SEQUENCE CRYSTALLIZATION DURING ISOTHERM EVAPORATION
OF SOUTHERN ALGERIA CHOTT BAGHDAD NATURAL BRINE

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ABSTRACT
Southern Algerian's natural brine sampled from chott Baghdad may be a source of mineral
salts with a high economic value. These salts are recoverable by simple solar evaporation.
Indeed, during isothermal solar evaporation, it is possible to recover mineral salts and to
determine the precipitation sequences of different salts as a function of the chemical
composition and the density of the brine. In this study, the variation of ionic composition of
concentrated brine during isothermal evaporation was measured; then the experimental
pathway of the point representing its composition on the oceanic fivefold diagram Na⁺, K⁺,
Mg²⁺, Cl⁻, SO₄²⁻/H₂O was plotted. In order to follow the precipitation sequences of mineral
salts, during solar evaporation at 35°C, X-Ray diffraction was performed on the precipitated
and removed salts from the brine during evaporation.

Keywords: seawater quinary diagram, Jänecke projection, isothermal evaporation, mineral
salts, chott Baghdad brine.

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1. INTRODUCTION
The Sebkhs of south Algeria contain a high volume of natural brine, which represents enormous resources for extraction of dissolved salts [1]. Sodium chloride has already recovered from Melghir and Merouane chotts. Potassium and magnesium salts may be recovered. Indeed, during isothermal solar evaporation it is possible to recover mineral salts and to determine the precipitation sequences of the different salts as a function of the composition and the density of the brine. These observations lead us to examine the case of chott Baghdad natural brine. With regard of this aim, the variations of the ionic compositions of chott Baghdad brine during isothermal evaporation at 35°C will be studied. XRD was performed to determine the precipitated salts during evaporation. The precipitation sequences of mineral salts, during isothermal evaporation at 35°C, are established and the experimental pathway of this brine during its evolution was plotted on the oceanic fivefold seawater diagram (Na\(^+\), K\(^+\), Mg\(^{2+}\), Cl\(^-\), SO\(_4^{2-}\)//H\(_2\)O).

2. MATERIALS AND METHODS
2.1. Experimental
The ionic composition of the chott Baghdad natural brine studied is given in Table 1.

| Density | Na\(^+\) | Mg\(^{2+}\) | K\(^+\) | Ca\(^{2+}\) | Cl\(^-\) | SO\(_4^{2-}\) |
|---------|---------|------------|--------|-----------|--------|-----------|
| 1,223   | 89,00   | 8,14       | 9,60   | 0,40      | 155,43 | 15,00     |

Four liters of the above brine were evaporated in a glass vessel and placed in a thermostatic bath, which was maintained at 35°C. The crystallized salts were separated by filtration, as well as they were formed. Then the solid and the liquid phases were characterized (solutions were designated Si, i=1 à 6 and solids were designated Salt-i). X Rays powder diffraction was used to characterize the solid phase, and chemical analysis and density measures were used to characterize the liquid.

The experimental pathway representing the evolution of the evaporated brine’s composition was then plotted on the Jänecke projection at 35°C of the quinary diagram Na\(^+\), K\(^+\), Mg\(^{2+}\), Cl\(^-\), SO\(_4^{2-}\)//H\(_2\)O.
2.2. Location of the chott Baghdad brine on the quinary diagram

The Jänecke coordinates of the point representing the chott Baghdad natural brine on the oceanic quinary diagram Na\(^+\), K\(^+\), Mg\(^{2+}\), Cl\(^-\), SO\(_4^{2-}\)/H\(_2\)O [2] were calculated using the following equations:

$$\% \text{Mg} = \frac{100 \times n_{Mg^{2+}}}{D}$$
$$\% \text{K}_2 = \frac{100 \times n_{K^{+}}}{D}$$
$$\% \text{SO}_4 = \frac{100 \times n_{SO_4^{2-}}}{D}$$

with, $D = n_{Mg^{2+}} + \frac{n_{K^{+}}}{2} + n_{SO_4^{2-}}$ and $n_i$ the mole number of the $i$ entity

