MINERALOGY, RADIOACTIVITY AND REE-GEOCHEMISTRY OF BATOGA MONZOGRA NITE AND ASSOCIATED PEGMATITE, SOUTHEASTERN DESERT, EGYPT

ABDEL MOEZ, A. SADEK and ABDEL HADI A. ABBAS

Nuclear Materials Authority, P.O. Box 530, El-Maadi, Cairo, Egypt

ABSTRACT

The studied area lies in the southern part of the Eastern Desert of Egypt to the southwest of Ras Benas. This area is covered by metagabbros, tonalite and monzogranite. The pegmatites occur as elongated, small pockets and lenses in the monzogranite and display sharp contacts with the host granite. In all cases, the pegmatites are characterized by obvious simple zonation.

The radiometric studies of monzogranite and associated pegmatite samples show that these rocks are uraniferous, where their average equivalent uranium contents are 12 ppm and 134 ppm, respectively. The uraniferous character was confirmed by the average eTh/eU ratios (1.13 and 0.96, respectively). The average ratios of $P_{\text{factor}}$ of the studied granites and associated pegmatites are more than one (1.89 and 16.35, respectively), indicating disequilibrium in U-decay series due to addition of uranium to these rocks. The present study proved that the presence of uranothorite, uranophane and the uranium-bearing minerals such as zircon, monazite, allanite and fluorite are responsible for both U and Th contents in the studied granite and associated pegmatite.

The normalized REE pattern of the monzogranite and associated pegmatite have similar geochemical characteristics. Where the REE patterns show a LREE relatively enriched feature with strong negative Eu-anomaly. The characteristic high LREE relative to HREE of the analyzed samples is attributed to the presence of the allanite, monazite and apatite minerals. The slightly positive Tb anomaly in the studied granites attributed to its preferential accommodation within the fluorite structure.

INTRODUCTION

The younger granites are widely distributed all over the Egyptian Shield, constituting approximately 30% of the plutonic assemblage. Their abundance relative to the older granitoids increases from 1 to 4 in the south to approximately 1 to 1 in the north of the Eastern Desert (Stern 1979). The granitic rocks are known to have much higher uranium content than the other common rock types. Some younger granites in Egypt host potential uranium mineralization. Such uranium mineralization seems to have an intimate relationship with the hydrothermal alteration (Mohamed, 1995, Shalaby, 1996, El Bayoumi et al., 1999, Assaf et al., 1999, Ammar and Abd El Wahed, 2000 and Dawood and Abd El Naby, 2001, Sadek, 2005).

The studied area lies in the southern part of the Eastern Desert of Egypt to the southwest of Ras Benas. It is located between longitudes 35° 20’ and 35° 25’ E, and latitudes 23° 47’ and 23° 52’ N covering an area of about 80 km². The area of study is covered by metagabbros, tonalite and monzogranites (Fig.1).

The present study deals with the geology,
petrography, radioactivity, mineralogy and rare earth geochemistry of the granitic rocks of Gabal Batoga in order to discuss the genesis of the radioactive anomalies.

FIELD AND PETROGRAPHIC OBSERVATION

The study area is covered by metagabbros, tonalite and younger granites.

The Tonalite show low resistance to weathering. They are characterized by cavernous, exfoliation and boudary weathering and monumental like -shape. Also in places, some irregular shaped enclosed of microgranular mafic rocks enclosed in these rocks.

Monzogranite is marked by relatively higher mountain, with prominent peak up to 801 m above sea level. The rocks are equigranular, medium-grained with pale buff to white colors, and have rounded pluton and show intrusive contacts to the older rocks. Generally, the monzogranite intrudes the older rocks by sending offshoots invading them. Sometimes, it carries the tonalite as roof pendants (Fig.2).

The studied monzogranite is highly weathered, showing exfoliation bouldary with characteristic monumental shapes. The pluton of younger granite is traversed by many dyke swarms of mafic (andesite) and felsic (aplite and microgranite) composition, trending mainly in the NW-SE and NE-SW directions (Fig.3).

Pegmatites are commonly encountered in the studied monzogranite. They are found as small lenticular and oval-shaped pockets of limited extension near the intersections of the NE-SW and NW-SE faulting systems (Figs4&5). Their dimensions are about 2 to 5m in width and 10 to 300 m in length. They are very coarse-grained rocks of pink color and are composed mainly of megacrysts of potash feldspar crystals (70%), milky quartz (20%), and plagioclase (8%), together with subordinate mafic minerals mainly biotite and muscovite. Radiometric field measurements indicated that the radioactivity of pegmatites is more than twice that of their enclosing granitic rocks.

Fig.1: Geologic map of Batoga Area, Central Eastern Desert, Egypt
rocks. They are characterized by high contents of radiometric measurements with visible mineralizations mainly of metallic black and brown color minerals, as well as yellow colored uranium minerals and disseminated green and violet fluorite. The studied area are traversed by many faults in different directions and types. These are either concomitant with wadis and drainage lines or cutting the country rocks and having NE-SW (older) and NW-SE (younger) directions.

