Geochemical inverse modeling of chemical and isotopic data from groundwaters in Sahara (Ouargla basin, Algeria)

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Abstract

New samples were collected in the three major Saharan aquifers namely, the "Continental Intercalaire" (CI), the "Complexe Terminal" (CT) and the Phreatic aquifer (Phr) and completed with unpublished more ancient chemical and isotopic data. Instead of classical Debye-Hückel extended law, Specific Interaction Theory (SIT) model, recently incorporated in Phreeqc 3.0 was used. Inverse modeling of hydrochemical data constrained by isotopic data was used here to quantitatively assess the influence of geochemical processes: at depth, the dissolution of salts from the geological formations during upward leakage without evaporation explains the transitions from CI to CT and to a first pole of Phr (pole I); near the surface, the dissolution of salts from sebkhas by rainwater explains another pole of Phr (pole II). In every case, secondary precipitation of calcite occurs during dissolution. All Phr waters result from the mixing of these two poles together with calcite precipitation and ion exchange processes. These processes are quantitatively assessed by Phreeqc model. Globally, gypsum dissolution and calcite precipitation were found to act as a carbon sink.

Keywords: hydrochemistry, stable isotopes, Sahara, Algeria

1. INTRODUCTION

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A scientific study published in 2008 showed that 85% of the world population lives in the driest half of the Earth. More than 1 billion people residing in arid and semi-arid areas of the world have only access to little or no renewable water resources (OECD, 2008). In many arid regions such as Sahara, groundwater is the only source of water supply for domestic, agricultural or industrial purposes, causing most of the time-overuse and / or degradation of water quality.

The groundwater resources of Ouargla basin (Lower-Sahara, Algerian) (Fig. 1) are contained in three main reservoirs (UNESCO, 1972; Eckstein and Eckstein, 2003; OSS, 2003, 2008):

• at the top, the phreatic aquifer (Phr), located in sandy gypsum permeable formations of Quaternary, is almost unexploited (only north of Ouargla) due to its salinity (50 g/L);

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- in the middle, the "Complexe Terminal" (CT) aquifer, (Cornet and Gouscov, 1952; UN-ESCO, 1972) which is the most exploited, and includes several aquifers in different geological formations. He circulates in one or two lithostratigraphic formations of the Eocene and Senonian carbonates or Miopliocene sands;
- at the bottom, the "Continental Intercalaire" (CI) aquifer, where water is contained in the lower Cretaceous continental formations (Barremian and Albian), mainly composed of sandstones, sands and clays. It is only partially exploited because of its significant depth.

After use, waters are discharged in a closed system (endorheic basin) and constitute a potential hazard to the environment, to public health and may jeopardize the sustainability of agriculture (rising of the phreatic aquifer watertable, extension of soil salinization and so on) (Hamdi-Aïssa et al., 2004; Slimani, 2006). Several previous studies (Guendouz, 1985; Fontes et al., 1986; Guendouz and Moulla, 1996; Edmunds et al., 2003; Guendouz et al., 2003; Hamdi-Aïssa et al., 2004; Foster et al., 2006; OSS, 2008; Al-Gamal, 2011) tried, starting from chemical and isotopic information (²H, ¹⁸O, ²³⁴U, ²³⁸U, ³⁶Cl) to best characterize the relationships between aquifers. They were more specifically tackling the issue of the Continental Intercalaire recharge. These investigations dealt particularly with water chemical facies, mapped isocontents of various parameters, and reported typical geochemical ratios ([SO₄²⁻]/[Cl⁻], [Mg²⁺]/[Ca²⁺]) as well as other correlations. Minerals / solutions equilibria were checked by computing saturation indices with respect to calcite, gypsum, anhydrite and halite, but processes were only qualitatively assessed.

In the present study, new data were collected in order to characterize the hydrochemical and the isotopic composition of the major aquifers in Ouargla's region. They also aimed at identifying the origin of the mineralization and water-rock interactions that occur along the flow. New possibilities offered by progress in geochemical simulations were used. More specifically, the inverse modeling of chemical reactions allows us to select the best conceptual model for the interpretation of the geochemical evolution of the Ouargla aquifer. The stepwise inversion strategy involves designing a list of the scenarios that includes the most plausible combinations of geochemical processes, solving scenarios in a stepwise manner, and selecting the scenario that provides the best conceptual geochemical model (Dai et al., 2006). Inverse modeling with Phreeqc 3.0 was used to quantitatively assess the influence of the processes that explain the acquisition of solutes for the different aquifers: dissolution, precipitation, mixing and ion exchange. This results in constraints on mass balances as well as on the exchange of matter between aquifers.

2. METHODOLOGY

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2.1. Presentation of the study area

The study area is located in the northeastern desert of Algeria "Lower-Sahara" (Le Houérou, 2009) near the city of Ouargla (Fig. 1), 31°54′ to 32°1′ N and 5°15′ to 5°27′ E, with a mean elevation of 134 (masl). It is located in the quaternary fossil valley of Oued Mya basin. Present climate belongs to the arid Mediterranean-type (Dubief, 1963; Le Houérou, 2009; ONM, 1975/2013). This climate is characterized by a mean annual temperature of 22.5 °C, a yearly rainfall of 43.6 mm/yr and a very high evaporation rate of 2,138 mm/yr.

Ouargla's region and the entire Lower Sahara has experienced during its long geological history alternating marine and continental sedimentation phases. During Secondary era, vertical

movements affected the Precambrian basement and Primary causing particularly progressive collapse of its central part, along an axis passing <u>substantially</u> through the Oued Righ valley and the upper portion of the valley oued Mya. According to (Furon, 1960), a epicontinental sea <u>spread</u> to the Lower Eocene of northern Sahara. After the Oligocene, the sea gradually withdrew. It is estimated at present that this sea did not reach Ouargla and transgression stopped at the edge of the bowl (Lelièvre, 1969). The basin is carved into Miopliocene (MP) deposits, which alternate with red sands, clays and sometimes marls; gypsum is not abundant and dated from Pontian (MP) (Cornet and Gouscov, 1952; Dubief, 1953; Ould Baba Sy and Besbes, 2006). The continental Pliocene consists of a local limestone crust with puddingstone or lacustrine limestone (Fig. 2), shaped by eolian erosion into flat areas (regs). The Quaternary formations are lithologically composed of alternating layers of permeable sand and relatively impermeable <u>marl</u> (Aumassip et al., 1972; Chellat et al., 2014).

The exploitation of Miopliocene aquifer is ancient and at the origin of the creation of the oasis (Lelièvre, 1969; Moulias, 1927). The piezometric level was higher (145 m a.s.l.) but over-exploitation at the end of the XIXth century led to a catastrophic decrease of the resource, with presently more than 900 boreholes (ANRH, 2011).

The exploitation of Senonian aquifer dates back to 1953 at a depth 140 m to 200 m depth, with a small initial rate ca. $540 \,\mathrm{L}\,\mathrm{mn}^{-1}$; two boreholes have been exploited since 1965 and 1969, with a total flowrate ca. $2,500 \,\mathrm{L}\,\mathrm{mn}^{-1}$, for drinking water and irrigation.

The exploitation of Albian aquifer dates back to 1956, with <u>a piezometric level 405 m</u> and a <u>pressure 22 kg cm</u>⁼². Presently, two boreholes are exploited:

- El Hedeb I, 1,335 m $\frac{\text{depth}_{3}}{\text{g}}$ with a flowrate 141 L s⁻¹;
- El Hedeb II, 1,400 m depth, with a flowrate 68 L s⁻¹.

2.2. Sampling and analytical methods

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The sampling scheme complies with the flow directions of the two formations (Phr and CT aquifers); for the CI aquifer only five points are available, so it is impossible to choose a transect (Fig. 3). Groundwater samples (n = 107) were collected during a field campaign in 2013, along the main flow line of Oued Mya, 67 piezometers tap the phreatic aquifer, 32 wells tap the CT aquifer and 8 boreholes tap the CI aquifer (Fig. 3). Analyses of Na⁺, K⁺, Ca²⁺, Mg²⁺, Cl⁻, SO₄²⁻ and HCO₃⁻ were performed by ion chromatography at Algiers Nuclear Research Center (CRNA). Previous and yet unpublished data (Guendouz and Moulla, 1996) sampled in 1996 are used here too: 59 samples for Phr aquifer, 15 samples for CT aquifer and 3 samples for the CI aquifer for chemical analyses, data ¹⁸O and ³H (Guendouz and Moulla, 1996).

2.3. Geochemical method

Phreeqc (Parkhurst and Appelo, 2013) was used to check minerals / solution equilibria using the specific interaction theory (SIT), *i.e.* the extension of Debye-Hückel law by Scatchard and Guggenheim incorporated recently in Phreeqc 3.0. Inverse modeling was used to calculate the number of minerals and gases' moles that must respectively dissolve or precipitate/degas to account for the difference in composition between initial and final water end members (Plummer and Back, 1980; Kenoyer and Bowser, 1992; Deutsch, 1997; Plummer and Sprinckle, 2001; Güler and Thyne, 2004; Parkhurst and Appelo, 2013). This mass balance technique has been used to quantify reactions controlling water chemistry along flow paths (Thomas et al., 1989). It is also used to quantify the mixing proportions of end-member components in a flow system

(Kuells et al., 2000; Belkhiri et al., 2010, 2012). The Inverse modeling involves designing a list of the scenarios that includes the most plausible combinations of geochemical processes. For example, the way to identify whether calcite dissolution/precipitation is relevant or not consists of solving the inverse problem under two alternate scenarios: (1) considering a geochemical system in which calcite is present, and (2) considering a geochemical system without calcite. After solving the two scenarios, it is usually possible to select the better result as the solution of the inverse problems and conclude whether calcite dissolution/precipitation is relevant or not. This stepwise strategy allows us to identify the relevance of a given chemical process by solving the inverse problem under alternative scenarios in which the process is either occurring or not.

3. RESULTS AND DISCUSSION

Tables 1 to 5 illustrate the results of the chemical and the isotopic analyses. Samples are ordered according to an increasing salt content that was estimated from their specific electric conductivity (EC). In both phreatic and CT aquifers, temperature is close to 25 °C, while for CI aquifer, temperature is close to 50 °C. The results presented in those tables are raw analytical data that were corrected for defects of charge balance before computing activities with Phreeqc. As analytical errors could not be ascribed to a specific analyte, the correction was made proportionally. The corrections do not affect the anions to anions mole ratios such as for $[HCO_3^-]/([Cl^-] + 2[SO_4^{2-}])$ or $[SO_4^{2-}]/[Cl^-]$, whereas they affect the cation to anion ratio such as for $[Na^+]/[Cl^-]$.

3.1. Characterization of chemical facies of the groundwater

Piper diagrams drawn for the studied groundwaters (Fig. 4) broadly show a scatter plot dominated by a Chloride-Sodium facies. However, when going into small details, the widespread chemical facies of the Phr aquifer is closer to the NaCl <u>pole</u> than those of CI and CT aquifers. The facies of the Phreatic aquifer most concentrated samples are in the following order: Ca-sulfate \leq Na-sulfate \leq Na-chloride. This sequential order of solutes is comparable to that of other groundwater occurring in North Africa, and especially in the neighboring area of the chotts (depressions where salts concentrate by evaporation) Merouane and Melrhir (Vallès et al., 1997; Hamdi-Aïssa et al., 2004).

3.2. Spatial distribution of the mineralization

The salinity of the phreatic aquifer varies considerably depending on the location (near wells or drains) and time (influence of irrigation) (Fig. 5a).

