Geochemistry of Carbon, Oxygen and Sulfur Isotopes in Soils along a Climotoposequence in Kerman Province, Central Iran

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ABSTRACT

Carbon and sulfur isotopic data are helpful in understanding environmental conditions under which soils are formed. Soils from 18 sampling sites along a climotoposequence from higher Laleh Zar Mountains (4,351 m above sea level with xeric-mesic soil moisture-temperature regimes) to Shahdad depression (250 m above sea level with arid-hyperthermic soil moisture-temperature regimes) in Kerman Province, central Iran, were studied. δ13C values of the soil organic carbon varied between -20.6‰ and -26.9‰ PDB. δ13C values of pedogenic carbonates varied between -2.2‰ and +2.2‰ PDB and a decreasing trend was observed downward in the climotoposequence. A positive correlation (r= 0.6) was found between δ13C and δ18O values in the lower arid landscape, but a different relationship (r= 0.37) in the upper Mediterranean climatic positions was observed. It seems that the presence of more C3 plants under stress and more decomposition of organic matter in lower positions of the landscape has taken place. δ13C values of soil organic carbon and pedogenic carbonates and their trend along the gradient showed that higher proportion of C3 plants at lower elevations and presence of more C4 plants in the past history of upper landscape positions may have been experienced. Mean δ34S value of 11.3‰ showed that lower Cretaceous sea sulfate may have been the source of sulfur in the area. Mean δD and δ18O values of -51.2‰ and +6.0‰ in gypsum crystallization water showed that direct deposition from a solution saturated with gypsum is the only probable mechanism of gypsum formation in the area.

Keywords: Gypsum crystallization water, Pedogenic carbonate, Soil organic carbon, Stable isotope geochemistry.

INTRODUCTION

Information on the amount and accumulation rate of pedogenic carbonate in soils helps estimation of carbon pools and fluxes and past climatic conditions in terrestrial environments, which also reflects the influence of paleoclimate in a given area (Landi et al., 2003). Carbon isotope ratios in both organic and inorganic soil materials could help to establish the kind of processes and the environmental conditions involved in soil formation (Boutton, 1996; Cerling and Hay, 1986). Carbon isotopic data were also used to study the mechanisms of carbon dissolution and precipitation (Mermut et al., 2000; Nordt et al., 1996) and to understand the carbon cycle (Salehi et al., 2004).

There is a substantial difference in 13C discrimination in the photosynthetic pathways of C3 vs. C4 plants with mean δ
$^{13}$C values of -27 and -12 $\%e$, respectively (Cerling and Quade, 1993; O’Leary, 1988). In the absence of C3-C4 vegetation shifts, other ecological factors affect carbon isotope values in the soil (Stevenson et al., 2005).

Mode of pedogenic carbonate formation studied by Khademi and Mermut (1999) in Isfahan area of central Iran, with an arid soil moisture regime, showed that carbonates were formed in an environment with more available moisture and higher proportion of C4 plants than observed today. In a similar study in the west central Iran, with a xeric soil moisture regime, Salehi et al. (2004) found that current vegetation consists of C3 plants, but soil pedogenic carbonate was formed in conditions with a lower proportion of C3 plant species and the shift of C3-C4 vegetation was attributed to human induced land degradation. The stable carbon isotope geochemistry of organic carbon and pedogenic carbonate in soils along a climosequence in Saskatchewan soils of Canada (Landi et al., 2003) and soils of Washington state, USA (Stevenson et al., 2005) showed a good relation between $\delta^{13}$C values and mean annual precipitation.

Gypsiferous soils cover about two hundred million hectares in the world (Nettelton, 1991). Origin of gypsum and more soluble salts in these types of soils has been studied by several researchers. Sofer (1978) described three important geochemical processes of gypsum formation including oxidation of sulfide minerals, hydration of anhydrite, and direct deposition from an evaporating solution saturated with gypsum. Pyrite oxidation has been reported as the major source of gypsum by Mermut and Arshad (1987) and Dowuona et al. (1992a, b). Khademi et al. (1997a) found that lower Cretaceous sea sediment is the source of sulfate in Isfahan region, central Iran. Farpoor et al. (2004) suggested that in southeastern Iranian soils, gypsum was formed through direct deposition of the Thetian sea lagoons, similar to gypsum formation in central Iran. Using the stable isotope geochemistry of sulfur bearing minerals in soils and sediments from different geomorphic surfaces in Loot Desert of central Iran, Farpoor and Krouse (2008) attempted to explain the evolution of the geomorphic surfaces in that area. Genesis of gypsum enriched soils in north-west Isfahan, central Iran was also studied by Toomanian et al. (2001).

