

Petrogenesis of fractionated calc-alkaline A-type granites and related fluorite, North Eastern Desert, Egypt

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ABSTRACT

This study focuses on the petrogenesis of A-type granites and associated fluorite in the Qattar area, Northeastern Desert, Egypt. These granitoids show a geochemical signature typical for an A-type granite, characterized by a slightly peraluminous, highly SiO₂, alkalis (Na₂O+K₂O), U, Rb, Zr, contents, and belong to a group of Ferroan granites with low MgO, CaO, and MnO contents and high FeO^{tot}/FeO^{tot}+MgO ratios. These rocks emplaced during a late-collisional stage at a temperature of ~755.3 °C and pressures from 2.33 to 2.95 kbars under oxidizing conditions ($\log fO_2$ -14.85 to -14.91). Moreover, a high 10,000×Ga/Al ratio, negative Eu anomaly, and the tetrad pattern of Qattar alkali-feldspar granites backed up their formation by fractional crystallization and fluid-melt interactions during the late stage of magmatic differentiation. Fluorite mineralization is recorded as four fluorite types (sky blue-, light-yellow-, colorless-, and yellow-type) formed by hydrothermal processes. These fluorites show a wide range of rare earth element variations (ΣREE from 76.05 to 166.15) and negative Eu anomaly, which indicates crystallization temperatures above 200 °C. Moreover, Gd / Yb ratios are < 1 in sky blue and light-yellow fluorite and >1 in colorless and yellow fluorites, reflecting various conditions of REE leaching.

1. Introduction

Most of the northern region of the East African Orogen is covered by the Arabian-Nubian Shield (ANS), which has evolved as a result of a complex amalgamation of intra-oceanic island arcs± oceanic plateaus (880 - 630 Ma, [1]. Three main stages can distinguish in the ANS's magmatic and tectonic evolution, which is produced due to various geodynamic events. These stages are: (i) The subduction stage was formed from the amalgamation of island-arc volcano-sedimentary sequences and plutonic rocks [2]–[5] (~870–670 Ma); (ii) continental collision stage (~650–640 Ma), included the formation of calc-alkaline gabbro's and granodiorites which, formed from the convergence between the juvenile accrete ANS crust with continental blocks of west Gondwana to form the East African orogen [6]–[9], and (iii) post-collisional stage (630-550 Ma), as defined by the calc-alkaline to alkaline/peralkaline volcanic and granitic rocks development [10]–[13].

According to [14], three main lithologic terrains make up Egypt's Eastern Desert (ED): i) the northern one, which is abundant in massive granites; ii) the central one, which has the greatest concentration of rocks with oceanic affinities; and iii) the southern one, which is abundant in gneissic granites. The Eastern Desert granitoid rocks of Egypt have been classified as syn- to late-orogenic granitoid assemblages (610-880Ma) and post-orogenic to anorogenic granitoids assemblages (475-600 Ma) [15]–[18]. The younger granites covered ≈ 22% of the total exposed Proterozoic belt, which fractionated to calc-alkaline I- and A-type granites such as alkali-feldspar granites, syenogranites, and monzogranites.

A-type granites are highly evolved and alkaline to peralkaline granites, and they have ages between 550 and 610 Ma based on the Rb-Sr and U-Pb age dating, respectively [19]–[21]. Rare metals bearing granitoids are volatile-rich rocks and highly evolved A-type granites, widespread in most Orogenic belts worldwide. They include granitoids that host the mineralization of one or more metal elements like U, Th, Nb, Ta, Sn, W, Zr and REEs. Moreover, they occasionally host rare metal minerals, such as uraninite, thorite, columbite, tantalite, pyrochlore, fergusonite, and cassiterite. Some of the Egyptian A-type granitoids host most of the known rare-metals (U, Th, W, Sn, Ta, Nb, Mo, and REE) such as Gabel Qattar (the subject of this study) and El-Erediya areas [22], [23]. Qattar younger granitoids classified as meta-aluminous

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mineralized A-type granites, which are affected by metasomatic alterations. Alterations lead to enrichment in the Y, W, U, Nb content and the chalcophile elements [24]–[29] in Qattar granitoids.

The significance of studying bulk geochemistry of granitoids and hosted fluorites from the Qattar pluton lies in its potential to provide valuable insights into mineral exploration and resource assessment. Major, trace, and rare-earth elements found in these geological materials can offer crucial information about the processes involved in their formation and their potential mineralization. By analyzing the bulk geochemistry, researchers can identify the source of the granitic rocks and evaluate their magmatic evolution, which can help determine the tectonic setting and the potential for associated mineral deposits. Moreover, studying trace and rare-earth elements can help identify geochemical anomalies and establish geochemical signatures associated with economically significant mineralization. Understanding the distribution and concentration of economically essential elements in granitoids and fluorites from the Qattar pluton can aid in targeting areas with high mineralization potential, guiding exploration efforts, and optimizing resource assessment strategies. The findings from these studies can contribute to informed decision-making in the mining industry, promoting efficient and sustainable resource extraction practices. The petrogenesis of A-type granitoids in ANS is still controversial, as various of their chemical characteristics do not reflect a proper crystal-melt balance. In this work, bulk geochemistry studies (major, trace, and rare-earth elements) of granitoids and hosted fluorites from Qattar pluton are presented, which investigate the origin and evolution of the post-collisional A-type Qattar granitic magmatism and its relevance to associated ore deposits.

2. Geological Background

The Qattar area, located in the Northeastern Desert of Egypt, exhibits a complex geological background shaped by various tectonic and magmatic events. The region comprises metamorphic and igneous rocks that provide valuable insights into the geological evolution of the area. The Qattar area is underlain by a basement complex composed of Precambrian rocks, including gneisses, schists, and granitic intrusions, indicating the presence of ancient crustal materials[25]. This basement complex has undergone multiple episodes of deformation and metamorphism, resulting in the formation of various structural features such as folds, faults, and shear zones. Also, this region is characterized by the presence of several granite plutons, including the Qattar pluton, which represents a significant magmatic event. These granitoid intrusions are commonly associated with tectonic processes such as subduction and collision, indicating the involvement of convergent plate boundaries in the

geological history of the Qattar area. The emplacement of these granitoids has not only influenced the regional tectonic framework but has also contributed to the development of mineralization in the area.

Qattar granite intrusion covers around 35 km² between latitudes 27° 05' - 27° 07' N and longitudes 33° 14' 42'' - 33° 19' E (Fig. 1). These granites formed isolated masses of older diorite and granodiorite as well as rugged mountain peaks of pink to red medium-grained younger granites [25], [30], [31]. These granites are divided into the three phases G1, G2, and G3[25]. In addition to uranium mineralization, the younger granite has undergone several post-magmatic hydrothermal modifications, including fluoritization, hematitization, kaolinization, and epidotization [24]. Qattar granites are represented by syenogranites and alkali feldspar granites[22]. These granites comprise perthite, alkali-feldspar, quartz, plagioclase, biotite, and hornblende (found only in syenogranite). Magnetite, ilmenite, monazite, apatite, allanite, xenotime, and zircon are the main accessory minerals, whereas fluorite, muscovite, hematite, and clay minerals are found as secondary minerals[29], [32], [33].

The younger granite under study has undergone several post-magmatic hydrothermal alterations [24]. There are two types of mineralization in Gabal Qattar: 1) the U–fluorite and 2) molybdenum–bismuth–silver ores. The fluorite associated with U mineralization is exposed through mining works at more profound levels of granite [22]. Fluorite mineralization is hosted in alkali-feldspar granites, controlled by two conjugate fault systems trending NW–SE and NNE–SSW faults [34] (Fig. 1).The alkali-feldspar Qattar granite is medium to coarse-grained. These granitoids were composed of K-feldspars, quartz, plagioclase, and biotite. Fluorite, calcite, muscovite, magnetite, apatite, monazite and zircon found as minor and accessory minerals [22], [29].

