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Hydrochemical and isotopic characteristics of groundwater in the Continental Intercalaire aquifer system: Insights from Mzab Ridge and surrounding regions, North of the Algerian Sahara

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ABSTRACT

Study region: North of Sahara, Algeria.

Study focus: The effects of water/rock interactions and hydrodynamic conditions on the characteristics of groundwater in the Continental Intercalaire aquifer (CI) are demonstrated using hydrochemical and isotopic data from the Mzab Ridge.

New hydrological insights: The results confirm that CI groundwater in the oriental basin is highly mineralized (1010 > EC > 3500 $\mu S/cm)$ and is of the SO_4 -Cl-Na type. However, it is poorly mineralized (340 > EC > 960) in the occidental basin and predominantly belongs to the HCO $_3$ -Ca type. The important contribution of evaporite minerals has been deduced using saturation indices and minor element ratios of Sr/Ca and Br/Cl. This is explained by lithological heterogeneities within the CI aquifer and corroborates the observations from regional piezometric features which suggest a groundwater divide located between 31.55° and 31.57 °N latitude. $\delta^{18}\text{O-NO}_3$ and $\delta^{15}\text{N-NO}_3$ analyses show that NO $_3$ has a predominantly natural origin, which is soil nitrification even for the high concentrations exceeding 50 mg/L. CI groundwaters are highly depleted in both $\delta^{18}\text{O}$ and $\delta^2\text{H}$ compared to modern rainfall isotopic signatures. The most depleted ones are those situated in the Great Oriental Erg sub-basin. Isotopic values as low as $\delta^{18}\text{O}=-6\%$ can be taken as the limit for groundwater that is typically old. In the El Golea region, isotopic enrichment seems to indicate mixing between old groundwater and evaporation-affected groundwater infiltrating through the dunes of the Great Occidental Erg.

1. Introduction

The North-Western Sahara Aquifer System (NWSAS) is one of the largest transboundary aquifer (TBA) systems and the second in size in the world after the Nubian Sandstone (IGRAC, 2015). Shared between Algeria, Tunisia, and Libya, the NWSAS covers an area of more than 1 million km² (OSS, 2003). Important groundwater reserves of this multilayer aquifer, estimated to be 31 000 × 10⁹ m³ (Baba Sy, 2005), are divided into two aquifers: the lower Continental Intercalaire (CI) and the upper Complexe Terminal (CT) (Castany, 1982). Excessive groundwater mining from these aquifers, mainly for irrigated oasis agriculture, has led to a general piezometric head decline, as well as groundwater quality degradation (OSS, 2003). In many parts of the aquifer system, groundwater shows high

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salinities, above the quality guidelines for domestic or irrigation uses.

Because of its strategic importance for the remote semi-arid and arid areas of Algeria, Tunisia, and Libya, a transboundary cooperation through the Sahara and Sahel Observatory (OSS) was launched between these countries in the mid-90 s (Nijsten et al., 2018). Several scientific programmes have been developed to improve the understanding of the hydrologic functions of the NWSAS and provide sustainable management scenarios of this resource (Besbes et al., 2003; OSS, 2003; UNESCO, 1972).

One of the pivotal parameters to be defined for sustainability analysis is the estimation of the groundwater budget of the system, in particular, the recharge rates of the CT and CI aquifers. For the NWSAS, several approaches have been used to assess the recharge (including rate, localization, and mean residence time), such as hydrogeological modelling (Baba Sy, 2005; Baba Sy et al., 2007; Gonçalvès et al., 2020; Zammouri and Ribeiro, 2017), geophysical approaches based on satellite-based gravity data (Gonçalvès et al., 2013, 2020), and geochemical and isotopic approaches (Darling et al., 2018; Gonçalvès et al., 2015; Gonfiantini et al., 1974; Guendouz and Michelot, 2006; Matsumoto et al., 2020; Petersen et al., 2013, 2018; Trabelsi et al., 2009).

The identification of geochemical processes affecting groundwater chemistry and quality is also pivotal to the elaboration of a sustainable water resource management programme, particularly in areas where high salinities are observed. Several studies have focused on processes controlling the mineralization of groundwater, either of the CT or the CI aquifers, mainly in Algeria and Tunisia (Abdelali et al., 2020; Abid et al., 2009, 2010, 2012a, 2012b; Ben Alaya et al., 2014; Edmunds et al., 2003; Elliot et al., 2014; Guendouz et al., 2003; Hadj Ammar et al., 2014, 2020; Moulla and Guendouz, 2003; Moulla et al., 2012; Zammouri et al., 2007). Multiple studies have shown that the hydrogeochemical signatures (chemical and isotopic) and residence times of groundwater in these aquifer systems significantly differ (lithology, geometry and tectonic) (Edmunds et al., 2003; Guendouz, 1985; Guendouz and Michelot, 2006; Hadj Ammar et al., 2014; Moulla et al., 2012; Petersen et al., 2018).

Although all these studies cover a significant part of the NWSAS, the region that extends between the Great Occidental and Oriental Erg sub-basins (Fig. 1) has received much less attention than the south-western part of the Great Occidental Erg sub-basin or the Great Oriental Erg sub-basin. However, this region is important in the global functioning of these aquifers because it corresponds to the Mzab Ridge (see geomorphological units in Fig. 1), which marks the groundwater divide between these sub-basins (Baba Sy, 2005). In addition, the CI is the principal water resource in Ghardaia and El Golea regions (Achour, 2013), which during the last decades, has become a preferred area for intensive agricultural investments.

