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Recharge, groundwater flow pattern and contamination processes in an arid volcanic area: Insights from isotopic and geochemical tracers (Bara aquifer system, Republic of Djibouti)

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*Original*

Recharge, groundwater flow pattern and contamination processes in an arid volcanic area: Insights from isotopic and geochemical tracers (Bara aquifer system, Republic of Djibouti) / Awaleh, Mohamed Osman; Baudron, Paul; Soubaneh, Youssef Djibril; Boschetti, Tiziano; Hoch, Farhan Bouraleh; Egueh, Nima Moussa; Mohamed, Jalludin; Dabar, Omar Assowe; Masse Dufresne, Janie; Gassani, Jean. - In: JOURNAL OF GEOCHEMICAL EXPLORATION. - ISSN 0375-6742. - 175:(2017), pp. 82-98. [10.1016/j.gexplo.2017.01.005]

*Availability:*

This version is available at: 11381/2820477 since: 2018-09-17T15:04:11Z

*Publisher:*

Elsevier B.V.

*Published*

DOI:10.1016/j.gexplo.2017.01.005

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note finali coverpage

(Article begins on next page)

13 August 2025

## Accepted Manuscript

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PII: S0375-6742(17)30018-3

DOI: doi: [10.1016/j.gexplo.2017.01.005](https://doi.org/10.1016/j.gexplo.2017.01.005)

Reference: GEXPLO 5885

To appear in: *Journal of Geochemical Exploration*

Received date: 27 June 2016

Revised date: 30 November 2016

Accepted date: 6 January 2017

Please cite this article as: Mohamed Osman Awaleh, Paul Baudron, Youssouf Djibril Soubaneh, Tiziano Boschetti, Farhan Bouraleh Hoch, Nima Moussa Egueh, Jalludin Mohamed, Omar Assowe Dabar, Janie Masse-Dufresne, Jean Gassani , Recharge, groundwater flow pattern and contamination processes in an arid volcanic area: Insights from isotopic and geochemical tracers (Bara aquifer system, Republic of Djibouti). The address for the corresponding author was captured as affiliation for all authors. Please check if appropriate. Gexplo(2017), doi: [10.1016/j.gexplo.2017.01.005](https://doi.org/10.1016/j.gexplo.2017.01.005)

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# **Recharge, groundwater flow pattern and contamination processes in an arid volcanic area: insights from isotopic and geochemical tracers (Bara aquifer system, Republic of Djibouti)**

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## **Abstract**

Fractured volcanic aquifers are the main water resources in the arid climate of the Republic of Djibouti. Nonetheless, these strategic reservoirs are overexploited and their comprehensive understanding is therefore a pre-requisite for a sustainable use. A geochemical and isotopic survey, including major ion chemistry, <sup>2</sup>H, <sup>18</sup>O, <sup>13</sup>C, <sup>3</sup>H, <sup>87</sup>Sr/<sup>86</sup>Sr, <sup>15</sup>N was performed and

combined with existing  $^{14}\text{C}$  data to study recharge, contamination processes and groundwater flow patterns inside and between the compartments of a complex aquifer system composed by basaltic rocks and by alluvium located in Petit Bara, Grand Bara, and Mouloud areas (Southwest of Djibouti). A main feature was the common trend from a fresh  $\text{Na-Cl-HCO}_3$  water type (alluvium groundwaters) to an intermediate water type (alluvium and basalt groundwaters) and finally to a  $\text{Na-Cl-SO}_4$  water type (most mineralized basalt groundwater). Elementary and isotopic nitrate evidenced and located anthropogenic and geogenic origins of nitrate. Alluvium groundwaters had  $\delta^2\text{H}$  and  $\delta^{18}\text{O}$  signature of modern precipitation while basalt groundwaters were significantly depleted and enriched in  $\delta^{13}\text{C}$  due to water-rock interactions. Modern radiocarbon and tritium were evidenced in the alluvium groundwaters, while recalculated radiocarbon ages located recharge of the basalt groundwaters in the early to mid-Holocene. These features revealed a common evolutionary pattern, with a recharge from wadi-rivers to the alluvium and a downward circulation to the basalt through major faults, combined with a mixing with a more geochemically evolved groundwater. Accordingly, highly saline groundwater at the outlet of the Petit Bara plain was found to be diluted by modern recharge in the alluvium. Two major basaltic aquifer compartments were found to be connected (Grand Bara and Mouloud), revealing a global northeastward flowpath below the endorheic Grand Bara plain.

*Keywords:* Djibouti, Hydrochemistry, Volcanic aquifer, Stable isotopes, Nitrogen isotopes

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## 1. Introduction

The republic of Djibouti (23,000 km<sup>2</sup>) is located in the Horn of Africa (Fig. 1), which is an emerged triple junction of the Red Sea, the Gulf of Aden and the East African Rift that correspond to large seismic, tectonic and volcanic activities (Barberi et al., 1975; Mlynarski and Zlotnicki, 2001).

On the other hand, the republic of Djibouti is subjected to an arid climate with an annual mean rainfall of 150 mm. This harsh climate explains the lack of permanent rivers, and has led the country to exploit the only available water resources, which are represented by groundwater in the volcanic aquifers (Jalludin and Razack, 1994). The continuous and drastic increases in water demand has led to an intensive exploitation of volcanic aquifers and has severely depleted its reserves and led to quality deterioration (Houssein and Jalludin, 1996), as found in several arid to semi-arid aquifers worldwide (e.g. Jimenez-Martinez et al., 2016).

The population of Djibouti relies heavily on groundwater resources for drinking water and irrigation purposes. Therefore, knowledge of the origin of groundwater resources and their renewal forms as well as the consideration of their vulnerability facing the anthropogenic pressure is essential to the rational management of volcano-sedimentary aquifers in Djibouti. However, few studies have been carried out on the hydrogeology (Jalludin and Razack, 1994; Jalludin and Razack, 2004) and hydrochemistry (Aboubaker et al., 2013) of these complex volcano-sedimentary aquifers. These studies address only briefly recharge conditions, groundwater residence times and groundwater flow pattern inside these aquifer systems. In order to fulfill this gap, environmental and radiogenic isotopes ( $\delta^2\text{H}$ ,  $\delta^{18}\text{O}$ ,  $^3\text{H}$ ,  $\delta^{13}\text{C}$ ,  $^{14}\text{C}$ ,  $^{87}\text{Sr}/^{86}\text{Sr}$ ) as well as major ion chemistry combined with hydrogeological and geological data are applied to

assess groundwater dynamics of the Bara aquifer systems located on the north of the East Africa Rift.

Furthermore, nitrate dual isotopes ( $\delta^{15}\text{N-NO}_3^-$  and  $\delta^{18}\text{O-NO}_3^-$ ) are applied to identify the major sources of  $\text{NO}_3^-$  in groundwaters from this volcano-sedimentary aquifer. Indeed, the isotopic composition of dissolved nitrate in waters ( $\delta^{15}\text{N-NO}_3^-$  and  $\delta^{18}\text{O-NO}_3^-$ ) has been used extensively to better constrain the sources of nitrate in groundwater (Edmunds and Gaye, 1997; Jackson et al., 2015; Kendall, 1998; Kendall et al., 2007; Stadler et al., 2008; Walvoord et al., 2003). However, to the best of our knowledge, no nitrate dual isotopes were used to investigate the origin of nitrate in groundwater in arid or semi-arid rift systems. The present study reports the first results of nitrogen isotopes of groundwater from semi-arid environment located in the East Africa Rift.

The results of this study will contribute to a comprehensive knowledge of the complex volcano-sedimentary aquifer systems in the Bara basin, by defining the groundwater flow pattern and the relations between aquifer compartments, assessing groundwater residence time and deciphering the origin of high nitrate content in semi-arid environment in the East Africa Rift areas. This study would serve as a valuable base for other integrated hydrodynamic, geochemical and isotopic studies of volcano-sedimentary aquifers in similar arid environments.

## **2. Site description**

### *2.1 Climate and hydrology*

The republic of Djibouti has a low precipitation regime, with annual mean rainfall of 150 mm. Overall, two seasons predominate: a cool season (winter) from October to April and a hot season (summer) from May to September. In winter, the climate is characterized by northeast trade

winds coming from Saudi and Gulf of Aden and mean temperature comprised between 20°C and 30°C. In summer dominates equatorial westerly wind zone and mean temperatures comprised between 30°C and 45°C with high rate of evapotranspiration amounting to 2000 mm per year (BGR, 1982).

The study area is localized in two main surface watersheds, which are Grand Bara and Ambouli (Fig. S1, see supporting information). The Grand Bara watershed, with an area of 869 km<sup>2</sup>, is characterized by an average altitude of 630 meters and average slopes of 9%. Due to the low permeability of outcropping rocks, this endorheic basin plays a major role in the concentrated runoff process and flash flood of different wadis to its localized downstream portion in the center of this depression. The Ambouli surface watershed, with an area of 774 km<sup>2</sup> and an average altitude of 470 meters and average slopes of 10%, is an exoreic watershed with its outlet in Djibouti city. The Petit Bara area (140 km<sup>2</sup>) is located in the upstream part of the Ambouli surface watershed.

## *2.2 Geological settings*

The Republic of Djibouti is located in the Southeastern part of the Afar rift which resulted from the intersection of two nascent mid-oceanic ridges (Red Sea and Gulf of Aden) and a continental rift (East African Rift) during the Miocene (Fig. 1). The first evidence of synrift magmatism in SE Afar is the 28–19 Ma Ali Sabieh mafic complex located in the South of the country and the more widespread Mablás rhyolites. These basalts are the effusive component of a magmatic complex that includes a laccolithic intrusion, the emplacement of which is assumed to have caused the NE-plunging antiform structure of the Ali Sabieh ranges by arching the Mesozoic sedimentary rocks (Le Gall et al., 2010). The Mablás rhyolites form the outer rim of this antiform and consist of rhyolites, ignimbrites and minor basaltic lava flows, intruded by felsic dykes

(Gadalia, 1980; Audin et al., 2004). These rhyolites were followed by the emplacement of the Dalha series, in which about 1000 m of basaltic flows erupted at 8.3-4.3 Ma, just before the initial opening of the Tadjoura Gulf (Gasse et al., 1986). These basalts are correlative with the Somali basalts emplaced approximately during the same period of time (7.2-3 Ma) (Audin et al., 2004).

Gasse et al. (1986) are reported for the Dalha series an aphyric facies with plagioclase olivine and pyroxene minerals. In addition, sedimentary rocks (clays, alluviums, diatomites) interbedded in Dalha basalts and fill tectonic basins.

The Stratoid basalts (3-1 Ma) cover two-third of the Afar depression. They unconformably overlie the Dalha basalts series and the sedimentary formation. During the last 3 Ma, the SE Afar margin recorded several episodes of extensional faulting in various tectonic setting (Daoud et al., 2011).

The study area is located in the south-eastern part of the country. The Petit Bara and Grand Bara zones correspond to sedimentary basins limited to the East by the Dalha basalts (8.3-4.3 Ma) and to the West by the Stratoid basalts (3-1 Ma) (Fig. 1).

The Petit Bara and the Grand Bara zones describe clearly different fracture trends of the Dalha basalts (Arthaud and Jalludin, 1993). A common zone separates both plains and shows a relatively regular fault network with EW and N160°-180° fractures. All the western part of the Grand Bara plain is characterized by mainly EW and N110°-130° faults (Arthaud and Jalludin, 1993). Dalha basalts fractures have a particular behavior with elliptical faults lines in the Northern part of the Petit Bara (Arthaud and Jalludin, 1993).

The #1 – #6 boreholes are located mainly in the Didjan Der valley at the north of the Petit Bara area, while #7 borehole is drilled in the south-east of the Petit Bara plain (Fig. 1). The depth of these boreholes ranges from 92 and 170 meters (Table 1). The lithologies of the #2 – #5 boreholes consist of alluvia and altered scoria interbedded with clays, this latter is associated with



calcite, while the #6 borehole consists of alternation of basalts, alluvia interbedded with silts and calcareous clays.

It is of interest to note that the #8 borehole was located on the basalt outcrop between the Petit Bara zone and the Grand Bara zone. That is why we have categorized #8 borehole as Central area in the Table 1. Since we didn't want to classify this central area apart, we decided to put #8 borehole in the Petit Bara zone in the Table 1.

In the Grand Bara zone, the study area is focused on the western flank of the Dalha basalts (#9, #11, #13, #14, #15, #16 and #17 boreholes); whereas only one borehole (#18) is located in eastern part of the Stratoid basalts, while borehole #12 is located in the Grand Bara plain. In this area, the Dalha basalts are affected by two distinct sets of faults at N160-180°E crosscutted by E-W faults-fractures and to a lesser extent N110 -130°E crosscutted by NS and N60-70°E directions. The basalts are delimited by several wadi valleys (Dagahdamer, Awraoussa, Dadin, and Mouloud). Depth of the drilled boreholes ranges between 90 and 181 meters, except for the #13 shallow dugwell which has a depth of about 19 m (Table 1). The following vertical successions were observed: (1) alternation of calcareous silts, sands, alluvia and clayish silts (#10); (2) calcareous sands and alluvium interbedded with altered basalts and scoria (#17); and finally (3) alternation of sands, alluvia associated with pebbles of basalts and rhyolites, silts, altered basalts and scoria (#15 and #16).

The Mouloud-Dadin area, which is part of the Dikhil basin, is located between the plain of Grand Bara and Dalha basalts (Fig. 1). Depth of the drilled boreholes in this area ranges between 115 and 150 meters (Table 1). The drilled sections exhibit complex deposits with an alternation of basaltic lava flows interbedded with alluvia and argillaceous sediments for Dadin (#22). Aboubaker (2012) reported that Dalha basalts appear on approximately ten meters. The Mouloud boreholes (#20, #21) are characterized by alluvia and basalts interbedded with sedimentary rocks

(alluvia and clays). Calcite occurs in vesicles of scoria and fissures. The alternation of alluvium, altered basalts, scoria with calcite, argillaceous sediments in the different wells hardens the correlation between them.

### *2.3 Hydrogeological settings*

The groundwater in Djibouti is controlled by volcanic and sedimentary aquifers. Volcanic aquifers systems are mainly represented by the Dalha basalts, the stratoid basalts and the Mabila rhyolites (BGR 1982; Jalludin and Razack 1994; Jalludin and Razack 2004).

