ABSTRACT: Burial diagenesis of the Upper Jurassic Torrecilla Reef Complex is recorded by a complex paragenetic sequence initiated with emplacement of ferroan calcite cements and followed by the precipitation of ferroan saddle dolomite. Ferroan calcite cements contain micro-sized dolomite inclusions (MDIs). Elemental and isotopic compositions of MDIs are virtually indistinguishable from those of ferroan saddle dolomite cements. In contrast to other scenarios for the formation of microdolomite inclusions that invoke either a magnesian calcite precursor or incomplete dedolomitization, the paragenetic relations of MDIs to their host ferroan calcite and their geochemical composition implies formation by fine-scale burial replacement of ferroan calcite by fluids associated with the emplacement of ferroan saddle dolomite.

Petrographic observations and mass-balance considerations suggest that it is unlikely that Fe and Mg incorporated into ferroan dolomite could have been derived from the cannibalization of the reefal host rock. Rather, dolomitizing fluids were likely associated with the hydrothermal, low-grade metamorphic event that affected the Cameros Basin during the middle to Late Cretaceous. Interaction of these fluids with preexisting burial cements produced MDIs in the ferroan calcites.

INTRODUCTION

The occurrence of microdolomite inclusions (MDIs) in calcite crystals has been widely reported in the literature, and two explanations have been invoked for their mode of formation and geologic significance: (1) MDIs have been interpreted as evidence for a high-magnesium calcite precursor that was subsequently converted into low-magnesium calcite and MDIs (e.g., Lohmann and Meyers 1978; Meyers and Lohmann 1978; Leutloff and Meyers 1984; Mazzullo 1994; Tobin and Walker 1998; Dickson 2001) or (2) MDIs are relics of a dolomite precursor that has undergone partial calcitization (e.g., Evamy 1963; Goldberg 1967; Munn and Jackson 1980; Frank 1981; Budai et al. 1984; Jones et al. 1989; Rossi and Cañaveras 1999).

Micro-sized dolomite inclusions in ferroan calcites of the early Kimmeridgian coral reef sequence (Cameros Basin, Northern Spain) (Fig. 1), however, cannot be explained with either of these hypotheses. Their occurrence in ferroan calcite cements, which are interpreted to have precipitated during progressive burial, and the absence of evidence to suggest association with HMC precursors or intervals of dedolomitization, requires an alternative mechanism for their formation. In the Kimmeridgian sequence, it is possible to reconstruct the process by which MDIs formed, as much as they represent an intermediate paragenetic stage in a sequence of burial cements. Although occurring as intracrystalline euherdral to anhedral inclusions within ferroan calcite, their association with saddle dolomite cement, which postdates and overlie ferroan calcite, suggests an origin unrelated to the ferroan calcite cement which hosts them. Rather, MDIs may have formed through a process of partial replacive dolomitization coincident with the emplacement of the saddle dolomites during the late stages of burial and the low-grade metamorphism that affected deposits of the northwestern Cameros Basin (Fig. 1A).

To test this hypothesis, we employ detailed petrographic and fine-scale elemental and isotopic analyses of burial ferroan calcite and later saddle dolomite to define the sequence and timing of their formation, the nature and source of the diagenetic fluids, and the process by which these MDIs were formed.

METHODS

A total of 150 samples were used for the petrographic and geochemical study of sedimentary and diagenetic features of the Torrecilla Reef Complex, utilizing standard petrographic techniques, cathodoluminescence, SEM petrography, and elemental and isotopic analyses. For each sample, a polished 30 μm thin section and matching 150–200 μm thick section for microsampling was prepared. Cathodoluminescence (CL) examination was carried out using a Technosyn® cold cathodoluminescence unit operating at 14–17 kV with 350–450 μA beam current. CL microphotographs were performed using 800–1600 ASA, 35 mm slide film. Following examination with CL, all thin sections were stained with Alizarin Red S and potassium ferricyanide (Dickson 1966). For SEM studies, thin sections were etched with 0.2% HCl solution for 1–2 minutes.

