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Constraints on Hf and Zr mobility in high-sulfidation epithermal systems: formation of kosnarite, KZr2(PO4)3, in the Chaquicocha gold deposit, Yanacocha district, Peru

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We report the first occurrence of Hf-rich kosnarite \([K(Hf,Zr)\_2(PO_4)\_3]\), space group \(R-3c\), \(Z=6\), in the giant Chaquicocha high-sulfidation epithermal gold deposit in the Yanacocha mining district, Peru. Kosnarite crystals are small (<100 \(\mu\)m) and occur in 2–3-mm-thick veins that cut intensively silicified rocks. The paragenesis includes a first stage of As-free pyrite and quartz (plus gratonite and rutile), followed by trace metal-rich pyrite \([(Fe,As,Pb,Au)S\_2]\) and secondary Fe-sulfates. Kosnarite is associated with quartz and is clearly late within the paragenetic sequence. Electron microprobe analyses (EMPA) of kosnarite show relatively high concentrations of HfO\(_2\) and Rb\(_2\)O (7.61 and 1.05 wt.\%, respectively). The re-calculated chemical formulas of kosnarite vary from \(K_{\Sigma 1.00}(Zr_{1.93}Na_{0.01}Hf_{0.01}Mn_{0.01})_{\Sigma 1.96}(P_{3.04}O_{4})_{\Sigma 3}\) to \((K_{0.92}Rb_{0.05}Na_{0.03})_{\Sigma 1.00}(Zr_{1.81}Hf_{0.19})_{\Sigma 2.00}[(P_{2.98}Si_{0.02}As_{0.01})_{\Sigma 3.01}O_{4}]_{\Sigma 3}\), where Hf and Rb are most likely incorporated according to a coupled substitution of Hf\(^{4+}\) + Rb\(^{+}\) \(\Leftrightarrow\) Zr\(^{4+}\) + K\(^{+}\). Back-scattered electron (BSE) images and elemental mapping of kosnarite reveal that Hf and Rb are enriched in 2–10-\(\mu\)m-wide oscillatory and/or sector zones. High-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) observations of such zones reveal a pattern of alternating, 5–50-nm-thick, Hf-rich and Zr-rich nanozones. These high-resolution observations indicate that the incorporation of Hf does not appear to cause significant distortion in the kosnarite structure. Semiquantitative TEM-energy-dispersive X-ray spectrometry (EDS) analyses of the nano-layers show up to 22 wt.% of HfO\(_2\), which corresponds to 31 mol% of the hypothetical, 

\[KHf_2(PO_4)_3\], end-member. The presence of kosnarite in the advanced argillic alteration zone at Yanacocha is indicative of Hf and Zr mobility under highly acidic conditions and points towards an unforeseen role of phosphates as sinks of Zr and Hf in high-sulfidation epithermal environments. Finally, potentially new geochronological applications of highly insoluble vein kosnarite,
including Rb-Sr dating, may provide further age constraints in pervasively altered areas where other isotopic systems might have been reset.

Kosnarite
Chaquicocha
Yanacocha
Hafnium
Zirconium
Phosphate

Editorial handling: R. Linnen

Introduction

High field strength elements (HFSE), including Zr, Hf, Nb, Ta, U, Th, and the rare earth elements (REEs) are generally incompatible in magmatic-hydrothermal systems and typically concentrate in accessory phases such as zircon. Despite the fact that they are relatively insoluble in most hydrothermal fluids and considered immobile, there is abundant evidence that these elements are mobile in aqueous fluids with specific hard ligands such as phosphate (PO₄)³⁻, among others (Linnen et al. 2014). In this paper, we report the first occurrence of the Hf-rich zirconium phosphate kosnarite [K(Hf,Zr)₂(PO₄)₃] in the giant Chaquicocha high-sulfidation epithermal gold deposit, Yanacocha district, Peru, and provide mineralogical evidence of Hf and Zr mobility in high-sulfidation epithermal systems. Kosnarite is believed to form only in pegmatitic environments during hydrothermal alteration of primary zircon, apatite, and other Zr-bearing minerals (More et al. 1983; Brownfield et al. 1993; Birch et al. 1994, 1995; Huminicki and Hawthorne 2002). Previous studies have confirmed the importance of aluminum-sulfate-phosphates (ASP) as minerals limiting the mobility of phosphorus during dissolution of primary apatite under highly acidic
conditions, generally pH <4 (Dill 2003). However, no secondary Zr- and/or Hf-rich minerals have been reported to form after dissolution of primary and accessory silicates such as zircon in extensively hydrothermally altered rocks. This paper combines mineral chemistry data, high-resolution observations, and thermodynamic modeling to confirm that kosnarite at Yanacocha is a sink for Hf and Zr in the advanced argillic alteration zone.

