Spheroidal weathering of basalt from Gebel Qatrani, Fayum Depression, Egypt

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Abstract

Background: Three stages of basalt weathering are known: early or incipient weathering, intermediate weathering and advanced weathering. The Late Oligocene basalt of Gebel Qatrani in Fayum Depression, Egypt, shows signs of early weathering, particularly exhibited in basalt spheroids found at the top of the basalt flow. The present paper gives the results of detailed petrographical, mineralogical and geochemical study of the weathering of these basalt spheroids.

Results: The core-stones of the basalt spheroids are composed of phenocrystals of plagioclase feldspars and clinopyroxenes set in a groundmass of tiny feldspars and pyroxenes, relatively altered olivine and opaque minerals. The basalt is subalkali (tholeiitic). The outer weathered shells surrounding the core-stones are composed of partly altered feldspars and pyroxenes. The calculated weathering indices show that there is marked weathering trend from the core-stones of the spheroids to the outer shells. The chemical mobility of the elements shows marked depletion of Mg, Ca, Na and K from the core-stones to the outer shells due to the weathering of olivine, pyroxene and feldspars. The trace elements Rb, Sr, Ni, V, Cr and Zn are also depleted.

Conclusions: The weathering of basalt spheroids from Gebel Qatrani, Fayum Depression, Egypt, is of the incipient type. The degree of weathering from the core-stones of the basalt spheroids to the corresponding weathered shells indicates that the weathering occurred under predominantly semiarid to arid conditions.

Keywords: Basalt spheroids, Gebel Qatrani, Fayum depression, Weathering indices, Element mobility

Introduction

The Fayum Depression has an area of about 12,000 km² and lies about 100 km south-west of Cairo (Fig. 1). The geology of the area was studied in detail by Beadnell (1905), and his work has remained the basis for subsequent studies (Said 1962; Vondra 1974; Bown and Vondra 1974 1974; Bown and Kraus 1988; Gingerich 1992, 1993, among others). These studies indicate that the sedimentary successions encountered in Fayum Depression are of middle and late Eocene and Oligocene age and can be divided into the following, formations, from base to top (Fig. 1):

1 Wadi Rayan Formation This formation is of middle Eocene age and forms the base of the Fayum Depression and is exposed in its southern part, reaching a thickness of about 130 m. It is mainly composed of limestone, with *Nummulites gizehensis*, argillaceous sand and sandy shale (Beadnell 1905; Said 1962). The formation was deposited on an open marine continental shelf. Abd-Elshafy et al. (2007) analyzed the faunal community of the formation at three locations in Wadi Rayan and concluded that the faunal content, as well as the lithology, indicates the alternation between two transgressive sedimentary cycles enclosing a regressive phase.

2 Gehannam Formation Also known as ravine beds, this formation, which is of middle Eocene age, conformably overlies the Wadi Rayan Formation and attains a thickness of about 70 m. It is mainly com-
posed of gypsiferous shale, marl, limestone and sands and was deposited on a shallow, but open marine shelf (Gimgerich 1992).

3 **Birket Qarun Formation** This formation is of late Eocene age and conformably overlies the Gehannam Formation. It has a thickness of about 50 m and is composed of sandstones and shale, with a few bands of limestone. Lithologically the Birket Qarun Formation is indistinguishable from the underlying one, but is paleontologically distinct, as it carries different fossil species (Beadnell 1905). The Birket Qarun Formation was a submerged barrier bar, rather than a beach complex (Bown and Kraus 1988; Gingerich 1992).

4 **Qasr El-Sagha Formation** This formation is also of late Eocene age and conformably overlies the Birket Qarun Formation. It varies in thickness from one location to another in Fayum Depression and is generally in the range of 175–200 m. Four distinct facies have been recognized in Qasr El-Sagha Formation: (a) interbedded claystone, siltstone and quartz sandstone facies; (b) quartz sandstone facies; (c) arenaceous bioclastic carbonate facies; and (d) gypsiferous and carbonaceous laminated claystone and siltstone facies (Vondra 1974). (Facies (a) and (b) have been called the Dir Abu Lifa Member, and overlies facies (c) and (d) which have been called the Temple Member (Bown and Kraus 1988). Both the Dir Abu Lifa and the Temple Members represent the upper part of Qasr El-Sagha Formation. At locations south and west of Qasr El-Sagha, Gingerich (1992) introduced two additional members: the Harab Member which
forms the middle part of Qasr El-Sagha Formation, and the Um Rigl Member which forms the lower part of the Formation. The Harab Member is composed of brown shale, whereas the Umm Rigl Member is of similar facies as the Temple Member but is separated from it by Harab Member. The Umm Rigl Member was deposited in a shallow outer lagoonal environment; the Harab Member in a deeper central lagoonal environment; the Temple Member in a shallow inner lagoonal environment; and the Dir Abu Lifa Member in a non-lagoonal (deltaic or interdeltaic) environment (Vondra 1974; Bown and Kraus 1988; Gingerich 1992).