Its salinity is low around irrigated and fairly well-drained areas, such as the palm groves of Hassi Miloud, just north of Ouargla (Fig. 3) that benefit from freshwater and are drained to the sebkha Oum el Raneb. However, the three lowest salinity values are observed in the wells of Ouargla palm-grove itself, where the Phr aquifer watertable is deeper than 2 m.

Conversely, the highest salinity waters are found in wells drilled in the chotts and sebkhas (a sebkha is the central part of a chott where salinity is the largest) (Safioune and Oum er Raneb) where the aquifer is often shallower than 50 cm.

The salinity of the Complexe Terminal (Miopliocene) aquifer (Fig. 5b) is much lower than that of the Phr aquifer, and ranges from 1 to 2 g/L; however, its hardness is larger and it contains more sulfate, chloride and sodium than the waters of the Senonian formations and those of the

CI aquifer. The salinity of the Senonian aquifer ranges from 1.1 to 1.7 g/L, while the average salinity of the Continental Intercalaire is 0.7 g/L (Fig. 5c).

A likely contamination of the Miopliocene aquifer by phreatic groundwaters through casing leakage in an area where water is heavily loaded with salt and therefore particularly aggressive cannot be excluded.

3.3. Saturation Indices

The calculated saturation indices reveal that waters from CI at 50 °C are close to equilibrium with respect to calcite, except for 3 samples that are slightly oversaturated. They are however all undersaturated with respect to gypsum (Fig. 6).

Moreover, they are oversaturated with respect to dolomite and undersaturated with respect to anhydrite and halite (Fig. 7).

Waters from CT and phreatic aquifers show the same pattern, but some of them are more largely oversaturated with respect to calcite, at 25 °C.

However, several phreatic waters (P031, P566, PLX4, PL18, P002, P023, P116, P066, P162 and P036) that are located in the sebkhas of Sefioune, Oum-er-Raneb, Bamendil and Ain el Beida's chott are saturated with gypsum and anhydrite. This is in accordance with high evaporative environments found elsewhere (UNESCO, 1972; Hamdi-Aïssa et al., 2004; Slimani, 2006).

No significant saturation indices' evolution from the south to the north upstream and down-stream of Oued Mya (Fig. 7) is observed. This suggests that the acquisition of mineralization is due to geochemical processes that have already reached equilibrium or steady state in the upstream areas of Ouargla.

3.4. Change of facies from the carbonated pole to the evaporites' pole

The facies shifts progressively from the carbonated (CI and CT aquifers) to the evaporites' one (Phr aquifer) with an increase in sulfates and chlorides at the expense of carbonates (SI of gypsum, anhydrite and halite). This is illustrated by a decrease of the following two ratios: $[HCO_3^-]/([CI^-] + 2[SO_4^{2-}])$ (Fig. 8) from 0.2 to 0 and of the ratio $[SO_4^{2-}]/[CI^-]$ from 0.8 to values ranging from 0.3 and 0 (Fig. 9) while salinity increases. Carbonate concentrations tend towards very small values, while it is not the case for sulfates. This is due to both gypsum dissolution and calcite precipitation.

Chlorides in groundwater may come from three different sources: (i) ancient sea water entrapped in sediments; (ii) dissolution of halite and related minerals that are present in evaporite deposits and (iii) dissolution of dry fallout from the atmosphere, particularly in these arid regions (Matiatos et al., 2014; Hadj-Ammar et al., 2014).

For most of the sampled points the [Na⁺]/[Cl⁻] ratio <u>remains close to 1</u>, but <u>significant ranges</u> are observed: from 0.85 to 1.26 for CI aquifer, from 0.40 to 1.02 for the CT aquifer and from 0.13 to 2.15 for the Phr aquifer. All the measured points from the three considered aquifers are more or less linearly scattered around the unity slope straight line that stands for halite dissolution (Fig. 10). The latter appears as the most dominant reaction occurring in the medium. However, at very high salinity, Na⁺ seems to swerve from the straight line, towards smaller values.

A further scrutiny of (Fig. 10) shows that CI waters are very close to the 1:1 line. CT waters are enriched in both Na⁺ and Cl⁻ but slightly lower than the 1:1 line while phreatic waters are largely enriched and much more scattered. CT waters are closer to the <u>seawater mole ratio</u> (0.858), but some lower values imply a contribution from another source of chloride than halite or from entrapped seawater. Conversely, a [Na⁺]/[Cl⁻] ratio larger than 1 is observed for phreatic

waters, which implies the contribution of another source of sodium, most likely sodium sulfate, that is present as mirabilite or thenardite in the chotts and the sebkhas areas.

[Br⁻]/[Cl⁻] ratio ranges from 2×10^{-3} to 3×10^{-3} . The value of this molar ratio for halite is around 2.5×10^{-3} , which matches the aforementioned range and confirms that halite dissolution is the most dominant reaction taking place in the studied medium.

In <u>these</u> aquifers, calcium originates both from carbonate and sulfate (Fig. 11 and 12). Three samples from CI aquifer are close to the [Ca²⁺]/[HCO₃⁻] 1:2 line, while calcium sulfate dissolution explains the excess of calcium. However, a small but significant number of samples (9) from phreatic aquifer are depleted in calcium, and plot under the [Ca²⁺]/[HCO₃⁻] 1:2 line. This cannot be explained by precipitation of calcite, as some are undersaturated with respect to that mineral, while others are oversaturated.

In this case, a cation exchange process seems to occur leading to a preferential adsorption of divalent cations, with a release of Na⁺. This is confirmed by the inverse modeling that is developed below and which implies Mg²⁺ fixation and Na⁺ and K⁺ releases.

Larger sulfate values observed in the phreatic aquifer (Fig. 12) with $[Ca^{2+}]/[SO_4^{2-}] < 1$ can be attributed to a sodium magnesium sulfate dissolution from a mineral bearing such elements. This is for instance the case of bloedite.

3.5. Isotope geochemistry

CT and CI aquifer exhibit depleted and homogeneous 18 O contents, ranging from -8.32 % to -7.85 %. This was already previously reported by many authors (Edmunds et al., 2003; Guendouz et al., 2003; Moulla et al., 2012). On the other hand, 18 O values for the phreatic aquifer are widely dispersed and vary between -8.84 % to 3.42 % (Table 6).

Waters located north of the <u>Hassi Miloud to Sebkhet Safioune axis</u> are more enriched in heavy isotopes and therefore more evaporated. In that area, water table is close to the surface and mixing of both CI and CT groundwaters with phreatic ones through irrigation is nonexistent. Conversely, waters located south of Hassi Miloud up to Ouargla city show depleted values. This is the clear fingerprint of a contribution to the Phr waters from the underlying CI and CT aquifers (Gonfiantini et al., 1975; Guendouz, 1985; Fontes et al., 1986; Guendouz and Moulla, 1996).

Phreatic waters result from a mixing of two end-members. An evidence for this is given by considering the ([Cl⁻], ¹⁸O) relationship (Fig. 13). The two poles are: i) a first pole of ¹⁸O depleted groundwater (Fig. 14), and ii) another pole of ¹⁸O enriched groundwater with positive values and a high salinity. The latter is composed of phreatic waters occurring in the northern part of the study region.

Pole I represents the waters from CI and CT whose isotopic composition is depleted in ¹⁸O (average value around -8.2%) (Fig. 13). They correspond to an old water recharge (palæorecharge); whose age estimated by means of ¹⁴C, exceeds 15.000 years BP (Guendouz, 1985; Guendouz and Michelot, 2006). So, it is not a water body that is recharged by recent precipitation. It consists of CI and CT groundwaters and partly of phreatic waters, and can be ascribed to an upward leakage favored by the extension of faults near Amguid El-Biod dorsal.

Pole II, observed in Sebkhet Safioune, can be ascribed to the direct dissolution of surficial evaporitic deposits conveyed by evaporated rainwater.

Evaporation alone cannot explain the distribution of data that is observed (Fig. 13). An evidence for this is given in a semi-logarithmic plot (Fig. 14), as classically obtained according to the simple approximation of Rayleigh equation (cf. Appendix):

$$\delta^{18}O \approx 1000 \times (1 - \alpha) \log[\text{CI}^-] + \underline{cte},$$
 (1)

$$\approx -\epsilon \log[\text{Cl}^-] + cte,$$
 (2)

where α is the fractionation factor during evaporation, and $\epsilon = -1000 \times (1 - \alpha)$ is the enrichment factor (Ma et al., 2010; Chkir et al., 2009).

CI and CT waters are better separated in the semi-logarithmic plot because they are differentiated by their chloride content. According to equation (1), simple evaporation gives a straight line (solid line in Fig. 14). The value of ϵ used is the value at 25 °C, which is equal to -73.5. There is only one sample (P115) on the evaporation straight line, which could be considered as an outlier in Fig. 13 ([Cl⁼] \simeq 0). All other samples fit on the logarithmic curve derived from the mixing line illustrated by Fig. 13.

The phreatic waters that are close to pole I (Fig. 13) correspond to groundwaters occurring in the edges of the basin (Hassi Miloud, piezometer P433) (Fig. 14). They are low-mineralized and acquire their salinity via two processes namely: dissolution of evaporites along their underground transit up to Sebkhet Safioune and dilution through upward leakage by the less-mineralized waters of CI and CT aquifers (for example Hedeb I for CI and D7F4 for CT) (Fig. 14) (Guendouz, 1985; Guendouz and Moulla, 1996).

The rates of the mixing that are due to upward leakage from CI to CT towards the phreatic aquifer can be calculated by means of a mass balance equation. It only requires knowing the δ values of each fraction that is involved in the mixing process.

values of each fraction that is involved in the mixing process.

The
$$\delta$$
 value of the mixture is given by:
$$f_1 \times \delta_1 + f_2 \times \delta_2$$
(3)

where f_1 is the fraction of CI aquifer, f_2 the fraction of the CT and δ_1 , δ_2 are the respective isotope contents.

Average values of mixing fractions from each aquifer to the phreatic waters computed by means of equation (3) gave the rates of 65 % for CI aquifer and 35% for CT aquifer.

A mixture of a phreatic water component that is close to pole I (*i.e.* P433) with another one which is rather close to pole II (*i.e.* P039) (Fig. 13 and 14), for an intermediate water with a δ^{18} O signature ranging from -5% to -2% gives mixture fraction values of 52 % for pole I and 48 % for pole II. Isotope results will be used to independently cross-check the validity of the mixing fractions derived from an inverse modeling involving chemical data (*cf. infra*).

Turonian evaporites are found to lie in between CI deep aquifer, and the Senonian and Miocene formations bearing CT aquifer. CT waters can thus simply originate from ascending CI waters that dissolve Turonian evaporites, a process which does involve any change in ¹⁸O content. Conversely, phreatic waters result to a minor degree from evaporation, and mostly from dissolution of sebkhas evaporites by ¹⁸O enriched rainwater and mixing with CI-CT waters.

3.5.1. Tritium content of water

Tritium contents of Phr aquifer are relatively small (Table 6), they vary between 0 and 8 TU. Piezometers PZ12, P036 and P068 show values close to 8 TU, piezometers P018, P019, P416, P034, P042 and P093 exhibit values ranging between 5 and 6 TU, and the rest of the samples' concentrations are lower than 2 TU.

This values are dated back to November 1992 so they are old values. This is the main reason why they are considered high comparatively to what is expected to be found nowadays. In fact,

at present times, tritium figures have fallen lower than 5 TU in precipitation measured in the northern part of the country.

The comparison of these results with that of precipitation which was 16 TU in 1992 was collected from the National Agency for Water Resources station from Ouargla).

