

# Facies and geochemistry of non-marine gypsum, EMISAL, Egypt

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## ABSTRACT

Diverse gypsum facies in terms of crystal size and morphology are deposited in a shallow, non-marine, sulfate-enriched, semi-closed, perennial evaporite basin. These gypsum facies were deposited in intimate association with cyanobacterial mats. Facies analyses and geochemistry revealed two distinct primary facies, based on gypsum fabric; massive selenite and laminated gypsum varieties. The massive selenite facies is composed of clusters of single and twinned crystals along (100), with upward pointing re-entrant angle. The laminated gypsum varies from rhythmically laminated grass-like selenite to gypsum microbialite (stromatolite). The change from massive selenite to laminated gypsum facies implies fluctuation in lateral and vertical salinity profiles, seasonal brine concentrations and wind action. Post-depositional changes in the form of dissolution, slight to moderate crystal deformation and recrystallization to anhydrite are evaluated. We develop a qualitative model for the depositional basin and facies distribution in order to simulate ancient analogues.

**KEYWORDS** | Gypsum. Lithofacies. Non-marine. Geochemistry.

## INTRODUCTION

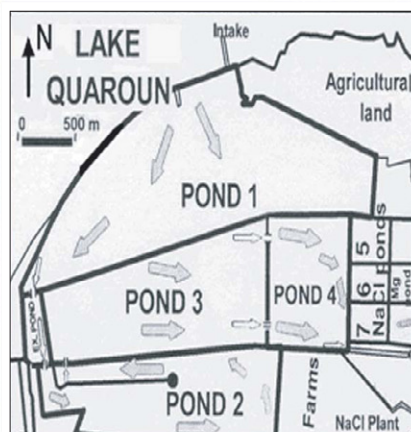
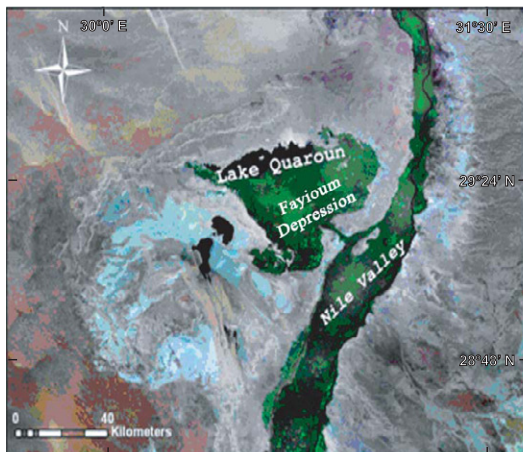
Marine and non-marine evaporites have formed in various modern and ancient depositional settings including marginal (mixed shallow-subaqueous and subaerial), shallow- and deep-subaqueous environments (*e.g.* Schreiber and El Tabakh, 2000). Evaporite basins vary from small lake/lagoon to large intracratonic/rift scale (Tucker, 2001). Data of evaporite deposits from artificial salinas is crucial in understanding shallow-water evaporite depositional environments. The current interpretations of ancient sulfate lithofacies are mainly based on data from modern analogues, where original sedimentary and early diagenetic features are

traceable (*e.g.* Warren and Kendall, 1985; Cody and Cody, 1988; Cornée *et al.*, 1992; Schreiber and El Tabakh, 2000; Kasprzyk, 2003) and on experimental data from chemically precipitated phases (Cody and Cody, 1989). In general, the observed gypsum facies in recent shallow subaqueous environments represent the most common rock assemblages of ancient deposits (*e.g.* Ortí Cabo *et al.*, 1984a; Schreiber and El Tabakh, 2000); hence, they can serve as reasonable representations of ancient analogues. Detailed studies of modern evaporites that evolved under known depositional and physico-chemical conditions are important for deciphering the origin of ancient analogues. A number of depositional models for selenite facies have been postulated

(*e.g.* Hardie and Eugster, 1971; Warren, 1982; Ortí Cabo *et al.*, 1984a; 1984b; Babel, 2004).

Facies analysis, in general, is a fundamental tool in inferring the paleo-topographic and climatic conditions that prevailed during the formation of ancient primary evaporites. Hence, detailed macroscopic and microscopic investigations can elucidate the depositional history and environmental characteristics of such deposits (*e.g.* Hardie and Eugster, 1971; Shearman and Ortí Cabo, 1976; Ortí Cabo and Shearman, 1977; Schreiber, 1978; Ortí Cabo *et al.*, 1984a; Ciarapica *et al.*, 1985; Kasprzyk, 1999 in Kasprzyk, 2003; Peryt, 2001; Babel, 2005; 2007; Becker, 2005; Babel *et al.*, 2011).

The concentration pond 3 at EMISAL Co. (longitude 30°14'E, latitude 29°26'N) in the continental Lake Quaroun (Fig. 1) provides a good example of different primary gypsum facies deposited under controlled physico-chemical conditions. These gypsum facies were accumulated over the past few decades by the evaporation of non-marine, sulfate-enriched brines, in variable environments. Hence,



**FIGURE 1** | Satellite image of Fayoum depression in Lake Quaroun (Mosaic\_Landsat/N-36/N-36-20.ETM-EarthSat-MrSID) and a sketch of the concentration ponds of EMISAL.

the detailed study of these deposits gives a good opportunity to evaluate the formation conditions of ancient analogues.

In the present work, facies analysis, post-depositional changes (diagenesis) and geochemistry of the different gypsum facies are given. In addition, a qualitative model of the basin and distribution of the gypsum facies is postulated.

## SETTING

The rocky, low altitude platform of the Egyptian Western Desert is characterized by conspicuous natural depressions of various sizes (*e.g.* Said, 1962; Embabi, 2004). The latter author distinguished nine mega depressions (>1000km<sup>2</sup>), among them the Fayoum depression (~2580km<sup>2</sup>) which reflects both desert and Nile Valley signatures is outstanding (Fig. 1). In spite of the debate about the origin of these depressions, the role of erosion and wind action in their morphology is widely accepted (*e.g.* Said, 1960; Embabi, 2004). Owing to differential erosion of interbedded resistant and soft rocks, the borders of these depressions are mainly outlined by mesas and cuestas.

According to Beadnell (1905), the lithostratigraphic section of the Fayoum depression covers a time-span from middle Eocene to Pleistocene. It comprises marine, fluvio-marine and lacustrine deposits. Generally, the depression is characterized by the presence of recycled ancient lake-beaches and lacustrine sediments, implying sporadic connections with the Nile since middle Pleistocene (Embabi, 2004 and references therein).

