GEOCHEMICAL CHARACTERISTICS OF A LATERITE: THE JURASSIC ZAN DEPOSIT, IRAN

Ali ASGHAR CALAGARI 1) *, Fatemeh KANGARANI FARAHANI 1) and Ali ABEDINI 2)

1) Department of Earth Sciences, Faculty of Natural Sciences, University of Tabriz, 5166616471 Tabriz, Iran (Tel: +98-411-3392699; Fax: +98-41-33392703)
2) Department of Geology, Faculty of Sciences, University of Urmia, 57153165, Urmia, Iran (Tel: +98-441-2972134; Fax: +98-44-32753172)

*Corresponding author’s e-mail: calagaria@yahoo.com and calagari@tabrizu.ac.ir

ARTICLE INFO

Article history:
Received 25 October 2014
Accepted 13 December 2014
Available online 4 February 2015

Keywords:
Geochemistry
Laterite
Absolute weathering index
Parent rock
Zan
Iran

ABSTRACT

The Zan laterite deposit is located in ~25 km southeast of Damavand, Tehran province, Iran. This deposit was developed as stratiform at the base of Shemshak (sandstone and shale) Formation of Jurassic age. Mineralogical data shows that rock-forming minerals in this deposit include kaolinite, diaspore, hematite, goethite, anatase, boehmite, siderite, rutile, quartz, titanomagnetite, zircon and pyrite. The values of absolute weathering index calculated on the assumption of Th as an immobile element indicate that lateritization processes at the Zan were accompanied by enrichment of elements like Al, Fe, Zr, Nb, Ni, Cr, Co, Y, V, and REEs and by leaching-fixation of elements such as U, Mn, Sr, Ba, and Ca. Combination of the data obtained from mineralogical and geochemical investigations reveals that pH variation of weathering solutions, oxidation potential, adsorption, presence of organic matters, preferential sorption by metallic oxides, presence in resistant minerals and fixation in neomorph phases played prominent role in distribution of elements in the Zan lateritic deposit. With considering of correlation coefficients among elements, the minerals such as zircon and rutile can be regarded as potential hosts for REE in the ores. Palaeogeography evidence as well as geochemical indices like TiO2/Al2O3 and Eu/Eu* anomaly suggests that this deposit is a weathering product of basalt.

1. INTRODUCTION

Chemical weathering of the rocks is one of important surficial process that causes geochemical redistribution of elements and is controlled by various factors such as topography, parent rock type, climate conditions, plant and biological activities (Bardossy, 1982; Bogatyrev et al., 2009). Physico-chemical characteristics like Eh and pH are determinative factors for the type of behavior of elements in weathering environment (Yalcin and Ilhan, 2008, 2013; Calagari and Abedini, 2007; Abedini and Calagari, 2013a; Gu et al., 2013; Li et al., 2013; Giorgis et al., 2014; Berger et al., 2014). The main problem concerning the laterites that were not developed in alumino-silicate rocks is to find the parent rocks and to get a proper perception of their mode of formation. In many researches done so far various methods were proposed by workers to solve this problem. Some are based upon using the immobile elements like Ti, Zr, Hf, Ta, Cr, and Nb (Kurtz et al., 2000; Calagari et al., 2010; Abedini and Calagari, 2013a, b, c; Haniçli, 2013) while others apply the pattern of rare earth elements (REEs) along with their anomaly indices (e.g., Eu anomaly) (Nyakairu et al., 2001; Mameli et al., 2007; Karadağ et al., 2009; Liu et al., 2010).

Depositional cessation during Permain, Permo-Triassic, Triassic, Jurassic, and middle Cretaceous brought about development of many lateritic-bauxitic horizons in various parts of Alborz and Zagros mountain chains as well as in Central and northwestern zones of Iran. The lateritic and bauxitic horizons of many parts of Iran have been mineralogically, geochemically, and genetically investigated by various workers (Esmaeily et al., 2010; Calagari et al., 2010; Zarasvandi et al., 2010, 2012; Abedini and Calagari, 2013a, b, c; Abedini and Calagari, 2014; Abedini et al., 2014). Temporally, the lateritic and bauxitic deposits of Alborz mountain chain were widely and principally developed during Triassic and Jurassic and to lesser extent during Permo-Triassic period. The Zan laterite is one of the Central Alborz lateritic deposits that is located ~ 25 km southeast of Damavand (Tehran province), Iran. However, no investigation on geochemical characteristics of this deposit was carried out previously up to 2012. In 2012, relatively comprehensive studies on field relations, petrography, and geochemistry (mass change geochemical techniques and correlation coefficients) to consider the parental affinity and the factors affecting the distribution of major, minor, trace, and rare earth elements during lateritization processes at Zan.
2. GEOLOGY