By combining geochemical and isotopic methods, the objective is to identify processes governing the hydrochemistry of CI groundwater and explain the spatial variations in groundwater quality in the Mzab Ridge and surrounding regions. Here, we present major elemental analyses to describe dominant geochemical processes, stable isotopic analyses (δ^{18} O and δ^{2} H) to discuss the origin of groundwater, and NO₃ stable isotope analyses to identify the sources of high nitrate concentration in CI groundwaters. For the sake of obtaining a complete overview of the CI groundwater characteristics in the north of Algerian Sahara, available data from Edmunds et al. (2003); Moulla et al. (2012) and Darling et al. (2018) were also combined with groundwater chemistry and isotopic data collected in the studied area. Finally, in the scope of our investigations, the implementation of a multivariate statistical analysis using the Self-Organizing Maps (SOMs) method is targeted (Kohonen, 1995; Peeters et al., 2007).

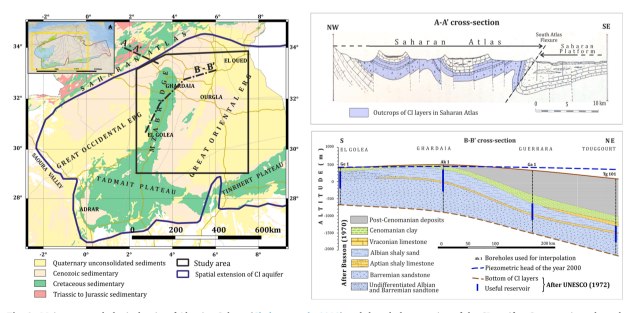


Fig. 1. Main geomorphological units of Algerian Sahara (Chabour et al., 2018) and the whole extension of the CI aquifer. Cross-sections show the geological characteristics in the study area. A-A' cross-section shows the tectonic transition between the Saharan Atlas and Saharan Platform (modified after Chettih, 1995). B-B' shows the variations of CI lithology and its depth and thickness across the Mzab Ridge.

2. Geographical, geological, and hydrogeological contexts of the study area

The study area is located between 2.8° and 6.1° E longitude and between 30.1° and 33.1° N latitude in the north of the Algerian Sahara between the towns of Ghardaia and El Golea (Fig. 1). It covers the Mzab Ridge and partially the basins located on either side, the Great Occidental and Oriental Ergs. Temperatures range between a minimum of -5° C in January and a maximum of 50° C in July. Annual rainfall is less than 65° mm, on average (FAO CLIMWAT, 2006), and the rainfall is very irregularly distributed between months and years.

From a geological point of view, the study area is located northwest of the "Saharan Platform" domain, which extends over the Algerian Sahara and corresponds to thick Phanerozoic sub-horizontal and quasi-monotonic sedimentary formations (1000–8000 m) (Askri et al., 1995; Fabre et al., 2005).

The "Continental Intercalaire" term was introduced for the first time by Kilian (1931). The meaning has evolved throughout history. Currently, it designates the Lower Cretaceous shaly-sand and sandstone formations (from Barremian to Albian) wrapped by clay formations of the Upper Jurassic below and Upper Cretaceous above. These clastic deposits are continuous over the whole basin (Figs. 1 and 2), from the Saharan Atlas in the north to the Tinrhert Plateau in the south and further to the Tassili mountains of the Hoggar and, west to east, from the Guir-Saoura valley in western Algeria to the Libyan desert (Edmunds et al., 2003; Moulla et al., 2012; OSS, 2003). It has a thickness of 350 m in average, varying from 0 m in its southern border to 1000 m under the Oriental Erg and towards Southern Tunisia (OSS, 2003).

The A-A' cross-section in Fig. 1 (modified from Chettih, 1995) shows that the "South Atlas Flexure" separates the CI layers outcropping in the Saharan Atlas from their equivalent in the Saharan Platform in the south where CI layers are buried at great depths (Askri et al., 1995). So, hydraulic continuity between these two parts of the CI aquifer is unexpected (OSS, 2003).

The B-B' cross-section in Fig. 1 is situated within the study area, which is a part of the Saharan Platform. It was obtained by a simple interpolation between the geological data given in Busson (1970) completed by geophysical logging (UNESCO, 1972) in the same boreholes. Piezometric heads for year 2000 were given in the SASS project (OSS, 2003). From a lithological point of view, heterogeneities can be observed within the CI aquifer. In the south (El Golea), it is mainly composed of detrital materials and sandstones sometimes cemented by carbonate minerals. However, it is clay-rich in the northwest (Ghardaia, Guerrara). Towards the northeast (Touggourt), the lithology progressively changes to limestone but, in the study area, limestone deposits are not so important compared to the rest of the Eastern Algerian Sahara (Busson, 1970). In the Great Oriental Erg sub-basin, Busson (1970) reported that the Albian layers contain gypsum and anhydrite in various forms (cement, fine crystallized intercalations etc.).

The first regional piezometric map was established during the ERESS project conducted by UNESCO in 1972, and then it was updated in 2000 as part of the SASS project accomplished by the OSS in 2003. The latter was added as background for most of our figures. It shows that the main recharge area of the CI aquifer is the Saharan Atlas in the Great Occidental Erg sub-basin (Baba Sy,

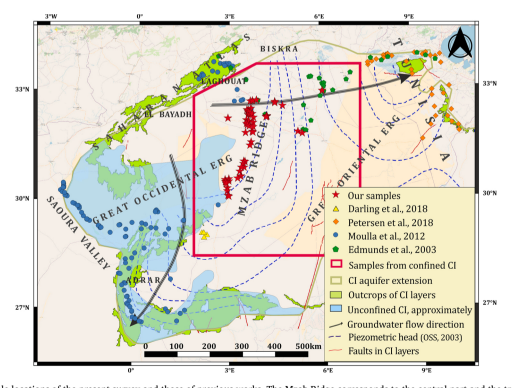


Fig. 2. Sample locations of the present survey and those of previous works. The Mzab Ridge corresponds to the central part and the transition zone between the Great Occidental and Oriental Erg sub-basins.