This value seems to be high but we can note that we are in an arid area (desert) where precipitation is very scarce and irregular. Precipitation takes place in the form of sudden thunderstorms in an unsaturated atmosphere and a great part of this precipitation evaporates back into the moisture unsaturated atmosphere sometimes during many cycles. Consequently, an enrichment in tritium happens because when water evaporates back, the lightest fractions (isotopes) are the ones that evaporate first causing an enrichment in Tritium in the remaining fraction. The 16 TU value would thus correspond then to a rainy event that had happened during the same sampling period (Nov. 1992). It's the only available value and it is not a weighted mean for a long period of time. It is the most representative value for that region and for that time. Unfortunately, all the other stations (Algiers, Ankara, and Tenerife) (Martinelli et al., 2014) are subject to a completely different climatic regime and besides the fact that they have more recent values, can absolutely not be used for our case. Therefore all the assumptions based on recent tritium rain values do not apply to this study.

Depleted contents in ¹⁸O and low tritium concentrations for phreatic waters fit well the mixing scheme and confirm the contribution from the older and deeper CI/CT groundwaters. The affected areas were clearly identified in the field and correspond to locations that are subject to a recycling and a return of irrigation waters whose origin are CI/CT boreholes. Moreover, the mixing that is clearly brought to light by the Cl⁻ vs. ¹⁸O diagrams (Fig. 13 and 14) could partly derive from an ascending drainage from the deep and confined CI aquifer (exhibiting depleted homogenous ¹⁸O contents and very low tritium), a vertical leakage that is favoured by the Amguid El-biod highly faulted area (Guendouz and Moulla, 1996; Edmunds et al., 2003; Guendouz et al., 2003; Moulla et al., 2012).

3.6. Inverse modeling

We assume that the relationship between ^{18}O and Cl^- data obtained in 1996 is stable with time, which is a logical assumption as times of transfer from CI to both CT and Phr are very long. Considering both ^{18}O and Cl^- data, CI, CT and Phr data populations can be categorized. The CI and CT do not show appreciable ^{18}O variations, and can be considered as a single population. The Phr samples consist however of different populations: Pole I, with $\delta^{18}O$ values close to -8, and small Cl^- concentrations, more specifically less than 35 mmol L^{-1} ; Pole II, with $\delta^{18}O$ values larger than 3, and very large Cl^- concentrations, more specifically larger than 4,000 mmol L^{-1} (Table 7); intermediate Phr samples result from mixing between poles I and II (mixing line in Fig. 13, mixing curve in Fig. 14) and from evaporation of pole I (evaporation line in Fig. 14).

The mass-balance modeling has shown that relatively few phases are required to derive observed changes in water chemistry and to account for the hydrochemical evolution in Ouargla's region. The mineral phases' selection is based upon geological descriptions and analysis of rocks and sediments from the area (OSS, 2003; Hamdi-Aïssa et al., 2004).

The inverse model was constrained so that mineral phases from evaporites including gypsum, halite, mirabilite, glauberite, sylvite and bloedite were set to dissolve until they reach saturation, and calcite, dolomite were set to precipitate once they reached saturation. Cation exchange reactions of Ca²⁺, Mg²⁺, K⁺ and Na⁺ on exchange sites were included in the model to check which cations are adsorbed or desorbed during the process. Dissolution and desorption contribute as

positive terms in the mass balance, as elements are released in solution. On the other hand, precipitation and adsorption contribute as negative terms, while elements removed from the solution. $CO_{2(g)}$ dissolution is considered by Phreeqc as a dissolution of a mineral, whereas $CO_{2(g)}$ degassing is dealt with as if it were a mineral precipitation.

Inverse modelling leads to a quantitative assessment of the different solutes' acquisition processes and a mass balance for the salts that are dissolved or precipitated from CI, CT and Phr groundwaters (Fig. 14, Table 8), as follows:

- transition from CI to CT involves gypsum, halite and sylvite dissolution, and some ion
 exchange namely calcium and potassium fixation on exchange sites against magnesium
 release, with a very small and quite negligible amount of CO_{2(g)} degassing. The maximum
 elemental concentration fractional error equals 1%. The model consists of a minimum
 number of phases (i.e. 6 solid phases and CO_{2(g)}); Another model implies as well dolomite
 precipitation with the same fractional error;
- transition from CT to an average water component of pole I involves dissolution of halite, sylvite, and bloedite from Turonian evaporites, with a very tiny calcite precipitation. The maximum fractional error in elemental concentration is 4%. Another model implies CO_{2(g)} escape from the solution, with the same fractional error. Large amounts of Mg²⁺ and SO₄²⁻ are released within the solution (Sharif et al., 2008; Li et al., 2010; Carucci et al., 2012);
- the formation of Phr pole II can be modeled as being a direct dissolution of salts from the sebkha by rainwater with positive δ¹⁸O; the most concentrated water (P036 from Sebkhet Safioune) is taken here for pole II, and pure water as rainwater. In a decreasing order of amounts respectively involved in that process, halite, sylvite, gypsum and huntite dissolve, and little calcite precipitates while some Mg²⁺ are released versus K⁺ fixation on exchange sites. The maximum elemental fractional error in the concentration is equal to 0.004%. Another model implies dolomite precipitation with some more huntite dissolving, instead of calcite precipitation, but salt dissolution and ion exchange are the same. Huntite, dolomite and calcite stoichiometries are linearly related, so both models can fit field data, but calcite precipitation is preferred compared to dolomite precipitation at low temperature;
- the origin of all phreatic waters can be explained by a mixing in variable proportions of pole I and pole II. For instance, waters from pole I and pole II can easily be separated by their $\delta^{18}O$ respectively close to -8% and 3.5 % (Fig. 13 and 14). Mixing the two poles is of course not an inert reaction, but rather results in the dissolution and the precipitation of minerals. Inverse modeling is then used to compute both mixing rates and the extent of matter exchange between soil and solution. For example, a phreatic water (piezometer P068) with intermediate values ($\delta^{18}O = -3$ and $[Cl^-] \simeq 2 M$) is explained by the mixing of 58% water from pole I and 42% from pole II. In addition, calcite precipitates, Mg^{2+} fixes on exchange sites, against Na^+ and K^+ , gypsum dissolves as well as a minor amount of huntite (Table 8). The maximum elemental concentration fractional error is 2.5% and the mixing fractions' weighted the $\delta^{18}O$ is -3.17%, which is is very close to the measured value (-3.04%). All the other models, making use of a minimum number of phases, and not taking into consideration ion exchange reactions are not found compatible with isotope data. Mixing rates obtained with such models are for example 98% of pole I and 0.9% of

pole II, which leads to a $\delta^{18}O = (-7.80\%)$ which is quite far for the real measured value (-3.04%).

The main types of groundwaters occurring in Ouargla basin are thus explained and could quantitatively be reconstructed. An exception is however sample P115, which is located exactly on the evaporation line of Phr pole I. Despite numerous attempts, it could not be quantitatively rebuilt. Its ³H value (6.8) indicates that it is derived from a more or less recent water component with very small salt content, most possibly affected by rainwater and some preferential flow within the piezometer. As this is the only sample on this evaporation line, there remains a doubt on its significance.

Globally, the summary of mass transfer reactions occurring in the studied system (Table 8) shows that gypsum dissolution results in calcite precipitation and $CO_{2(g)}$ dissolution, thus acting as an inorganic carbon sink.

4. CONCLUSIONS

 Groundwater hydrochemistry is a good record indicator for the water-rock interactions that occur along the groundwater flowpath. The mineral load reflects well the complex processes taking place while water circulates underground since its point of infiltration.

The hydrochemical study of the aquifer system occurring in Ouargla's basin allowed us to identify the origin of its mineralization. Waters exhibit two different facies: sodium chloride and sodium sulfate for the phreatic aquifer (Phr), sodium sulfate for the Complexe Terminal (CT) aquifer and sodium chloride for the Continental Intercalaire (CI) aquifer. Calcium carbonate precipitation and evaporite dissolution explain the facies change from carbonate to sodium chloride or sodium sulfate. However reactions imply many minerals with common ions, deep reactions without evaporation as well as shallow processes affected by both evaporation and mixing. Those processes are separated by considering both chemical and isotopic data, and quantitatively explained making use of an inverse geochemical modeling. The main result is that Phr waters do not originate simply from infiltration of rainwater and dissolution of salts from the sebkhas. Conversely, Phr waters are largely influenced by the upwardly mobile deep CT and CI groundwaters, fractions of the latter interacting with evaporites from Turonian formations. Phreatic waters occurrence is explained as a mixing of two end-member components: pole I, which is very close to CI and CT, and pole II, which is highly mineralized and results from the dissolution by rainwater of salts from the sebkhas.

At depth, CI leaks upwardly and dissolves gypsum, halite and sylvite, with some ion exchange, to give waters of CT aquifer composition.

CT transformation into Phr pole I waters involves the dissolution of Turonian evaporites (halite, sylvite and bloedite) with minor calcite precipitation.

At the surface, direct dissolution by rainwater of salts from sebkhas (halite, sylvite, gypsum and some huntite) with precipitation of calcite and Mg^{2+}/K^+ ion exchange results in pole II Phr composition.

All phreatic groundwaters result from a mixing of pole I and pole II water that is accompanied by calcite precipitation, fixation of Mg^{2+} on ion exchange sites against the release of K^+ and Na^+ .

Moreover, some $CO_{2(g)}$ escapes from the solution at depth, but dissolves much more at the surface. The most complex phenomena occur during the dissolution of Turonian evaporites while CI leaks upwardly towards CT, and from Phr I to Phr II, while the transition from CT to Phr I

implies a very limited number of phases. Globally, gypsum dissolution and calcite precipitation processes both act as an inorganic carbon sink. 420

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APPENDIX 427

According to a simple Rayleigh equation, the evolution of the heavy isotope ratio in the 428 remaining liquid R_l is given by: 429

$$R_l \approx R_{l,0} \times f_l^{\alpha - 1},$$
 (4)

where f_l is the fraction remaining liquid and α the fractionation factor. 430

The fraction remaining liquid is derived from chloride concentration, as chloride can be con-431 sidered as conservative during evaporation: all phreatic waters are undersaturated with respect to 432 halite, that precipitates only in the last stage. Hence, the following equation holds:

$$f_l \equiv \frac{n_{w,1}}{n_{w,0}} = \frac{[\text{Cl}^-]_0}{[\text{Cl}^-]_1}.$$
 (5)

By taking natural logarithms, one obtains:

$$\ln R_l \approx (1 - \alpha) \times \ln[\text{Cl}^-] + cte, \tag{6}$$

As, by definition, 435

$$R_l \equiv R_{std.} \times (1 + \frac{\delta^{18}O}{1000}),$$
 (7)

one has: 436

434

437

438

$$ln R_l \equiv ln R_{std.} + ln(1 + \frac{\delta^{18}O}{1000}),$$
(8)
$$\approx ln R_{std.} + \frac{\delta^{18}O}{1000},$$
(9)

$$\approx \ln R_{std.} + \frac{\delta^{18}O}{1000},\tag{9}$$

hence, with base 10 logarithms:

$$\delta^{18}O \approx 1000(1-\alpha)\log[\text{Cl}^-] + cte, \tag{10}$$

$$\approx -\epsilon \log[\text{Cl}^-] + cte, \tag{11}$$

$$\approx -\epsilon \log[\text{Cl}^-] + cte,$$
 (11)

where as classically defined $\epsilon = 100(\alpha - 1)$ is the enrichment factor.

