Unusual sedimentary geodes filled by gold-bearing hematite laths

M. A. BUSTILLO*†, J. GARCIA-GUINEA* J. MARTINEZ-FRIAS* & A. DELGADO‡

*Departamento de Geología, Museo Nacional de Ciencias Naturales, José Gutiérrez Abascal 2, Madrid 28006, Spain
‡Departamento de Ciencias de la Tierra y Química Ambiental, Estación Experimental del Zaidin, CSIC, Prof. Albareda 1, 18008 Granada, Spain

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Abstract – Unusual mineralized and silicified carbonate-rich geodes were found hosted in Lower Triassic red mudstones in Central Spain. From their borders to their centres, the geodes display (a) a millimetric carbonate crust, (b) a quartz rim, (c) massive dolomite, (d) gold-bearing iron-rich infills (hematite laths and goethite with up to 7000 ppm of gold) and (e) calcite cements that sometimes seal the central hollow. Textural evidence indicates that the geodes were originally anhydrite, which was subsequently replaced by quartz and dolomite. The resultant porosity from this replacement, or by later dissolution, has been filled by epithermal gold-bearing iron-oxide hydroxides, romanechite and calcite.

$\delta^{13}C$ values indicate the participation of meteoric waters in an environment which was characterized by both a sub-desert climate and a temperate–tropical climate. Oxygen signatures reflect very variable temperatures for all minerals, with the exception of calcite, which appears to have precipitated at $< 38 \, ^\circC$. Iron-oxide temperature values can reach up to 85 °C (epithermal stage). The mineralogical assemblages of these unusual geodes denote early diagenetic replacement followed by the epithermal activity. This mineralization is linked to the Late Hercynian, calc-alkaline volcanism of Central Spain (the Hiendelaencina mining district).

1. Introduction

Silicified carbonate-rich nodules with relics of evaporites are hosted in different types of sedimentary rocks that range from Precambrian to Tertiary in age, and reveal diverse diagenetic histories (Chowns & Elkins, 1974; Milliken, 1979; Elorza & Rodriguez-Lazaro, 1984; Maliva, 1987; Ulmer-Scholle, Scholle & Brady, 1993). In Spain, this type of carbonate–silica nodule occurs in the Buntsandstein–Muschelkalk red mudstones (Muñoz et al. 1992; Marfil et al. 1998) from the Central Range. The nodules range from 2 to 20 cm in diameter, and often display sub-spherical shapes and external rounded protuberances, resulting in an appearance resembling the head of a cauliflower.

The nodules in the study area (Fig. 1) occur in Lower Triassic red mudstones. They range from 2 to 12 cm in diameter, and are often hollow, and can be termed ‘geodes’ in the sense determined by Farris (1990). Most geodes are silicified and rimmed by a millimetre-sized carbonate crust (Fig. 2a). The remainder of each geode is made up of a complex assemblage of quartz and dolomite, along with variable, but subsidiary, proportions of iron oxide and hydroxides. Euhedral hematite lath-like crystals have precipitated in the geode core as fibrous–radial aggregates. With the exception of the so called ‘Bristol diamonds or potato stones’ (Tucker, 1976), other Triassic occurrences of comparable diagenetic structures are very uncommon. Besides the remarkable diagenetic associations found in the geodes, the occurrence of iron-rich infills, displaying an anomalous gold content, makes these structures unique and perfect witnesses to the saline diagenesis and mineralizing processes that have occurred in this area.

2. Regional setting

The Triassic red mudstones, where these geodes occur, are located near Tordelrabano (Guadalajara province) in the eastern part of the Spanish Central Range, where the Central and Iberian ranges meet (Fig. 1). Extensional faulting controlled Permian calc-alkaline volcanism (Hernan, Perni & Ancochea, 1981; Doblas et al. 1988) and Late Hercynian epithermal mineralization (Martinez-Frias, 1987).

The volcanism is well represented in the Atienza area (Fig. 1) by the presence of small (up to 1.5 km in diameter) circular or elongated andesite–dacite outcrops (Hernan, Perni & Ancochea, 1981). Its greatest development coincides with the intersection of two systems of extensional faults running northeast–southwest and northwest–southeast, which are contemporaneous (De Vicente et al. 1986) with the faults controlling the main lodes of the mineralized area (the Hiendelaencina mining district; see Fig. 1). At the surface, the andesite–dacite rocks display a late pervasive and propylitic alteration (chlorite and calcite) and
host minor veins of quartz, calcite and barite. Jasperoid silica also fills small pods and vugs. The Hiendelaencina mineralization displays a complex paragenesis, which is rich in base metal sulphides and Ag–Sb sulphosalts. The mineralization is characterized by the presence of quartz–gold veins in the cap of the whole hydrothermal system (e.g. in the Navas de Jadraque area; see Instituto Geológico y Minero de España, 1974; Concha et al. 1992).