Sedimentary aquifers are located in the two plains of Petit Bara and Grand Bara and also along the main wadi beds (Fig. 1). The plains are generally filled by very heterogeneous sediments with coarse materials as gravels, sands, shales and fine sediments. The substratum is represented by the Dalha basalts. Pleistocene sediments mainly fill the plains and the upper part is irregularly covered by holocene deposits. The thickness might exceed 150 meters in "Grand Bara" and "Petit Bara" sediments should be above 100 meters. In addition, sediments along the narrow wadi bed form of a few to several tens of meters thick aquifer systems that are considered to play a significant role in the recharge mechanism. Given the geomorphological conditions and the showery type of rainfall, recharge may occur mainly in these aquifers during runoff generated by short showery periods. Recharge of the underlying Dalha basalts would then be ensured by these wadi aquifers through faults caused by intense tectonics (Aboubaker, 2012; BGR 1982). Therefore, one may possibly consider hydraulic continuity between sedimentary and volcanic systems through tectonic faults, fissured volcanic flows and horizontal levels of slag and lava breccias.

A cross section of the Petit Bara area is shown in Fig. 2A. On the other hand, the substratum of the Grand Bara sedimentary basin is supposed to consist essentially of basalts (Fig. 2B), and the

basaltic aquifer may continue under the sedimentary cover of the Grand Bara plain (Aboubaker, 2012). Dalha basalts and sediments unconfined aquifers are considered potential aquifers for water supply in the study area. The transmissivity of Dalha basalts ranges between  $1.5 \cdot 10^{-4} \text{ m}^2/\text{s}$  and  $2.4 \cdot 10^{-3} \text{ m}^2/\text{s}$ , and for the sediments between  $5.6 \cdot 10^{-6} \text{ m}^2/\text{s}$  and  $1.3 \cdot 10^{-3} \text{ m}^2/\text{s}$ . A cross section of the Mouloud-Dadin area highlights the geometry of the alluvial aquifer, mainly determined by the fracturing system that shift the basalt panels vertically several meters by a set of normal faults (Fig. 2C). In Mouloud-Dadin area, the transmissivities range between  $10^{-3} \text{ m}^2/\text{s}$  and  $10^{-4} \text{ m}^2/\text{s}$  due to the presence of clay (CERD, 2014).

### 3. Material and methods

Twenty two borehole waters were collected in April – June 2014. Whenever possible, water level in boreholes was measured manually with a dip meter. Temperature ( $\pm 0.1 \text{ }^\circ\text{C}$ ), pH ( $\pm 0.01$  unit), electrical conductivity ( $\pm 1 \mu\text{S}/\text{cm}$ ), redox potential ( $\pm 0.1 \text{ mV}$ ), and Dissolved Oxygen ( $\pm 0.1 \text{ mg O}_2/\text{l}$ ) were measured on site using portable instruments, i.e. CheckTemp (Hanna), pH 610 (EutechInstruments), COND 610 (Eutech Instruments), WTW multi 3410, and YSI 550A DO Instrument respectively. Each was calibrated in the field prior to sampling. Water samples were collected in polyethylene containers after filtration through  $0.45 \mu\text{m}$  membrane filters. All samples used for determination of cations were acidified after collection through addition of Suprapure®  $\text{HNO}_3$  (Merck). Analyses of anions and major cations were carried out using a Dionex ICS 3000 Ion Chromatograph calibrated through repeated analysis of five working anions and cations standards, with concentrations within the range of analyses. Analyses of boron and strontium elements were conducted using an Ultima 2 (Horiba Jobin Yvon) Inductively Couple Plasma Atomic Emission Spectrometer (ICP-AES). National Institute of Standards and

Technology traceable commercial standards were used for quality control. These standards were analyzed to within  $\pm 3\%$  of the known values. For the analysis of aqueous  $\text{SiO}_2$ , the water samples were diluted fivefold using deionized water to prevent  $\text{SiO}_2$  precipitation.  $\text{SiO}_2$  contents were determined by colorimetry and analyzed using a Jenway 6300 spectrophotometer, while  $\text{HCO}_3^-$  was analyzed by titration with 0.1 M HCl. The charge balance between anions and cations ( $(\Sigma[\text{Anions}] - \Sigma[\text{Cations}]) / (\Sigma[\text{Anions}] + \Sigma[\text{Cations}])$ ; concentrations in  $\text{meq l}^{-1}$ ) was assessed and analyses were accepted for deviations of less than 3%.

Additional samples of untreated waters were collected in 50 mL glass bottles (Quorpak) for analyses of stable isotopes of the water molecule ( $\delta^2\text{H-H}_2\text{O}$  and  $\delta^{18}\text{O-H}_2\text{O}$ ), in 1000 mL plastic bottles for tritium ( $^3\text{H}$ ) analysis, in 1000 mL plastic bottles for carbon-13 of dissolved inorganic carbon ( $^{13}\text{C-DIC}$ ) and in 250 mL plastic bottles for  $^{87}\text{Sr}/^{86}\text{Sr}$ . The isotope ratios of hydrogen and oxygen were analyzed in BRGM (Orleans, France) using a Finnigan MAT 252 mass spectrometer and converted in per mil delta values ( $\delta\text{‰}$ ) versus the Vienna Standard Mean Ocean Water (V-SMOW) standard following  $\delta\text{ (‰)} = [(R_{\text{sample}} / R_{\text{standard}}) - 1] \times 10^3$ , where R is the isotopic ratio of interest ( $^2\text{H}/^1\text{H}$  or  $^{18}\text{O}/^{16}\text{O}$ ). The average precision, based on multiple analyses of various samples and laboratory standards was  $\pm 0.1\text{ ‰}$  for  $\delta^{18}\text{O-H}_2\text{O}$  and  $\pm 0.8\text{ ‰}$  for  $\delta^2\text{H-H}_2\text{O}$ . Tritium activity measurements were done in the BRGM (Orleans, France) by direct liquid scintillation counting. The detection limit was 0.6 TU (Tritium Unit with 1 TU equal to 1 tritium atom in  $10^{18}$  hydrogen atoms).

Sr-isotopes were measured on a VG sector 54 in dynamic mode at the GEOTOP laboratory at the Université du Québec à Montreal (Canada).  $^{87}\text{Sr}/^{86}\text{Sr}$  ratios were normalized to  $^{86}\text{Sr}/^{88}\text{Sr} = 0.1194$ . Repeated analyses of the NIST-987 standard yielded values of 0.710294 ( $\pm 0.000022$ ,  $2\sigma$  reproducibility). Furthermore, these results were corrected to the accepted value for NIST-987 of

0.710248. The  $^{13}\text{C}$  contents were measured using a mass spectrometer (Micromass, Isoprime model with triple universal collectors) at the GEOTOP laboratory at the Université du Québec à Montreal (Canada).  $^{13}\text{C}$  contents are reported using conventional  $\delta$  (‰) notation as a deviation from V-PDB, and the error of  $\delta^{13}\text{C}$  is 0.05‰ (mean error obtained from replicate analyses). The  $\delta^{15}\text{N-NO}_3^-$  and  $\delta^{18}\text{O-NO}_3^-$  values were measured at the University of Waterloo Environmental Isotope Laboratory (Canada).  $\delta^{15}\text{N-NO}_3^-$  values were determined using a Carlo Erba 1108 CNOS Elemental Analyzer coupled to a Fisons Instrument Isochrom-EA mass spectrometer (GV Instruments, Manchester, UK).  $\delta^{18}\text{O-NO}_3^-$  values were measured by breakseal combustion using a VG PRISM Series II mass spectrometer (GV instruments, Manchester, UK). Isotopic ratios are reported in delta ( $\delta$ ) notation in units of per mil (‰) relative to the reference standards of atmospheric  $\text{N}_2$  for  $\delta^{15}\text{N}$  and Vienna Standard Mean Ocean Water (VSMOW) for  $\delta^{18}\text{O}$ . The analytical precision for  $\delta^{15}\text{N-NO}_3^-$  and  $\delta^{18}\text{O-NO}_3^-$  values are  $\pm 0.3\text{‰}$  and  $\pm 0.8\text{‰}$ , respectively. Mineral saturation indices,  $P_{\text{CO}_2}$  and  $\delta^{13}\text{C}$  of aqueous  $\text{CO}_2$  were calculated by PhreeqcI software using *lnl.dat* and *iso.dat* databases (Parkhurst and Appelo 2015; Thorstenson and Parkhurst 2004).

#### 4. Results and discussion

Samples were separated into categories according to i) the regional sub-area and possible sub-compartment considered, namely “Petit Bara”, “Grand Bara”, “Dadin” and “Mouloud” and ii) the aquifer type the borehole corresponded to, mostly “basalt” and “alluvium”. Two additional categories corresponding to boreholes lacking stratigraphic description or in specific sub-areas are proposed, namely “Grand Bara - central” (#12, located in the central Grand Bara plain) and “Petit Bara - central” (#7 and #8, located between Petit Bara and Grand Bara).

## 4.1 Hydrogeochemical pattern

### 4.1.1 Overall geochemical evolution

The geochemistry of analyzed groundwater samples from the study area is detailed in Table 2 and classification of the waters samples in Table 1 was made according to principles of IAH (1979). According to the salinity classification of Kharaka and Hanor (2014), most of sampled groundwaters are brackish ( $10 \text{ g/l} > \text{TDS} > 1 \text{ g/l}$ ) with the exception of several fresh waters ( $\text{TDS} < 1 \text{ g/l}$ ) from North of Petit Bara (#1-#4), Grand Bara (#14) and Dadin (#22). All samples are slightly alkaline (pH from 7.2 to 8.4), close to saturation or oversaturated with respect to calcite and dolomite, and under saturation with respect to anhydrite and halite (Table 2).

The groundwater temperatures in the present study range from 28 to 43°C with an average of about 37°C (Table 1) and are typical for this arid region (Abdoulkader et al., 2013). On the other hand, studied waters may not be considered “hot or thermal”, neither in sense of Pentecost (2005), nor in sense of Schoeller (1962).

As the cation-anion relation  $(\text{Ca}+\text{Mg})-(\text{SO}_4+\text{HCO}_3)$  shows an excellent correlation with electric conductivity in Fig. 3a, where only #12 (center of the Bara plain) and #13 (shallow dugwell in alluviums) outstand, this relation appears as relevant indicator of the degree of geochemical evolution of groundwater. Indeed, the  $(\text{Ca}+\text{Mg})-(\text{SO}_4+\text{HCO}_3)$  vs  $(\text{Na}+\text{K})-(\text{Cl}+\text{NO}_3)$  diagram (Fig. 3b) offers a graphical overview revealing a similar overall evolutionary trend for groundwater issued from both the alluvium and basaltic geological formations. The term evolutionary refers here to both water-rock interaction and mixing. This trend starts with a Na-Cl- $\text{HCO}_3$  water type in the Dadin area (#22) and most alluvium groundwater (#1, #2, #3, #4) (-5 to -4 on the horizontal axis). In this group, the excess of Na compared to Cl and depletion in Mg+Ca compared to  $\text{HCO}_3+\text{SO}_4$  is explained by re-freshening through cation exchange, that typically

occurs when freshwater with high Ca and  $\text{HCO}_3$  flushes sediments that had been in prolonged contact with Na (e.g. Baudron et al., 2016). The geochemical trend ends with a Na-Cl- $\text{SO}_4$  water type in the most mineralized basalt groundwaters, namely #7 and #8 (16 to 19 on the horizontal axis), located in the junction between the Grand Bara and the Petit Bara plains (see below). The origin of these highly mineralized groundwaters will be discussed further in this paper. Samples from #10 and #16 (9 on the horizontal axis) are displayed in-between the intermediate water type and the most mineralized groundwaters, most probably revealing a mixing between them (see Fig. 3c). Finally, the correlation with the mean depth of the well screens and the aforementioned cation-anion relation, as showed by Fig. 3d, shows the vertical component of this geochemical trend inside both aquifers, although samples from boreholes of the central area (#7 and #8), as well as geochemically intermediate samples (boreholes #10 and #16), show a slightly different tendency.

#### 4.1.2 Insights into water origin

To better understand the origin of the sampled waters, anions have been isolated in a ternary diagram (Fig. 4A). According to Fig. 3a, waters evolve from  $\text{HCO}_3$  corner to Cl vertex following a linear trend, also depicted by shallow groundwater interacting with Stratoid basalts (Awaleh et al. 2016). In the same manner, most of previously studied samples coming from surrounding areas are here clustered, but two water samples which interact with evaporites (mainly gypsum; Aboubaker 2012, Aboubaker et al. 2013). Although these sulfate samples were collected by the authors in sedimentary formation westward of the Dikhil region, they show similar effects to those previously highlighted. For example, waters interacting with gypsum do not result in a Ca- $\text{SO}_4$  composition, but Na- $\text{SO}_4$  probably due to Ca-Na exchange with clays and/or mixing with Na-Cl waters followed by precipitation of calcium as  $\text{CaCO}_3$ . Finally, our high salinity samples

deviate from such common path, and point towards a Na-Cl geothermal waters of meteoric origin as Abhe (Fig. 4A), which are compositionally very different from Na-HCO<sub>3</sub> geothermal waters from Main Ethiopian Rift (Püschel et al. 2013; Rango et al. 2009). The Langelier-Ludwig diagram (Fig. 4B) is particularly adapted to shows evolutionary trend and mixing, in particular if Na-HCO<sub>3</sub> waters are involved in the processes as in this case (Boschetti 2011). However, the fluid origin information gained from ternary anion diagram could be confused due to the overlap of water-rock interaction processes which involve mainly cations (Giggenbach 1991). Indeed: i) the Na-HCO<sub>3</sub> towards Na-Cl evolutionary trend of the samples is more scattered in comparison to that depicted in Fig. 4A; ii) the fields of geothermal waters of meteoric origin (Abhe) and seawater-derived (Asal) are interdigitated; iii) differences between Na-SO<sub>4</sub> and Na-Cl is not resolved. To better discern between the cations variations related to high temperature effects, possibly inherited by geothermal waters, and low temperature effects, samples are plotted in the geothermal square plot (Fig. 5A). In this diagram is evident how most of our samples fall on a path which joins two hypothetical fluids (Fig. 5A) corresponding to the isochemical dissolution of Dahla basalt and to the fluid equilibration with rock at shallow temperatures (20-40°C, Giggenbach 1988; Fig. 5A). According to Giggenbach (1995), this low temperature shift of the samples might also involve K-uptake or Na-release from clays. Therefore local geothermal waters, if involved in fluid generation, changed their cationic proportions due water-rock interactions at more supragenetic conditions.