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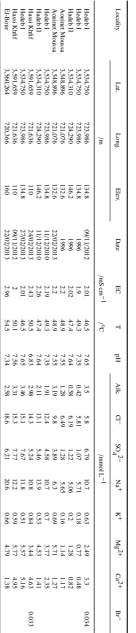


Table 1: Field and analytical data for the Continental Intercalaire aquifer.

Table 2: Field and analytical data for the Complexe Terminal aquifer.

Locality	Site	Aquifer	Lat.	Long.	Elev.	Date	BC	F	Hd	Alk.	G10	so_4^{2-}	Na+	K ⁺	Mg ²⁺	Ca ² +	Br-
				/m			$/\mathrm{mS}\mathrm{cm}^{-1}$	/°C					$/\text{mmol L}^{-1}$	-1			
Bamendil Bamendil Ifri Said Otba Oglat Larbaâ El-Bour Said Otba I	D7F4 D7F4 D1F151 D2F66 D6F64 D4F94 D2F71	MMSSMMS	3,560,759,6 3,560,759,6 3,588,91,7 3,506,501,4 3,556,501,4 3,556,412,4 3,557,112,4	720,586.2 720,586.2 720,085.4 729,369.3 722,641.7 718,722.8	296 296 204 216 177 100.6 211.9	20/01/2013 1996 1996 1996 1996 27/01/2013 26/01/2013	2.02 2.02 2.67 2.33 2.23 2.23 2.23	20.1 21.1 23.5 24 26.2 26.2 24.2 24.2	7.86 8.2 7 8 8 7.9 7.37 8.2 7.37	1.63 0.96 1.26 1.43 1.41 1.61 1.54	10.1 10.6 10.75 11.02 11.36 12.8 13.53	5.79 3.54 4.73 6.85 5.72 8.41	9.88 10.61 7.99 11.47 5.15 5.15 14.99	0.68 0.09 0.73 0.16 2.31 1.94 0.33	3.92 2.33 2.32 2.07 1.96 3.28 5.38	2.51 1.8 2.12 3.33 4.58 9.13 4.43	
Rouissat III Said Otba I Souid Otba I Iffi Said Otba Oglat Larbaâ SAR Mekhadma	D3F10 D2F71 D3F10 D1F151 D2F66 D6F64 D1F91	×××××××××××××××××××××××××××××××××××××	3,335,068.1 3,335,068.1 3,338,891.7 3,338,891.7 3,540,257.3 3,566,501.4 3,336,757.7	722,352.1 718,272.8 721,060.5 720,085.4 729,369.3	248 211.9 204 216 177	1996 26/01/2013 20/01/2013 27/01/2013 31/01/2013 03/02/2013	2.33 2.33 2.33 2.33 4.4 4.4	252 253 253 253 253 253 253 253 253 253	7.27 7.34 7.79 7.62 7.62	2.39 1.65 2.19 2.33 3.43	14.27 15.2 15.4 16.1 16.3	6.89 6.86 8.64 8.65 8.65	13.05 13.1 13.7 16.5 16.5	0.04 0.022 0.022 0.074 0.071	5.33 5.79 5.17 5.86 5.27	4.25 4.25 4.29 4.97	0.034
Sidi Kouiled Ain N'sara A.Louise	D9F12 D6F50 D4F73	ννν ;	3,540,855.1 3,559,323.6 3,537,523.4	729,055.4 716,868.4 721,904.6	329 255 310	24/01/2013 25/01/2013 26/01/2013	2.57 2.57 2.57	25.7 24.7 25.7	8.05 7.36	2.1.98 8.1.98 7.00 7.00 7.00 7.00 7.00 7.00 7.00 7.0	16.8 17.4	8.85 9.71 9.04	16.1 15.9 13.9	0.00 5.80 5.80 6.80 6.80 6.80 6.80 6.80 6.80 6.80 6	339 578 578	5.01 5.05	0.033
Christian A.H. Ain Mussa II Ain N'sara H.Miloud H.Miloud H.Miloud M. goussa El Hou El Koum	D0F/9 D0F30 D0F30 D1F135 D0F97 D0F67 D0F67	≅∾∾∑∾∑∾∾;	3,537,814, 3,593,32,6 3,547,557,1 3,540,936,5 3,547,557,1 3,573,694,1 3,573,694,1	720,536.8 716,868.4 717,067.1 717,067.1 718,979.5 721,639.7	220.6 255 173 169 1143 1143	02/02/2013 02/02/2013 02/02/2013 03/02/2013 1996 31/01/2013 21/01/2013	7.7.7.7.7.7.7.7.7.7.7.7.7.7.7.7.7.7.7.	22 23 23 23 23 23 23 23 23 23 23 23 23 2	7.53 7.55 7.55 8.02 8.03 8.09 7.52	2.23.47 2.23.33.1 3.25.03 3.55.03	4.5.7.7.7.9 17.9 17.9 18.0 18.4 18.4 18.7 18.7 18.7 18.7 18.7	923 924 922 922 923 971 717	15.5 15.5 15.8 16.61 17.9 10.18	0.39 0.39 1.13 1.01 0.51 0.32 3.33 3.33 3.34 3.34	6.11 6.11 6.11 5.84 3.65 6.17 6.49	6,46 6,46 4,73 4,73 5,84 5,84 5,84 5,84 5,84 5,84 5,84 5,84	0.033
I I AS Ain moussa V El-Bour Rouissat I Rouissat I St. pompage chott Chott Palmeraie	DIFISO D9F13 D3F18 D3F18 D5F80 D5F77	ZZZZZSS	3,536,186,6 3,538,499,2 3,535,544,2 3,535,564,2 3,541,656,9 3,538,219,3 3,545,533,1	718,680.2 722,641.7 722,498.9 722,498.9 725,541.3	293.1 100.6 80.4 80.4 242.1 86.8	21/01/2013 08/02/2013 1996 26/01/2013 1996 04/02/2013 05/02/2013 05/02/2013	98,828 13,388 14,378 13,388	252 252 254 254 254 254 254 254 254 254	7.7.2.2 7.7.2.2 7.7.2.3 8.2.3 7.2.3 7.3.3	2.58 2.38 2.38 2.38 2.38 2.38 2.38 2.38 2.3	222.56 222.56 223.3.3.1.66 23.3.3.1.66	707 945 11.1 11.9 12.1 12.1 13.0 14.0 15.0 16.0 16.0 16.0 16.0 16.0 16.0 16.0 16	10.1 11.2.09 11.7.2 20.9 20.9	20.20.3.4 20.1.2.1.1.2.2.3.4 3.4.2.3.3.4 3.4.3.3.3.4 3.4.3.3.3.4 3.4.3.3.4 3.4.3.3.4 3.4.3.3.4 3.4.3.4	4.94 9.33 7.08 7.08 8.25 1.38 1.39 1.30 1.30 1.30 1.30 1.30 1.30 1.30 1.30	55.77 6.01 6.28 5.78 5.78	0.034
Abazat Garet Chemia Frane Oum Raneb	D2F69 D1F113 D6F62 D6 F69 D6F51	E S Z Z S ;	3,552,504.9 3,536,174.1 3,570,175.8 3,556,256.7	712,786.3 716,808.5 721,919.8 721,979.5	137.1 213.7 167.5 215.8 198	03/02/2013 28/01/2013 27/01/2013 25/01/2013 1996	4.8.4.8.4.8.4.8.4.2.4.8.4.2.4.4.3.7.9	23.1.2 23.1.2 23.1.2	7.95 7.03 8	2 2 2 2 2 2 2 3 2 4 3 2 4 3 3 3 3 3 3 3	2835 2835 3935 3955 3955 3955 3955 3955 3955 3	13.5 8.67 8.61	222.6 222.6 23.14	0.57 0.00 0.00 0.00 0.00 0.00 0.00 0.00	8.8.8.3.8.4.5.4.4.2.4.4.2.4.4.2.4.4.2.4.4.2.4.4.2.4.4.2.4	6.47 7.17 7.16 8.01	0.037
H.Miloud Benyaza Ain Laarab H.Miloud Benyaza Rouissat Rouissat Ain El Arch St. pompage chott	DIF138 DIF138 D3F8 D3F8 D3F26 D5F80	SZZZZZ	3,531,192.5 3,558,822.6 3,551,192.5 3,545,470.7 3,534,843.9 3,541,656.9	717,042.1 716,042.1 732,837.6 732,837.6 723,381.8 723,521.9	186.5 186.5 332.4 332.4 93.6 224.1	28/01/2013 28/01/2013 1996 03/02/2013 1996 1996	3.83 2.99 6.16 3.69 3.69	088888 0584614	7.61 7.33 7.51 7.22 7.45	2.16 2.16 1.71 1.71 2.28	28.9 28.9 29.8 29.8 42.22	9.01 9.03 8.33 8.33 8.34 13.53	23.9 22.8 22.8 22.8 36.77	0.53 0.53 1.23 0.87 1.12	10.01 5 4.99 6.23 8.38 7.43	7.12 7.72 7.7 6.08 6.08 6.5	0.037

= Miopliocene aquifer; S = Senonian aquifer

Table 3: Field and analytical data for the Phreatic aquifer.