2005), which is the origin of two groundwater flow directions. In the Great Oriental Erg sub-basin, groundwater flows from the west to the east until Gabes Gulf in Tunisia and, in the Great Occidental Erg sub-basin, from north to south and southwest until Adrar region. Baba Sy (2005) showed that the CI aquifer is partially unconfined in the Great Occidental Erg sub-basin and shallow towards the southwest (Fig. 2), where it is one of the discharge areas. The CI aquifer is confined in the other areas of its extension over the Algerian Sahara and South Tunisia. The equipotential lines are almost orthogonal to the outcrops of CI layers in the Saharan Atlas in the north (Laghouat and Biskra regions), which means that there is no ongoing flux through the northern limit of the CI aquifer as discussed in the B-B' cross-section (Fig. 1).

Transmissivity values range from 10 to 40×10^{-3} m²/s in El Golea region south and from 3 to 5×10^{-3} m²/s in Ghardaia, Ouargla and El Oued regions north (UNESCO, 1972). This is reflected by the hydraulic gradient which is bigger in the north (piezometric curves are closer). In addition, the hydrochemistry of the CI groundwater is significantly affected by this variation in the lithology. This is discussed below.

The lithological variations and their impact on hydrochemical characteristics of groundwater have led to the definition of "useful reservoir" (B-B' cross-section in Fig. 1). It corresponds to the parts of the CI aquifer where sandstones and shally sandstone strata are dominating (-50 % of shale) and contain fresh groundwater (dry residue <6 mg/L) (UNESCO, 1972).

Hydrochemical and isotopic characteristics of the CI aquifer in Algeria have been primarily investigated by Edmunds et al. (2003); Moulla et al. (2012), and Darling et al. (2018). They demonstrated that the CI aquifer is hydraulically continuous from west to the east in the Great Oriental Erg sub-basin in accordance with the piezometric map. Using stable isotopes of water, they observed large variations between different regions and gave some conclusions that are mentioned and discussed below. It can be observed that the heterogeneities are more pronounced within the unconfined CI area than in the confined CI area both for hydrochemical and isotopic characteristics. However, the causes in differences of the chemical characteristics of CI groundwater throughout the Algerian Sahara, particularly between the north and the south of the Mzab Ridge, have not been explicitly discussed.

In a previous work, we focused on the north of Ghardaia region where high concentrations of NO_3 (Hakimi-Dedjell et al., 2019) were noticed. For the present work, nitrate content in the CI groundwater ranges from 0 to 58 mg/L, which is high for such confined aquifer system. To identify the nitrate content origin, we analysed NO_3 -stable isotopes.

3. Groundwater sampling and analysis campaign

The sampling campaign took place from February to May 2017 as the first detailed work addressing this problem of developing a high-density sampling network in the northwest of the Algerian Sahara (Ghardaia and El Golea regions). A total of 58 wells, public and private (agricultural farms), were sampled (Fig. 2). Some localities, such as Hassi Fehal, had been never covered before. In addition, many samples have been taken from recently drilled boreholes.

A subset of 20 wells were also sampled for the analysis of stable isotopes ($\delta^{18}\text{O-H}_2\text{O}$) $\delta^{2}\text{H-H}_2\text{O}$, $\delta^{18}\text{O-NO}_3$, and $\delta^{15}\text{N-NO}_3$). These wells were selected along groundwater flow directions, 2 W-E transects in the north and 1 N–S transects in the south, while two intermediate samples were situated in the middle of the Mzab Ridge.

The temperature and electrical conductivity (EC) of groundwater were measured in the field using a portable multiparameter YSI 556 MPS. pH measurements were not performed in the field due to technical issues. Groundwater samples were collected in 200 mL polyethylene vials for the analyses of major and minor ions, and in 140 mL polyethylene vials for the determination of dissolved iron and manganese. The second bottle was filtered using a 0.45 μ m membrane and acidified in the field. For water molecule isotope, δ^{18} O-NO₃, analyses, 60 mL polyethylene bottles were filled and filtered in the field using a 0.20 μ m membrane to eliminate bacterial activity and avoid NO₃ transformation before analysis.

Analyses were performed by ion chromatography for major elements and atomic absorption spectrometry for trace elements at the Aqueous Phase Analysis Laboratory of the University of Liège. Stable isotope analyses were accomplished by the Stable Isotope Laboratory of the Helmholtz Centre for Environmental Research in Halle, Germany. $\delta^{18}\text{O-H}_2\text{O}$ and $\delta^{2}\text{H-H}_2\text{O}$ analyses were carried out using a Laser Cavity Ring-down spectrometer. $\delta^{15}\text{N-NO}_3$ and $\delta^{18}\text{O-NO}_3$ isotopes of NO₃ were determined by the denitrifier method, which converts all sampled NO₃ to N₂O (Jurado et al., 2018) and are measured by a Delta V Plus IRMS coupled to the Thermo Scientific GasBench II. The results are expressed in delta " δ " notation and reported in per mil " δ " relative to the corresponding Vienna international standards, the Standard Mean Ocean Water (VSMOW) for $\delta^{18}\text{O-H}_2\text{O}$, $\delta^2\text{H-H}_2\text{O}$, and $\delta^{18}\text{O-NO}_3$ and Air-N₂ for $\delta^{15}\text{N-NO}_3$.

