Groundwater physico-chemical properties and water quality changes in shallow aquifers in arid saline wetlands, Ouargla, Algeria

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Abstract
Shallow aquifers are vulnerable to natural geogenic processes as well as anthropogenic influences, and this is especially apparent in desert regions. Within arid and hyperarid climates, evaporation is a controlling hydrologic process leads to an important increase in the concentration of dissolved minerals of both surface water and groundwater. In groundwater, this increase is not only dependent on shallow water table depth, but also on the hydraulic properties of sediments present within the unsaturated zone of the aquifer itself. The main objective of this research is to investigate possible mechanisms that might influence water quality changes under seasonal conditions in shallow aquifers situated within the Saharan desert region of Algeria. In this work, we focus on observed changes in hydrogeochemical characteristics, and the possible responsible processes. Under arid conditions, high water mineralization results in hypersaline water or brine solution formation within shallow aquifers. Due to active physico-chemical mechanisms such as Na+/Ca2+ ion exchange, the successive precipitation of calcite, gypsum, mirabilite or blödite and halite is induced. Biological processes were also observed as prevalent; evidenced by large measured variations in CO2 load concentrations. These processes contributed to an inverse relationship between CO2 and O2 concentrations within the shallow aquifers studied.

Keywords Biogeochemistry · Dissolution/precipitation · Saturation · Salt paragenesis · Sahara

Introduction

In hydrogeological studies, natural groundwater chemical characteristics vary both temporally and spatially; with the sources of these variances being quite numerous. Among many cited, the proportionality between increased water mineralization and prolonged water–rock interactions, whether upstream to downstream, is a common source of hydrochemical composition change (Langbein and Dawdy 1964; Meybeck 1976; Peters 1984; Kilham 1990). Aerated water conditions and watershed lithological heterogeneity also produce spatial variations in aquifer hydrochemical characteristics, as well (Peters 1984; O’Connor 1967; Phillips 1971; Cook et al. 1989).

The study of shallow aquifer vulnerability in arid regions with respect to high evaporation rates, and increased contamination from agricultural, domestic or industrial sources has been the subject of several recently published research (Amharref et al. 2007; Omosuyi and Osehale 2012; Mohamed et al. 2015). It is currently accepted that the combination of the three aforementioned effects (i.e., water–rock interactions, aerated water conditions and lithological heterogeneity within the watershed), coupled with low precipitation, lead to increased water mineralization and eventual salt paragenesis within the endorheic region beneath the stream bed in the unsaturated zone of a shallow groundwater system. Resulting groundwater vulnerability is dependent on the thickness of the capillary zone above the shallow aquifer water table, and zone proximity to the soil–atmosphere interface. Groundwater vulnerability, however, is not exclusively limited to shallow water table depths. In certain regions of the world, deep groundwater may also become vulnerable under high surface temperatures, as fluctuations may cause changes in hydraulic conductivity, viscosity and groundwater flow velocity within that deep reservoir system (Anderson 2005).
The aim of this work is to study both mechanisms responsible for water quality change, and processes affecting observed hydrogeochemical characteristic variations in shallow Algerian Saharan aquifers. A literature search was initially performed to identify recent studies along three lines of interest: (1) the impact of carbonate mineral equilibria, (2) mineral dissolution or precipitation with resulting ion exchange, and finally (3) the effect of CO₂ dissolution on groundwater chemistry within saline aquifer systems. First, two sequential alluvial aquifer geochemical studies were reviewed. Research studies conducted highlighted the relationship and behavior exhibited at different saturation indexes (SI) of carbonate minerals in coastal aquifers located in Pondicherry, India (Chidambaran et al. 2011, 2012). These SIs showed that increasing values of log pCO₂ during water–rock interaction directly influenced mineral dissolution processes. Secondly, a descriptive statistics and geochemical modeling study was reviewed to better understand hydrogeochemical processes involving silicate weathering or cation exchange, and the origins of groundwater chemical components in the coastal part of a similarly arid region in Calabar, Nigeria (Edet et al. 2011). Finally, a numerical model of a CO₂ leakage scenario occurring during geological storage and sequestration within the Jurassic Dogger Limestone formation in the Paris Basin, Paris, France, was reviewed to determine the various impacts CO₂ might have on groundwater quality and hydrochemical characteristics. This aforementioned geologic situation was of interest to our purposes, because of calculations regarding both the CO₂ rate of migration, and the amount of released CO₂ gas from pressurized solutions. The analysis of said sequestered CO₂ gas, stored in close proximity to overlying groundwater, showed the subsequent dissolution of CO₂ in the adjacent aquifer; resulting in an overall drop in pH within that underground system (Humez et al. 2011).

