Fracture dolomite as an archive of continental palaeo-environmental conditions

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The origin of Quaternary dolomites in continental environments (e.g. karst and lakes) is barely constrained compared to marine dolomites in sedimentary records. Here we present a study of dolomite and aragonite formations infilling young fractures of the ‘Erzberg’ iron ore deposit, Austria, under continental-meteoric and low temperature conditions. Two dolomite generations formed shortly after the Last Glacial Maximum (~20 kyr BP): dolomite spheroids and matrix dolomite. Clumped isotope measurements and U/Th disequilibrium ages reveal formation temperatures of 0–3 °C (±6 °C) and 3–20 °C (±5 °C) for both dolomite types, and depositional ages around 19.21 ± 0.10 kyr BP and 13.97 ± 0.08 kyr BP or younger, respectively. Meteoric solution and carbonate isotope compositions (δ18O, δ13C and 87Sr/86Sr) indicate the dolomites formed via aragonite and high-Mg calcite precursors from CO2-degassed, Mg-rich solutions. Our study introduces low temperature dolomite formations and their application as a sedimentary-chemical archive.
Factors that control the origin and distribution of sedimentary dolomite in geologic records are intensely debated in the literature due to the much higher abundance of dolomites in ancient sedimentary rocks compared to modern marine and continental sediments. This enigma is referred to as the "dolomite problem." Numerous mechanisms and models to explain the kinetic constraints on low-temperature dolomite formation in nature have been postulated, with the recognition of mineralogical, hydrogeochemical, and microbiological factors.

Modern dolomite-forming environments are mainly marine associated, such as sabkha flats, methane seeps, playa lakes and hypersaline lagoons, where elevated water temperatures (up to ~60 °C), high Mg/Ca ratios and carbonate alkalinity as well as the presence of reactive templates (e.g., nuclei or seeds), sulfate-reducing bacteria and extracellular polymeric substances (EPS) secreted by microorganisms have been reasonably suggested to promote and accelerate dolomitization. Occurrences of modern and ancient sedimentary dolomites have also been observed in "exotic" continental surroundings, such as in speleothems, in palustrine-pedogenic sediments, in ephemeral wetlands, and in dolomitic lakes where dolomite frequently precipitates at low to ambient temperature (~4 °C to 35 °C). These continental dolomites are often interpreted to have been formed under the influence of micro- or macrobiota, but without the direct control of physicochemical conditions archetypically thought to facilitate dolomite precipitation, such as high Mg/Ca ratios, elevated alkalinity and temperature. Details on the nature, timing and processes governing the formation of continental dolomite at low temperature remain largely elusive and analytical evidence of the proposed cool formation temperatures, for example by clumped isotope thermometry, is often missing.

Here, we present a case study of dolomite and prominent CaCO₃ precipitates (i.e., laminated successions of aragonite and minor calcite) in infilling vertical fractures within the Erzberg iron ore deposit, Eastern Alps, Austria (Fig. 1). This locality hosts the largest siderite [FeCO₃] deposit worldwide. Herein, carbonate-based U/Th ages, formation temperatures derived from clumped isotope (Δ₁₅⁰₃) measurements and high-resolution chemical, petrographic and isotope (δ¹³C, δ¹⁸O and ⁸⁷Sr/⁸⁶Sr) records of authigenic dolomite and erzbergite fracture mineralizations were investigated. Local host rocks and modern meteoric water, collected from different sections of the open pit mine, were also assessed to constrain the nature, timing and driving factors of dolomite formation in a continental-meteoric setting at low temperature. Our results reveal a complex evolutionary history of the dolomite and erzbergite fracture mineralizations as well as the potential of these continental carbonate archives to be used in palaeo-environmental and palaeo-climate studies, complementary to the widespread modern and ancient carbonate deposits of the marine realm.

**Results**

**Occurrence of authigenic dolomite and erzbergite in young fractures.** The sedimentary and tectonic evolution of the Erzberg iron ore deposit is marked by Variscan age nappe stacking of Silurian and Carboniferous siliceous shales sandwiching lower Devonian limestones, which host the main iron ore body (Fig. 1c). Discordant Permo-Triassic transgressive sedimentary rocks and Late Cretaceous Alpine deformation follow this sequence. The sedimentary succession is cross-cut by numerous vertical fractures of variable width (few centimetres to several decimeters) and length (tens of meters) (Figs. 1d and 2a) that are interpreted to have been developed during fault tectonics or gravitational mass movements up to the Late Pleistocene. These fractures are partly healed with authigenic erzbergite and more rarely dolomite, especially in the southern part of the Erzberg, e.g., encountered on the pit level "Schuchart" (Fig. 2a-c). The young U/Th ages of the erzbergite (19.21 ± 0.10 to 13.11 ± 0.09 kyr BP, see Supplementary Table 1) indicate the healing of the fractures by carbonate precipitation started after the Last Glacial Maximum (~20 kyr BP) onwards (Fig. 2c). The formation of the erzbergite is post-dated by the occurrence of two generations of brecciated dolomite-hosting layers of either dark brown or light beige color (Fig. 2c), which replace the erzbergite and exhibit unique petrographic, mineralogical and isotope chemical signatures.