Finally, fluid origin is tentatively inferred by B/Cl and Br/Cl ratios plot (Fig. 5B). In their previous study, Aboubaker et al. (2013) detected a Br/Cl molar ratio range from 0.0045 to 0.0086, excluding in this manner contribution from halite or other seawater evaporites. These ratios are higher than detected in our samples:  $0.0020 \pm 0.0005$  (Table 2). This range includes that of local seawater (Br/Cl = 0.0017; Sanjuan et al. 1990). However, Fig. 5B shows clearly that



most of the sampled waters mimic the wide B/Cl variations detected in other Afar fluids. In particular, our samples are within a ternary mixing which includes basalt (# 14), meteoric-derived geothermal waters (e.g. Abhe; Awaleh et al. 2015) and seawater-derived geothermal waters (e.g. Asal; Sanjuan et al. 1990). Two samples from Bara alluvium show the highest Br/Cl ratios (#13 = 0.0039; #18 = 0.0028; Table 2), which in combination with their B/Cl could be explained by interaction with an Afar evaporite, originated by seawater-derived geothermal fluid (Fig. 5B).

#### 4.1.3 Origin of CO<sub>2</sub>

One surprising feature is how bicarbonate severely decreases along the flow path (Fig. 4c): from 6 to 7 meq/l in the less mineralized groundwater sample of the Dadin area (#22) and of Petit Bara alluvium (#2 and 3) to 1 meq/l in the most evolved groundwater samples, located in the central area (#8). As well, logPCO<sub>2</sub> is found to decrease from -1.7, typical of soils, down to -3.1, while pH increases from 7.26 to 8.38 (see Table 1). Atmospheric CO<sub>2</sub> and the subsequent enrichment in the soil layer is therefore a major source of HCO<sub>3</sub> in the aquifer system, although other subsurface sources of HCO<sub>3</sub> like the release of HCO<sub>3</sub> following water-rock interaction or a geothermal enriched CO<sub>2</sub> cannot be misconsidered (Awaleh et al. 2016 and reference therein). Geothermal waters in Djiboutian Afar and Main Ethiopian Rift have similar mantle-like  $\delta^{13}\text{C}(\text{CO}_2)$  of approximatively -3.5 ‰ (e.g. Demlie et al., 2008, Alemayehu et al., 2011; Awaleh et al. 2016); however, the former have acid pH and no-HCO<sub>3</sub> in comparison to the alkaline high HCO<sub>3</sub> of the latter (Awaleh et al. 2016; D'Amore et al. 1998; Teklemariam and Beyene 2001). As illustrated by Fig. 6a, studied samples evolve from -18.5‰ vs PDB (at a HCO<sub>3</sub> value of 6.6 meq/l) which is quite close to the value of soil CO<sub>2</sub> (e.g. Cerling et al. 1991), to -12.7‰ vs PDB at 1.8 meq/l. The high HCO<sub>3</sub> – low  $\delta^{13}\text{C}(\text{CO}_2)$  end-member could be due to the weathering in

open system CO<sub>2</sub> condition (e.g. Clark 2015). Differently, taking into account the alkaline pH of the sampled waters and their saturation indexes (Table 1 and Table 2), the low HCO<sub>3</sub><sup>-</sup> - high  $\delta^{13}\text{C}(\text{CO}_2)$  side could be originated by calcite equilibrium/oversaturation rather than deep CO<sub>2</sub> contributions.

#### 4.1.4 Origin of nitrate

It is worth noting that all boreholes are used for drinking purpose, although for many of them, NO<sub>3</sub><sup>-</sup> contents lie around the maximum permissible value prescribed by the World Health Organization standard set for drinking waters (50 ppm; WHO, 2011; see Table 2). The lowest values are found in Petit Bara and Dadin (Fig. 7a), while the highest values are found in Grand Bara (up to 250 mg/l in alluvium groundwaters from boreholes #13 and #18; up to 150 mg/l in basalt groundwaters from #11 and #12) and in the central area (up to 120 ppm in #7 and #8). It should be noted that people living in the Republic of Djibouti are mostly nomad, and hence have no agricultural tradition. Furthermore, the local nomads, who raise goats, cheeps, cows and camels, do not have a fixed settlement, hence not septic tanks. Therefore, in many cases, natural soil might contribute mainly for nitrate concentrations in the study area as observed in some semi-arid or arid regions over the world (Stadler et al., 2008; Walvoord et al., 2003). Exceptions concerned samples from boreholes like #13 and #18 whose nitrate content might feature anthropogenic forcing (Fig. 7a). To better constrain its origin we used the isotopic composition of nitrate ( $\delta^{15}\text{N}-\text{NO}_3^-$  and  $\delta^{18}\text{O}-\text{NO}_3^-$ ). As shown in Table 3 and illustrated by Fig. 7b, most values are located in a narrow range between 6.47 and 9.31 ‰ vs AIR for  $\delta^{15}\text{N}-\text{NO}_3^-$  and 7.15 and 12.74 ‰ vs VSMOW for  $\delta^{18}\text{O}-\text{NO}_3^-$  (Table 3). According to published N-isotope ranges

(Kendall, 1998; Kendall et al., 2007; Aravena and Mayer, 2010), these groundwaters may all be affected by either sewage/manure and/or soil nitrogen. Taking into account the local oxygen isotope effect during nitrification (2/3 of O from H<sub>2</sub>O between -4 and -1 from aquifer and +6‰ from evaporated rain, see next paragraph, and 1/3 from atmospheric  $\delta^{18}\text{O}-\text{O}_2 = +23.5\text{‰}$ ; Kendall et al., 2007) and denitrification, the high nitrate concentration of borehole #18 (255 mg/l) could be derived from a denitrified manure source. However, because of the overlapped isotopic values of reduced nitrogen precursors (Fig. 7b) and the not well defined fractionation effects on oxygen during nitrification (Casciotti et al., 2011), nitrate isotopes can hardly differentiate different sources (Fenech et al., 2012). Anyway, considering the similar trend of #18 and #13 in Fig. 7a, the same sources could be hypothesized for the shallow borehole #13 (19m deep). Similarly, the dual isotopes ( $\delta^{15}\text{N}-\text{NO}_3^-$  and  $\delta^{18}\text{O}-\text{NO}_3^-$ ) are not suitable in differentiating closely related sources of nitrate contamination, such as sewage and manure, for Mouloud aquifer (boreholes #19 and #20) (Fig. 7b). These nitrate and chloride contents of the basalt aquifer in the studied area are very similar to the mean value revealed in the bordering Dalha basalt aquifer (Aboubaker, 2012; Fig. 7a). On the other hand, the  $\delta^{15}\text{N}-\text{NO}_3^-$  and  $\delta^{18}\text{O}-\text{NO}_3^-$  of boreholes #9, #11, #14, #17, and #22 indicate that the nitrate in those groundwaters is mainly derived from naturally occurring soil (Fig. 7b).

The #7 borehole, drilled in 2014 in the south of Wea valley, displays high nitrate content (119 mg/l) and appears as an outlier due to its high  $\delta^{18}\text{O}-\text{NO}_3^-$  value of 16.8 ‰, that places it in the typical desert field trend as depicted by Jackson et al. (2015). Hydrogeochemical and isotopic findings further described in this paper evidenced that its mineralization was influenced by palaeowater of geothermal origin. Likely, the deep nitrogen signature of  $\delta^{15}\text{N} = -5\text{‰}$  (e.g. Li et al. 2016) was obliterated by paleoenvironmental factors related to former vegetation cover (e.g;

Stadler et al., 2008) and/or accumulation of atmospheric nitrate in caliche and calcrete, which given their imprint for concomitant high  $\text{NO}_3\text{-Cl}$  contents and  $\delta^{18}\text{O-NO}_3^-$  values to the recharging waters, as observed in several semi-arid and arid regions of the world (Fig. 7b; Al-Taani and Al-Qudah, 2013; Edmunds and Gaye, 1997; Jackson et al., 2015; Walvoord et al., 2003). As future perspective, and according to Dietzel et al. (2014) and Jackson et al. (2015), the analysis of perchlorate and  $^{17}\text{O}$  in nitrate would be better clarify and trace the cross contamination between different nitrate sources.

#### 4.1.5 Water isotope and recharge conditions

The oxygen  $\delta^{18}\text{O}$  and hydrogen  $\delta^2\text{H}$  stable isotopes are useful tracers for determining the origin of groundwater and are widely used in studying the natural water circulation and groundwater movement. One main task before interpreting such tracers is to set up a meteoric water line that will serve as a reference regarding fractionation and mixing processes. Since no systematic isotopic dataset concerning rainfall is available for Djibouti, most local studies choose the local meteoric water line (LMWL) from Addis Ababa station (e.g. Adam and Fontes, 1982; Aboubaker et al., 2013). Nonetheless, Addis Ababa is separated from the study area by more than 600 km of distance and is located almost 1800 m higher in altitude (2355 m a.s.l compared to around 600 m a.s.l). Furthermore, the study area is drier and hotter (about  $150 \text{ mm.y}^{-1}$  and  $34^\circ\text{C}$ , respectively) than Addis Ababa ( $> 2500 \text{ mm.y}^{-1}$  and  $16.5^\circ\text{C}$ , respectively). Therefore, data from Addis Ababa might not provide realistic data for the study area. Local rainfall data from Djibouti is scarce (Adam and Fontes, 1982; Houmed-Gaba 2009) but provides interesting insights in accordance with a deuterium excess value of  $d = 2$  (Fig. 8). This latter value coincides with that of the LMWL proposed by Fontes et al. (1979) and with a fractionation effect caused by secondary

evaporation of the falling raindrops leading to a deuterium excess in rainfall, originated from both Atlantic monsoon (“pseudo-altitude effect”; Kebede, 2004) and from Indian monsoons (Pang et al., 2004). In the present study, despite the very little number of isotope measurements in Djibouti rainfall and to the high temporal variability of rainfall isotope composition in arid climate, we selected the aforementioned LMWL from Fontes et al. (1979), based on three rainfall samples, as the most probable orientative proxy of the isotopic composition of local precipitation, although it cannot be used as an absolute reference for evaporative effects.

As illustrated by Fig. 8, groundwater samples are mostly located along a line parallel to both the global meteoric water line (GMWL) and the LMWL, with a deuterium excess value of approximately 6. Therefore, no evaporation effect is evidenced, although sample from borehole #7 is located below this line and might reveal a slight fractionation process. Two samples from Fontes et al. (1980) are also located below this line, but it could simply be partly due to the difference between analytical laboratory techniques. Previous sections revealed the common geochemical evolution pattern of alluvium and basalt groundwaters. By difference, the  $\delta^2\text{H}$  and  $\delta^{18}\text{O}$  composition of groundwater (see Table 3 and Fig. 8) show that alluvium and basalt groundwater samples can be separated into two groups. The first cited have  $\delta^{18}\text{O}$  values ranging from -1.9 to -1.1 ‰, i.e. in the range of wadi-recharged aquifers according to Fontes et al. (1979) and Awaleh et al. (2015) and in the range of modern precipitation for the area, according to Clark and Fritz (1997). Basalt groundwaters range between -3.9 and -2.6 ‰, rather close to Lake Abhe hot springs (Fontes et al., 1980; Awaleh et al., 2015). Two outsiders, alluvium groundwater from borehole #10 (northwestern Grand Bara) and basalt groundwater from borehole #22 (Dadin area), are located in the respective opposite categories. The  $\delta^2\text{H}$  and  $\delta^{18}\text{O}$  difference between alluvium and basalt groundwater can be explained by two hypotheses, or their combination: (i) recharge

water is isotopically depleted in the highest basaltic outcrops (up to 1000 m a.s.l) compared to the plains (down to 600m a.s.l) because of altitude effects and (ii) recharge to the basalt groundwater occurred under a past climate, with cooler temperatures than present (e.g. Jirakova et al., 2011, Baudron et al., 2014). A  $\delta^{18}\text{O}$  altitudinal gradients for Djibouti can be calculated based on the one proposed by Kebede et al. (2008) for Ethiopia: 0.1‰ per 100m. In such case, a 0.4‰ difference between alluvium and basalt groundwater would be found, much less than the mean difference between both (around 2.4‰) and rather in the range of variability inside one category. An altitude gradient value of 0.6‰ per 100m would be required to support such difference. Nonetheless, it would represent a very high value compared to global literature, while Levin (2008) considered the altitude effect to be mostly muted in Eastern Africa. Therefore, in addition to a possible slight altitude effect, the basalt groundwater is considered to be recharged under a past climate, in comparison to a recharge by more recent precipitation for the alluvium. Meaningful information to this respect is provided by time tracers in the next section.

#### **4.1.6 Groundwater residence time**

The only available tritium data for Djibouti precipitation are in the mid-eighties (Grand Bara area:  $7.3 \pm 0.7$  TU; Dikhil area:  $8.6 \pm 1.3$  TU and Mouloud area  $< 2.3$  TU) (BGR, 1982). Coastal/low latitude tritium table of Clark and Fritz (1997) was therefore used to distinguish between modern and sub-modern (pre-bomb) waters. In the mid-seventies and early-eighties, the basalt groundwaters from #20 (Mouloud area), #16 (Grand Bara) and #12 (central Grand-Bara) boreholes displayed no measurable tritium ( $< 4\text{T.U}$  in Fontes et al., 1980 and  $< 2.2\text{T.U}$  in BGR, 1982). By difference, alluvium groundwater from #3 (Petit Bara area) featured in 1975 up to 7 T.U. (Fontes et al., 1980) and therefore revealed a significant modern recharge component.

Updated tritium data covering a wider spatial scale (11 samples) is brought by the present study. Similarly to previous campaigns by Fontes et al. (1980) and BGR (1982), no measurable tritium was found in any of the basalt groundwaters ( $<0.6$  T.U), sampled in five boreholes from both the Mouloud and Grand Bara area. By comparison, tritium was detected in three of the six alluvium groundwater samples, issued from both the Petit Bara (#2 and #6 boreholes; 0.7 and 1.2 T.U, respectively) and Grand Bara areas (#13 borehole; 0.8 T.U). These results confirm an elder age for basalt groundwater than for the alluvium, where significant recharge from modern precipitation occurs.