**Study area**

The study area is located in the city of Ouargla, in the northeastern part of the Algerian lower-Sahara (Fig. 1). This area is 800 km away from the capital city of Algiers and bounded by the following wilayas or provinces: El Oued, Biskra and Djelfa to the North, Ghardaia in the West, Illizi and Tamanrasset to the South, and the Tunisian border in the East. The Ouargla region is characterized by a hyperarid desert climate. Data from the National Meteorological Office “NMO” (Ouargla station) in the period between 1996 and 2010 have shown an average annual temperature of 23.4 °C, with a minimum mean of 4.8 °C in January, and a maximum

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**Fig. 1** Location map of the study area
mean of 43 °C in July. Rainfall is very sparse and irregular with an average value of around 39.1 mm/year. These intermittent rains are marked by low or varying quantities, such as 117.8 mm in 2004 and 5.9 mm in 2001. Evaporation is very high that is materialized by an average of 3085 mm/year. These intermittent rains are marked by low or varying quantities, such as 117.8 mm in 2004 and 5.9 mm in 2001. Evaporation is very high that is materialized by an average of 3085 mm/year. It represents an evaporation rate of about 9 mm/day, when it gets its maximum during July 16 mm/day (Idder et al. 2011).

The Ouargla basin is a large geologic depression with an area of approximately 750 km². It is located in the bed of the low fossil valley of the M’ya wadi which extends from the Tademaït plateau in the In-Salah region, Tamanrasset in the south; to Sebka Safioune located 40 km north of the city of Ouargla. The Oued Mya converges with Oued Igharghar from a source south of the Ahaggar mountains in the Oued Rich valley; finally draining into Chott Melrhir (Ballais 2010).

In the Ouargla region, the width of the Oued M’ya varies between 4 and 30 km. The average valley gradient is very low at 1‰, gently sloping northward from Ouargla to Sebka Safioune.

The western boundary is marked by Cenozoic sandstone plateau cliffs covered with gravel and pebbles, and associated reddish sands overlooking the valley at about 220 m in elevation (Aumassip et al. 1972). The eastern boundary is much lower in elevation 160 m, and intensively eroded. It is made of sandstone overlain by a huge Aeolian dune. Where, the basin is mainly dominated by alluvial deposits.

Wetlands appear as flooded depressions. Chotts and Sebkhas, as they are called, are endorheic structures composed of gypsum saline soils. These wetland areas encompass about 25,000 ha, whereas wetlands occupied by palm groves does not exceed 5,000 ha (Idder et al. 2011).

During the Neolithic age, these Chotts and Sebkhas were fed naturally by groundwater contributions (Aumassip et al. 1972). Currently, they receive waters of anthropogenic origin via infiltration of water surpluses from palm plantations and wastewater, causing a marked water table rise in the shallow aquifer. During hot and dry periods, areas flooded in Sebkhas during winter months dry up as a result of evaporation; forming gypsum and other salt-bearing crusts.

