Auriferous pyrite formed by episodic fluid inputs in the Akeshi and Kasuga high-sulfidation deposits, Southern Kyushu, Japan

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Abstract
Pyrite geochemistry has proven useful for tracking changes in the composition and physico-chemical conditions of hydrothermal fluids in ore-forming environments. Here, we investigated the microtextural features and chemical composition of pyrite, a main Au-bearing phase in the Akeshi and Kasuga deposits (Southern Kyushu, Japan), to better constrain the ore-forming processes in these high-sulfidation epithermal Au deposits. Despite the widespread distribution of Au-bearing pyrite in both deposits, no visible Au minerals coexist with pyrite. However, in situ laser ablation inductively coupled plasma mass spectrometry results show that Au concentrations in pyrite vary from below the detection limit to 41 ppm and are positively correlated with Cu ($r = 0.4$; up to 7400 ppm) and Bi concentrations ($r = 0.44$; up to 640 ppm). In both deposits, high Cu and Au concentrations occur in small (< 25 μm) anhedral grains of pyrite, which are interpreted to have rapidly crystallized from the ore-forming hydrothermal fluid. In addition, dissolution–reprecipitation textures and thin, concentric, Cu-rich overgrowths were identified in a number of larger (> 25 μm) pyrite grains and aggregates. These abrupt changes in the trace element compositions of pyrite grains likely record episodic metal-rich fluid inputs. We also propose that gold adsorption onto growing pyrite surfaces played a key role in the mineralization of these deposits.

Keywords Epithermal Au deposits · Pyrite geochemistry · LA-ICP-MS · EPMA · Nansatsu district

Introduction
Southern Kyushu, Japan, is known for its intense volcanic–hydrothermal activities related to the subduction of the Philippine Sea plate over the past 5 million years (Watanabe 2005; Garwin et al. 2005). Abundant epithermal Au mineralization is associated with this volcanism, including all four currently mined epithermal Au deposits in Japan (Watanabe 2005; Garwin et al. 2005). Active hydrothermal systems, including Satsuma-Iwojima (Shinohara et al. 1993; Hedenquist et al. 1994a) and Kirishima (Hedenquist and Aoki 1991), have been studied recently, plus the Quaternary
Hishikari low-sulfidation Au deposit (e.g., Izawa et al. 1990; Faure et al. 2002; Morishita et al. 2018; Fig. 1). Other studies have focused on the Pliocene high-sulfidation Au deposits in the Nansatsu district, southern-most Kyushu. Earlier studies have reported ubiquitous pyrite in these deposits (Tokunaga 1954; Nakamura et al. 1994; Hedenquist et al. 1994b) as an important host of primary Au (Imai et al. 1996; Morishita et al. 2019). However, there has been few studies on the microtextural characteristics and the trace element composition of pyrite in these deposits.

Detailed microtextural observations of pyrite combined with in situ geochemical analyses have proven useful in unraveling Au mineralization processes in ore-forming environments and, particularly, for tracking changes in the composition and physico-chemical conditions of hydrothermal fluids (e.g., Deditius et al. 2008; Reich et al. 2013; Tardani et al. 2017; Román et al. 2019). This is because pyrite (i) is ubiquitous and stable over a wide range of physico-chemical conditions in ore deposits (Deditius et al. 2008, 2014; Reich et al. 2013) and (ii) can incorporate a variety of metals and semi-metals (Cu, Co, Pb, Sb, As, Ag, Au, Ni, Zn, Se, Te, and Hg, among others) either in solid solution or as micro- to nano-sized inclusions depending on its formation conditions (Deditius et al. 2011, 2014; Román et al. 2019). Moreover, pyrite has been recognized as an important host of Au in ore deposits because pyrite can absorb and incorporate Au from fluids undersaturated with respect to native Au, increasing its scavenging potential in ore systems (Reich et al. 2005; Cook et al. 2009; Gopon et al. 2019; Pokrovski et al. 2019; Kusebauch et al. 2019).

Here, we present the microtextural occurrence and in situ chemical analyses of pyrite in the Akeshi and Kasuga high-sulfidation epithermal Au deposits. Microtextural observations using scanning electron microscopy (SEM) were integrated with major, minor, and trace element analyses of pyrite performed by electron probe microanalysis (EPMA) and laser ablation inductively coupled plasma mass spectrometry (LA-ICP-MS). By combining these data, we contribute additional information on the hydrothermal activities that led to Au mineralization in the Nansatsu district.