5 Jebel Qatrani Formation This formation is of Oligocene age and is separated from the underlying Qasr El-Sagha Formation by an unconformity that involved erosion of up to 70 m of Qasr El-Sagha strata in places before the deposition of Jebel Qatrani Formation (Gingerich 1992). The lithology of Jebel Qatrani Formation is rather complex and is composed of about 340 m of variegated alluvial rocks: fine to coarse sandstone, granule and pebble conglomerate, sandy mudstone, carbonaceous mudstone and limestone. The lower part of Jebel Qatrani Formation is assumed to have been deposited by meandering streams, whereas the upper part of the formation was deposited under the influence of an encroaching marine strandline (Bown and Vondra 1974, 1974; Bown and Kraus 1988).

In later Oligocene times, a tensional tectonic episode occurred and was accompanied by northwest-trending normal faults throughout the northern parts of the Eastern and Western Deserts of Egypt. Basaltic lavas were extruded from these fissures at different locations (Rittmann 1954). In the northern part of Fayum Depression, these basaltic lavas capped the upper Qatrani escarpment varying in thickness from 2 to 25 m. (The latter thickness has been recorded at Widan El Faras.) In outcrops where the basalt is thinnest, it appears that it was a single flow; however, some authors believe that there were two or more flows (Fleagle et al. 1986; Bown and Kraus 1988). The flows overlie the Jebel Qatrani Formation with a pronounced erosional unconformity. Meneisy and Abdel Aal (1984) determined the whole rock K/Ar ages of two samples of these basalts as 23.8 ± 0.7 Ma and 24.4 ± 9.7 Ma, and Bown and Kraus (1988) reported two ages: 24.7 ± 0.4 Ma and 27.0 ± 3 Ma. Meneisy (1990) compiled other K/Ar age data of 27 ± 1 Ma, 25 ± 1 Ma and 23 ± Ma. However, Bown and Kraus (1988) indicated that a sample of basalt taken from the lowest part of the flow near the top of Jebel Qatrani Formation gave an age of 31 ± 1 Ma and, therefore, stated that the basalt might be of late Early Oligocene to Late Oligocene. However, on average the previously mentioned data show that the age of Gebel Qatrani basalt is 25.7 Ma, i.e., of Late Oligocene age, according to the International Chronostratigraphic Chart (Cohen et al. 2013; updated 2017).

The petrography and geochemistry of Gebel Qatrani basalt are described in several studies (El-Hinnawi 1965; El-Hinnawi and Abdel Maksoud 1968, 1972; Heikal et al. 1983; Sharara 1984; Abdel Meguid et al. 1992; Endress et al. 2011). The basalt has generally an ophitic texture, with plagioclase feldspars clinopyroxene phenocrysts set in a fine groundmass of plagioclase, clinopyroxene, occasional olivine (mostly altered), opaque minerals and altered glass.

The basalt of Gebel Qatrani is unconformably overlain by Miocene alluvial sediments (the Khashab Formation). The contact between the basalt and the Khashab Formation is markedly erosional, and scours in the top of the basalt are filled with coarse sand, sandstones containing basalt debris and other pebble conglomerate (Said 1962; Bown and Kraus 1988). The basalt itself is not markedly weathered; no weathering profiles have been detected and the weathering observed is predominantly rind weathering. At the top of the basalt flows of Gebel Qatrani, spheroids of basalt are occasionally encountered.

The present paper describes the mineralogy and geochemistry of the cores and weathered rims of these basalt spheroids and discusses the paleo-environmental conditions of their weathering.