**Mineralogy, geochemistry, and petrography of dolomite fracture mineralizations.** The light beige dolomite-hosting layer consists of poorly ordered, calcian dolomite (79 ± 6 wt.%, 54 ± 1 mol% CaCO₃, cation ordering ratio of 0.25 ± 0.06; n = 6) erzbergite (8 ± 5 wt.% of aragonite), quartz (5 ± 1 wt.%), illite (4 ± 1 wt.%), Fe-(hydr)oxide grains (4 ± 2 wt.% of goethite and hematite) and rarely ankerite (see Fig. 3a and Supplementary Fig. 1a for bulk mineralogy). On the contrary, the dark brown dolomite-hosting layer contains more near-stoichiometric and better crystallized dolomite (74 ± 5 wt.% 51 ± 1 mol% CaCO₃, cation ordering ratio of 0.35 ± 0.08; n = 4) than the light beige layer. The dark brown dolomite-hosting layer has a similar proportion of quartz (5 ± 1 wt.% and illite (4 ± 2 wt.%), but a higher content of finely disseminated Fe-(hydr)oxides (13 ± 4 wt.%) and sporadically ankerite clasts, explaining its darker color. Clumped isotope data obtained from the two dolomite-bearing layers, and U/Th ages from the adjacent erzbergite deposits (labeled as EB1 to EB3 in Fig. 3a) hosting the dolomite mineralizations, reveal formation temperatures of 3 to 20 °C (± 5 °C) for the light beige colored layer and 0 to 3 °C (± 6 °C) for the dark brown colored layer (see Supplementary Table 2 for carbonate formation temperatures), with depositional ages younger than 13.97 ± 0.08 kyr BP and 19.21 ± 0.10 kyr BP, respectively.

Petrographically, both dolomite-hosting layers contain two types of sedimentary dolomite with distinct abundance, texture, crystal shape and composition. The light beige dolomite-hosting layer is dominated by very fine-grained (−5–15 μm in size), subhedral dolomite rhombs with a composition of Ca₁⁺₀.₀⁻₀.₁₉Mg₀.₇₈⁻₀.₉₉(⁸₀Ca₃₀₄) (Fig. 3b, d, f, g). This dolomite type (hereafter referred to as matrix dolomite) shows a diffuse contact to fibrous aragonite needles and corroded aragonite relics (Fig. 3d), exhibiting embayments and interpenetrated features, which are typical for a replacement process. The matrix dolomite is characterized by moderate amounts of Fe (1700 ± 1400 mg/kg) and Mn (400 ± 600 mg/kg) and low amounts of Na (10 ± 5 mg/kg) and Sr (20 ± 5 mg/kg), on average. Neither a cloudy-center-clear rim texture nor zonation patterns are observable in this dolomite type (see Supplementary Fig. 2 for dolomite composition and microstructural features). Rarely, spheroidal dolomite crystals interwoven with illitic clays are seen, accounting for <5 % of the total dolomite content, judged from optical microscopy of thick sections. In contrast, the dark brown dolomite-hosting layer appears as a mixture of matrix dolomite growing onto spheroidal dolomite, which presumably corrodes and replaces former acicular aragonite crystals and aragonite fans (Fig. 3c, e). Spheroidal dolomite locally accounts for up to 30% of the total dolomite content and exhibits, on average, lower concentrations of Sr (15 ± 5 mg/kg), similar Mn (400 ± 500 mg/kg) contents and slightly higher Fe (2500 ± 1700 mg/kg) and Na (15 ± 5 mg/kg) contents compared to the matrix dolomite. The dolomite spheroids are
~50–300 µm across and typically show a higher Ca content in the inner part and a higher stoichiometry toward the rim, in the range Ca$_{0.89–1.19}$Mg$_{0.80–1.10}$(CO$_3$)$_2$ (Fig. 3f, g).