3. Material and Methods

Twenty-four samples, representing 8 samples of Qattar granites and 16 of hosted fluorites, investigated in detail. The major, trace, and REE element concentrations (Tables 1 and 2) were carried out at the Earth Science Institute, Vienna University. The major and trace element concentrations were carried out by X-ray fluorescent (XRF) technique using a Philips PW 2400 series spectrometer on fused and pressed powder discs, respectively. Loss on ignition (LOI) was determined after igniting 5 g of rock sample powder in porcelain crucibles at 1050 °C for 1 h. Analytical precision is better than 1% and 2 - 5% for major and trace elements, respectively. REE concentrations were carried out with the inductively coupled plasma-mass spectrometer (ICP-MS) Elan 6100 with Hg Flow Injection System "FIMS 100" and Laser Sampler "LSX 200" has used for REE analyses.

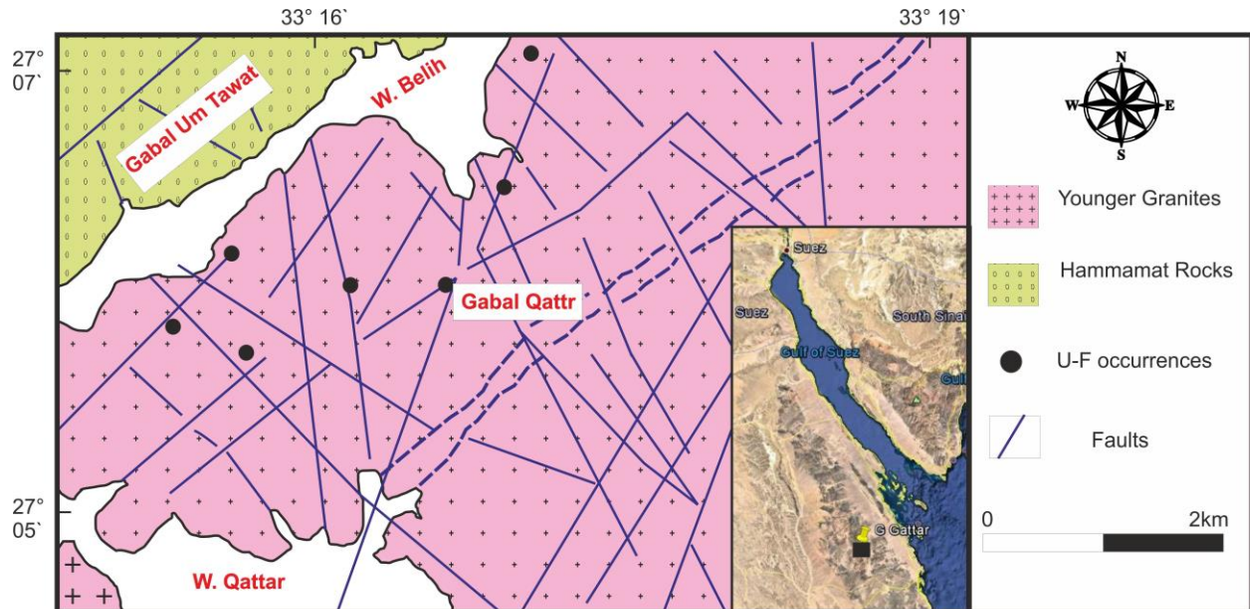


Fig. 1 Geological map of Qattar granite, North Eastern Desert, Egypt, modified after [22].

4. Result and discussion

4.1 Geochemical characteristics and tectonic setting

The major-, trace- and rare earth elements for Qattar granites listed in Table 1. These rocks are highly and limited variations of SiO_2 content ranging from 74.17 to 74.74 wt.% and have low abundances of Al_2O_3 (avg. 13.06 wt. %), MgO (avg. 0.03 wt. %), CaO (avg. 0.27 wt. %), Fe_2O_3 (avg. 0.92 wt. %), and high alkalis contents ($\text{Na}_2\text{O}+\text{K}_2\text{O}=8.79\text{--}9.02$ wt.%). Moreover, there are rich in high field-strength element contents (HFSE, Rb 341.33 ppm, Cs 14.6 ppm, Nb 156.9 ppm, Ta 10.14 ppm, Th 37.28 ppm, U 25.1 ppm, and Pb 18.13 ppm, on average; Table 1). All these features are characteristic of A-type granites [35], [36]. In the diagrams of ANNOR–Q'-F' and A/NK–A/CNK, the Qattar granites are plotted in the alkali-feldspar granite and peraluminous fields [37], [38] (Fig. 2a, b; respectively), reliable with their mineralogical characteristics. These rocks show high $\text{FeO}^{\text{tot}}/(\text{FeO}^{\text{tot}}+\text{MgO}=0.96\text{--}0.98)$ and are plotted in Ferron A-type granites (Fig. 2c) [39]. They investigated high K-calc-alkaline affinity with highly fractionated calc-alkaline (HFCA) rocks [40], [41] (Fig. 2d, e; respectively) and classified them as calc-alkaline Egyptian younger granitoids (Fig. 2f) [42]. Studied granites have K/Rb ratios (112.85-124.90) <140, which indicates the Rb separated during the aqueous liquid phase segregation stage at highly evolved melt [43], [44]. Moreover, they plotted enormously evolved and fractionated granitoids (Fig. 3a) [45] with barren nature [46] (Fig. 3b).

Qattar alkali feldspar granites have low total REE abundances (104.5–123.90 ppm), and display tetrad chondrite-normalized REE pattern [47] (Fig. 3c), characterized by slight enrichment in light REE (LREE; $(\text{La}/\text{Yb})_{\text{N}}=0.32\text{--}0.61$) with characteristic negative Eu anomalies (0.10–0.28). They display insignificant fractionations LREE $(\text{La}/\text{Sm})_{\text{N}}=0.61\text{--}1.57$) and (high REE)

HREE $(\text{Gd}/\text{Yb})_{\text{N}}=0.45\text{--}0.50$), accepted with those of the calc-alkaline granitoids of Sinai [12], indicating these rocks have resulted from volatiles transfer of REEs by F complexes [48]. The primitive mantel-normalized diagram [47] (Fig. 3d) exhibits negative Sr, P, and Ti anomalies and weak to moderate La, Ba, Ce, Pr, P, Eu, and Yb anomalies, which indicate plagioclase, K-feldspar, titanate, and apatite fractionated in that order of the magma. Moreover, Ti, Nb, P, and Sr depletion indicate granite derivation from crustal protolith [49]. Qattar granites have high $10,000\times\text{Ga}/\text{Al}$ ratios (6.04 to 6.42), higher than the S- and I-type granites (avg. 2.28 and 2.1028, respectively; [36]. Moreover, they fall in the A-type granite fields (Fig. 3e, f) [36] and fall in the fraction crystallization granites (A1 subtype; Fig. 4a) [50].

4.2 Crystallization Signatures

By using the Al-Ti thermometer [51], Qattar granitoids show high emplacement temperature (avg. 755.3 °C), which indicates highly fractionated granites [52]. The crystallization pressures were determined by using the equation proposed by [53]:

$$P = -980.74 \times (\text{Qz}) + 26.392 \times (\text{Qz})^2 - 0.2426 \times (\text{Qz})^3 + 12563 \quad (1)$$

Where (P) is the determined pressure and (Qz) is normative quartz value.

