CONCENTRATION VARIATIONS OF MAJOR AND MINOR ELEMENTS ACROSS VARIOUS ALTERATION ZONES IN PORPHYRY COPPER DEPOSIT AT SUNGUN, EAST AZARBAIDJAN, IRAN

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Abstract

The porphyry copper deposit at Sungun is located in East Azarbaidjan, NW of Iran. The Sungun porphyries (Oligo-Miocene) occur as stocks and dikes and intruded a series of Oligocene pyroclastics and lavas, Eocene arenaceous-argillaceous rocks, and Upper Cretaceous carbonates. The porphyry stocks are divided into two groups, 1) porphyry stocks I (quartz monzo-diorite) and 2) porphyry stock II (quartz monzonite, granodiorite, and granite). Porphyry stock II hosts the copper and molybdenum ores. Numerous and various cross-cutting veinlets and micro-veinlets of quartz, sulfides, sericite, carbonates, and sulfates were developed in porphyry stock II. Three distinct types of hydrothermal alterations and sulfide mineralizations are recognized at Sungun; 1) hypogene, 2) contact metasomatic, and 3) supergene. The vertical variations in concentration of 12 elements (S, Cu, Mo, K, Rb, Ca, Sr, Na, Mg, Fe, Al, and Si) across various hypogene alteration zones at Sungun demonstrate that S, Cu, Mo, Fe, K, and Rb are mainly concentrated in the phyllic and potassic-phyllic zones where the abundance of Ca, Sr, Mg, and Na are appreciably low relative to the potassic zone. The values of Al and Si are relatively high in the supergene zones and their abundance gradually decrease downward toward the potassic zone. The copper and molybdenum are substantially localized in the phyllic and potassic-phyllic zones particularly in the central part of the mineralized porphyry stock where the rocks underwent severe stockwork-type fracturing and micro-fracturing. Their concentrations are relatively low in the potassic zone.

Keywords: Porphyry copper; Veinlets and micro-veinlets; Concentration variation; Hypogene alterations; Major and trace elements

Introduction

The earliest work on geochemistry (fluid inclusions) of the porphyry copper deposit (PCD) at Sungun was

done by Etminan [4]. The later studies on geochemistry, alteration, and mineralization was carried out by Momenzadeh *et al.* [9], Emami and Babakhani [3], and Ghaderi [5]. Mehrpartou [8] studied many geologic

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aspects including petrology, alteration, mineralization, and geochemistry at Sungun. In 1994, over 100 diamond drill holes (DDH) in a grid pattern were dug at Sungun main ore body. Calagari [1], based upon systematic subsurface data, carried out relatively more detailed studies on various geological features at Sungun. Additional studies on "controls of alteration and mineralization" and "factors controlling copper stability on Sungun PCD" were presented by Hezarkhani and William-Jones [6] and Hezarkhani *et al.* [7], respectively. The following study considers the migration, concentration, and re-distribution of major and some minor elements (e.g., Rb, Sr, S, Cu, and Mo) in relation to the alteration and mineralization processes at Sungun.

Geology (background)

The Sungun PCD is located ~100 km NE of Tabriz in East Azarbaidjan, NW of Iran (see inset in Fig. 1). The Sungun porphyries are of Oligo-Miocene age, and intruded Upper Cretaceous carbonate rocks, a series of Eocene arenaceous-argillaceous rocks, and a series of Oligocene pyroclastics and lavas (Fig. 2; [3,8]). The Sungun hypabyssal intrusives, calc-alkaline in character, occur as stocks and dikes. The stocks are divided into two groups, 1) porphyry stocks I (principally quartz monzo-diorite) and 2) porphyry stock II (quartz monzonite through granodiorite to granite) which hosts the Sungun PCD. Dikes are categorized into four series (DK₁, DK₂, DK₃, and DK₄) varying in composition from quartz monzonite to granodiorite [1]. Late Tertiary-Quaternary volcanic rocks ranging in composition from latite to andesite cover some parts of Sungun porphyries (Fig. 2; [3,8]).