**Discussion**

**The dolomitization process.** Fracture opening, host rock dissolution and prior aragonite precipitation: The host rocks surrounding the main carbonate fracture mineralizations are reddish-brown to light gray ferruginous limestones (the so-called “Sauberger Kalk”), which were deposited during the lower Devonian (Fig. 1a). These limestones are partially mineralized by iron ore carbonates, such as ankerite [Ca(Fe,Mg,Mn)(CO$_3$)$_2$] and siderite (Fig. 2a and Supplementary Fig. 1b for bulk mineralogy), which precipitated from Fe$^{2+}$-rich hypersaline solutions at temperatures between 200 °C and 350 °C in the Late Triassic.$^{32,34}$ Owing to this metasomatic overprinting, the ferruginous limestones and the iron ore carbonates are isotopically offset from mean Devonian marine limestones and further display metasomatic-epigenetic structures. The proposed (neo)tectonic activity in the Late Pleistocene (Stage i) in Fig. 4) led to the
formation of open fractures. Geologically young vertical fractures at Erzberg provide fresh chemical reaction surfaces and important water flow paths, which promote the infiltration of meteoric water-derived solutions from the adjacent catchment areas. Subsequent fluid-rock interactions cause oxidation of accessory (Fe,Cu,Zn)-sulfide minerals exposed in the iron ore host rocks and related generation of sulfuric acid, as supported by sulfur isotopic compositions of modern surface and fracture water (−4.9 to +2.5‰ of δ34SSO₄, VCDT) and iron sulfides (−2.2 to +5.1‰ of δ34S_Pyrite, VCDT) at Erzberg. Sulfuric acid attack facilitates an efficient dissolution of the partly brecciated carbonate host rocks (Stage ii) in Fig. 4), which leads to the mobilization and subsequent enrichment of Ca, Mg, trace elements (Fe, Sr, Mn) and alkalinity along the water flow paths. These processes, coupled with fast outgassing of CO₂ in the fissured aquifer (Fig. 5), generate highly differentiated meteoric water-derived solutions with distinct hydrogeochemical signatures (see Supplementary Table 3 for modern water composition). These include high Mg/Ca molar ratios (up to 26) and alkaline conditions (pH: 8.1 to 8.8), thus, favorable for the precipitation of the first generations of the erzbergite (EB 1 and 2) shortly after the Last Glacial Maximum (Stage iii) in Fig. 4). Such precipitation of aragonite from meteoric water-derived solutions at very low temperatures is indicated by the clumped isotope data (3 ± 4°C for EB1), the positive δ¹³CCarb values due to enhanced CO₂ outgassing and isotopically light palaeofluid δ¹⁸O values (inferred from carbonate δ¹⁸O values and clumped isotope temperatures) in the range of −11.8 ± 0.9‰ (VSMOW, Supplementary Table 2). These constraints are typical for continental carbonates formed at low temperatures. The formation of aragonite is initially favored over other Ca-Mg carbonates, because of the strong kinetic barriers induced by the attachment of partially dehydrated Mg²⁺ ions onto the surfaces of incipient calcite nuclei inhibiting their growth and favoring the precipitation of aragonite under oxic-vadose conditions, which incorporates virtually no Mg. Prior aragonite precipitation along the water flow path under predominantly open system conditions with respect to gas-solution exchange causes an increase of the aqueous Mg/Ca molar ratio downstream, which promotes the precipitation of (very) high Mg-calcite (or an amorphous carbonate precursor) in open conditions.
fractures and in the fissured and brecciated host rocks (see Supplementary Fig. 1b for bulk mineralogy), thereby providing the precursors for subsequent transformation to dolomite (see Supplementary Fig. 3 for systematics in carbonate vein mineralizations). Carbonate mineral precipitation under closed system conditions is unlikely, because the high $\delta^{13}C$ values of the precipitates indicate CO$_2$ degassing and the low $\delta^{18}O$ values indicate a meteoric water origin. We exclude effects of mineral transition$^{41}$ due to lack of recent hydrothermal activity at Erzberg and experimental evidence against low temperature influence of carbonate mineral transition on clumped isotope signatures$^{42,43}$.

**Main dolomitization event(s):** The fracture dolomites at Erzberg represent the final product of coupled dissolution and re-precipitation reactions involving metastable aragonite and
high-Mg calcite (or amorphous precursor phases), as well as CO₂-degassed, meteoric water-derived solutions rich in Mg⁴³. Petrographic analyses of Ca-Mg carbonate deposits from modern caves indicate that increased aqueous Mg/Ca molar ratios and CO₂ outgassing promote sequential carbonate mineral formation, which follows the succession: aragonite, high-Mg calcite, huntite [Mg₃Ca(CO₃)₄], disordered dolomite and finally ordered dolomite⁴⁴. This sequence of mineral growth and transformation can also be seen in the carbonate fracture mineralizations at Erzberg, except for huntite, which was not detected: A first generation of “old and cool” dolomite spheroids formed at the expense of aragonite and high-Mg calcite precursors (EB 1 and 2 in Fig. 3a, c) at very low temperatures (0 to 3 ± 6 °C) and under vadose conditions (palaeo-fluid δ¹⁸O: −14.1 ± 1.3 to −12.7 ± 1.6‰, VSMOW) between 19.21 ± 0.10 kyr BP and 13.97 ± 0.08 kyr BP (Stage iv) in Fig. 4). The establishment of generally warmer temperatures (3 to 20 ± 5 °C) and more humid climatic conditions with increased precipitation shortly after the last glaciation caused shifts in the water recharge and flow regime (palaeo-fluid δ¹⁸O: −12.9 ± 1.8 to −8.5 ± 0.9‰, VSMOW). This is seen in the
transition to phreatic conditions\textsuperscript{45}, and thus in the carbonate mineral precipitation mode, which is reflected by the formation of younger erzbergite deposits (EB 2 and 3 in Fig. 3a), followed by high-Mg calcite and then “young and warm” matrix-replacive dolomite (Fig. 3b) from about 14 kyr BP onwards (Stage v) in Fig. 4. Extrapolation of kinetic datasets obtained from dolomite formation through Ca-Mg carbonate intermediates and estimation of the time needed for the ordering of the dolomite crystal lattice (cf. Eq. 7 in\textsuperscript{46}) indicate that \(-3.0 \pm 0.2\) kyr and \(-2.8\) to 1.8 kyr are required to form a near-stoichiometric dolomite with a cation ordering ratio of 0.3 at 0 to 3 °C and 3 to 20 °C (see Supplementary Table 2 for clumped isotope-based carbonate formation temperatures), respectively, which is consistent with the dark and light brown dolomite-bearing layers at Erzberg. The two dolomite-bearing layers are cross-cut by a youngest aragonite-calcite vein generation, which shows a diffuse transition zone with partial dolomitization and dolomite overgrowths (see Supplementary Fig. 3 for systematics in carbonate vein mineralizations), indicating that dolomite precipitation has probably continued until recently (Stage vi) in Fig. 4). Hints for ongoing erzbergite formation and dolomitization are provided by a few occurrences of sub-recent aragonite veins (1.03 \pm 0.04 kyr BP\textsuperscript{30}), the modern hydrochemistry at Erzberg promoting aragonite and dolomite formation (see Supplementary Table 3), and (palaeo-)fluid \(\delta^{18}O\) values obtained from erzbergite and matrix dolomite resembling \(\delta^{18}O\) compositions measured for modern fracture and surface water (see Supplementary Table 2).