	Ecole Okba B. Nafaa	Route Ain Moussa	Route Ain Moussa	Bour El Haicha	PARC SONACOM	Hôpital	DSA	Hassi Debich	Station d'épuration	Route El Goléa	Route El Goléa	DSA	Ecole paramédicale	Route Ain Moussa	Route El Goléa	Gherbouz	Route El Goléa	H Chegga	Polyclinique Belabès	Mekmahad	Route El Goléa	Route Ain Moussa	Hassi Miloud	H Chegga	Route Ain Bida	Frane Ank Djemel	Station d'épuration	Bour El Haicha	Gherbouz	Frane El Koum	Maison de culture	Oglet Larbaa	Hassi Miloud	El Bour	Maison de culture	,	Hassi Naga	Ain Kheir	Hassi Miloud	Khezana	Khezana		Locality
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	PL41	P056	P056	P408	PL28	LTPSN2	PL10	P416	PL30	P116	P117	PL10	PL32	P057	P117	PL15	P116	PLX4	PL18	PL05	P115	P057	P058	PLX4	PLX2	P422	PL30	P408	PL15	P401	PL31	P430	P059	P006	PL31	LTP 30	PLX3	PL06	P059	P433	P433		Site
Best Date EC T pH Alk. CT SQ_2 Na ⁺ K ⁺ Mg ²⁺ Ca ²⁺	3,538,660	3,549,933	3,549,933	3,544,999.3	3,536,077	3,538,292.9	3,537,055	3,581,097	3,538,398	3,532,463	3,531,435	3,537,055	3,538,478	3,548,943	3,531,435	3,537,962	3,532,463	3,577,944.8	3,537,270	3,537,109.4	3,533,586	3,548,943	3,547,329.7	3,577,944.8	3,537,323.9	3,575,339	3,538,398	3,544,999.3	3,537,962	3,572,820.2	3,537,988	3,567,287.5	3,547,216	3.564.272	3,537,988		3,584,761.4		3,547,216	3,597,046	3,597,046		Lat
Date EC T pH Alk CT 804 Ne ² Ne ²	719,831	717,022	717,022	719,930.6	719,558	720,442.9	719,746	730,922	721,404	713,715	713,298	719,746	720,170	717,353	713,298	718,744	713,715	714,428.5	721,119	718,419.1	714,060	717,353	716,520.7	714,428.5	724,063.3	718,875	721,404	719,930.6	718,744	719,721.4	720,114	730,058.8	718,358	719,421	720,114		717,604.5		718,358	719,626	719,626	/m	Long.
Fig. 1. Pig. Mile Cit No.4 Nit	127	128	128	110	134	132	114	106	130	117	111	114	131	133	111	134	117	111	119	137	141.6	133	129	H	127	109	130	110	134	112	124	139	124	161	124		125		124	118	118		Elev.
T pH Alk. CI 80^2 Na^+ K† M_8^{2+} C_a^{2+} PC	31/01/2013	26/01/2013	1996	27/01/2013	21/01/2013	27/01/2013	28/01/2013	24/01/2013	31/01/2013	03/02/2013	03/02/2013	1996	21/01/2013	26/01/2013	1996	21/01/2013	1996	1996	31/01/2013	1996	1996	1996	27/01/2013	20/01/2013	1996	20/01/2013	1996	1996	1996	20/01/2013	28/01/2013	24/01/2013	1996	1996	1996	1996	20/01/2013	1996	27/01/2013	1996	20/01/2013		Date
pH Alk. Cl ² Na ² Na ² K ² Og. ² Ca ² hmmolL ⁻¹ hmmolL ⁻¹ p.18 1.56 12.02 7.3 13 0.99 4.34 2.8 8.86 1.46 12.0 687 11.57 0.93 4.4 2.9 8.115 1.86 12.0 687 11.57 0.93 4.43 3.43 7.52 1.36 14.15 17.39 12.80 0.61 10.61 7.5 8.09 2.04 14.15 17.39 12.80 0.61 10.61 7.5 8.13 1.46 18.91 7.78 12.60 0.03 5.75 5 7.12 2.93 2.21 2.93 2.41 2.24 2.60 2.84 2.93 2.51 3.47 8.23 4.22 2.26 8.84 2.21 2.21 4.25 0.83 0.24 7.75 2.33 2	6.26	5.98	7.62	6.22	6.08	6.09	5.51	5.5	5.29	5.8	5.5	6.08	5.72	5.7	4.77	4.65	5.62	4.49	4.67		2.62	5.3	3.66	4.1	4.7	4.08	5.51	2.43	2.47	3.44	3.7	4.5	2.77	2.96	2.51	4.08	2.93	4.01	2.1	2	2.09	/mS cm ⁻¹	EC
Alk. CIT SO ₄ ² Na [±] K [±] Mg ^{2±} Ca ^{2±} /mmolL-1 1.56 12.02 7.3 13 0.99 4.34 2.8 1.86 12.02 7.3 13 0.99 4.44 2.9 1.86 13.2 6.87 11.57 0.93 4.4 2.9 1.86 14.15 17.89 15.89 0.61 10.61 7.5 2.24 11.77 9.4 16.6 0.93 5.75 5 5.25 18.21 9.97 24.29 0.41 1.43 3.43 1.26 18.91 7.74 12.44 2.60 0.62 2.21 2.3 1.27 22.12 22.3 3.41 2.29 8.61 3.47 4.22 22.2 22.3 13.34 12.84 2.21 4.01 3.17 4.22 22.3 13.94 13.22 24.26 0.88 0.23 <	24.1	24.6	23.65	23.1	24.5	25.4	24.6	23.7	25.1	22.5	25	23.71	22.9	26.2	23.70	23.3	23.69	23.67	22.2	23.87	23.68	23.44	24.6	25.2	23.61	24.2	23.80	23.46	23.47	27.5	22.2	27.5	23.45	23,43	23.83	23.73	23	23.79	23.9	22.1	22.7	/°C	Т
CIT SO ₄ ² Na [±] K [±] Mg ^{2±} Ca ^{2±} /mmolL-1 /mmolL-1 /mmolL-1 5 12.02 7.3 13 0.99 4.34 2.8 6 12.2 6.87 11.57 0.93 4.4 2.9 5 13.2 6.87 12.50 0.61 10.61 7.5 18.21 9.9 24.26 0.093 5.75 5 18.21 9.9 24.26 0.02 2.213 8.13 18.21 9.9 24.66 0.03 5.75 5 4 17.7 9.4 16.66 0.03 5.75 5 4 17.7 9.4 16.66 2.04 2.21 2.21 2.29 20.83 13.4 12.44 2.24 2.29 0.83 0.25 21.2 22.6 8.4 2.21 1.24 2.21 3.37 3.43 22.3 </th <th>7.68</th> <th>7.63</th> <th>7.93</th> <th>8.07</th> <th>8.13</th> <th>7.78</th> <th>8.44</th> <th>8.86</th> <th>7.84</th> <th>8.04</th> <th>7.72</th> <th>7.69</th> <th>8.21</th> <th>7.64</th> <th>7.70</th> <th>8.16</th> <th>7.62</th> <th>7.58</th> <th>7.89</th> <th>7.76</th> <th>7.65</th> <th>7.69</th> <th>8.1</th> <th>7.61</th> <th>7.22</th> <th>8.38</th> <th>7.39</th> <th>7.75</th> <th>7.72</th> <th>7.52</th> <th>8.23</th> <th>8.29</th> <th>7.83</th> <th>7.88</th> <th>8.08</th> <th>7.12</th> <th>8.09</th> <th>7.52</th> <th>8.15</th> <th>8.86</th> <th>9.18</th> <th></th> <th>pH</th>	7.68	7.63	7.93	8.07	8.13	7.78	8.44	8.86	7.84	8.04	7.72	7.69	8.21	7.64	7.70	8.16	7.62	7.58	7.89	7.76	7.65	7.69	8.1	7.61	7.22	8.38	7.39	7.75	7.72	7.52	8.23	8.29	7.83	7.88	8.08	7.12	8.09	7.52	8.15	8.86	9.18		pH
Na ⁺ K ⁺ Mg ²⁺ C ₃ 2 ⁺	2.11	2.16	0.56	1.82	1.82	1.62	2.37	0.35	4.1	1.66	3.25	1.32	1.96	2.48	1.55	1.78	1.45	1.5	1.78	1.75	2.84	1.34	ω	3.03	2.02	4.39	3.01	2.39	2.99	2.21	4.22	3.29	2.27	1.27	1.46	5.25	2.04	1.86	1.86	1.46	1.56		Alk.
Mat K+ Mg2+ Ca2+ //mmoll1 13	44.9	42.5	42.14	42	39.8	39.7	38.8	38.6	38.4	36.3	35.4	35.01	33.6	33.5	32.81	32.4	31.94	31.52	31.2	30.87	28.77	28.21	27.7	26.2	25.68	25.3	24.32	24.16	23.54	23.3	22.6	22.1	20.83	18.98	18.91	18.21	17.7	14.15	13	12	12.02		CI-
mmolL ⁻¹ Wg ²⁺ C _u ²⁺ 0.99 4.34 2.8 0.93 4.43 2.9 0.93 4.43 3.43 3.64 0.61 0.62 2.69 2.79 2.69 2.79	13.2	17.9	10.72	19.1	11.8	11.7	16.9	18	14.6	11.6	13.8	13.52	12.1	11.9	12.85	14.6	12.83	10.08	15.4	16.66	14.52	11.48	10.6	9.8	10.36	9.5	21.22	13.23	13.97	13.4	8.6	12.4	9.366	7.74	7.8	9.97	9.4	17.89	7.3	6.87	7.3		SO ₄ ²⁻
K+ Mg ²⁺ C _a ²⁺ 0.99 1.25 1.43 0.61 0.61 0.62 1.35 0.62 2.13 2.99 2.69 2.43 2.53 0.61 1.86 8.25 0.81 1.86 2.21 2.91 2.91 2.91 2.91 3.17 2.21 4.91 3.17 2.21 4.91 3.17 2.21 4.91 3.17 2.21 4.93 7.91 2.22 9.39 6.38 7.36 6.38 7.36 6.38 7.36 6.38 7.36 6.38 7.36 6.38 7.36 7.36 7.36 7.37 7.33 7.36 7.37 7.37 7.37 7.38 7.39 7.37 7.39 7.30	36.2	32.1	18.87	27.5	30.6	36	36.9	22.3	28.5	28.5	37.1	8.6	29.2	27.7	30.18	27.8	22.23	20.05	21.3	24.9	58.74	17.58	19	24	14.83	23.7	24.26	41.89	50.56	21.8	28.4	21.8	34.17	12.41	26.05	24.29	16.6	15.89	12.6	11.57	13	/mmo	Na+
C _A 2+ 2.8 2.9 3.43 3.53 8.13 8.13 8.13 8.13 9.29 9.31 7.36 6.28 0.28 0.24 0.24 7.40 6.28 0.25 0.26 6.28 0.27 6.28 0.28 1.10 1.20	11.8	8.03	1.86	13.21	5.2	8.43	1.93	0.89	4.45	3.21	3.04	1.92	3.35	5.93	0.96	0.8	0.8	5.87	3.87	0.97	0.03	2.03	2.29	2.32	0.24	1.77	0.88	6.08	2.82	1.86	2.21	2.59	4.25	2.69	0.62	0.41	0.93	0.61	1.25	0.93	0.99	IL-1	K+
	6.32	12.49	12.63	13.39	7.14	5.11	9.03	4.8	11.62	6.75	8.45	19.37	6.36	5.98	9.19	6.76	10.55	7.53	11.17	15.69	0.83	11.48	9.09	4.96	9.33	4.18	20.16	2.34	0.98	8.25	4.01	8.61	1.35	5.32	2.13	1.43	5.75	10.61	4.43	4.4	4.34		Mg ²⁺
0.024 0.032 0.033 0.033	6.68	8.07	9.32	8.12	8.46	5.97	9.21	21.26	8.14	8.37	5.68	7.23	8.17	7.57	5.74	10.83	7.89	6.5	8.37	4.49	0.73	5.8	6.55	7.46	7.36	7.91	2.23	0.84	0.25	6.28	3.17	5.47	0.86	5.31	2.99	8.13	S	7.5	3.43	2.9	2.8		Ca ²⁺
I I																							0.033	0.033		0.025				0.032							0.031		0.024				Br-

Table 4: Field and analytical data for the Phreatic aquifer (continued).