The Lake Quaroun is the lowest topographic part of the Fayoum depression (~45m below sea level). It represents the remains of the pre-historic Lake Moeris (Said, 1962), which covered a large part of the Fayoum depression. The Lake Quaroun collects the excess of irrigation and the agricultural waste-water from the surrounding cultivated lands (Fig. 1) and receives about 60.000 tons of dissolved salts per year from leached surface crusts and fertilizers (Embabi, 2004). Na<sup>+</sup>, Ca<sup>2+</sup>, Mg<sup>2+</sup>, Cl<sup>-</sup> and SO<sub>4</sub><sup>2-</sup> are the major cations and anions in the lake water. Local climatic conditions of the area have been discussed in detail by many authors (*e.g.* Samy, 2000; Al-Kordy, 2003; Ali and El Sheikh, 2004). Generally, the region is characterized by hot daytime temperatures (35°C–45°C during summer months) and dry conditions (relative humidity ranges from 64 to 33%) with scarce rainfall (≤10mm, annually). The velocity of the dominant wind in the area (from north-northwest to south-southeast), ranges from 2.74 to 5.16m/s. Owing to the arid and dry climate of the area, considerable amounts of water are lost annually by evaporation. Thus, the salinity of the lake has been increasing dramatically, reaching about

35g/l at present. In comparison with seawater, Lake Quaroun water is richer in  $\text{SO}_4^{2-}$  and  $\text{HCO}_3^-$  by factors of 3.56 and 2, respectively, and depleted in  $\text{K}^+$  and  $\text{Cl}^-$  by factors of 0.57 and 0.67, respectively (Ali and El Sheikh, 2004).

Based on environmental and economic reasons (White Young and Partners, 1976; DSS-Engineers, 1979), the EMISAL project (Fig. 1) was developed in the Lake Quaroun to extract  $\text{Na}_2\text{SO}_4$ , edible  $\text{NaCl}$  and  $\text{Mg}$ -salts. The salt-extraction process is based on brine-concentration technology via controlled evaporation-precipitation cycles using the solar energy, throughout four successive ponds of about  $5.3\text{km}^2$ . Based on local meteorological information, four successive concentration ponds were designed with adjusted depths of 1.7m, 1.80m, 2.3m and 2.8m deep, to allow natural flow of the brine from one pond to another, via controlled gates (Fig. 1).

Concentration ponds 1, 2 and 3 cover areas of about  $4.82\text{km}^2$ , representing successive evaporation basins with different degrees of salinity. Clays and other suspended materials, as well as part of the carbonate ions, are eliminated in pond 1, by settling and crystallization, respectively. The complete precipitation of carbonates (mainly in the form of aragonite) is achieved in pond 2, while, the precipitation of calcium sulfate in the form of primary gypsum is confined to pond 3. The latter ( $\sim 1.403\text{km}^2$ ) represents a perennial, shallow, sub-aqueous, semi-closed basin, wherein the seasonal water level is governed mainly by the evaporation rate. Its extent, depth and configuration, enhances brine flow from west to east (Fig. 1), generating a horizontal salinity gradient. The brine column is about 2.30m deep with an average salinity of about 231.1g/l (Metwally, 2002); the brine entering from pond 2 has a salinity approaching the saturation of primary gypsum ( $130\pm 20\text{g/l}$ ), and salinity increases gradually eastward to about 260g/l at the eastern end of the pond (close to pond 4).

## MATERIALS AND METHODS

During the past few decades, a considerable amount of gypsum has accumulated along the banks (up to 3m wide x 1.5m thick) of pond 3 (Fig. 2). Sampling was carried out during the spring of 2006, above and below the temporarily brine on both northern and southern banks of pond 3 at regular intervals, from west to east, in order to register the direct effect of both salinity gradient throughout the pond and wind action on the growth of the gypsum crystals. Five brine samples representing the expected lateral salinity variation were collected. In addition, three bottom sediments covering the entire bottom surface of the pond and a sample of recently precipitated gypsum from the northwestern portion of the pond were collected. These samples represent the region of lowest salinity and minimal wind effect.



FIGURE 2 | Gypsum exposure at the southern bank of pond 3.

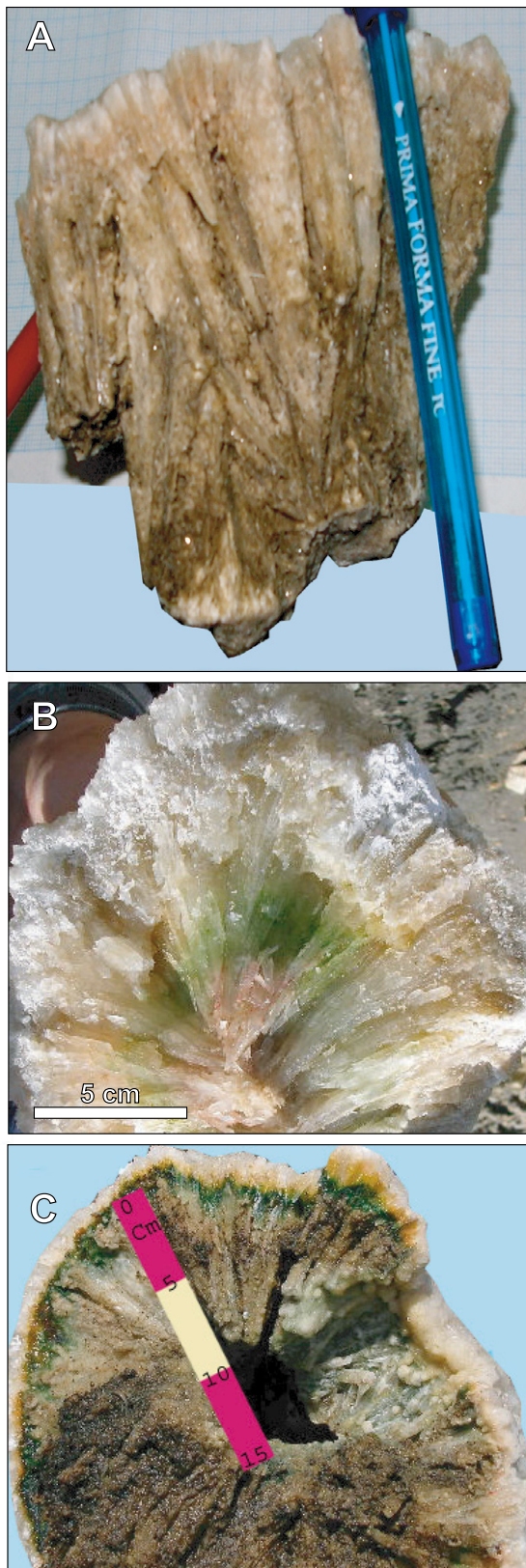
Gypsum samples were impregnated by epofix and left to dry and consolidate. Large (7.5x5cm) thin sections were prepared and studied with a polarizing microscope. Representative gypsum samples were then coated by gold and examined using a SEM (model Philips XL 30). Representative bulk samples were chemically investigated using X-Ray Fluorescence (XRF). The analyses were carried out by a wavelength-dispersive spectrometer (Axios PANalytical 2005), housed at National Research Centre, Dokki, Cairo. Gypsum standards (GYP-A and GYP-C) were used in calibration process.