**Fig. 1** Location of the Akeshi and Kasuga deposits, Kagoshima, Japan
Deposit geology and samples

There are several Au-bearing silicified rocks and high-sulfidation epithermal Au deposits in the southern part of the Satsuma peninsula in southern Kyushu, Japan (Nansatsu district; Fig. 2). These deposits, including Akeshi and Kasuga, are referred to as the Nansatsu-type deposits. The district is also considered to have potential for porphyry copper mineralization, although no deposit has been discovered to date (Izawa and Hayashi 2018; Watanabe et al. 2018). Alunite K-Ar ages and paleomagnetic data from previous studies indicate an approximate age of 4 ± 1 Ma for the formation of the Nansatsu-type deposits (Izawa et al. 1984; Takeda et al. 2001; Watanabe 2005).

Gold mineralization in the Nansatsu district was closely associated with the eastward-shifting calc-alkaline volcanism in this region (Izawa 1992; Watanabe 2005). The Kasuga deposit is hosted by the Upper Formation of Nansatsu Group, which comprises pyroxene andesite lavas and pyroclastic rocks, whereas the Akeshi deposit is hosted by the Nansatsu Middle Volcanics, composed of hornblende andesite lavas and pyroclastic rocks (Nakamura et al. 1994; Metal Mining Agency of Japan 1993).

Total gold production in the Akeshi deposit has reached 12.7 metric tonnes (t) since 1912 (average Au grade of 5.8 g/t; H. Kono per. commun. 2020) and 10.7 t since 1910 in the Kasuga deposit (average Au grade of 2.7 g/t; K. Shinagawa per. commun. 2020). Hydrothermal activity contemporaneous with volcanism resulted in strong leaching of the volcanic host rocks by acidic fluids (Izawa 1992; Hedenquist et al. 1994b). The preferential dissolution of phenocrysts and pumice fragments produced a vuggy texture in residual quartz (Hedenquist et al. 1994b). Secondary deposition of quartz is common in vugs and veinlets, locally resulting in massive siliceous rock (Hedenquist et al. 1994b). The ore zone, with gold grades of 1 g/t or more, is restricted to mushroom-shaped residual quartz bodies (Online Resource 1, Figs. S1 and S2) on the order of 100 m wide, not including the advanced argillic alteration zone around the orebody (Tokunaga 1954). Published works provided detailed descriptions of the hydro-

Fig. 2 Geological map of the Nansatsu district. Modified from the Ministry of International Trade and Industry (1985)
thermal alteration and gangue minerals associated with the orebody (Tokunaga 1954; Nakamura et al. 1994; Hedenquist et al. 1994b).

Limited fluid inclusion studies have been conducted in the Akeshi and Kasuga deposits due to the small size of quartz grains. All these analyses were performed on secondary quartz, formed after the initial leaching event (Hedenquist et al. 1994b; Nakamura et al. 1994). Homogenization temperatures range from 160 to 350 °C, and the occurrence of vapor-rich inclusions suggests that fluids were locally trapped from boiling solutions (Takenouchi 1983; Nakamura et al. 1994; Hedenquist et al. 1994b). Generally, the salinities are low (< 1 wt% NaCl$_{eq}$) in both deposits, but higher salinities—based on the presence of daughter minerals—were reported in one sample from the Kasuga deposit (Hedenquist et al. 1994b).

The Akeshi and Kasuga ore zones typically comprise around 90 wt% SiO$_2$ as fine-grained quartz (Nakamura et al. 1994; Fig. 3). The pyrite content of the ore is reported to increase with depth, partly reflecting a decrease in post depositional oxidation (Hedenquist et al. 1994b). Today, pyrite is ubiquitous throughout the orebody, possibly because the upper part of the orebody has already been completely mined. Alunite and native sulfur are also common in the orebody (Hedenquist et al. 1994b; Nakamura et al. 1994). According to Imai et al. (1996), the Akeshi and Kasuga ores can be classified into three types: (i) unoxidized pyrite–gold ore, (ii) unoxidized gold–copper ore, and (iii) oxidized high-grade gold ore. The unoxidized pyrite–gold ore type is the most abundant, and is composed of pyrite and rutile disseminated in fine-grained quartz. The unoxidized gold–copper ore, which comprises pyrite, enargite (in Kasuga) or luzonite (in Akeshi), covellite, scorodite, and rare native gold within a fine-grained quartz matrix, is found mainly in the upper section of the Akeshi deposit (Tokunaga 1954; Nakamura et al. 1994). The relation between these first two ore types is not well constrained. The oxidized high-grade Au ore is related to white quartz veinlets with native gold and rust-colored silicified rock with iron oxides (goethite, scorodite, and triphyte), barite, clay, and native gold (Nakamura et al. 1994; Imai et al. 1996). This ore type has the highest Au grades locally reaching 1000 g/t or more (Imai et al. 1996) and is likely related to hypogene and/or supergene oxidation and weathering of unoxidized ore types.