In the early eighties, six groundwater samples were collected for radiocarbon and  $^{13}\text{C}$  analysis by Fontes et al. (1980) and BGR (1982). They were collected from four basalt boreholes from the Mouloud, Grand Bara and central Grand Bara area and from one alluvium borehole from the Petit Bara area (Table 3). Since little detail was provided on the assumptions related to the correction of radiocarbon ages, they were recalculated in the present study. Calculating groundwater ages requires taking into account the initial composition of soil gas and correcting the measured activities with respect to water-rock interactions (e.g. Fontes, 1992; Baudron et al., 2013) if necessary. The progressive enrichment in  $\delta^{13}\text{C}$  with decreasing  $A^{14}\text{C}$  (Fig. 5a) appeared as an indicator of isotopic exchange with rock-matrix carbonate (e.g. Gillon et al., 2009). Therefore, the IAEA (Gonfiantini, 1972; Salem et al., 1980) age correction method was plebiscited to simpler methods like Tamers (Tamers, 1967). In fact, Table S1 (see supporting information) reveals irrelevant results for most data from other correction methods when using the criteria mentioned below. The  $\delta^{13}\text{C}$  and  $A^{14}\text{C}$  of the solid carbonates were set up to values of 0. Regarding the  $\delta^{13}\text{C}$  of soil gas, Table 3 suggests an effect of both C3 and C4 plants, as supposed by Bocherens et al. (1996). Since no precise information was available on their respective location regarding recharge processes (e.g. basaltic outcrops, alluvium), a  $\delta^{13}\text{C}(\text{CO}_2)$  value for

soil gas of  $-21\text{‰}$  vs PDB was set, as also proposed by Fontes et al. (1980) and close to the most depleted  $\delta^{13}\text{C}(\text{CO}_2)$  calculated in groundwater (Table 3). On this basis, indicative rounded radiocarbon groundwater ages of 6250 years BP (before present, borehole #16 from the Grand Bara basalts), 2250 and 3100 years BP (boreholes #20 and #21 from the Mouloud area) and 10100 years (borehole #12 from the central Grand-Bara area) are obtained for basalt groundwater. By difference, alluvium groundwater (#3) age is 23 years BP (i.e., modern, sampling by Fontes, 1980) and 1050 years BP (sampling by BGR, 1982). The last value is probably impacted by intra-borehole mixing with older water, since this borehole features a long screen catching both alluvium and basalt and its geochemical and isotopic composition varies along time between both end-members (CERD, 2014). Based on the calculated radiocarbon groundwater ages, an important modern (post nuclear bomb tests) component of recharge is confirmed in the alluvium groundwater, while recharge is dated between the early and mid-Holocene in the basalt. These radiocarbon results only refer to a few discrete samples. Nonetheless, since  $\delta^{13}\text{C}(\text{CO}_2)$  is linked to both decreasing  $\text{HCO}_3^-$  (Fig. 6b) - as an indicator of the geochemical evolution of groundwater - and  $\delta^{18}\text{O}$  (Fig. 6c) - as an indicator of modern and old recharge -  $\delta^{13}\text{C}$  might be used as a qualitative indicator of the evolution of groundwater age, thus generalizing the groundwater age pattern to the whole study area.

## 4.2 Insights into groundwater flow pattern

### 4.2.1 Overall groundwater flow pattern inside basalt groundwater

The #15 – #17 boreholes from the South of the Grand Bara area present comparable isotopic contents ( $-3.7\text{‰} \leq \delta^{18}\text{O} \leq -3.1\text{‰}$ ;  $-24.8\text{‰} \leq \delta^2\text{H} \leq -18.8\text{‰}$ ) to boreholes tapping the Mouloud volcanic aquifer (#19 and #20). Besides, groundwater samples from boreholes #15 – #17 and #19



– #20 have almost the same Na–Cl–Mg–SO<sub>4</sub> facies (Table 1) and are located in the same range of Ca+Mg-(SO<sub>4</sub>+HCO<sub>3</sub>) values (Fig. 3a). Therefore, the Grand Bara and Mouloud basalt aquifers are considered to be interconnected, forming one single volcanic aquifer (namely Grand Bara – Mouloud aquifer). Furthermore, according to potentiometric data (Fig. 2; Table S2 see supporting information), the moderately mineralized groundwaters from the Grand Bara – Mouloud area flows towards the North of the Grand Bara plain where the most mineralized groundwaters (#8, #12) are localized (Fig. 1). The high mineralization found in the central area (#8), i.e. the subterranean outlet of this endorheic plain, located at the geological threshold that separates Grand Bara and Petit Bara plains, is therefore most probably linked to the high residence time of groundwater and a subsequent higher level of water-rock interaction. A slight contribution of hydrothermal exchange with the host-rock also probably occurred for this sample (Fig. 8). Although they do not reach such high electric conductivity and show slightly different mineralization pattern, water sampled from the Abhe hot springs (Awaleh et al., 2015) are other examples of long-term water-rock interaction in the basalt formations, including a similar isotopic component of hydrothermal exchange with the host-rock (Awaleh et al., 2015).

#### 4.2.2 From wadi-recharged alluvium to basalt aquifers

Continuity at the scale of the study area between the alluvium and basalt aquifers in terms of geochemical evolution and residence time of groundwater was shown in previous sections. Potentiometric levels bring complementary information regarding such connection. On the southwestern side of the Grand Bara plain (Mouloud), boreholes screened in an alluvial deltaic formation (#23, #24, #25) show higher potentiometric levels (by 20m, see Table S2) than deeper ones screened in the underlying basalt (#20, #26, #27). Such downward potentiometric relation is also observed in narrower wadi valleys like the one where is located #13, from the alluvium (#13)

to the basalt (#14). There, the difference between the geochemical and isotopic signatures of both aquifer types is also clearly visible. As well, groundwater from Dadin basalts, located along a wadi-valley, features the isotopic and geochemical signature of modern wadi recharge water and therefore provides a confirmation of the connection between both compartments proposed by Aboubaker et al. (2013) at this location. This data demonstrates i) that recharge to alluvium is consistent in wadi valleys and in deltaic formation on the sides of the Grand Bara plain and ii) that it is transmitted downwards to the basaltic aquifers, probably through major faults.

Regarding the central Grand Bara plain, no direct interpretation can be made on the intensity of recharge processes to the alluvium and their transmission to basalt groundwater, since no data from the alluvium groundwater quality and hydrodynamics is available. Still, since the hydrology of Grand Bara plain is controlled by ephemeral floods caused by extreme rainfall events on a clayey surface, any significative contribution of recharge from the alluvium to the basalt groundwater would involve high evaporated stable isotopes of water. Nonetheless, as illustrated by #12, such feature is not found, while a high degree of geochemical evolution and thus long residence time is evidenced in basalt groundwater. Therefore, recharge from the alluvium in the central Grand Bara area is most probably insignificant, although it is consistent in the wadi valleys.

#### **4.2.3 Refreshening by modern recharge at Petit Bara**

The flowpath from Petit Bara plain to the Didjan Der valley, located northwestward and acting as its main outlet, is characterized by a continuous refreshening from 6.5 mS/cm (#7) to 2.8-2.9 mS/cm (#6 and #5), to around 1 mS/cm (#4, #3, #2) and below 1 mS/cm (#1), as shown in Table 1. Such process is confirmed by the  $\text{Ca}+\text{Mg}-(\text{SO}_4+\text{HCO}_3)$  proxy of geochemical evolution (Fig. 9a) that decreases from values higher than 15 meq/l (#7) to slightly above 0 meq/l (#6 and #5)

and finally around -5 meq/l (#4, #3, #2, #1). A similar tendency of decreasing evolution is observed with isotopic tracers  $\delta^{18}\text{H}$  (Fig. 9b) and  $\delta^{13}\text{C}(\text{DIC})$  (Fig. 9c) whose compositions evolve from the isotopic signature of basalt groundwater (-3.7 ‰ vs V-SMOW and -12.7 ‰ vs PDB, respectively) to the one of freshly recharged alluvium groundwater (-1.1 vs V-SMOW and -18.3 ‰ vs PDB, respectively). As a consequence, recharge from the Didjan Der valley is consistent and induces a refreshing process of groundwater flowing out from the Petit Bara plain. This feature is an additional evidence of the consistent recharge occurring in the riverbed of wadi rivers as well as it provides insightful information on groundwater flowpath in the Petit Bara area. Regarding the salinity increase of the #6 groundwater over time (CERD, 2014), it should be due to an overexploitation that may bring back #7 or #8 highly mineralized waters by upcoming (Lorenzen et al., 2011), as proposed by Fig. 3d.

#### 4.2.4 Strontium isotope ratio: complementary insight on groundwater origin

The  $^{87}\text{Sr}/^{86}\text{Sr}$  ratio measured in water reflects the distinct isotopic composition of the interacting rocks and can furnish information on groundwater flowpaths and hydrodynamics (Drever, 1997; Barbieri et al., 2005). The  $^{87}\text{Sr}/^{86}\text{Sr}$  isotopic ratios of the investigated groundwater samples range from 0.70556 to 0.70694 (Table 3). These data are compared with the  $^{87}\text{Sr}/^{86}\text{Sr}$  values of local lithotypes available in the literature in order to define the nature of the interacting rocks. The reported Sr isotopic values of basalt samples from the Republic of Djibouti show considerable variation from 0.70350 to 0.70664, with the lower value corresponding to the most recent Afar mantle plume (Vidal et al., 1991; Barrat et al., 1993; Deniel et al., 1994; Rogers, 2006). Barrat et al. (1993) reported that the pre-Dalha basalts, which are often weathered, show a large variation in  $^{87}\text{Sr}/^{86}\text{Sr}$  ratios from 0.7037 to 0.7067. On the other hand, for the non weathered basalts of the

Dalha series (8.3-4.3 Ma), the mean Sr isotopic composition is quite uniform:  $0.70411 \pm 0.00047$  (Barrat et al., 1993; Deniel et al., 1994; Vidal et al., 1991). Almost all the investigated groundwater lies outside the non weathered Dalha basalt range, suggesting contributions from more radiogenic rocks along the groundwater flow path, for example: i) lacustrine sediments; ii) weathered Dalha basalt or more radiogenic rocks (e.g. rhyolites). Some evidences support this latter hypothesis. For examples, drilling logs revealed that most of the Dalha basalts in the study area are weathered. Moreover, the highest  $^{87}\text{Sr}/^{86}\text{Sr}$  values of the water samples were revealed in the southern side of the Bara plain (boreholes #22, #13, #14, #11). It is reasonable to assume that some amounts of the radiogenic strontium in the local weathered basalt or alluvium come from the older Mablas rhyolite (up to 0.722; Deniel et al., 1994).

Most of the  $^{87}\text{Sr}/^{86}\text{Sr}$  of dissolved Sr in Djibouti (SanJuan et al., 1990; Dekov et al., 2014; Awaleh et al., 2015; Awaleh et al., 2016) pertain to geothermal waters issuing from Stratoid and Asal rift basalts (which have not so different strontium isotope ratio from Dalha basalt, Faure 2001; Fig. 10a) which have a relatively low  $^{87}\text{Sr}/^{86}\text{Sr}$ , testifying a full or partial equilibration with the source rock (Fig. 9a; Awaleh et al., 2015; Sanjuan et al., 1990; unpublished data). Exceptions include shallow groundwater (Awaleh et al., 2016) and waters interacting with alluvial deposits. As an example of this latter case, Oued-Kalou springs near to Lake Asal area show similar composition to the waters from this study (Fig. 10a; Sanjuan et al. 1990). Moreover, the statistically significative linear relationship between sulfate and strontium of the Bara groundwater testify the big effect of alluvial component on the Sr isotope ratio (Fig. 10b). Therefore, the high  $^{87}\text{Sr}/^{86}\text{Sr}$  ratios observed for the investigated groundwater from volcanic aquifers might be related to different contributions: not only Dalha basalt and its alteration products in this study area, but also the recharge of the volcanic aquifers through major faults by

waters interacting with alluvial deposits which mediate the contribution of surrounding formations.

## 5. Conclusion

This integrated study aimed at revealing the recharge conditions, groundwater age distribution, origin of contaminants and groundwater flow pattern inside the complex volcanic aquifer system of Bara (Djibouti). To this end, an integrated hydrodynamic, geochemical and isotopic approach was set up. Geochemistry showed an evolutionary trend from alluvium aquifers located in wadi-valleys and deltaic formations bordering the main sedimentary basins (Petit Bara and Grand Bara), with low salinity groundwater and Na-Cl-HCO<sub>3</sub> water type, to a more saline and intermediate water type in most basalt groundwaters (Grand Bara and Mouloud area) and finally to a Na-Cl-SO<sub>4</sub> water type in the most mineralized groundwaters located at the border between Grand Bara and Petit Bara. Elementary and isotopic nitrate evidenced anthropogenic and geogenic origins of nitrate. Stable isotopes of water revealed modern recharge in the alluvium through infiltration of precipitation in wadi aquifers and revealed significantly depleted values in the second. No evaporation effects were evidenced, since most groundwater samples plotted on a  $\delta^{18}\text{O}$  vs  $\delta^2\text{H}$  line parallel to both global and local precipitation lines. Modern radiocarbon and tritium were evidenced in the alluvium groundwaters, while recalculated radiocarbon ages located recharge of the basalt groundwaters in the early to mid-Holocene. These features revealed a common evolutionary pattern, with a significant recharge from the alluvium, transmitted downward to the basalt aquifer through major faults in the wadi valleys or through the sediments of the alluvial fans, followed by mixing with ancient Na-Cl water. This latter endmember with depleted content of  $^2\text{H}$  and  $^{18}\text{O}$  isotopes could be a deep and old geothermal water from the

Stratoid basalts, then cooled and evolved after interaction with Dalha basalts. This study also revealed strategic characteristic of the groundwater flow pattern. Two major basaltic aquifer compartments, the Grand Bara and Mouloud aquifers, were found to form one single aquifer flowing northeastward below the Grand Bara endorheic plain. Highly mineralized groundwater flowing out from the Petit Bara basin was found to be rapidly diluted by modern recharge in the alluvium of the Didjan Der valley. In addition to providing significant insights for a groundwater management of the Bara area, this study will also serve as a valuable base for other integrated hydrodynamic, geochemical and isotopic studies of volcanic aquifers in arid areas.

### **Acknowledgements**

This research work was financially supported by the United Nation Development Program (UNDP), the Centre d'Etudes et de Recherche de Djibouti (CERD), and the International Foundation for science, Stockholm, Sweden, through a grant N° W/5800-1. We would like to thank Mr Aden Atteyeh for fruitful discussion and Mr Samaleh Ahmed Idriss and Abdi Abdillahi Djibril for their assistance in the field works. We would also like to thank two anonymous reviewers for their constructive comments that improved the manuscript.