Three distinct types of hydrothermal alterations and sulfide mineralizations occurred at Sungun, 1) hypogene, 2) contact metasomatic, and 3) supergene [1]. The development of pervasive hypogene alteration and sulfide mineralization were intimately associated with intense stockwork-type hydro-fracturing manifested by numerous cross-cutting veinlets and micro-veinlets of quartz, sulfides, sericite, carbonates, and sulfates [2]. The hypogene alterations are divided into four types, 1) potassic, 2) potassic-phyllic, 3) phyllic, and 4) propylitic. The types 1, 2, and 3 were mainly developed within porphyry stock II, whereas the propylitic is present chiefly in peripheral porphyry stocks I and some dike series (e.g., DK₂ and DK₃). Contact metasomatic (skarn) occurred as patches along the contact of porphyry stock II with the Upper Cretaceous carbonates. Two distinct supergene alteration zones were recognized at Sungun, 1) oxidized and leached zone and

2) supergene sulfide zone. Hypogene mineralization occurred primarily as disseminations and as fillings in the fractures and micro-fractures. The sulfide minerals (mainly pyrite, chalcopyrite, and molybdenite) are principally concentrated in the phyllic and potassicphyllic alteration zones.

Method of Investigation

To investigate the chemical variations of the rocks associated with hydrothermal alteration and mineralization along B-B' direction in the porphyry stock II (Fig. 1), 15 samples from surface and 70 from diamond drill cores were chosen for chemical analysis (Fig. 2). The pulverized samples were chemically analyzed for 12 major and minor elements using X-ray fluorescence (XRF) technique at the department of Earth Sciences, Manchester University. Dike series were excluded in this study.

Geochemistry

Vertical Concentration Variations

The concentration variations of 12 elements (S, Cu, Mo, Rb, K, Sr, Ca, Na, Mg, Fe, Al, and Si) in samples from three DDH (B_{99} , B_7 , and B_{13}) along the B-B' direction against elevations (from sea level) were studied (Fig. 3). The distribution of these elements were affected primarily by hypogene alterations, supergene processes, and local shear and fault zones.

$DDH B_{99}$

Two thin skarn patches are present in DDH B_{99} (Fig. 3a). The sample from the skarn zone (at elevation ~1800 m) shows a sudden increase in S, Cu, Fe, Ca, and Mg, and a decrease in K, Al, and Si.

The sulfur concentration curve shows two conspicuous peaks at elevations $\sim 1800m$ ($\sim 11.2\%$) near the skarn zone and 1760m ($\sim 8.5\%$) in the phyllic zone which are due to the high sulfides (principally pyrite) content. The samples below the elevation 1760m have sulfur content within the range of 2.7% to 5.6%.

Disregarding the very low concentration (<0.1%) at elevation 1760m in the phyllic zone and the anomalous peak (~1.4%) within the elevations 1795-1815m in the skarn and supergene sulfide zones, the copper concentration in the rest of the samples in this DDH varies from ~0.25% to ~0.6%.

The overall concentration of Mo remains relatively low (<90 ppm) across the various alteration zones.

The Rb and K₂O curves typically match one another. The K₂O content in the skarn ($\sim 0.06\%$) and kaolinized rocks in the supergene sulfide zone (~ 1.5) is low, but

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Figure 1. A modified and simplified geologic map (after Mehrpartou, 1993) and a cross section along A-A' profile in the Sungun area. The dashed rectangle at the top shows the area where over 100 DDHs were dug. The filled circles on the B-B' and C-C' cross lines are the locations of the studied DDHs. The inset is an index map of Iran showing the location of Sungun PCD.



Figure 2. An E-W profile along B-B' direction in porphyry stock II showing the locations of surface and sub-surface samples from which XRF data were acquired. Shown on this figure are also the positions of the 13 diamond drill holes, various hypogene and supergene alteration zones, and hydrothermally developed stockwork-type breccia zone.

remains high (~3.5-5.5%) across phyllic and potassic-phyllic zones.

The Sr abundance is low in the supergene sulfide, skarn, and phyllic zones, and displays sudden increase in the potassic-phyllic zone.

Irrespective of the skarn zone, the Ca abundance is low (<0.2%) in the supergene sulfide and phyllic zones, and shows an increase in the potassic-phyllic zone. Its abrupt increase at elevation 1545m in the potassic-phyllic zone is due to the presence of gypsum veinlets which also caused a corresponding increase in sulfur content at this level.

The Na abundance in the skarn, supergene sulfide, and phyllic zones is low (<0.5%) with no noticeable variation. However, its concentration shows an increase in the potassic-phyllic zone.