The studied lateritic deposit is a part of the Irano-Himalayan karst bauxite belt. The study area, based upon structural divisions of Iran (Nabavi, 1976) lies in the Alborz-Azarbaidjan zone (Fig. 1a). Zaigun (silty shale), Lalun (shale and sandstone), and Mila (dolomite and limestone) Formations are the oldest sedimentary rocks, which are of Cambrian age. Ordovician-Silurian and early Devonian sediments do not crop out in this area. The Mila Formation is disconformably overlain by the Jeyrud (sandstone, shale and limestone) Formation of upper Devonian age that in turn is covered by the Mobarak (calcareous shale and carbonate) Formation. There was a depositional cessation during middle to Upper Carboniferous period (Aghanabati, 2004). Lower Permian sandstone and shales are overlain disconformably by the Elika (limestone and dolomite) Formation, which is of early Triassic age. There was depositional cessation during middle and upper Triassic and no sedimentation was occurred during this period. Erosion and weathering during this period of regression caused the development of karstic topography on the top of the Elika Formation. Basaltic
rocks cover the Elika Formation in the study area as well as in many other places in the Alborz mountain chain (Aghanabati, 2004). In early Jurassic, continental-lagoonal sediments of Shemshak (coal-bearing shale and sandstone) Formation were formed. The Zan lateritic horizon lies at the basal section of the Shemshak Formation. The lithological sequence continues with Fajan (conglomerate and sandstone) Formation of Paleocene, Karaj (tuff, marl, and shale) Formation, which is of Eocene age. These are followed by Quaternary alluviums (Amini and Khalatbari, 1972; Fig. 1b).

The Zan lateritic horizon occurs as stratiform layer trending NE-SW and extends about 1 km with variable thickness ranging from 5 to 15 meters (Fig. 1b). A selected profile shows ten lithologic layers defined by various colors. They are, from the bottom to the top, (1) brown (BL), (2) red (RL), (3) yellow (YL), (4) layered gray (LGL), (5) olive gray (OGL), (6) greenish gray (GGL), (7) reddish gray (RGL), (8) dark gray (DGL), (9) bluish gray (BGL), and (10) gray (GL) (Fig. 2). The horizon and the enclosing rocks were affected by structural forces and display fracturing and brecciation in almost all units within the horizon.

3. METHODS

Field work was carried out in spring 2012. About 30 samples from various units across lateritic horizon and the enclosing rocks were taken. After mesoscopic examination, six samples of lateritic horizon were selected to be analyzed by X-ray diffraction (XRD) (SIEMENS Diffractometer, model D-5000, CuKα radiation, fixed graphite chromators, voltage 40 kV, current 40 mA, scanning speed 2° per minute, scan range 2-70°) for identification of unknown mineral phases at Geological Survey of Iran. For identification of clay minerals, the samples were treated with KCl (at 550 °C), MgCl₂, and ethylene glycol. Complementary investigations on mineralogy of the ores were carried out by scanning electron microscopy (SEM; a XLC Philips at Amirkabir University in Tehran). For determination of chemical composition, ten representative samples taken from the ores across the studied profile were selected and chemically analyzed at ACME laboratories in Canada (analytic code 4A4B) by using ICP-ES (for major and minor oxides) and ICP-MS (for trace and rare earth elements) methods. Loss on ignition (LOI) values were measured by weighing the samples (3 to 4 grams each) before and after five minutes of heating at 1040-1060 °C. Detection limit for major and minor oxides is 0.01 wt% (except for Fe₂O₃, Cr₂O₃, and LOI which are 0.04, 0.002, and 0.1 wt %, respectively). Detection limit for trace elements is 0.1 ppm and for rare earth elements 0.05 ppm (except for Ba, Ga, Sr, Pr, Nd, Eu, Ho, Er, and Tm which are 1, 0.5, 0.5, 0.02, 0.3, 0.02, 0.01, 0.02, 0.03, and 0.01, respectively). For determination of REE values, about 0.2 gram of each sample is treated with metaborate and lithium tetraborate and then is dissolved in HNO₃ acid. The results of the chemical analyses are presented in Table 1.