Saturation indices (SIs) were calculated using PHREEQC code (Parkhurst and Appelo, 2013). Self-organizing maps were generated using the SOM Matlab toolbox (Vesanto et al., 1999).

4. Results and discussion

4.1. Physical and chemical characteristics of CI groundwaters

In the SASS project (OSS, 2003), it was postulated to divide the CI groundwater samples from the Mzab ridge between the Great Oriental and Occidental Erg sub-basins, with the Hassi Fehal locality as the limit. In this paper, this hypothesis has been considered as well for consistency with these previous works. So, the samples from the north of the Mzab Ridge (mainly Ghardaia region) are considered in conjunction with those from the other localities in the east (Ouargla, Touggourt and El Oued) as the "oriental group" and those from its south (Hassi Fehal and El Golea regions) as the "occidental group". In the next sections, our samples, together with those from Edmunds et al. (2003); Moulla et al. (2012), and Darling et al. (2018) belonging to the confined CI, were compiled and discussed but represented in figures with a different symbology that is respected throughout the table containing all data added as a

supplementary material. The oriental group contained 72 samples and the occidental group contained 28.

EC ranges from 340 to 960 μ S/cm for samples belonging to the occidental group and from 1010 to 3500 μ S/cm for the oriental group (Fig. 3). Temperature values range between 25–34 °C and between 25–56 °C, respectively. As can be seen for both parameters, the values are lower in the occidental group than in the oriental one. The central region (Hassi Fehal) is characterized with intermediate values for both parameters.

Fig. 3 shows increasing temperature and EC along the WE transect, which was also demonstrated by Edmunds et al. (2003). The electrical conductivity increases along the main groundwater flow directions. The *in situ* temperature increases with depth (see B-B' cross-section in Fig. 1).

For the oriental group, Ca concentrations range from 80 to 280 mg/L, Mg from 25 to 100 mg/L, Na from 15 to 630 mg/L and K from 2 to 50 mg/L. Regarding anions, HCO_3 concentrations range from 100 to 250 mg/L, SO_4 from 80 to 1000 mg/L, SO_4 from 65 to 650 mg/L. For the occidental group, Ca, Mg, Na and K range from 30, 5, 25, and 5 mg/L to 80, 85, 115 and 15 mg/L, respectively. HCO_3 , SO_4 , and SO_4 and SO_4 from 80, 30, and 10 mg/L to 240, 200, and 140 mg/L (Fig. 4).

In El Golea region, the EC ranges around $450 \pm 100 \,\mu\text{S/cm}$. However, the EC of the sample Hg02 is $1300 \,\mu\text{S/cm}$. It is characterized by the highest concentrations for all ions compared to other samples in El Golea. This can be seen in the box plot of the occidental group where the maxima values, which represent Hg02 characteristics, are much bigger than the 3rd quartile. In addition, Hg02 is situated close to the El Golea Lake. This probably indicates that there are some leaks in this well from the shallow aquifer draining the evaporated water from the lake. For instance, NO_3 exceeds the WHO standard guideline for drinking water fixed at 50 mg/L (WHO, 2017).

Fig. 3 presents the spatial repartition of chemical type according to the degree of mineralization using a Stiff diagram (Stiff, 1951). There are two main water types: for the oriental group, the samples can be characterized as SO₄-Cl-Na type, and for the occidental group, they predominantly belong to the HCO₃-SO₄-Cl-Ca type.

In order to better describe the interaction between groundwater and the geological matrix, there are multiple approaches allowing a better understanding of the origin of mineralization and even to deduce which minerals are present in the aquifer. The most common are saturation indices (SI), as defined by Alley (1993) and Plummer et al. (1976).

The SI values range from -8.1 to -5.2 for halite, from -2.5 to -0.6 for gypsum (anhydrite in same range), and from -0.4 to 0.6 for calcite (aragonite in same range). Therefore, the CI groundwater is undersaturated with respect to halite and gypsum and in equilibrium with respect to calcite (Fig. 5). It can also be observed that SI values of the occidental group (El Golea) are smaller than those reported in the oriental group (Ghardaia, Ouargla and El Oued) for all minerals.

The new data shows clearly the variations in hydrochemical characteristics of CI groundwater between both sides of the Mzab Ridge. This difference is due to either lithological characteristics of the aquifer and/or to residence time and groundwater age. As explained in "Section 2", the CI aquifer is more sand-enriched, so more conductive, in El Golea region.

Such low values of saturation indices for halite and gypsum seem to indicate their absence in the geological matrix. However, the logarithmic tendency curve of their SI's are strongly correlated with EC ($R^2 > 0.9$, SI is calculated with logarithmic equation). Actually, Tellam (1995) has calculated SI for saline groundwater in Permo-Triassic sandstone aquifer. He found that in samples with gypsum and halite SI's equal to -0.5 and -1, respectively, SO₄ and Cl concentrations were approximately 3600 and 10,000 mg/L, respectively. Clearly, extremely high concentrations are necessary to approach equilibrium with respect to gypsum and halite. Thus, we deduce that

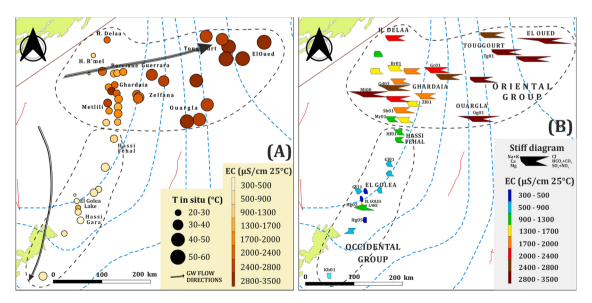


Fig. 3. Spatial evolution of physical parameters and chemical water type of the CI groundwater. In (A), the symbol size is referring to in situ-temperature and the colour scale is based on the electrical conductivity. In (B), the water type is showed using Stiff diagram which allows well visualizing of dominant ions and the mineralization.