**PETROGRAPHY**

The microscopic investigations of the studied monzogranites indicate they are mainly composed of quartz (~30%), plagioclase (~32%), K-feldspars (~26%), biotite, hornblende and muscovite as mafic minerals (~10%). The common accessory minerals are zircon, allanite, fluorite, monazite, apatite and opaques.

Quartz, commonly occurs as anhedral crystals of variable sizes and show undulose extinction. Minute quartz crystals occupy the interstitial spaces between feldspars. Sometimes, cataclastic deformation, granulation, recrystallization are observed in many crystals. The presence of myrmekitic and graphic texture (Fig.6) is a reflection of mineral replacement and reaction features affecting of this monzogranite (Shelley, 1993).

*K-feldspar* occurs as subhedral to anhedral crystals of microcline, microcline perthite and orthoclase perthites as well as antiperthite. Orthoclase perthite occurs as patchy and flame
types. Microcline perthite has both albite and periclase twinning which together form what is known tartan pattern (Fig.7).

Plagioclase occurs as oligoclase which is the most abundant essential mineral. It is represented by subhedral to euhedral prismatic crystals of variable sizes with distinct carlsbad and albite twinning and sometimes occurs as zoned crystals (Fig.8). It is suassuritized, and muscovitized along core and some crystal altered along peripheries are observed (Fig.9).

Biotite and muscovite are the most abundant mafic minerals. Biotite occurs as subhedral flakes of brown color. Some biotite crystals poikilitically enclose zircon that is surrounded by the pleochroic haloes due to radiogenic effects (Fig.10). In some places biotite altered to chlorite. Muscovite is subordinate mineral and found as minute amount as a primary mineral associated with opaques and also as a second-ary mineral resulting from the alteration of plagioclase and biotite. Hornblende occurs as greenish subhedral crystals, showing simple twinning, partially altered to chlorite and also corroded by quartz and feldspar (Fig.11).

Zircon occurs as fine prismatic metamict crystals intimately associated with biotite and feldspars (see Fig.10). Apatite occurs as fine rod–like and/or prismatic crystals (Fig.12). Allanite occurs as euhedral crystals, yellowish brown masked interference colors (Fig.13). The petrographic investigation indicates
that the monzogranite of Batoga granites belong to the subsolvus type (Deer et al., 1962), which was crystallized under relatively high water pressure conditions.

**Pegmatite** is mainly composed of K-feldspar, plagioclase, and quartz in addition to muscovite and biotite. The accessory minerals are mainly zircon, monazite, allanite, and fluorite. K-Feldspars occur as orthoclase, microcline and microcline perthite. Perthite and antiperthite form subhedral to euhedral prismatic megacrysts corroded by quartz (Fig.14). Plagioclase (An_{20}) exhibits slight deformation, suassuratization and muscovitization in some crystals (Fig.15). Quartz is cracked, in places filled with iron oxides and yellowish color radioactive minerals (uranophane?) (Fig.15). Biotite occurs as euhedral flaky crystals of brownish green color (Fig. 16). The muscovite flakes exhibit different interference color (Fig. 17).

**RADIOACTIVITY**

Uranium occurs in the nature in several valance states, in which tetravalent (U^{4+}) and hexavalent (U^{6+}) are the most common ones, whereas thorium exists only in the tetravalent (Th^{4+}) state. The U^{4+} has an ionic radius close to the ionic radii of Th, Ca, Na, Zr, Nb, Y, Ce, Pb^{2+} and the rare earth elements. Therefore, U^{4+} can substitute these elements in the
U$^{+4}$ is insoluble in water and occurs mainly in primary minerals, but when oxidized to U$^{+6}$ becomes soluble in water and forms secondary minerals. Thorium is insoluble in water, occurs mainly in primary minerals, and is rarely leached by secondary processes.

Field measurements show a slight increase in radioactivity of the examined Batoga monzogranite and associated pegmatite. Uranium, thorium, radium and potassium were measured by using four energy regions representing $^{238}$Th, $^{212}$Pb, $^{214}$Pb and $^{40}$K at 93 kV, 239 kV, 352 kV, and 1460 kV for uranium, thorium, radium and potassium, respectively. The measurements were carried out in sample plastic containers, cylindrical in shape, 12.6 cm$^3$ in volume with 9.5cm average diameter and 3cm height. The rock sample was crushed as fine as about 1 mm in grain size, and then the container was filled with about 300–400 gm of the crushed sample sealed well and left for at least 21 days to accumulate free radon to attain radioactive equilibrium. The relation between the percentage of $^{222}$Rn accumulation and time increased till reaching the steady stage after about 38 days (Matolin, 1991).