The MgO content of the rocks in the supergene

sulfide (~9.75%) and skarn (~5.35%) zones is relatively high which is due to the high chlorite content in the former and presence of considerable amounts of Mgbearing silicates (e.g., chlorite and diopside) in the latter. Its concentration is low (<1.35%) in the phyllic zone, but shows an overall increase toward the potassicphyllic zone.

The high amounts of pyrite and chalcopyrite in the supergene sulfide zone and considerable amounts of pyrite and Fe-bearing silicates in the skarn zone account for the high Fe₂O₃ content in these alteration zones. The Fe concentration in the phyllic zone is relatively high and shows a decrease toward the potassic-phyllic zone.

Except the low concentration in the supergene sulfide and skarn zones, the Al and Si seem not to have undergone any noticeable enrichment or depletion across the hypogene alteration zones.

$DDH B_7$

This DDH is located within the intense stockworktype brecciation zone in porphyry stock II (see Fig. 2). The rocks were heavily hydro-fractured and contain high density of the veinlets and micro-veinlets.

Except the sample taken from the surface, almost all the subsurface ones from the hypogene and supergene alteration zones in this DDH (Fig. 3b) show sulfur content >2%. The sulfur curve displays two conspicuous positive anomalies at elevations ~2070 m (~6%) in the supergene sulfide zone (where the sulfide content is very high), and at 1700 m (~7.6%) in the potassicphyllic zone (where the gypsum veinlets and microveinlets are present).

The abundance of copper is relatively high in the phyllic and potassic-phyllic zones. Its curve shows two positive peaks ($\sim 2\%$) in the supergene sulfide zone (where supergene Cu-bearing sulfides such as covellite and chalcocite are present) and at the contact of potassic-phyllic and phyllic zones (where the chalcopyrite content is high).

The Mo curve features relatively two striking anomalies (>1100 ppm) in the middle of the oxidized and leached zone and at elevation \sim 2025 m in the lower part of supergene sulfide zone.

The amounts of K_2O , in general, is high (>5%) and show minor local fluctuations. Rb curve well mimics the K_2O curve.

Disregarding the effect of gypsum veinlets and micro-veinlets in the potassic-phyllic zone, the CaO curve remains almost flat across alteration assemblages.

The Na₂O content is generally low (<1%) and the majority of the samples show Na₂O <0.5%.

Except a few samples at elevations \sim 1840 m (\sim 6%), \sim 1790m (\sim 3.4%), and \sim 1720m (\sim 3%), almost all samples show the MgO content <2.5%.

The Fe₂O₃ curve shows iron enrichment in the middle of the oxidized and leached zone where considerable amounts of jarosite and minor amounts of Fe-oxides are present. However, the highest Fe₂O₃ value (6.5%) belongs to the heavily fractured rocks at elevation ~2000 m where they contain high amounts of sulfides (principally pyrite and chalcopyrite) and minor amounts of Fe-oxides (goethite).

Although displaying minor fluctuations, the Al_2O_3 content of the rocks remain within the range of 17.5-19.5%. Its concentration is also relatively high (19.5-21%) in the oxidized and leached zone, and increases up to ~25.5% in the upper part of the supergene sulfide zone where the rocks suffered intense kaolinization. The kaolinized rocks in the oxidized and leached zone contain relatively high SiO₂ (~65-70%). Except minor variations, the SiO₂ content in the supergene sulfide and

phyllic zones ranges from 62-66%, however, its concentration shows a decrease in the potassic-phyllic zone.

$DDH B_{13}$

There is no noticeable variations in concentration of the elements at the border of the phyllic and potassic-phyllic zones in DDH B_{13} (Fig. 3c). However, at the border of the potassic and the potassic-phyllic zones, the abundance of S, Cu, Sr, Na₂O, and MgO displays a shift.

The abundance of sulfur in the oxidized and leached zone is negligible (<0.1%). Its overall concentration in the phyllic, potassic-phyllic, and potassic zones is low (<2.5%), and its local anomalies are due to the realtively high sulfide (mainly pyrite and chalcopyrite) content of the samples.

The copper concentration in the supergene zone is very low (<0.1%), and gradually increases downward toward the phyllic and potassic-phyllic zones. Its curve shows two local anomalies at elevations ~1905 m (~1.4%) and ~1820 m (~1.8%) in the potassic-phyllic zone where the Cu-bearing sulfides (chiefly chalcopyrite) are relatively high. Its abundance decreases in the potassic zone.