Applying of this equation indicates that Qattar granites crystallized under pressures ranging from 2.33 to 2.95 kbars. Moreover, the emplacement depth could calculate using the equation of [53] depending on the estimated pressures and density of continental crust (2.85 g/cm³), equivalent to 8.33-10.57-km depth.

The Oxygen fugacity buffers ($\log f\text{O}_2$) can be calculated by using [54] equilibrium expression:

$$\text{Log}(fO_2) = 9.36 - [24930 / T \text{ (k)}] + 0.046[(P \text{ (kb)} - 1/T \text{ (k)})] \text{ (2)}$$

Where (P) is the pressure in kbar and (T) is the temperature in K

The occurrence of magnetite minerals compared to Fe-Ti oxides [33] in Qattar granitoid support their crystallization under oxidizing conditions and controlled by NNO fugacity buffer (log fO_2 from -14.91 to -14.85, Fig. 4b) [55].

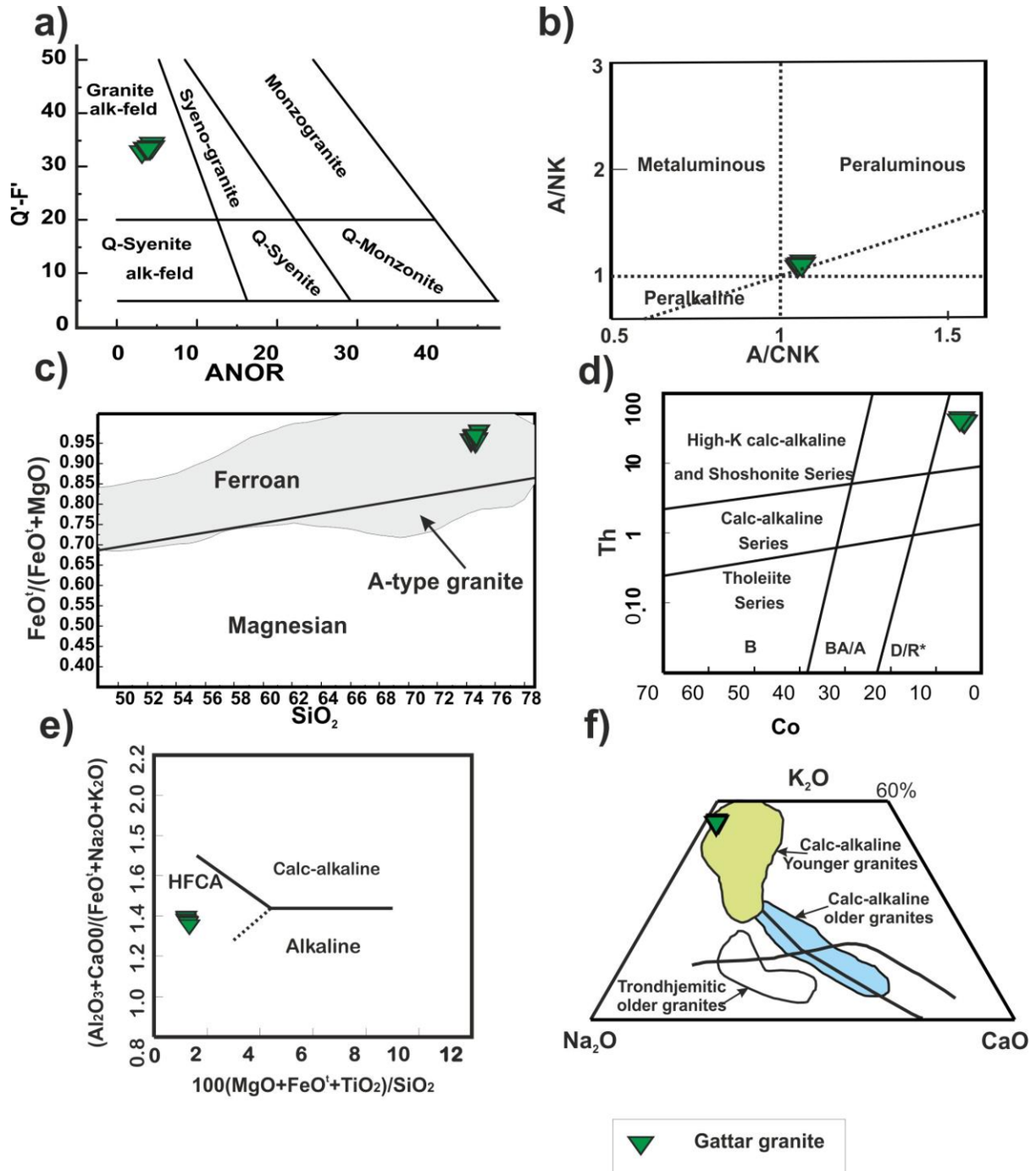


Fig. 2 (a) Q' - F' - ANOR diagram [38]; (b) $Al_2O_3/CaO+Na_2O+K_2O - Al_2O_3/Na_2O+K_2O$ diagram [37]; (c) $SiO_2 - FeO^{tot}/FeO^{tot} + MgO$ after [39]; (d) Co-Th variation diagram after [40]; (e) $Al_2O_3+CaO/(FeO^{tot}+Na_2O+K_2O) - 100*(TiO_2+FeO^{tot}+MgO)/SiO_2$ variation diagram [41]; (f) $Na_2O-CaO-K_2O$ ternary diagram showing the Egyptian younger and older granitoids fields after [42].