The objectives of the present research were to: (1) determine the carbon isotope geochemistry of pedogenic carbonates and soil organic matter along a climotoposequence from Laleh Zar elevations to Shahdad depression, (2) study the S and O isotope geochemistry in the area, (3) investigate the $\delta^{18}$O and $\delta D$ values of gypsum crystallization water, and (4) establish the mode of gypsum and carbonate formation in the area.

**MATERIALS AND METHODS**

**Study Area and Sampling**

The study site is located in the Kerman Province, southeastern part of central Iran (Figure 1). The climotoposequence under study extends from Laleh Zar with an elevation of 4351 m asl (above sea level) in the south west to the Shahdad depression, 250 m asl in the north-east (Figure 2).

Mean annual precipitation in the area ranges between 250 mm (in the south-west) and 50 mm (in the north-east). Soil moisture and temperature regimes change from xeric and mesic in the Laleh Zar Mountain to aridic and hyperthermic in the Shahdad area near the Loot playa.

Geological formations are dominated by igneous formations and volcanic tuffs in the south-west to limestone and Neogene Formations in the north-east of the transect. Soil samples, secondary carbonates, and gypsum crystals were collected from 18 sites along the transect (Figure 2).

Due to the extent of the area and soil variations along the studied transect, sampling for organic carbon, pedogenic carbonates, soluble sulfates, and crystallization water of gypsum analyses...
was performed in appropriate sites. Site 1 was located at higher elevations of the transect. Histosols were dominant soils in this site (Table 1). Moving down the slope, soil moisture decreased, so that in site 6, xeric moisture regime shifted to aridic. Organic matter content showed also a decreasing trend toward downslope positions (about 0 %). That is why organic carbon samples were only collected from sites 1-9 on the transect.

On the other hand, C and O isotopes of pedogenic carbonates were measured in the soft mass and carbonate nodules were sampled only from sites containing Bk horizons. Moving down the slope toward site 18 (near Loot Desert), the amount of soluble sulfates increased, but S and O isotopes determination were performed on soils of all 18 sites studied. Besides, four pure gypsum samples from sites 8, 11, and 12 were also collected for gypsum crystallization water analyses and no pure gypsum crystals were found in other sites studied.

**Laboratory Studies**

For inorganic carbon isotopic composition analysis, a sample containing approximately 10 mg of pedogenic carbonate was placed in the main portion of the Y-shaped reaction vessel, keeping the sample off the side walls and out of the side arm. An extra amount of 100% phosphoric acid (about 4 mL) was placed in the side arm. The vacuum system was used to evacuate the reaction vessel. The phosphoric acid was tipped into the sample chamber after evacuation and left overnight for the reaction with soil carbonates (Boutton, 1991; Khademi and
Mermut, 1999). Using the vacuum system on the next day, the liberated CO$_2$ gas was purified and utilized for analysis. The CO$_2$ gas derived from the above mentioned procedure can be used for both $\delta^{18}$O and $\delta^{13}$C measurements of the samples, because the oxygen in the CO$_2$ generated from carbonate acidification is derived from carbonates.

Soil organic carbon samples were also analyzed for carbon isotopic composition. Samples were soaked in an excess amount of 3M HCl at room temperature overnight to remove the carbonates (conversion of...
carbonate to CO$_2$). Using a millipore filter and deionized water, the excess acid was then removed from the soil sample. The carbonate-free soil samples were dried at 110°C after leaching and used for stable isotope analysis of organic carbon (Khademi and Mermut, 1999).

Dry combustion procedure described by Boutton (1991) in an excess of oxygen was used to convert organic carbon of the samples to CO$_2$ for isotopic analysis. This procedure involved the combustion of samples in sealed, evacuated vycor tubes containing CuO as the source of oxygen at 850°C for 3 hours. Cryogenic distillation was used to separate the CO$_2$ from other combustion products. The separated CO$_2$ was collected for δ$^{13}$C analysis.

All forms of sulfates (soluble and mineral forms) were converted to pure BaSO$_4$ for δ$^{34}$S and δ$^{18}$O analysis according to the method described by Dowouna et al. (1992a).