Samples analyzed in this study were collected from the residual quartz orebodies in both deposits. Nine samples were obtained from three different locations in the open pit in the Akeshi deposit, and seven samples were selected from two different areas in the open pit in the Kasuga deposit (Table 1). Sampling locations, which were the active mining sites at the time of sample collection, are shown in Online Resource 1 (Figs. S1 and S2). Pyrite-rich parts of each sample (Fig. 3) was selected to avoid the effect of supergene oxidation and prepared as a 2-cm-diameter polished sections. Analytical methods are reported in the Appendix.
Results

Scanning electron microscope observations

Pyrite is the main sulfide mineral in all samples analyzed (Fig. 4, Table 1). In some samples, pyrite is spatially associated with enargite (Fig. 4a, Table 1) or rutile (Fig. 4b; Table 1), indicating that the studied samples are dominated by unoxidized pyrite–gold ore (Imai et al. 1996). Pyrite has various morphologies, from euhedral crystals (Fig. 4b, d, f) to anhedral aggregates (Fig. 4c, e). Their grain sizes also varied significantly, from a few microns to up to hundreds of microns (Fig. 4). Since pyrite grains are finely disseminated within the vuggy quartz matrix, it was not possible to discriminate pyrite associated with the initial leaching stage from pyrite formed during secondary quartz deposition. Textural features observed in eight out of 16 samples (Table 1, Fig. 5a–c, Online Resource 1 Fig. S3) indicate coupled dissolution–reprecipitation (CDR) processes (i.e., fluid-mediated mineral replacement reactions; Putnis 2009). Two common characteristics of CDR were recognized in the studied pyrite grains: (i) a porous phase that preserves the external morphology of the original grain and (ii) sharp reaction fronts between the primary grain and product phases (Fig. 5c). Some grains have multiple reaction fronts that are parallel to each other (Fig. 5a, b).

Electron probe microanalysis and X-ray maps

EPMA measurements were carried out on primary, inclusion- and pore-free pyrite crystals. All EPMA analyses are reported in Online Resource 2. Cobalt and Bi data are not discussed here, because the measured concentrations may have been affected by spectral interferences from Fe and S, respectively. Overall, trace element concentrations in pyrite as detected by EPMA range from a few wt% to detection limits (~100–900 ppm). Copper, Se, and Sb were detected in more than a quarter of the total analyzed points. For other elements, concentrations were below detection in more than three quarters of the analyzed points. Copper showed the greatest variation (median: 0.06 wt%), ranging from a maximum concentration of 3.9 wt% to below the detection limit (251 ppm). Selenium also showed variation (median: 0.06 wt%), ranging from 0.6 wt% to below the detection limit (216 ppm); higher Se concentrations were mainly attributed to two samples from the Kasuga deposits (samples KA15 and KA19). The maximum concentration of As, a key element commonly contained in high abundance in pyrite from a large variety of Au deposits, was 0.4 wt% (median: 0.05 wt%).

WDS X-ray maps for selected elements in pyrite grains are shown in Online Resource 3. Due to the relatively high detection limits of the EPMA technique, no clear zonation was observed in the WDS maps for most elements. However, Cu

| Sample name | Deposit | Orebody | Depth (mL) | Pyrite | CDR | Rutile | Enargite |
|-------------|---------|---------|------------|--------|-----|--------|----------|
| AK01 01-2   | Akeshi  | #1      | 15         | +      | +   | +      |          |
| AK01-9      | Akeshi  | #1      | 15         | +      | +   | +      |          |
| AK02-03     | Akeshi  | #2      | 5          | +      | +   | +++    | +        |
| AK02-06     | Akeshi  | #2      | 5          | +      | +   | +      |          |
| AK04-0      | Akeshi  | #4      | 20         | +      |     |        |          |
| AK04-04     | Akeshi  | #4      | 15         | +++    | +   | +      |          |
| AK15112705  | Akeshi  | #4      | 10         | +      | +   |        |          |
| AK15112706  | Akeshi  | #4      | 10         | +++    | +   | +      |          |
| AK15112710  | Akeshi  | #4      | 20         | +      |     |        | +        |
| KA05        | Kasuga  | Honko   | 75         | +      | +   | +      |          |
| KA08        | Kasuga  | Honko   | 75         | +      | +   | +      |          |
| KA09        | Kasuga  | Honko   | 75         | +      |     |        |          |
| KA10        | Kasuga  | Honko   | 75         | +      |     | +++    | +        |
| KA15        | Kasuga  | Honko   | 75         | +      | +   | +      |          |
| KA19        | Kasuga  | Honko   | 75         | +      | +   | +      |          |
| KA27        | Kasuga  | Honko   | 130        | +      |     | +++    | +        |