4. RESULTS AND DISCUSSION

4.1. MINERALOGY AND CHEMICAL COMPOSITION

X-ray diffraction (XRD) analyses show that the Zan laterites consist of kaolinite, diaspore, hematite, goethite, boehmite, anatase, siderite, rutile, and quartz. However, scanning electron microscopy (SEM) examinations of ore samples from the bluish gray (BGL) and the layered gray (LGL) units revealed that in addition to the mentioned mineral phases (Fig. 3a), there also exist crystalline particles of titanomagnetite (Figs. 3b, c, d), pyrite (within goethite) (Fig. 3e), and zircon (Fig. 3f) in this deposit.

4.2. INDUSTRIAL APPLICATIONS

The Al₂O₃/SiO₂ ratio of lateritic-bauxitic ores (so-called module) plays a crucial role in determining their industrial applications (Yalcin and Ilhan, 2013). The module values of Zan laterites have a range of
Table 1 Results of chemical analyses (ICP-ES and ICP-MS) for major, minor, trace and rare earth elements along with Eu anomaly in the ores of the Zan latectic deposit. The order of each number on samples is from the bottom to the top of the horizon.

<table>
<thead>
<tr>
<th>Sample</th>
<th>GL (Z-1)</th>
<th>BGL (Z-2)</th>
<th>DGL (Z-3)</th>
<th>RGL (Z-4)</th>
<th>GGL (Z-5)</th>
<th>OGL (Z-6)</th>
<th>LGL (Z-7)</th>
<th>YL (Z-8)</th>
<th>RL (Z-9)</th>
<th>BL (Z-10)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SiO2 (WT%)</td>
<td>40.34</td>
<td>31.98</td>
<td>23.04</td>
<td>33.20</td>
<td>26.22</td>
<td>30.61</td>
<td>19.12</td>
<td>19.37</td>
<td>25.32</td>
<td>27.03</td>
</tr>
<tr>
<td>Fe2O3</td>
<td>5.09</td>
<td>16.20</td>
<td>12.49</td>
<td>16.43</td>
<td>24.59</td>
<td>20.32</td>
<td>35.19</td>
<td>43.77</td>
<td>35.78</td>
<td>24.07</td>
</tr>
<tr>
<td>MgO</td>
<td>0.59</td>
<td>0.79</td>
<td>0.47</td>
<td>0.63</td>
<td>0.68</td>
<td>0.59</td>
<td>1.40</td>
<td>0.43</td>
<td>0.58</td>
<td>0.65</td>
</tr>
<tr>
<td>CaO</td>
<td>0.46</td>
<td>0.40</td>
<td>11.68</td>
<td>0.32</td>
<td>0.35</td>
<td>0.35</td>
<td>0.65</td>
<td>0.25</td>
<td>0.34</td>
<td>0.86</td>
</tr>
<tr>
<td>Na2O</td>
<td>0.15</td>
<td>0.12</td>
<td>0.12</td>
<td>0.11</td>
<td>0.09</td>
<td>0.12</td>
<td>0.18</td>
<td>0.08</td>
<td>0.14</td>
<td>0.08</td>
</tr>
<tr>
<td>K2O</td>
<td>0.24</td>
<td>0.05</td>
<td>11.68</td>
<td>0.32</td>
<td>0.35</td>
<td>0.35</td>
<td>0.65</td>
<td>0.25</td>
<td>0.34</td>
<td>0.86</td>
</tr>
<tr>
<td>TiO2</td>
<td>3.01</td>
<td>4.79</td>
<td>4.86</td>
<td>4.21</td>
<td>3.51</td>
<td>4.76</td>
<td>3.31</td>
<td>3.35</td>
<td>3.00</td>
<td>3.55</td>
</tr>
<tr>
<td>P2O5</td>
<td>0.42</td>
<td>0.14</td>
<td>10.13</td>
<td>0.13</td>
<td>0.17</td>
<td>0.27</td>
<td>0.10</td>
<td>0.27</td>
<td>0.09</td>
<td>1.33</td>
</tr>
<tr>
<td>MnO</td>
<td>0.02</td>
<td>0.11</td>
<td>0.09</td>
<td>0.07</td>
<td>0.55</td>
<td>0.05</td>
<td>0.04</td>
<td>0.08</td>
<td>0.06</td>
<td>0.10</td>
</tr>
<tr>
<td>Cr2O3</td>
<td>0.01</td>
<td>0.01</td>
<td>0.01</td>
<td>0.01</td>
<td>0.01</td>
<td>0.01</td>
<td>0.01</td>
<td>0.01</td>
<td>0.02</td>
<td>0.02</td>
</tr>
<tr>
<td>LOI</td>
<td>17.60</td>
<td>19.10</td>
<td>14.60</td>
<td>17.00</td>
<td>21.70</td>
<td>16.90</td>
<td>23.30</td>
<td>15.20</td>
<td>14.10</td>
<td>17.70</td>
</tr>
</tbody>
</table>