## FACIES DESCRIPTION

In terms of fabric, texture and crystal morphology and size, the studied primary gypsum is classified into two main facies: i) massive selenite and ii) laminated gypsum. The latter is subdivided into two subfacies, namely, laminated grass-like selenite and microbialite (stromatolite).

### Massive selenite

This facies is mainly composed of clusters of upward-growing selenite crystals of various crystal sizes (2 – 12cm) and growth fabric patterns (Figs. 3A, B). Clusters are formed of single and twinned crystals along (100), with upward pointing re-entrant angle. Based on morphology and style arrangement of sub-parallel gypsum crystals, different fabrics may be distinguished (*e.g.* Hardie and Eugster, 1971; Schreiber, 1978). For example, the clustering of twinned crystals (Fig. 3A), characterized by an acute twin angle, exhibits in cross section (perpendicular to the twin plane) a radial pattern around a cavity (Fig. 3C). Such structure either formed around a stick or a bulge “bubble” in the mats (before



**FIGURE 3** | A) Single and twinned selenite crystals forming the massive selenite facies. B) A palmate pattern (fan-like) of upward-grown selenite crystals in the massive selenite facies. C) A cross section (perpendicular to the twin plane) exhibiting a radial pattern.

it was encrusted) that lithified due to gypsum growth (B.C. Schreiber, pers. commun., 2012). Commonly, multi-twin stacks may develop either simultaneously or subsequently, giving rise to a competent, displacement, growth nature.

Upward-growing crystals that exhibit a palmate pattern (fanning clusters) is yet another common growth pattern (*cf.* Fig. 3B). The (100) twin plane is commonly the nucleus of such a fabric, and the curved parts of each crystal are developed by regular zones of growing (120) prism faces (Schreiber, 1978). However, the palmate structure may develop by splitting individual members of the twin stacks of the crystals that have obtuse re-entrants (*e.g.* Shearman and Ortí Cabo, 1976; Ortí Cabo and Shearman, 1977; Schreiber, 1978). As these gypsum crystals grow outward and upward, successive split segments may develop quasi-triangular or polygonal cross-section outlines. The spaces created by splitting are commonly filled by either clastic substances or in situ skeletal, broken and lenticular gypsum. Upward curvature, sutured ends and thickening of the growing selenite crystals are conspicuous features of the selenite crystals. Hardie and Eugster (1971) attributed such upward arcing of selenite to the thickening of an individual crystal growing perpendicular to the bedding.

In general, the gypsum crystals exhibits several colors, varying from colorless with a pearly luster to green, black, purple, brown or even deep grey. Such coloration is attributed to the bacterial assemblages contained within the gypsum crystals (Schreiber and El Tabakh, 2000; Panieri *et al.*, 2008; 2010; Ali-Bik *et al.*, 2011). Based on FTIR data, Ali-Bik *et al.* (2011) inferred the relative abundance of intercalated cyanobacterial mats within the gypsum colored zones as: green > black > brown > white. They reflect the signature of the prevailed vertical redox conditions during the continuous growth of gypsum crystals and the seasonal fluctuation of brine salinity (Ali-Bik *et al.*, 2011).

For example, green and purple colors represent photosynthetic bacterial forms, filamentous and coccoidal, respectively (Schreiber and El Tabakh, 2000), indicating oxidizing conditions, in contrast to black crystals near the bottom of some dark beds due to the predominance of nonphotosynthetic, sulfate-reducing bacteria. The most characteristic feature of this facies is the continuous crystal growth that gives the selenite its massive aspect with little macroscopically discernible bedding. According to Ciarapica *et al.* (1985), the observed structure is nearly the same as the idiotopic massive bank type.

Under the polarizing microscope, the selenite crystals are relatively large (~4cm) and have interlocking boundaries. They form clusters of mainly sub-vertically arranged crystals. The crystals exhibit perfect crystal

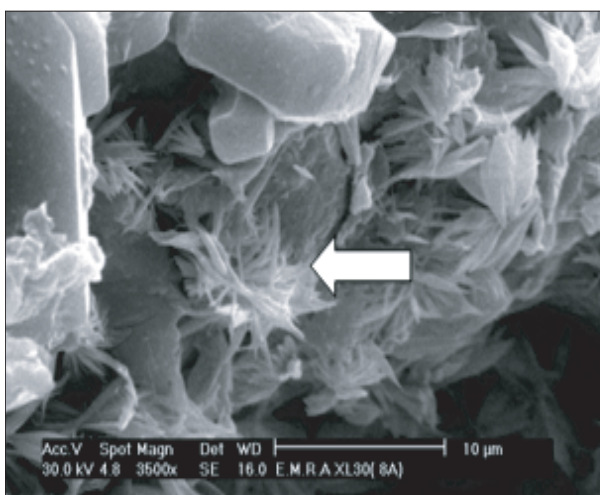
faces and distinct cleavages with the intercalation of thin gypsified algal-mat filaments. Similar filamentous bacterial forms were identified in upper Miocene (Messinian) primary gypsum in Italy (Panieri *et al.*, 2008; 2010).

### Laminated gypsum

This facies exhibits conspicuous lamination and varies from rhythmically laminated grass-like selenite to microbialite (stromatolite) subfacies. In contrast to the selenite layers that indicate periods of gypsum saturation (spring and summer), the rhythmic darker lamellae imply a flourishing bacterial and algal population during the winter wet seasons. Thus, as long as the salinity is lower than the gypsum saturation limit, the development of carbonate patches and fibrous glauberite crystals between gypsum lamellae is expected in intimate association with the bacterial mats (Fig. 4).

### Grass-like selenite

Macroscopically, the grass-like subfacies exhibits successive gypsum lamellae (up to 1cm thick) of various colors, *i.e.* white, brown, green, buff and black, depending on the prevailing bacterial communities. Thin gypsified algal mats and dark microcrystalline veneers of clay and minute fragmented gypsum grains separate the successive lamellae (Fig. 5A). However, the considered variety could be classified as rhythmic primary gypsum (Carozzi, 1960), laminas and beds (Ciarapica *et al.*, 1985) or a grass-like type (Babel, 2005; Becker, 2005 and many others). Under the polarizing microscope, the successive selenite lamellae are of planar contacts and comparable size. Each selenite lamella is composed of twinned (V-shape) crystals and is covered by a thin veneer of gypsified algal mats that acted as a substratum for the next growing selenite lamella during the gypsum saturation seasons (spring and summer).