Eighteen samples (8 samples from monzogranite and 10 samples from associated pegmatite) were radiometrically analyzed.
to determine the equivalent uranium (eU) equivalent thorium (eTh), radium and potassium contents (Table 1). The eU contents of the analyzed monzogranites range from 8 ppm to 20 ppm with an average 12 and the eTh contents range from 10 ppm to 23 ppm with an average 13.3. Mean value of uranium contents in the studied granites is slightly higher than the average of granitic rocks (4.75 ppm) given by Rogers and Adams (1969). eTh/eU ratios of these granites range from 0.93 to 1.25 with an average 1.13.

The eU contents of the analyzed pegmatite samples range from 28 ppm to 355 ppm with an average 134 and the eTh contents range from 24 ppm to 210 ppm with an average 97.6 as given in Table 1. It is worthy to mention that the altered granites belong to high-thorium, high-uranium granitic rocks where uranium is mostly located in the Th-rich accessory minerals as monazite, zircon and allanite. The eTh/eU ratios of these samples range from 0.52 to 1.78 ppm with an average 0.96. The very low eTh/eU < 1 in pegmatites indicate strong addition of uranium during post-magmatic stages (Charbonneau, 1982).

Table 1: eU and eTh contents in (ppm) of the studied granite and pegmatite

<table>
<thead>
<tr>
<th>R.T</th>
<th>S.no.</th>
<th>eU(ppm)</th>
<th>eTh(ppm)</th>
<th>eTh/eU</th>
<th>eU/Ra</th>
<th>K%</th>
<th>Ra(eU)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Monzogranite</td>
<td>1</td>
<td>14</td>
<td>13</td>
<td>0.93</td>
<td>2.0</td>
<td>3.5</td>
<td>7.0</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>10</td>
<td>12</td>
<td>1.2</td>
<td>1.37</td>
<td>3.0</td>
<td>7.3</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>8.0</td>
<td>10</td>
<td>1.25</td>
<td>1.60</td>
<td>3.2</td>
<td>5.0</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>9.7</td>
<td>11</td>
<td>1.13</td>
<td>1.76</td>
<td>3.4</td>
<td>5.5</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>11</td>
<td>13</td>
<td>1.18</td>
<td>1.77</td>
<td>3.3</td>
<td>6.2</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>20</td>
<td>23</td>
<td>1.1</td>
<td>2.94</td>
<td>4.0</td>
<td>6.8</td>
</tr>
<tr>
<td></td>
<td>7</td>
<td>9.0</td>
<td>10</td>
<td>1.11</td>
<td>1.55</td>
<td>3.0</td>
<td>5.8</td>
</tr>
<tr>
<td></td>
<td>8</td>
<td>13</td>
<td>14.5</td>
<td>1.12</td>
<td>2.10</td>
<td>3.6</td>
<td>6.2</td>
</tr>
<tr>
<td>Av.</td>
<td>11.8</td>
<td>13.3</td>
<td>1.13</td>
<td>1.89</td>
<td>3.38</td>
<td>6.23</td>
<td></td>
</tr>
<tr>
<td>Pegmatite</td>
<td>1</td>
<td>35</td>
<td>28</td>
<td>0.8</td>
<td>4.86</td>
<td>4.0</td>
<td>7.2</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>28</td>
<td>36</td>
<td>1.29</td>
<td>4.00</td>
<td>4.2</td>
<td>7.0</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>18</td>
<td>24</td>
<td>1.33</td>
<td>2.77</td>
<td>3.8</td>
<td>6.5</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>54</td>
<td>56</td>
<td>1.04</td>
<td>7.20</td>
<td>3.6</td>
<td>7.5</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>63</td>
<td>112</td>
<td>1.78</td>
<td>8.29</td>
<td>4.0</td>
<td>7.6</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>98</td>
<td>80</td>
<td>0.82</td>
<td>12.25</td>
<td>3.6</td>
<td>8.0</td>
</tr>
<tr>
<td></td>
<td>7</td>
<td>345</td>
<td>180</td>
<td>0.52</td>
<td>40.59</td>
<td>3.8</td>
<td>8.5</td>
</tr>
<tr>
<td></td>
<td>8</td>
<td>225</td>
<td>169</td>
<td>0.75</td>
<td>27.44</td>
<td>3.0</td>
<td>8.2</td>
</tr>
<tr>
<td></td>
<td>9</td>
<td>355</td>
<td>210</td>
<td>0.59</td>
<td>40.80</td>
<td>3.4</td>
<td>8.7</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>122</td>
<td>80.5</td>
<td>0.66</td>
<td>15.25</td>
<td>3.1</td>
<td>8.0</td>
</tr>
<tr>
<td>Av.</td>
<td>134</td>
<td>97.6</td>
<td>0.96</td>
<td>16.35</td>
<td>3.65</td>
<td>7.72</td>
<td></td>
</tr>
</tbody>
</table>