Samples and methods
The term ”spheroidal weathering” is used if concentric shells completely surround the core-stone (Ollier 1971). A difference exists between spheroidal weathering and exfoliation. In spheroidal weathering, the weathering process acts all around the spheroid, weathering the underside as well as the top. Patino et al. (2003) pointed out that basalt flows have cooling features that serve as hydrologic discontinuities which function as major pathways for fluids that are the agents of weathering. In incipient and intermediate weathering stages, the concentric shells surrounding the basalt spheroids still contain part of the primary minerals and/or secondary derivatives. In advanced stages of weathering, however, the shells are completely transformed into argillaceous material.

In Gebel Qatrani the basalt spheroids (Fig. 2) encountered vary in size and have gray to gray-greenish color. In the present work, several samples were collected, taking care that their outer weathered shells are more or less complete surrounding the core-stones. The collected samples were generally egg-shaped, the longest dimension of which varied from 8 to 15 cm, the intermediate dimension from 6 to 12 cm, and shortest dimension
from 6 to 10 cm. The weathered shells varied in thickness between 2 and 5 mm.

In laboratory, the outer weathered shells were carefully detached from the core-stones. Thin sections of the latter were prepared for petrographic study. The shell material was powdered for analysis by X-ray diffraction. (The analyses were carried out at the Metallurgical Institute, Tebbin, Cairo, using a Philips X-ray diffractometer; scanning was carried out at 2 Ø of 1° per minute between 4° and 60°.) The major and trace element composition of both the powdered core-stones and the shells were determined by X-ray fluorescence (XRF), at the Central Laboratories of the Egyptian Geological Survey, Dokki, Cairo.

Results

Petrography and mineralogy of the basalt spheroids

The examination of thin sections of the core-stones of the basalt spheroids under the polarizing microscope revealed that they exhibit an ophitic texture, with plagioclase feldspar and clinopyroxene phenocrysts within a fine groundmass of plagioclase, clinopyroxene, opaque minerals, glassy and altered minerals. The modal composition of the core-stones was determined by point counting. It consists of an average of 56% plagioclase feldspars, 27% clinopyroxene, 6% opaque minerals and 11% glass and alteration products.

The plagioclase phenocrysts occur as tabular subhedral to euhedral crystals with an average size of 1 × 0.35 mm. The crystals exhibit albite and albite-carlsbad twinning, often with oscillatory zoning. Some of the crystals show tiny inclusions of pyroxene. The composition of the plagioclase varies from one crystal to another (El-Hinnawi and Abdel Maksoud 1968; Endress et al. 2011). On the average, it is composed of An_{48} Ab_{30} Or_{2}. In contrast to the plagioclase phenocrysts, the plagioclase in the groundmass is present as tiny laths, about 0.5 × 0.1 mm, mingled with clinopyroxene, opaque and altered grains.

Clinopyroxene is present as anhedral to subhedral tabular phenocrysts about 1.7 × 1.3 mm. The crystals exhibit lamellar twinning, some are zoned. The composition of the clinopyroxene was determined optically and found to range from pigeonite to augite. This is in agreement with the findings of Endress et al. (2011). Some clinopyroxene crystals contain tiny inclusions of plagioclase, and some are corroded at the edges. The clinopyroxene in the groundmass is in the form of tiny grains intermixed with plagioclase, opaque and altered material. Few samples show the presence of altered olivine (iddingsite) in the groundmass.

X-ray powder diffraction analysis of the weathered shells separated from the basalt spheroids showed the presence of sharp peaks of plagioclase feldspars as the predominant minerals present. In one sample (sample No. 1) peaks of pyroxene have been detected. This indicates that the weathering of the core-stones of the basalt spheroids was at an initial (incipient) stage; the main minerals (the plagioclase and the ferromagnesian minerals) were not completely altered as in advanced weathering stages. No marked peaks of clay minerals were detected in the studied samples, although Morsy and Attia (1983) reported the presence of montmorillonite-vermiculite, chlorite and illite in two samples.

Geochemistry of the basalt spheroids

Table 1 gives the chemical analyses of core-stones and shells of seven representative basalt spheroids, and Table 2 gives the distribution of some trace elements in the studied samples. Figure 3 shows the relationship between the total alkali (Na_2O + K_2O) and silica for these samples and indicates that all samples fall below the Macdonald and Katsura boundary line (1964) in the subalkali (i.e., tholeiitic) field. The figure indicates that shells have lower content of alkalis which illustrates the loss of these elements due to weathering. On the other hand, there is a slight increase in the silica content indicating the conservative behavior of silica during incipient weathering.

