Trace element and stable isotope data from a flowstone in a natural cave of the mining district of SW Sardinia (Italy): evidence for Zn\(^{2+}\)-induced aragonite precipitation in comparatively wet climatic conditions

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Abstract:


A speleothem from Crovassa Azzurra, a mine cave in SW Sardinia (Italy), has been analysed for mineralogy, minor and trace elements and stable isotopes. It is composed of layers of primary calcite and aragonite, with a region of secondary calcite. The primary carbonate is strikingly rich in Zn and Pb, presumably as the result of transport in solution from overlying Pb-Zn deposits. Immediately below the transition between calcite and aragonite, concentrations of Zn, Cd and P increase. At the transition between aragonite and Pb-rich aragonite, concentrations of Pb and P increase. Stable isotopes do not indicate an evolution toward drier periods for these two transitions, conversely to what is normally observed in calcite-aragonite speleothems. On the contrary the combined observation of increase in P and metals derived from oxidation of sulphides and the variation of isotopic composition of aragonite and calcite suggests that in this mine cave aragonite was deposited with increasing flowrate and thus more humid conditions. In addition, the effect of Zn\(^{2+}\) or Pb\(^{2+}\) in inhibiting precipitation of calcite appears to have been more important than that of Mg\(^{2+}\).

Keywords: mine cave; speleothems; trace elements; stable isotopes; mineralogy; calcite; aragonite; kinetics; environmental reconstruction

Received 15 March 2011; Revised 23 April 2011; Accepted 15 May 2011

INTRODUCTION

Many natural caves in the Lower Cambrian limestones and dolostones of SW Sardinia (Italy) have been discovered during mine excavations in several Pb-Zn sulphide deposits. These “mine caves” host a wide variety of speleothems, both from a morphological and mineralogical point of view. Besides the rather uncommon cave minerals such as hemimorphite, hydrozincite, barite and cerussite, most of the speleothems are composed of aragonite and calcite. Another anomaly of the region is that aragonite is found here, in the stability field of calcite, in much larger quantities than in normal karstic settings.

Many authors agree that the principal cause of metastable precipitation of aragonite in cave environments is related to the presence of Mg\(^{2+}\) in the percolating fluids (Hill & Forti, 1997 and references therein). The inhibiting effect of Mg\(^{2+}\) on the deposition of calcite has been confirmed by many experimental studies (Murray, 1954; Curl, 1962; Roques, 1965; Bischoff, 1968; Berner, 1975; Morse, 1983; Mucci, 1986; Fernandez-Diaz et al., 1996; Davis et al., 2000; De Choudens-Sanchez & Gonzalez, 2009). On the other hand, many studies have shown that the growth rate of aragonite is not influenced by the presence of Mg in solution, since this element is not incorporated in the aragonite structure (Berner, 1975; Mucci & Morse, 1983; Jimenez-Lopez et al., 2004).

Studies carried out in karst areas have confirmed the role of the Mg\(^{2+}\) ion as inhibitor of calcite precipitation (Cabrol, 1978; Gonzales & Lohmann, 1988; Bar-Matthews et al., 1991; Railsback et al., 1994). Besides, in the majority of cases, the calcite-aragonite transitions in speleothems appear to be related to changes in environmental and climatic conditions towards aridity, since the reduction of water flow feeding the speleothem favors the increase in Mg/Ca (Cabrol, 1978; Gonzales & Lohmann, 1988; Bar-Matthews et al., 1991; Railsback et al., 1994; Frisia et al., 2002; Pagliara et al., 2010). Mg\(^{2+}\), in fact, tends to concentrate in residual solutions, because its partition coefficient is less than 1 with respect to both aragonite and calcite.

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These findings have a general validity, since Mg$^{2+}$ is by far the most abundant solute in karst waters that is able to inhibit the growth rate of calcite in caves. However, many other solutes are supposed to inhibit precipitation of calcite (Cabrol, 1978; Meyer, 1984), even though the details of such inhibition are not well described and understood. Thus, in some particular cases like the mine caves of Iglesiente (SW Sardinia, Italy), where other cations such as Pb$^{2+}$ and Zn$^{2+}$ are present in significant concentrations, the precipitation of aragonite might also be related to other factors inhibiting calcite precipitation.

If the increase in concentration of these metals in the water inside mine caves has an important role in the genesis of aragonite, the appearance of this mineral phase would be related to environmental processes that favor the increase of these solutes in the solutions. These processes, however, can be very different from those that cause an increase in Mg$^{2+}$.