The U-Th diagram of the studied granites and pegmatites shows a positive correlation which indicates that the behavior of U and Th was controlled by post-magmatic processes (Fig.18) (Simpson et al., 1979; Raslan et al., 2010a and b). The variation of Th and U against Th/U ratio shows a negative correlation with U and Th, which has confirmed the idea that U and Th enrichment in these anomalies was controlled by magmatic processes or deuteric ones (Fig.19 and 20) (Simpson Op.Cit). It is important to observe a slight increase in uranium concentration with increasing potassium concentration in monzogranites on the contrary to pegmatites (Fig.21), which is mostly due to the mobility of uranium as reported by Anjos et al. (2005). Thorium displays positive correlation with potassium in the studied monzogranites suggesting magmatic controlling processes and negative correlation with associated pegmatites probably due to the effect of hydrothermal alteration processes which led to the formation of clay minerals at the expense of feldspars (El-Feky et al., 2011). In general, low eTh/K ratios are an excellent indicator of potassium alterations (Shives et al., 1995). Ratios of eTh/eU <2 are highly suggestive of relative uranium enrichment but ratios >7 indicate preferential removal of uranium, possibly by leaching (Adams and Weaver, 1958). The cross plot of eTh/eU and eTh/K ratios presented in (Fig.23). exhibit that granite samples lie in the fixed-U sector while pegmatite samples lie in the leached-U sector with the exception of one sample (Anjos et al., 2005).

Radioactive Equilibrium

According to Reeves and Brooks (1978), uranium (U\textsuperscript{238}) series attains the equilibrium state in nearly 1.5 M.a. Cathelineau and Holliger (1987) stated that uranium mineralization is affected by different processes. Leaching, mobility and redistribution of uranium are affected by hydrothermal solutions and/or supergene fluids, which cause disequilibrium in the radioactive decay series in U- bearing rocks. The radioactive equilibrium of the
Fig. 18: Variation diagram between $e^{\text{Th}}$ vs. $e^{\text{U}}$ of the studied monzogranite (♦) and associated pegmatite (■) in Batoga area.

Fig. 19: Variation diagram between $e^{\text{U}}$ vs. $e^{\text{Th}}/u$ of the studied monzogranite (♦) and associated pegmatite (■) in Batoga area.

Fig. 20: Variation diagram between $e^{\text{Th}}$ vs $e^{\text{Th}}/u$ of the studied monzogranite (♦) and associated pegmatite (■) in Batoga area.

Fig. 21: Variation diagram between $e^{\text{U}}$ vs K% of the studied monzogranite (♦) and associated pegmatite (■) in Batoga area.

Fig. 22: Variation diagram between $e^{\text{Th}}$ vs K% of the studied monzogranite (♦) and associated pegmatite (■) in Batoga area.

Fig. 23: Variation diagram between $e^{\text{Th}}/K$ vs. $e^{\text{Th}}/u$ of the studied monzogranite (♦) and associated pegmatite (■) in Batoga area.
studied granitic rocks can be determined by the calculation of equilibrium factor (P) which is the ratio of radiometric uranium contents (eU) to the radium content Ra(eU); \( P_{\text{factor}} = \frac{eU}{Ra(eU)} \) (Hussein, 1978; El-Galy, 1998; Surour et al., 2001).

The average ratios of \( P_{\text{factor}} \) of the studied granites and associated pegmatites are 1.89 and 16.35, respectively. These ratios are more than one (Table 1) indicating disequilibrium in U-decay series due to uranium addition.

MINERALOGY

The heavy minerals were separated from the monzogranite and associated pegmatite by heavy liquid separation and picked as individual minerals under the binocular microscope. These minerals were identified by Environmental Scanning Electron Microscope (ESEM) back-scattered images, EDAX analyses and confirmed by X-ray diffraction (XRD) techniques. The EDAX examination revealed the presence of uranothorite, uranophane, Nb-Ta minerals (columbite and exunite) and REE-bearing minerals (allanite, monazite, apatite), zircon and fluorite.

**Uranothorite** (U,\( \text{Th} \))\( \text{SiO}_4 \)

It occurs as fine subhedral to anhedral bright inclusions. The ESEM analysis indicates that it consists essentially of \( \text{ThO}_2 \), \( \text{SiO}_2 \) and significant amount of \( \text{UO}_2 \), reaching up to 7%. Heinrich (1958) stated that, uranium is usually present in amount up to 10% in uranothorite. Other elements present in small to minor amounts such as Ca, Al, Zr and Fe Mg (Fig. 24).