Special features of continental-meteoric dolomite formation at Erzberg: In contrast to previously published studies on sedimentary dolomite formation in continental-meteoric or marine-lacustrine environments at low temperatures\textsuperscript{20,31,32,25,26,46,47}, we found no evidence for detrital dolomite in the local carbonate host rocks and in the adjacent sedimentary rocks from the catchment areas, demonstrating that the dolomites at Erzberg are authigenic. This argues against the possibility that young fault-activity has pulverized any host rock dolomites and that such reactive (nano-scale) grains could have acted as nuclei for the later (over)growth of the fracture dolomites at a low temperature\textsuperscript{48}. On the contrary, a participation of ankerite nuclei in the dolomitization process cannot be completely excluded, because of the structural similarity between the two minerals.

Modern or relict microbial biofilms, laminated crusts, fossilized algae, EPS or bacterial forms have not been identified in the dolomite-bearing layers at Erzberg, which is consistent with the low total organic carbon contents (TOC of \(<0.1\) wt.%) of the carbonate rocks. Similar features were derived from the study of Quaternary speleothem-associated dolomites of the Snežna Jama cave, Slovenia, i.e., no evidence for the participation of microorganisms during dolomite formation was found\textsuperscript{20}. However, genetic relations between dolomite crystal growth and microbes have been observed in other karst environments, such as in speleothems of the Cayman Brac and in speleothems of lava tubes of Hawai\textsuperscript{i}\textsuperscript{20}. It has been argued that disordered dolomite, that is, mineralogically speaking, a very high-Mg calcite, can biotically precipitate at ambient temperature from solutions containing polysaccharides, which is readily produced from degrading organic matter\textsuperscript{19}. Recently, it was hypothesized that such disordered Ca-Mg carbonate precursors could serve as a reactive template to recrystallize to dolomite over time\textsuperscript{10,14}. Thus, the involvement of microbes, whether direct or indirect, in the cold fracture system at Erzberg occupied by carbonate minerals that has been active since the last glaciation cannot be completely ruled out.

Arguably low to moderate concentrations of reduct-sensitive elements (2100 to 2800 ± 2200 mg/kg of Fe and Mn) and the virtual absence of any transition metals in the dolomites from Erzberg indicate that dominantly oxic conditions prevailed during the majority of the crystal growth of the dolomites. This contrasts with the occurrences of modern and ancient dolomites in mostly reducing sedimentary environments and deep-marine dolomites formed alongside anaerobic oxidation of methane, which frequently contain much higher Fe and Mn contents (e.g., several mole percent\textsuperscript{13,50–53}). Moreover, the extremely low Na and Sr contents (30 ± 10 mg/kg) of the dolomites from Erzberg greatly differ from common marine and lacustrine dolomites (e.g., several hundreds of mg/kg\textsuperscript{13}).

Finally, it seems unlikely that the presence of illicite clays or carboxyl functional groups had a strong influence on dolomitization at Erzberg, i.e., through electrostatic binding, desolvation and carbonation of Mg\textsuperscript{2+} and Ca\textsuperscript{2+} ions at the fluid-mineral interface\textsuperscript{16,19}. This is indicated by the low contents of illite in the carbonate fracture mineralizations and low concentrations of dissolved organic carbon species (DOC: 0.4 mg/l) present in the modern meteoric water from southern sections at Erzberg.

It can be concluded that high Mg/Ca ratios and carbonate alkalinity, elevated pH, carbonate precursors and reaction time\textsuperscript{5,8,46} are likely the most critical parameters for controlling the formation of the fracture dolomites at Erzberg.
physicochemical conditions in the continental-meteoric setting at Erzberg have indeed led to the formation of fracture dolomites that are encountered unique, e.g., see Supplementary Tables 4–8 for mineralogical, elemental, and isotopic signature.