0.79 to 0.94. Based on the classification of Voros (1979), the ores at the Zan in light of aluminum extraction are considered as low quality. However, comparison of the mean values of major constituent oxides of the studied ores with the ideal chemical composition of the lateritic-bauxitic ores for industrial applications (Edwards and Atkinson, 1986; Manning, 1995) shows that the Fe-poor and Fe-rich parts of the horizon at Zan have great potential for uses (as raw materials) in refractory and cement industries, respectively.
Fig. 3 Scanning electron microscopic microphotographs of the Zan lateritic ores. (a) Tabular fragment of diaspore along with Fe-oxide and – hydroxide in the ore of GL unit. (b) The presence of titanomagnetite crystal in LGL unit. (c) Concentration of iron in titanomagnetite. (d) Concentration of titanium in titanomagnetite. (e) Very fine-grained pyrites within goethite in BGL unit. (f) A solitary crystal of zircon in unit BGL. Abbreviations: Zr = zircon, Dsp = Diaspore, Ti-Mag = titanomagnetite, Gt = goethite, and Py = pyrite

4.3. MASS CHANGE CALCULATIONS

Customarily, calculations of mass change for elements during weathering are carried out by employing two methods, (1) iso-volume (Brimhall and Dietrich, 1987; Brimhall et al., 1991) and (2) immobile element (Nesbitt, 1979; MacLean and Kranidiotis, 1987; MacLean, 1990; MacLean and Barrett, 1993). In this study, for determination of degree of mobility and enrichment of elements during lateritization processes, absolute weathering index (AWI) of immobile element (Nesbitt, 1979) was used. The calculations were done with an assumption of Th as immobile element (Wimpenny et al., 2007) and of taking the mean composition of upper continental crust (UCC) (Taylor and McLennan, 1985) as parent rock for the Zan deposit by using the following equation:

$$%\text{AWI} = \left[\frac{\text{Element}_{\text{Laterite}}}{\text{Th}_{\text{Laterite}}} \right] \left/ \left[ \frac{\text{Element}_{\text{UCC}}}{\text{Th}_{\text{UCC}}} \right] \right. \times 100$$

This investigation was carried out for five groups, (1) major and minor elements, (2) large ion lithophile elements (LILE), (3) transition trace elements (TTE), (4) high field strength elements (HFSE), and (5) rare earth elements (REEs).

Major and minor elements (Al, Fe, Ti, Si, Na, K, Ca, Mg, Mn, and P): The mass change calculations show that silica relative to alkali and alkali earth elements suffered moderate leaching (Figs. 4a, b). This is likely owing to formation of secondary clay minerals like kaolinite during lateritization. Enrichment of Al and Ti in the profile (Fig. 4a) could be due to the coupled alteration of silicates minerals (feldspars and ferromagnesians) and formation of Al
Fig. 4 Patterns of mass changes for major and minor oxides (a, b) and large ion lithophile elements (c) in various units of the Zan studied profile.