**Uranophane** \( \text{Ca(UO}_2)_2\text{Si}_2\text{O}_7 \)

Uranophane polymorphs of calcium uranium hydrate silicates; they are light yellow to orange in fibrous crystals with vitreous luster. It occurs as fissure filling or coating the altered K-feldspars. Fairly common in oxidation zones of most uranium deposits (Cesborn et al., 1993).
The ESEM-EDAX semiquantitative analysis of uranophane was composed of Ca, U and Si confirmed by X-ray diffraction (Fig.25&26) respectively. The uranium may be derived from altered primary radioactive minerals, while calcium and silica are available in the aqueous solutions penetrating shear zone. Calcium can be liberated from altered plagioclase, calcite and silica which liberated during kaolinite initiation of feldspars.

\[
\text{Ca}^{2+} + 2\text{UO}_2^{2+} + 2\text{SiO}_2 + 6\text{H}_2\text{O} + 3\text{O}^{2-} \leftrightarrow \text{Ca(UO}_2\text{)}_2\text{Si}_2\text{O}_7 \cdot 6\text{H}_2\text{O} 
\]

In the study area, the absence of any trace of primary U-minerals and the abundance of the metamictized accessory minerals reflect that the metamictization is the main controlling process in the accumulation of the U which may be leached and redistributed along the fractures. Also, the leaching of the U-bearing minerals from the host granites is controlled by hydrothermal system operating in response to regional tectonism as well as the interaction with hydrothermal fluids (Cuneey 1978; Abdel-Monem et al. 1998 and El-Kammar et al. 2001) which become enriched in U.

**Rare Earth-Bearing Minerals**

**Columbite** (Fe,Mn)(Nb,Ta)\(_2\)O\(_6\)

Ferrocolumbite grains were detected in the studied sample of pegmatite. The grains are generally black in color and possess metallic luster under binocular microscope. In the studied area, columbite found as blackish color; composed mainly of Fe-columbite (Ta<Nb, Mn<Fe) as revealed from the ESEM analysis. Several columbite crystals have been subjected to semiquantitative analyses using EDAX-scanning electron microscope. The EDAX-ESEM analyses show that both Nb and Fe are the essential components together with subordinate amount of Ta, U, Th and Mn (Fig.27).

**Euxenite** (Y,Ca,Ce,U,Th)(Ti,Ta,Nb)\(_2\)O\(_6\)

It is the predominant REE-bearing mineral in the pegmatite body occurring as disseminated stout prismatic crystals and sometimes occurs as flattened black crystals commonly metamict. Chemically, euxenite is considered as oxide or titanite-columbite of the type AB\(_2\)O\(_6\). In the EDAX analyzed samples, ΣNb+Ta always exceeds Ti (Fig.28). In these minerals, U varies between 23.5 to 28.5% and Th from nil to 2.3, indicating the strong radioactive nature and their metamict state.

**Monazite** (Ce,La,Th,...)(Po\(_4\),SiO\(_4\))

It is a phosphate mineral, composed essentially of Ce, La and Th. Th is usually present in substitution for Ce and/or La where ThO\(_2\) is about 10 to 12 wt%. Y substitutes in small amount for Ce & La together with minor amounts of Ca and less abundantly Mg, Fe"′, Fe" due to oxidation. Al, Mn can be found...
and U is generally present in amount up to ~1 wt% $\text{U}_3\text{O}_8$. The Si has been found and ascribed to alteration or to admixed thorite but may be in part due to substitution of Si for P (Dana, 1963). Certain minerals such as monazite, bastnaesite and allanite are Ce- selective and prefer the LREE. Thus the Ce-group or LREE tend to be concentrated in highly fractionated acid rocks such as alkaline granites and pegmatites. In the studied granite, monazite occurs in minor amounts as minute crystals. The EDAX-ESEM analysis of monazite showed that U is low. The U may also replace some of the REE sites in the monazite (Hughes et al., 1991). The mineral composition contains a considerable amount of Si, La, Ce, Nd, Al, Ca, and Mn (Fig.29).

**Allanite** $(\text{Ce,Ca,Y})_2(\text{Al,Fe})_3(\text{SiO}_4)_3(\text{OH})$

It is a member of the epidote group minerals of both hydrothermal and magmatic origin with rare earth elements substituting Ca. It varies in color from light brown to black. Because of their similarity to the rare earth elements in ionic radius, U and Th may be incorporated in the structure of allanite. The metamictization of allanite produces an amorphous alteration product due to the destruction of the crystalline structure of allanite by the radioactive decay of its uranium and thorium contents. In the studied granites and pegmatites, allanite has been found as accessory mineral. They form euhedral to subhedral zoned tabular prismatic crystals (may attain 4-6 mm in length) of reddish brown, intimately associated with biotite and zircon. Scan Electron Microscope (EDAX) show that the allanite is composed essentially of LREE, Ca, Fe, K, Al and Si, as well as some U and Th content (Fig.30). The radioactivity of allanite in the studied granites and pegmatites is much less important than that of zircon.