Fracture dolomite as a continental palaeo-environmental archive. The reconstruction of palaeo-environmental conditions recorded in continental and marine carbonate archives is the key to the understanding of the Earth’s climatic evolution over time, and for the prediction of the potential responses of terrestrial ecosystems to modern climate change33,34. At present, the reconstruction of continental depositional environments of the past and evidence for particularly rapid climate change in modern times are largely derived from the study of speleothems, lake deposits, travertine, fracture carbonates and soil carbonates, i.e., calcrite55,56. Our study demonstrates the potential of authigenic carbonates infilling young fractures in the low temperature and high-altitude alpine environment of Erzberg to be used as a sensitive palaeo-environmental archive. Apart from the prominent erzbergite fracture mineralization, the applicability in palaeo-environmental reconstruction studies has been recently evaluated30, the carbonate fracture mineralizations exposed in the southern section at Erzberg host two generations of authigenic fracture dolomites. Both dolomite generations carry important environmental information on varying palaeo-fluid δ18O compositions (−14.1 to −8.5 ‰, VSMOW), water flow paths and related crystallization modes (phreatic vs. vadose), palaeo-redox conditions (dominantly oxic), dolomite depositional ages (Pleistocene to recent) and formation temperatures (glacial to ambient interglacial), which have not yet been been reported in dolomite research.

Complementary analyses of radiogenic Sr (87Sr/86Sr) isotopic compositions of the erzbergite, dolomite-bearing layers, carbonate host rocks and meteoric water-derived solutions provide additional clues on the provenance of Sr in the carbonates, the origin of water and fluid-rock interactions37, with implications for the interpretation of modern hydrochemistry at Erzberg. Specifically, from the cross-plot of the 87Sr/86Sr composition vs. 1/Sr (Fig. 6) the following conclusions can be drawn: (i) The origin of the carbonate fracture mineralizations at Erzberg can be explained by contributions from two sources, with iron ore carbonates and Devonian limestones (Sauberger Kalke) as end-members. (ii) The 87Sr/86Sr isotopic composition of the spheroidal dolomite is mainly sourced from the dissolution of the iron ore carbonates, which suggests that intensified weathering and restricted water flow prevailed at around the Last Glacial Maximum (cf. Stage iv) in Fig. 4). (iii) Erzbergite and matrix dolomite have a mixed 87Sr/88Sr isotopic composition, indicative of changes in provenance and related water mixing regimes in more recent times (cf. Stage iv) and (v) in Fig. 4). (iv) The modern hydrochemistry at Erzberg still holds a great potential for recent erzbergite formation and even dolomitization, as inferred from the similar 87Sr/86Sr isotopic compositions of geologically young erzbergite, matrix dolomite and meteoric water from southern sections at Erzberg. Most of the modern fracture water and surface water at Erzberg are strongly oversaturated with respect to aragonite, calcite and dolomite (see Supplementary Table 3 for modern water composition), which suggests that the carbonate vein mineralizations still proceed until recently. The latter findings make this study site interesting for future studies on thermodynamic and isotopic equilibria between modern meteoric water-derived solutions and young Ca-Mg carbonates.

Taking the valuable palaeo-environmental and climatic information recorded in the carbonate fracture mineralizations at Erzberg, we call for a change in the long-standing conceptual view that fracture and gangue carbonates formed in continental surroundings are not particularly useful in regional-scale environmental reconstruction studies58,59. Considering the vast expanse of cool and high-latitude settings and high-altitude mountain regions worldwide, well known to be particularly sensitive to climate variations, and the extensive occurrence of carbonates therein60,61, a critical re-evaluation of the proxy potential of these mineral archives (including the observation of authigeneic dolomite) is needed to enlarge both the quality and quantity of currently available carbonate-based continental-sedimentary records. In this line, already known occurrences of speleothem-associated dolomites, for example, in the Titus Canyon Cave (California, USA), the Carlsbad Caverns (New Mexico, USA), the Baruta Cave (Venezuela), the Sumidero Tenejapa Cave (Mexico), the Soreq Cave (Israel), the Cueva de Nerja Cave (Spain), the Castañar Cave (Spain), and the Cayman Brac (UK) (see ref. 28 and references therein), should be revisited regarding their applicability in palaeo-environmental reconstructions. Furthermore, if one could demonstrate that Quaternary, post-glacial and deep marine dolomite deposits are more widespread than previously thought, recent uncertainties in the oceanic Mg budget could be much better constrained: Based on an extensive δ26Mg isotope dataset it has been proposed that a significant, yet undetermined, dolomitic sink should exist ever since the Neogene62. Our study results demonstrate dolomite formation via Ca-carbonate precursors at cool to ambient Earth surface temperatures within a geologically short timeframe, which is consistent with the sub-recent dolomite formation in deep-marine basins proposed recently62. It is up to future studies to quantify the fluxes of Mg linked to the formation of dolomite and its precursors in different continental and marine sedimentary basins and to estimate the relative contribution of such environments to the global Mg cycling and budget. We conclude that fracture dolomites precipitating in continental-meteoric settings at low temperature can create unique and environmentally restrictive chemical-sedimentary archives, which should be considered in future climate and environmental studies.