The study of the chemical variation along profiles perpendicular to the growth bands of calcite towards aragonite can be useful in understanding the possible inhibiting role of minor elements on the precipitation of both minerals. Variation in concentration of a chemical constituent in the solid phase, in absence of relevant variations in partition coefficient, reflects change in that constituent’s concentration in the original fluid. At the transition from calcite to aragonite, concentrations in the solid commonly change because of the difference of the partition coefficients for any given element with respect to the two minerals. However, extrapolating that trend into the following aragonite layer allows evaluation of the extent to which that constituent was more concentrated in solutions from which aragonite was formed, relative to the earlier solution from which calcite precipitated.

To investigate the possible role played by different solutes in the genesis of aragonite and to understand the environmental significance of calcite-aragonite banding in caves associated with sulphide mines, a flowstone from the Crovassa Azzurra in the San Giovanni Mine (Sardinia, Italy) has been sampled and studied.

**STUDY AREA**

Crovassa Azzurra is a small natural cave intercepted by the Peloggio mine tunnel at 218 m asl in the San Giovanni Mine, in SW Sardinia (Fig. 1). These underground works have exploited a very rich Pb-Zn deposit of the Mississippi Valley Type since the middle of the XIX$^\text{th}$ century until the late 1980s (Bechstadt & Boni, 1996). The host rock consists of limestones and dolostones of Lower Cambrian age.

The mountain in which the mine was cut hosts over 50 mine caves with no known natural entrance and thus accessible only via the mine voids. The most famous of these is the Santa Barbara cave that, like many other caves of this area, is of hypogenic origin (Pagliara et al., 2010). These caves often contain rather uncommon cave minerals such as hydrozincite, hemimorphite and barite. Many have widespread aragonite speleothems in the form of helicitites, but there are also aragonitic flowstones.

The climate of the area is of the Mediterranean type, with hot and dry summers and relatively wet winters and autumns. Average air temperature in the caves of Mount San Giovanni is around 15.5ºC (Chiesi, 2005).

The Crovassa Azzurra cave consists of two underground rooms connected by a small and narrow shaft, which is dissected by the mine tunnel. Most of the walls in both rooms are composed of a bluish flowstone, composed of an alternation of white-grey-
bluish calcite and bluish aragonite layers. The lower room shows clear evidence of a standing water level, with calcite dogtooth spar at the bottom. The sample SGV9 described and studied in this paper derives from the flowstone in the upper room. The sample SGV9 was taken from a flowstone in the Crovassa Azzurra in the San Giovanni Mine, close to the city of Iglesias in SW Sardinia. Two thin sections (transparent and polished) were prepared for the study via polarising optical microscopy and laser ablation. Samples for X-ray diffraction and stable isotope (C and O) analyses were obtained by drilling at several points using a dental drill.

The mineral phases have been determined using a Panalytical X’Pert Pro X-ray diffractometer (Cu Kα radiation, operating at 40 mA and 40 kV) at the Department of Earth Sciences of the University of Cagliari (Sardinia, Italy).

Carbon and oxygen stable isotope analyses were performed by a method modified from McCrea (1950). Samples weighing roughly 5 mg were reacted under vacuum in 100% phosphoric acid at 50°C. The resulting CO2 was extracted on a conventional vacuum line and analyzed on Finnigan MAT Delta E and MAT 252 mass spectrometers at the Department of Geology of the University of Georgia (Athens, USA). Laboratory standards were prepared and analyzed with each batch of samples. These standards have been calibrated to NBS-19 (δ13C = +1.95, δ18O = -2.2‰ relative to VPDB) and NBS-18 (δ13C = -5.0 and δ18O = -23.0‰ relative to VPDB). Sample isotopic results are normalized to the lab standards using a two-point scale, so that all δ13C and δ18O values herein are reported relative to VPDB. The 2-sigma error of the combined extraction and analysis is 0.04‰ for δ13C and 0.05‰ for δ18O.

The minor and trace elements have been studied along the banding of polished sections using Laser Ablation Inductively Coupled Plasma Mass Spectrometry (LA-ICP-MS). This instrument at the Centro Grandi Strumenti of the University of Cagliari consists of a Perkin Elmer Elan DRC ICP-MS coupled with a New Wave Research Nd:YAG Laser, working at a wavelength of 213 nm. All analyses have been acquired at a frequency of 10 Hz and a spot size of 50 μm. Inside the ablation chamber, He has been used as a carrier gas, mixed with Ar as makeup gas, before entering the ICP system.