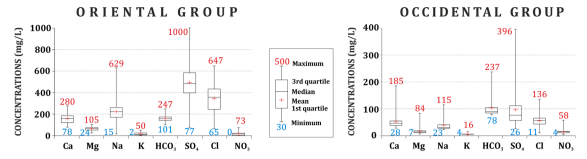


Fig. 4. Concentrations of major elements and their variation in CI groundwater in the confined area of the CI aquifer. Variabilities between the oriental and the occidental groups are much contrasted.

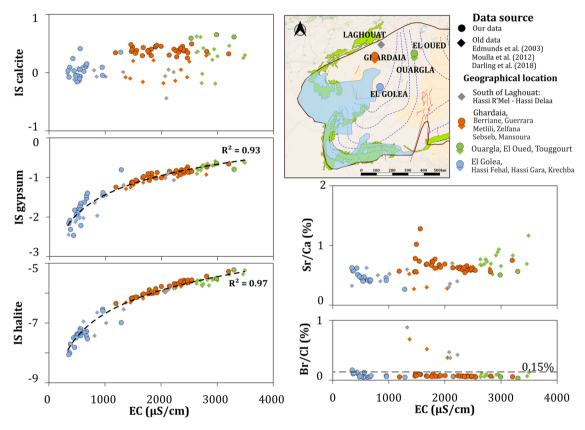


Fig. 5. Saturation indices for the main minerals. Anhydrite is in the same range as gypsum and aragonite is in same range as calcite. Sr/Ca and Br/Cl ratios are calculated using meq/L concentrations. The map shows spatial localization of samples. The samples collected during this survey are represented with dots. The diamonds are those from the old studies. This symbology is employed in the next figures.

the existence of evaporitic minerals in the CI aquifer can be supposed even in small proportion.

Those results show the major contribution of evaporitic minerals in CI groundwater mineralization all over its extension but their concentrations in groundwater are diluted. However, the bicarbonate type in El Golea region can be explained by the interaction of CI groundwater with the calcite cement of sandstone.

To verify this hypothesis, molar ratios using minor elements such as strontium and bromide are widely used in the literature to deduce the aquifer lithology (Alcalá and Custodio, 2008; Ben Alaya et al., 2014; Bouchaou et al., 2008; Hem, 1985; Herczeg et al., 1991; Négrel and Petelet-Giraud, 2005; Ouhamdouch et al., 2017).

Sr/Ca ratios range from 0.2 to 2.5 % (Fig. 5). According to Edmunds et al. (2003), this ratio is almost constant in the Great Oriental Erg sub-basin which should imply the dissolution of gypsum. This is in good agreement with the aforementioned geological data.

In addition, bromide is a good conservative anion behaving similarly as chloride. Most of the samples have a Br/Cl ratio less than 0.15 % (Fig. 5), which is the lower limit for seawater salinization (Herczeg and Edmunds, 2000). So, CI groundwater salinity is derived from dissolution of continental halite that is contained in the aquifer.

4.2. Stable isotopes of water molecule

Stable isotopes of the water molecule ($\delta^{18}O$ vs δ^2H) are presented in Fig. 6. The isotopic composition of CI groundwater in Algeria and Tunisia ranges from -9.2 to -2.3% for $\delta^{18}O$ and from -69.2 to -34.8% for δ^2H . Most of the samples from the CI aquifer lay below both the GMWL of Craig (Craig, 1961) and Sfax local meteoric lines (South Tunisia) defined by Maliki et al. (2000) and Abid et al. (2009), the closest dataset for rainwater available in the literature. The scatter plot of CI groundwater situated inside of the SASS basin in Algeria follows the trendline $\delta^2H = 3.01$ $\delta^{18}O$ - 36.70 showing that the CI groundwater has undergone evaporative processes. In addition, the samples from bordering regions (Saharan Atlas and Saoura Valley) follow the trendline $\delta^2H = 3.07$ $\delta^{18}O$ - 31.39.

Groundwater from the confined and unconfined CI aquifer show different behaviours. On the one hand, groundwater from the confined part (except for the samples from south of El Golea) are the most depleted ($\delta^{18}O < -6\%$), while $\delta^{18}O$ for groundwater from the unconfined part of the aquifer is more variable (from -8 to -2.5%).

Despite the big distances separating samples from the confined CI situated in the Great Oriental Erg sub-basin, their isotopic composition is more homogeneous than that from the unconfined CI situated in the Great Occidental Erg sub-basin. This has been reported since the first studies on the stable isotope composition of the CI groundwaters (Conrad and Fontes, 1970; Gonfiantini et al., 1974). They concluded that more variability in isotopic composition induces more variation in groundwater origins and recharge conditions and vice versa. Gonfiantini et al. (1974) and Edmunds et al. (2003) confirmed that CI isotopic composition under the Great Oriental Erg sub-basin is the initial signature of an old recharge because the CI groundwater there is protected from the external input or evaporation.