Using the above mentioned results in Table 1 and these equations, the Jänecke coordinates are as follows:

$$\% \text{Mg} = 29.13$$
$$\% \text{K}_2 = 17.18$$
$$\% \text{SO}_4 = 53.69$$

**Fig.1.** indicates the location of the chott Baghdad natural brine on the oceanic quinary diagram Na\(^+\), K\(^+\), Mg\(^{2+}\), Cl\(^-\), SO\(_4^{2-}\)/H\(_2\)O. The chott Baghdad natural brine is located in the Glaserite crystallization field. This position helps us to predict its theoretical pathway.

![Chott Baghdad natural brine diagram](image-url)
2.3. Theoretical pathway

According to the composition and the geometrical position of the chott Baghdad natural brine’s, its theoretical pathway is represented on the oceanic fivefold diagram \( \text{Na}^+, \text{K}^+, \text{Mg}^{2+}, \text{Cl}, \text{SO}_4^{2-}/\text{H}_2\text{O} \) at 35°C, given in Fig.2. Our prediction is based on Gibbs law: \( v = c - \Phi \)

- \( v \): system variance
- \( \Phi \): phase number
- \( c \): independents constituents number

In the case of oceanic quinary system, all the solutions are permanently saturated with NaCl. So, when a representative point is situated on a given plan (crystallization field), that means \( v = 2 = 5 - \Phi \); thus, \( \Phi = 3 \), Two salts and one solution. Consequently, the evaporation of chott Baghdad natural brine will give the following theoretical sequences:

| Sequence number | Pathway     | Crystallized salts |
|-----------------|-------------|--------------------|
| 1               | \( \alpha \alpha' \) (\( v = 3 \)) | Halite (NaCl)      |
| 2               | \( \alpha' \beta \) (\( v = 2 \)) | Halite + Glasérite (\( \text{Na}_2\text{SO}_4 \cdot 3\text{K}_2\text{SO}_4 \)) |
| 3               | \( \beta \text{T} \) | \( \text{NaCl} + \text{Glaserite} + \text{Astrakanite} \) ((\( \text{Na}_2\text{SO}_4 \cdot \text{MgSO}_4 \cdot 4\text{H}_2\text{O} \))  |
| 4               | Point T     | Decomposition of Glaserite and apparition of Leonite \( (\text{K}_2\text{SO}_4 \cdot \text{MgSO}_4 \cdot 4\text{H}_2\text{O}) \) |
| 5               | TV          | Halite + Astrakanite + Leonite. |
| 6               | V Point     | Dissolution of Leonite and apparition of Kaïte \( (\text{KCl} \cdot \text{MgSO}_4 \cdot 3\text{H}_2\text{O}) \) |
| 7               | VW          | Halite + Astrakanite + Kaïte |
| 8               |             | This pathway continued till Z point if the amount of water is enough |

Table 2. Predicted salts sequences
Fig. 3. Theoretical pathway of the studied brine on the oceanic quinary diagram

\[ \text{Na}^+, \text{K}^+, \text{Mg}^{2+}, \text{Cl}^-, \text{SO}_4^{2-}/\text{H}_2\text{O} \text{ at } 35^\circ\text{C}. \]

At Z point (true invariant point), the solution does not move and will carry on the evaporation till drying. Fig. 2 represents the above pathway followed by the brine on the quinary diagram. It is important to mention that this theoretical crystallization sequence is not almost respected due to metastable phenomena and supersaturated solutions [3, 4, 5].

3. RESULTS AND DISCUSSION

3.1. Results

The ionic composition and densities of the brine during the evaporation process are presented on Table 3. The ionic composition of the precipitated salts is given on Table 4.

As made for natural brine coordinates calculations (§ 2.2), we have determined those of the different solutions in evolution.

The obtained points were projected in Fig. 3. This corresponds to the evaporation pathway followed by the brine.