In this work the data on eight elements are considered: Mg, P, Cu, Zn, Sr, Cd, Ba and Pb. The calibration strategy was achieved using the NIST 612 glass as external standard and 44Ca as internal standard. For each spot analysis the weight concentrations of the internal standard were previously determined in the Department of Geology of the University of Georgia (Athens, USA) by a wavelength dispersive system of a JEOL JXA 8600 electron microprobe. Data elaboration, regarding the concentrations of the different elements for each spot analysis, has been performed using the software GLITTER (Van Achterbergh et al., 2001). Only the obtained values of concentration over the Minimum Detection Limit, which was always less than 7 ppm for P and less than 1 ppm for Mg, Zn, Pb, Ba, Cd, Sr and Cu, was considered. Quality control of the data was monitored during each analytical session by repeated analyses of a natural standard glass (BCR-2). This standard indicated a precision (RSD%) within 10% for all the investigated elements and better than 3% for Mg, Ba and Sr.

The polished section has also been studied with a Scanning Electron Microscope (SEM) Quanta 200 at the Centro Grandi Strumenti of the University of Cagliari. This SEM is coupled with an Energy Dispersive System (EDS) EDAX with which some spectra have been acquired relative to some specific analyses. The identification of the peaks and the semi-quantitative analyses has been carried out for each spectrum using the elaboration software GENESIS. The observations have been done using electron backscattering and working at low vacuum conditions.

RESULTS

Mineralogical composition

Analysis via X-ray diffraction shows that the sample consists, from bottom to top, of brown and grey layered calcite and then a wide interval of layers of blue, green and white aragonite (Fig. 2). Laterally equivalent to the uppermost layers of aragonite is a region of light and less distinctly layered calcite.

The aragonite can be divided into three sub-layers on the basis of colours and Pb content as determined by SEM-EDX. A middle green aragonite sub-layer (PbA) having some white horizons is much richer in Pb than lower (BA1) and upper (BA2) blue aragonite sub-layers. In backscatter-SEM images (Fig. 3a) the transition from the lower to the middle sub-layer of
aragonite is very evident because of a sharp increase in brightness. Backscatter-SEM images and EDX microanalysis also showed the presence of some cerussite (PbCO$_3$) crystals having a size between 1 and 50 μm (Fig. 3b). The cerussite crystals are enclosed in the white horizons of the PbA aragonite sub-layer, where maximum Pb concentrations in the aragonite matrix are also observed.

In the upper part of the sample, an irregular region of light calcite designated SC is present in an area laterally equivalent to the uppermost aragonite. Petrographic observation of this calcite reveals some aragonite relics and an equant fabric. The SC calcite is therefore interpreted to have replaced some portion of blue BA2 aragonite and to have a secondary origin (Fig. 4a).

Beneath the aragonite, at the base of the sample, a layered region of brown and grey calcite is found and is designated PbC. This calcite has a parallel-columnar fabric typical of speleal calcite and thus is interpreted to be primary (Figs. 4b and 4c).

**Minor and Trace Elements**

**Introduction**

Analyses with LA-ICP-MS reveal the presence of Mg, Zn, Cu, Sr, Ba, Pb, and P as minor and trace elements both in calcite and aragonite. The concentration of these elements is variable along traverses perpendicular to the flowstone layers, and these geochemical variations can provide insight into the mineralogical and environmental changes.

**Secondary calcite**

Secondary calcite (SC) shows lower Mg concentration that the primary calcite (PC). This is probably due to the low Mg content of the precursor aragonite and perhaps also to the slow rate of secondary precipitation, and thus decreased partition coefficient of Mg$^{2+}$ (Railsback, 1999), during the transformation of aragonite to calcite. On the other hand, Sr concentrations are much higher in secondary calcite, probably because they were inherited from the precursor aragonite (Fig. 5). The same holds true, but less strikingly, for Ba (Fig. 5), and presumably for the same reason. Conversely, Zn content is greater in primary calcite but probably only because this element was present in lower concentration in the aragonite precursor to the secondary calcite.