### Materials and methods

#### Experimental protocol: sampling locations and water analysis

Groundwater samples were collected during two contrasting temperature periods in both June (Summer) and February (Winter). Surface water piezometers were utilized and situated in various locations differentiated by natural geogenic and anthropogenic conditions. Electrical conductivity (EC), temperature (°C), pH and dissolved oxygen (DO) were measured in situ immediately before filtration. Initially, 18 groundwater samples, taken from the surface, were collected in February during a seasonally high groundwater dilution period. Five months later, 27 additional samples were collected in June during a seasonally high groundwater evaporation period. All groundwater samples were stored in the dark for less than 24 h at low temperatures (< 4 °C) in polyethylene bottles. Ca²⁺, Mg²⁺, Cl⁻ ions and alkalinity (HCO₃⁻) were determined by volumetric method (Rodier et al. 1996) at Sonatrach Haoudh Berkaoui Physico-chemical Analysis Laboratory located in Ouargla, Algeria. Na⁺, K⁺ ions were analyzed by IC9200 flame photometer, and SO₄²⁻ ion was analyzed by HACH-DR/2000 spectrophotometer. All measured groundwater physico-chemical data may be found on Table 1 for Summer and Winter.

| Samples | Winter measured groundwater data (February) | Summer measured groundwater data (June) |
|---------|--------------------------------------------|---------------------------------------|
| Depth (m) | Min 0.3 Max 10.2 Average 2.164 SD 1.567 | Min 0.3 Max 10.6 Average 1.91 SD 1.24 |
| T°C | 19.2 23.2 Average 21.7 SD 1.01 | 25 37 Average 28.56 SD 2.16 |
| pH | 6 7.8 Average 7.1 SD 0.4 | 6.38 8.14 Average 7.34 SD 0.28 |
| DO (mg/l) | 0.9 7.7 Average 2.7 SD 1.2 | 0.5 4.5 Average 2.24 SD 0.93 |
| EC (mS/cm) | 3 174.5 Average 40.7 SD 38.1 | 3.76 233.61 Average 47.32 SD 43.87 |
| SO₄²⁻(mg/l) | 1150 31,700 Average 8208 SD 5718 | 1180 83,600 Average 47.32 SD 8916 |
| Cl⁻(mg/l) | 426 186,659 Average 32,775 SD 37,378 | 547 198,148 Average 30,697 SD 35,601 |
| HCO₃⁻(mg/l) | 49 1090 Average 304 SD 211 | 137 1494 Average 395 SD 262 |
| Na⁺(mg/l) | 420 110,330 Average 21,384 SD 23,773 | 510 118,000 Average 20,733 SD 24,198 |
| K⁺ (mg/l) | 20 10,560 Average 1320 SD 1498 | 22.2 11,860 Average 1290 SD 1394 |
| Mg²⁺(mg/l) | 73 21,388 Average 2583 SD 3006 | 85 21,684 Average 2639 SD 2678 |
| Ca²⁺(mg/l) | 200 1122 Average 595 SD 218 | 210 1420 Average 737 SD 257 |
The majority of the piezometers are shown in (Fig. 2), where static level depths of the groundwater in shallow aquifer range from 0.2 up to 10 m. Most of these water gauge points were localized at urban agglomerations, chotts, sebkhas and palm grove elevations.

Experimental protocol: activity calculations and concentration factors

Calculation of activities and equilibrium diagram

Solution ion activities were calculated from total...
concentrations using the "Aqua" Ion Pair software program, and an association model from “GYPSOL” (Valles 1987; Valles et al. 1992, 1997) based on the Debye–Hückel law and extended by the Scatchard Deviation function (Scatchard 1936).

The equilibration level for the partial pressure of CO$_2$ was estimated from pH measurements and carbonate alkalinity (Bourrié 1976; Keller et al. 1987).