Lithologically, the vertical section of the Triassic...
Gold-bearing sedimentary geodes

sediments in the northwestern Iberian Range displays the typical Germanic Triassic Facies, which consists in ascending order of (a) Buntsandstein continental siliciclastics, (b) Muschelkalk carbonate, sand and evaporites of shallow marine origin, and (c) Keuper evaporites, clays and subordinate sandstone of coastal sabkhas. The geodes occur in the red mudstones that are included in the uppermost facies of the Buntsandstein unit (Sopeña & Sanchez-Moya, 1997; see Fig. 1). This whole upper unit extends more than 80 km, and displays variable thickness (30–110 m). It is mainly composed of sandstones and mudstones, along with uncommon beds of fluvial conglomerates. Alonso-Zarza, Soneña & Sanchez-Moya (1998) described dolomitic pedogenic palaeosols in the massive red mudstone of the uppermost sandy braided fluvial system. Marfil et al. (1998) point out the existence of silica and carbonate nodules at the top of the Buntsandstein in zones close to the study area. They also record pedogenic iron crusts, calcretes and silcretes.

According to Ramos, Soneña & Perez-Arlucea, (1986) and Soneña & Sanchez-Moya (1997) these Buntsandstein red beds form the base of a sequence that passes up into siliciclastic and carbonate tidal sediments (the Muschelkalk facies). In short, a mixed-load braided system evolved upwards to an ephemeral stream system. Subsequently, shallow siliciclastic marine conditions transgressed from the east, overlying the fluvial sequences.

3. Methodology

Geodes were selected taking into account the different types of mineralogical zonations. Mineralogy was determined by X-ray powder diffraction using a Phillips powder diffractometer with CuKα radiation. The patterns were obtained by step scanning from 2° to 64° 2θ, in steps of 0.02° with a count of six seconds per step. The texture and chemistry of the authigenic minerals were studied using optical (Figs 2a, b, c) and scanning electron microscopy (SEM) (Figs 2d, e, f) and an electron microprobe (EM) (Figs 2h, i). For SEM studies the specimens were coated with gold (20 nm) in a Bio-Rad SC515 sputter coating unit. General SEM observations were carried out in a Philips XL20 SEM at accelerating voltages of 20–30 kV. Energy-dispersive X-ray microanalyses (EDX) were obtained using a Philips EDAX PV9900 with a light-element detector type ECON. The crystal-chemical characteristics of iron-oxide hydroxides were determined on the basis of a large data series of electron microprobe analyses (Jeol Superprobe JXA-8900M), bulk and channel-selected (TAP, PETJ, LIF, PETH) X-ray spectra search and identification routines. The standards used were natural and synthetic crystals from a collection stored at the Servicio de Microscopia Electrónica Lluis Bru, Complutense University, Madrid.

Isotope analyses were carried out at the Laboratorio de Isótopos Estables of the Estación Experimental del Zaidin in Granada, Spain. All samples were ground to a <200 mesh. Carbon dioxide was generated from the carbonates using 100% phosphoric acid for 12 hours in a thermostatic bath at 25 °C for the calcite and 50 °C for the dolomite (see McCrea, 1950). Quartz samples were treated with cold 1:1 HCl to remove carbonates. Oxygen was extracted from the quartz and the goethite by reaction with ClF3, using the method described by Borthwick & Harmon (1982) and Venneman & Smith (1990). The released oxygen was converted to CO2 by reaction with a hot platinitized graphite rod (Clayton & Mayeda, 1963). The isotope ratios were measured in a Finnigan MAT 251 mass spectrometer. Commercial CO2 was used as an internal standard for the oxygen analysis of silicates. The CO2 was contrasted with the V-SMOW, SLAP and GIPS water standards. The measured δ18O values of the international NBS-30 (biotite) and NBS-28 (quartz) standards were + 5.1 ± 0.20 and + 9.6 ± 0.10 (V-SMOW), respectively. The experimental error found in carbonates (δ13C and δ18O) was < ± 0.050‰ using Carrara and EEZ-1 as internal standards, previously compared with NBS-18 and NBS-19.

4. Composition of the geodes

Nodules and geodes were sampled from the S3M stratigraphic unit (Fig. 1), which is composed mainly of red mudstones. These mudstones are rich in illite, and contain dispersed silt-sized quartz, K-feldspar and mica grains. The geodes are zoned from the border to the centre as follows.