Using tritium content, Moulla et al. (2012) highlight the existence of modern recharge in Saoura Valley shallow aquifer, which shows also an evaporative signature. Moreover, Gonfiantini et al. (1974) and Edmunds et al. (2003) showed, that CI groundwater contains around 55 pmc of 14 C in Gourara region and the Saharan Atlas, successively, where the CI layers are unconfined. These activities indicate the ingoing of relatively modern recharge in those regions. So, the trendline equation " $\delta^2 H = 3.07 \delta^{18} O - 31.39$ ", which includes the samples from the unconfined aquifer, should represent modern Saharan groundwater.

In addition, the CI groundwater isotopic signature is significantly depleted relative to weighted mean modern rainfall in Sahara ($\delta^{18}O = -5.1\%$, $\delta^2H = -26\%$) calculated by Edmunds et al. (2003). So, the CI groundwater cannot be in any way derived from the actual rainfall as concluded by Darling et al. (2018). According to Rozanski et al. (1993), these depleted values seem to indicate that CI groundwaters were infiltrated under a cooler and/or wetter climate regime than today. This is in agreement with recharge temperatures calculated by Guendouz et al. (1997) using noble gases, which were 2–3 °C cooler than the present mean annual temperature of 21 °C. Moreover, several studies have showed that the most important recharge occurred during the Late Pleistocene when the paleoclimate in the North Sahara and Sahel regions was wetter than today (e.g., Beyerle et al., 2003; Edmunds, 2009).

Knowing that the CI aquifer under the Great Occidental Erg sub-basin is unconfined and sometimes covered by pervious layers of Mio-Plio-Quaternary (almost dunes), the samples from El Golea region should be more exposed to evaporation and more enriched in stable isotopes. This corroborates with the work of Gonfiantini et al. (1974) who concluded that CI groundwaters in the Great Occidental Erg sub-basin indicate "an evaporation mechanism, probably peculiar to the dune systems, which gives rise to heavy-isotope enrichment compared with the recharge of other types of formations."

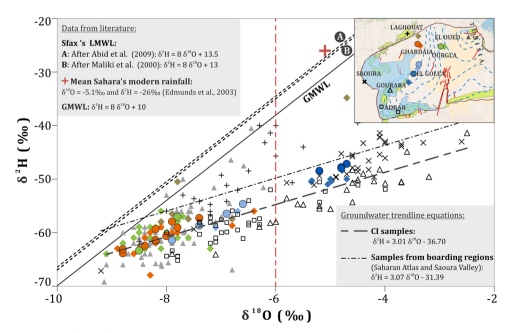


Fig. 6. Isotopic composition (δ^2 H vs δ^{18} O) of CI groundwaters. Data from the unconfined CI are also plotted. The isotopic composition of CI in South Tunisia after Petersen et al. (2018) is represented by the grey triangle for comparison.

However, samples situated downgradient in the south of El Golea region (Hassi Gara and Krechba), where CI is confined, are the most enriched samples of all over the confined CI ($\delta^{18}O > -6\%$). These groundwaters could be the result of a mixture between old groundwater and recently evaporated water. Another possible explanation is that the recharge area for CI in El Golea region is different between the north (El Golea city) and the south (Hassi Gara and Krechba).

In summary, end-members of the scatter plot correspond to the most depleted samples from the confined CI under the Great Oriental Erg (Ouargla, El Oued) and the most enriched samples from the occidental basin (Gourara region).

Finally, Edmunds et al. (2003) concluded that there is no correlation between $\delta^{18}O$ and $\delta^{2}H$ of CI groundwater and its age in the Great Oriental Erg sub-basin. However, this global vision demonstrates that there is an isotopic depletion that is proportional to the distance from the recharge area and thus to the age of groundwater. The most enriched groundwaters are situated in the Great Occidental Erg sub-basin, where CI is partially unconfined.

To verify the hypotheses and the conclusions of this section, it is likely that the use of radioactive isotopes and noble gases would provide more clarification. It is the next step of our research project.

4.3. Multivariate analysis with a self-organizing map (SOM)

The self-organizing map (SOM) technique is an artificial neural network (ANN) algorithm based on unsupervised learning. It is a powerful tool for exploring and clustering large and multivariate datasets (Kohonen, 1995). This technique has been widely applied is various fields of groundwater quality (e.g., Hong and Rosen, 2001; Jurado et al., 2018; Tobiszewski et al., 2012).

The main objective of SOM is to reduce the dimensionality of datasets for easier visualization of their variability. First, all samples are normalized and introduced as "input vectors", and then the algorithm runs an iterative "training process" to join each sample to one "node" of the artificially created network corresponding to "output vectors". The "Euclidean distance" between these two vectors should be as small as possible. Unlike ordinary multivariate statistical techniques, SOM is able to consider non-linear relationships between variables and run with missing data. As a result, the SOM produces two kinds of outputs: (1) A "U-matrix" showing the observation's position and defining clusters using K-mean clustering methods, where the number is chosen by the user. The closer the two individuals are, the more they look alike. (2) The 2D "component planes" showing the values of each variable, according to the precedent matrix. The observation of the component planes allows deductive relationships and correlations between variables, based on their similarities or dissimilarities (Peeters et al., 2007).

We chose the SOM technique to discuss two phenomena (Fig. 7). (i) First, defining redox conditions at the captive part of the CI aquifer using results of the available indicators of these processes, namely, NO_3 and Fe + Mn. (ii) In addition, we included other susceptible variables to explain the evolution of fluoride concentrations in CI groundwater.