										4			0	
		m/			$/\mathrm{mS}\mathrm{cm}^{-1}$)°C					$/\text{mmolL}^{-1}$			
P419	3,539,494	725,605	132	31/01/2013	7.03	26.4	7.84	2.05	45.1	14.4	41.4	10.78	5.95	6.91
PL13	3,536,550	720,200	123	21/01/2013	7.22	24.5	7.51	3.24	87.8	14.5	4.4	10.55	6.35	6.59
PL25	3,536,230	718,708	129	21/01/2013	7.64	27.1	3 .	1.78	84	14.5	42.9	6.56	7.4	7.61
P506	3,535,528.1	725,075.1	126	04/02/2013	8.32	24.3	8.12	1.71	52.6	14.6	42.8	10.97	7.51	7.83
P506	3,535,528.1	725,075.1	126	1996	6.7	23.28	7.46	1.8	54.39	17.58	33.32	4.11	22.16	5.17
P566	3,540,433.1	719,661.3	115	27/01/2013	6	24.6	7.64	1.72	62.5	15.2	71.6	3.03	4.61	90'9
PL17	3,536,908	718,511	130	21/01/2013	9.4	24.5	8.06	3.39	63.2	15.6	77.2	2.51	4.08	5.11
P413	3,530,116.2	722,775.1	130	04/02/2013	10.09	30.2	7.91	1.63	63.6	21.5	88.3	4.08	4.21	4.65
PL25	3.536.230	718.708	129	1996	9.5	23.72	7.96	0.63	75.57	10.62	10.22	2.64	32.94	9.54
P066	3,542,636,5	718,957.4	126	1996	7.75	23.48	7.62	1.51	80.23	12.45	45.87	2.46	23.59	5.91
PI 03	3.540.010.9	725.738.1	130	9661	7.34	23.86	7.60	3.04	84.14	30.58	108.55	2.23	10.17	8.99
PI 21	3.536.074	721.268	128	9661	9.73	23.82	7.25	4.46	84.26	23.68	61.62	3.75	33.53	88
PI 23	3 538 419	720.950	126	28/01/2013		24.2	, « , c	4.53	86.6	16.7	79.0	3.72	14.54	589
PI 21	3 536 074	721.268	2 2 8	28/01/2013	16.41	75.7	7.45	1 97	6 66	17.4	85.5	5.7	15.66	76
PI 05	3 537 109 4	718 419 1	13.7	21/01/2013	891	24.8	25	200	1013	17.7	85.9	282	16.69	7.59
PI 44	3 536 039 3	721 673 9	134	9661	4 68	23.85	7 19	2.74	109.75	17 21	134.67	5.71	20 07	8.77
NS Id		719.416	12.5	22/01/2013	17.08	24.9	×	3.41	1142	181	6.06	12.8	16.85	781
PI 03		725 738 1	130	27/01/2013	10.84	23.1	2	3.29	1173	14.7	116.4	2.06	8 99	7.24
P006	3.564.272	719,421	191	03/02/2013	18.31	23.6	7.76	6.26	131.9	182	96.3	8.61	27.11	7.99
P015	3,551,711	720,591	103	1996	12.42	23.62	7.71	2.38	134.68	28.2	72.98	3.1	52.44	6.25
PL04	3,541,410.1	723,501.1	138	27/01/2013	19.01	26.4	7.85	4.03	138	16.7	108.8	13.06	19.51	8.72
D.Ch				1996		23.88	79.7	2.68	142.22	24.5	96.31	3.16	44.22	3.02
PL44	3,536,039.3	721,673.9	134	28/01/2013	20.18	25.8	7.8	4.96	153	17.7	125.9	6.29	22.83	808
PL27	3,535,474	718,407	126	21/01/2013	21.23	24.8	8.11	1.7	169.4	18.4	130.3	4.89	27.81	8.63
P076	3,540,137	716,721	118	26/01/2013	22.31	27.2	7.57	4.33	171.5	17.1	130.8	6.32	28.01	8.83
P041	3,559,563	716,543	135	26/01/2013	25.94	24.5	8.18	7.95	208.6	13.4	198.9	3.61	11.81	8.75
P009	3,559,388	717,707	123	26/01/2013	27.51	28.4	8.39	11.45	208.8	15.8	195.1	2.65	18.7	9.01
LTP16				1996	11.53	23.78	7.48	3.84	213.35	48.63	147.9	7.46	75.31	4.25
P100				1996	17.18	23.64	7.59	3.37	235.01	46.44	264.84	4.74	25.57	5.56
PLX1	3,540,758.8	726,115.6	132	28/01/2013	32.93	23.4	7.95	4.44	245.6	20.9	141.4	26.88	44.56	17.66
P003	3,569,043	721,496	134	02/02/2013	31.03	23.5	8.01	6.91	252.7	17.9	208.2	9.41	29.99	10.03
P007		718,651	129	26/01/2013	30.07	28.4	7.76	5.42	254.7	15.5	209.2	10.43	28.82	7.51
PLX2	3,537,323.9	724,063.3	127	21/01/2013	43.25	25.7	8.07	5.15	262.2	93	270.4	15.5	62.77	21.46
P015	3,551,711	720,591	103	25/01/2013	32.02	7.22	8.03	2.95	263	15.4	206.9	92'9	32.12	9.95
P402	3,549,503	721,514	138	25/01/2013	09	28.7	9.8	7.69	313.2	93.9	442.8	23.26	12.56	10.17
P001	3,572,148	722,366	127	1996		23.63	8.37	4	323.62	58.13	331.43	5.01	49.77	3.97
P014	3,551,466	719,339	131	1996		23.40	7.31	3.98	336.96	64.29	328.67	5.53	62.37	5.45
P019	3,562,960	717,719	113	02/02/2013	60.58	27.8	7.65	6.02	356.2	96	432.5	29.77	21.02	26.23
P018	3,562,122	716,590	110	26/01/2013	90'19	26.2	8.42	6.46	372.4	82.3	347.1	22.64	60.71	26.63
P014	3,551,466	719,339	131	25/01/2013	49.04	25.2	7.89	1.8	399.7	21.1	389.3	2.41	18.97	7.39
P113	3,535,586	714.576	105	03/02/2013	62.24	24.8	8.2	5.96	414.8	83.8	362.7	33.34	70.23	26.51

Table 5: Field and analytical data for the Phreatic aquifer (continued).

Sebkhet Safioune	Sebkhet Safioune	Sebkhet Safioune		Sebkhet Safioune	N'Goussa	Hassi Debich	Oum Raneb					Sebkhet Safioune	El Bour-N'gouca	Bamendil		Sebkhet Safioune	Chott Adiadia	Sebkhet Safioune	Sebkhet Safioune	Sebkhet Safioune	Hôtel Transat	Oum Raneb	Route Sedrata	Oum Raneb	N'Goussa	ANK Djemel	Said Otba(Bab sbaa)	N'Goussa	Said Otba-Chott	Route Frane	Station de pompage	ANK Djemel	N'Goussa	Sebkhet Safioune	Said Otba-Chott	ANK Djemel	Oum Raneb	Oum Raneb	Sebkhet Safioune	Route Frane	Sebkhet Safioune	Sebkhet Safioune	Route Frane		Locality
P036	P037	P074	P039	P034	P041	P416	PZ12	P068	P042	P093	P044	P063	P007	P076	LTP06	P063	PLXI	P023	P034	P023	PL 23	PZ 12	P113	P162	P018) P066	P019	P096	P002	PL04	P021	P017	P030	P096	P423	P012	P012	P030	P002	P031	P031	P001		Site
				3,579,698	3,559,563	3,581,097	3,547,234					3,545,586.8	3,562,236	3,540,137	1	3.545.586.8	3.540,758.8	3.577.198	3,579,698	3,577,198	3,538,419	3,547,234	3,535,586	3,546,133	3,562,122	3,573,943	3,542,636.5	3,562,960	3,540,265	3,570,523	3,541,410.1	3,573,943	3,560,256	3,577,253	3,540,265	3,540,881	3,554,089	3,554,089	3,577,253	3,570,523	3,577,804	3,577,804	3,572,148		Lat.
				725,633	716,543	730,922	722,931					725,667.4	718,651	716,721		725.667.4	726,115.6	725,726	725,633	725,726	720,950	722,931	714,576	725,129	716,590	723,161	718,957.4	717,719	724,729	722,028	723,501.1	723,161	715,781	721,936	724,729	723,178	718,612	718,612	721,936	722,028	720,172	720,172	722,366	/m	Long.
				97	135	106	110					99	129	118		99	132	99	97	99	126	110	105	98	110	105	126	113	Ξ	108	138	105	130	130	Ξ	102	114	114	130	108	120	120	127		Elev.
1996	1996	1996	1996	1996	1996	1996	1996	1996	1996	1996	1996	05/02/2013	1996	1996	1996	1996	1996	05/02/2013	05/02/2013	1996	1996	05/02/2013	1996	25/01/2013	1996	24/01/2013	03/02/2013	1996	03/02/2013	02/02/2013	1996	1996	26/01/2013	03/02/2013	1996	31/01/2013	1996	25/01/2013	1996	1996	02/02/2013	1996	02/02/2013		Date
												178.9						117.9	130			114.9		160		82.28	150.6		68.31	62.82			100.1	64.66		90.8		64.05			75.96		66.16	/mS cm ⁻¹	EC
23.35	23.36	23.54	23.37	23.34	23.38	23.33	23.31	23.51	23.42	23.58	23.39	26.7	23.26	23.53	23.77	23.50	23.60	29.4	34.9	23.32	23.49	27.4	23.66	30.7	23.29	29.6	26.2	23.30	25.9	26.9	23.57	23.55	31	23.1	23.59	23.5	23.41	30.3	23.52	23.81	27.9	23.75	28.3	/°C	7
7.54	6.92	6.47	6.87	7.85	7.94	7.84	7.59	7.54	7.59	7.49	7.79	7.67	7.67	7.71	7.64	7.45	8.02	8.19	8.08	7.42	7.37	7.44	7.70	7.15	7.46	7.64	7.18	7.72	8.7	7.57	7.42	7.43	7.13	7.83	7.71	7.48	7.46	7.83	7.72	7.76	8.06	7.31	7.24		рH
1.4	1.52	4.17	1.94	1.95	2.13	4.33	2.21	3.35	1.1	1.49	4.53	1.43	1.41	5.72	7.84	1.94	3.82	1.85	1.76	2.25	3	2.88	2.81	2.43	1.24	2.35	12.29	2.42	1.24	1.65	2.37	4.24	3.78	3.71	3.69	6.19	2.72	7.77	4.43	6.29	5.85	6.32	6.49		Alk.
4,972.75	4,953.84	4,356.48	4,189.51	2,752	2,599.74	2,433.73	2,405.55	2,335.67	2,330.85	2,198.58	2,106.07	1,887.9	1,860.53	1,743.55	1.638.66	1.379.35	1,296,65	1.209.3	1,189.1	1,176.99	1,103.31	980.1	954.89	842.8	818.67	800.4	799.1	779.13	771	748.5	716.27	700.77	679.3	671.8	645.07	636.5	539.35	534.3	527.7	522.39	500.3	481.83	468.7		CI-
108.12	184.54	180.88	201.44	134.14	324.58	178.87	109.92	222.08	101.22	182.08	18.27	92.9	91.55	143.36	712.09	139.61	134.01	15.6	14.7	91.14	94.49	15.5	124.85	289.9	81	94.4	283	77.13	53.1	62.6	34.75	154.45	114.1	90.3	78.46	101.3	60.64	20.9	123.48	182.95	110.3	43.35	101.5		so ₄ ²⁻
4,692.23	4,611.06	2,759.9	4,042.62	2,616.77	2,878.99	2,361.09	2,178.55	2,302.25	1,963.71	1,957.53	1,765.47	1,455.8	1,434.73	1,321.87	2.621.61	1.257.42	1,458.73	1.129.4	1,055.1	1,058.21	707.81	930.8	997.52	1,309.9	244.21	824	1,249.7	711.46	615.9	651.5	560.07	605.68	597.8	742.9	357.28	495.5	413.55	529.6	533.79	653.78	470.5	326.82	350.3	/mmol/L	Na+
36.84	2.9	57.4	17.9	24.42	44.57	24.34	25.23	26.84	52.19	29.49	27.33	26.66	26.2	26.85	41.55	18.6	5.24	8.38	18.27	11.72	19.14	7.53	13.3	13.3	49.54	10.99	18.95	9.23	23.46	14.72	7.04	53.6	10.71	15.97	5.89	38.31	5.55	6.41	11.59	9.97	28.67	12.61	25.96		K+
221.13	347.57	930.06	257.81	180.14	152.83	196.07	199.35	219.9	248.1	278.18	171.23	282.88	278.77	331.38	190.51	182.26	47.98	42.85	56.37	133.47	270.91	23.9	86.69	33.47	319.35	53.35	37.63	95.59	69.64	77.72	99.58	163.08	125.85	41.46	208.4	125.81	112.77	19.73	106.21	104.7	79.12	94.15	116.21		Mg ²⁺
9.63	7.86	22.63	9.23	10.48	10.97	9.2	12.65	7.19	11.24	10.44	6.54	13.44	13.25	12.26	13.34	10.03	4.34	10.15	17.38	12.41	13.3	14.24	11.67	17.74	24.76	25.39	18.06	12.05	50.39	27.29	11.04	14.24	26.29	7.65	12.86	30.32	9.42	4.73	10.65	10.99	35.47	23.56	35.31		Ca ²⁺

Table 6: Isotopic data $^{18}\mathrm{O}$ and $^{3}\mathrm{H}$ and chloride concentration in Continental Intercalaire, Complexe Terminal and Phreatic aquifers (sampling campaign in 1992).