The Mo curve exhibits a peak (670 ppm) in the lower part of the phyllic zone (at elevation around 2050 m) where the rocks were severely fractured and contain molybdenite and quartz-molybdenite micro-veinlets. Besides the peak (~340 ppm) at elevation ~1905 m, the Mo content in the rest of the samples is <130 ppm.

The K_2O content of the rocks in all supergene and hypogene alteration zones ranges from 3.7% to 5.65%. Its curve mimics very well the Rb one.

Disregarding the sample from surface outcrop, almost all samples in the supergene, phyllic, and potassic-phyllic zones show Sr content <600ppm, however, its concentration increases toward the potassic zone.

The CaO and MgO curves at elevations \sim 2290 m, \sim 2200 m, and \sim 1850 m show positive peaks which are due to the presence of calcite-dolomite content of the rocks. Apart from these peaks, their concentrations in the supergene and hypogene alteration zones remain low and almost constant. However, the MgO curve shows an overall increase toward the potassic zone.

Besides three peaks belonging to the rocks from the surface (~2.9%) and elevations ~2230 m (~3.15%) and ~1930 m (~3%), the Na₂O content of the rest of the samples in the supergene, phyllic, and potassic-phyllic zones are relatively low (<2%), however, its concentration abruptly increases (up to 4%) in the potassic zone.



Figure 3. (b) Explanation on next page.



Figure 3. Vertical concentration variations (wt%) of 12 elements across various hypogene and supergene alteration zones in three diamond drill holes along B-B' direction in porphyry stock II; a) in DDH $B_{7,}$ b) in DDH B_{13} , and c) in DDH B_{16} . The filled circles on the right side of the alteration column are the locations of the analyzed samples.

The Fe₂O₃ curve features one conspicuous positive peak (~5.5%) at elevation ~2140 m in the phyllic zone where the sulfide content (particularly pyrite) is high. The Fe₂O₃ concentration, in general, is relatively low with minor fluctuations across the alteration assemblages.

The Al_2O_3 and SiO_2 curves do not show discernible variations in various alteration zones, however, samples showing relatively low values of Al and Si, contain higher amounts of sulfides and carbonates instead.

Two-Dimensional Vertical Variations of Cu and Mo

The abundance variations of copper and molybdenum along B-B' profile are presented in the form of contoured 2-dimensional patterns.

The distribution pattern of copper (Fig. 4a) demonstrates that it is entirely depleted in the leached and oxidized zone, and shows an abrupt increase in abundance in the supergene sulfide zone. In the hypogene alteration zones, it is mainly concentrated in the phyllic and potassic-phyllic zones and its abundance decreases downward toward the center of the potassic zone.

The distribution pattern of molybdenum (Fig. 4b)

also exhibits enrichment mainly in the phyllic and potassic-phyllic zones. The density of Mo contours of equal concentration in DDH B_7 , B_6 , and B_5 clearly indicates that Mo is mainly localized in the intensely hydro-fractured zone where relatively high density of molybdenite and quartz-molybdenite micro-veinlets exists. Its abundance decreases toward the center of the potassic zone. It is depleted in the upper parts of the phyllic zone as well as in the supergene zones. However, It shows an anomalous site in the oxidized and leached zone in DDH B_7 which may be due to the presence of ferrimolybdite [Fe₂(MoO₄)₃.15H₂O].

Summary and Conclusions

The vertical distribution pattern of the 12 elements (S, Cu, Mo, Rb, Sr, K, Na, Ca, Fe, Mg, Al, and Si) in E-W profile along the B-B' direction in the porphyry stock II at Sungun showed that their abundance were highly affected by the hypogene and supergene alteration and mineralization. Due to the widespread and intensive oxidation and leaching effects, some elements such as Cu, Mo, S, Fe, K, and Si underwent re-distribution in the leached and oxidized zone.



Figure 4. Contoured 2-dimensional pattern of concentration variations for a) copper (wt%) and b) molybdenum (ppm) along B-B' direction in porphyry stock II. The locations of diamond drill holes and the pattern of hypogene and supervenes alteration zones are also shown. Contours calculated using Sigma Plot computer program with banding (X-Y) = 15 and weight = 7.