**FIGURE 4** | SEM micrograph showing fibrous glauberite crystals (white arrow).

### Gypsum microbialite (stromatolite)

This is the microcrystalline laminated gypsum subfacies with vertically stratified colored zones (white, green, brown and black) of few mm to ~1cm thick. It exhibits characteristic doming and crenulation (Fig. 5B), due to combined effects of buckling of the growing gypsum crystals and the action of accumulated biogas (methane) beneath the algal mats, giving rise to a stromatolitic structure (Krumbein, 1983). In general, the considered subfacies exhibits a marked porosity and well established fenestral structure and could be considered as gypsum microbialite.

A gypsum microbialite is an organo-sedimentary deposit, formed by the interaction of a microbial community (Babel, 2005; Babel *et al.*, 2011). Under the polarizing microscope, and compared to the other petrographic types, gypsum crystals of this variety are finer in size. It exhibits characteristic bedding, with the development of thin streaks of algal mats and carbonate patches between gypsum lamellae. Characteristic lenticular gypsum crystals (< 1mm size) are recorded (Fig. 5C) in intimate association with the cyanobacterial mats.

In fact, the observed organic lamination with different colorations and grain-size gradation reflects, on one hand the prevailing oxidation/reduction conditions during gypsum crystallization and then the seasonal variation in brine concentration on the other hand. The role of bacterial mats in mineralization and development of various gypsum types are discussed in detail by Ali-Bik *et al.* (2011 and references therein). In general, these microorganisms and their metabolism have effective role in i) building organo-sedimentary outcrops (microbilitate and laminated stromatolite), ii) biogenic mineralization by regulating the physico-chemical conditions on a micro-scale; and iii) shaping crystal morphology and mineral texture. Thus, the development of lenticular gypsum morphology in the present example (*cf.* Fig. 5C) is attributed to metabolic action of cyanobacteria (Ali-Bik *et al.*, 2011). According to Cody and Cody (1989) the development of lenticular gypsum morphology is attributed to the interaction of certain organic molecules, such as enzyme  $\alpha$ -amylase which act as crystallization inhibitor by decreasing the growth gypsum crystals in c [001] direction and inducing nucleation of d [101] Montmartre twins.

### DIAGENETIC ALTERATION

The diagenetic impact on sediments is generally enhanced over time. However, in the studied recent gypsum deposits slight post-depositional changes are clearly apparent. Destruction of gypsum mounds, methane

emission (due to decay of organic matter) at the western side of the pond (close to pond 2) and dissolution of gypsum crystals are reported. Different generations of gypsum precipitation that imply varied growth rates under fluctuating physio-chemical conditions have been observed. Accumulations of the new gypsum precipitation accompanied by pressure-overgrowth may affect the early-formed gypsum crystals leading to undulose extinction (Fig. 6A) and recrystallization. Deformation may also lead

to cracking and disintegration of selenite crystals (Fig. 6B) or may proceed to complete fragmentation with changes in optical continuity of the deformed crystal.

Recrystallization of subaqueous primary gypsum might occur in response to pressure overgrowth, temperature rise and the fluctuation of brine composition and saturation level. As a result, the smaller fragmented gypsum crystals may transform to hemihydrate, bassanite or even to anhydrite (Fig. 6C). The latter phase exhibits higher interference colors than gypsum. Alteration on the peripheries of the gypsum crystals in response to dissolution is also recorded.

## DISTRIBUTION OF FACIES

The distribution of gypsum facies in pond 3 is illustrated in Figure 7, whereas close to pond 2, the outcrop is relatively soft with an offensive odor due to the decay of organic matter. The thickness of accumulated gypsum deposits at the bottom of pond 3 was about 10cm as checked through dragging and coring. They form a dark-grey, friable crust and more or less resemble the microbialite subfacies. Both of laminated gypsum subfacies (microbialite and grass-like selenite) are distributed along the northern and the southern banks of pond 3. On the southern bank, the massive selenite facies is reported, and commonly lays in between the two common subfacies (microbialite and grass-like selenite).

## CHEMICAL COMPOSITION

### Water chemistry

The chemical composition of Lake Quaroun water and of concentration pond 3 (averages for 8 months from January to August, 1999), as well as seawater as given by Ali and El Sheikh (2004) are summarized in Table 1. Other parameters, including pH, electrical conductivity and total dissolved salts of the collected brine samples are given in Table 2.

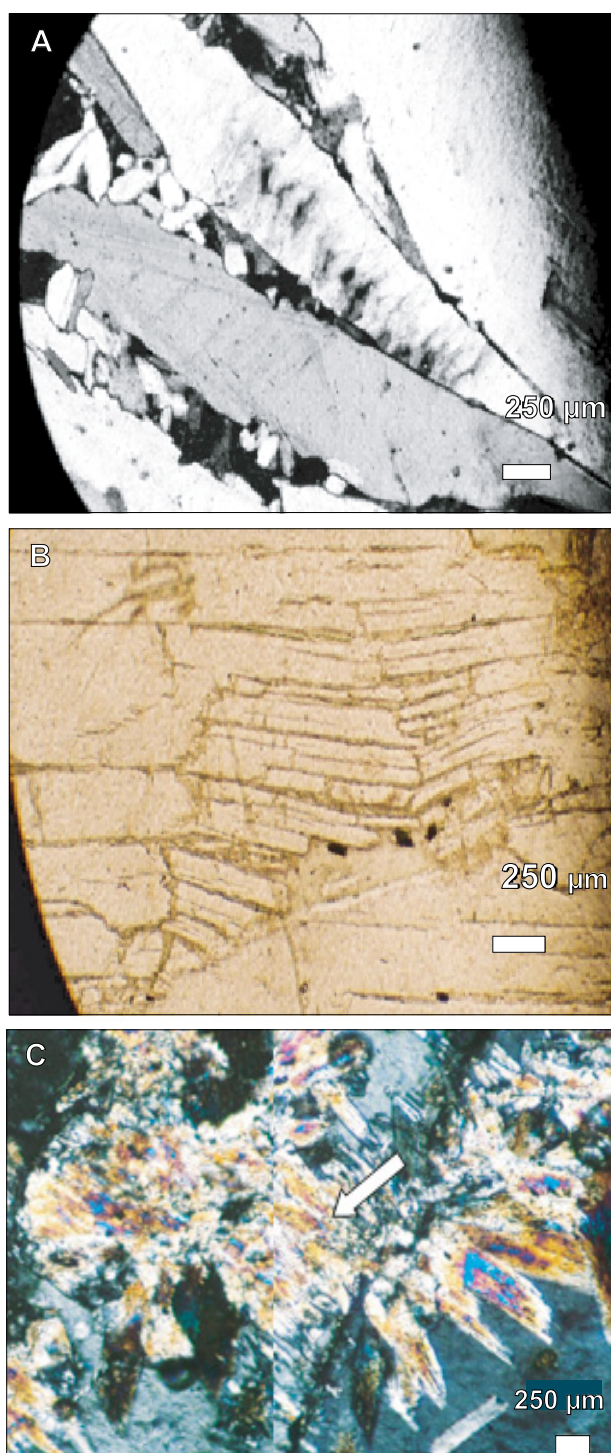
In terms of Total Dissolved Salts (TDS), Lake Quaroun and seawater are more or less comparable. In comparison with seawater, Lake Quaroun water is conspicuously enriched in sulfate, bicarbonate and to lesser extent in calcium. However, the concentrations of K and Cl are low in Lake Quaroun water.