**Primary calcite: traverse X**

The results along traverse X are reported in Fig. 6. In the interval from X1 to X40, the concentrations of Zn are always less than 1.3 mole % in ZnCO$_3$. From X41 to X43 the concentration of Zn increases significantly and, in points X41 and X43, is twice that of the mean value of the first 40 analyses. Zn concentration reaches a maximum value of 3 mole % in the final point immediately before entering the aragonite layer (Fig. 6a). X42, instead, has a Zn concentration similar to that of the first 40 points, but it contains the greatest concentration of Mg (over 5 mole %). In general, however, concentrations in Mg never exceed 4 mole %, and in the last point (X43), where Zn concentration is greatest, Mg concentration is less than 3.5 mole %. In the final part of the traverse, where the variations in Zn are greater, Mg and Zn are negatively correlated (Fig. 7).
In some points along traverse X, large concentrations of Pb are visible (X4, X15, X18, X21, X24, and X27), generally accompanied by large concentrations of Zn. This becomes clear also from Figure 6b, where a correlation between P and Pb is also visible, while Mg seems to be negatively correlated. Maxima in Pb also correspond to maxima in P (X1, X4, X7, X11, X15, X18, X21, X24, X34, X41, and X43). Also Cd increases in the final part, resulting in covariance with Pb, P, and Zn. In general it appears that P and the metals derived from the oxidation of sulphides (Zn, Pb, and Cd) have a similar behavior, and show a negative correlation with Mg.

**Upper primary calcite and aragonite: traverse Y**

The results of traverse Y are shown in Fig. 8. The transition from calcite to aragonite is shown by the interruption of the lines. The behavior of the elements in the first part of this traverse, in the upper part of the primary calcite, confirms the observations done on the terminal part of traverse X. Here again the concentration of Zn increases, reaching its maximum value in the uppermost calcite layer, very close to the beginning of aragonite. Also the maximum Mg concentration is reached at Y8, just below the transition to aragonite, with a slight decrease in the final part (Y9 and Y10) (Fig. 8a). Also, the covariance of P with the metals derived from oxidation of sulphides (Pb, Zn, and Cd) and the negative correlation between these elements and Mg are confirmed (Fig. 8b).

Further along traverse Y, variations in minor and trace elements in the transition from calcite to aragonite are readily explained by the change in mineralogy. At the transition to aragonite, concentrations in Mg, Zn, and Cu decrease, because the relatively small cations of these bivalent elements have smaller distribution coefficients in aragonite than in calcite. Nevertheless, Zn and Cu show minor decreases with respect to Mg, staying at concentrations between 720 and 328 ppm and 179 and 82 ppm, respectively. The concentration of ions with greater radius instead increases, since these are more readily incorporated into aragonite than into calcite. Among these ions are Sr$^{2+}$, which increases at this mineral transition from 60 to 1360 ppm; Ba$^{2+}$, which increases from 68 to 505 ppm; and Pb$^{2+}$, which increases from 138 to 427 ppm. P and Pb persist in covarying in the aragonite layer as they did in the calcite layer. For instance, the rapid increase in P is accompanied by a rapid increase in Pb at the transition from the underlying aragonite layer BA1 to the plumbiferous aragonite above (PbA), at points Y24 and Y25.

**Carbon and Oxygen stable isotopes**

Stable isotope analyses have been carried out along the growth axis (orthogonal to the layers) and are reported in Figure 9. Of the 7 analyses, three (A, B, and C) were in primary calcite, two (D and E) in aragonite without exceptional Pb concentrations and the remaining two (F and G) in the plumbiferous aragonite. Each analysis sampled an area much larger than the laser ablation spots, because they have been obtained by a dental drill taking 10-15 mg of powder, and thus can include multiple LA analysis points.
The three analyses in calcite reveal a small but consistent increase in $\delta^{18}O$. Also $\delta^{13}C$ increases slightly from B to C but decreases between A and B. The increase in $\delta^{13}C$ is much greater from C to D, corresponding to the passage from primary calcite to aragonite. In normal conditions, however, $\delta^{13}C$ of aragonite is greater than that of calcite by 1.68‰, for values close to -10‰ (Romanek et al., 1992). Also $\delta^{18}O$, in normal conditions, suffers a greater fractionation in aragonite with respect to calcite. At a temperature of 15°C this difference is 0.8‰ (Kim et al., 2007).

To be able to compare the isotopic fractionation of the waters giving rise to calcite and aragonite, assuming there have not been sensible temperature variations, it is thus necessary to make a correction. The dashed line in Figure 7 shows these corrected values with respect to the measured ones (full lines). From these corrected values it becomes evident that the mean isotopic composition of aragonite shows lower values in $\delta^{18}O$ and $\delta^{13}C$ with respect to those obtained from calcite, and that this is especially true for the Pb-rich aragonite of isotope samples F and G.