The Yanacocha district

The Yanacocha district straddles a portion of the continental divide in the high Andes of northern Peru and consists of eight large high-sulfidation epithermal gold deposits hosted by intense acid-leached and quartz-rich alteration in the Miocene Yanacocha Volcanic Complex. Ages of volcanism and hydrothermal advanced argillic alteration are well documented, and eruptions spanned ~6 million years from 14.5 to 8.4 Ma interspersed with discrete pulses of hydrothermal alunite from 13.6 to 8.2 Ma (Longo et al. 2010). Gold-bearing ore at Yanacocha is associated with intense quartz-rich, advanced argillic alteration in andesitic to dacitic subvolcanic domes, dikes, and pyroclastic rocks. The hydrothermal advanced argillic alteration assemblage is characterized by vuggy and granular quartz, alunite, pyrophyllite, ±dickite, ±kaolinite, and gold-bearing ores contain pyrite, covellite, enargite-digenite, tennantite, and coarse visible gold intergrown with barite, boitriodal limonite, and quartz in vugs and fractures (Longo et al. 2010).

Chaquicocha is one of the Yanacocha gold deposits, 1700 t of Au, located in the central part of the district. This deposit developed along near-vertical fracture zones in breccia and as stratiform bodies in acid-leached pyroclastic rocks below a barren “steam-heated” granular silica cap (Longo 2000; Teal and Benavides 2010). Gold ore is hosted by quartz-rich alteration with pyrite, enargite, tennantite, covellite, minor amounts of Pb-sulfides, and native gold (Deditius et al. 2008; Longo et al. 2010; and references therein). Local zones with bonanza-grade gold ore host native gold in silica-oxide breccia and vuggy quartz intergrown with barite and limonite (Longo 2000).
Samples and methods

Kosnarite was found in a fragment of a drill core from the Chaquicocha mine (CHQ-470/N4—depth, 343 m) in the Chaquicocha gold deposit in the Yanacocha district, Peru. The sample was examined in hand specimen, polished thin sections, and polished bulk samples by optical and scanning electron microscopy (SEM, Hitachi S3200N equipped with energy-dispersive X-ray spectrometry, EDS). The chemical composition of kosnarite was quantitatively determined by an electron microprobe analyzer (EMPA, Cameca SX100) using wavelength-dispersive spectroscopy (WDS) at the Electron Microbeam Analysis Laboratory (EMAL) at the University of Michigan. Accelerating voltage and beam current were 15 kV and 10 nA, respectively. The data acquisition time was 40 s (20 s on the peak and 10 s on each of the background positions) and spot ~1 μm analyses were performed. In order to increase the analytical total, the time of acquisition for Hf was set to 60 s. For F and Na, counting time of 20 s (10 s on the peak and 5 s on each of the background positions) and raster 5 μm were used to avoid Na and F migration under the beam. The PAP correction procedure was used to calculate the analyses. The standards used for calibration and detection limits (in ppm) are given in parenthesis: zircon (ZrSiO₄) for Zr(Lα) (4292) and Hf (Mα) (697), synthetic KTaO₃ for K (Kα) (305), CePO₄ for P(Kα) (1183) and Ce(Lα) (1184), pollucite for Rb (Lα) (510) and Cs (Lα) (844), albite for Al(Kα) (165) and Na(Kα) (349), CaTiO₃ for Ti(Kα) (384), andradite for Si(Kα) (312) and Fe(Kα) (493), manganotantallite for Mn (Kα) (486), arsenopyrite for As(Lα) (390), and Evans apatite for F(Kα) (745).