CDR: coupled dissolution and reprecipitation
+ : the mineral phase or the feature is observed in the sample
+++ : the mineral phase or the feature was very common in the sample
is an exception and shows sharp compositional contrasts (Figs. 6 and 7). For example, zones enriched in Cu were observed in 14 out of 21 EPMA pyrite maps. These include thin (< 10 μm) Cu- and Co-rich rims in pyrite grains (e.g., Fig. 6b, d), as well as Cu-rich zones with porous, CDR-like textures (Fig. 7a, b).

LA-ICP-MS trace element analysis

We conducted LA-ICP-MS spot analyses on primary, inclusion- and pore-free pyrite crystals, as well as in pyrite aggregates and pyrite grains showing porous CDR textures. Thus, our LA-ICP-MS data represent an “efficient microsampling of small volumes” (Cook et al. 2009) of pyrite in the Akeshi and Kasuga deposits, and hence, we interpret the laser ablation signal profiles considering contributions from elements incorporated in solid solution and as (micro-/nano-)inclusions within pyrite.

A statistical summary of LA-ICP-MS analyses is shown in Fig. 8 (results are listed in Online Resource 4). Copper, Ni, Pb, and Co were above detection in more than three quarters of the analyzed spots. Among these four elements, Cu showed the highest concentration (above detection median of 138.5 ppm; maximum: 7400 ppm). Arsenic, Au, Bi, Tl, Te, and Sb were detected in more than half of the spots. The maximum concentration of As (6500 ppm) was comparable to that of Cu, but the overall concentrations were lower than Cu (above detection median of As concentration was 61.5 ppm).

Scatter diagrams of selected elements are shown in Fig. 9. Gold shows positive correlations with Bi ($r = 0.44$) and Cu ($r = 0.4$; Fig. 9a). Bismuth, Te, and Sb, on the other hand, display positive correlations between each other (Fig. 9b, d). Cobalt also shows a positive correlation with Ni ($r = 0.57$).
Arsenic and Tl did not show clear correlations with any of these elements. Correlation coefficients of Cu, Ni, Pb, Co, As, Au, Bi, Tl, Te, and Sb are shown in Online Resource 1 Fig. S4.

Discussion

Polymetallic Au-bearing nanoparticles in pyrite

Our LA-ICP-MS depth profiles suggest that Au in the analyzed pyrites is present both in solid solution and as submicron polymetallic Au-bearing inclusions (Figs. 10 and 11). Although LA-ICP-MS depth profiles alone cannot confirm the presence of metal nanoparticles, the composition of the apparent inclusions (discussed below) seems to be similar to those reported from other high-sulfidation epithermal Au deposits (Deditius et al. 2011).

Gold and Ag signals appear at about half of the ablation time, coinciding with increasing Cu and Bi signals (gray band in Fig. 10a, at ~20–35 s; see Online Resource 1 Fig. S5 for BSE image of this analysis spot). This profile indicates that Au is distributed heterogeneously in pyrite, probably forming clusters of Au–Ag–Bi–Cu-bearing nanoparticles.

Profiles of Au and Te (Fig. 10b) display matching sharp peaks (gray bands). For example, at ~35 s, signal peaks of Au and Te correspond to peaks in the Cu and Sb profile. These observations suggest the presence of Au–Te and Au–Te–Sb–Cu nanoparticles at the analyzed spot. These nanoparticles might be related to CDR processes (Cook et al. 2009; Deditius et al. 2011; Tooth et al. 2011), since several CDR-like textures are observed near this spot (See Online Resource 1 Fig. S6 for BSE image of this analysis spot).

The presence of Pb and Sb–Au nanoparticles is inferred based on similar patterns (gray bands; Fig. 11b). These nanoparticles may also have been formed via CDR processes, although there is no conclusive evidence—for instance, preservation of the external morphology of the parent phase—to support our suggestion that porous zones in the corresponding spot (Fig. 11a) were actually formed via CDR processes.
Au–As relation in pyrite

Gold and As in pyrite commonly show a positive correlation in many hydrothermal Au deposits (Reich et al. 2005; Deditius et al. 2008, 2014; Morishita et al. 2018), which has led previous studies to propose a key role of As to facilitate Au incorporation into the pyrite structure (e.g., Kusebauch et al. 2019; Pokrovski et al. 2019). The Au–As correlation in pyrite was also examined by Morishita et al. (2019) for the Akeshi and Kasuga deposits. Based on 21 secondary ion mass spectrometry (SIMS) analyses, they concluded that the Au–As relationship is not clear.