**DISCUSSION**

Change from calcite to aragonite in speleothems is often explained by environmental changes towards aridity, and one controlling factor seems to be the Mg/Ca ratio in the depositing fluids (Railsback et al., 1994; Frisia et al., 2002; Baker et al., 2008). The increase in this ratio can in fact be connected to a decrease in recharge of the speleothem depositing waters. The more residual a water becomes, the more prior precipitation has occurred and the greater the Mg/Ca ratio becomes, since the repartition coefficient of Mg with respect to calcite is <1.

In the flowstone from Crovassa Azzurra, stable isotope analyses of the primary calcite and overlying aragonite do not seem to indicate a change from more humid to arid conditions or reductions of flowrate at the passage between calcite and aragonite. On the contrary, the preliminary isotope data indicate the possibility that aragonite might have deposited in slightly more humid conditions than calcite.

To make a correct palaeoclimate interpretation of these data, from a general point of view at these mid latitudes, presently characterised by semi-arid climate, an enrichment in heavy isotopes along the growth layers of stalagmites or flowstones is normally considered a sign of increase in aridity (Bar-Matthews et al., 1999). An important factor that makes the $\delta^{13}C$ in speleothems increase during less humid periods is the prolonged degassing to which cave waters are subjected (Frisia et al., 2002). If a seeping water, along a certain distance, reaches more rapidly the surface of a flowstone or a stalagmite, it will also have had less time to release CO$_2$ to the cave atmosphere, resulting in an enrichment in the heavy isotope $^{13}C$.

With regard to oxygen, the increase in isotopic ratio $^{18}O/^{16}O$ in the feeding waters is generally associated to a shift to more arid climate, since it is in these conditions that evaporation increases significantly. Evaporation can be extremely efficient, especially during the hot and dry seasons (Bar-Matthews et al., 1999; Denniston et al., 1999). Evaporation can even be more important inside caves during the dry season, because of longer air-water interaction during the pathway of the solution towards the place of deposition. The decrease in $\delta^{18}O$ in the transition from calcite to aragonite does not support the hypothesis of the aragonite genesis in an increasingly arid climate. If we also assume that increases in Mg/Ca ratios are related to decreases in flowrate, the stable isotope data also contradict the hypothesis that aragonite formation was related to the inhibiting effect of Mg$^{2+}$ on calcite formation. The calcite-aragonite transition in the Crovassa Azzurra flowstone is thus not readily explained by changing saturation state and Mg/Ca ratios, and other controlling factors have to be considered.

**The Zn-Pb hypothesis**

A possible explanation for the precipitation of aragonite during an apparent increase of flowrate might be the inhibiting effect of dissolved Zn$^{2+}$ and/or Pb$^{2+}$ on calcite growth. Meyer (1984) showed that small concentrations of dissolved Zn$^{2+}$ and Pb$^{2+}$...
diminish the rate of growth of calcite from solution. Calculations by Meyer (1984) showed that, even at small concentrations, these ions could occupy all of the possible cation sites at steps on the surface of a growing calcite crystal. More recently, Rouff et al. (2005) and Elzinga et al. (2006) likewise concluded that adsorption of Pb\(^{2+}\), Zn\(^{2+}\), and Cu\(^{2+}\) on calcite surfaces can limit mineral-solution exchange and thus inhibit calcite growth. Such ions are relevant because they are relatively soluble in circumneutral oxidizing solutions, whereas Fe\(^{2+}\), for example, is readily oxidized to Fe\(^{3+}\) that then precipitates as a hydrous oxide of some sort and thus, like other trivalent ions (e.g., Al\(^{3+}\) and Sc\(^{3+}\)), is less relevant to solution-calcite interaction. Zn\(^{2+}\) and Pb\(^{2+}\) are especially relevant to the Crovassa Azzurra cave flowstone because they could readily enter solution as the result of oxidative weathering of the sulphides in the mine above the cave.