Estimate of the concentration factor (CF)

From the study of solution chemistry, followed by the concentration process, it is understood that any variance in exhibited chemical behavior is dependent upon the major elements or ions present (Valles et al. 1997; Al-Droubi 1976; Fritz 1981; Gac 1980). In the studied high concentration range, chloride ion is ubiquitous and conservative (Feth 1981). Therefore, it can be considered an inert solute tracer against the porous unsaturated zone sediment matrix above the shallow aquifer. Chloride ion may also serve as a criterion for concentration factor (CF) estimations and is equilibrated to the ratio between the chloride content of a shallow groundwater sample as compared to that of dilute groundwater.

Results and discussion

Water balance with calcite

Chemical activity calculations for aqueous species permitted the generation of an equilibrium diagram between groundwater solutions and the mineral calcite [CaCO$_3$(s)] (Fig. 3). Groundwater samples collected and analyzed are well within thermodynamic equilibrium range with calcite. This is consistent with the general presence of that particular carbonate mineral throughout the study area.

The partial pressure of CO$_2$ in the samples becomes variable at $10^{-3.5}$ or 0.000316 bars. Comparatively, atmospheric CO$_2$ can reach up to $10^{-1}$ or 0.1 bars: reflecting a high level of CO$_2$(g). Inverse relationship is observed between pCO$_2$ in equilibrated water and dissolved oxygen (Fig. 4). This behavior may be explained by dissolved oxygen consumption and CO$_2$ production mechanisms from biological activity within the shallow groundwater system.

Another important relationship considered was that between pressure and temperature. Generally, thermodynamic equilibrium with dissolved gases in groundwater relies on both pressure and temperature regimes in subjected aquifer system. Considering pressure alone, CO$_2$ solubility in water is a function of its partial pressure in an aquifer; with groundwater fluid pressure being proportional to aquifer depth. With a stated increase in pCO$_2$, groundwater pH would tend to decrease; and in an unbuffered aqueous system, carbonate mineral dissolution would eventually result. Conversely, pCO$_2$ (as well as dissolved oxygen) levels in water are inversely proportional to temperature. As a shallow aquifer would be more directly influenced by surface temperatures in an arid or hyperarid climate, higher temperatures affecting the capillary zone may cause lower pCO$_2$ (and dissolved oxygen) levels in those groundwaters. Therefore, the determination of the pCO$_2$ in consideration of these two important parameters is rather complex at best.

![Fig. 3 Water-calcite equilibrium diagram](image-url)
Equilibrium with sulfated and chlorinated minerals

Overall, analyzed groundwater samples mostly remained at equilibrium with gypsum [CaSO₄ • 2H₂O(s)] (Fig. 5a). However, some samples reached equilibrium with mirabilite [Na₂SO₄ • 10H₂O(s)], as well. This trend was found to be prevalent during the seasonally high groundwater dilution period in February, due to reduced solubility of this particular mineral at low temperatures (Fig. 5b). More saturated solutions reaching equilibrium with halite [NaCl(s)] were only found in samples P160 and P165 collected at sebkha Oum Raneb. Halite saturation was due to very shallow water table depths measured at 0.30 and 0.88 m in June (Summer), and 0.40 and 1.00 m in February (Winter); as well as water quality being affected by mineral dissolution from surrounding saline soils (Fig. 5c).
Diagrams of concentration (molality vs. CF)

As equilibrium with the mineral halite was achieved with a small number of highly saturated samples, chloride ion (Cl<sup>-</sup>) was considered the most conservative major element constituent present. This ion was subsequently used to estimate a concentration factor (CF) in the form of a ratio between chloride concentration measured in a sample solution compared to the minimum chloride ion concentration observed in a sample series (Valles 1987). The concentration factor calculated varied from 1 to 513 for surface waters. This CF variation signifies that water mineralization depends mainly on measured mineral constituent concentration in evaporating waters within an arid or hyperarid climate. Therefore, evaporation directly affected groundwaters located at the surface of the shallow water table.