(a) Up to 3 mm of carbonate crust (Fig. 2a). This is composed of wavy laminae of micrite (Fig. 2b), and includes claystones and detrital grains from the host rock. Most of the micrite laminae are recrystallized to sparite, partially masking the initial texture.

(b) Quartz rim from 2 to 5 mm. This penetrates the interior of the geodes through small fissures and cracks. The rim is composed of megaquartz crystals (100 µm to 2 mm in diameter), which occasionally contain inclusions of anhydrite (Fig. 2c). The anhydrite inclusions are either disorganized within the megaquartz crystals or parallel to the outer of the euhedral quartz crystals. The presence of anhydrite inclusions is evidence that the geodes had an anhydrite precursor.

(c) The core of the geodes is composed mostly of euhedral dolomite crystals that are 80 µm to 3 mm in diameter and sometimes also contain anhydrite inclusions (Fig. 2c). Some dolomite crystals have been replaced by calcite. Euhedral calcite cements are present in the centre of some geodes. The textural relationship between dolomite and quartz, where isolated quartz crystals grew in dolomite mosaics, could indicate that the dolomite was substituted by the quartz.
Figure 2. Photographs of the textural and compositional details of geodes. (a) View of a geode exterior. Note the remains of the millimetric carbonate crust (C) on the right. (b) Thin section (parallel nicols) of the outer part of the geode with a carbonate crust (C), quartz rim (Q) and dolomite (D). The carbonate crust is composed of wavy micrite laminae. Quartz crystals (Q) are idiomorphic and could replace dolomite crystals (D). (c) Thin section (crossed nicols) of the geode core: quartz and dolomite contains anhydrite relics. (d) Aggregates of hematite laths observed under the SEM. (e) Hematite laths show zonation (sandwich
However, the absence of dolomite inclusions in the megaquartz indicates that these textures more likely formed by the partial replacement of the original anhydrite nodule by quartz or by the subsequent replacement of residual anhydrite by dolomite, leaving the original quartz crystals intact.

5. The gold-bearing iron infills

The iron-rich infills occur as radial–fibrous crystals and spheres, which are sometimes enveloped by calcite. The infills are composed of lath-shaped crystals, which are often texturally zoned and show dissolution edge pits (Figs 2d, e, f). Minor romanechite [(Ba, H₂O)₂(Mn⁴⁺, Mn³⁺)O₁₀] is also present as millimetre-sized fibres (Fig. 2g). Diffraction patterns of the lath crystals show diffraction lines of two mineral phases: hematite (2.68₆₀–2.51₅₀–2.28₅₀, etc.) with Rietveld refined parameters of a:5.033 Å and c:13.749 Å and goethite (2.4₈⁶–4.1₈⁶, etc.) with cell parameters: a:4.596 Å, b:9.957 Å, c:3.021 Å.

Electron microprobe analysis of both laths (hematite, α-Fe₂O₃) and small crystal aggregates (10–80 µm in diameter) found on laths (goethite, α-FeO(OH)) indicates near-pure compositions that fit well within the theoretical stoichiometric formulae. Nevertheless, small amounts (never more than 1 wt %) of some elements (Co, Au) were detected. The experimental formula indicates the following mean average composition: Fe₁.₉₈Au₀.₄₈Co₀.₂₁O₃. Gold values range from 0.24 to 0.71 wt % and cobalt from 0.17 to 0.33 wt %. It is important to stress two points. Firstly, both elements display an erratic chemical distribution in the hematite host crystals, and this is verified by both specific point analysis and detailed 12 hour chemical mapping (Figs 2h, i); secondly, no geochemical anomalies in these elements were detected in the quartz grains.

The combination of the detailed close-up SEM–EDX images and the electron microprobe results reveal the following features: (a) The laths are ended by {021} faces. (b) ‘Sandwich’-zoned hematite crystals contain different layers (black, grey and white; Fig. 2e); these differences are not reflected by chemical variations. (c) Goethite and native gold patches are present in the hematite crystals. (d) There are etch-dissolution pits on {021} faces, and a series of rough-edged steps on the basal {001} planes. (e) Late deposition of romanechite needle aggregates has occurred; these are spread over the hematite crystals, partially covering the edges and apices of the crystals.