Regarding the abundances of cations and anions in the Lake Quaroun saline water, the following distribution is observed:  $\text{Na} > \text{Mg} > \text{Ca} > \text{K}$  and  $\text{Cl} > \text{SO}_4 > \text{HCO}_3$ . A similar ion distribution is traceable in concentration pond 3 (Table 1). TDS in the collected brine samples (1S - 6S) as shown in Table 2 vary from 110.55 to 192.36g/l. The lowest value represents the collected brine from the inlet (close



**FIGURE 5** | A) Laminated selenite (grass-like subfacies). B) Fine grained gypsum microbialite (stromatolite) subfacies. C) SEM micrograph of fine lenticular gypsum crystals embedded in microbial biofilm (gypsum microbialite).

to the pond 2) of pond 3; in general, the TDS increased in eastward direction. Electric conductivity ranges from 1424 to 2222 $\mu$ S, while pH values fluctuate between 7.86 and 8.19.



**FIGURE 6** | A) A selenite crystal exhibiting undulose extinction in response to a slight deformation. Crossed-polarized light. B) Disintegration and cracking of a selenite crystal. Plane -polarized light. C) Alteration of gypsum to anhydrite (white arrow). Crossed polarized light.

## Whole-rock chemistry

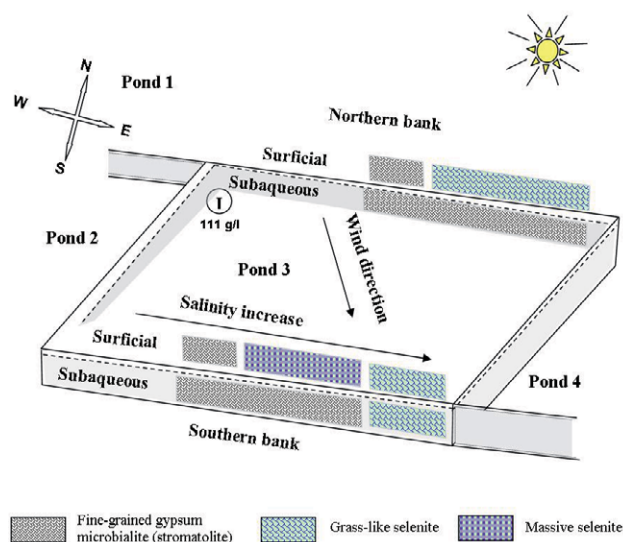
Bulk chemical composition of the studied gypsum and sediments and their calculated normative components are given in Table I, Electronic Appendix available at [www.geologica-acta.com](http://www.geologica-acta.com). Of the main constituents (gypsum-combined water, SO<sub>3</sub> and CaO), the massive selenite facies exhibits the highest contents, *i.e.* highest purity.

In term of major and trace element contents, both grass-like selenite and microbialite subfacies are more or less comparable as they contain small amount of clay fraction (aggregates). Compared to the massive selenite facies, both grass-like selenite and gypsum microbialite subfacies are depleted in Sr and Co contents; while Ba and Y are enriched.

Regarding the normative components, the massive selenite facies is more homogenous, as expected, exhibiting the highest gypsum and anhydrite content (up to 95% wt.). In the grass-like selenite and microbialite subfacies of the southern and northern banks, normative gypsum and anhydrite content is ranging from 93 to 86% by weight. The recently precipitated gypsum contains about 90% normative gypsum and anhydrite by weight. Turning to bottom sediments, the sample 3B (high-salinity zone) contains about 84% normative gypsum and the sample 1B (low-salinity) contains the highest amount of clay fraction (up to 26% by weight).

## DISCUSSION

Enhancing the wind-action effect, the gypsum outcrop on the southern bank is more mature in terms of crystal size



**FIGURE 7** | A sketch showing the distribution of gypsum facies in pond 3, above and below the temporarily brine level.

**TABLE 1** | Hydrochemistry of Lake Quaroun water and pond 3 (averages for 8 months from January to August, 1999), as well as sea water, as given by Ali and El Sheikh (2004). All values are given in g/l

	Seawater	Lake Quaroun	Pond 3
Na	10.56	9.89	73.10
Ca	0.40	0.48	0.38
Mg	1.27	1.31	8.70
K	0.38	0.22	0.79
Cl	18.98	12.74	96.30
SO <sub>4</sub>	2.56	9.41	57.30
HCO <sub>3</sub>	0.14	0.30	0.64
Salinity (TDS)	34.40	34.30	244.30

and degree of lithification, compared to that of the northern side. Similarly, the salinity gradients change eastwardly, and the sediment maturity has obviously increased in the same direction also. Hence, at the northwest corner (close to ponds 1 and 2), no gypsum mounds have been recorded; instead, newly formed, very fine gypsum crystals have been visually monitored (at the time of sampling), indicating a fast nucleation rate. These new gypsum crystals will more readily dissolve than accumulate as gypsum mounds. In general, the dissolution and subsequent destruction of incipient gypsum mounds are more common in the western part of the pond, where the lowest salinity values are expected (about 111 g/l at the time of sampling). Meanwhile, distribution of selenite varieties (both massive and laminated grass-like) is confined to the higher salinity area, in comparison with the fine-grained gypsum microbialitic (stromatolitic) type.