The sulfur is strongly leached out from the leached and oxidized zone, however, some (<2%) was fixed locally (e.g., in DDH B₇) as jarosite. Its local anomalies in the phyllic and potassic-phyllic zones are due to the presence of anomalous amounts of sulfides (mainly pyrite) and/or gypsum. Its overall concentration, which is intimately affiliated with sulfide content of the rocks, is greater in the east (DDH B₉₉) and middle (DDH B₇) of the porphyry stock II, and gradually decreases toward the west (DDH B₁₃).

The copper is concentrated highly in the phyllic and potassic-phyllic, and moderately in the potassic zones where Cu-bearing sulfides (mainly chalcopyrite) are present. It was almost entirely leached out from the oxidized and leached zone, but remarkably enriched in the supergene sulfide zone (particularly in the DDH B_7 , B_6 , B_5 , B_4 , B_3 , B_2 , and B_{99} .) where it was fixed as covellite and chalcocite.

The overall concentration of the Mo in the supergene zone is very low, however, the local anomaly in this zone may be due to the presence of ferrimolybdite, though it was not microscopically detected. It also shows low abundance in the potassic zone. It is mainly concentrated within the phyllic and the potassic-phyllic zones. Its abundance is relatively greater in the DDH B_7 , B_6 , and B_5 where the rocks suffered intense stockwork-type brecciation and contain high density of molybdenite and quartz-molybdenite micro-veinlets.

The K_2O is mainly enriched in the phyllic and potassic-phyllic zones and its abundance decreases toward the potassic zone. The Rb distribution pattern strikingly mimics that of the K_2O . Due to similarity of ionic radii, electronegativity, and ionization potential, Rb and K are intimately associated. Thus, Rb is preferentially incorporated in K-bearing minerals and any variation in concentration values of K will show a corresponding change in Rb values. This accounts well for the similarity of their abundance variations.

The overall abundance of CaO, Sr, MgO, and Na₂O in the phyllic, potassic-phyllic, and supergene zones are low and their abundance relatively increases toward the potassic zone.

Strontium is geochemically associated with Ca in igneous rocks, and its abundance variations are normally dependant on the Ca content of the rocks. This relationship is more pronounced in samples having relatively higher amounts of Ca-bearing silicates.

Sodium and calcium are the major components of plagioclase which was altered almost entirely (in the phyllic and potassic-phyllic zones) to moderately (In the potassic zone) to sericite. Sericitization caused extensive removal of Na and Ca from silicates and fixation of K in sericite. Weathering brought about further removal of Na and Ca from the supergene alteration zones. The local anomalies of CaO pertain to the presence of carbonates (calcite and dolomite) and sulfates (gypsum) veinlets. These veinlets caused the distribution pattern of CaO and Sr not to match one another in these localities.

The relatively low Na_2O content of the rocks (<2%) in the phyllic, potassic-phyllic, and supergene alteration zones indicates its substantial removal from Na-bearing silicates in these zones. Its abundance, however, exhibits a sharp increase in the potassic zone.

The Mg concentration in the supergene, phyllic, and potassic-phyllic zones remains low and almost constant which indicates that it was significantly removed during hypogene alterations, and the supergene processes did not play an effective role in its re-distribution. Its abundance increases toward the potassic zone where the primary and secondary biotites are present. Its local anomalous concentrations are due to the high content of Mg-bearing silicates (e.g., chlorite and/ or diopside) and/or dolomite.

The iron is relatively enriched in the phyllic zone where the amount of pyrite is relatively high, and its abundance gradually decreases downward toward the potassic-phyllic and potassic zones. Its local anomalies (Fig.3c) also pertain principally to the high pyrite content in the sample. The relatively high abundance of Fe_2O_3 (~2.5-5%) in the oxidized and leached zone (Fig. 3, b and c) is due to the presence of Fe-oxides (hematite, goethite, and limonite) and jarosite.

The Al_2O_3 and SiO_2 are relatively more abundant in the supergene zones (Fig. 3, b and c) where the rocks underwent supergene kaolinization process. Their concentrations in the hypogene alteration zones remain nearly constant. However, samples containing relatively high density of quartz veinlets and micro-veinlets display greater SiO₂ values. On the other hand, samples possessing substantial amounts of sulfides (mainly pyrite) and/or carbonates show lower amounts of Al and Si.

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