainly influenced by the superficial oxidation of sulfide and the low solubility of sulfate minerals at high temperatures (Giggenbach, 1986; Nicholson, 1993).

The Cl-Li-B ternary diagram (Fig. 6a) is mainly utilized in high-temperature geothermal systems, as both B and Cl$^-$ are volatile and mobilized by vapor at high temperatures. In addition, their presence can be linked with the formation of a deep brine layer or with an increase in temperature (Spivack et al., 1987; Arnórsson and Andrésdóttir, 1995). At low temperatures and in fluids rich in NaCl, B remains in volatile form, and the Cl/B ratio is elevated (Magaña, 1999). In fluids derived from subducted sediments, high concentrations of B have also been found in comparison to superficial water (Peacock and Hervig, 1999). In addition, elevated contents of B have been quantified in shale and marine sediments due to the adsorption of B by clay minerals (Goldschmidt and Peters, 1932; Williams et al., 2001). Furthermore, B is highly mobile in the aqueous phase in environments of sedimentary origin (Levinson, 1980). Several of these geological characteristics were present at IxS-T and can explain the high B content in spring water. Fig. 6b shows that the samples from IxS-THS have high Cl and B concentration, while samples from IxS-TW have low contents of Cl and B, it is possible that concentrations of samples from IxS-TS reflect a mixture process.

Relative concentrations of Cl, Li, and B indicate that the geothermal system of IxS-T could have a high degree of maturity because Cl$^-$ concentration is proportionally greater than those of B and Li. This could be due to the salinity of the water and the possible upward migration of deep water, (Fig. 6a). It would be necessary to perform an isotopic study of $^{14}$C, $^{18}$O and Deuterium to confirm or reject this theory. The concentrations of Cl$^-$ and B suggest that their presence is due to rock dissolution. There is no evidence of a magmatic contribution to the environment according to the comparative concentrations of Cl$^-$ and B (Fig. 6b). Water in IxS-T therefore originates from a deep environment, and its geothermal characteristics likely result from heating, wherein water is heated through its interaction with active and deep faults, and/or increasing temperatures along the geothermal gradient.

In the case of JR, the concentrations of Cl$^-$, Li, and B indicate that the water comes from a recent low-temperature geothermal system (Fig. 6a). The comparison of the Cl$^-$ and B ratio in the water samples does not indicate a magmatic contribution of heat (Fig. 6b). Therefore, the heat source in JR may be remnant heat, or result from radioactive decay, or the interaction of the water flow with deep, active faults. Considering the characteristics of the aquifer (basin filled with sedimentary material and presence of clay), boron may be adsorbed by clay during lateral flow, and this phenomenon can affect the Cl and B concentrations if materials of sedimentary origin are present in the groundwater flow environment (Kharaka and Hanor, 2007), this effect complicates the evaluation of mixing processes using concentrations of some elements.

To confirm some of the hypotheses raised here (mainly mixing

Table 4

<table>
<thead>
<tr>
<th>Sample</th>
<th>Chalcedony$^a$</th>
<th>Chalcedony$^b$</th>
<th>Sample</th>
<th>IxS-T</th>
<th>$K^A/Mg$</th>
<th>Na-K-Ca, Mg correction$^d$</th>
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$^a$ Fournier (1977).
$^b$ Arnórsson (1975).
$^c$ Giggenbach (1986).

Fig. 4. Identification of the most suitable geothermometer for calculating the water reservoir temperatures of JR and IxS-T (Giggenbach and Glover, 1992).
Fig. 5. Graphical representation of the saturation index of the different minerals that are in equilibrium with the reservoir from JR and IxS-T. 

(a) Sampled wells in JR-LowT that were selected to represent the equilibrium conditions (05 JR, 10JR, 11JR) and their calculated saturation index (using PHREEQC; Qz = quartz, Trid = tridymite, Chalc = chalcedony, Crist = cristobalite). 