**Zircon** $(\text{ZrSiO}_4)$

It occurs as euhedral prismatic grains characterized by pale to deep brown color and generally sub-translucent to opaque with dull luster. The EDAX data reflect the chemical composition of zircon is $\text{ZrO}_2$ and $\text{SiO}_2$ with significant amounts of Th, U, Fe, Al, Ca and Hf (Fig.31). The present study revealed that zircon and thorite can exist in partially solid solution state due to close up in the ionic radii of Zr and Th (Förster, 2006). Sometime, there is a clear intergrowth between zircon and thorite.

**Apatite** $\text{Ca}_5(\text{PO}_4)_3(\text{OH,F,Cl})$

It is a common accessory mineral in almost all igneous rocks. The common substitutions for Ca are Sr and rare earth elements and the lighter rare earth elements may be important in apatite of alkaline rocks. In the studied monzogranite and associated pegmatite, apa-
ABDEL MOEZ, A. SADEK and ABDEL HADI A. ABBAS

Fig. 31: EDX and back scattered images of the studied zircon

Fig. 32: EDX and back scattered images of the studied apatite

tite occurs as massive and pale green rode-like crystals. The EDAX analysis of the studied samples shows that apatite is composed essentially of Ca and P with minor amount of Si, Al, U and Fe. U substitutes for Ca in apatite structure (ionic radii are close to each other Ca=1.06) (Fig. 32).

Fluorite (CaF$_2$)

It is found as anhedral crystals vary from colorless, violet to blackish color. In both colorless and violet fluorite varieties more than 70% of the rare earths content in fluorite is of HREE. The ESEM analysis of fluorite reveals that it is composed of Ca, F, Si Al, and U (Fig. 33).

RARE EARTH ELEMENTS (REES)
GEOCHEMISTRY

The REE are the most useful of all trace elements in the interpretation of the igneous rocks petrogenesis. They include the elements with atomic numbers in between 57-71 (La-Lu) and tend to concentrate in the silicate melts. So, the REEs tend to concentrate in the residual magma and occur in the accessory minerals such as allanite, zircon, monazite, xenotime, apatite and sphene. The distribution of europium (Eu) is particularly important because it unlike the other REEs. It occurs in two oxidation states, Eu$^{2+}$ and Eu$^{3+}$ depending on oxygen fugacity. Under the oxidizing conditions, Eu is trivalent Eu$^{3+}$ and behaves as the other REEs, while under the reducing conditions, it occurs as divalent Eu$^{2+}$ and has larger ionic radius replacing mainly Ca$^{2+}$ in plagioclases and rarely K$^+$ in K-feldspars.

The REE normalized patterns of the igneous rocks are controlled by the REE chemistry of their sources and the crystal–melt equilibria that have taken place during their evolution. So, the shape of the normalized REEs pattern is mainly related to the rock forming minerals, as well as the processes that affected the rock formation, like fractional crystallization, partial melting and/or assimilation. REE distribution patterns are usually considered as an important tracer for source rocks because...
the REE concentrations have few changes and little deformation to lost during rock weathering, sedimentation and diagenetic processes (Byrne and Sholkovitz, 1996; Nozaki et al., 1997; Lawrence and Kamber, 2006; Frimmel, 2009).

Eleven samples are collected from the investigated granitic samples (seven samples from monzogranite and four samples from associated pegmatite) were analyzed for REEs in the Egyptian nuclear materials authority labs (NMA) using ICP-AES, Perkin Elmer, and Optima 3000 instrument. The operating conditions are: nebulizer flow (1L/min), auxiliary flow (1L/min), plasma flow (15L/min), viewing light (15mm) and RF power (1200 watts). The accuracy of the results was found to be in the range 5-10 % for the rare earth elements. The results of these analyses are listed in Table (2).

In order to evaluate the relative fractionation trend within the REE group, their contents in the studied granites are normalized to chondrite values of Anders and Grevesse (1989) and plotted against REE atomic numbers. The normalized REE pattern of monzogranite and pegmatite samples exhibit a remarkable similarity in REE contents; revealed by the superposition of different trends of both LREE and HREE (Fig.34 and 35). Monzogranite and pegmatite samples show similar chondrite-normalized REE patterns and have clearly fractionated LREE relative to HREEs and show distinct negative Eu anomaly.