To assess the extent of weathering, several weathering indices have been proposed, e.g., the weathering index of Parker (WIP, Parker 1970), the chemical index of alteration (CIA, Nesbitt and Young 1982), the chemical index of weathering (CIW, Harnois 1988) and more recently, the mafic index of alteration under oxidizing and reducing conditions (MIA(O), MIA(R)) which includes Fe and Mg-oxides (Babechuk et al. 2014). Each uses the molecular proportions of major element oxides, with variations on which oxides are included, and assumptions about element mobility. In the present work, the following indices have been calculated and used to describe the changes that accompanied the spheroidal weathering of Gebel Qatrani basalt:
The weathering index of Parker (WIP) is considered relatively sensitive to the chemical variations in the early stages of weathering because it allows for differential mobility of cations based on their bond strength with oxygen. WIP values range between 100 and 0, with smaller numbers indicating enhanced weathering. In all the studied samples (Table 1), there is a decrease of WIP from the core-stones to the corresponding shells, with varying degrees. Sample No. 3 shows the highest difference in WIP from the core-stone to the weathered

### Table 1 Chemical analyses and weathering indices

|          | 1   | 2   | 3   | 4   | 5   | 6   | 7   | Av  |
|----------|-----|-----|-----|-----|-----|-----|-----|-----|
| **Core-stone** |     |     |     |     |     |     |     |     |
| SiO$_2$   | 49.66 | 49.28 | 49.76 | 49.75 | 49.96 | 48.75 | 49.08 | 49.46 |
| TiO$_2$   | 2.19 | 2.08 | 2.12 | 2.12 | 2.1 | 2.33 | 2.89 | 2.26 |
| Al$_2$O$_3$ | 13.67 | 14.63 | 14.41 | 14.42 | 14.7 | 12.9 | 15.15 | 14.27 |
| Fe$_2$O$_3$ | 14.98 | 13.91 | 13.82 | 13.88 | 13.58 | 13.62 | 13.58 | 13.91 |
| MnO       | 0.13 | 0.15 | 0.15 | 0.15 | 0.16 | 0.15 | 0.15 | 0.15 |
| MgO       | 6.68 | 6.92 | 6.95 | 6.96 | 6.62 | 6.19 | 6.33 | 6.66 |
| CaO       | 9.72 | 9.41 | 9.94 | 9.84 | 10.03 | 9.84 | 9.76 | 9.79 |
| Na$_2$O   | 2.39 | 2.13 | 2.19 | 2.17 | 2.21 | 2.34 | 2.58 | 2.29 |
| K$_2$O    | 0.78 | 0.73 | 0.81 | 0.8 | 0.79 | 0.78 | 0.78 | 0.78 |
| P$_2$O$_5$| 0.26 | 0.21 | 0.21 | 0.21 | 0.22 | 0.24 | 0.3 | 0.24 |
| LOI       | 0.50 | 0.61 | 0.31 | 0.25 | 0.21 | 0.85 | 0.66 | 0.48 |
| **Total** | 100.97 | 100.07 | 100.66 | 100.54 | 100.56 | 99.92 | 100.24 | 100.42 |
| **Weathering indices** |     |     |     |     |     |     |     |     |
| WIP       | 71.5 | 69.3 | 71.3 | 70.1 | 70.6 | 72.9 | 72.4 | 71.16 |
| CIA       | 38.4 | 41.2 | 39.3 | 39.8 | 39.8 | 36.9 | 40.4 | 39.40 |
| CIW       | 38.7 | 41.3 | 39.9 | 40.2 | 40.2 | 37.2 | 40.8 | 39.76 |
| MIA(O)    | 37.4 | 38.0 | 37.0 | 37.1 | 37.5 | 36.0 | 38.3 | 37.33 |
| **Shell around core-stone** |     |     |     |     |     |     |     |     |
| SiO$_2$   | 50.68 | 52.21 | 52.68 | 48.73 | 48.90 | 50.70 | 50.11 | 50.57 |
| TiO$_2$   | 2.00 | 2.00 | 2.10 | 2.20 | 2.00 | 2.20 | 2.80 | 2.19 |
| Al$_2$O$_3$ | 14.70 | 16.32 | 16.57 | 16.44 | 16.70 | 15.90 | 17.15 | 16.25 |
| Fe$_2$O$_3$ | 15.00 | 12.08 | 12.18 | 14.88 | 14.50 | 14.62 | 15.58 | 14.12 |
| MnO       | 0.14 | 0.15 | 0.15 | 0.14 | 0.13 | 0.13 | 0.13 | 0.14 |
| MgO       | 5.80 | 4.20 | 4.29 | 5.96 | 4.62 | 4.19 | 4.38 | 4.78 |
| CaO       | 8.10 | 8.00 | 7.16 | 7.80 | 8.03 | 7.81 | 7.50 | 7.77 |
| Na$_2$O   | 2.00 | 2.00 | 2.00 | 2.01 | 2.00 | 1.90 | 1.99 | 1.99 |
| K$_2$O    | 0.72 | 0.70 | 0.65 | 0.70 | 0.70 | 0.72 | 0.72 | 0.69 |
| P$_2$O$_5$| 0.24 | 0.41 | 0.30 | 0.22 | 0.20 | 0.23 | 0.25 | 0.26 |
| LOI       | 0.80 | 1.20 | 1.50 | 1.20 | 1.90 | 1.50 | 0.38 | 1.21 |
| **Total** | 100.16 | 99.27 | 99.76 | 100.73 | 100.72 | 99.87 | 100.99 | 100.21 |
| **Weathering indices** |     |     |     |     |     |     |     |     |
| WIP       | 62.7 | 58.0 | 50.9 | 61.0 | 59.2 | 55.8 | 55.3 | 57.56 |
| CIA       | 44.7 | 48.1 | 50.4 | 48.2 | 47.8 | 47.6 | 50.1 | 48.13 |
| CIW       | 44.9 | 47.8 | 50.4 | 48.5 | 48.3 | 47.9 | 50.9 | 48.39 |
| MIA(O)    | 41.2 | 46.0 | 47.3 | 44.2 | 46.4 | 47.3 | 49.0 | 45.91 |