and Ti-bearing minerals. Non-rhythmic distribution of Fe in the profile and its increase toward bedrocks may be explicative of fluctuation of underground water table level. It seems that decomposition of organic matters in the upper parts of the profile provided locally acidic and reducing conditions which resulted in movement of iron downwardly and its subsequent fixation in the lower parts of the profile. The mass change calculations reveal that lateritization processes at Zan were accompanied by moderate to strong leaching of Ca, Na, K, and Mg and to a lesser extent by Mn (Fig. 4b). It seems that Na, Ca, and K were released into weathering solutions. On the other hand, depletion of Mg and Mn from the system is related to decomposition of ferromagnesian minerals. Enrichment of Mn, however, in the GL-unit exhibits that Mn$^{2+}$ was oxidized to Mn$^{4+}$ and was fixed as insoluble oxides and hydroxides (Koppi et al., 1996). Enrichment of CaO in the DGL-unit is due to deposition of CaCO$_3$ by supergene solutions in this part of the profile.

Large ion lithophile elements (Rb, Ba, Sr, and U): Depletion or enrichment of trace elements in the lateritic profiles depends on their host minerals in the parent rocks. Mobile trace elements, if their host minerals are insoluble, can become enriched in laterites (Meshram and Randive, 2011). Rb was released during feldspar weathering (break-down) causing its depletion in all of the lateritic units (Fig. 4c). The mass changes of Ba, Sr and U show their enrichment in some units (Fig. 4c). The positive correlation between pairs of U-Y ($r = 0.85$) and U - P$_2$O$_5$ ($r = 0.7$) could be due to the presence of U in the crystal structure of Y-rich minerals like xenotime (YPO$_4$) (Gonzales–Lopez et al., 2005) that was not identified by XRD analysis as the result of its low concentration. The positive correlation between pairs of Sr-P$_2$O$_5$ ($r = 0.82$) and Sr-Ca ($r = 0.73$) may be suggestive of likely hosting of crandallite mineral group for Sr. These minerals are commonly formed by alteration of apatite in karst bauxites (Bardossy, 1982). The moderate to strong positive correlation between Ba and P$_2$O$_5$ ($r = 0.83$) and between Ba and REEs ($r > 0.7$) may provide reason for the presence of the Al-bearing secondary phosphates like gorceixite [BaAl$_2$(LREE,PO$_4$)$_2$(OH)$_5$H$_2$O] (Costa and Araujo, 1996) that was not detected by XRD analysis.

Transition trace elements (Ni, Co, Cr, and V): The variation of AWI values for Co, Cr, Ni, and V at Zan was irregular but incremental (Fig. 5a). Iron colloids have effective role in transportation and concentration of a group of trace elements, especially transitional trace elements, during weathering processes (Pokrovsky et al., 2006). Considering the moderate and positive correlation between pairs of Fe-Co ($r = 0.75$), Fe-V ($r = 0.69$), and Fe-Cr ($r = 0.72$), the role of Fe-oxides and -hydroxides in fixation and concentration of these elements (as the result of scavenging and adsorption) can be established. Among the transition trace elements, Ni has very weak correlation with Fe but moderate and positive correlation with SiO$_2$ ($r = 0.68$). This may be indicative of being clay minerals as potential fixing agent for Ni in the studied profile.

High field strength elements (Y, Nb, Zr, and Hf): These elements were enriched in all of the lateritic units (Fig. 5b). It seems that the enrichment of these almost immobile elements took place in an authigenic
Rare earth elements: The pattern of mass changes of LREEs and HREEs in the Zan lateritic deposit shows an incremental trend (Figs. 6a, b). The remarkable mass increase of REEs in the DGL unit is likely in connection with pH increase and formation of REE-bearing carbonate minerals. The good and positive correlation between lanthanides and Zr ($r > 0.74$) and between lanthanides and Ti ($r > 0.72$)
reveals that titanium oxides and zircon are the major hosts for REEs. LREEs have weak but positive correlation with Mn ($r > 0.35$), Al ($0.25 < r < 0.4$), and Si ($0.1 < r < 0.4$) indicating the weak role of Mn-oxides and clay minerals in hosting LREEs.