Table 2: Rare earth elements contents (ppm) in the study Batoga monzogranite and associated pegmatite

<table>
<thead>
<tr>
<th>S. No.</th>
<th>S1</th>
<th>S2</th>
<th>S3</th>
<th>S4</th>
<th>S5</th>
<th>S6</th>
<th>Av. n=7</th>
<th>Pegmatite</th>
</tr>
</thead>
<tbody>
<tr>
<td>La</td>
<td>30.8</td>
<td>12.7</td>
<td>15.0</td>
<td>21.0</td>
<td>25.8</td>
<td>20.8</td>
<td>21.0</td>
<td>21.01</td>
</tr>
<tr>
<td>Ce</td>
<td>59.0</td>
<td>31.0</td>
<td>37.2</td>
<td>50.8</td>
<td>55.7</td>
<td>43.</td>
<td>58.8</td>
<td>47.93</td>
</tr>
<tr>
<td>Pr</td>
<td>7.4</td>
<td>7.8</td>
<td>6.5</td>
<td>7.0</td>
<td>7.3</td>
<td>6.8</td>
<td>8.7</td>
<td>7.36</td>
</tr>
<tr>
<td>Nd</td>
<td>116.5</td>
<td>63.0</td>
<td>60.8</td>
<td>76.3</td>
<td>94.7</td>
<td>85.3</td>
<td>73.4</td>
<td>81.43</td>
</tr>
<tr>
<td>Sm</td>
<td>13.4</td>
<td>9.4</td>
<td>9.8</td>
<td>9.4</td>
<td>11.4</td>
<td>11.4</td>
<td>13.4</td>
<td>11.17</td>
</tr>
<tr>
<td>Eu</td>
<td>0.7</td>
<td>0.3</td>
<td>0.5</td>
<td>0.6</td>
<td>0.8</td>
<td>0.9</td>
<td>0.6</td>
<td>0.63</td>
</tr>
<tr>
<td>Gd</td>
<td>6.8</td>
<td>5.6</td>
<td>4.4</td>
<td>6.7</td>
<td>8.6</td>
<td>9.0</td>
<td>6.5</td>
<td>6.80</td>
</tr>
<tr>
<td>Tb</td>
<td>1.9</td>
<td>1.74</td>
<td>1.8</td>
<td>1.5</td>
<td>1.7</td>
<td>1.8</td>
<td>1.35</td>
<td>1.68</td>
</tr>
<tr>
<td>Dy</td>
<td>9.6</td>
<td>6.5</td>
<td>6.2</td>
<td>8.7</td>
<td>11</td>
<td>11.8</td>
<td>7.6</td>
<td>8.77</td>
</tr>
<tr>
<td>Ho</td>
<td>2.8</td>
<td>1.4</td>
<td>1.4</td>
<td>1.8</td>
<td>2.2</td>
<td>2.5</td>
<td>1.6</td>
<td>1.96</td>
</tr>
<tr>
<td>Er</td>
<td>6.4</td>
<td>4.0</td>
<td>3.6</td>
<td>5.1</td>
<td>7.0</td>
<td>8.0</td>
<td>5.0</td>
<td>5.59</td>
</tr>
<tr>
<td>Tm</td>
<td>1.2</td>
<td>0.65</td>
<td>0.6</td>
<td>0.75</td>
<td>0.9</td>
<td>1.2</td>
<td>0.65</td>
<td>0.85</td>
</tr>
<tr>
<td>Yb</td>
<td>6.6</td>
<td>5.5</td>
<td>5.0</td>
<td>6.2</td>
<td>8.35</td>
<td>8.0</td>
<td>5.25</td>
<td>6.41</td>
</tr>
<tr>
<td>Lu</td>
<td>1.4</td>
<td>1.0</td>
<td>0.7</td>
<td>1.25</td>
<td>1.25</td>
<td>1.4</td>
<td>0.76</td>
<td>1.11</td>
</tr>
<tr>
<td>∑REE</td>
<td>264.5</td>
<td>147.99</td>
<td>153.5</td>
<td>197.1</td>
<td>236.7</td>
<td>211.9</td>
<td>204.61</td>
<td>202.3</td>
</tr>
<tr>
<td>∑LREE</td>
<td>227.8</td>
<td>124.2</td>
<td>129.8</td>
<td>165.1</td>
<td>195.7</td>
<td>168.2</td>
<td>175.9</td>
<td>169.5</td>
</tr>
<tr>
<td>∑HREE</td>
<td>36.7</td>
<td>23.7</td>
<td>32.0</td>
<td>41.0</td>
<td>43.7</td>
<td>28.71</td>
<td>32.80</td>
<td>24.8</td>
</tr>
</tbody>
</table>

*(La/Lu)*n = normalized values after Anders and Grevesse (1989)
The analyzed monzogranite shows REE fractionation where (La/Lu)_N vary from 1.29 to 2.8 with an average of 1.97 and 1.3 to 1.86 with an average of 1.63 for associated pegmatite samples (Table 2). They have relative enrichment of LREE contents in the analyzed monzogranite varying between ∑LREE 124.2 to 227.8ppm with an average 169.95. While the pegmatite, the LREE vary between ∑LREE 98.3 to 162.95ppm with an average 146.2; (the average of (La/Sm)_N range from 1.5 and 1.13 for these rocks, respectively (Table 2). The contribution of the REE bearing accessory minerals such as monazite, apatite, allanite and sphene represents potential contribution to the LREE concentration (Miller and Mittlefehldt 1982). This is confirmed by the petrographic study which shows that monzogranite contains sphene, allanite and apatite.