During the precipitation of calcite, the behavior of Pb\(^{2+}\) should be analogous to that of Mg\(^{2+}\), in that both have partition coefficients in calcite much less than 1. Thus, precipitation of calcite should increase the Pb/Ca ratio of the precipitating solution, as much as it causes an increase of the Mg/Ca ratio. In contrast, the behavior of Zn\(^{2+}\) should be the opposite inasmuch the partition coefficient for Zn\(^{2+}\) in calcite is >1 because the cation size of Zn\(^{2+}\) provides a nearly ideal fit for the calcite cation site (Railsback, 1999). Thus precipitation of calcite preferentially removes Zn\(^{2+}\) from the solution and leads to a smaller Zn/Ca ratio in the residual solution. This may in part account for the inverse correlation of Mg and Zn in the last analyses of Traverse X (Fig. 7). The observed increase in the Zn/Ca ratio of calcite (Fig. 6), and the inferred increase in the Zn/Ca ratio in the solution, may reflect a major input of Zn at its source linked to more humid periods. This would be compatible with the observed isotopic signal.

Evidence of aragonite deposition during wetter conditions

Evidence of a humid period during precipitation of aragonite in the Crovassa Azzurra flowstone may come from the LA-ICP-MS data regarding phosphorus. Huang et al. (2001) have demonstrated that P in speleothems is mainly present as phosphate, and that its concentration increases systematically in response to more humid periods. P, according to these authors, can thus be taken as an indicator of flow rate changes at the depositing speleothem. The covariance between P and the metals related to the oxidation of sulphides (Pb, Zn, and Cd) suggests that oxidative weathering and the transport in solution of the weathering products is more efficient during wet periods, when these metals increase their concentrations in the percolating waters (Fig. 6). Cidu & Frau (2009), in a study of surface waters in mine areas of Sardinia, have demonstrated that the concentration of metals, such as Pb, Zn, and Cd, tends to increase in conditions of major flow. This behavior might also be valid for a karst system.

Further evidence for evolution toward wetter conditions during deposition of the Crovassa Azzurra flowstone comes from the stable isotope data. Both corrected \(\delta^{13}C\) and \(\delta^{18}O\) decreased during aragonite deposition from their corresponding levels during earlier calcite deposition (Fig. 9). Variation in both of these parameters can result from many environmental changes, but the most fundamental control on \(\delta^{13}C\) of spelean calcite is the abundance of overlying vegetation (e.g., Hesterberg & Siegenthaler, 1991; Baldini et al., 2005), with lower values corresponding to more vegetation and thus, in most cases, wetter conditions. Likewise, the major controls on \(\delta^{18}O\) of spelean calcite commonly include amount effects of rainfall (McDermott, 2004) and evaporation of soil water (Bar-Matthews et al., 1996), and in both cases smaller \(\delta^{18}O\) values correspond to wetter conditions. Thus, the C and O stable isotope data reinforce the

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Fig. 6a. Molar percentages of Mg and Zn along traverse X.

Fig. 6b. Comparison in logarithmic scale of concentrations of various minor and trace elements along traverse X.

Fig. 7. Plot of Zn and Mg concentrations in calcite along Traverse X. The various lines for linear regression show that, within the entire set of data, Zn and Mg are positively correlated, but among the analyses richer in Zn and Mg there is an inverse correlation.
P concentration data in suggesting wetter conditions during the precipitation of aragonite, and in particular the Pb-rich aragonite, of the Crovassa Azzurra flowstone.

Origin of the Crovassa Azzurra flowstone aragonite

The considerations above combine to present a strong argument that precipitation of aragonite in the Crovassa Azzurra flowstone was not driven by Mg\(^{2+}\)-inhibition of calcite or by more arid conditions. Instead, multiple lines of evidence suggest that inhibition of precipitation of calcite by Zn\(^{2+}\) and perhaps Pb\(^{2+}\) during wetter conditions allowed precipitation of aragonite. With regard to climate, both the increasing P concentration in calcite underlying aragonite and the lower values of “corrected” \(^{8}\)C and \(^{18}\)O in aragonite relative to calcite suggest a trend to wetter rather than drier conditions during aragonite deposition. The increase in Zn concentration observed in calcite underlying aragonite and the large Pb concentrations additionally suggest greater oxidation of sulphides such as sphalerite and galena in waters descending from the overlying Pb-Zn ore deposits to the cave itself. This argument is supported by the greater concentration of Pb, Zn, and Cd during greater flow of surface waters in the region (Cidu & Frau, 2009).