### **Table and Figure captions**

**Table 1.** Sampling locations, T, depth, pH, EC, TDS and hydrochemical types of the sampled waters.

**Table 2.** Hydrochemical characteristics (in mg/l) of water samples in the study area and saturation index (SI) of selected minerals.

**Table 3.** Isotope data of Bara groundwaters from this study and literature.

**Figure 1.** Geological map of the study area (EAR = East African Rift).

**Figure 2.** Geological cross-sections in Petit Bara (a), Mouloud-Grand Bara (b) and Mouloud (c) areas.

**Figure 3.** Variations of EC (a), (Na+K)-(Cl+NO<sub>3</sub>) (b), HCO<sub>3</sub> (c) and half depth of the screen (d) versus the (Ca+Mg)-(HCO<sub>3</sub>+SO<sub>4</sub>) indicator of geochemical evolution.

**Figure 4.** Anions ternary diagram, mg/l basis (a) and Langelier-Ludwig square plot, meq/l basis (b). Volcanic, geothermal and peripheral/shallow fields are from Giggenbach (1988); dotted field depicts samples collected by Aboubaker (2012) and Aboubaker et al (2013); fields with different hatches distinguish meteoric-derived (Abhe; Awaleh et al. 2015) and seawater-derived (Sakalol, Asal; Awaleh et al. 2016) geothermal waters, respectively.

**Figure 5.** (a) Geothermal square plot (Giggenbach 1988): dashed lines describe the evolution of the waters from isochemical dissolution of Dahla basalt (Mohr and Zanettin 1988) towards low temperature equilibrium with average crust (full equilibrium line, numbers are temperature in Celsius degrees; Giggenbach 1988). Dotted field as in Fig.4. (b) B/Cl vs. Br/Cl diagram. Agriculture drainage, sewage and seawater evaporites fields are drawn according to Vengosh (2014). Afar Hydrothermal waters describes waters of meteoric origin, whereas mean  $\pm$  standard deviation crosses depict geothermal or regional waters from Afar with variable seawater contributions (Awaleh et al. 2016, Lowenstern et al. 1999, Sanjuan et al. 1990; UNDP 1973). Afar evaporites (Binega 2006, Martini 1977), Basalts (Awaleh et al. 2016) and Main Ethiopian Rift (M.E.R.; Gianelli and Teklemariam 1993, Pittalis et al. 2016) fields are also shown for comparison.

**Figure 6.** Variations of  $\text{HCO}_3^-$  (a),  $\delta^{14}\text{C}$  (b) and  $\delta^{18}\text{O}$  (c) versus  $\delta^{13}\text{C}-\text{CO}_2$ . Samples with a circle are taken from the literature (Fontes et al., 1979; BGR, 1982). In (b), rounded mean apparent ages calculated with the IAEA correction method.

**Figure 7.**  $\delta^{15}\text{N}-\text{NO}_3^-$  versus  $\delta^{18}\text{O}-\text{NO}_3^-$  diagram (a) of the groundwaters from the study area, in comparison with nitrates derived from typical: i) N sources after nitrification by local waters (Aravena and Mayer 2010; Kendall 1998; Kendall et al., 2007); ii) desert nitrates (Jackson et al., 2015). Nitrate vs. chloride plot (b).

**Figure 8.**  $\delta^{18}\text{O}$  versus  $\delta^2\text{H}$  diagram of the groundwaters from the study area showing the global meteoric water line (GMWL after Craig, 1961) and local meteoric water line (LMWL after Fontes et al. 1979). d accounts for the deuterium excess. Samples with a circle are taken from the literature (Fontes et al., 1979; BGR, 1982; Aboubaker et al., 2013).

**Figure 9.** Variations of  $\text{Ca}+\text{Mg}-(\text{HCO}_3^-+\text{SO}_4^{2-})$  (a),  $\delta^{18}\text{O}$  (b) and  $\delta^{13}\text{C}-\text{CO}_2$  (c) along the flowpath from Petit Bara plain to Didjan Der valley. Samples with a circle are taken from the literature (Fontes et al., 1979; BGR, 1982).

**Figure 10.** Relationship between  $^{87}\text{Sr}/^{86}\text{Sr}$  and  $1/\text{Sr}$  (a). Curves depict the best fits of hot springs issuing from active geothermal systems from Asal rift (“marine” curve; data from Sanjuan et al., 1990) and Stratoid basalts (“meteoric” curve; Abhe waters: Awaleh et al., 2015; Awaleh et al. 2016). It is noteworthy that values detected in the Main Ethiopian Rift (Rango et al. 2010) fall on the high  $1/\text{Sr}$  - high  $^{87}\text{Sr}/^{86}\text{Sr}$  range of the “meteoric curve”. Open circles: hot springs from alluvial deposits, SW of Lake Asal (Oued-Kalou; Sanjuan et al., 1990). Strontium isotope ratios of Quaternary basalts and Afar rift are from Faure (2001) and Rooney et al. (2012). Strontium vs. sulfate plot (b), with an additional sample from Oued-Kalou (Bosch et al., 1977).



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Table 1. Sampling locations. T, Eh, depth, pH, EC, TDS, calculated partial pressure and hydrochemical types of the sampled waters.

Samples	Latitude	Longitude	T (°C)	Eh (mV)	Depth (m)	DO (mg/l)	Filters depth (m)	EC (μS/cm)	TDS (mg/l)	logP <sub>CO2</sub>	pH	Local categorie	Hydrochemical types
<i>Petit Bara Zone</i>													
1 PK44	11.44416	42.81727	38	229	135	4.4	45-65; 85-110	956.9	666	-2.2	7.8	Alluvium - Petit Bara	Na-HCO <sub>3</sub> -Cl
2 PK48	11.43917	42.80333	36	238	133	4.3	40-64; 83-116	1099	688	-2.12	7.6	Alluvium - Petit Bara	Na-HCO <sub>3</sub> -Cl
3 PK50	11.43761	42.78342	38	231	92	4.1	60-90	1127	795	-2.29	7.9	Alluvium - Petit Bara	Na-HCO <sub>3</sub> -Cl
4 PK51	11.42322	42.76103	39	241	132	4.5	69-78; 100-118; 120-129	1204	811	-2.14	7.7	Alluvium - Petit Bara	Na-HCO <sub>3</sub> -Cl
5 Didjander	11.39842	42.75795	28	236	170	5.2	55-155	2876	1617	-2.57	7.9	Alluvium - Petit Bara	Na-Cl
6 Omar Jakka	11.3815	42.75508	38	228	160	3.4	100-130	2958	2015	-2.44	7.8	Alluvium - Petit Bara	Na-Cl
7 Naheel	11.35122	42.75233	39	234	155	6	100-140	6564	3668	-2.24	7.2	Alluvium - Petit Bara	Na-Cl-Mg
8 Kourtimaley ♠	11.33	42.67222	37	-	161	-	120-155	7731	-	-3.08	7.8	Central area	Na-Cl-Ca
<i>Grand Bara Zone</i>													
9 Iskoutir	11.30153	42.69911	32	232	181	5.5	80-103 ; 136-160; 162-171	3258	2363	-2.82	7.8	Basalt - Grand Bara	Na-Cl-Ca-SO <sub>4</sub>
10 Dagahdamere	11.28267	42.6963	39	237	168	6.2	55-120; 130-155	4846	3484	-2.73	7.7	Basalt - Grand Bara	Na-Cl-Mg-Ca-SO <sub>4</sub>
11 Gor Galo	11.25814	42.71881	43	219	114	6.7	65-70; 80-90; 100-106	2871	1879	-2.42	7.8	Alluvium-Grand Bara	Na-Cl-SO <sub>4</sub>
12 Goroja ♥	11.24917	42.66389	42	-	90	-	60-85	6300	-	-2.66	7.8	Basalt - Central Grand Bara	Na-Cl-SO <sub>4</sub>
13 Hambocto F2	11.20783	42.67158	35	220	19	6.1	-	1896	1329	-1.96	7.3	Alluvium-Grand Bara	Na-Cl-Mg-Ca-HCO <sub>3</sub>
14 Hambocto F1	11.21311	42.66786	38	270	150	4	70-125	1464	988	-1.87	7.3	Basalt - Grand	Na-Cl-Mg-

15 Doudoub Bololé F2	11.21308	42.65172	36	216	103	4.5	65-88	2337	1576	-2.36	7.6	Bara Basalt - Grand Bara	Ca-HCO <sub>3</sub> Na-Cl-Mg- SO <sub>4</sub>
16 Doudoub Bololé F1	11.21483	42.64633	41	197	93	2.4	58-93	4143	3173	-2.45	7.5	Basalt - Grand Bara	Na-Cl-Mg- SO <sub>4</sub>
17 Awraoussa 1	11.19594	42.61178	36	217	195	4.8	114-152; 166- 182	2692	1842	-2.65	7.8	Basalt - Grand Bara	Na-Cl-Mg- Ca-SO <sub>4</sub>
18 Gabla-galan	11.29444	42.58325	30	165	156	4.6	90-144	2200	1519	-3.07	8.4	Alluvium-Grand Bara	Na-Cl- HCO <sub>3</sub> -SO <sub>4</sub>
<i>Mouloud- Dadin Zone</i>													
19 Mindil	11.20667	42.43175	36	205	150	4.6	65-120	2630	1824	-2.03	7.5	Basalt - Mouloud	Na-Cl-Mg- SO <sub>4</sub>
# Mouloud 3	11.16525	42.50456	39	227	115	6.1	72-106	3035	2075	-2.19	7.5	Basalt - Mouloud	Na-Cl-Mg- SO <sub>4</sub>
21 Mouloud 4	11.1654	42.50442	39	230	140	5.7	100-122	2880	1965	-2.02	7.3	Basalt - Mouloud	Na-Cl-Mg- SO <sub>4</sub>
# Dadin 3	11.14511	42.52781	34	262	139	4.7	36-50; 60-80; 96-134	1388	980	-1.72	7.4	Basalt - Dadin	Na-Cl-HCO <sub>3</sub>

\*Closed after drilling in 1985 because of the high salinity of its groundwater; \*Closed after drilling in 1978 because of the high salinity of its groundwater.

**Table 2.** Hydrochemical characteristics (in mg/l) of water samples in the study area and saturation index (SI) of selected minerals

	Site	Ca (mg/l)	Mg (mg/l)	Na (mg/l)	K (mg/l)	Li (mg/l)	NH <sub>4</sub> (mg/l)	HCO <sub>3</sub> (mg/l)	Cl (mg/l)	SO <sub>4</sub> (mg/l)	NO <sub>3</sub> (mg/l)	F (mg/l)	Br (mg/l)	B (mg/l)	Sr (mg/l)	♦SiO <sub>2</sub> (aq) (mg/l)	NO <sub>2</sub> (mg/l)	IBE (%)	Calcite (SI)	Dolomite (SI)	Anhydrite (SI)	Halite (SI)
	<i>Petit Bara Zone</i>																					
1	PK 44	28.8	22.8	165.3	4.9	≤ DL	≤ DL	267.2	131.9	84.3	29.6	0.6	0.5	–	0.69	39.8	0.02	1	0.12	0.6	-2.28	-6.29
2	PK 48	26.8	22.7	151.6	4.1	0.02	0.01	347.8	100.1	65.7	12.4	0.83	0.41	–	0.54	40.1	0.01	-1	0.33	1.04	-2.42	-6.44
3	PK 50	24.9	22.2	187.7	4.4	0.04	≤ DL	352.9	147	77.9	17.7	0.71	0.72	0.06	0.57	41	≤ DL	-2	0.52	1.47	-2.4	-6.19
4	PK 51	23.1	21.1	222.9	4.3	0.01	≤ DL	307.9	189.1	88.9	14.8	0.78	0.85	0.15	0.53	40.6	0.03	1	0.23	1.18	-2.37	-6.01
5	Didjan Der	88	63.2	384	18.5	0.9	0.13	237.1	609.5	198	32.5	0.01	2.88	–	–	–	0.03	2	0.66	1.55	-1.7	-5.28
6	Omar Jakka	89.9	68.9	435.7	8.5	0.02	≤ DL	219	697.8	234	54.3	0.43	2.69	0.36	2.24	34.3	0.02	0	0.57	1.49	-1.61	-5.2
7	Naheel	215	231	852.5	11	0.2	0.43	95	1874	608	119	0.9	7.74	0.89	6.5	46.0	0.15	-1	-0.13	0.24	-1.1	-4.54
8	Kourtimalay ♠	368	165	1080	32	–	–	61	2410	572	122	–	–	–	–	–	–	-2	0.45	1.01	-0.92	-4.33
	<i>Grand Bara Zone</i>																					
9	Iskoutir	145.7	65.2	440.1	8.2	0.02	≤ DL	100.9	818.5	319	50.8	0.28	3.5	1.95	3.80	–	0.09	0	0.37	0.8	-1.33	-5.11
10	Dagahdamere	199.3	123	653	15.3	≤ DL	0.8	95	1272	487	62	0.09	5.29	0.22	6.71	37.4	0.21	0	0.39	1.04	-1.11	-4.8
11	Gor Galo	88.5	57.4	453	8.3	0.04	≤ DL	213.3	572.1	334	142	0.89	2.5	1.18	2.58	38.2	≤ DL	0	0.6	1.49	-1.38	-5.28
12	Goroja ♥	180	132	1180	24	–	–	122	1990	553	130	–	–	–	–	–	–	0	0.53	1.39	-1.1	-4.4
13	Hambocto F2	115	66.8	169.5	5	0.01	≤ DL	209.7	269.2	131	255.7	0.60	2.35	0.07	2.37	43.7	≤ DL	2	0.14	0.49	-1.68	-6
14	Hambocto F1	58.4	38.7	186	4.1	0.01	≤ DL	239.7	249.8	113	49.8	1.01	1.27	4.31	2.6	38.6	≤ DL	0	0.05	0.39	-1.94	-5.98
15	Doudoub Bololeh F2	71	61.9	327.5	7.7	0.02	≤ DL	166.1	517.2	240	55.1	0.68	2.09	0.34	2.38	32.8	≤ DL	0	0.19	0.76	-1.64	-5.43
16	Doudoub Bololeh F1	165	134.8	507.5	13.1	0.03	≤ DL	107.7	1120	419	59.1	0.61	4.13	0.37	4.27	29.4	≤ DL	-2	0.19	0.77	-1.17	-4.96
17	Awraoussa 1	116.9	65.9	321.2	11.7	0.04	≤ DL	140.8	519	295	71	0.64	2.59	0.48	2.46	35.7	≤ DL	2	0.49	1.18	-1.37	-5.45
18	Gabla-galan	36.2	50.3	382.5	7.9	0.04	0.13	235.1	356	193	247.5	0.39	2.26	0.11	0.94	28.7	≤ DL	2	0.56	1.82	-2.1	-5.41
	<i>Mouloud Zone</i>																					
19	Mindil	55.1	92.6	361.2	9.8	0.02	≤ DL	283.7	481.4	301	47.3	0.52	2.51	1.91	2.82	32.8	≤ DL	2	0.13	0.92	-1.72	-5.43
20	Mouloud 3	82.2	79.8	442.9	10.9	0.02	≤ DL	188.9	686.9	337	79.6	0.43	2.73	0.31	2.39	32.6	≤ DL	-1	0.17	0.78	-1.51	-5.2
21	Mouloud 4	96.5	92.2	377.3	8.3	0.03	≤ DL	177.5	625.5	329	75.8	0.62	2.56	–	–	32.4	≤ DL	1	0.06	0.56	-1.45	-5.31
22	Dadin 3	46.9	26.4	217.5	5.1	0.03	≤ DL	443.6	157.3	103	33.8	1.51	0.4	≤ DL	1.02	30.9	0.07	-1	0.29	0.77	-2.08	-6.1