(b) and (c) Set of mineral selected for the geothermometrical calculations considering local geology and the chemical equilibrium with minerals such as chalcedony and quartz in JR-LowT. 

d) Sampled hot springs in IxS-THS that were selected to represent the equilibrium conditions (M1, M2, M3, M4, M6) and their calculated saturation index (Using PHREEQC. Anhy = anhydrite; Qz = quartz; Talc = talc; Celes = celestine). 

e) Set of mineral selected for the geothermometrical calculations considering local geology and the chemical equilibrium with minerals such as anhydrite, quartz and Celestine in IxS-THS.

Fig. 6. 

(a) Cl-B-Li ternary plot of the water samples from IxS-T (square and triangle) and JR (diamonds), 

(b) Cl vs B diagram of the water samples from IxS-T (squares, well water; hollow circles, thermal water; filled circles, non-thermal water) and from JR (diamonds); (B/Clsw = seawater ratio between B and Cl).
processes and water from a deep environment) it is necessary to develop an study based on groundwater flow modeling in combination with major ion chemistry considering the isotopic evaluation of $^2$H, $^{18}$O, $^3$He, $^4$He, $^{13}$C, $^{14}$C or $^{86}$Sr and $^{87}$Sr.

10. Arsenic and fluorides

In IxS-T, presence of $A_{\text{tot}}$ and $F^-$ may be related with the chemical weathering of sedimentary rocks. As shown in Fig. 7a and b, the HCO$_3$/SiO$_2$ relation which can discriminate between the chemical weathering of carbonates ($>10$) and silicates ($<5$), presents values higher than 10 (Hounslow, 1995).

In Fig. 7c and d, the relationships of $A_{\text{tot}}$ and $F^-$ to Cl/$\text{sum anions}$ can differentiate between brine-water of marine evaporite rocks ($>0.8$ and TDS $>500$ ppm), rainwater ($>0.8$ and TDS $<100$ ppm), and the chemical weathering of rocks ($<0.8$). In these figures, the samples from thermal and non-thermal springs presented values higher than 0.1 (except in the case of wells). These values in addition to those of the TDS, which were $>500$ ppm, indicate the presence of brine-water in a marine evaporite rock environment. With respect to Fig. 7e and f, the TDS concentrations allows to differentiate fresh water ($<1000$ ppm), drinking water ($<500$ ppm), brackish water ($>1000$ ppm), salt water ($35,000$ ppm), and brine ($>35,000$ ppm). According to these categories, the samples of thermal water (IxS-T THS) may be classified as brackish water and the rest of the samples as fresh water (Levy et al., 1999).

The salinity levels (Fig. 7c and d), TDS content (Fig. 7e and f), and increase in temperature (Table 2) represent a common set of characteristics of geothermal water according to Mukherjee et al. (2009).

In the groundwater of JR, presence of $A_{\text{tot}}$ and $F^-$ may be related to the chemical weathering of sodium silicate, probably of albite, biotite, or potassium feldspar, as well as the alteration of volcanic glass, this information has been previously reported (Morales et al., 2015; Morales-Arredondo et al., 2016a; Morales-Arredondo et al., 2016b). The Na/Ca relationship presented values $>1$, indicating that the chemical weathering of sodium silicates (similar to the plagioclase from which they originate) is occurring. In contrast, if the weathering of calcium silicates were occurring, the Na/Ca relationship would present very low values. Also, ion exchange processes could result in high $A_{\text{tot}}$ and $F^-$ contents, as deduced from Fig. 8b and c. In Fig. 8b, $\text{Cl}^-$ was subtracted from Na$^+$, and any remaining sodium would likely be due to the chemical weathering of albite or ion exchange. Also, $\text{Cl}^-$ was subtracted from K$^+$, wherein K$^+$ likely originates from the chemical weathering of biotite or potassium feldspar. In Fig. 8c, the (Na + K-Cl)/SiO$_2$ $>1$ relationship suggests that cation exchange is the main source of Na$^+$ (if the Ca$^{2+}$ content is low). Also, the $F^-$ content increases with temperature as has been observed at other sites like San Luis Potosí and Los Altos de Jalisco (Carrillo and Armienta, 1989; Hurtado-Jimenez and...
This process may be related to silicates alteration and volcanic glass dissolution. However, this increase was not observed above 30 °C, more research is needed on specific geochemical processes and kinetics to explain this behavior (Fig. 8d). In addition, As$_{tot}$ vs F$^-$ show an increasing concentration trend (Fig. 8e) probably indicating the same source.