With regard to inhibiting effects of various cations, the decrease in Mg concentration immediately below the transition to aragonite (Figs. 6 and 8) suggests that inhibition of calcite precipitation by dissolved Mg\(^{2+}\) was not the controlling factor in triggering deposition of aragonite. Instead, the known capacity of Zn\(^{2+}\) and Pb\(^{2+}\) to inhibit precipitation of calcite (Meyer, 1984), combined with observed increases in Zn concentration in the calcite immediately below the transition to aragonite, suggest that these metal cations, and especially Zn\(^{2+}\), were the significant inhibiting agents. This is not surprising in light of the large Zn concentrations in the Crovassa Azzurra flowstone and Meyer’s (1984) observation that Zn\(^{2+}\) and Pb\(^{2+}\) are much more effective inhibitors of calcite precipitation than is Mg\(^{2+}\).

In addition to the cation-driven inhibitions of calcite precipitation considered above, some anions can also inhibit precipitation of calcite. For example, SO\(_4^{2-}\) can inhibit precipitation of both calcite and aragonite, with a greater inhibition of calcite (Walter, 1986). However, SEM-EDX analyses yielded no evidence of S within the flowstone, suggesting that SO\(_4^{2-}\) was of little significance to precipitation of the flowstone. PO\(_4^{3-}\) can also inhibit precipitation of both calcite and aragonite, but it causes a greater inhibition of aragonite (Walter, 1986; Burton & Walter, 1990). Thus, the presence of PO\(_4^{3-}\) in Crovassa Azzurra waters, as inferred from flowstone composition, cannot explain inhibition of calcite coeval with precipitation of aragonite. These thoughts combine to suggest that cations like Zn\(^{2+}\) and Pb\(^{2+}\) were more likely inhibiting agents than the dissolved anions.

The paragraphs above focus on changes within the latest calcite, or differences in isotopic composition between the calcite and aragonite, as evidence of the changes that induced precipitation of aragonite rather than calcite. In addition, a good example of the effectiveness of this unusual model of aragonite deposition comes from stable isotope sample F and the coeval LA-ICP-MS data from analyses 25 to 30 of transect Y. In this interval of aragonite, corrected values of \(^{8}\)C and \(^{18}\)O reach their smallest values, and P and Pb concentrations reach their greatest values. This interval thus epitomizes the characteristics of the aragonite, suggesting deposition in wet, rather than dry, conditions.

CONCLUSION

Multiple lines of evidence combine to suggest that aragonite in a flowstone from Crovassa Azzurra below the San Giovanni Mine in SW Sardinia (Italy) has formed not because of Mg\(^{2+}\)-inhibition of calcite and/
or dry conditions but instead because of inhibition of calcite precipitation by Zn$^{2+}$, and perhaps Pb$^{2+}$, in wetter conditions than those under which calcite previously formed. This interpretation is unusual, because the factors most commonly cited as favoring speleothem deposition of aragonite rather than calcite are presence of Mg$^{2+}$ in drip water, higher temperature, and drier conditions (e.g., Railsback et al., 1994). This unusual interpretation comes from a speleothem exceptional for its large concentrations of Zn$^{2+}$ and Pb$^{2+}$, where Zn$^{2+}$ concentrations by weight in calcite are locally greater than those of Mg$^{2+}$, and where Pb$^{2+}$ concentrations in aragonite are locally greater than those of Sr$^{2+}$. These unusual aragonite and calcite chemistries, and presumably the unusual chemistries of the waters that precipitated the flowstone, are the result of the presence of sulphide mineral deposits in the overlying bedrock. Thus, the findings of this study are not generally applicable to all occurrences of aragonite. However, they may be applicable in other regions where chemistries of karst waters are anomalous, particularly as the result of input from weathering of sulphides. Furthermore, this study provides evidence that many factors can control the polymorphism of CaCO$_3$ in speleothem deposits. These factors should be taken into account when obtaining paleoclimatic information from speleothems.

ACKNOWLEDGEMENTS

This research was funded by Università degli Studi di Cagliari (Italy), as part of a PhD program. The research was produced in collaboration with the Department of Geology of the University of Georgia (USA). The authors acknowledge the Centro Grandi Strumenti di Università di Cagliari for granting access to the use of SEM/EDX.

We also would like to thank Julie Cox, Chris Fleisher and Francesca Podda for their assistance in the laboratory respectively of Stable Isotopes, SEM/WDS and LA-ICP-MS.

Finally, this manuscript was improved by constructive comments from two anonymous reviewers and Giovanni Zanchetta of the Dipartimento di Scienze della Terra di Università di Pisa.

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