X-Ray diffraction of recovered mineral salts during evaporation is presented by Fig. 4. to Fig. 7.
Table 3. Ionic compositions (g.l\(^{-1}\)) of the studied brine at 35°C

| Solution | Density | Ca\(^{2+}\) | Mg\(^{2+}\) | Na\(^+\) | K\(^+\) | Cl\(^-\) | SO\(_4^{2-}\) |
|----------|---------|-------------|-------------|----------|--------|--------|----------|
| NS       | 1.223   | 0.40        | 8.14        | 89.00    | 9.60   | 155.43 | 15.00    |
| S1       | 1.235   | 0.12        | 14.45       | 72.00    | 13.60  | 152.70 | 23.00    |
| S2       | 1.254   | 0.00        | 21.68       | 58.00    | 18.07  | 150.00 | 31.82    |
| S3       | 1.275   | 0.00        | 28.00       | 52.00    | 23.43  | 147.13 | 37.93    |
| S4       | 1.289   | 0.00        | 37.98       | 34.00    | 33.79  | 145.78 | 47.27    |
| S5       | 1.293   | 0.00        | 42.90       | 20.00    | 36.34  | 159.45 | 39.45    |
| S6       | 1.305   | 0.00        | 45.00       | 18.00    | 33.45  | 165.89 | 33.67    |

Table 4. Ionic composition (%) of the solid phase salts recovered during isothermal evaporation at 35°C.

| Salt   | Density | Ca\(^{2+}\) | Mg\(^{2+}\) | Na\(^+\) | K\(^+\) | Cl\(^-\) | SO\(_4^{2-}\) |
|--------|---------|-------------|-------------|----------|--------|--------|----------|
| Salt-1 | 1.235   | 0.42        | 0.15        | 37.56    | 0.10   | 60.76  | 1.41     |
| Salt-2 | 1.254   | 0.08        | 0.19        | 37.23    | 0.22   | 61.18  | 1.17     |
| Salt-3 | 1.275   | 0.00        | 0.12        | 35.25    | 0.24   | 53.58  | 3.99     |
| Salt-4 | 1.289   | 0.00        | 0.14        | 33.00    | 0.93   | 39.70  | 20.30    |
| Salt-5 | 1.293   | 0.00        | 0.60        | 22.86    | 18.10  | 20.77  | 43.57    |
| Salt-6 | 1.305   | 0.00        | 0.78        | 22.86    | 21.27  | 22.92  | 47.71    |
Fig. 3. Experimental pathway of chott Baghdad brine on the oceanic quinary diagram Na\(^+\), K\(^+\), Mg\(^{2+}\), Cl\(^-\), SO\(_4\)\(^{2-}\)/H\(_2\)O at 35°C.

This pathway is completely different from the theoretical one mentioned in Fig. 2.

Fig. 4. XR Diffraction of Salt-2: Halite NaCl and Gypsum
Fig. 5. XR Diffraction of Salt-4: Halite.

Fig. 6. XR Diffraction of Salt-5: Halite, Picromerite and Epsomite.
The above diffractograms allow us to establish the mineralogical composition of the recovered salts. The results are listed on Table 5. The obtained salts sequences are completely different from those predicated on Table 2.

**Table 5.** Mineralogical identification of recovered salts

| Salt   | Density | Solid phase salts (identified by XRD)                                                                 |
|--------|---------|-----------------------------------------------------------------------------------------------------|
| Salt-1 | 1.235   | Gypsum + Halite                                                                                     |
| Salt-2 | 1.254   | Gypsum + Halite                                                                                     |
| Salt-3 | 1.275   | Halite                                                                                              |
| Salt-4 | 1.289   | Halite                                                                                              |
| Salt-5 | 1.293   | Halite + Picromerite (MgSO₄K₂SO₄, 6H₂O) + Epsomite (MgSO₄.7H₂O)                                    |
| Salt-6 | 1.305   | Halite + Sylvite (KCl) + Epsomite                                                                    |

### 3. DISCUSSION

Table 5. shows the apparition of Sodium Chloride between densities from 1.235 to 1.289. In
the first stage, Halite was accompanied with Gypsum (d=1,235±1,254) and then became alone (d=1,254±1,289), the gypsum apparition is explained by the presence of minor amounts of Calcium in the initial solution as mentioned in Table 1. This solution is concentrated in Sulfate ion. Consequently and with regard to its very low solubility, Gypsum will precipitate. These Two sequences represent only one which agrees with the predicted one (Table 2. sequence number 1).