### 11. Water potability

In the majority of sites evaluated in IxS-T, the concentrations of As exceeded up to 10 times the permissible limit established in NOM-127 (As = 0.025 mg/L), whereas the F$^-$ content only presented values above the established limit in thermal springs (F$^-$ = 1.5 mg/L). In JR, several wells of potable water presented As$_{tot}$ and F$^-$ values up to double the permissible limits in NOM-127, thereby presenting a risk to the local population, especially if water is consumed or used for daily activities such as cooking or watering edible crops. In both regions, the presence of As$_{tot}$ and F$^-$ is associated with geothermal water (Morales et al., 2015; Rodríguez et al., 2016; Martínez-Florentino, 2015) because as water temperature rises; the As$_{tot}$ and F$^-$ concentrations also increase (Tables 2 and 3). In the evaluated sites (rural communities) the problem has not been remediated. Water treatment by osmosis (and other treatments) is mainly used in cities. Most rural communities with As$_{tot}$ and F$^-$ presence lack of water treatment systems. Only in the state of Chihuahua (northern Mexico) use of osmosis systems in rural areas is widespread.

To evaluate whether the populations of both sites are experiencing negative effects resulting from exposure to As and F$^-$ and to determine the latent risk, an epidemiological study is required (ASTDR, 2008).

### 12. Conclusions

The IxS-T and the JR regions display characteristics of low-temperature thermal systems according to results obtained with geothermometers and the saturation indexes calculated at different temperatures. In both regions, an interaction is occurring between metamorphic sedimentary rocks and volcanic material. However, water in IxS-T has mainly interacted with sedimentary rocks (carbonate), while the aquifer matrix in JR corresponds to volcanic rocks. Based on the chemistry of sampled water, the dominant hydrogeochemical processes of IxS-T are associated with the dissolution of sedimentary rocks (e.g., limestones and dolomites) or minerals related with evaporite deposits, leading to the liberation of high concentrations of mainly Na$^+$, Ca$^{2+}$, Cl$^-$, As, and F$^-$. The salinity of these samples could be due to the hydrogeochemical evolution of water occurring at great depths, which is affected by the geothermal environment. The dominant hydrogeochemical processes in JR were associated with the dissolution of silicates present in volcanic rocks and with cation exchange processes that are accelerated by rising temperature and thereby lead to an increase in the Na$^+$, HCO$_3^-$, As, and F$^-$ contents.

To estimate the temperature of geothermal water in high salinity sedimentary aquifers the K$^2$/Mg and Na-K-Ca with Mg correction geothermometers are adequate. On the other hand, to estimate the temperature of geothermal water from altered silicates felsic volcanic
reservoirs, chalcedony and quartz geothermometers are adequate.

The presence of ASAs and F− in the groundwater of JR and IxS-T can be associated with particular hydrogeochemical characteristics and could result from the dissolution of encased rock and the migration of hydrothermal flows through fault and fracture systems, which contain high quantities of these elements, geochemical comparison between Cl−, CO3^2−, HCO3^− and SiO2− concentrations; while for JR, elements like Na and HCO3− related with the alteration of sodium silicates should be considered.

In the IxS-T water samples, the concentrations of As surpassed up to 10 times the permissible limits of NOM-127. In JR, wells of potable water with hydrothermal characteristics presented up to double the permissible limit of ASAs. These high concentrations constitute a risk for local populations.

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