Concerning the HREE, The ∑HREE vary from 23.7 to 43.7 ppm with an average 32.8 for monzogranite and ∑HREE varying between 27.75 and 44.1ppm with an average of 34.4 for associated pegmatite. Where the average of (Gd/Lu)_N vary from 0.4 to 1.06 with an average of 0.74 for monzogranite and 0.67 to 1.15 with an average of 0.78 for the pegmatite samples (Table 2).

Strong negative Eu-anomaly which proposed by Henderson (1984) as (Eu/Eu*)_N/(Sm/(Gd)), when the value is >1.0 indicates positive anomaly while the value <1.0 is a negative anomaly. The rocks under consideration have (Eu/Eu*)_N varying between 0.13 and 0.26 with an average of 0.2 for the monzogranite. While the analyzed pegmatite samples vary from 0.046 to 0.21ppm (Table2).

In fact the negative Eu anomalies is related to Eu^{2+} which compatible with plagioclase. These are explained by the crystal fractionation of plagioclase from the felsic granitic melt at low pressures (Neiva 1992), or partitioning into feldspar during fractionation (Singh et al. 2006). Another alternative explanation for the negative Eu-anomaly is based on the high oxygen fugacity (fO_2) in the granitic melt due to volatile saturation (Grenne and Roberts 1998). The oxygen activity of melt would be sufficiently high to keep Eu at the trivalent state and thus detains its incorporation into the accumulating plagioclase.

The depletion of the total REEs contents in monzogranites and associated pegmatite are 202.3 and 180.6 ppm respectively which is less than the average of the world wide granites (270 ppm) as given by Hermann (1970). The depletion of REEs has been attributed to various processes including magmatic differentiation (Cuney and Friedrich, 1987), hydrothermal leaching (Cathelineau & Holliger, 1989).
1987) and or a combination of both. The metamorphic process may also cause severe leaching of REE from the albitized granite. For example the REE depletion may be related to the transformation of accessory minerals such as monazite to apatite and thorite in some albitized granites (Boulvais et al., 2007).

Also, the slightly positive Tb anomaly (Fig.8a and b) in the studied samples is attributed to its preferential accommodation within the fluorite structure (Yonan, 1999). This is confirmed by the petrographic study which shows the monzogranite and pegmatite contains fluorite.

CONCLUSIONS

The studied monzogranite and associated pegmatites represent a part of the basement rocks of the Central Eastern Desert of Egypt. The pegmatites occur as elongated, small pockets and lenses exhibiting variation in color and display sharp contacts with the host granite. The pegmatites are characterized by obvious simple zonation. The common accessory minerals are zircon, allanite, fluorite, monazite, apatite and iron oxides.

The eU contents of the analyzed monzogranite samples range from 8 ppm to 20 ppm with an average 12 and the eTh contents range from 10 ppm to 23 ppm with an average 13.3. It is slightly higher than the average of granitic rocks, while the associated pegmatites, eU contents range from 28 ppm to 355 ppm with an average 134 and the eTh contents range from 24 ppm to 210 ppm with an average 97.6. It is worthy to mention that the altered granites belong to high-thorium, high-uranium granitic rocks where uranium is mostly located in the Th-rich accessory minerals as monazite, zircon and allanite. The studied monzogranites and associated pegmatites have low eTh/eU ratios indicating uranium enrichment.

The present study proved that radioactive minerals responsible for both U and Th contents in the studied monzogranite and associated pegmatite are uranothorite, uranophane, zircon, monazite, allanite and fluorite.

The normalized REE pattern of monzogranite and associated pegmatite samples exhibit a remarkable similarity in REE contents that revealed by the superposition of different trends of both LREE and HREE. These samples show similar chondrite-normalized REE patterns and have clearly fractionated LREEs relative to HREEs, with distinct negative Eu anomaly.

REE bearing accessory minerals such as monazite, apatite, allanite and sphene represents potential source of LREEs. This is confirmed by the petrographic study. Strong negative Eu-anomaly could be explained by the crystal fractionation of plagioclase from the felsic granitic melts at low pressures or partitioning into feldspar during fractionation. Another alternative explanation for the negative Eu anomaly is based on the high oxygen fugacity (fO2) in the granitic melt due to volatile saturation. The depletion of REEs has been attributed to various processes including magmatic differentiation, hydrothermal leaching and or a combination of both. The slightly positive Tb anomaly in the studied samples is attributed to its preferential accommodation within the fluorite structure.

REFERENCES


Egypt with special emphasis to the origin of the uranophane mineralization. Egypt. Mineral., 12, 107-126


ABDEL MOEZ, A. SADEK and ABDEL HADI A. ABBAS

405p.