6. Carbon- and oxygen-isotopic composition

Oxygen- and carbon-isotope ratios were measured for the three types of carbonate occurrences in the geodes: calcite crust, massive dolomite and euhedral calcite crystals that partially fill the geode core (Fig. 3). The δ¹³C values ranged from −5 to −9.50 ‰ (PDB) and the δ¹⁸O values ranged from +23.70 to +25.70 ‰ (SMOW). The δ¹⁸O values of quartz samples ranged from +26.7 to +28.80 ‰ (SMOW) (Fig. 3). The δ¹⁸O values of the iron-rich infills of hematite ranged from +1.1 to –1.60 ‰ (SMOW).

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Figure 3. Oxygen- and carbon-isotope ratios in different sections of and different minerals in the geodes.
Negative $\delta^{13}C$ values indicate significant contributions of pedogenic carbon, and, therefore, the participation of meteoric waters. This same interpretation was proposed by Morad et al. (1992) for similar values ($-6$ to $-9.50$‰) in other carbonate cements in Triassic Buntsandstein sandstones from Central Spain. Dolomite $\delta^{13}C$ values range from $-5$ to $-6.70$‰ (PDB), which indicates a significant contribution of atmospheric CO$_2$. In as much as the atmospheric $P(CO_2)$ was likely to have been 3–7 times higher during the Upper Triassic and Lower Jurassic than at present (Berner, 1990), the $\delta^{13}C$ values of the geodes indicate that they were formed in areas characterized by climates between sub-desert and temperate-tropical (Cerling, 1991).

During the Triassic, the area studied in Central Spain was at a low latitude (Ziegler, Scotese & Barret, 1983). At present, the isotopic composition of rainwater at latitudes between 0° and 15° typically ranges between $-0$ and $-3$‰ (SMOW) (Rozansky, Araguas & Gonfiantini, 1993). Ocean waters during the Triassic appear to have been 2.3‰ more negative than those of the present day (Scherrer, 1977). Therefore, meteoric water derived mainly from the evaporation of Triassic marine waters would probably have been between $-2$ and $-6$‰ for these latitudes. The equilibrium temperature and water isotopic-ratio curves for the $\delta^{18}O$ for each mineral of the geodes are plotted in Figure 4 to estimate the theoretical precipitation temperatures. Assuming pore-water $\delta^{18}O$ values of between $-2$ and $-6$‰ (SMOW), the oxygen-isotopic data indicate that calcite precipitated at temperatures < 38 °C, and that the rest of the minerals, especially the iron oxides and hydroxides, formed at higher temperatures.

7. Discussion and genetic model

The cauliflower shape of the geodes, the presence of anhydrite inclusions and the occurrence of the geodes in continental red mudstones suggest the possibility that these geodes initially may have been anhydrite nodules that grew in a continental sabkha environment. The high evaporation rates in sabkha environments create favourable conditions for the formation of anhydrite nodules, which may later be replaced by carbonate and silicified (Chowns & Elkins, 1974; Tucker, 1976). However, Elorza & Rodriguez-Lazaro (1984) and Maliva (1987) indicate that silicified anhydrite nodules cannot be used as indicators of a sabkha environment. These researchers conclude that anhydrite nodules only display evidence of hypersaline pore waters during early diagenesis.

In the study area, geodes were generated after several replacement, dissolution and cementation processes. If the anhydrite dissolution was faster than the precipitation of silica and/or carbonate minerals, a central hollow formed that later allowed the euhedral crystals to grow. An alternative hypothesis is that anhydrite remained in the nodule core after quartz and dolomite replacements terminated, and the remaining anhydrite was later dissolved. The resultant porosity in the core of geodes was either preserved or was filled by other mineral phases (diagenetic and epithermal). A theoretical genetic sequence for the Tordelrabano geodes is shown in Figure 5.

The isotopic composition of $\delta^{13}C$ in the dolomite and calcite crystals indicates that the replacement of anhydrite by dolomite and the precipitation of calcite likely took place in equilibrium with meteoric water. The temperatures of precipitation that were estimated from the $(\delta^{18}O/\delta^{16}O)$ ratios of all minerals in the geodes (Fig. 4), with the exception of calcite, do not indicate whether the origins were diagenetic or hydrothermal. In addition, the participation of marine waters during diagenesis should not be disregarded, because shallow siliciclastic marine deposits (Muschelkalk) overlie the fluvial sequences of the study area. If marine pore waters were present then the isotopic data from the goethite can be interpreted as indicative of a temperature of up to 116 °C. Finally, the
carbonate crust that rims the geodes can be interpreted as a pedogenetic crust linked with the general process of palaeosol formation in the continental red mudstones (Alonso-Zarza, Sopeña & Sanchez-Moya, 1998). The isotopic composition of this crust, displaying the most negative $^{13}$C values, could indicate a significant edaphic influence (Salomons & Mook, 1986).