Methods

Materials. Field work and related collection of oriented, hand-sized specimens of erzbergite, carbonate host rocks (Sauberg Kalk and iron ore carbonates) and dolomite-hosting mineralizations was carried out during several field trips from November 2016 to June 2018. Samples were taken explicitly from the southern section of Erzberg, because the locally more widespread authigenic carbonate fracture mineralizations contain abundant dolomite (for sampling sites see Figs. 1 and 2). The more restricted occurrences of fracture carbonates from the central and northern sections are typically composed exclusively of erzbergite. The authigenic dolomite-hosting layers from this site are of particular interest, because of their continental-meteoric and low temperature origin and the young depositional age. Making these carbonate fracture mineralizations interesting as archive for climatic and palaeo-environmental conditions. In addition, modern fracture and surface water from northern, central and southern sections at Erzberg were analyzed for their chemical and isotopic composition (for sampling locations see Fig. 1b). The samples were initially recovered during two field campaigns in July 2015 and April 2016 in the different places at Erzberg68. Access to the different sampling sites across the open pit mine was permitted by DI Hannes M. Pluch (Geologie, Bergbauplanung & Qualitätsteuerung, VA Erzberg GmbH).

Characterization of modern aqueous solutions. Water temperature, specific electric conductivity and pH were measured on-site using a WTW Multi350i combi instrument equipped with SenTix41 and TetraCon325 electrodes. Liquid samples were filtered via 0.45 μm cellulose acetate membranes. One aliquot per sample was kept untreated, whereas another aliquot was acidified to 2% HNO3 matrix for subsequent hydrogeochemical analysis (see below).

Major cations and anions in solution were measured by ion chromatography ( Dionex ICS-3000; analytical error: <3%). Carbonate alkalinity (expressed as HCO3−) was measured by automatic titration with a 0.02 M HCl solution (Schott Titriloop; more: analytical error: <3%). Minor and trace elemental concentrations were determined by inductively coupled plasma—optical emission spectrometry (ICP-OES, Optima 8300 DV; analytical error: <5%). The dissolved organic carbon
Fig. 6 $^{87}$Sr/$^{86}$Sr vs. $^{1}$Sr diagram indicating a mixed origin of the authigenic carbonate fracture mineralizations at Erzberg. The "old and cool" dark brown dolomite spheroids are mainly sourced from dissolution of iron ore carbonates, whereas the "young and warm" light beige dolomite and the erzbergite carbonate have a mixed origin, being strongly affected by the Sauberger Kalk aquifer host rock. Modern water draining the Erzberg (right site), especially from the southern area, are beneficial for (sub-)recent carbonate precipitation and dolomitization. Note that all carbonates have a non-marine $^{87}$Sr/$^{86}$Sr isotopic signature due to the variable enrichment of radiogenic Sr during multiple recrystallization events, which are most likely related to the mineralization of the iron ore carbonates. Analytical error is smaller than symbol size unless stated otherwise.

(DOC) concentration was analyzed using a Shimadzu TOC-V-CPH Total Organic Carbon Analyzer (analytical precision: ±0.1 mg/l).

Hydrochemical modeling based on the measured water parameters was carried out using the PHREEQC computer code with its water.dat database. This included the calculation of aqueous speciation, ion activities, ionic strength, ion charge balance and saturation indices of the surface and fracture water at Erzberg with respect to calcite, aragonite and dolomite, as well as CO$_3^2-$ partial pressure (internal pCO$_2$).

Stable isotope analyses of meteoric water-derived solutions included their hydrogen ($\delta^2$H) and oxygen ($\delta^{18}$O) isotopic values, carbon isotopes of dissolved inorganic carbon ($\delta^{13}$C$_{\text{DIC}}$) and sulfur isotopes of dissolved sulfate ($\delta^{34}$S$_{\text{SO}_4}$). The $\delta^2$H and $\delta^{18}$O values of water were analyzed by wavelength-scanned cavity ring-down spectroscopy (WS-CRDS) using a Picarro L1102-i system at JR-AquaConSol GmbH (Graz, Austria). Water isotope values are reported relative to the Vienna Standard Mean Ocean Water (VSMOW) reference material (analytical precision: ±0.8‰). The $\delta^{13}$C$_{\text{DIC}}$ compositions were analyzed accordingly and are reported relative to the Vienna Pee-Dee Belemnite (VPDB) reference material (analytical precision: ±0.1‰ for $\delta^{13}$C$_{\text{DIC}}$). Isotope ratios of $\delta^{34}$S$_{\text{SO}_4}$ were obtained on a Finnigan MAT 253 mass spectrometer coupled to a Thermo Flash 2000 elemental analyzer through a Thermo Finnigan Conquest interface at the Leibniz-Institute for Baltic Research Warnemünde (Germany). Isotope values are reported relative to the Vienna Canyon Diablo Troilite (VCDT) reference material (analytical precision: ±0.2‰).

The radiogenic Sr ($^{87}$Sr/$^{86}$Sr) isotopic composition of selected water samples (six in total) was analyzed with a Plasma II multi-collector inductively coupled plasma mass spectrometer (Nu Instruments, Wrexham, UK; MC-ICP-MS; analytical reproducibility of NIST NR 987 standard: ±0.00001 for $^{87}$Sr/$^{86}$Sr, see Supplementary Table 6) at the NAWI Graz Central Laboratory for Water, Minerals and Rocks, Graz University of Technology (Graz, Austria).

Hydrogeological and isotopic compositions of the modern water draining the Erzberg are reported in Supplementary Table 3. An extended discussion of the modern hydrochemistry at Erzberg and a full description of the analytical methods used is provided elsewhere.