Our significantly larger dataset (617 LA-ICP-MS analyses) complements a previous study of the Akeshi and Kasuga deposits (Morishita et al. 2019), allowing a broader evaluation of the Au–As relation. Our LA-ICP-MS data for Au and As in pyrite (Fig. 12) are consistent with concentrations reported in previous SIMS analyses (gray area in Fig. 12, from Morishita et al. 2019). Gold and As do not show a clear correlation, which may be due to the variable forms of incorporation of these two elements in the studied pyrite grains. Gold incorporation has been thought to be enhanced by As\(^{1-}\), which substitutes for S in pyrite (e.g., Fleet and Mumin 1997). Recently, more oxidized forms of As in pyrite, such as As\(^{3+}\), which substitutes for Fe (Deditius et al. 2008), and As\(^0\), which exists as “liquid-like” inclusions (Deditius et al. 2009b), have also been reported in pyrite from other high-sulfidation deposits (e.g., Pueblo Viejo, Dominican Republic). As\(^{3+}\) is reported to correlate with Au in the Yanacocha high-sulfidation epithermal gold deposit in Peru (Deditius et al. 2008), but it remains unclear whether this correlation truly reflects the ability of As\(^{3+}\) in pyrite to enhance Au incorporation (i.e., higher pyrite-fluid partition coefficient for Au). Instead, it may simply reflect the composition of the mineralizing fluid and/or the depositional environment (e.g., pressure, temperature) as suggested by several other studies (e.g., Reich and Becker 2006; Deditius et al. 2009a; Reich et al. 2013; Tardani et al. 2017; Román et al. 2019; Herazo et al. 2021). Since we do not have...
clear evidence for As speciation in pyrite in the studied deposits, we consider it difficult to evaluate the role of As in Au uptake in the Akeshi and the Kasuga deposits. Future spectroscopic studies (e.g., synchrotron XANES and EXFAS) should provide further evidence on As oxidation state(s) and incorporation into pyrite (e.g., Simon et al. 1999a; Savage et al. 2000; Manceau et al. 2020) and its influence on Au enrichment.

**Cu enrichment events recorded in pyrite**

**Cu as an indicator of changes in ore-forming environments**

In the Akeshi and Kasuga pyrites, the Cu content shows a larger variation than other elements. In addition, Au and Bi contents show positive correlations with Cu content (Fig. 9). Here, we discuss the Cu concentration and distribution in pyrite and evaluate its use to track changes in elemental supply and/or physico-chemical conditions during the genesis of these deposits.

Copper negatively correlates with the Fe content in pyrite (Fig. 13; $r = -0.52$). This negative correlation between Fe and Cu is a common feature between the Akeshi (orange triangles) and Kasuga (blue circles) deposits. Since inclusion-free pyrite grains were selected for microprobe analyses, the effect of nanoparticles, especially those related to CDR and porous textures, is small. Therefore, simple substitution of Fe$^{2+}$ for Cu$^{2+}$ in pyrite is likely to be the major mechanism for Cu incorporation into the Akeshi and Kasuga pyrite, as has been reported in several hydrothermal deposits (e.g., Pacevski et al. 2008; Reich et al. 2013). However, the negative correlation between Fe and Cu is not necessarily conclusive evidence for the elemental substitution process, as it is common for Cu to occur as both solid solution and micro- to nano-sized inclusions (Pacevski et al. 2008; Reich et al. 2013), and hence, their influence cannot be disregarded.

Previous studies have suggested that the trace element content in the pyrite structure reflects the trace element composition of the pyrite-forming fluid (Deditius et al. 2009a; Reich et al. 2013; Tardani et al. 2017). Based on these findings, the Cu content in the Akeshi and Kasuga pyrites may reflect changes in the Cu content of the ore-forming fluid. High-temperature fluids may be responsible for the Cu enrichment, considering that the solubility of Cu in Cl-bearing hydrothermal fluids is enhanced at higher fluid temperature (Liu and McPhail 2005; Kouzmanov and Pokrovski 2012).