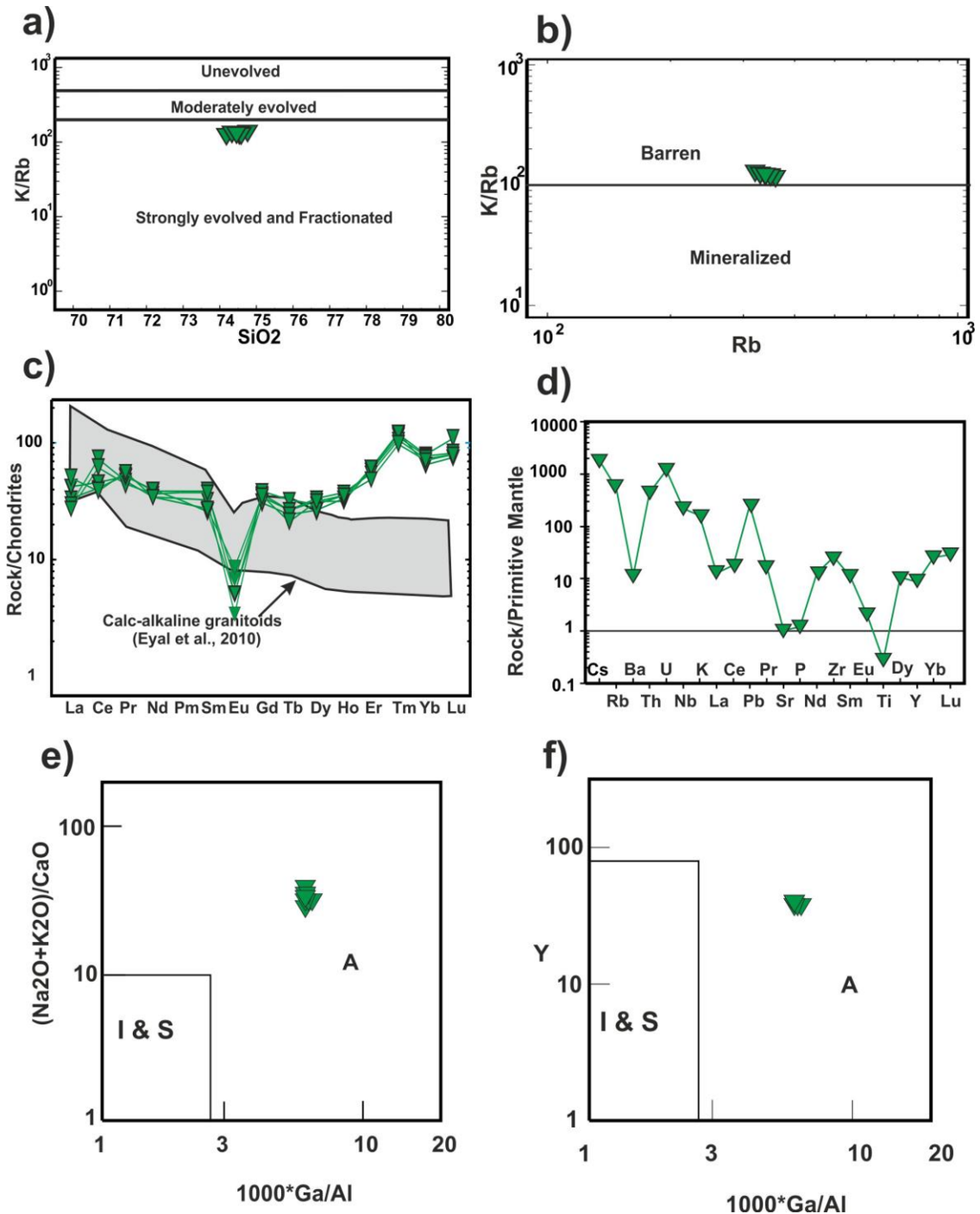


Fig. 3 (a) Blevin's K/Rb–SiO₂ diagram [45]; (b) Variations of K/Rb - Rb ratios of the studied granites [46]; (c) REE chondrite-normalized diagram [47] of Qattar granitoids. (d) Primitive mantle -normalized diagram [47] of Qattar granitoids; (e; f) Na₂O+K₂O- 10,000 Ga/Al, and FeO^{tot}/MgO-Zr+Nb+Ce+Y diagrams for A type granites [36].

4.3 Fluorine-uranium (F-U) contents and REEs Tetrad Effect

The highly differentiated Qattar granitoids exhibit an M-type tetrad effect (Fig. 3c), which is comparable to the REEs tetrad effects produced in the highly F-rich differentiated granites from the El-Erediya pluton in Egypt and Qianlishan composite plutons, S. China [23], [56], respectively). In characteristic magmatic systems, CHARAC (CHArge- and RAdius-Controlled) trace-element

behaviors predominated, but the lanthanides tetrad effects were commonly associated with non-CHARAC trace-element behaviors. Y-Ho, Zr-Hf, etc. pairs (elements with similar charge and radius) were controlled by chemical integration processes that led to trace element fractionations characterized by a geochemical system of non-CHARAC behavior [57], [58]. The Qattar granites show non-CHARAC behavior [57] (Fig. 4c). Besides, it occurs in highly evolved magmatic systems (rich in CO₂, H₂O, B, Li,

F, and/or Cl), which may be developed between hydrothermal fluids and pure silicate-melts in late-stage of crystallization [57], [59]. The REEs tetrad effect and non-

CHARAC trace element behavior of Qattar granitoids are often related to F-rich granitic magmas [58], [60], [61].

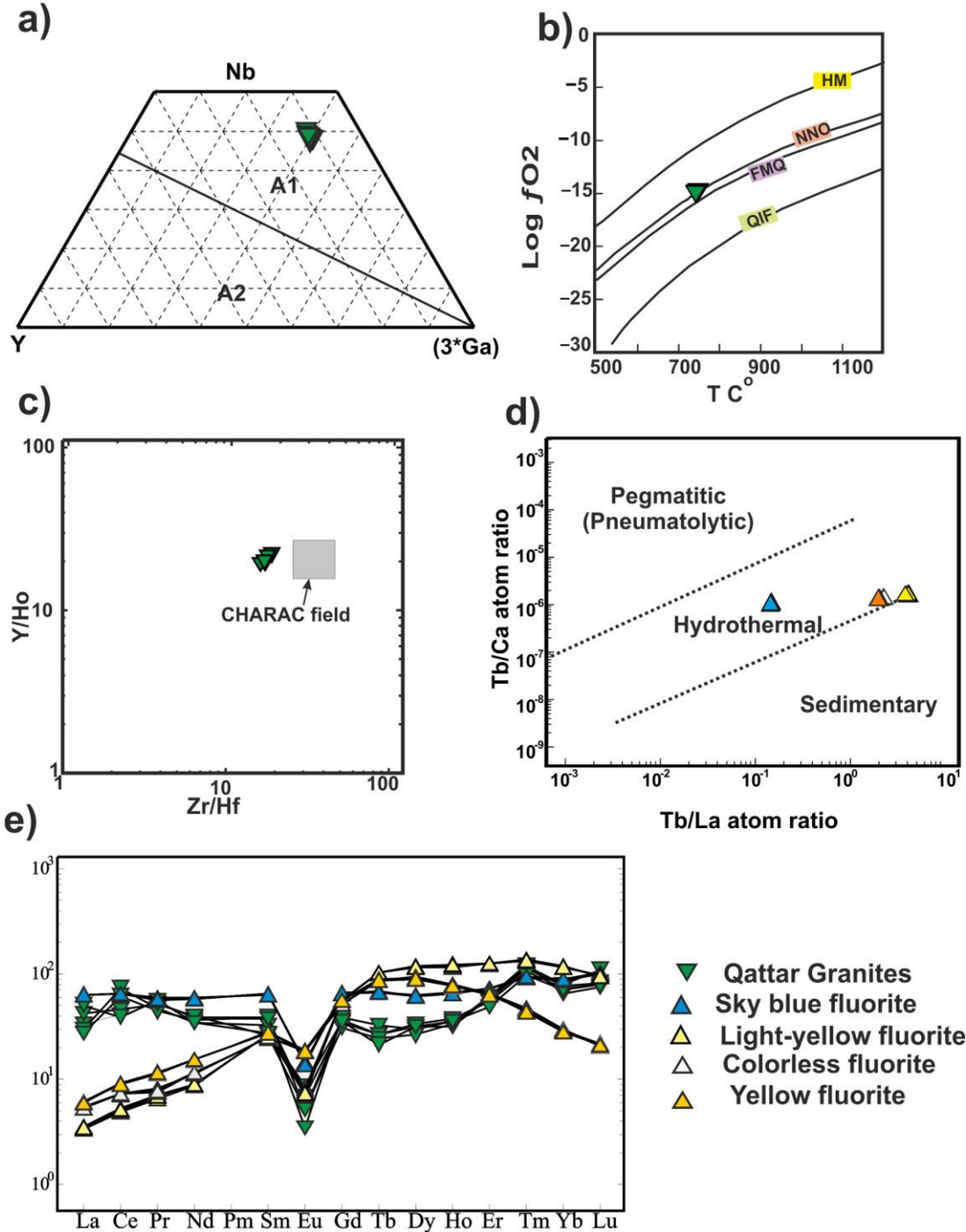


Fig. 4 (a) Nb-Y-3*Ga triangular diagram of[50]; (b) Temperature-oxygen fugacity (fO_2) diagram [55]; (c) Y/Ho-Zr/Hf diagram with (CHARAC) field after[57] ; (d) Tb/Ca-Tb/La variation ratios of Qattar fluorites[69]; (e) REE chondrite-normalized patterns of Qattar granite and their hosted fluorite [47].