A distribution of the samples into two distinct groups (oriental and occidental) was clearly shown by the U matrix. The "OCD" cluster groups the samples belonging to the Great Occidental Erg sub-basin. In addition, a set of clusters "OR1-4" includes the samples belonging to the Great Oriental Erg sub-basin with an arrangement according to the direction of flow mentioned by the arrows.

The component matrix shows that δ^{18} O of the CI groundwater is inversely correlated with the electrical conductivities (ECs). As demonstrated in the previous section, the enrichment of δ^{18} O is inversely correlated with the residence time as well as with EC. Samples from the occidental cluster (OCD) are the most enriched and the less mineralized.

 NO_3 concentrations range between 3 and 75 mg/L for OCD, OR1, OR2, and OR3 clusters, and they are less than 1 mg/L in OR4 cluster. It can be noticed that they significantly decrease downgradient. The component matrix shows that Fe + Mn is inversely correlated with NO_3 . These elements have the highest concentrations in the downgradient OR4 cluster. This corresponds to residence time and the predominance of reduced conditions downgradient of the confined CI. Actually, after consuming of NO_3 by denitrification, Fe and Mn become the terminal electron acceptors, they are reduced and solubilized in groundwater (Appelo and Postma,

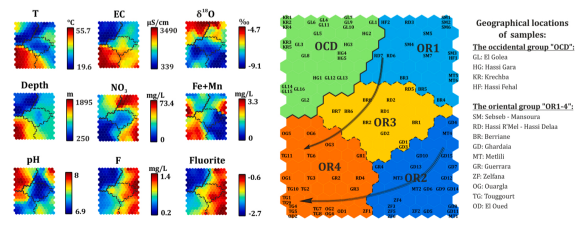


Fig. 7. SOM results for samples belonging to the confined CI. On the right, the U-matrix defines the clusters: four clusters from the oriental group "OR1-4" and one cluster from the occidental group "OCD". On the left, the component planes show the variables and explain the correlations between them. The samples' names indicate their geographical locations.

2005). This is in agreement with the results of Edmunds et al. (2003) who showed that even if the oxidizing conditions dominate in the captive part of the CI south of Laghouat (RD samples) and approximately on the whole region of Ghardaia (samples: BR, GR, GD and ZF), they begin reducing in the east of the sub-basin of Grand Erg Oriental (samples: TG, OG and OD).

It can be noted that NO_3 concentrations are higher upgradient of the oriental clusters "OR1-2" than the occidental clusters "OCD" that are also characterized by a depletion in $\delta^{18}O$. The previous section had already concluded that the palaeoclimatic conditions of the CI recharge were wetter and colder in the Great Oriental Erg sub-basin than in the Great Occidental Erg. In a similar context, Edmunds and Gaye (1997) studying high nitrate concentrations in groundwater in the Sahel countries, concluded: "There is also evidence that high ratios relate to wetter and low values to more arid climatic periods".

Another phenomenon is the decrease in fluoride concentrations downgradient from 1.4 to 0.4 mg/L within the clusters of the oriental clusters "OR1-4". Paradoxically, this decrease is not accompanied by an increase in the saturation index of fluorite, the main mineral controlling its concentration in water. This would thus indicate an evolution towards the precipitation of this mineral. In contrast, there was a decrease from -0.6 to -1.7, which implies a tendency towards dissolution if the mineral would have been present.

Moreover, in the literature, fluoride mobility is considered a "pH-dependent" process. The fixation of F following anionic exchange induces the release of OH, which would increase the pH. Paradoxically, for the oriental clusters, a decrease in pH can be observed downgradient with a decrease in F concentrations. Thus, the hypothesis of anionic F/OH change as a cause of decrease in F levels can be excluded.

The negative correlation of the F with δ^{18} O is also observable in our dataset. This reinforces the hypothesis of Edmunds et al. (2003) that F contents are influenced by palaeoclimatic conditions during recharge. Samples located upgradient of the oriental clusters (OR1 and OR2) were exposed to more intense evaporation than those located downgradient (OR4). On the other hand, the lowest levels were observed in the occidental cluster (OCD), whereas the stable isotope data show that these waters are the most evaporated. According to Edmunds and Smedley (2013), these concentrations are specific to the rainfall signature and their variations are related to evaporation.

4.4. Nitrate origin

Fig. 8 shows that δ^{15} N-NO₃ ranges from -1.9% to 9.5% and that δ^{18} O-NO₃ ranges from 2.6% to 12.6%, except for three samples, HgO3, GrO1 and OgO2 (Fig. 8). Based on the classification of Mayer (2005), which is an updated version of that of Kendall (1998), such low values reflect soil nitrification processes, a natural origin of NO₃, although reported concentrations are high.

Og01 and Tg01 samples contain less than 0.3 mg/L of NO₃, which is not enough to analyse the NO₃ isotopes. They are situated downgradient, northeast of the study area, in the reducing zone according to Edmunds et al. (2003). Therefore, such low concentrations can be explained by an ongoing denitrification process of reaction.

Until now, many studies have reported high values of NO_3 in the sandstone aquifers of arid zones in Australia (Barnes et al., 1992), in Sahelian countries (Edmunds and Gaye, 1997) and in the Kalahari Desert (Heaton et al., 1983; Stadler et al., 2008, 2012). They concluded by using different methods that nitrates originate from natural sources. Some studies have also proven that soil can contain naturally high values of NO_3 in arid zones (Schwiede et al., 2005; Stone and Edmunds, 2014; Walvoord et al., 2003).