4.4. PARENT ROCK COMPOSITION

Commonly lateritic-bauxitic deposits have compositions and textures that are related to underlying host rocks (Bardossy and Aleva, 1990). However, this can hardly be observed for bauxites developed on limestone. A wide range of parent rocks for karst bauxites have been proposed including argillaceous materials (MacLean et al., 1997), mafic igneous rocks (Calagari and Abedini, 2007; Esmaeily et al., 2010); volcanic ashes (MacLean et al., 1997), and wind-drifted materials (Brimhall et al., 1988; Pyk, 1988).

Eu anomaly remains commonly constant during weathering processes (Mongelli, 1993; Mameli et al., 2007; Abedini and Calagari, 2013a). Al and Ti also remain immobile during lateritization (MacLean et al., 1997; Mameli et al., 2007) and have a great potential to get into sediments. Therefore, TiO$_2$/Al$_2$O$_3$ and Eu anomaly are sensitive index for determination of parent rocks. The mean composition values of upper continental crust (UCC), basalt, granite (Taylor and McLennan, 1985), and various units of the Zan lateritic ores are illustrated on Eu anomaly versus TiO$_2$/Al$_2$O$_3$ plot (Fig. 7a). According to Figure 7a, the Zan data plots in the basalt (as precursor) domain. Further investigations revealed that the Zr/TiO$_2$ is commonly similar to that of parent rocks and can provide favorable geochemical information for determination of parent rocks (Hallberg, 1984). On the basis of this premise, the position of the Zan data on a TiO$_2$ versus Zr bivariate plot also suggests a basaltic composition for the parent rocks (Fig. 7b). Floyd and Winchester (1978) used the bivariate plot of Nb/Y versus Zr/TiO$_2$ for determination of parent rocks of lateritic deposits (this method, especially when the parent rocks are extrusive igneous rocks, is very applicable). The Zr/Ti of the parent rock remains constant during
lateritization processes (Hill et al., 2000). The Zan data on Nb/Y versus Zr/TiO₂ plot reflects the alkali basalt composition for the precursor of the horizon (Fig. 7c).

Interpretation of a basic source material for Zan laterite deposit is supported by palaeogeography evidence of Iran. In the Central Alborz structural zone, the early Jurassic andesite to basaltic volcanic rocks cover an eroded and often karstified surface, at the base of the Shemshak Formation (Annelles et al., 1975; Nabavi and Seyed-Emami, 1977; Nabavi, 1987; Esmaeily et al., 2010). The volcanic activity occasionally continued into the lower part of the Shemshak Formation.

5. CONCLUSIONS

Calculations of the degree of absolute weathering index (AWI) demonstrate that lateritization of the Zan horizon was resulted by desilification and removal of alkali and alkaline-earth elements coupled with Al and Fe enrichment. Despite remaining in the residual system, Fe shows strong variation within the horizon. Based upon correlation coefficients among various elements, it can be deduced that the distribution of trace and rare earth elements in various unit within this deposit was controlled essentially by neomorph accessory and resistant mineral phases. In general, distribution of REEs, Hf, and Nb in the lateritic ores was controlled substantially by minerals like Ti-oxides and zircon. Fe-oxides and -hydroxides host elements such as Co, V, and Cr. Clay minerals are the chief controlling agents for Ni distribution in this deposit. Distribution of Y was mainly controlled by Al-bearing mineral phases. The phosphate minerals played crucial roles for concentration of Ba, U, and Sr during lateritization processes. Palaeogeography evidence as well as geochemical indices such as TiO₂/Al₂O₃ and values of Eu anomaly reveals that the Zan lateritic horizon is a weathering product of basalt.

ACKNOWLEDGEMENTS

This work was enjoyed financial support by the Research Deputy Bureau of the University of Tabriz. The authors would like to express their thanks and appreciations to the authorities of this bureau. Special thanks are also expressed to Prof. Josef Stemberk for his editorial assistance and to two anonymous reviewers for making critical comments on this manuscript.

REFERENCES


MacLean, W.H. and Kramidiotis, P.: 1987, Immobile elements as monitors of mass transfer in hydrothermal alteration: phelps Dodge massive sulfide deposit, Matagami, Quebec. Economic Geology, 82, No. 4, 951–962. DOI: 10.2113/geosecsop.82.4.951


Pokrovsky, O.S., Schott, J. and Dupre, B.: 2006, Trace element fractionation and transport in boreal rivers