**Rapid crystallization of Cu-rich pyrite**

The trace element composition of pyrite can be influenced by its growth rate, in addition to the composition of the pyrite-forming fluid (e.g., Chouinard et al. 2005a). The fact that relatively large euhedral grains (Fig. 4b, d, f) and aggregates of small (< 25 μm; smaller than our LA-ICP-MS spot) anhedral pyrite grains (Fig. 4c, e) both exist in the Akeshi and Kasuga deposits suggests that the pyrite growth rate fluctuated during ore formation. Among these morphologies, the small grain aggregates are suggestive of rapid pyrite crystallization from the hydrothermal fluid (Huston et al. 1995; Simon et al. 1999b; Román et al. 2019). All the aggregates analyzed...
in this study \((n = 55)\) show higher Cu concentrations (median, 1900 ppm) than the other pyrite types (Fig. 14). Moreover, the range of Cu contents in the aggregates is two orders of magnitude less than in all other pyrite types analyzed herein, which is not observed for other elements (Fig. 14). Therefore, the processes that formed the pyrite aggregates (i.e., rapid crystallization) in the Akeshi and Kasuga deposits likely triggered efficient Cu precipitation.

Fig. 8  Graphical summary of LA-ICP-MS spot analyses \((n = 617)\) of pyrite from the Akeshi and Kasuga deposits. Each boxplot illustrates the minimum, median, and maximum concentrations. The number of analyses above detection limit for each element is labeled above each box.

Fig. 9  Scatter diagrams of \(a\) Cu vs. Au, \(b\) Sb vs. Bi, \(c\) Cu vs. Bi, and \(d\) Sb vs. Te for pyrite from the Akeshi and Kasuga deposits. See text for the discussion.
In addition to Cu, the pyrite aggregates contain higher concentrations of Pb, Sb, Te, Bi, and Au than other pyrite types (Fig. 14). The 1st quartile elemental contents of pyrite aggregates are higher than the 3rd quartile elemental contents of other pyrite types. This may reflect abundant micro- to nano-sized sulfosalt inclusions and polymetallic nanoparticles in pyrite (e.g., Deditius et al. 2011; Ciobanu et al. 2012), probably due to supersaturation of the fluid with respect to these elements. In contrast, the concentration of Co, Ni, As, and Tl in all pyrites are similar (Fig. 14).

Cu-rich overgrowth in pyrite

Sector and concentric zoning in pyrite can be related to two different types of control on trace element incorporation into pyrite (Chouinard et al. 2005a), either stereochemical influence of the crystal surface structure on the incorporation of trace elements, or concentric zoning that is believed to reflect temporal variation of the pyrite-forming fluid condition (e.g., temperature and/or fluid composition) (Chouinard et al. 2005a; Deditius et al. 2009a; Reich et al. 2013; Tardani et al. 2017).

The WDS X-ray elemental maps (Fig. 6) illustrate pyrite with Cu-rich overgrowths. Concentric micrometer-wide zones enriched in Cu were identified in some pyrite grains (Fig. 6b, d). Such sharp transition from the trace element-poor core...
to micron-scale, Cu-rich rim in pyrite from Au deposits in epithermal environments has been suggested to form as a consequence of rapid changes in the composition and/or physicochemical condition of the pyrite-forming fluid (Deditius et al. 2009a; Peterson and Mavrogenes 2014; Sanchez-Alfaro et al. 2016; Tardani et al. 2017; Wu et al. 2019b). Hence, it seems reasonable to assume that the Cu content of the ore-forming fluid in the Akeshi and Kasuga deposits experienced a relatively sudden increase, which we refer to as a Cu enrichment event. Since Au in pyrite positively correlates with Cu (Fig. 9a), the Cu enrichment event is likely related to at least a portion of the Au mineralization in the Akeshi and Kasuga deposits. In addition, the rapid crystallization of Cu-rich pyrite suggested by the anhedral aggregates also seems to be closely associated with the Cu enrichment event.

Replacement of primary pyrite by Cu-rich pyrite

The porous zones in some pyrite grains probably formed via CDR (Fig. 7a), as they usually match the Cu-rich zones identified in the EPMA elemental maps (Fig. 7b). This suggests that primary (early) pyrite was replaced by Cu-rich pyrite via CDR. However, a caveat is that our WDS X-ray maps yield relative concentrations, not absolute values. In addition, we were not able to measure the actual composition of the porous zones because most were not smooth enough for microprobe analysis; also, other mineral inclusions or nanoparticles may also be present (e.g., the extremely Cu-rich area in Fig. 6b).

In the Akeshi and Kasuga deposits, multiple CDR textures can be recognized in a single pyrite grain of several hundred microns. This observation suggests repetitive Cu enrichment events in a relatively short period. Although no detailed data regarding the timescale of pyrite CDR are available, previous experimental studies showed that CDR could be a relatively fast process, on the order of seconds to minutes (Renard et al. 2019) or days to weeks (Putnis 2009).