Cd	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1
Sb	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1
Bi	7.3	7.7	7.7	7.1	7.9	7.3	7.7	7.5	7.525
Ag	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1
U	22.5	25.5	27.5	24	26	24	26.3	25	25.1
Th	36.3	34	40	36.3	40	35.5	38	38.1	37.28
K/Rb	124.9	116.9	112.85	120.5	116.68	120.53	115.36	118.43	118.27
Na₂O+K₂O	8.79	8.82	9.02	8.87	8.94	8.82	8.92	8.9	8.89
FeO^{tot}/FeO^{tot}+MgO	0.98	0.96	0.96	0.97	0.96	0.97	0.96	0.97	0.97
10000*Ga/Al	6.09	6.42	6.07	6.04	6.11	6.19	6.20	6.08	6.15

REEs (ppm)

La	8	7	6.50	10.00	12.00			8.7
Ce	25	45	38.00	28.00	24.00			32
Pr	5	4.5	4.20	4.90	5.30			4.78
Nd	18	16	17.00	18.50	16.00			17.1
Sm	6	5	5.80	4.00	4.20			5
Eu	0.3	0.2	0.40	0.30	0.50			0.34
Gd	7.5	8	7.00	6.40	7.20			7.22
∑LREEs	69.8	85.7	78.9	72.1	69.2			75.14
Tb	1	1.2	1.00	0.90	0.80			0.98
Dy	8	7.5	8.50	6.70	8.00			7.74
Ho	1.8	2	2.10	1.90	2.00			1.96
Er	10	9.5	9.00	10.20	8.00			9.34
Tm	3	2.9	2.70	3.10	2.50			2.84
Yb	13.5	13	12.50	11.00	12.00			12.4
Lu	2.8	2.1	2.00	1.90	2.00			2.16
∑LREEs	40.1	38.2	37.8	35.7	35.3			37.42
∑REEs	109.90	123.90	116.70	107.8	104.5			112.56

Some geochemical ratios

Eu/Eu*	0.14	0.10	0.19	0.18	0.28			0.18
(La/Sm)_N	0.73	0.77	0.61	1.37	1.57			1.01
(La/Yb)_N	0.36	0.33	0.32	0.55	0.61			0.43
(Gd / Yb)_N	0.45	0.5	0.45	0.47	0.49			0.47
T_{Al}/Ti.exp.C	756.6	754.9	753.90	754.7	755.8	755.4	754.9	755.30
log fO₂-	-14.85	-14.89	-14.91	-14.89	-14.87	-14.88	-14.89	-14.88
P	2.33	2.49	2.95	2.65	2.60	2.48	2.60	2.61
Depth	8.33	8.90	10.57	9.49	9.29	8.87	9.32	9.35

The normalization is based on the chondrite-normalized data given by [47] (1989): Eu anomaly is defined by $Eu/Eu^ = E_{UN} / (\sqrt{Sm_N \times Gd_N})$.

Qattar granites considered as uraniferous granites (U avg. 25.1 ppm > 4 ppm Clark value 4 ppm). Most U is found in high-temperature magma-hydrothermal systems as a compound with F [62]. F tends to change the liquidus phase and is also suited to produce melt depolymerization, which increases the solubility of U in the magma [63], [64]. The U fluid-magma partition coefficients are extremely low, ranging from 0.03 to 0.04 [65]. As a result, F-rich-magma tends to have a higher U concentration [66]. Besides, Qattar alkali-feldspar granites contain insignificant amounts of biotite and apatite minerals, which F in magma tends to join in its crystal lattices in the post-magmatic stages [31], [67]. So, these granites are considered as highly evolved F-enriched granites [33]. Additionally, according to [68] F and U are two elements accumulating in residual melts during crystal differentiation. In oxidizing conditions, F can change the valency of uranium from U^{4+} to U^{6+}

and produce soluble fluoride or hydroxyfluoride complexes as the U-F combination.

4.4 Geochemistry of Fluorite

Qattar fluorites are classified into four types (sky blue-, light-yellow-, colorless- and yellow-type), according to the mineral chemistry analysis (Table 2). From the Tb/Ca-Tb/La diagram [69], these fluorites were formed by hydrothermal processes (Fig. 4d). The colorless fluorite is abundant in trace elements, particularly Ti, Ba, Pb, Sr, and Zn (Table 2). The trace element enrichments in these fluorite-bearing granites compared to barren fresh granites (Fig. 3b) are suitable for leaching from major and accessory minerals of Qattar granite during fluid-mineral interactions [22].

Table 2: Chemical analyses of the Qattar fluorites

	F01	F01a	F01b	F01d	Avg.	F02	F02b	F02c	F02e	Avg.
	sky blue fluorites					Light-yellow fluorites				
Ca wt %	56.71	58.42	57.96	56.35	57.36	56.69	56.85	57.91	55.49	56.74
Na	0.11	0.11	0.11	0.11	0.11	0.17	0.17	0.17	0.17	0.17
	0.004	0.004	0.004	0.004	0.004	0.02	0.02	0.02	0.02	0.02
Mg	0.005	0.004	0.005	0.004	0.004	0.02	0.02	0.02	0.02	0.02
Fe	0.25	0.24	0.24	0.24	0.24	0.27	0.27	0.27	0.27	0.27
Al	0.02	0.02	0.02	0.02	0.02	0.03	0.03	0.03	0.03	0.03
Mn	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01
Ti	0.04	0.04	0.04	0.04	0.04	0.05	0.05	0.05	0.05	0.05
F	42.12	40.29	41.06	42.12	41.40	41.25	42.36	40.28	43.21	41.78
Sl	0.01	0.02	0.01	0.11	0.04	0.09	0.08	0.07	0.10	0.09
total	99.28	99.17	99.47	99.02	99.23	98.61	99.87	98.84	99.38	99.17
Trace elements and REE (ppm)										
Ba	2.50	2.60	2.55	2.58	2.56	4.00	4.20	4.10	4.15	4.11
Co	0.78	0.77	0.78	0.77	0.77	0.95	1.00	0.98	0.99	0.98
Cr	6.50	6.30	6.40	6.35	6.39	4.80	4.60	4.70	4.65	4.69
Cu	13.60	12.90	13.25	13.08	13.21	98.40	98.20	98.30	98.25	98.29
Li	2.67	2.59	2.63	2.61	2.63	0.34	0.31	0.33	0.32	0.32
Mo	15.27	15.23	15.25	15.24	15.25	0.40	0.35	0.38	0.36	0.37
Ni	1.50	1.30	1.40	1.35	1.39	11.70	12.10	11.90	12.00	11.93
Pb	43.10	42.20	42.65	42.43	42.59	6.00	5.80	5.90	5.85	5.89
Rb	0.12	0.10	0.11	0.11	0.11	0.37	0.32	0.35	0.33	0.34
Sr	33.70	33.10	33.40	33.25	33.36	15.70	15.20	15.45	15.33	15.42
Zn	34.20	35.00	34.60	34.80	34.65	61.50	62.50	62.00	62.25	62.06
Sb	0.13	0.10	0.12	0.11	0.11	0.21	0.20	0.21	0.20	0.20
V	0.90	0.90	0.80	0.85	0.86	0.80	1.00	1.00	1.00	0.95