Gr01 and Og02 samples were strongly enriched in both δ^{15} N-NO₃ and δ^{18} O-NO₃. Knowing that their NO₃ concentrations are 2.7 and 3.9 mg/L, respectively, and they are situated downgradient, close to Og01 and Tg01, their positions in the plot show that they are situated in the denitrification trajectory.

Sample Hg03 was strongly enriched in δ^{18} O-NO₃, with isotope signatures close to NO₃ derived from synthetic nitrate-containing fertilizers. The sampled borehole was located on agricultural land and was used for irrigation, where pollution may have occurred.

The position of sample Hg02 in the boxes representing the origin of NO₃ (with respect to the isotopic composition) confirms that its high mineralization is due to a mixing with natural evaporated water that does not originate from polluted groundwater.

The oriental group is more enriched in NO₃ isotopes than the occidental group. This is most likely to be due to partial denitrification

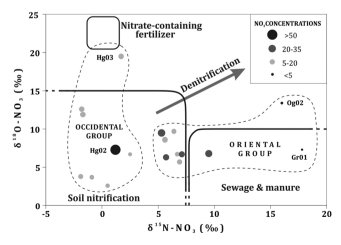


Fig. 8. Identification of NO₃ origin using the classification of Mayer (2005).

occurring in this part of the study area or to differences in recharge conditions.

5. Conclusions

This paper fills the gap in the knowledge of Continental Intercalaire aquifer hydrogeochemistry on both sides of the Mzab Ridge immediate vicinity.

It can be concluded that groundwater in the confined area of the CI aquifer under the Mzab Ridge and the Great Oriental Erg sub-basin behaves differently than that in the unconfined zone in the Great Occidental Erg sub-basin. In the first area, the characteristics are primarily controlled by water-rock interactions that involve residence time, but in the second zone, some external factors disturb those interactions.

Using classical methods of hydrochemistry, isotope evidence, and classification techniques, CI groundwater in the confined area is proven to belongs to two different groups according to their geographical location, namely, oriental (Ghardaia, Ouargla and El Oued regions) and occidental (Hassi Fehal and El Golea regions) groups with distinct and specific hydrochemical characteristics for each region.

The observed divide between these two groups is not strict. The samples obtained from locations situated between 31.55° and 31.57 °N latitude (Hf01 and Hf02) represent a transitional member from the hydrochemical and the isotopic perspective. This finding corroborates the observations from the regional piezometric map suggesting a groundwater divide situated in this zone.

It was clear that the variations in hydrochemistry of the CI groundwater were largely attributed to the impact of aquifer lithology. Moreover, as could be expected in such vast transboundary aquifer, the residence time also has a pronounced influence.

On the one hand, in the Great Occidental Erg sub-basin, the CI aquifer is sand-enriched, and the groundwater flowing through is poorly mineralized. Mainly, it could be attributed to the HCO_3 -Ca water type, though it becomes progressively enriched in SO_4 and CI downgradient to the south. On the other hand, in the Great Oriental Erg sub-basin, the CI aquifer is clay and gypsum-enriched. Here, the groundwater is highly mineralized and predominantly belongs to the SO_4 -Cl-Na water type. Minor element ratios indicate that the salinization originates from dissolution of continental evaporates.

 NO_3 -stable isotope analysis shows that the high nitrate content has a predominantly natural origin of soil nitrification. However, the denitrification processes are insignificant in the study area even though the CI aquifer is confined, except downgradient in the northeastern part of the study area.

Stable isotopes of water molecules from the CI groundwater have shown that the most depleted samples ($\approx \delta^{18} O < -6\%$) are those situated in the confined area within the Great Oriental Erg sub-basin (including north of the Mzab Ridge and South Tunisia). Their isotopic composition is homogeneous representing, apparently, the older meteoric water infiltrated under palaeoclimatic regime cooler and wetter than the present day climate. Moreover, the most enriched and heterogeneous samples are those belonging to unconfined CI (mainly the Great Occidental Erg sub-basin). The CI groundwater lies along the mixing line $\delta^2 H = 3.01 \ \delta^{18} O - 36.70$. The end-members correspond to the most depleted groundwater from the confined CI under the Great Oriental Erg (Ouargla, El Oued) in one side and in another side to the most enriched samples from Gourara region in the Great Occidental Erg sub-basin. In El Golea, some samples are relatively enriched in both $\delta^{18} O$ and $\delta^2 H$, which suggested the existence of mixing of the typical old CI groundwater with younger and evaporated water infiltrated under the dune of the Great Occidental Erg.

As a follow-up for this work, we are planning to use cosmogenic isotopes (¹⁴C, ³⁶Cl) to determine the residence time of the CI groundwater and noble gases to understand the climatic conditions during its recharge in order to verify some of the abovementioned hypotheses.

More generally, we recommend placing greater focus on the extreme northwestern zone near the recharge area (El Bayadh region), both from the quantitative and the qualitative viewpoint. This would allow a better understanding of modern recharge reinforcing thus the strategies aiming at a sustainable exploitation of this invaluable resource.

CRediT authorship contribution statement

Youcef Hakimi: Conceptualization, Data curation, Formal analysis, Methodology, Software, Writing - original draft, Writing - review & editing. Philippe Orban: Conceptualization, Formal analysis, Methodology, Project administration, Supervision, Validation, Writing - review & editing. Pierre Deschamps: Conceptualization, Formal analysis, Methodology, Project administration, Supervision, Validation, Writing - review & editing. Serge Brouyere: Conceptualization, Formal analysis, Methodology, Project administration, Supervision, Validation, Writing - review & editing.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary material related to this article can be found, in the online version, at doi:https://doi.org/10.1016/j.ejrh.2021. 100791.

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