♠ Closed after drilling in 1985 because of the high salinity of its groundwater; ♥ Closed after drilling in 1978 because of the high salinity of its groundwater;

DL: Detection Limit (DL(B) = 0.3 µg/l; DL(NO<sub>2</sub>) = 1.6 µg/l); IBE: Ionic Balance Error.; ♦ SiO<sub>2</sub>(aq) ≡ H<sub>4</sub>SiO<sub>4</sub><sup>0</sup>

**Table 3.** Isotope data of Bara groundwaters from this study and literature.

Table 3: Isotope data of Bara groundwater from this study and literature.												
Samples		$\delta^{18}\text{O}(\text{H}_2\text{O})$ (‰ vs V-SMOW)	$\delta^2\text{H}(\text{H}_2\text{O})$ (‰ vs V-SMOW)	Tritium (TU)	$\delta^{13}\text{C}(\text{DIC})$ (‰ vs PDB)	$\delta^{13}\text{C}(\text{CO}_2)_{\text{aq}}^{\clubsuit}$ (‰ vs PDB)	$\text{A}^{14}\text{C}$ (pmC)	$^{14}\text{C}$ age <sup>♣</sup> (years BP)	$^{87}\text{Sr}/^{86}\text{Sr}$	$\delta^{15}\text{N}(\text{NO}_3)$ (‰ vs AIR)	$\delta^{18}\text{O}(\text{NO}_3)$ (‰ vs V-SMOW)	reference
Petit Bara Zone												
2	PK 48	-1.10	-4.8	$0.7 \pm 0.5$	-	-	-	-	-	-	-	This study
3	PK 50	-1.10	-4.8	$< 0.6$	-11.81	-18.2	-	-	0.70600	8.45	8.46	This study
3	PK 50	-	-	$< 2.2$	-11.60*	-17.9	$69.9\pm0.9$	$1061\pm107$	-	-	-	BGR (1982)
3	PK 50	-1.29	-	$7 \pm 1$	-11.60	-17.8	$\leq 60$	-	-	-	-	Fontes et al. (1980)
3	PK 50	-1.57	-	-	-11.69	-17.9	$80.4\pm1.0$	$23\pm104$	-	-	-	Fontes et al. (1980)
4	PK 51	-1.10	-5.0	$< 0.6$	-11.41	-17.7	-	-	0.70606	8.84	7.25	This study
6	Omar Jakka	-1.70	-11.0	$1.2 \pm 0.6$	-10.78	-17.1	-	-	0.70610	8.15	11.13	This study
7	Naheel	-3.24	-24.5	-	-8.71	-14.4	-	-	0.70596	7.57	16.79	This study
Grand Bara Zone												
9	Iskoutir	-	-	-	-	-	-	-	-	6.70	7.04	This study
10	Dagah Damer	-3.65	-24.8	-	-	-	-	-	0.70594	-	-	This study
11	Gor Galo	-2.82	-17.1	-	-9.67	-15.6	-	-	0.70662	6.53	9.43	This study
12	Goroja	-3.20	-19.6	$< 2.2$	-8.40	-14.4	$17.2\pm1.1$	$8943\pm546$	-	-	-	BGR (1982)
13	Hambocto F2	-1.80	-8.9	$0.8 \pm 0.7$	-9.83	-15.9	-	-	0.70668	8.58	11.53	This study
14	Hambocto F1	-2.94	-16.8	-	-7.99	-13.9	-	-	0.70657	6.78	7.37	This study
15	Doudoubolole F2	-3.30	-21.4	$< 0.6$	-6.89	-12.7	-	-	0.70648	8.18	10.66	This study
16	Doudoubolole F1	-3.70	-24.8	$< 0.6$	-7.09	-13.5	-	-	0.70637	9.31	12.74	This study
16	Doudoubolole	-4.06	-32.0	$< 4$	-8.40	-14.8	$27.6\pm2.7$	$6266\pm850$	-	-	-	Fontes et al. (1980)
17	Awraoussa 1	-3.10	-18.8	$< 0.6$	-8.74	-15.3	-	-	-	7.70	12.03	This study
18	Gabla Galan	-1.90	-9.4	$< 0.6$	-6.85	-14.5	-	-	0.70556	11.35	10.59	This study
Mouloud-Dadine Zone												
19	Mindil	-3.90	-25.5	$< 0.6$	-10.75	-17.0	-	-	0.70647	9.24	10.09	This study
20	Mouloud 3	-3.40	-21.5	$< 0.6$	-10.31	-16.5	-	-	0.70613	8.47	11.16	This study
	Mouloud 5	-3.60	-21.3	-	-	-	-	-	-	-	-	Aboubaker et al. (2013)
20	Mouloud 3	-3.05	-20.1	$< 2.1$	-9.50	-15.9	$49.6\pm0.8$	$2246\pm134$	-	-	-	BGR (1982)
21	Mouloud	-3.93	-32.0	$< 4$	-10.79	-17.0	$52.2\pm2.5$	$3085\pm406$	-	-	-	Fontes et al. (1980)
22	Dadin 3	-2.40	-1.3	$< 0.6$	-12.21	-18.5	-	-	0.70694	6.47	7.15	This study
	Dadin 6	-2.50	-1.2	-	-	-	-	-	-	-	-	Aboubaker et al. (2013)

- not measured or not calculated
- ♠  $\delta^{13}\text{C}$  of aqueous  $\text{CO}_2$  was calculated by PHREEQCI software and *iso.dat* database (Parkhurst and Appelo 2015).
- ♥ Corrected with the IAEA method
- ♣ Based on data from Fontes et al. (1980)