Solid-phase characterization. Polished thick sections (~50 µm thick) were prepared from the collected rock specimens to study (i) the mineralogy and petrography of the authigenic carbonate fracture mineralizations as well as (ii) possible indications of post-depositional alteration, such as recrystallization of aragonite and dolomite (over)growth features, by optical microscopy using a Leica DMLP Polarizing Microscope connected to an Olympus DP26 camera.

Rietveld-based analyses of powder X-ray diffraction (XRD) patterns (PANalytical XPert PRO diffractometer equipped with a Cu-Kα X-ray radiation source and a Scientific X’Celerator detector) obtained from bulk rock samples (cf. Fig. 2a) were made for mineral identification and quantification using the PANalytical HighScore Plus software package and pdf-4 database (analytical error: <3 wt.% for all mineral phases, see Supplementary Table 5). The measuring conditions were as follows: 40 kV, 40 mA, range: 4–85° 2θ, step size: 0.008° 2θ and count time: 40 s per step. Dolomite stoichiometry was calculated from the XRD patterns using the linear expression: mol% CaCO$_3$ = 333.33 × ($d_{(040)/d_{(104)}}$) $^{\times}$ 911.99.

Therefore, the position of the $d_{(104)}$-reflection of dolomite was fixed to the position of the main two quartz reflections used (analytical error: <0.6 mol% CaCO$_3$). The cation ordering ratio in dolomite was calculated from the intensity ratio of the $d_{(021)/d_{(110)}}$ peaks. This approach is semi-quantitative and its accuracy depends on particle orientation and size (analytical precision: ±3%)..

The rock textures, microfabrics, and geochemical compositions of the erzbergite and dolomite-hosting layers from Erzberg were obtained on polished thick sections by electron microprobe analyses (EMPMA) conducted on a JEOL JXA8530F Plus Hyper Probe (JEOL, Tokyo, Japan) at Karl-Franzens-University Graz. Element distribution maps of 1624 x 1024 pixel resolution were acquired using a focused beam, 15 keV accelerating voltage, 30 nA beam current and a dwell time of 12 ms/step. For the single spot analyses of the geochemical composition of the carbonates the analytical conditions were as follows: 15 keV accelerating voltage, 0.5–3.0 nA beam current and defocused beam, ~1.0–3.0 µm in size. The quantitative chemical analyses were standardized against natural and synthetic crystals, and included the following elements with characteristic spectral lines: Kα of Ca, Mg and Sr (dolomite), Fe-Kα (siderite), Mn-Kα (rhodochrosite) and Sr-Lα ( celestine). Counting times of 20–40 s on peak and 10–20 s on the background position on each side of the Ca, Mg, Fe, Mn, and Sr peaks were used. Only compositions with an analytical error of ±3% were included into the further consideration (see Supplementary Table 8).

Note here that the Na and Sr contents of the authigenic fracture carbonates from Erzberg were obtained by inductively coupled plasma optical emission spectroscopy (PerkinElmer Optima 8300) on acid-digested bulk rock samples (6% HNO$_3$ matrix, ultrasonic treatment at 70 °C for 12 hours). The estimated analytical error is better than ±3% (2 SD, $n = 3$) for the Na and Sr analyses.

The total organic carbon (TOC) content of the carbonates was measured on a Shimadzu TOC-VCPH – ASI-V Total Organic Carbon Analyzer at an analytical error of ±3%.

For $^{238}$U–$^{234}$U–$^{230}$Th isotopic analyses, small sample specimens (few tens of milligrams of powder) were milled along selected aragonite (erzbergite) growth layers with a tungsten-carbide handheld drill. Subsamples were selected from positions dependent on the anticipated degree of detrital mineral contamination (identified by coloring) or diagenetic alteration (aragonite-to-calcite transition), i.e., all the selected subsamples consist of pure, snow-white aragonite (cf. Fig. 2). Pure dolomite subsamples were not extracted, because abundant Fe-(oxy)hydrates and potential aragonite contents can cause large uncertainties and shifts in the conventional U/Th age determination. Moreover, application of the U/Pb small scale isochron dating method by laser ablation—sector field—ICP-MS$^{31}$ to our samples is erroneous, making it impossible to precisely date dolomite phases. After
acid digestion of the selected erzbergite subsamples using nitric acid and the addition of a 239Th tracer (spike), U and Th aliquots were separated and purified from the carbonate matrix after applying iron co-precipitation and anion-exchange chromatography. This wet-chemical operating procedure is accommodated in a clean laboratory environment. Routinely, chemical yields can reach ~90% or better for both the U and Th fractions. The U and Th isotope ratios relevant for the radiometric age determinations were measured via a ThermoFisher Scientific Scientific ScienTopp Neptune Plus MC-ICP-MS at the Central Laboratory for Water, Minerals and Rocks, Graz University of Technology. Sample solutions were introduced to the instrument via a desolvating nebulizer. Apart from relatively abundant 232Th and 232U measured by Faraday collectors, the different U and Th isotopes were analyzed in peak-jumping mode applying a secondary electron multiplier installed after a retarding potential quadrupole lens, i.e., an energy filter to improve abundance sensitivity. The mass spectrometric data yielded 238U concentrations from 102 to 1379 ng/g (mean 753 ng/g), i.e., a broad range of intermediate to high U contents favorable for precise age constraints. Likewise, the analyzed 232Th concentrations ranged from 360 to 1330 pg/g and, therefore, a relatively large range of 238U/232Th atomic ratios, from 65 to 4032 (±190), was measured. Absolute U–Th ages were calculated using published 238U and 230Th decay constants87 and all the ages were corrected using a bulk Earth 230Th/232U atomic ratio of 4.4 ± 2.2 mg/kg. In the vast majority of the samples analyzed, the influence of the initial Th correction was of minor relevance (cf. Supplementary Table 1, uncorr. vs. corr. ages). The measured isotope ratios and resulting age uncertainties are reported at the 2σ level. Typical relative age errors of the Ca-carbonate precipitates range between 0.4 and 1%. Hence, absolute age uncertainties range from decades to few hundreds of years, indicative of U–Th ages of high precision. Finally, the age data were referenced to “years Before Present” (yr BP), where the present is defined as the year 1950 A.D.