Gypsum crystallization water was extracted using the method described by Halas and Krouse (1982) and Gonfiantini and Fontes (1963). The CO$_2$-H$_2$O isotopic equilibration method of Epstein and Mayeda (1953) with the modification of Roether (1970) was used for O isotopic analysis. For H/D analyses, H$_2$ was prepared by reduction of H$_2$O with Zn as described by Coleman et al. (1982).

The isotopic composition of gypsum mother water was calculated using the following equation:

$$\alpha = (\delta_{wc} + 1000)/(\delta_{mw} + 1000)$$

Where $\delta_{wc}$ is the isotopic composition (δ$^{18}$O or δD) of crystallization water and $\delta_{mw}$ is the isotopic composition for gypsum mother water. According to Gonfiantini and Fontes (1963), the $\delta^{18}$O is 1.004 and according to Pradhananga and Matsuo (1985), the $\alpha D$ is 0.98.

## RESULTS AND DISCUSSION

### Stable Isotope Composition of Organic and Inorganic Carbon

**Organic Carbon**

δ$^{13}$C values of the soil organic carbon varied between -20.6‰ (site 7) and -26.9‰ (site 1) PDB (Table 2). In the climotoposequence studied, the δ$^{13}$C values showed an increase with decreasing elevation and precipitation (Figures 1 and 2). The δ$^{13}$C values of C$_3$ and C$_4$ plants are reported to be between -25‰ and -32‰ and -10‰ and -14‰, respectively (Cerling and Quade, 1993). Under soil moisture stress conditions, the isotopic composition of C$_3$ plants are several per mil enriched in $^{13}$C (Stevenson et al., 2005; Salehi et al., 2004; Delucia et al., 1988; Ehleringer et al., 1986). The δ$^{13}$C values of -20‰ have been reported for C$_3$ plants growing in arid environments where stomatal conductance is limited (Ehleringer et al., 1991). This could be one plausible reason for increasing δ$^{13}$C values toward arid parts of the transect studied. Thus, the carbon isotopic

<table>
<thead>
<tr>
<th>Site</th>
<th>Horizon/layer</th>
<th>Depth (cm)</th>
<th>Organic carbon δ$^{13}$C (‰), PDB</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Oe1</td>
<td>0-60</td>
<td>-25.9</td>
</tr>
<tr>
<td>1</td>
<td>Oe2</td>
<td>60-90</td>
<td>-26.3</td>
</tr>
<tr>
<td>1</td>
<td>Bwg1</td>
<td>90-120</td>
<td>-26.9</td>
</tr>
<tr>
<td>1</td>
<td>Bwg2</td>
<td>120-150</td>
<td>-26.9</td>
</tr>
<tr>
<td>2</td>
<td>A</td>
<td>0-60</td>
<td>-26.7</td>
</tr>
<tr>
<td>3</td>
<td>Bkz</td>
<td>40-70</td>
<td>-24.0</td>
</tr>
<tr>
<td>4</td>
<td>Bkz</td>
<td>5-30</td>
<td>-22.0</td>
</tr>
<tr>
<td>5</td>
<td>Bkz</td>
<td>70-100</td>
<td>-21.1</td>
</tr>
<tr>
<td>7</td>
<td>Bkt1</td>
<td>50-90</td>
<td>-20.6</td>
</tr>
<tr>
<td>9</td>
<td>Bk1</td>
<td>30-50</td>
<td>-23.9</td>
</tr>
</tbody>
</table>

Table 2. Isotopic composition of soil organic matter in selected soils from area studied.
composition of soil organic carbon in the area studied seems to be predominantly derived from C3 plants. Stevenson et al. (2005) have also found a similar trend in \(\delta^{13}C\) values of soil organic carbon in a gradient from 200 to 500 mm precipitation.

Decomposition processes and discrimination against \(^{13}C\) and preferential preservation of organic matter rich in \(^{13}C\) would also result in a slight fractionation process (Monger et al., 1998; Stevenson et al., 2005). This could be a second possible reason for increased trend of \(\delta^{13}C\) in soil organic carbon and could be attributed to the above mentioned fractionation process. Higher temperature and lower precipitation in the arid parts of the transect can cause a faster organic matter decomposition and hence enrichment in \(\delta^{13}C\).