High-resolution transmission electron microscopy (HRTEM), analytical electron microscopy (AEM), and high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) were conducted using a JEOL JEM2010F at EMAL. HAADF-STEM images provided submicron Z-contrast resolution (in a way analogue to BSE images were obtained using EMPA/SEM). Analytical parameters/conditions were spherical aberration coefficient Cₛ = 1.0 mm, probe size = 1.0 nm
(STEM mode), and collection angle of the HAADF detector = 50–110 mrad. The size of condenser aperture was 20 μm. Samples were cut out of the thin sections and mounted on the 3-mm-diameter Cu grid. Subsequently, samples were polished using tripod method and the final thinning was performed by milling with an Ar ion beam (4.0 keV) in a Gatan precise ion-polishing system.

Speciation and stability diagrams were computed with the Geochemist’s Workbench® (Bethke 1996) using the SUPCRT92 data compilation (Johnson et al. 1992) complemented for Zr aqueous species and mineral phases with the chemical thermodynamics of zirconium (Brown et al. 2005) that critically reviews most of the thermodynamic Zr database available.

Mineral paragenesis and the chemical composition of kosnarite

Euhedral crystals of kosnarite occur in 2–3-mm-wide veins that cut an intensively silicified rock. These 10–100-μm crystals of kosnarite coexist with the pre-existing minerals like quartz, rutile, pyrite, Au-bearing arsenian pyrite, and Fe-sulfates (Fig. 1a). The observed paragenetic sequence was determined for one sample of drill core and it is as follows: (1) pyrite, + quartz, + gratonite [Pb₉As₄S₁₃], (2) As-pyrite₂ [(Fe,As,Pb,Au)S₂], (3) kosnarite + quartz, and (4) Fe-sulfates. Pyrite, pre-dates Au and most probably precipitated during quartz-alunite event (though, no alunite was found in the sample). Two generations of quartz were found in the sample, anhedral quartz₁ and euhedral quartz₂. The latter one formed in the pores and is most probably contemporaneous with kosnarite. However, the distinction between these two generations of quartz is not always straightforward, particularly in thin section. No porcelainous-textured silica was found in the sample. The back-scattered electron (BSE) images show three different types of Hf-rich zoning in the crystals of kosnarite: (i) fine, oscillatory zoning; dominant in the samples; (ii) crystallographically oriented sector zones that mimic the terminating faces of the prism; and (iii) irregularly distributed grains with patchy-type zoning (Fig. 1a–d).
Back-scattered electron (BSE) images of kosnarite (kos) and associated minerals. **a** Euhedral aggregates of kosnarite crystals deposited on the surface of quartz (qtz). Note the platy crystals of secondary Fe-sulfates (szomolnokite?) forming at the expense of porous arsenian pyrite (As-py) and gratonite (g). **b** Hf-rich patchy zoning in kosnarite formed on the surface of pyrite (py) and arsenian pyrite. **c** Fine, oscillatory Hf-rich zoning in kosnarite (*white arrows*). **d** BSE image of Hf sector zoning in kosnarite associated with elemental map of Hf (d1).
The EMPA analyses of kosnarite from the Chaquicocha deposit at Yanacocha show relatively high concentrations of HfO₂ and Rb₂O at 7.61 and 1.05 wt.%, respectively. The concentrations of other detected elements are ≤0.27 wt.% As₂O₃, ≤0.23 wt.% TiO₂, ≤0.22 wt.% Na₂O, ≤0.17 wt.% Al₂O₃, ≤0.10 wt.% FeO, and ≤0.05 wt.% MnO,
respectively. The concentrations of F, Cs₂O₃, and Ce₂O₃ are below detection limits (bdl) (Table 1). The re-calculated chemical formulas of kosnarite vary from (with 1σ standard deviation) \((K_{1.00}Na_{0.01})Σ_{1.01±0.04} (Zr_{1.93}Hf_{0.01}Mn_{0.01})Σ_{1.95±0.02} (P_{3.04±0.15}O_{3})\) to \((K_{0.92}Rb_{0.05}Na_{0.03})Σ_{1.00±0.04} (Zr_{1.81}Hf_{0.19})Σ_{2.00±0.02} [ (P_{2.98}Si_{0.02}As_{0.01})Σ_{3.01±0.02} O_{4} ] \). Based on the EMPA data, it is likely that Hf was incorporated into kosnarite according to coupled substitution: \(Hf^{4+} + Rb^+ \Leftrightarrow Zr^{4+} + K^+\). Additional, minor isovalent substitutions include, e.g., \(P^{5+} \Leftrightarrow As^{5+}\) and \(K^+ \Leftrightarrow Na^+\). It is important to note that Hf and Rb are heterogeneously distributed: EMPA analyses, BSE images, and elemental mapping reveal that these elements are enriched in oscillatory or sector zones between 2 and 10 μm thick, as shown in Fig. 1b–d.