* See text

WIP = \( \left( 2 \text{Na}_2\text{O}/0.35 \right) + \left( \text{MgO}/0.9 \right) + \left( 2\text{K}_2\text{O}/0.25 \right) + \left( \text{CaO}/0.7 \right) \times 100 \) (Parker 1970).

CIA = \( \left[ \text{Al}_2\text{O}_3/ \left( \text{Al}_2\text{O}_3 + \text{CaO} \times + \text{Na}_2\text{O} + \text{K}_2\text{O} \right) \right] \times 100 \) (Nesbitt and Young 1982).

CIW = \( \left[ \text{Al}_2\text{O}_3/ \left( \text{Al}_2\text{O}_3 + \text{CaO} \times + \text{Na}_2\text{O} \right) \right] \times 100 \) (Harnois 1988).

MIA(O) = \( \left( \text{Al}_2\text{O}_3 + \text{Fe}_2\text{O}_3T \right)/ \left( \text{Al}_2\text{O}_3 + \text{Fe}_2\text{O}_3T + \text{MgO} + \text{CaO} + \text{Na}_2\text{O} + \text{K}_2\text{O} \right) \times 100 \) (Babechuck et al. 2014).
shell, indicating higher weathering than in other samples, whereas sample No. 4 shows the least difference, indicating minimum weathering. On average, the WIP decreases from 71 in the core-stone to 58 in the weathered shell.

The CIA, monitors the progressive alteration of feldspars in the samples. The high CIA values reflect the removal of mobile cations (Ca, Na, K) relative to Al during weathering (Nesbitt and Young 1982). Table 1 shows that the CIA values of the core-stones of the basalts range from 37 to 41. The weathered shells have higher CIA values than their corresponding core-stones, ranging from 45 to 50. On average the CIA increases from 39 in the core-stone to 48 in the weathered shell. This slight increase indicates the low degree of weathering characteristic of incipient weathering. In this case, the feldspars in the core-stones were not excessively altered. As a matter of fact, the X-ray