**Inorganic Carbon**

The \(\delta^{13}C\) and \(\delta^{18}O\) values of pedogenic carbonates formed as soft masses and nodules (Table 3) showed a correlation with an \(r\) value of 0.40 which was insignificant, but the \(r\) value changed from 0.37 in the elevated and humid-arid parts of the transect (sites 3-7) to 0.6 in the really arid positions (sites 9-14). The positive correlation between \(\delta^{13}C\) and \(\delta^{18}O\) in pedogenic carbonates is already well established (Salomons and Mook, 1976). Because both isotopes are enriched in the remaining \(\text{HCO}_3^-\), during precipitation in such a system, \(\delta^{13}C\) and \(\delta^{18}O\) are, therefore, correlated. Schlesinger et al. (1989) found that the positive correlation of \(\delta^{13}C\) and \(\delta^{18}O\) could also be related to seasonal variations in biotic processes that affect the carbonate deposition. Khademi and Mermut (1999) also found a positive correlation with an \(r\) value of 0.7 between \(\delta^{13}C\) and \(\delta^{18}O\) in pedogenic carbonates in Isfahan region of Iran, which is almost similar to the area under study. However, the \(r\) value in the more humid parts of the transect with a Mediterranean climate is as low as 0.4. Similarly, the soil pedogenic carbonates in Mediterranean climate of Australia, Greece and Turkey, did not show a correlation between \(\delta^{13}C\) and \(\delta^{18}O\) in pedogenic carbonates (Cerling and Quade, 1993).

The \(\delta^{13}C\) values of pedogenic carbonates decreased downward on the transect (Table 3). One reason could be the higher proportion of C3 plants at lower altitudes, due to a change from mesic to aridic conditions. Most desert scrub species are among C3 plants (Volkoff and Cerri, 1987). It is also possible that the pedogenic carbonates at higher elevations may have formed under a different paleoenvironment. Mean \(\delta^{13}C\) values of pedogenic carbonates in the upslope positions (sites 1-7) of the area is much higher (+0.45‰), than the mean value reported by Salehi et al. (2004) in Chaharmahal Bakhtiari Province of Iran, with a Mediterranean climate (-5.02‰). In

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**Table 3.** Carbon and oxygen isotopic composition of pedogenic carbonates of some selected soils.

<table>
<thead>
<tr>
<th>Site</th>
<th>Horizon</th>
<th>Depth(cm)</th>
<th>(\delta^{13}C)(PDB)</th>
<th>(\delta^{18}O)(SMOW)</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>Bkz</td>
<td>40-70</td>
<td>-2.2</td>
<td>+28.36</td>
</tr>
<tr>
<td>4</td>
<td>Bkz</td>
<td>5-30</td>
<td>+1.5</td>
<td>+32.56</td>
</tr>
<tr>
<td>5</td>
<td>Bktn</td>
<td>65-100</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>6</td>
<td>Bktn</td>
<td>70-100</td>
<td>+2.2</td>
<td>+28.96</td>
</tr>
<tr>
<td>7</td>
<td>Bktn</td>
<td>50-90</td>
<td>+0.3</td>
<td>+32.56</td>
</tr>
<tr>
<td>8</td>
<td>C</td>
<td>140-160</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>9</td>
<td>Bk1</td>
<td>30-50</td>
<td>+1.3</td>
<td>+36.86</td>
</tr>
<tr>
<td>10</td>
<td>C</td>
<td>85-110</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>11</td>
<td>Bk1</td>
<td>30-70</td>
<td>-1.4</td>
<td>+32.76</td>
</tr>
<tr>
<td>12</td>
<td>Bk2</td>
<td>70-90</td>
<td>-1.7</td>
<td>+32.56</td>
</tr>
<tr>
<td>13</td>
<td>Bk</td>
<td>40-80</td>
<td>-1.6</td>
<td>+26.86</td>
</tr>
<tr>
<td>14</td>
<td>Bk</td>
<td>30-70</td>
<td>-1.1</td>
<td>+25.96</td>
</tr>
</tbody>
</table>

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the Isfahan region of Iran, with an arid climate, Khademi and Mermut (1999) reported a mean $\delta^{13}C$ value of -0.25‰. It seems, therefore, that higher $\delta^{13}C$ value of pedogenic carbonates in the study area is due to higher proportion of C4 plants in the past.