**Table 1**

Representative analyses of kosnarite from Chaquicocha

<table>
<thead>
<tr>
<th>Analyses</th>
<th>Det. Lim. (ppm of element)</th>
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<th>3</th>
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<th>5</th>
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<td>(P₂O₅) (wt.%)</td>
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<td>40.90</td>
<td>41.77</td>
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<td>ZrO₂</td>
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<td>8.75</td>
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MnO, Ce₂O₃, Cs₂O₃, and F are below detection limits 486, 1184, 844, and 745, respectively. The concentrations of F, Cs₂O₃, and Ce₂O₃ are below detection limits (bdl) (Table 1). The re-calculated chemical formulas of kosnarite vary from (with 1σ standard deviation) \((K_{1.00}Na_{0.01})Σ_{1.01±0.04} (Zr_{1.93}Hf_{0.01}Mn_{0.01})Σ_{1.95±0.02} (P_{3.04±0.15}O_{3})\) to \((K_{0.92}Rb_{0.05}Na_{0.03})Σ_{1.00±0.04} (Zr_{1.81}Hf_{0.19})Σ_{2.00±0.02} [ (P_{2.98}Si_{0.02}As_{0.01})Σ_{3.01±0.02} O_{4} ] \). Based on the EMPA data, it is likely that Hf was incorporated into kosnarite according to coupled substitution: \(Hf^{4+} + Rb^+ \Leftrightarrow Zr^{4+} + K^+\). Additional, minor isovalent substitutions include, e.g., \(P^{5+} \Leftrightarrow As^{5+}\) and \(K^+ \Leftrightarrow Na^+\). It is important to note that Hf and Rb are heterogeneously distributed: EMPA analyses, BSE images, and elemental mapping reveal that these elements are enriched in oscillatory or sector zones between 2 and 10 μm thick, as shown in Fig. 1b–d.

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MnO, Ce₂O₃, Cs₂O₃, and F are below detection limits 486, 1184, 844, and 745, r recalculate per 12 oxygen atoms

*apfu* atom per formula unit
Kosnarite from the Chaquicocha deposit contains the highest amounts of Hf and Rb reported in the literature, significantly higher than kosnarite samples from pegmatites at Mount Mica, Black Mountains (USA), and Wycheproof (Australia). The maximum concentrations in oxides of Hf and Rb in the North-American pegmatites reach 1.4 and 0.3 wt.%, respectively (Brownfield et al. 1993), while slightly higher HfO₂ amounts of 1.7 wt.% were reported in the aforementioned Australian locality (Birch et al. 1994). The Hf-rich kosnarite reported in this study contains much lower concentrations of Mn and Na,
compared with kosnarite from Mount Mica, which are up to 1.3 MnO and 1.7 wt.% Na₂O, respectively.