### Modeling of pond 3

In general, the varied evaporite facies in a particular deposition basin reflect the characteristics of their respective stratified brine zones (*e.g.* Ortí Cabo *et al.*, 1984a; 1984b; Peryt, 1996; Babel, 2004; 2007; Schreiber *et al.*, 2007). Regarding pond 3 as a gypsum deposition basin, its main features argue in favor of both the selenite shoal and monomictic selenite pan models of Babel (2004). The used terminology in the proposed models follows mainly Babel (2004 and references therein) and Schreiber *et al.* (2007):

- i) Monomictic: a lake, which is stratified only in one season (winter or summer) of the year and becomes homogeneous through entire mixing in other seasons.
- ii) Polymictic: a lake, with frequent periods of mixing per year.
- iii) Epilimnion: an upper layer of a seasonally stratified water body.
- iv) Hypolimnion: represents the lower layer of a seasonally stratified water body.
- v) Pycnocline: the boundary between two different brines (low and high salinities).

**TABLE 2** | Total Dissolved Salts (TDS), pH and electrical conductivity of the collected brine samples (April 2006) from pond 3

Sample	TDS	pH	EC (μS)
1 S	110.55	8.11	1424
3 S	170.85	8.19	2220
4 S	192.36	7.86	2222
5 S	150.15	8.14	1939
6 S	161.00	8.10	2074

In both shoal and monomictic pans models, gypsum microbialite and selenite reflect different salinity environments, whereas the latter is confined only to the relatively higher salinity brines and more resistant to dissolution processes (*e.g.* Ortí Cabo *et al.*, 1984a; 1984b; Babel, 2004). In the selenite shoal model, the salinity gradient is horizontal; hence, the selenite crust on an evaporite shoal is thin and more susceptible to dissolution. In contrast, the salinity gradient is vertical in the monomictic pan model during the wet seasons; hence, the level of the pycnocline is crucial, whereas the selenite growth is mainly related to evaporation of the epilimnion and occasionally to interaction with the hypolimnion. The latter case is maintained in shallow and large pans when the hypolimnion is partially exposed to the surface, and hence, to evaporation by the action of strong unidirectional wind. Steady strong wind acting in this way, may lead to complete mixing (polymictic condition).

Regarding the water of Lake Quaroun (~3.5% salts), the gypsum saturation level is approached by reducing the water budget via evaporation to about 20% of its original volume (Ali *et al.*, 2007). Concentration pond 3, as a basin of gypsum deposition, experienced two evolutionary stages: i) the first stage extended from the onset of gypsum saturation until 1995, where the pond acted as a natural gypsum precipitation basin; and ii) the second stage continued from 1995 to the present, during which time the pond management policy was adopted. The aim of this process was to maximize the productivity of thenardite (Na<sub>2</sub>SO<sub>4</sub>) and edible NaCl by preventing any salt precipitation in pond 4 by controlling the concentration levels and salinities of the other ponds.

The pond management process of Lake Quaroun involves two complementary steps. The first concerns decreasing the concentration of Ca ions in ponds 3 and 4 by enhancing the aragonite precipitation in pond 2 using specific types of algae (Wali *et al.*, 1997). In the second step, a brine-mixing technique (back drift) is employed by pushing the hyper-saline brine from the end of pond 4 back to pond 3 to maintain the salinity of pond 4 as planned.

Metwally (2002) recorded the fluctuation of brine levels and subsequent concentrations in pond 3 for three successive years (1996-1998), and he noted a substantial fluctuation in



brine level and subsequent seasonal concentrations during the year 1998 compared to those of years 1996 and 1997 (Fig. 8). In winter season of 1998, pond 3 received excess amounts of low-salinity water (under-saturated with respect to gypsum) from Lake Quaroun through pond 2, representing the wet season of the perennial basin. During the dry season (spring and summer), there was a conspicuous increase in the reported concentration, providing favorable conditions for gypsum precipitation. According to Ali *et al.* (2007 and references therein) the biota of pond 3 include algae, diatoms and brine shrimps; the collected algae from pond 3 were identified as cyanobacteria and chlorophyta (Ali-Bik *et al.*, 2011).

The relatively shallow brine depth (2.30m) of pond 3 and the surface salinity gradient towards pond 4 argue in favor of a selenite shoal model, except for the region close to pond 4. Hence, the relatively persistent brine budget near pond 4 (higher salinity zone), provides the opportunity for pycnocline development (stratification) in this part of the pond during wet seasons when low salinity influx is supplied from pond 2 (*cf.* Fig. 8).

However, the western portion of pond 3, acts as a low-salinity zone, following either the selenite shoal or the monomictic pan models. Hence, fine-grained gypsum

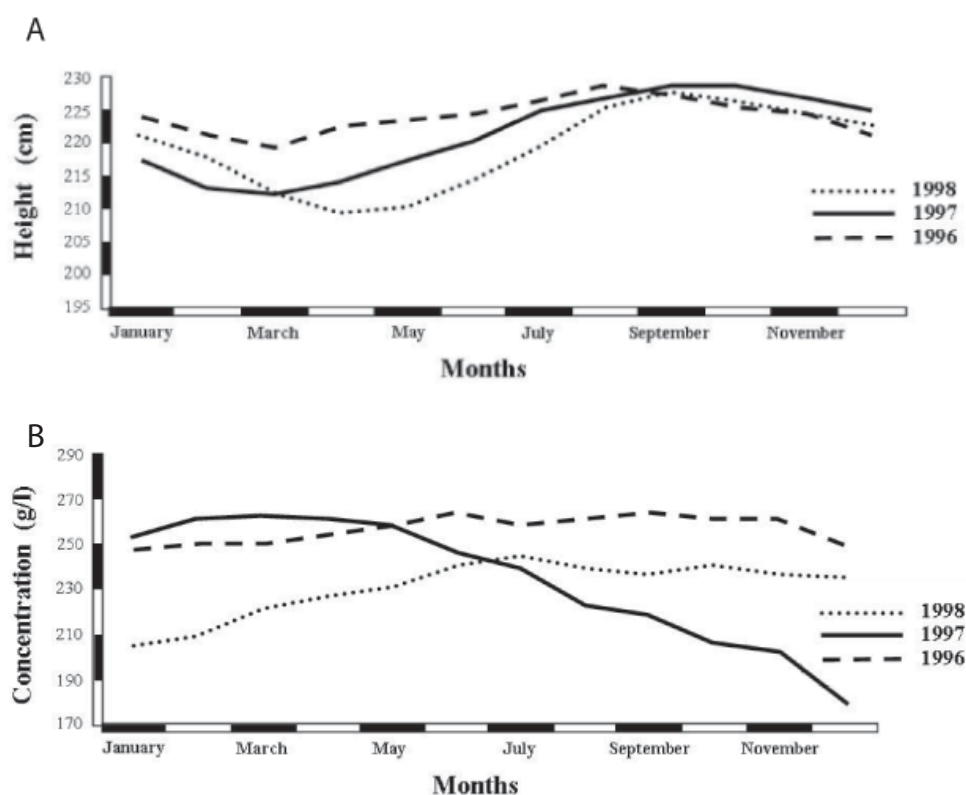
microbialite with a thin selenite crust is expected to be developed during the dry seasons of such environments (*e.g.* Ortí Cabo *et al.*, 1984a; 1984b; Babel, 2004). The area (I) at the northwestern part of the pond (Fig. 7) is characterized by the lowest degree of salinity and minimal wind action; hence, dissolution of recently formed gypsum is typical, *i.e.*, the opportunity to build gypsum mounds is greatly decreased. Generally, during wet seasons (dilution process from pond 2), dissolution of the pre-existing gypsum is common.