Stable O and C isotope compositions of the carbonate fracture mineralizations as well as of the carbonate host rocks were obtained from powdered subsamples (0.3 to 0.6 mg; handheld drill) by continuous-flow isotope-ratio mass spectrometry (ThermoFisher Scientific GasBench II connected to a Finnigan Delta Plus XP spectrometer) in the laboratory facilities of the JRAquaConSol GmbH (Graz, Austria). Isotope ratios are reported relative to the Vienna Pee-Dee Belemnite (VPDB) reference material (analytical precision: ±0.1% for δ13C and ±0.08‰ for δ18O, see Supplementary Table 7).

The δ56Sr/86Sr isotopic compositions of the carbonate fracture mineralizations and host rocks were measured on a Plasma II MC-ICP-MS at the NAWI Graz Central Laboratory for Water, Minerals and Rocks, Graz University of Technology (Graz, Austria) at an analytical precision of 0.00001% for δ56Sr/86Sr, NBS 987 (see Supplementary Table 6).

Samples for clumped isotope analyses were obtained following the procedure described above for the U–Th age determinations, i.e., pure, snow-white aragonite was drilled with a tungsten-carbide handheld drill. In addition, areas containing pure dolomite were drilled to obtain a few milligrams of sample powder. The mineralogies of the powders was investigated by XRD, which yielded almost pure samples, i.e., aragonite samples free of dolomite, as well as dolomite samples containing less than 1% aragonite by weight. Carbonation temperatures of the carbonate fracture mineralizations were then derived from clumped isotope (i.e., multiply-substituted isotopologues; Δ34C-values) analyses performed on ThermoFisher 253 Plus stable isotope-ratio mass spectrometers at ETH Zurich (Switzerland) and Heidelberg University (Germany). At Heidelberg University, all samples were digested manually, using phosphoric acid for 10 min and subsequently cleaned through a Porapak trap.88 Typical measurements consisted of eight acquisitions with ten cycles per sample acquisition. The empirical transfer function (ETF) to project the samples in the carbon dioxide equilibration scale (CDES89) was determined utilizing a combination of equilibrated gases and internal carbon standards. The data from Heidelberg are presented for a reaction temperature of 90 °C. Standard deviations (SD) of the analyzed samples were 0.007–0.019‰. At ETH Zurich, 100–120 µg aliquots of the dolomite samples were reacted in a Kiel device with phosphoric acid at 70 °C, cleaned by transfer through a Porapak trap held at −40 °C and analyzed in LID1 mode with 400 s integration time. All data are presented in the CDES for a reaction temperature of 70 °C and were calculated with the IUPAC parameters and the revised values of the ETH standards used for normalization71 using the software “Easotope.”

Carbonate precipitation temperatures (see Supplementary Table 2) were calculated for dolomite with published Δ34C–relationships for samples reacted at 90 °C (Heidelberg University)72 and at 70 °C (ETH Zurich).73 The use of calibrations that were determined at exactly the same temperature and same procedures during acid reaction avoids uncertainties due to acid fractionation corrections. For aragonite, we applied Δ34C–relationships published elsewhere96, consistent with previous approaches on erzbergite formations at Erzberg97.

A full description of the analytical methods used is provided elsewhere.95,96 A summary of the mineralogical, geochemical, and isotopic compositions of the authigenic carbonate fracture mineralizations, as well as of the local carbonate host rocks at Erzberg is reported in Supplementary Table 4.

Data availability Raw data are provided in the Supplementary Information of this Article, and are available in the Zenodo repository, https://zenodo.org/record/4061740#.X3W_yqu2xU-U.
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Author contributions
A.B. wrote the manuscript. A.B., A.Z., M.D., and R.B. provided the hydrogeochemical and mineralogical data. F.M. and C.G. conducted the EMP analyses. S.M.B. and T.K. provided the clumped isotope data. K.L. and X.W. measured the U/Th ages. D.H. provided the $87Sr/86Sr$ data, and A.L. measured the $\delta^{13}C$ and $\delta^{18}O$ values. F.M. and R.B. discovered the dolomite. All authors contributed to the writing of the manuscript.

Competing interests
The authors declare no competing interests.

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