Several possible explanations can be proposed for the gold anomalies. These include mechanisms of concentration during weathering and diagenesis (Mann, 1984; Benedetti & Boulegue, 1991) or mineralizing hydrothermal processes. In the study area, no references exist regarding gold in the Triassic detrital host rocks or other underlying formations. Nevertheless, this precious metal is common in epithermal quartz veins in the Hiedelaencina mining district. Concha et al. (1992) stress the importance of the Atienza volcanics as a potential target for hidden Au–Ag epithermal mineralization.

Interpretation of the metallogenetic episode in geodes suggests the following possible hypogenic sequence: (1) precipitation of gold-bearing hematite laths, and (2) subsequent dissolution of the hematite (etch pits) and precipitation of iron hydroxide as goethite. Both processes can be explained by the decrease in pH of the mineralizing fluids (see Maurice et al. 1995). The hydroxylation of hematite laths can produce both the dissolution features and the epitaxial overgrowth of goethite on the hematite $\{001\}$ planes. From a crystallographic point of view, the growth of goethite can be explained in terms of the atomic configuration of the hematite $\{001\}$ and the goethite $\{001\}$, which match reasonably well (Barron et al. 1997). The tracing of dissolution steps on the $\{001\}$ plane and the etch pits on the $\{021\}$ faces display the characteristic flat shapes of the hematite hydration–hydroxylation. The adsorption of water withdraws the Fe$^{3+}$ ions approximately 0.18 Å with respect to the superficial Fe positions (Becker, Hochella & Apra, 1996). A final supergenic stage was responsible for the late formation of calcite and romanechite.

From a metallogenetic point of view, the unusual geode infills represent the last gold- and iron-rich episode that is linked to the Late Hercynian volcanic-related mineral deposits in Central Spain (the Hiedelaencina mining district). These deposits are especially rich in silver and base metals, and have been assigned to the epithermal type (Martinez-Frias, 1987; Martinez-Frias et al. 1988; Doblas et al. 1988) using the classification of Eimon (1981). It is important to note that gold has also been reported in other nearby mineralized areas (e.g. the gold–quartz veins of La Nava de Jandraque), which share the same ore-bearing epithermal system.

In a regional metallogenetic scenario, this mineralizing episode would correspond to an almost-barren late episode, given that the epithermal fluids lost their ore charge during the three previous precious- and base-

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**Figure 5. Genetic model of the Tordelrabano geodes.**

Abbreviations: AN, anhydrite; D, dolomite; Fe, oxides and hydroxides of iron; AC, carbonate crust; Q, quartz; V, voids; CA, calcite; CM, mudstones. (a) Formation of anhydrite nodules in the mudstones during early diagenesis from hypersaline fluids. Later, a crust (probably of edaphic origin) formed around the anhydrite nodule. (b) Replacement of the anhydrite by dolomite and quartz under continental groundwater. Formation of voids by dissolution of anhydrite or dolomite. (c) Filling of the voids formed in the nodules by hydrothermal minerals (gold-bearing hematite–goethite). Subsequent precipitation of calcite under meteoric water.
metal-rich stages (Martinez-Frias, 1987). Nevertheless, small amounts of gold, barium, manganese and iron would still have been present in the mineralizing system, resulting in the precipitation of gold-bearing hematite, goethite and romanechite in different pulses in the geode cores, making up the cap of the whole system (Lower Triassic).

According to the fluid inclusion studies carried out by Sierra et al. (1988) and Concha et al. (1992) this episode is characterized by mineralizing solutions that show both monophase (liquid) and two-phase (vapour + liquid) aqueous fluid inclusions, and that evolved following a decreasing trend in temperature and salinity ($T: 260–90 \degree C$, and from 18 to 7 eq. wt% NaCl). These same authors also indicate that one of the main features of this last mineralization episode is that it was marked by complex interactions between fluids of a different nature. Two sets of sulphur-isotopic data were reported for the whole hydrothermal system: (1) $\delta^{34}S$ values around +6‰, and (2) $\delta^{34}S$ values around 25‰, which probably indicates two different sulphur sources and supports the hypothesis of mixing of fluids (Concha et al. 1992).

Considering these epithermal conditions, the discovery of the mineralized geodes in Lower Triassic sediments indicates (Concha et al. 1992) the participation of Triassic sediments during the last metallogenetnic stage. The crystallization sequence of a primary gold–hematite assemblage followed by secondary goethite matches the experimentally demonstrated evidence that whereas gold does not affect the formation of hematite {001} surfaces: applications to the interpretation of STM images and heterogeneous surface reactions. American Mineralogist 81, 1301–14.


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