Implications for the mineralization processes

Possible cause of Cu enrichment events

According to Deditius et al. (2009a), abrupt and intermittent changes in the composition of the pyrite-forming fluid, the Cu enrichment event, are unlikely to occur by wall-rock reaction or mineral deposition; rather, such changes are more likely to occur when a (magmatic) fluid separates into two different fluids or by mixing with another fluid. Within this framework, one possible scenario is intermittent introduction of magmatic fluids into the brittle, lower-pressured domain, where meteoric water circulates at hydrostatic pressure (Fournier 1999). The fluid injection itself can also promote hydraulic fracturing and brecciation in the brittle domain and reduce fluid pressure and temperature, which further enhances phase separation and/or fluid mixing between magmatic and meteoric fluid (Sillitoe 1985; Fournier 1999). Hence, steep pressure and temperature gradients are required to explain the Cu enrichment events. Although direct evidence of magmatic fluid is lacking, most previous studies agree that magmatic fluids supply a major

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![Fig. 12](scatter_diagram_of_As_versus_Au_content_in_pyrites_from_the_Akeshi_and_Kasuga_deposits_black_curve_represents_the_solubility_limit_of_Au_as_a_function_of_As_reich_et_al_2005_gray_shaded_area_is_the_range_of_Au_and_As_concentration_in_pyrite_from_the_Akeshi_and_Kasuga_deposits_reported_by_Morishita_et_al_2019.png)

![Fig. 13](scatter_diagram_of_Fe_versus_Cu_content_wt_in_pyrites_of_the_Akeshi_and_Kasuga_deposits_data_from_electron_probe_microanalysis_EPMA.png)
portion of the Au and Cu in high-sulfidation deposits (Vennemann et al. 1993; Arribas 1995; Heinrich 2005). This interpretation may also be consistent with findings in other hydrothermal Au deposits that pressure-driven, episodic hydrothermal fluid movements in brittle rheological zones are recorded in pyrite grains, sometimes bearing CDR textures (Peterson and Mavrogenes 2014; Rottier et al. 2016; Sanchez-Alfaro et al. 2016; Wüth et al. 2019a).

Role of pyrite in mineralization

Deposition of Au in epithermal deposits is commonly regarded as a consequence of H₂S removal, destabilization of Au hydrosulfide complexes (the dominant Au species in epithermal environments), and supersaturation of the fluid with respect to Au⁰ or other Au minerals (e.g., Drummond and Ohmoto 1985; Krupp and Seward 1987; Stoffregen 1987; Spycher and Reed 1989). Major processes that remove H₂S in epithermal environments include boiling, fluid oxidation, and sulfide precipitation (Pokrovski et al. 2019).

Adsorption on growing pyrite surfaces, a potentially effective mechanism for scavenging Au from a relatively Au-poor fluid, is also proposed to have played a key role in mineralization in other high-sulfidation deposits where pyrite is the main host of Au (Chouinard et al. 2005b; King et al. 2014). Therefore, we suggest that the same mechanism may also have contributed to Au mineralization in the Akeshi and Kasuga deposits. Our findings indicate, at least qualitatively, that the hydrothermal fluid was largely undersaturated (e.g., Simmons et al. 2016 and references therein) with respect to native Au for a considerable period during the mineralization process.

When the fluid is undersaturated with respect to Au, processes operating at the mineral surfaces can play an important role in Au mineralization (e.g., Kusebauch et al. 2019); the Au hydrosulfide complexes in the fluid can be adsorbed onto the growing pyrite surface, and consequently, Au can be incorporated in pyrite either as solid solution or nanoparticles (Widler and Seward 2002; Reich et al. 2005; Pokrovski et al. 2019).

Conclusions

This study combined SEM, EPMA, and LA-ICP-MS analyses to unravel the microtextural characteristics and in situ chemical composition of pyrite, a main Au-bearing phase in the Akeshi and Kasuga high-sulfidation deposits in Kagoshima,
Japan. Overall, pyrite from the Akeshi and Kasuga deposits shows various morphologies, from euhedral crystals to anhedral aggregates of small pyrite grains; the latter indicating rapid crystallization from a hydrothermal fluid. In addition, multiple coupled dissolution–precipitation (CDR) textures are commonly recorded within single pyrite grains a few hundred micrometers in diameter. The CDR textures indicate the replacement of primary pyrite by Cu-rich pyrite.

Several trace elements including As, Au, Bi, Co, Cu, Ni, Pb, Sb, Te, and Tl were detected in more than half of the LA-ICP-MS analyses of all pyrite types. Pyrite aggregates formed by rapid crystallization have higher Cu, Au, Pb, Sb, Te, and Bi concentrations than large, euhedral pyrite grains, whereas Cu-rich overgrowths likely reflect rapid changes in the pyrite-forming fluid composition. In general, the Cu content in pyrite has a large variation compared to other elements and positively correlates with the Au content in pyrite. Most Au in pyrite is likely to exist as solid solution, whereas Au-bearing, polymetallic micron- to nanometer-sized inclusions may be present in some portions of pyrite grains. Gold and As in pyrite of the Akeshi and Kasuga deposits do not show a clear correlation, presumably reflecting multiple valence states of these elements.