In contrast, the eastern portion of pond 3, close to pond 4, is characterized by relatively higher salinity environment; hence, the development of selenite, following any of the proposed models is the case. In both models, this zone is characterized by continual growth (syntaxial) of selenite due to permanent brine saturation with gypsum (Babel, 2004). In this manner, pond 3 acts as a shallow-brine selenite pan.

Regarding the evolution of brines in pond 3, two stages are distinguished:

#### ***From the onset of gypsum saturation until 1995***

During this period, pond 3 acted as a natural deposition basin for selenite, more or less simulating the entire process



**FIGURE 8** Fluctuation of brine levels (A) and ionic concentrations (B) in pond 3 after Metwally (2002).

of selenite deposition in the selenite shoal model of Babel (2004). In this situation, the pond received from time to time (wet seasons) the inflow of brine from pond 2, with a salinity of about 130g/l. Then, the brine flowed towards pond 4 with a gradual increase of salinity (up to about 260 ±20g/l), due to evaporation (dry seasons) and subsequent gypsum precipitation. In general, the precipitation of gypsum begins at a salinity of about 150‰.

Hence, the distribution of gypsum facies in the pond reflects the lateral variation in the salinity environments. Fine-grained gypsum microbialite subfacies is confined to the low-salinity zone (western portion of the pond) and selenite deposition to the higher salinity environment.

### **From 1995 to the present**

Pond management policy and repeated dilution of the pond turned it into a true shallow-brine selenite pan, with development of a seasonal pycnocline and stratification (particularly at the eastern end, close to pond 4). In such an environment, laminated selenite (grass-like type) may develop. For this to happen, the mixing (back-drift) process, *i.e.* pumping of the heavier hyper-saline brine with salinity that exceeds the upper salinity limit of gypsum crystallization (> 320g/l salinity) from pond 4 to the lower (hypolimnion) zone of pond 3, is crucial. Thus, the pre-existing selenite (pre-management) will be partially or totally dissolved by the hyper-saline brine of pond 4.

In addition, the conspicuous deficiency of Ca ions in pond 3 due to pond management greatly diminishes the opportunity of developing large selenite crystals, as it is a matter of concentration rather than salinity. Accordingly, the reported massive selenite facies represents an earlier gypsum outcrop that developed in conditions of high salinity (hypolimnion zone) in presence of excess of aqueous Ca and absence of mixing, *i.e.* during pre pond-management. Simulation of such conditions in nature may give rise to giant and saber gypsum facies, for example, the northern Carpathian Foredeep (Babel, 2005).

## **CONCLUSIONS**

From a geologic point of view, the clay fraction that settled and salts precipitated throughout the concentration ponds at EMISAL (1, 2 and 3) represent a thin sedimentary succession. Clays occur at the base of the section (pond 1), followed upward by calcium carbonates (ponds 1 and 2) and topped by primary gypsum in pond 3. In the latter, gypsum precipitates formed with variable fabrics, crystal morphologies and sizes on the floor and along the banks. Accordingly, the primary gypsum is classified into two main facies: i) massive selenite; and ii) laminated gypsum. The second facies is subdivided into two subfacies, namely

grass-like selenite and fine-grained gypsum microbialite (stromatolite). The massive selenite facies is composed of clusters of single and twinned crystals along (100), with upward pointing re-entrant angle. The laminated gypsum facies exhibits conspicuous bedding and varied from rhythmically laminated grass-like selenite to microbialite (stromatolite). The variation from massive selenite to laminated-gypsum facies implies fluctuation in lateral and vertical salinity profiles, seasonal brine concentrations and wind action.

Many signs of diagenetic modification are apparent including: i) dissolution; ii) slight to moderate crystal deformation and finally; and iii) recrystallization, including the dehydration of primary gypsum. Chemically, the massive selenite facies exhibits the highest purity; both grass-like selenite and fine-grained gypsum microbialite subfacies are more or less comparable.

Modeling of pond 3 should be considered in the light of the evolution history of the brine (pre- and post-management of the pond). During the first stage, the pond acted as a natural basin of selenite crystallization (selenite-shoal model), and the distribution of gypsum facies in the pond reflects the laterally varying salinity. The fine-grained gypsum microbialite subfacies is confined to the low-salinity zone (western portion of pond 3) and selenite to the higher salinity environment. Dissolution and fragmentation of the selenite crust at the western portion of the pond (low-salinity area) is common in the wet seasons. Meanwhile, the relatively higher salinity environment in the eastern portion, close to pond 4 (higher salinity zone) is characterized by continual growth of selenite (syntaxial growth) with little dissolution signature.

After adopting pond-management policy and repeated dilution, the pond has been turned to a true shallow-brine selenite pan. Hence, due to different gradients of Ca concentration, the laminated grass-like selenite subfacies may develop at higher salinity zones instead of massive selenite facies, *i.e.*, it is the matter of continuous concentration rather than salinity.

Therefore, the pre-existing (pre-management) large selenite crystals will partially or totally dissolve during the back-drift processes due to contrast with the hyper-saline brines (>320g/l salinity) of pond 4. Accordingly, the reported, massive selenite facies might represent an earlier gypsum outcrop that developed in the presence of excess Ca ions (pre-management) perhaps under deeper water levels and without subsequent dilution processes.