Figure 1

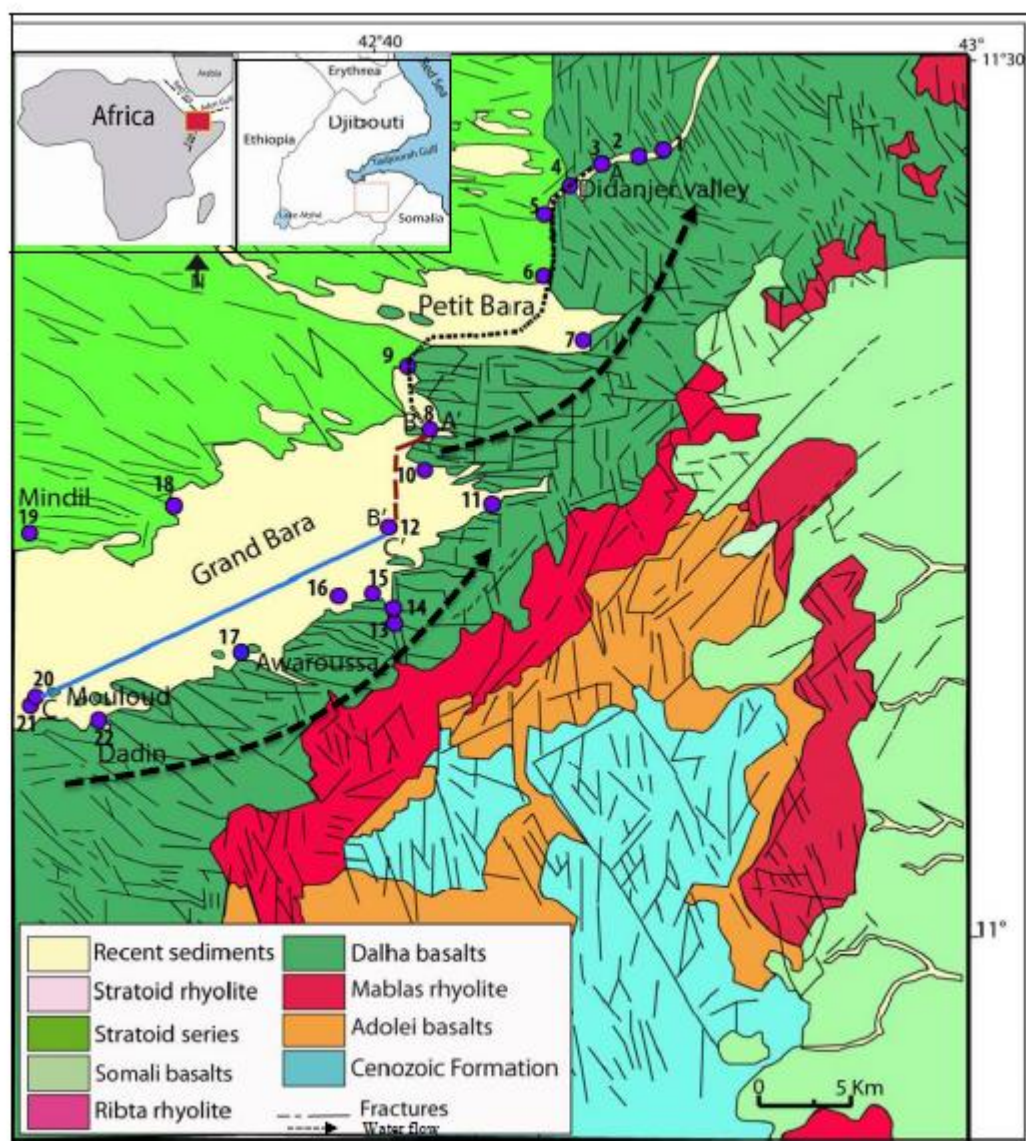


Figure 2

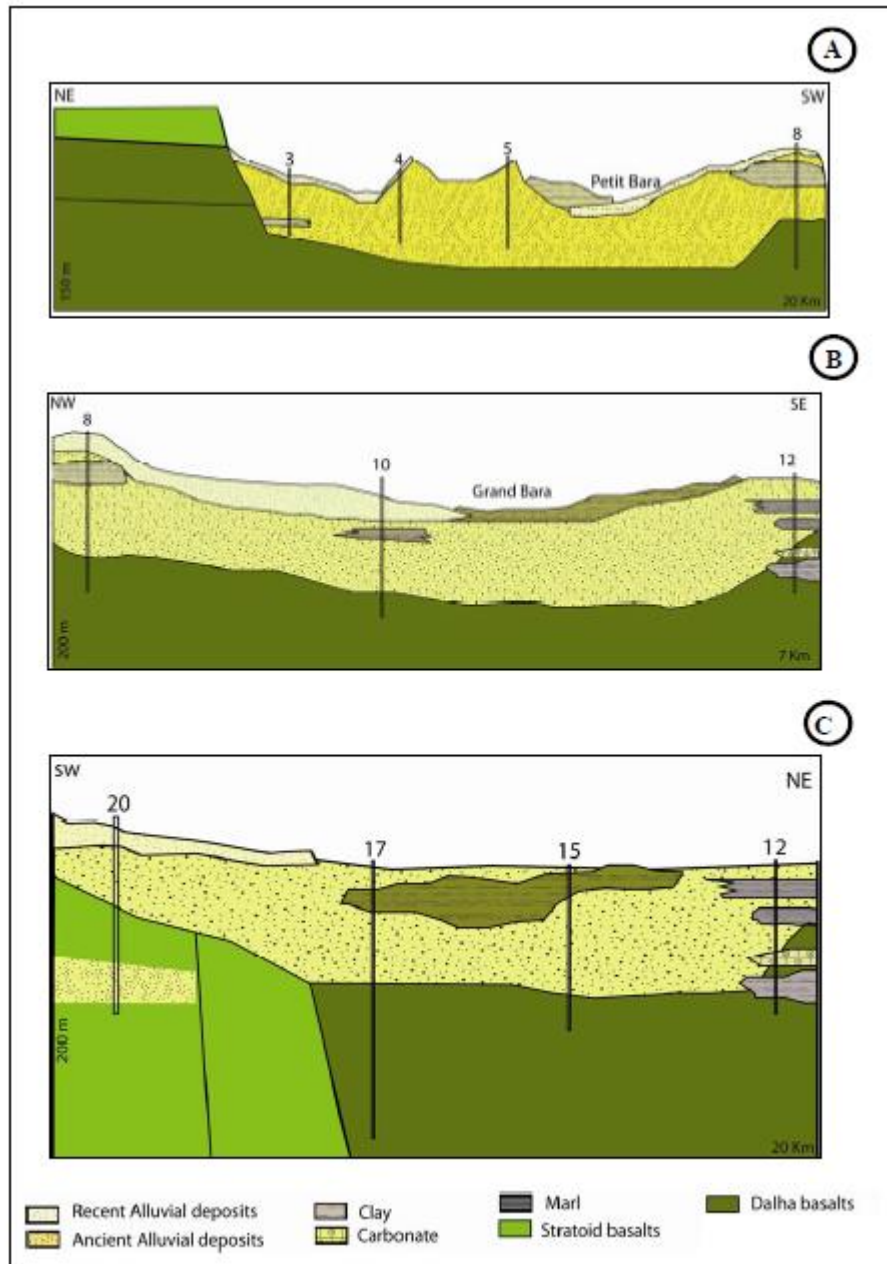


Figure 3

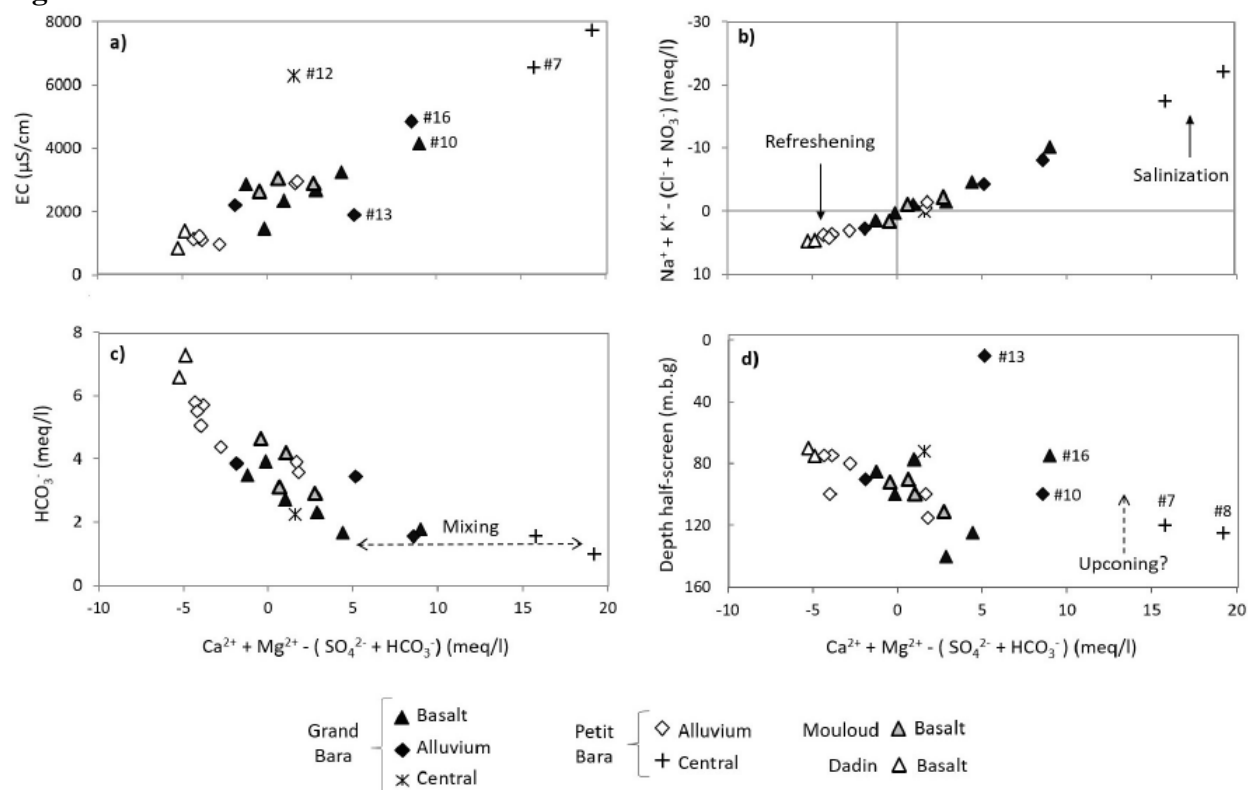




Figure 4

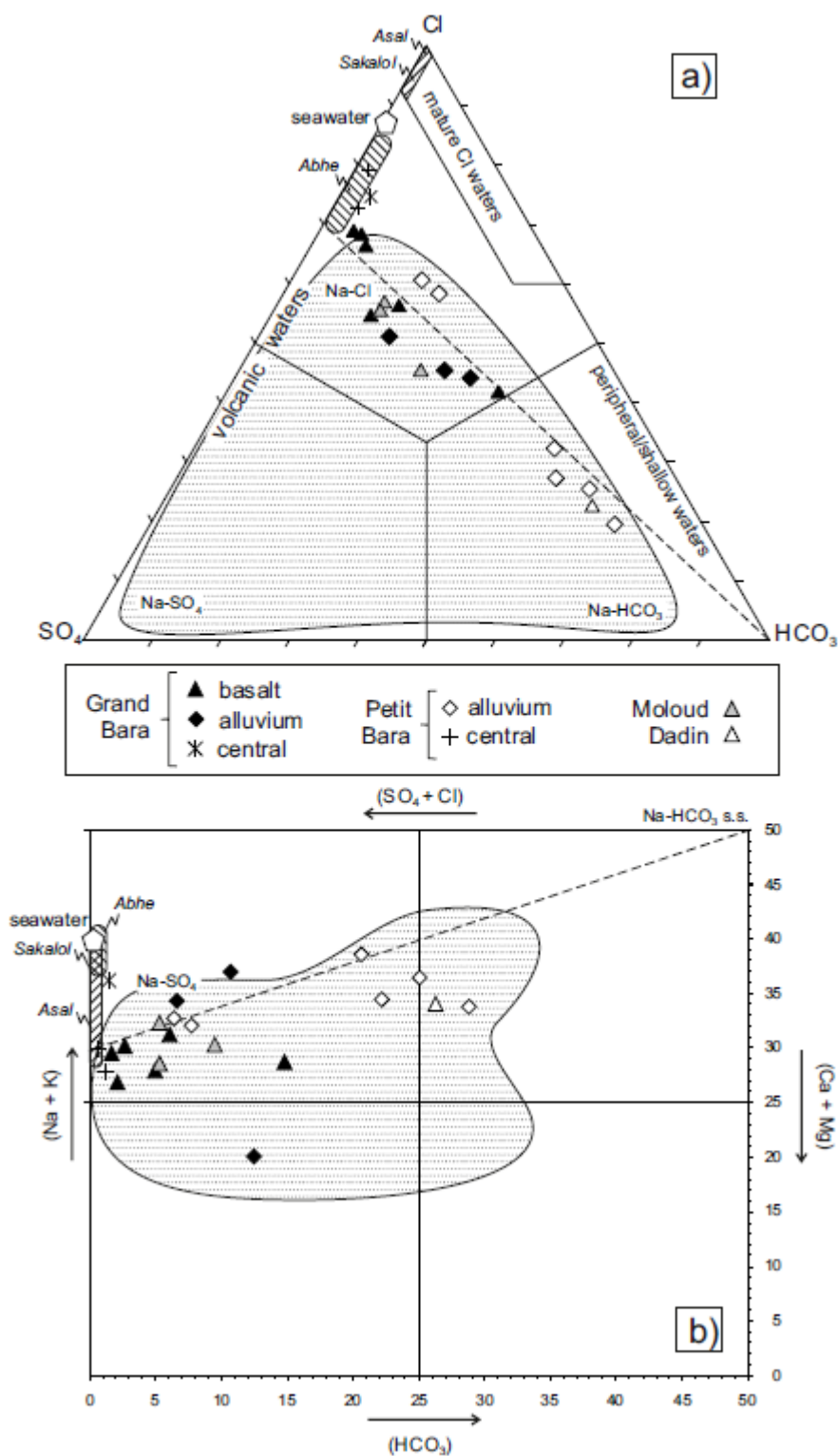


Figure 5

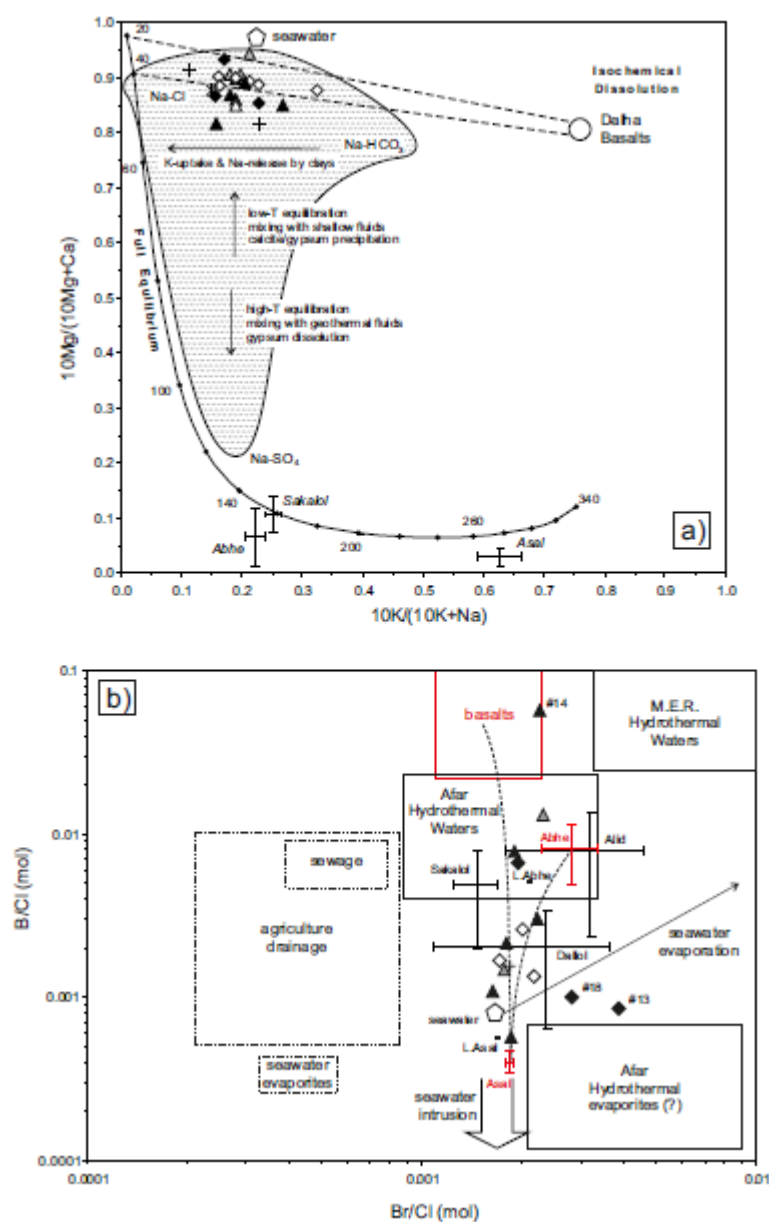


Figure 6

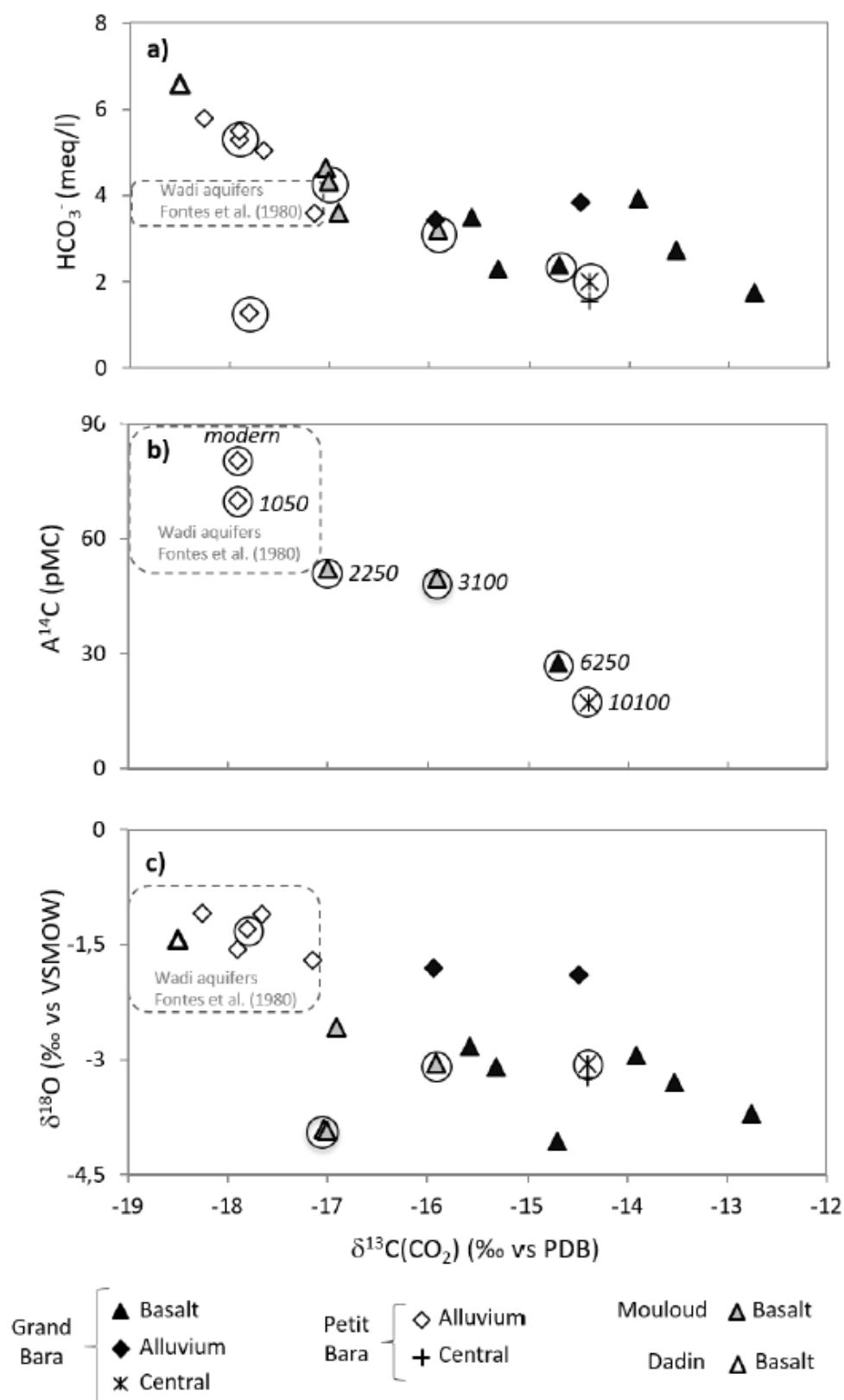


Figure 7

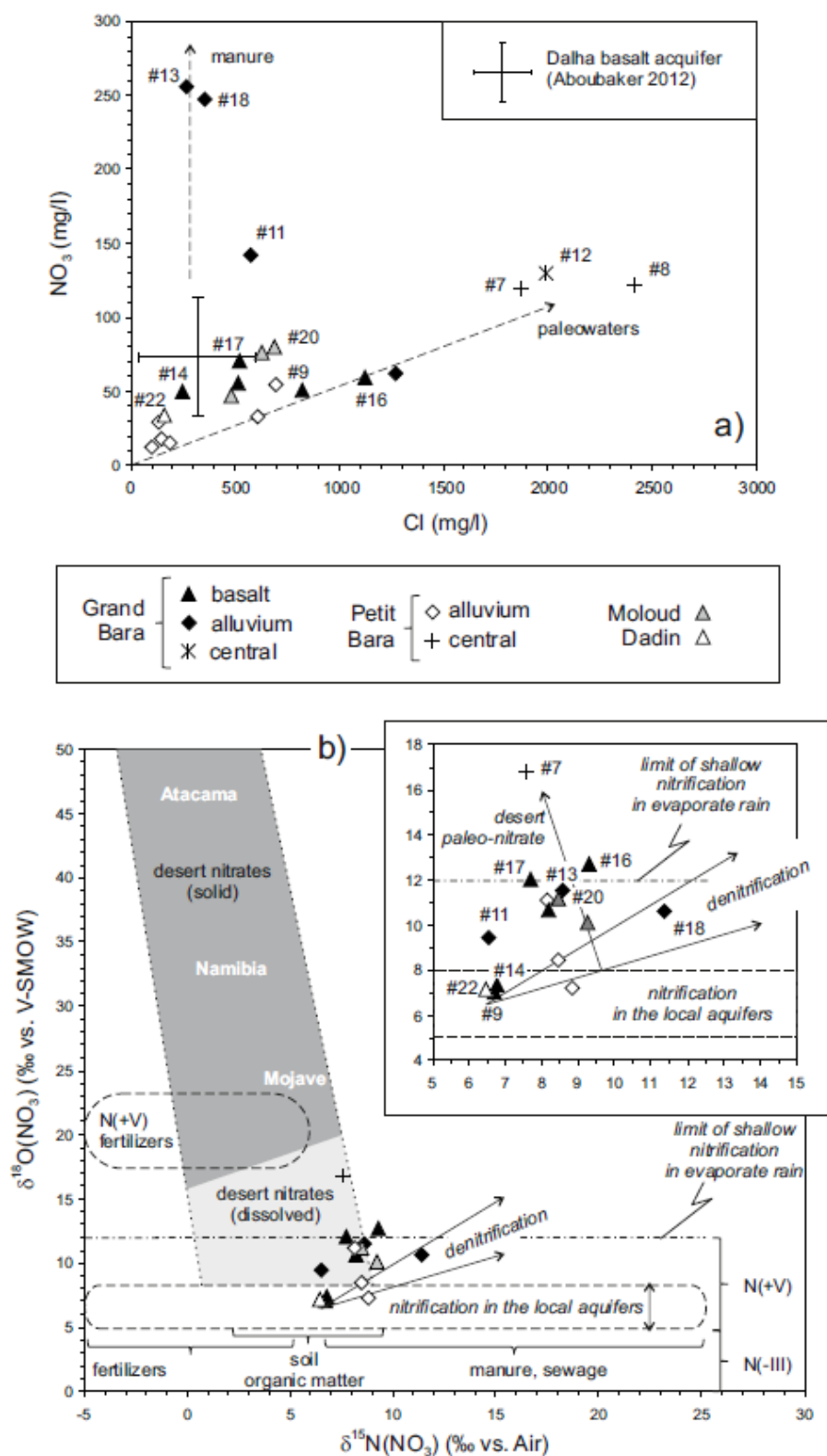


Figure 8

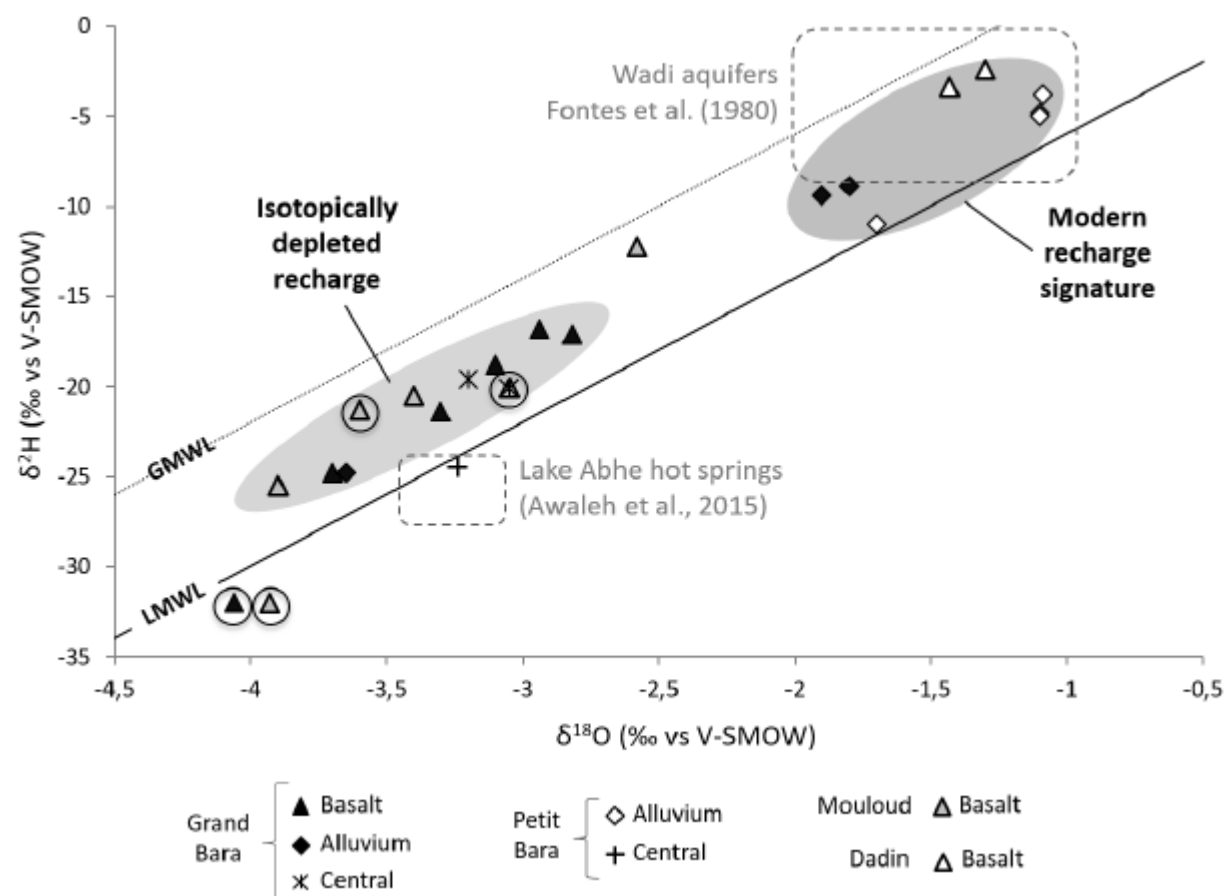


Figure 9

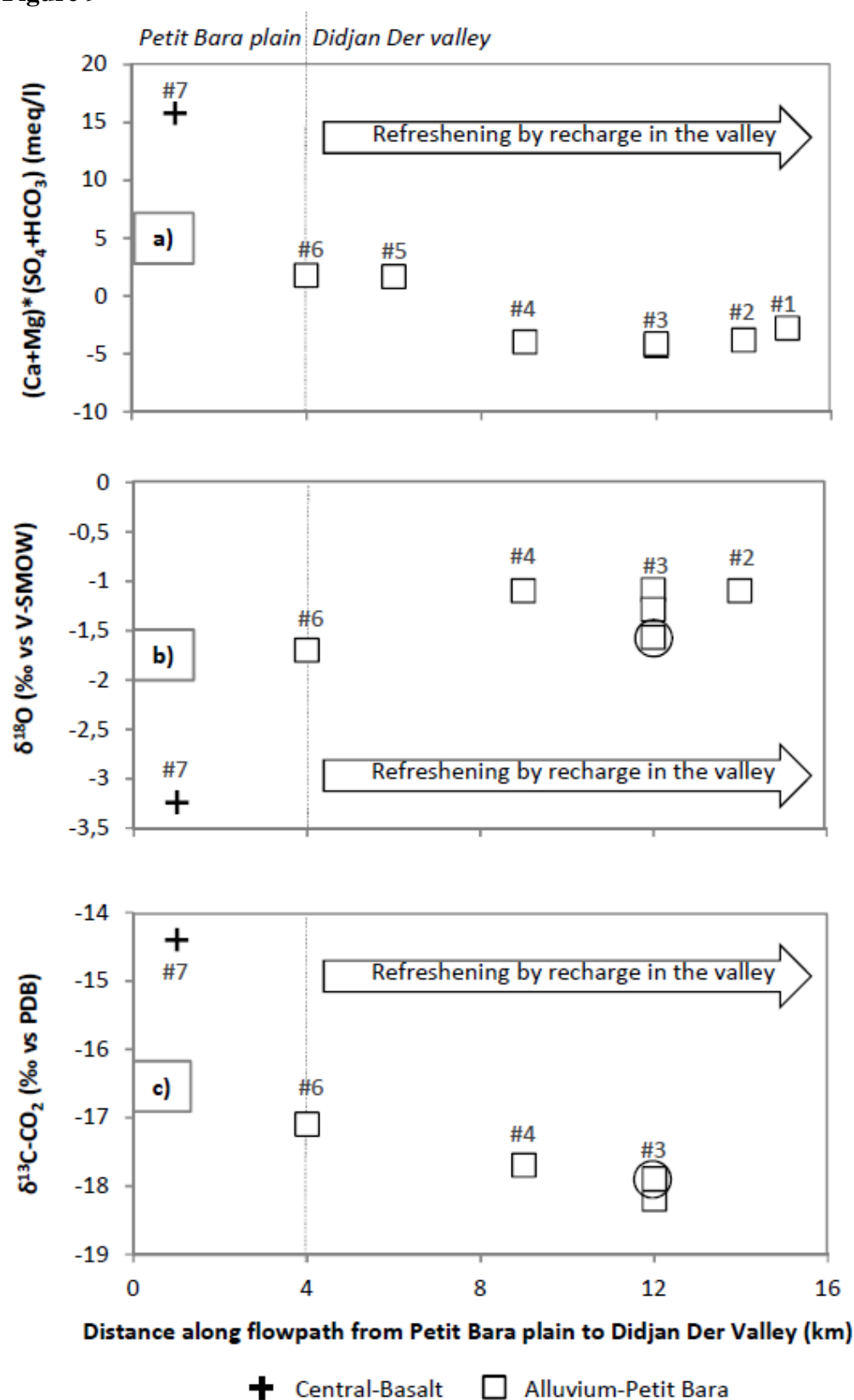
