Microbial Processes in Ore-Bearing Laterite at the Tomtor Nb-REE Deposit: Evidence from Carbon Isotope Composition in Carbonates

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Abstract. The unique Nb-REE deposit is located within the Tomtor complex of ultramafic alkaline and carbonatite rocks in the northern Sakha Republic (Yakutia) (Kravchenko and Pokrovsky 1995; Dobretsov and Pokhilenko 2010; Lazareva et al. 2015). Ores reside in three layers (Severny, Yuzhny, and Buranny sites) which fill depressions in subsided profiles of weathered carbonatites. Judging by stable isotope analysis, carbonates from laterite weathering profiles at the Tomtor Nb-REE-deposit formed by different mechanisms, including microbially mediated organic-clastic sulfate reduction and anaerobic oxidation of methane.

Keywords: Tomtor · Nb-REE-deposit · Laterite · Carbonates · C-isotopes · Microbial · Methane

1 Introduction

Rare earth elements have been broadly used in advanced technologies. They often occur in carbonatite deposits among which laterite profiles of weathered carbonatites are most attractive commercial targets. A unique Nb-REE deposit is located within the Tomtor complex of ultramafic alkaline and carbonatite rocks in the northern Sakha Republic (Yakutia) (Kravchenko and Pokrovsky 1995; Dobretsov and Pokhilenko 2010; Lazareva et al. 2015). Ores reside in three layers (Severny, Yuzhny, and Buranny sites) which fill depressions in subsided profiles of weathered carbonatites. Lately, evidence has been obtained that localization of elements in ore-bearing beds may have biotic controls (Lazareva et al. 2015). The possible role of microorganisms in ore formation has not been discussed yet in the literature on laterite profiles from carbonatite deposits, including the giants of Mountain Weld (Australia) or Araxa and Catalão (Brazil). In this respect, the Tomtor deposit is remarkable by a contribution of biogenic and bacterial processes to its formation, besides magmatism and high-temperature hydrothermalism that are common to all carbonatite deposits. Biogenic agents are usually identified from the carbon isotope composition of rocks controlled by their interaction with fluids. The discussed stable isotope compositions of carbonates from ore-bearing laterites at the Tomtor deposit provide evidence of possible microbial mediation.

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S. Glagolev (Ed.): ICAM 2019, SPEES, pp. 208–211, 2019.
https://doi.org/10.1007/978-3-030-22974-0_49
2 Methods and Approaches

The stable isotope compositions were analyzed in three samples (TM-592, TM-590, and BH 101) from laterite ore zones with ~1%, 0.6%, and 1.6% REE, respectively. Extraction of monofractions for analysis is difficult because the laterite samples are fine grained while the carbonates are zoned. The problem can be solved by using selective acid extraction based on progressively slower reactions of carbonates with H₃PO₄ in the stoichiometric series: calcite → dolomite → ankerite → siderite → rhodochrosite. According to experimental evidence (Al-Aasm et al. 1990), the reaction duration sufficient for δ¹³C determination is 1 h for calcite, 24 h for dolomite, and 5–7 days for siderite and rhodochrosite. The C and O isotope compositions of carbonates are determined by digestion in anhydrous H₃PO₄ followed by measurements on a FINNIGAN MAT-253 mass spectrometer with a GasBench II analyzer, in a stream of pure helium. The GasBench unit was also used for preconditioning of samples digested at 70 °C. The results are quoted in (‰) relative to the Vienna Peedee belemnite PDB standard for carbon and relative to SMOW for oxygen. The analytical errors were ±0.1‰ for carbon and 0.15‰ for oxygen. The composition and morphology of minerals were studied at the Analytical Center of IGM (Novosibirsk) on a Tescan MIRA 3 LMU scanning electron microscope with Oxford Instruments Nanoanalysis Aztec Energy/INCA Energy 450+ XMax 80 and INCA Wave 500 analyzers, applicable to scan nanometer particles.