Nanoscale Hf zoning in kosnarite

The HAADF-STEM images reveal that the individual 2–10-μm-thick oscillatory zoning identified by EMPA-BSE imaging consists of a set of alternating Hf-rich and Zr-rich nanozones with thickness varying between 1 and 50 nm (Fig. 2a, c). Some of the alternating Hf-rich and Zr-rich nanozones grew epitaxially on Zr-rich kosnarite that host inclusions of quartz (Fig. 2a). The semiquantitative TEM-EDS analyses of the individual nanozones reveal significant differences in elemental concentrations (Fig. 2c). The EDS analyses show concentrations of Hf as high as 22 wt.% of HfO₂, and within Hf-rich nanozones, Hf concentrations can be three times higher than the EMPA analyses. The Zr-rich, Hf-poor nanozones contains 7 wt.% of HfO₂, which is equivalent to the highest values obtained by EMPA analyses. This discrepancy is most probably due to the contamination of the excitation volume of the EMPA beam by submicron size Zr-rich zones. The combination of the HRTEM observations and EDS analyses of the areas next to the boundary between these two sets of nanozones (Fig. 2c–f) reveal that there is no detectable structural misfit or misorientation between the nanozones, in spite of significant differences in their chemical composition (Fig. 2e, f). Selected area electron diffraction (SAED) patterns across the nanozones reveal the single-crystal structure of kosnarite with sharp diffraction maxima (Fig. 2e).

Fig. 2

TEM images of nanozoning domains in kosnarite. a Hf-rich zone in the kosnarite matrix. Note the partial dissolution and replacement of Hf-rich area by Hf-depleted kosnarite. b Crystal boundary between Hf-rich and Zr-rich areas of kosnarite, patchy type of zoning. c, d TEM and HAADF-STEM images of kosnarite associated with the EDS spectra shown in image c. Note the position of the EDS analyses on the TEM image. e TEM image of the boundary between Hf-rich and Zr-rich zones. f HRTEM image and SAED pattern (inset, lower right) of the boundary between Hf and Zr zones (magnified area from e). No visible structural distortion is observed.
Discussion and implications

The presence of Hf-rich kosnarite in veins that cut massive silica and in voids of vuggy quartz is an evidence of mobilization and subsequent sequestration of Hf and Zr during extensive alteration of the volcanic rocks under acidic conditions, during intense quartz-rich advanced argillic alteration that hosts the high-sulfidation epithermal gold deposit at Chaquicocha. In contrast to pegmatitic environments, where kosnarite is contemporaneous with (Li,Be,Al,Ca)-hydrated secondary phosphates and Zr-bearing secondary phosphates (Brownfield et al. 1993; Birch et al. 1994, 1995), kosnarite from Chaquicocha coexists only with quartz (Fig. 1). The presence of earlier mineral assemblages in the analyzed samples, including Au-bearing As-pyrite and inclusions of fine-grained rutile in quartz (Fig. 1), suggests that kosnarite precipitated after the formation of quartz-rich advanced argillic alteration, massive quartz, and later Au deposition (cf., Longo et al. 2010). Because no relics of primary silicates and phosphates were found in the analyzed samples, it is plausible that Zr, Hf, P, K, and Rb were sourced from primary (Al) -silicates, apatite, and zircon after dissolution under acidic conditions, pH <4, characteristic for high-sulfidation epithermal deposits (Simmons et al. 2005). Consequently, the transport of chemical components throughout a network of fractures would lead to subsequent precipitation from aqueous fluids supersaturated with respect to kosnarite. Thermodynamic modeling based on the Brown et al. (2005) database indicates that the formation of kosnarite is controlled by physicochemical properties of the fluid that involves low pH conditions (<5); high contents of HPO₄²⁻ (log a[HPO₄²⁻] >15) that decrease with temperature; dominant Zr-OH complexes; low activities of F and Ca, log a[Ca²⁺] ≤−15 and log a[F⁻] ≤−5; and decreasing temperature (Fig. 3a). It is likely that the high content of HPO₄²⁻ can only be reached locally in epithermal conditions and might be the limiting condition for the formation of kosnarite. Based on the result of the thermodynamic modeling, the higher concentrations of Ca and F in the
hydrothermal fluid would promote precipitation of (CaF)-apatite instead of kosnarite. Our thermodynamic modeling is consistent with EMPA analyses showing concentrations of F in kosnarite below the detection limit for this element (745 ppm). Therefore, it is suggested that F was not a major ligand that enhanced migration of the relatively immobile Zr and Hf, a feature that was noted in other hydrothermal systems (Rubin et al. 1993). However, the presence of ZrF(OH)₃ and ZrF(OH)₂ complexes (Magdisov et al. 2011) cannot be ruled out. Moderately hard ligands like PO₄³⁻ could also complex Zr and Hf to facilitate their transport particularly under lower pH < 4 and lower temperature conditions, < 200 °C (Linnen et al. 2014).