As reported by Romanek et al. (1992), the $\delta^{13}C$ value of CaCO$_3$ is about 10% greater than that of the CO$_2$ in the soil atmosphere at 20°C. Adding the least amount of 4.4 ‰ diffusional enrichment to the above mentioned value, Cerling and Quade (1993) concluded that the total enrichment between the soil respired CO$_2$ and pedogenic carbonate is about 14.4‰ at 20°C. Khademi and Mermut (1999) showed that the $\delta^{13}C$ values of pedogenic carbonates in Isfahan region, central Iran, ranged between -2.1‰ and +0.9‰. With the 14.4‰ enrichment discussed above, they concluded that the respired CO$_2$ from which the pedogenic carbonate formed, had $\delta^{13}C$ values ranging between -16.5‰ and -13.5‰, which is much higher than the $\delta^{13}C$ values of soil organic carbon in that studied area. A great proportion of more C4 plants in the history of the area, and a great contribution of a parent material with an isotopically heavier carbon are two possible reasons mentioned by Khademi and Mermut (1999) for the higher $\delta^{13}C$ values of pedogenic carbonates in the Isfahan region, although the former reason is much more emphasized.

Considering the range of $\delta^{13}C$ values of -2.2‰ to +2.2‰ for the pedogenic carbonate of this research area (Table 3), it is calculated that the respired CO$_2$ has had the $\delta^{13}C$ values ranging between -16.6‰ and -12.2‰, which are much higher than the $\delta^{13}C$ values of soil organic carbon (Table 2). This is even higher than the value reported in the Isfahan region. It is concluded that much of the pedogenic carbonates in this study area were precipitated in a much drier period. The $\delta^{13}C$ values of pedogenic carbonates are 21.9‰, heavier than the mean $\delta^{13}C$ value (-24.1‰) of soil organic carbon in the area.

### Sulfur Isotope Geochemistry

Figure 2 shows the sampling sites for sulfur geochemistry studies. The summary of isotopic composition of sulfur and oxygen is given in Table 4. Although soil samples from all sites were used for sulfur isotope investigations, no gypsum or other soluble sulfates was found at the higher elevations of the transect (sites 1-5). Samples with gypsum taken from the lower slope position and depression (sites 6-18) had $\delta^{34}S$ values ranging between 7.7‰ and 12.6‰ (with a mean value of 11.3 ‰) and $\delta^{18}O$ values ranging between 7.2‰ and 17.8‰ (with a mean value of 11.3‰).

No trend in sulfur and oxygen isotope data regarding the change in elevation could be observed. The $\delta^{34}S$ values of samples (Table 4) are quite similar to 13.5‰ reported for Lower Cretaceous sea sulfate (Claypool et al., 1980; Cecile et al., 1983). Khademi et al. (1997a) and Farpoor et al. (2004) also reported that sulfate in Isfahan and Rafsanjan areas, central Iran originates from the Lower Cretaceous sea sulfate. That is why the same origin for soil sulfate in this research is supported.

### Table 4. Summary of the sulfur and oxygen isotopic data of some selected sites.

<table>
<thead>
<tr>
<th></th>
<th>Mean (‰)</th>
<th>SD (‰)</th>
<th>No. of samples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sites 6-8</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\delta^{34}S$</td>
<td>+11.8</td>
<td>0.23</td>
<td>5</td>
</tr>
<tr>
<td>$\delta^{18}O$</td>
<td>+11.2</td>
<td>0.24</td>
<td>5</td>
</tr>
<tr>
<td>Sites 10-14</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\delta^{34}S$</td>
<td>+11.9</td>
<td>0.73</td>
<td>4</td>
</tr>
<tr>
<td>$\delta^{18}O$</td>
<td>+11.4</td>
<td>0.73</td>
<td>3</td>
</tr>
<tr>
<td>Sites 15-18</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\delta^{34}S$</td>
<td>+10.2</td>
<td>1.54</td>
<td>11</td>
</tr>
<tr>
<td>$\delta^{18}O$</td>
<td>+11.1</td>
<td>2.8</td>
<td>10</td>
</tr>
</tbody>
</table>
The oxygen and hydrogen isotopic composition of the crystallization water of soil gypsum from Isfahan, Rafsanjan and Loot areas, in central Iran, are compared with results of this study in Table 5. The δD and δ18O values in this study are lighter than the values of the Rafsanjan and heavier than Isfahan and Loot areas.

Since precipitation during cooler temperatures is depleted in heavier isotopes (Craig, 1961; Fontes, 1980; Gat, 1980; Khademi et al., 1997a; Farpoor et al., 2004; and Farpoor and Krouse, 2008), it seems that the study area has experienced a cooler temperature than Rafsanjan area (which is located 120 km west of the studied area) at the time of gypsum crystallization.