Salt-5, precipitated between d=1,289 and d=1,293 is a mixture of Sodium Chloride, Picromerite (MgSO$_4$ K$_2$SO$_4$. 6H$_2$O) and Epsomite (MgSO$_4$.7H$_2$O). This sequence is different from the predicted one. Also salt-6 which crystallized between d=1,293 and d=1,305 consists of Halite, Sylvite (KCl) and Epsomite, is different from the foreseeable last sequences as indicated in Table 2.

So, the plotted experimental crystallization pathway didn’t confirm the predicted one. Effectively, the different points of chott Baghdad natural brine evaporated were inside the Glaserite (Na$_2$SO$_4$. 3K$_2$SO$_4$) crystallization field (Fig.3.), but experimentally this salt (Glaserite) was missed and replaced by Picromerite and Epsomite. Moreover, the last point, which represents the last solution S6, still in the Glaserite field and the final salt is formed by Halite, Sylvite and Epsomite, instead of Halite, Astrakanite (Na$_2$SO$_4$. MgSO$_4$. 4H$_2$O) and Glaserite (Na$_2$SO$_4$. 3K$_2$SO$_4$).

The exam of recovered salts shows that only the first sequence agrees with the predicted one. Glaserite and Astrakanite are missed. Similar observations were made by F. Khalissa and all [3] according to their work on the Zarzis Sabkhat El Mellah south Tunisia brine, where Epsomite takes place of Astrakanite and Glaserite. Moreover, the same phenomenon was observed by A. M’nif and R. Rokbani [4] when they have studied the Sabkhat El Adhibate and Sabkhat El Mellah natural brines, where Epsomite takes place of Astrakanite.

The above established differences between foreseeable and experimental crystallization sequences corroborate perfectly with the result reached by Kurnakow and Nikolaew when they have studied the solar diagram as it is reported by Balarew [5]. Indeed, on the solar diagram, the solubility fields of astakranite, kainite and kieserite are missed, that’s what makes the Epsomite replace Astrakanite. The same observation was made by Neitzel as it is
reported by A. M’nif [4]. Indeed, according to his work on the Great Salt Lake brine, he mentioned that Astrakanite and some others double salts do crystallize only when respective nuclei are present in the system. Furthermore, M’nif and Rokbani [4] observe that the recovered mixture of halite and Epsomite convert completely into Astrakanite after a long period. Thus, astrakanite is a secondary mineral.

Glaserite didn’t crystallize because of its slower kinetic comparing with Epsomite or Sylvite [4]. It is established that when seconds are needed for KCl and NaCl crystallization, minutes are needed for Picromerite (MgSO₄ K₂SO₄. 6H₂O), and hours are needed for Leonite (MgSO₄ K₂SO₄. 4H₂O) and Glaserite (Na₂SO₄. 3K₂SO₄) crystallization [6].

4. CONCLUSION

The obtained results confirm that Jaénecke projection is indicated to study the high concentrated brine because of its pertinence and simplicity. The crystallization sequences determined to chott Baghdad natural brine are: Gypsum, Halite, Picromerite, Epsomite and Sylvite. Only for the first sequence, a good correlation is observed between predictable evolution and experimental one. After that, a clear deviation is reported. It is mainly caused by kinetic considerations as explained before. It is well known that high concentrated Magnesium solutions form a [Mg(H₂O)₆]²⁺ complex [7] which steers the future pathway salts crystallization. In this case, some crystallization surfaces, which didn’t have the octahedral crystalline form like the complex already mentioned become so flexible and penetrate the juxtaposed salt field’s to precipitate it, before crystallizing its own salt. This is the case of Glaserite. Astrakanite and other double salts do crystallize only when respective nuclei are present in the system, they are secondary minerals [4].

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