In dilute waters, the molality of sodium ion (Na<sup>+</sup>) increased generally at a slower rate than chloride ion (Cl<sup>-</sup>) due to preferential ion adsorption onto clay particles. However, in this study, the line progression of both chloride and sodium ion concentrations eventually became identical (Fig. 6), due to high concentrations in water.

Figure 6 shows that water sample solution alkalinity was found to be lower in moles of dissolved calcium. Calcite residual alkalinity was therefore negative when sample water solutions were saturated. Alkalinity remained stable, however, with calcium content increase (Gac et al. 1979; Ribolzi et al. 1993) due to calcium equilibrium with fluoride ion (Barbiero et al. 1995; Nezli et al. 2009).

With gypsum precipitation, calcite + gypsum residual alkalinity was positive, where SO₄<sup>2-</sup> + alkalinity > Ca²⁺ loads in moles. Sulfate content increased while calcium content stabilized, and the chemical activity of calcium decreased with an increase in ionic strength. Successive precipitation of calcite and gypsum led to an increase in the Mg²⁺/Ca²⁺ ratio when water salinity increased.

During the winter season, when lower temperatures prevail and sodium sulfate mineral is less soluble, the precipitation of mirabilite, magnesium sulfates (hexahydrate) and sodium + magnesium sulfates, such as blödite [Na₂Mg(SO₄)₂ • 4H₂O(s)] takes place. Halite mineral precipitation is observed in the most concentrated or supersaturated saline waters, and these mineral precipitation processes have been described at length by Valles (Valles 1987). However, within the expected general variation by dissolved element constituents, some divergence does appear within said trends.

H⁺ ion activity was highly variable around the trend curve (Fig. 7). This variability was less remarkable with alkalinity, but remained somewhat important for calcium and other major elements measured. These results may be attributed to pCO₂ heterogeneity; as shown by chemical equilibrium calculations, and the layout of the sample water-calcite equilibrium diagram (Fig. 3).

H⁺ ion activity distribution was observed to be high when pCO₂ fluctuated strongly. This result may be attributed to high aerated conditions of ground water, and spatial variability of biological activity producing CO₂ concentrations within the shallow aquifer environment.
Conclusions

In this Ouargla region shallow aquifer study, major element concentrations were controlled by a complex interaction among numerous factors and various mechanisms. Arid climate induced a high concentration of shallow water mineral precipitates, resulting in the formation of saturated saline solutions and salt paragenesis. Physico-chemical mechanisms observed were Na⁺/Ca²⁺ ion exchange, and the successive precipitation of calcite, gypsum, mirabilite or blœdite and halite. Biogeochemical processes favored by anthropic effects (wastewater) were evidenced by CO₂ concentration variability and biological processes were proposed as responsible for the inverse relationship observed between measured CO₂ and O₂ concentrations in shallow groundwater samples. Other hydrogeochemical processes (e.g., mineral precipitation and dissolution) were also active within the endorheic zone of each shallow system, as represented by measured data variation from water samples collected from seasonally high evaporation and dilution periods in February (Winter) and June (Summer), respectively.

Mechanisms responsible for measured chemical composition variability in analyzed shallow groundwater samples were multiple, naturally occurring and highly variable in intensity. In this work, mechanisms associated with carbonate equilibrium, partial pressure of CO₂ and dissolved oxygen produced from biogeochemical reactions, aquifer temperature and depth; ion exchange and successive salt paragenesis were considered. As our goal was to identify mechanisms that caused variability, it was soon apparent that mechanism interdependence and complexities made it difficult to differentiate between different chemical composition sources within our measured groundwater samples independently. Although a challenging undertaking, it is suggested that in the future, an attempt should be made to establish a hierarchical organization of mechanism intensities, in spite of the many interactions that might affect the carbonate equilibria studied. This is beyond the scope of this paper, but it will be the basis for subsequent research issuing from this same study site.

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Declarations

Conflict of interest The authors declare that there is no conflict of interest.

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