Sc	0.50	0.60	0.50	0.55	0.54	0.50	0.60	0.50	0.55	0.54
Y	308.85	307.54	308.20	307.87	308.11	682.35	683.21	682.78	683.00	682.83
La	15.07	15.12	15.10	15.11	15.10	0.79	0.84	0.82	0.83	0.82
Ce	39.23	39.85	39.54	39.70	39.58	3.01	3.21	3.11	3.16	3.12
Pr	5.70	5.40	5.55	5.48	5.53	0.62	0.68	0.65	0.67	0.65
Nd	27.75	27.26	27.51	27.38	27.47	4.10	4.21	4.16	4.18	4.16
Sm	9.77	9.85	9.81	9.83	9.82	4.02	4.09	4.06	4.07	4.06
Eu	0.84	0.79	0.82	0.80	0.81	0.48	0.41	0.45	0.43	0.44
Gd	13.32	13.54	13.43	13.49	13.44	11.56	11.62	11.59	11.61	11.59
∑LREE	111.68	111.81	111.75	111.78	111.75	24.58	25.06	24.82	24.94	24.85
Tb	2.57	2.49	2.53	2.51	2.53	3.81	3.84	3.83	3.83	3.83
Dy	15.58	15.96	15.77	15.87	15.79	29.64	30.00	29.82	29.91	29.84
Ho	3.68	3.74	3.71	3.73	3.71	6.57	7.00	6.79	6.89	6.81
Er	12.13	12.02	12.08	12.05	12.07	20.56	21.00	20.78	20.89	20.81
Tm	2.33	2.41	2.37	2.39	2.38	3.36	3.48	3.42	3.45	3.43
Yb	14.93	15.10	15.02	15.06	15.03	19.76	20.00	19.88	19.94	19.90
Lu	2.53	2.62	2.58	2.60	2.58	2.46	2.41	2.44	2.42	2.43
∑HREE	53.75	54.34	54.05	54.19	54.08	86.16	87.73	86.95	87.34	87.04
REE	165.43	166.15	165.79	165.97	165.84	110.74	112.79	111.77	112.28	111.89
Th	1.10	1.00	0.90	0.95	0.99	0.40	0.50	0.40	0.45	0.44
U	1.30	1.40	1.35	1.38	1.36	0.80	0.70	0.75	0.73	0.74
Gd / Yb norm	0.72	0.73	0.72	0.72	0.72	0.47	0.47	0.47	0.47	0.47
Eu/Eu*	2.61	2.60	2.61	2.60	2.60	1.80	1.81	1.80	1.81	1.80
	F03	F03c	F03d	F03f	Avg.	F04	F04d	F04e	F04g	Avg.
	Colorless fluorites					Yellow fluorites				
Ca wt %	55.47	57.01	57.48	57.12	56.77	58.95	58.94	56.63	58.41	58.23
Na	0.13	0.13	0.13	0.13	0.13	0.14	0.14	0.14	0.14	0.14
K	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
Mg	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01
Fe	0.26	0.26	0.26	0.26	0.26	0.25	0.25	0.25	0.25	0.25
Al	0.02	0.02	0.02	0.02	0.02	0.01	0.01	0.01	0.01	0.01
Mn	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01
Ti	0.04	0.04	0.04	0.04	0.04	0.04	0.04	0.04	0.04	0.04
F	43.02	41.23	41.53	41.36	41.79	39.86	40.22	42.32	40.28	40.67
Si	0.12	0.14	0.05	0.17	0.12	0.07	0.05	0.12	0.18	0.11
total	99.08	98.85	99.53	99.12	99.14	99.34	99.67	99.53	99.33	99.47
Trace elements and REE (ppm)										
Ba	132.80	134.00	133.40	133.70	133.48	1.90	2.00	1.95	1.98	1.96
Co	0.80	0.75	0.78	0.76	0.77	0.85	0.82	0.84	0.83	0.83
Cr	9.80	9.50	9.65	9.58	9.63	9.60	9.60	9.60	9.60	9.60

Cu	32.10	32.20	32.15	32.18	32.16	14.60	14.40	14.50	14.45	14.49
Li	0.33	0.34	0.34	0.34	0.34	0.14	0.13	0.14	0.13	0.13
Mo	0.45	0.42	0.44	0.43	0.43	21.62	21.74	21.68	21.71	21.69
Ni	11.10	11.30	11.20	11.25	11.21	11.40	11.54	11.47	11.51	11.48
Pb	643.80	649.50	646.65	648.08	647.01	54.80	55.00	54.90	54.95	54.91
Rb	0.15	0.16	0.16	0.16	0.16	0.10	0.10	0.10	0.10	0.10
Sr	116.00	112.00	114.00	113.00	113.75	40.50	41.20	40.85	41.03	40.89
Zn	55.20	54.80	55.00	54.90	54.98	52.70	53.20	52.95	53.08	52.98
Sb	0.01	0.01	0.01	0.01	0.01	0.02	0.02	0.02	0.02	0.02
V	0.90	0.90	0.70	0.80	0.83	0.80	0.70	0.70	0.70	0.73
Sc	0.50	0.70	0.60	0.65	0.61	0.50	0.50	0.50	0.50	0.50
Y	302.36	301.54	301.95	301.75	301.90	352.82	353.24	353.03	353.14	353.06
La	1.27	1.28	1.28	1.28	1.28	1.46	1.41	1.44	1.42	1.43
Ce	4.49	4.42	4.46	4.44	4.45	5.50	5.58	5.54	5.56	5.55
Pr	0.78	0.71	0.75	0.73	0.74	1.07	1.11	1.09	1.10	1.09
Nd	5.23	5.42	5.33	5.37	5.34	7.12	7.14	7.13	7.14	7.13
Sm	3.79	3.82	3.81	3.81	3.81	4.27	4.25	4.26	4.26	4.26
Eu	1.07	1.09	1.08	1.09	1.08	1.06	1.09	1.08	1.08	1.08
Gd	10.71	10.74	10.73	10.73	10.73	11.47	11.51	11.49	11.50	11.49
∑LREE	27.34	27.48	27.41	27.45	27.42	31.95	32.09	32.02	32.06	32.03
Tb	3.24	3.26	3.25	3.26	3.25	3.25	3.29	3.27	3.28	3.27
Dy	23.52	23.64	23.58	23.61	23.59	22.89	23.00	22.95	22.97	22.95
Ho	4.52	4.48	4.50	4.49	4.50	4.37	4.32	4.35	4.33	4.34
Er	10.81	10.88	10.85	10.86	10.85	10.59	10.62	10.61	10.61	10.61
Tm	1.18	1.21	1.20	1.20	1.20	1.14	1.11	1.13	1.12	1.12
Yb	4.90	5.10	5.00	5.05	5.01	4.87	4.82	4.85	4.83	4.84
Lu	0.54	0.52	0.53	0.53	0.53	0.56	0.53	0.55	0.54	0.54
∑HREE	48.71	49.09	48.90	49.00	48.92	47.67	47.69	47.68	47.69	47.68
REE	76.05	76.57	76.31	76.44	76.34	79.62	79.78	79.70	79.74	79.71
Th	0.30	0.20	0.10	0.15	0.19	0.80	0.70	0.80	0.75	0.76
U	1.00	0.80	0.80	0.80	0.85	1.00	0.50	0.70	0.60	0.70
(Gd / Yb)_N	1.77	1.70	1.74	1.72	1.73	1.91	1.93	1.92	1.93	1.92
Eu/Eu*	1.81	1.82	1.82	1.82	1.82	1.86	1.85	1.86	1.85	1.86

REE of Qattar fluorites (Table 2) shows a wide variety of variation (\sum REE from 76.05 to 166.15) according to the variability of the parameters during the mineralization process [70]. The total REE in the sky-blue and light-yellow fluorites (avg. \sum REE 165.84 and 111.89, respectively) are higher than the colorless and yellow fluorites (avg. \sum REE 76.34 and 79.71, respectively) (Table 2). Figure 4e shows the chondrite-normalized REE patterns of Qattar granites and the hosted fluorite exhibiting a negative Eu anomaly

(1.81-2.60; Tables 2, 3), indicating the crystallization temperatures above 200 °C [71]–[74]. The Eu anomaly is controlled by thermal, chemical, and redox conditions at average temperatures < 200 °C [26]. Moreover, sky blue and light-yellow fluorites exhibit higher values of LREE compared to HREE and (Gd / Yb) ratios < 1, in contrast with colorless and yellow fluorite show higher values of HREE compared to LREE and (Gd / Yb) ratios >1. These different fluorite patterns denoted that: 1) strong effects of

chemical interaction and physiochemical conditions of REE leaching [74] and 2) the parent fluids interacted with host rocks under reduced or weakly oxidizing conditions, backed found amount of sulfides in Qattar granite [22], [26], [73], [75].