					Phreati	c aquifer					
Piezometer	Cl ⁻ /mmol L ⁻¹	δ ¹⁸ O /‰	3 _H /UT	Piezometer	Cl ⁻ /mmol L ⁻¹	δ ¹⁸ O /‰	³ H /UT	Piezometer	Cl ⁻ /mmol L ⁻¹	δ ¹⁸ O /‰	³ H /UT
P007	1,860.5	-2.49	0	PL15	23.54	-7.85	0.6(1)	P074	4,356.4	3.42	6.8(8
P009	426.85	-6.6	1.2(3)	P066	80.23	-8.14	0.8(1)	PL06	14.15	-8.13	1.0(2
P506	54.39	-6.83	1.6(3)	PL23	1,103.32	-6.1	0	PL30	24.32	-7.48	2.4(4
P018	818.67	-2.95	6.2(11)	P063	1,379.3	-3.4	8.7(15)	P002	522.39	-5.71	0.6(1
P019	779.13	-4.67	5.6(9)	P068	2,335.6	-3.04	8.8(14)	PL21	84.26	-7.65	1.2(2
PZ12	2,405.5	-2.31	8.1(13)	P030	527.7	-6.57	2.4(4)	PL31	18.91	-7.38	1.6(3
P023	1,176.9	-2.62	0.2(1)	P076	1,743.5	-5.56	2.8(5)	P433	12	-8.84	0
P416	2,433.7	-7.88	5.9(9)	P021	700.7	-5.16	2.6(4)	PL03	84.14	-7.35	1.7(3
P034	2,752	-1.77	5.7(9)	PL04	716.27	-2.89		PL44	109.75	-8.82	1.0(2
P036	4,972.7	3.33	2.1(4)	P093	2,198.5	-2.64	5.1(8)	PL05	30.87	-7.44	1.9(3
P037	4,953.8	3.12	1.8(3)	P096	645.07	-6.13	4.8(8)	P408	24.16	-7.92	0
P039	4,189.5	0.97	2.2(4)	PLX1	1,296.6	-5.6	1.1(2)	P116	31.94	-7.18	1.1(2
P041	2,599.7	-0.58	7.3(13)	PLX2	25.68	-7.6	1.3(2)	LTP 16	213.35	-7.48	1.6(3
P044	2,106.1	-4.46	2.7(5)	P015	134.68	-6.77	3.0(5)	P117	32.81	-6.92	0.1
P014	336.96	-6.9	2.8(5)	P001	323.62	-4.66	2.5(4)	PL10	35.01	-7.31	0.2(1
P012	539.3	-6.41	2.2(4)	P100	235.01	-5.81	0	PL25	75.57	-7.41	0.9(2
P042	2,330.8	2.05	6.0(10)	P056	42.14	-7.03	2.9(5)	LTP30	18.21	-7.5	1.1(2
P006	18.98	-6.64	0.5(1)	P113	954.89	-4.75	0.8(2)	LTP06	1,638.6	-1.97	2.8(5
P057	28.21	-7.33	1.1(2)	PLX4	31.52	-7.1	0.3(1)	P031	481.83	-6.06	3.0(5
P059	20.83	-7.81	0	P115	28.77	-2.54	6.8(12)				
					Complexe Te	rminal aquifer					
Borehole	Cl ⁻	$\delta^{18}O$	3 _H	Borehole	Cl ⁻	δ^{18} O	3 _H	Borehole	Cl ⁻	δ^{18} O	3 _H
	$/\text{mmol L}^{-1}$	/‰	/UT		$/\text{mmol L}^{-1}$	/‰	/UT		$/\text{mmol L}^{-1}$	/‰	/UT
D5F80	42.22	-7.85		D1F138	28.92	-8.13	0.7(1)	D2F71	13.53	-8.23	0.6(1
D3F8	29.81	-8.14	1.4(2)	D3F18	21.66	-8.23	0.2(1)	D7F4	10.6	-8.27	0.1(1
D3F26	34.68	-7.97	0.8(1)	D3F10	14.27	-7.88	1.5(2)	D2F66	11.02	-8.3	
D4F94	20.05	-8.18	0.6(1)	D6F51	28.39	-7.9	0.7(1)	D1F151	10.75	-8.32	0.4(
D6F67	18.79	-8.23	3.7(6)	D1F135	18.08	-7.97	1.1(2)	D6F64	11.36	-8.28	4.3(
					Continental Int	ercalaire aquife	er				
Borehole	Cl ⁻	$\delta^{18}O$	3 _H	Borehole	Cl ⁻	$\delta^{18}O$	$^{3}\mathrm{H}$	Borehole	Cl ⁻	$\delta^{18}O$	$^{3}\mathrm{H}$
	$/\text{mmol } L^{-1}$	/‰	/UT		$/\text{mmol L}^{-1}$	/‰	/UT		$/\text{mmol L}^{-1}$	/‰	/UT
Hadeb I	5.8	-8.02	0	Hadeb II	6.19	-7.93	0.1(1)	Aouinet Moussa	6.49	-7.88	1.1(

Table 7: Statistical parameters for Continental Intercalaire (CI), Complexe Terminal (CT) and Phreatic (Phr) aquifers samples selected on the basis of δ^{18} O and Cl⁻ data (see text).

Aquifer	Size	Parameter	EC /mS cm ⁻¹	T /°C	pН	Alk.	Cl ⁻	so ₄ ²⁻	Na ⁺ /mmol/L	K ⁺	Mg ²⁺	Ca ²⁺
CI	11	Average	2.2	49.0	7.5	2.3	11.0	4.7	10.3	0.51	3.6	2.4
CI	11	Stdd. dev.	0.3	2.0	0.2	1.0	4.6	2.5	4.6	0.23	2.0	1.8
CT	50	Average	3.2	23.0	7.8	2.3	20.0	8.9	17.0	1.0	5.5	5.6
CT	50	Stdd. dev.	1.1	2.4	0.4	0.8	7.0	2.6	6.0	0.8	2.2	1.7
Phr pole I	30	Average	3.9	24.0	7.9	2.3	24.7	11.8	24.2	2.1	7.2	5.3
Phr pole I	30	Stdd. dev.	1.3	1.3	0.4	1.0	6.9	3.4	11.0	1.7	5.0	2.7
Phr pole II Phr pole II	3 3	Average Stdd. dev.		23.4 0.1	7.0 0.5	2.4 1.6	4,761.0 350.0	158.0 43.0	4,021.0 1,093.0	32.4 28.0	500.0 378.0	13.0 8.0

Table 8: Summary of mass transfer for geochemical inverse modeling. Phases and thermodynamic database are from Phreeqc 3.0 (Parkhurst and Appelo, 2013).

Phases	Stoichiometry	CI/CT	CT/Phr I	Rainwater/P036	PhrI/PhrII 60%/40%
Calcite	CaCO ₃	_	-6.62×10^{-6}	-1.88×10^{-1}	-2.26×10^{-1}
$CO_2(g)$	CO ₂	-6.88×10^{-5}	_	8.42×10^{-4}	5.77×10^{-4}
Gypsum	$CaSO_4 \cdot 2H_2O$	4.33×10^{-3}	_	1.55×10^{-1}	1.67×10^{-1}
Halite	NaCl	7.05×10^{-3}	3.76×10^{-3}	6.72	1.28
Sylvite	KCl	2.18×10^{-3}	1.08×10^{-3}	4.02×10^{-1}	_
Bloedite	$Na_2Mg(SO_4)_2 \cdot 4H_2O$	_	1.44×10^{-3}	_	_
Huntite	$CaMg_3(CO_3)_4$	_	_	4.74×10^{-2}	5.65×10^{-2}
Ca ion exchange	CaX ₂	-1.11×10^{-3}	_	_	_
Mg ion exchange	MgX_2	1.96×10^{-3}	_	1.75×10^{-1}	-2.02×10^{-1}
Na ion exchange	NaX	_	_	_	3.92×10^{-1}
K ion exchange	KX	-1.69×10^{-3}	-	-3.49×10^{-1}	1.20×10^{-2}

Values are in mol/kg (H_2O) . Positive (mass entering solution) and negative (mass leaving solution) phase mole transfers indicate dissolution and precipitation, respectively; — indicates no mass transfer.

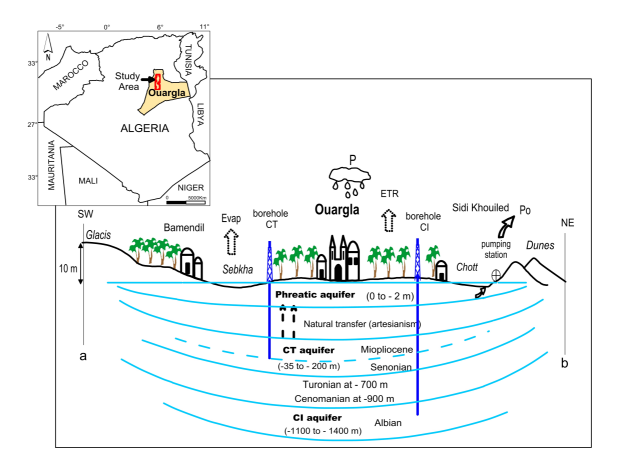


Figure 1: Localisation and schematic relations of aquifers in Ouargla. Blue lines represent limits between aquifers, and the names of aquifers are given in bold letters; as the limit between Senonian and Miopliocene aquifers is not well defined, a dashed blue line is used. Names of villages and cities are given in roman (Bamendil, Ouargla, Sidi Khouiled), while geological/geomorphological features are in italic (Glacis, Sebkha, Chott, Dunes). Depths are relative to the ground surface. Letters a and b refer to the cross section (fig. 2) and to the localisation map (fig. 3).

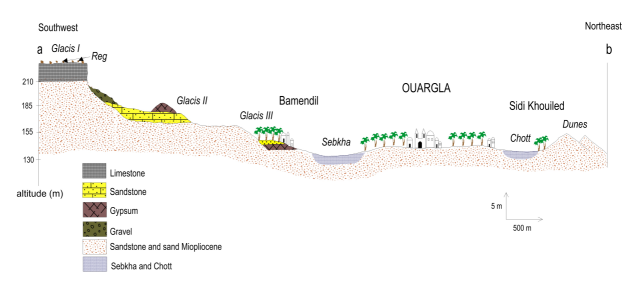


Figure 2: Geologic cross section in the region of Ouargla.

The blue pattern used for Chott and Sebkha correspond to the limit of the saturated zone.