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## ELECTRONIC APPENDIX

**TABLE I** | Bulk chemical composition of different gypsum facies along the banks and on the floor of pond 3. Averages are also given as AV<sub>1</sub> and AV<sub>2</sub>

Sample N°	Southern and northern banks							
	Massive selenite				Grass-like selenite			
	3A	4A	5A	AV <sub>1</sub>	6A	6Q	10A	AV <sub>2</sub>
wt. %								
Combined H <sub>2</sub> O(250C°)	19.11	19.26	19.11	19.16	18.74	17.14	16.36	17.41
SO <sub>3</sub>	44.66	44.68	44.82	44.72	43.46	40.14	41.07	41.56
CaO	31.49	30.80	32.10	31.46	30.89	29.05	29.52	29.82
Al <sub>2</sub> O <sub>3</sub>	0.09	0.05	0.08	0.19	0.53	0.57	0.43	0.28
Fe <sub>2</sub> O <sub>3</sub> Tot.	0.05	0.04	0.05	0.14	0.39	0.43	0.32	0.20
MgO	0.12	0.19	0.08	0.13	0.23	0.49	0.69	0.47
SiO <sub>2</sub>	0.45	0.31	0.34	0.37	1.36	3.15	2.45	2.32
TiO <sub>2</sub>	--	--	--	--	--	0.08	0.06	0.07
P <sub>2</sub> O <sub>5</sub>	0.01	0.01	0.01	0.01	--	0.01	0.02	0.02
Na	0.70	0.45	0.33	0.49	0.53	1.28	1.05	0.95
K	0.02	0.02	--	0.02	0.05	1.10	0.10	0.42
Cl	--	0.63	0.24	0.44	0.95	2.26	1.44	1.55
Sr	--	4678	4300	4489	3675	3473	4717	3955
ppm								
Ba	--	119	113	116	109	154	161	141
Ni	--	49	67	58	38	60	65	54
Rb	--	7	9	8	12	10	9	10
Zn	--	18	--	18	20	11	26	19
Mn	--	35	1	18	9	140	278	142
Cu	--	--	10	10	46	14	28	29
Br	--	--	--	--	--	58	53	55
wt. %								
Combined H <sub>2</sub> O(550C°)	1.89	1.85	0.83	1.52	2.03	2.79	3.75	2.86
Moisture (H <sub>2</sub> O-45C°)	0.06	0.07	0.13	0.09	0.13	0.18	0.10	0.14
L.O.I	21.60	22.80	21.40	21.93	21.90	22.50	21.80	22.07
Estimated CO <sub>2</sub>	0.54	1.62	--	1.08	--	2.39	1.59	1.99
Normative wt. %								
Gypsum	91	92	91	91.3	90	82	78	83.3
Anhydrite	4	3	4	3.75	3	3.5	8	4.8
Calcite	0.07	--	1	0.5	0.5	0.5	--	0.50
Dolomite	0.5	--	0.30	0.4	1	2	--	1.5
Halite	0.30	--	0.30	0.30	1	2.37	2.13	
Aggregates	0.6	0.4	0.47	0.5	1.69	4.07	3.45	3.07

Estimated CO<sub>2</sub> = L.O.I. - (Moisture + Combined H<sub>2</sub>O at 250 C° + Combined H<sub>2</sub>O at 550 C°).

Combined H<sub>2</sub>O at 250 C° denotes the crystalline water in gypsum structure.

Combined H<sub>2</sub>O at 550 C° denotes the released water from clay structure.

Normative components are mainly calculated according to Ragab (1967).

TABLE I | Continued

Sample N°	Gypsum microbialite (stromatolite)								New precipitates		Gypsum microbialite	
	1A	1Q	2Q	3Q	4Q	5Q	8A	9A	11Q	1B	2B	3B
Wt.%												
Combined H <sub>2</sub> O(250C°)	19.08	17.67	16.58	17.68	17.64	18.11	17.59	16.82	16.85	8.12	14.07	15.93
SO <sub>3</sub>	43.48	40.82	39.54	41.35	41.86	41.11	40.83	39.74	42.80	24.25	39.98	40.13
CaO	30.10	29.52	29.62	29.45	30.29	29.44	29.83	27.99	29.88	21.79	27.61	27.58
Al <sub>2</sub> O <sub>3</sub>	0.17	0.22	0.50	0.29	0.17	0.20	0.83	0.87	0.75	3.98	0.92	0.79
Fe <sub>2</sub> O <sub>3</sub>	0.12	0.16	0.37	0.20	0.10	0.15	0.62	0.64	0.54	2.30	0.62	0.46
MgO	0.37	0.49	0.44	0.45	0.42	0.41	0.45	0.64	0.39	1.67	0.53	0.72
SiO <sub>2</sub>	0.86	1.50	3.80	2.54	1.26	1.94	3.35	0.44	2.75	19.80	4.17	4.35
TiO <sub>2</sub>	--	--	0.06	--	--	--	0.10	0.10	--	--	--	--
P <sub>2</sub> O <sub>5</sub>	0.02	0.02	0.02	--	0.01	0.01	0.02	0.02	0.03	0.10	0.04	0.03
Na	0.78	1.23	1.07	1.11	1.38	1.36	0.79	1.60	0.71	1.97	1.27	1.38
K	0.05	0.06	0.10	0.07	0.07	--	0.10	0.12	0.08	0.45	0.12	0.12
Cl	1.24	2.00	1.88	--	2.35	2.50	1.34	--	0.9	2.54	2.16	2.09
ppm												
Sr	4561	3534	3557	--	2940	3385	3283	--	--	3071	2716	3611
Ba	166	166	104	--	138	143	127	--	--	240	116	154
Ni	41	40	8	--	41	55	42	--	--	--	31	36
Rb	13	3	9	--	13	12	7	--	--	--	12	4
Zn	13	8	8	--	14	11	15	--	--	47	30	11
Mn	192	100	84	--	--	18	212	--	--	393	218	63
Cu	36	41	38	--	9	43	----	--	--	--	22	70
Br	49	46	155	--	55	60	150	--	--	255	62	62
Wt.%												
Combined H <sub>2</sub> O(550C°)	2.32	3.35	3.34	3.79	2.99	2.24	2.70	2.89	2.89	7.67	5.55	3.37
Moisture (H <sub>2</sub> O-45C°)	0.19	0.15	0.17	0.15	.22	0.07	0.14	0.18	0.26	1.09	0.26	0.79
L.O.I	22.8	23.70	22.30	22.40	22.90	22.80	21.60	21.90	21.49	22.59	23.99	23.59
Estimated CO <sub>2</sub>	1.21	2.53	2.21	0.78	2.05	2.38	1.17	2.01	4.38	--	4.11	3.50
Normative Wt.%												
Gypsum	91	84	79	85	84	87	84	80	81	39	67	76
Anhydrite	1	3	5	3.5	4.5	1.5	3	4	9	10.5	15	8
Calcite	--	0.5	2	--	0.7	0.5	1	--	--	4	--	--
Dolomite	--	2	2	--	2	1	2	--	--	--	--	--
Halite	0.74	2.39	2.39	2.24	1.10	2.8	1.1	3.12	--	--	--	--
Aggregates	1.15	2.15	4.67	3.03	1.53	2.29	4.8	3.95	4.04	26.08	5.71	5.60