4.5 Petrogenesis

The Qattar highly differentiated granites had numerous A-type granite features, including 1) high SiO_2 , $\text{Na}_2\text{O}+\text{K}_2\text{O}$, Zr, Nb, and Rb; 2) high $10000 \cdot \text{Ga}/\text{Al}$ ratio (avg. 6.15); 3) high K-calc-alkaline natures and weakly peraluminous; 4) MgO , P_2O_5 , TiO_2 , CaO , Eu, and Sr contents depletion [35], [36], [76], [77]. On the other hand, the low Eu/Eu^* ratio (avg. 0.18) showed that the Qattar A-type granites are highly differentiated [57]. Comparing these ratios with the lithospheric root propose contributions of slab break-off and/or lower crust [78], [79] at depth > 30km (Fig. 5a) [80]. U and Th have negative correlations with Zr/U and Zr/Th ratios (Fig. 5b, c; respectively), indicating that U-Th elements are controlled by Qattar granitoids magmatic accessory minerals [29]. However, the U and Th contents showed that magmatic fractionation was applied to the Qattar melt, which affects magmatic and late- to post-magmatic processes related to the exsolution of a fluid phase (Fig. 5d) [81]. Moreover, in the H_2O -F haplogranite Qz-Ab-Or- system (Fig. 5e) Qattar granitoids were plotted close to the minimum melt composition in the existence of fluorine at temperature ≈ 780 °C and pressure 2-5 kbar [82]–[84]. Moreover, Qattar granites are post-collisional tectonic settings (Fig. 5f) [37], supported by [85].

4.6 Geodynamic implications

The origin of post-collision magmatism in the ANS can be used to explain the extensional collapse, followed by continental collision from 600 to 650 Ma between West Gondwana and the juvenile ANS crust [86] (Fig. 6a). According to many studies [4], [8]–[10], [87], this continental collision affected significant crustal thickening, but the extensional collapse formed at 550 to 600Ma and caused the thicker lithosphere to weaken (Fig. 6b, c). The extensional collapse was regulated by lithospheric delamination and slab break-off [10], [12], [88], [89]. During the late collisional stage (630–610 Ma), a slab window opened, and unusually elevated temperatures supplied by the upwelling asthenosphere (Fig. 6 b, c), which caused an extensive melting process [90], (Fig. 6d). At late-collisional to early post-orogenic (610–280 Ma) stages, calc-alkaline and alkaline magmas were obtained in the ANS [12], [91], which included rich K calc-alkaline Qattar granitoids.

5. The study contribution to Sustainable Development Goals (SDGs)

Studying the bulk geochemistry of granitoids and hosted fluorites from the Qattar pluton aligns with several Sustainable Development Goals (SDGs), promoting sustainable economic development and responsible resource management.

This research contributes to SDG 9: Industry, Innovation, and Infrastructure, by providing valuable scientific knowledge that can support the development of sustainable mining practices. By understanding the

geochemical characteristics of these geological materials, mining companies can optimize their exploration efforts, reducing the environmental impact associated with mineral extraction and ensuring efficient resource use. Furthermore, this research also supports SDG 12: Responsible Consumption and Production. By studying the bulk geochemistry of granitoids and fluorites, it becomes possible to identify areas with high mineralization potential, leading to targeted mining activities and minimizing waste generation. This knowledge can help promote the sustainable production of mineral resources, reducing the environmental footprint and ensuring the responsible use of natural resources. Moreover, SDG 13: Climate Action can be linked to this research through its potential to inform sustainable resource management strategies. Understanding the geological processes and mineralization associated with granitoids and fluorites can aid in identifying mineral deposits, including those relevant to renewable energy technologies. Directing exploration efforts towards these resources supports the transition to a low-carbon economy and contributes to mitigating climate change.

Therefore, the scientific studies on the bulk geochemistry of granitoids and hosted fluorites from the Qattar pluton have the potential to positively impact multiple SDGs, promoting sustainable economic development, responsible resource extraction, and climate action.

6. Future directions

Several future recommendations and areas for further study can be proposed to advance the understanding of the bulk geochemistry of granitoids and hosted fluorites from the Qattar pluton.

Petrogenetic Modeling: Petrogenetic modeling techniques, such as geochemical modeling and thermodynamic calculations, can help reconstruct the magmatic history and source characteristics of the granitoids. These models can provide insights into the melting processes, magma evolution, and magma source regions, aiding in interpreting the geochemical signatures observed in the rocks[56].

Integration with geophysical surveys, remote sensing, and machine learning: Integrating bulk geochemical data with these techniques can facilitate the correlation between geochemical anomalies and subsurface structures or mineralization zones. This multidisciplinary approach can enhance the understanding of the spatial distribution of mineralization and assist in target selection for future exploration efforts [92], [93].

Environmental Implications: Investigating the potential environmental implications of the granitoids and hosted fluorites, especially regarding the presence of trace elements and rare-earth elements, is crucial. Assessing the mobility and behavior of these environmental elements can provide insights into potential environmental risks associated with mining activities and aid in developing sustainable mining practices.