Table 2 Trace elements (ppm)

|     | 1  | 2  | 3  | 4  | 5  | 6  | 7  | Av  |
|-----|----|----|----|----|----|----|----|-----|
| **Core-stone** |    |    |    |    |    |    |    |     |
| V   | 300| 280| 290| 300| 310| 290| 300| 295.7|
| Cr  | 85 | 88 | 90 | 88 | 85 | 90 | 85 | 87.3 |
| Co  | 55 | 52 | 53 | 55 | 54 | 55 | 54 | 54.0 |
| Ni  | 44 | 43 | 45 | 46 | 44 | 45 | 45 | 44.6 |
| Zn  | 75 | 80 | 77 | 75 | 80 | 78 | 80 | 77.9 |
| Rb  | 20 | 21 | 21 | 20 | 20 | 21 | 22 | 20.7 |
| Sr  | 360| 365| 370| 350| 365| 360| 385| 365.0|
| Y   | 35 | 35 | 36 | 35 | 36 | 36 | 35 | 35.4 |
| Zr  | 220| 219| 220| 219| 220| 220| 230| 221.1|
| Ba  | 225| 230| 230| 225| 220| 230| 222| 226.0|
| La  | 22 | 23 | 22 | 23 | 21 | 22 | 22 | 22.1 |
| Ce  | 50 | 51 | 50 | 51 | 52 | 50 | 51 | 50.7 |
| Pb  | 3  | 3  | 3  | 2  | 3  | 3  | 3  | 2.9  |
| Th  | 3  | 3  | 3  | 2  | 3  | 2  | 2  | 2.6  |

| **Shell around core-stone** |    |    |    |    |    |    |    |     |
| V   | 265| 248| 257| 265| 275| 257| 265| 261.7|
| Cr  | 79 | 82 | 74 | 82 | 79 | 78 | 85 | 79.9 |
| Co  | 55 | 52 | 53 | 55 | 54 | 55 | 54 | 54.0 |
| Ni  | 36 | 35 | 39 | 38 | 36 | 37 | 46 | 38.1 |
| Zn  | 58 | 73 | 70 | 68 | 73 | 71 | 81 | 72.0 |
| Rb  | 10 | 11 | 10 | 10 | 11 | 11 | 11 | 104  |
| Sr  | 334| 296| 301| 294| 286| 295| 295| 300.1|
| Y   | 35 | 35 | 36 | 35 | 36 | 35 | 35 | 35.4 |
| Zr  | 222| 221| 222| 221| 222| 222| 232| 222.9|
| Ba  | 251| 253| 254| 251| 249| 245| 250| 250.4|
| La  | 22 | 23 | 22 | 23 | 21 | 22 | 22 | 22.1 |
| Ce  | 51 | 52 | 51 | 52 | 53 | 51 | 52 | 51.2 |
| Pb  | 3  | 3  | 3  | 2  | 3  | 3  | 3  | 2.9  |
| Th  | 3  | 3  | 3  | 2  | 3  | 2  | 2  | 2.6  |

Fig. 3 Total alkali-silica diagram; the solid line is the boundary line of Macdonald and Katsura; the core-stones (black squares) and their shells (triangles) lie below the solid line in the field of subalkaline (tholeiitic) basalts.
diffraction analysis of the weathered shells indicated the presence of peaks of feldspars.

Harnois (1988) pointed out that potassium cations leached during weathering can be fixed by adsorption on weathering products, especially clays, and this may disturb the geochemical trend of potassium. He, therefore, proposed the chemical index of weathering (CIW) as an improved index of the degree of weathering to WIP or CIA. The CIW index increases with the degree of depletion in Na and Ca relative to Al. Table I shows that CIW increased from the core-stones of the basalt spheroids (average CIW of 40) to the corresponding weathered shells (average CIW of 48).

Figures 4 and 5 show the relationship between WIP and CIA, and between CIW and CIA, respectively. Figure 4 shows the decrease in WIP and increase in CIA from the core-stone samples to the corresponding weathered shells. Figure 5 shows that there is a linear correlation between CIW and CIA, both indices increase from the core-stone samples to the corresponding weathered shells.