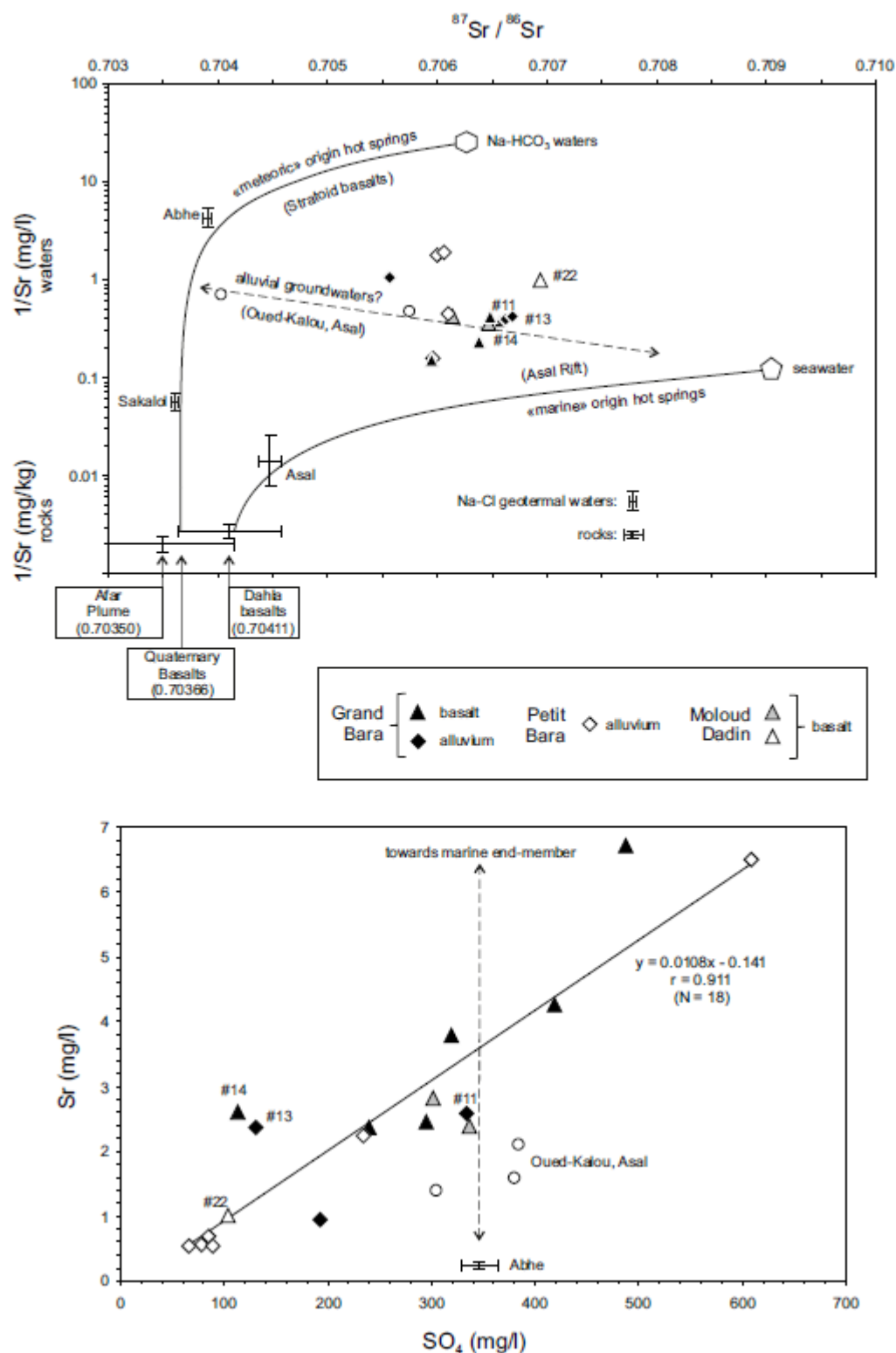


Figure 10



## Highlights

- Investigating an alluvium-basaltic aquifer system in arid eastern Africa
- Groundwater flow pattern and mixing revealed by geochemistry and isotopes
- Consistent recharge from wadis and downward circulation to the basalts
- Absence of a deep volcanic CO<sub>2</sub> contribution to basalt groundwater
- Soil, manure and caliche nitrate sources evidenced by isotopic nitrate