Finally, we suggest that episodic input of metal-rich fluid, likely magmatic in origin, may be recorded in pyrite grains as sharp changes in trace element composition. Gold adsorption onto growing pyrite surfaces may also have played a key role in mineralization by scavenging Au from a relatively Au-poor fluid.

**Appendix: Analytical methods**

Microtextural observations of the pyrite-bearing samples were carried out by using a JEOL JXA-8500F scanning electron microscope (SEM) at the Japan Agency for Marine-Earth Science and Technology (JAMSTEC) and a FEI Quanta 250 SEM at the Andean Geothermal Center of Excellence (CEGA) in the Department of Geology, University of Chile, Santiago.

Electron probe microanalysis (EPMA) of pyrite grains (362 spot analyses in total) was performed at the JAMSTEC using a field-emission JEOL JXA-8500F instrument. The beam energy was 20 keV, the beam current was 12 nA, and the beam size was 0.1 to 3 μm. Elements were detected using the following analyzing crystals: LiF for Mn Kα, Fe Kα, Cu Kα, Zn Kα; LiF for Co Kα, Ni Kα; PETJ for S Kα, Pb Mα, Sb Lα; PETH for Bi Mα, Ag Lα; and TAP for As Lα, Se Lα. The standards used (Astimex MINM-25-53 and METM-25-44) included natural and synthetic sulfides, arsenides, tellurides, selenides, and native metals. Wavelength-dispersive spectrometry (WDS) X-ray maps were also acquired, with beam energy 20 keV, beam current 50 nA, and beam size 0.1 to 3 μm.

LA-ICP-MS spot analyses of selected pyrite grains (617 points in total) were conducted using a 193 nm ArF excimer laser (Photon Machines Analyte 193) coupled to a quadrupole ICP-MS (Thermo Fisher Scientific iCAP Q) at the Mass Spectrometry Laboratory of the Andean Geothermal Center of Excellence (CEGA), Department of Geology, University of Chile. The analytical protocol is based on Román et al. (2019), with some modifications. Prior to each analytical session, the ICP-MS was tuned by ablating a NIST SRM 610 glass reference material (i.e., 238U/232Th between 0.95 and 1.05), oxide production (ThO+/Th+ < 0.5%), and double-charged production (22Mg/44Ca++ < 0.01%). Ablation was carried out using a laser pulse frequency of 4 Hz, an energy density of ~1.5 J/cm², and a spot size of 25, 30, or 40 μm. Pure He was used as a carrier gas. Each spot was ablated for 30 s following 30 s of gas background collection. The following isotopes were monitored: 34S, 51V, 52Cr, 53Cr, 55Mn, 57Fe, 59Co, 66Ni, 63Cu, 65Cu, 66Zn, 69Ga, 72Ge, 73Ge, 75As, 77Se, 82Se, 95Mo, 97Mo, 107Ag, 109Ag, 111Cd, 115In, 118Sn, 120Sn, 121Sb, 123Sb, 125Te, 182W, 197Au, 202Hg, 205Tl, 206Pb, 207Pb, 208Pb, and 209Bi, using a total quadrupole sweep time of 0.55 s.

Both external and internal standard calibrations were applied (Longerich et al. 1996). The MASS-1 pressed synthetic sulfide reference material (Wilson et al. 2002) was used as a primary standard, and total Fe concentrations obtained by EPMA were used as the internal standard. Additionally, the GSE-1G glass reference material was employed as a secondary standard using the concentration values reported by Jochum et al. (2005). External standard measurements were performed at the beginning and end of each analysis round of 20 spot analyses. Data integration and reduction was carried out using LAtools, an open-source Python package (Branson et al. 2019), with some modifications. Detection limits were calculated based on the equations of Longerich et al. (1996). For some elements with very low background, values from Heinrich et al. (2003) were used as the detection limit. The MASS-1 reference material may contain some heterogeneities, especially for Au (Wilson et al. 2002). Therefore, LA-ICP-MS depth profiles of the MASS-1 reference material were thoroughly inspected before calibration to avoid introducing Au heterogeneities, and MASS-1 readings were made in duplicate each analysis round. Except for the MASS-1 reference material, we did not exclude analyses affected by possible inclusions.

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Data and materials availability Not applicable.

Code availability Not applicable.

Declarations

Conflict of interest The authors declare no competing interests.

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