Babechuk et al (2014) proposed the mafic index of alteration (MIA) as a chemical weathering index that extends the equation of the CIA to include the mafic elements Mg and Fe. It is known that many of the mafic minerals (especially pyroxene and olivine) are susceptible to chemical weathering, resulting in the loss of Mg from the weathering products. The fate of Fe during the weathering of most mafic minerals is, however, redox-dependent. In reducing environments, ferrous iron can be mobile and is leached along with Mg. In oxidative weathering environments, however, Fe is usually retained as insoluble ferric iron oxide or oxyhydroxide and thus enriched along with Al. Two equations have been suggested by Babechuk et al. (2014) for the calculation of MIA, one for oxidizing environment and the second for reducing environment. Table 1 gives the calculated values of MIA(O), as it is expected that an oxidizing environment had prevailed during the weathering of Gebel Qatran basalt MIA(O) increases from the core-stones of the basalt spheroids (average MIA of 37) to the corresponding weathered shells (average MIA of 46), indicating that iron from the mafic minerals in the core-stones was conservatively preserved as ferric oxide and/or oxyhydroxide, with progressive weathering.

Because of the complexity of geological systems and the weathering process, Nesbitt and Young (1982) and Nesbitt and Wilson (1992) used the A-CN-K ternary diagram to explain the trend of weathering, rather than depending on single indices like PIW, CIA or CIW. This diagram portrays the molar proportions of Al₂O₃ (A apex), CaO*+Na₂O (CN apex) and K₂O (K apex). Figure 6 shows the weathering trend from the core-stones of the basalt spheroids to the corresponding weathered shells. This trend is adjacent and parallel to the CN-A axis, as predicted by Nesbitt and Wilson (1992) for basalts. If weathering had progressed further than that, clay minerals should have been produced at the expense of feldspars and the bulk composition of weathered samples would have moved up the trend in the diagram towards the A apex. Since the weathered shells of the basalt spheroids plot at the lower part of the weathering trend (Fig. 6), this indicates that the weathering had been in the initial (incipient) stage.

The depletion of the various elements from the weathering regime is related to the nature of the host primary minerals, the mobility of these elements once they are released by the weathering of these minerals, and the susceptibility of the elements to retention in new phases in their immediate environment. To evaluate the chemical changes during weathering, it is necessary to assume that
one chemical component is stable and not removed during the weathering process. For this purpose (e.g., Nesbitt et al. 1980; Nesbitt and Wilson 1992; Kurtz et al. 2000; Patino et al. 2003; Ma et al. 2007), TiO$_2$ was chosen by several authors (see, e.g., Nesbitt and Wilson 1992).

In the present work, we studied the variability of the element concentrations in the shells of the basalt spheroids after normalizing them to TiO$_2$ concentrations and then compared them to the core-stones. The percentage change in the TiO$_2$–normalized concentration of an element in the shell to that in the core-stone was calculated according to the following (Nesbitt 1979; Nesbitt and Wilson 1992):

$$\% \text{ change to ratio} = 100 \times \left[ \frac{(R_{\text{sh}} - R_{\text{cs}})}{R_{\text{cs}}} \right]$$

where $R_{\text{sh}}$ and $R_{\text{cs}}$ are the elemental ratios in the shell and the core-stone, respectively. The assumption for this type of calculation is that there is no volume change between the weathering product and the un-weathered rock (Nahon and Merino 1996).

Figure 7 shows the average percentage change in the ratios of the major elements in the shell to core-stones of the basalt spheroids. The Mg/Ti, Ca/Ti, Na/Ti and K/Ti ratios decrease from the core-stone to the weathered shell. The decrease in Mg is attributed to the rapid weathering of olivine in the core-stone. Ca is associated primarily in plagioclase, clinopyroxene and glassy material, all of which weather less rapidly than olivine. Therefore, Ca/Ti decreases less rapidly than Mg/Ti in incipient weathering. Na and K reside in plagioclase feldspars which are less readily weathered than olivine and clinopyroxenes. In contrast to Mg, Ca, Na and K, Al and Si are more conservative in incipient weathering and the Al/Ti, Si/Ti ratios show slight or no increase during weathering. Fe/Ti ratio shows an increase in early weathering, but rapid decrease during more advances weathering (Nesbitt and Wilson 1992). Mn/Ti ratio decreases with increased weathering, and P/Ti ratio increases in early weathering but decreases in advanced weathering.