3 Results and Discussion

Sample TM-590 is composed mainly of goethite, siderite (with a Mn impurity), calcite, and rhodochrosite. Goethite occurs either as dripstone, with concentric zonation, or as pseudomorphs after a disappeared mineral. The carbonates exist as zoned anhedral grains, with calcite being the latest phase. Fine apatite crystals, pyroxene, monazite, pyrochlore, and sphalerite are present in lesser amounts. The rock-forming phases in TM-592 are goethite, siderite, and calcite and the accessories are apatite and monazite. Apatite occurs as fine prismatic euhedral grains in calcite. Sample BH 101 consists of finely intergrown Fe-chlorite (chamosite) and apatite. The apatite-chamosite aggregate encloses clearly seen large rhodochrosite grains with siderite in their core. The two phases have a distinct boundary but have inherited orientations of the crystallographic axes. The accessory phases are TiO₂, submicrometer monazite platelets, sphalerite, and galena.

Synthesis of C and O isotope data from magmatic and postmagmatic (hydrothermal, metasomatic) carbonates in different carbonatite deposits worldwide shows a large range of values from −10 to +3‰ (PDB) for δ¹³C and from +6 to +30‰ (SMOW) for δ¹⁸O (Deines 1989).

The δ¹³C values obtained for the Tomtor samples (Table 1) are lower. The δ¹³C patterns in carbonates are controlled by the isotope composition of bicarbonate. Low δ¹³C in bicarbonate is due to oxidation of organic matter where δ¹³C can reach −32‰ (Oleary 1988). In the course of microbially mediated sulfate reduction of geopolymers and biopolymers in aerobic conditions, the δ¹³C values of bicarbonate that forms by
reaction (1) and those of carbonate produced later by reaction (2) are inherited from the precursor component.

\[
2\text{CH}_2\text{O} + \text{SO}_4^{2-} \rightarrow 2\text{HCO}_3^- + \text{H}_2\text{S} \quad (1)
\]

\[
2\text{HCO}_3^- + \text{Me}^2+ = \text{MeCO}_3^- + \text{CO}_2 + \text{H}_2\text{O} \quad (2)
\]

There are two facts that implicitly support the above considerations: laterite contains organic remnants (Lazareva et al. 2015) that are electron donors in reaction (1); the samples contain sulfides, including frambooidal pyrite, which formed at the account of \( \text{H}_2\text{S} \) released in reaction (1). However, this model can account only for the isotope composition of the calcite component in TM-590, while other \( \delta^{13}\text{C} \) values (\( -24.8\%\) in TM-590) are below \( -32\%\), and may have formed with participation of methane. Biogenic methane has \( -55\%\) to \( -80\% \) \( \delta^{13}\text{C} \), and the \( \delta^{13}\text{C} \) values in bicarbonate resulting from microbially mediated methane oxidation (react/3) and in carbonate that formed later by reaction (2) will be low.

\[
\text{CH}_4 + \text{SO}_4^{2-} \rightarrow \text{HCO}_3^- + \text{HS}^- + \text{H}_2\text{O} \quad (3)
\]

**Table 1.** \( \delta^{13}\text{C} \) and \( \delta^{18}\text{O} \) (‰) in different fractions of carbonates

| Sample  | \( \delta^{13}\text{C} \) | \( \delta^{18}\text{O} \) |
|---------|----------------|----------------|
| TM-590  | -29.6          | +12.0          |
|         | -39.2          | +16.1          |
|         | -24.8          | +14.6          |
| TM-592  | -31.2          | +7.6           |
|         | -44.1          | +13.6          |
|         | -37.1          | +15.1          |
| BH 101  | -59.0          | +9.4           |
|         | -54.6          | 10.5           |

Note that a low value of \( \delta^{13}\text{C} \) was previously reported (Pokrovsky 1990; Kravchenko and Pokrovsky 1995) at the Tomtor deposit, but the mineralogical and geochemical characterization of the sample is not given.

### 4 Conclusions

The reported isotope data show that organic clastic sulfate reduction and anaerobic methane oxidation with participation of microbial communities were among key mechanisms responsible for the formation of carbonates in laterite profiles of the Tomtor Nb-REE deposit.

**Acknowledgements.** The study was supported by grant 18-17-00120 from the Russian Science Foundation.
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