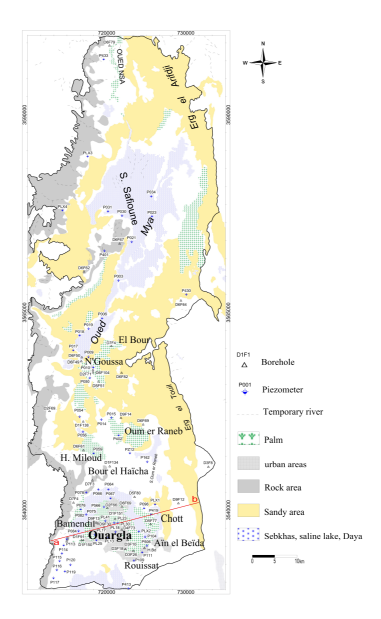


Figure 3: Localisation map of sampling point

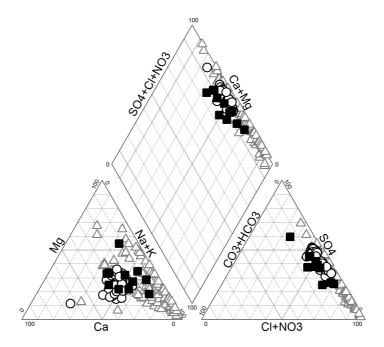


Figure 4: Piper diagram for Continental Intercalaire (filled squares), Complexe Terminal (open circles) and Phreatic aquifer (open triangles).

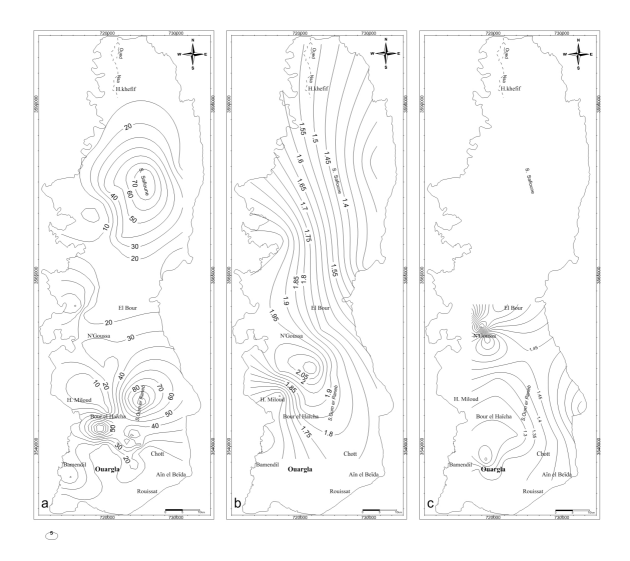


Figure 5: Contour maps of the salinity (expressed as global mineralization) in the aquifer system, (a) Phreatic aquifer; (b) and (c) Complexe Terminal [(b) Miopliocene and (c) Senonian]; figures are isovalues of global mineralization (values in g/L).

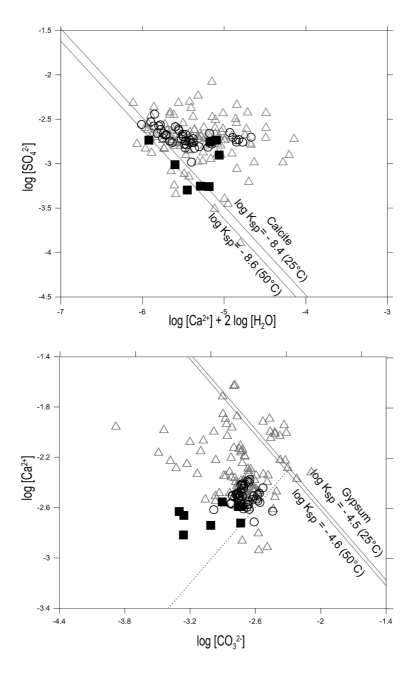


Figure 6: Equilibrium diagrams of calcite (top) and gypsum (bottom) for Continental Intercalaire (filled squares), Complexe Terminal (open circles) and Phreatic aquifer (open triangles). Equilibrium lines are defined as: $\log\{\mathrm{Ca^{2+}}\} + \log\{\mathrm{CO_3^{2-}}\} = \log K_{sp}$ for calcite, and $\log\{\mathrm{Ca^{2+}}\} + 2\log\{\mathrm{H_2O}\} + \log\{\mathrm{SO_4^{2-}}\} = \log K_{sp}$ for gypsum.

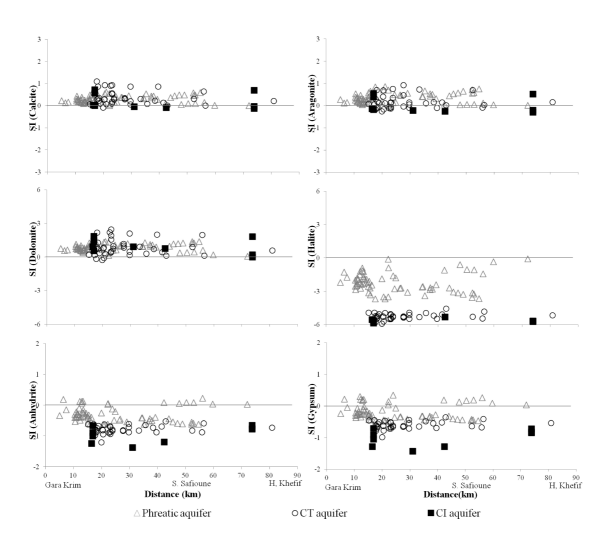
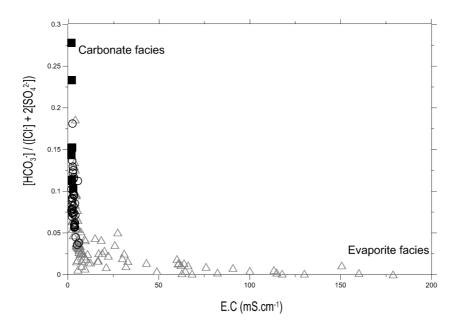
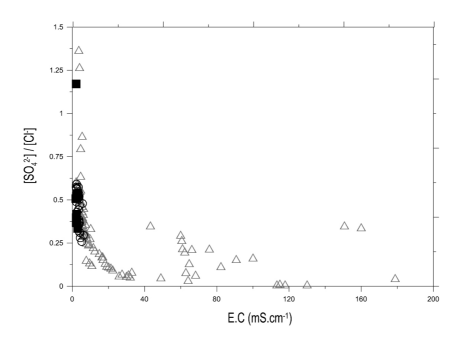


Figure 7: Variation of saturation indices with distance from south to north in the region of Ouargla.



<u>Figure 8:</u> Change from carbonate facies to evaporite from Continental Intercalaire (filled squares), Complexe Terminal (open circles) and Phreatic aquifer (open triangles).



<u>Figure 9:</u> Change from sulfate facies to chloride from Continental Intercalaire (filled squares), Complexe Terminal (open circles) and Phreatic aquifer (open triangles).

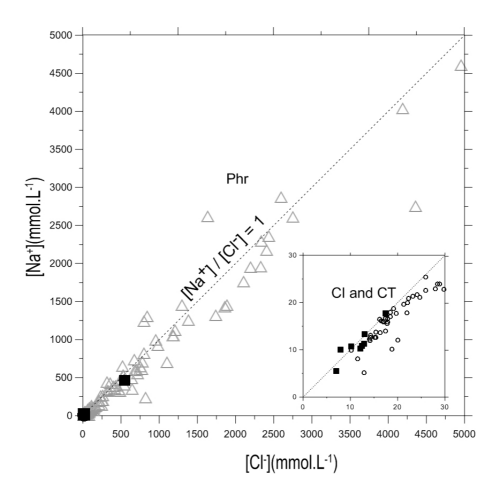


Figure 10: Correlation between Na^+ and Cl^- concentrations in Continental Intercalaire (filled squares), Complexe Terminal (open circles) and Phreatic aquifer (open triangles). Seawater composition (star) is $[Na^+] = 459.3 \text{ mmol } L^{-1}$ and $[Cl^-] = 535.3 \text{ mmol } L^{-1}$ (Stumm and Morgan, 1999, p.899).

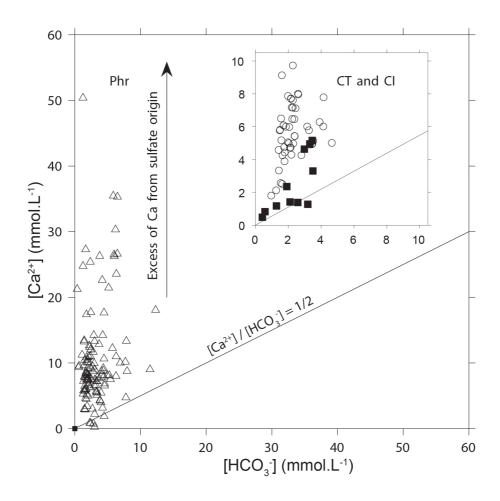


Figure 11: Calcium vs. HCO_3^- diagram in Continental Intercalaire (filled squares), Complexe Terminal (open circles), Phreatic aquifer (open triangles) and Seawater composition (star) is $[Ca^{2+}] = 10.2 \, \text{mmol} \, L^{-1}$ and $[HCO_3^-] = 2.38 \, \text{mmol} \, L^{-1}$ (Stumm and Morgan, 1999, p.899).

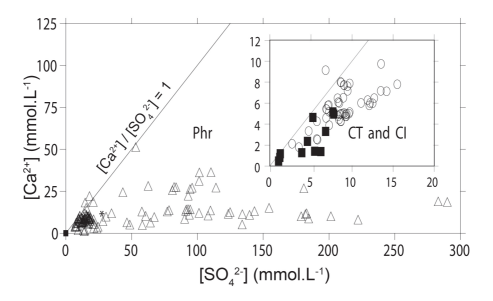


Figure 12: Calcium vs. SO_4^{2-} diagram in Continental Intercalaire (filled squares), Complexe Terminal (open circles), Phreatic aquifer (open triangles) and Seawater composition (star) is $[Ca^{2+}] = 10.2 \, \text{mmol} \, L^{-1}$ and $[SO_4^{2-}] = 28.2 \, \text{mmol} \, L^{-1}$ (Stumm and Morgan, 1999, p.899).

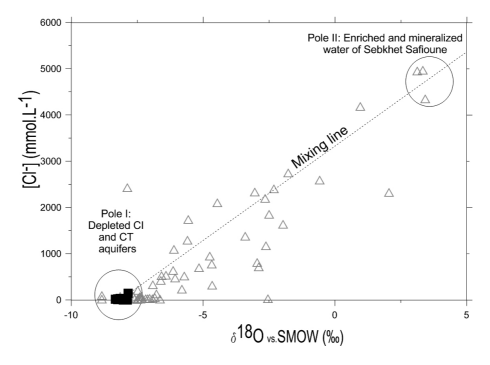


Figure 13: Chloride concentration versus δ^{18} O in Continental Intercalaire (filled squares), Complexe Terminal (open circles) and Phreatic aquifer (open triangles) from Ouargla.

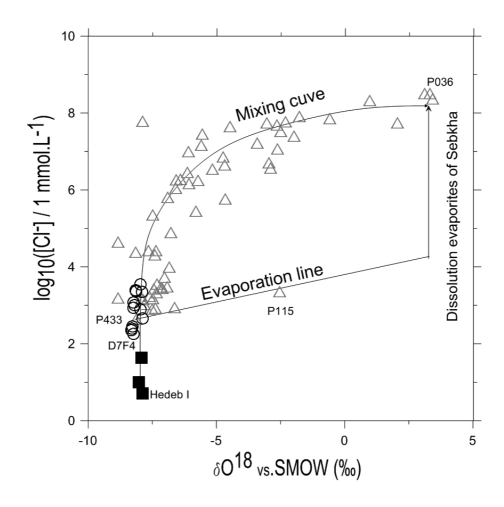


Figure 14: Log [Cl $^-$] concentration versus δ^{18} O in Continental Intercalaire (filled squares), Complexe Terminal (open circles) and Phreatic aquifer (open triangles) from Ouargla.