Three geochemical processes for gypsum formation have been described by Sofer (1978): oxidation of sulfide minerals, hydration of anhydrite, and direct deposition from an evaporating solution saturated with gypsum. Since the isotopic composition of local meteoric water for this study area was not available, data from Rafsanjan area, which is very similar in climate, was used. Table 6 shows the regression equations between δD and δ18O values from the study area as compared with Isfahan and Rafsanjan areas. Because gypsum crystallization water line and the equilibrium line are not parallel to the local meteoric water line (the intercept is at δ D= -78.5 and δ 18O= -15.5), oxidation of sulfide minerals could not have played a role in gypsum formation (Sofer, 1978; Dowuona et al., 1992c). On the other hand, gypsum has not been deposited by the hydration of anhydrite because the crystallization water and the mother water lines have not moved along a line with a negative ΔδD/Δδ 18O value (Matsubaya and Sakai, 1973; Sofer, 1978). The ΔδD/Δδ 18O value in the area is 2.4 and this means that evaporation from a solution saturated with gypsum was the major process for gypsum accumulation in the area. Sofer (1978), Khademi et al. (1997b), and Farpoor et al. (2004) found the ΔδD/Δδ 18O values of 2.6, 3.09, and 2.8, respectively, which are similar to the ratio obtained in the present research and could be a further support for the mode of gypsum formation in the area.

**CONCLUSIONS**

The δ13C values of soil organic carbon showed an increasing trend toward arid
downslope positions of the climotoposequence which is attributed to both the presence of C3 plants under stress and more decomposition of organic matter in lower positions.

The δ^{13}C and δ^{18}O values of pedogenic carbonates showed a positive correlation (r=0.6) in the arid downslope positions. This is probably attributed to the enrichment of both isotopes in the remaining HCO$_3^-$, during precipitation of CaCO$_3$ in the system (Salomon and Mook, 1976). However, this correlation (r= 0.37) was not observed in the upper landscape positions of the transect with a somewhat Mediterranean climate.

The δ^{13}C values of pedogenic carbonates decreased with decreasing elevation. Higher proportion of C3 plants in lower positions and presence of more C4 plants in the upper positions in the past could be considered as possible reasons.

The mean δ^{34}S value of the area (11.3‰) is very close to lower Cretaceous sea sulfate (13.5‰) which seems to indicate the sedimentary source of sulfur in the area studied. Mean δ D and δ^{18}O values of gypsum crystallization water (-51.2‰ and +6.0‰ respectively) show that direct deposition from an evaporating solution saturated with gypsum is the only possible mechanism of gypsum formation in the area studied.

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REFERENCES


چکیده
داده‌های ایزوتوپی کربن و اکسیژن به فهم بستر شیاطین محتویاتی زمان تشکیل خاک کمک می‌نمایند. خاک‌های ۱۸ محل نمونه‌برداری در طول یک رودخانه‌ای در ارتقایات اولیه زار (با ارتفاع ۴۳۵ متر از سطح دریا و رژیم رطوبتی-حرارتی زریک-مزیک) به مساحت ۱۵ هکتار در استان کرمان واقع در ایران مرکزی مورد مطالعه قرار گرفتند. مقادیر $^{13}$C مربوط به کربن آلی خاک از ۲۰۰۶-۱۳/۱۴ همزمان با تغییرات مقادیر $^{18}$O در زمین نمای همیشگی مثبت ($t=0$) بین $^{13}$C و $^{18}$O در حالتی ارتباط متقابل $t=0$ و در موقعیت‌های ناهایی شیب با اقلیم مدرن‌سازی نیز درست. به نظر می‌رسد که در موقعیت-۲/۳۷ PDB تغییرات $^{13}$C موجود در کربن آلی و نیز کربن‌های خاک‌زای و نیز روند تغییرات آنها در طول توالی نشان داد که مقادیر بیشتری از $^{13}$C در موقعیت-۳/۱۴ های ناهایی زمین‌های سیاه و مواد یافته در سطحی از $^{18}$O در تاریخچه بخش‌های فلوتی منطقه وجود داشته است. میانگین ۱۱/۳ مربوط به $^{34}$S نشان داد که سواحل با دریاهم‌سازی مربوط به رکنش $^{18}$O پایین به عنوان منبع $^{13}$C در منطقه می‌باشد. میانگین ۵۱/۷ و ۱/۸۰ پیمان به عنوان منبع $^{13}$C در منطقه می‌باشد. میانگین در آب شیب‌های بایوق گچ حاکی از آن است که رسم‌های مستقیم از محلول اشباع از $^{18}$O به عنوان نتیجه سازوکار تشکیل این کانال در منطقه به حساب می‌آید.