Figure 8 shows the average percentage change in the ratios of trace elements in the shell to core-stones of the basalt spheroids. V/Ti, Cr/Ti, Ni/Ti, Zn/Ti, Rb/Ti and Sr/Ti ratios show varying rates of decrease during weathering. V and Cr reside in clinopyroxenes, glass and oxides, and V/Ti and Cr/Ti show slight decrease in incipient weathering. Ni is primarily located in olivine and Zn in clinopyroxene and glass; both Ni/Ti and Zn/
Ti ratios decrease in early weathering. Sr resides primarily in plagioclase, and like Na, the Sr/Ti ratio decreases with increased weathering. Rb resides also in plagioclase and in glass and Rb/Ti ratio decreases like K/Ti ratio with weathering. On the other hand, the remaining trace elements (Co, Y, Zr, La, Ce, Pb and Th) are more conservative and their ratios show no change or slight increase. Ba released during weathering may be rapidly precipitated by sorption to secondary minerals, and hence the high Ba/Ti ratio (Nesbitt and Wilson 1992).

Discussion

The petrographical, mineralogical and chemical characteristics of the core-stones and the weathered shells of the basalt spheroids indicate that the weathering of Gebel Qatrani basalt is of the incipient stage. The chemical weathering indices, WIP, CIA and CIW, indicate slight weathering and rather low rates of release of mobile elements. Such weathering of the basalt must have taken place under semiarid to arid conditions. This interpretation is, more or less, in agreement with that reached Frey et al. (2013) in their study of the weathering of basalt in the semiarid Deschutes Basin of central Oregon. Novikov et al. (1993) studied the basalt crusts from Syria and pointed out that they reflect significant climate fluctuations in the region, from semiarid (Miocene basalt) to humid warm (Pliocene basalt) and finally to arid conditions (Quaternary basalt).

In Fayum area Bown et al. (1982) and Bown and Kraus (1988) pointed out that in Oligocene times, the Fayum area was subtropical to tropical lowland coastal plain with damp soils and seasonal rainfall that supported an abundance of and variety of vegetation, and large and varied vertebrate fauna. In post-Oligocene times, after the eruption of Gebel Qatrani basalt, the climatic conditions must have changed. El-Saadawi et al. (2014) studied the silicified fossil wood found in the Early Miocene Khashab Formation and found that the woods are characterized by having few wide vessels which are indicative of warm humid climate. However, they pointed out that the mode of occurrence of the fossil trunks and the absence of other plant remains such as twigs and roots indicates that fossil woods were not preserved in situ, but transported from where they grew before silicification. Consequently, the anatomical features of the fossil woods cannot be taken as an indication of the climate at the locality of Gebel Khashab. More recently, Zhang et al. (2014) studied the aridification of the Sahara desert in North Africa using different models and found out that North Africa experienced pronounced aridification from the Early Miocene to the Late Miocene. They indicated that in the Late Oligocene and Early Miocene, North Africa was dominated by semiarid steppe climate with restricted areas of arid desert climate. In the Late Miocene, the arid desert climate expanded across much of North Africa with a greater resemblance to today’s conditions. Zhang et al. (2014) pointed out that aridification was due to a reduction in North African precipitation.

If subtropical and tropical humid environment had prevailed in Gebel Qatrani area after the eruption of the basalt, this would have led to the formation of weathering soil profiles on the basalt substrate. No such profiles have been encountered on top of the Gebel Qatrani basalt. Therefore, it can be concluded that the incipient weathering of Gebel Qatrani basalt must have taken place under semiarid to arid conditions.
Conclusions

The present study on the spheroidal weathering of basalt from Gebel Qatrani, Fayum Depression, Egypt, indicates that weathering was at the initial (incipient) stage. The indices of chemical weathering (WIP, CIA, CIW and MIA(o)) and element mobility studies indicate that there was a loss of Mg, Ca, Na and K during the weathering of the core-stones of the basalt spheroids to the corresponding weathered outer shells. Fe mainly resided with Al, indicating that weathering was under oxidizing conditions. There was also a depletion of V, Cr, Ni, Zn, Rb and Sr due to the weathering of the core-stone. On the other hand, Co, Y, Zr, La, Ce, Pb and Th behaved essentially conservatively. The low degree of weathering of the basalt spheroids indicates that weathering took place mainly in arid to semiarid environment.

Authors' contributions

EE-H was responsible for field work and assistance in interpretation of results. SDA was responsible for analytical work and Khalil was responsible for petrography and mineralogy. All three authors were responsible for the preparation of the final manuscript submitted for publication. All authors read and approved the final manuscript.

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There are no competing interests in carrying out the present research.

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