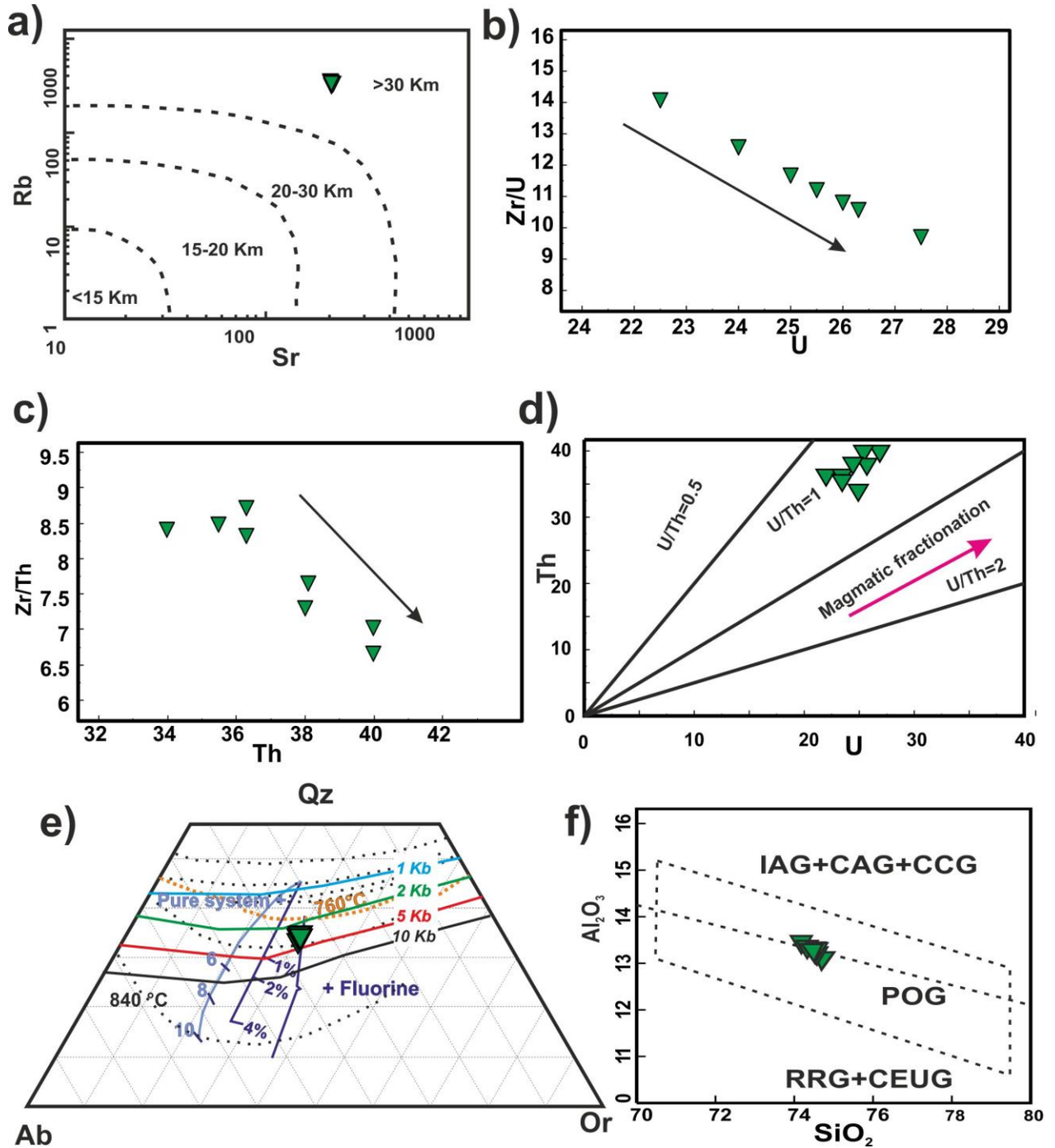


Fig. 5 (a) Sr-Rb diagram[80]; (b; c; d) U-Zr/U; Th- Zr/Th; U-Th for Qattar granitoids; (e) Qz-Ab-Or ternary diagram, temperature lines proposed by [82], pressure lines proposed by [84] and fluorine content proposed by [83]; (f) SiO₂-Al₂O₃ variation diagram [37].

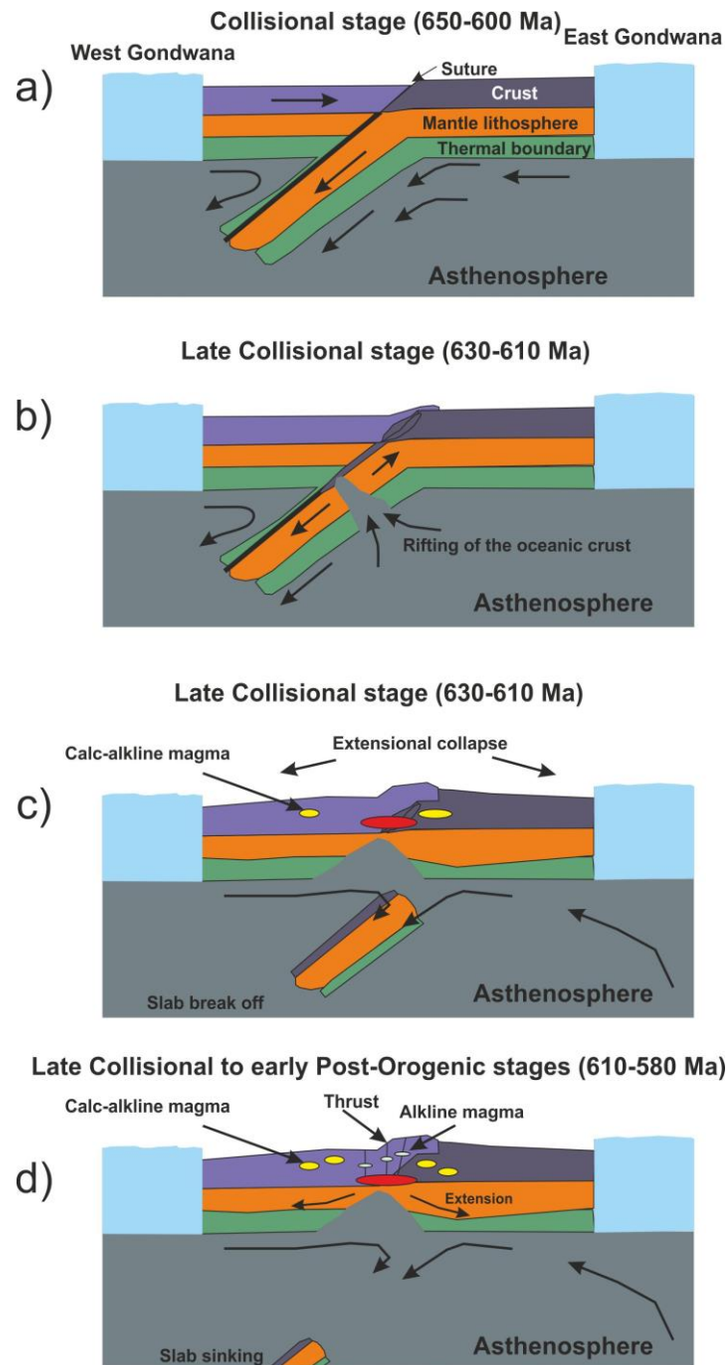


Fig. 6 Schematic tectono–magmatic evolution of the post-collisional stage of younger granite of Qattar area in the ANS (modified after; [10], [18], [87], [88], [91]) **(a)** The start of collision between East and West Gondwana, (650–600Ma); **(b)** Starting of extensional collapse and rifting, with raising the asthenosphere to fill the rift; **(c)** Slab breaks off and rises of hot asthenosphere lead to magmatism; **(d)** Slab sinks away and asthenosphere up wells, which caused common melting of the subducted oceanic crust and induce partial melting to produce a calc-alkaline and alkaline magmatism including high-K calc-alkaline Qattar granitic rocks in ANS.

6. Conclusions

Qattar alkali-feldspar granitoids originated from a highly fractionated rich K calc-alkaline magma source under oxidizing conditions (avg. $\log fO_2$ -14.88). These granitoids show a geochemical signature characteristic of A1-subtype granite, such as slightly peraluminous, highly SiO_2 , alkalis (Na_2O+K_2O), Rb, Zr, contents, rich K calc-alkaline magma, and belong to the ferroan granites group with low MgO ,

CaO , and MnO contents and high $FeO^{tot}/FeO^{tot}+MgO$ ratios. They have low REE (104.5–123.90 ppm), slight enrichment in LREE ($(La/Yb)_N = 0.32-0.61$), negative Eu ($Eu/Eu^*, 0.10-0.28$), and strong negative Ti, Pb, and Sr anomalies (primitive mantel-normalization) indicating magmatic fractional crystallization. They emplaced at depths between 8.33 to 10.57 km, at a temperature of 755.3 °C, and under pressure 2.33-2.95 kbars by

delaminating the lithospheric root and slab break-off. They are highly F-rich differentiated granites hosted by four types of fluorites, varying from sky blue, light-yellow, colorless, and yellow in color. Moreover, these fluorites show a wide range of REE variations and negative Eu anomaly, indicating crystallization temperatures above 200 °C (under reduced or weakly oxidizing conditions) and affected by chemical complexation and physiochemical conditions of REE leaching.

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