**Fig. 3**

Activity-pH diagrams showing stability conditions for a kosnarite and crystalline FeSO₄. a The log a[HPO₄²⁻] vs. pH diagram shows an expansion of the stability field of kosnarite when temperature decreases from 170 °C (bright gray) to 100 °C (dark gray). This change occurs at the expense of whitlockite and H₃P₂O₇⁻ and H₂P₂O₇²⁻ aqueous species. Dashed lines show the stability fields of aqueous phosphate species. b Log fO₂ vs. pH diagram showing the stability field for FeSO₄ (approximating stability field of szomolnokite) and other Fe-bearing species. The stability of FeSO₄ (c) expands with increase of sulfate activity. The minor shifts in the stability fields of pyrite and hematite were omitted for clarity. The dashed lines denote the stability of S species. Magnetite disappears when S concentration is increased. Stability diagram calculations are based on the review of the thermodynamic data base of zirconium and its compounds (Brown et al. 2005), where kosnarite stability field is approximated to Zr(HPO₄)₂·H₂O.
The presence of Fe-sulfates (Fig. 1) suggests that SO₄ complexes could also facilitate the transport of Zr and Hf, particularly under acidic, pH <4, and oxidizing conditions (log f_O₂ >−32) (Rubin et al. 1993; Aja et al. 1995), although geochemical modeling indicates that the dominating Zr species are hydroxides such as Zr(OH)₄ (Brown et al. 2005).

Formation of crystalline FeSO₄ is promoted by relatively high S concentrations (log a[SO₄²⁻] ≥0.5) (Fig. 3b), which are much higher than log m[SO₄²⁻] = −2, as proposed for szomolnokite by Chouinard et al. (2005). The conditions of kosnarite formation presented here are in agreement with those reported for aluminum-sulfate-phosphate (ASP) precipitation, formed in the vicinity of the feeder systems of epithermal gold and porphyry deposits (Dill 2003; and references therein).

The presence of alternating, oscillatory, and sector Hf-Zr zoning in kosnarite suggests local changes in physicochemical conditions during kosnarite growth (Fig. 1c, d). Therefore, the dominant oscillatory nanoscale zoning in kosnarite may be attributed either to cyclic changes in the Hf concentration in the parental hydrothermal fluid and/or non-equilibrium crystal growth with depletion in Hf⁴⁺ content in the adjacent solution that resulted in compositional gradients (Putnis et al. 1992; Shore and Fowler 1996; Stowell et al. 2011). Significant differences in the HfO₂ content in adjacent nanozones (22 vs. 7 wt.%) were observed in single crystal of kosnarite. This fact, in addition to the nanoscale size...
(1–50-nm width) of the alternating Hf-rich zones, Hf-poor zones supports a non-equilibrium growth scenario. The observed sector and patchy zonings in kosnarite suggest that some of the crystals may have precipitated from separated batches of fluid under local equilibrium conditions, with selective partition of Hf and Rb and crystal growth rates exceeding the lattice diffusion rates (Watson and Liang 1995; Shtukenberg et al. 2009; Stowell et al. 2011). Therefore, it is unlikely that the oscillatory Hf zoning observed in kosnarite from the Chaquicocha deposit at Yanacocha is a result of rapid changes in fluid composition, pressure, and/or temperature as reported for oscillatory-zoned pyrites in fractured and sealed discontinuities in the deeper parts of other epithermal systems (Peterson and Mavrogenes 2014).

Concluding remarks
The observations reported in this note document the first occurrence of kosnarite in hydrothermally altered volcanic rocks and point to the previously unforeseen role of phosphates in partitioning Hf, Zr, and Rb in areas of extensive acid leaching such as those occurring in epithermal systems, geothermal fields, and even porphyry deposits. Furthermore, (Hf,Zr)-phosphates may be of particular importance in geochemical studies attempting to understand the behavior of high-field strength elements (HFSE) in hydrothermal systems. Finally, potentially new geochronological applications of highly insoluble vein kosnarite, including Rb-Sr dating, may provide further age constraints in pervasively altered areas where other isotopic systems might have been reset.

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