Depositional micritic matrix and seven different diagenetic phases were then microsampled from thick sections using the microscope-mounted drilling system and 3 to 30 spot analyses were performed per phase. Sample areas ranged from 300 to 500 μm in diameter, resulting in the recovery of 100–150 μg powdered samples. Analyses for δ13C and δ18O values were performed in the Stable Isotope Laboratory at the University of Michigan. All sample powders were roasted in vacuo for one hour at 200°C, to remove volatile organic contaminants, then reacted at 73°C in an automated carbonate reaction system (CarboKiel-I) coupled directly
to the inlet of a Finnigan MAT 251 gas ratio mass spectrometer. Isotopic ratios were corrected for $^{17}$O contribution and are reported in per mil notation relative to the VPDB standard. Values were calibrated utilizing NBS 19 as the primary standard, and analytical precision was monitored by daily analysis of NBS powdered carbonate standards. Measured precision was maintained at better than 0.1% for $\delta^{13}$C and $\delta^{18}$O.

Elemental analyses for Ca, Mg, Sr, Mn, and Fe were performed on an electron microprobe with an accelerating voltage of 15 kV and a spot size of 5 µm. Measured precision was maintained at better than ± 0.14% for Mg, ± 0.09% for Ca, ± 0.08% for Sr, ± 0.08% for Fe, and ± 0.06% for Mn, and the detection limits were 100 ppm for Mg, 250 ppm for Sr, 200 ppm for Mn, and 250 ppm for Fe. Seventeen samples were analyzed, and 12 to 98 spot analyses were performed per phase. To allow direct comparison with isotopic analyses, all elemental analyses were made on the same thick sections.

In order to isolate MDIs from host calcite cement for isotopic analysis, four sample powders of calcite cement containing MDIs were taken from thick sections using the microscope-mounted drilling system. A 1M solution of acetic acid and sodium acetate with a pH of 5 was prepared (see Jackson 1969). Each powdered MDI/calcite sample was placed into the solution for 48 hours, and the undissolved powder was then analyzed by XRD to be sure that all calcite was dissolved before isotopic analyses of residual MDIs.

GEOLOGIC SETTING AND STRATIGRAPHIC FRAMEWORK

The study area is located in the Cameros Basin (Fig. 1), a rapidly subsiding extensional and continental basin developed during the Late Jurassic and Early Cretaceous, which was filled from the Tithonian to middle Albian by up to 5 km of fluvial and lacustrine deposits with rare marine incursions (Mas et al. 1993; Guimerá et al. 1995; Arribas et al. 2003). These deposits overlie to 1.5 km of Triassic and Jurassic rocks (Mas et al. 1993; Guimerá et al. 1995). The basin-fill succession embodies a large cycle or megasequence overlain by an intra-Albian unconformity.
that corresponds to the base of the Upper Cretaceous post-rift megacycle (Mas et al. 1993, Mas et al. 2003; Salas et al. 2001). The basal portion of the Upper Cretaceous megacycle consists of upper Albian sandstones overlain by large Upper Cretaceous carbonate-platform deposits (Alonso et al. 1993; García et al. 2004). The Cameros Basin was subsequently tectonically inverted during Paleogene to middle Miocene alpine contraction (Guimerá et al. 1995; Mas et al. 2003).

One of the main features of this basin is that a large proportion of the Upper Jurassic–Lower Cretaceous deposits of the northwestern area were affected by low- to very low-grade metamorphism (Guiraud and Seguret 1985; Casquet et al. 1992; Mas et al. 1993; Mas et al. 2003) (Fig. 1A). This metamorphism is manifest microscopically by the appearance of chloritoid (an index mineral of low-grade metamorphism) and slaty cleavage oblique to bedding in fine-grained incompetent layers. Pyrite ore deposits in the eastern part of the Cameros Basin are also associated with this metamorphism (Alonso-Azárate et al. 1999b; Alonso-Azárate et al. 2001b).

On the basis of a detailed study of the mineralogical, petrological, geochemical, and structural attributes of the basin, this metamorphic episode has been characterized as hydrothermal and allochemical (Casquet et al. 1992; Barrenechea et al. 1995, 2000; Barrenechea et al. 2001; Alonso-Azárate et al. 1995, 2005; Alonso-Azárate et al. 1999a, Alonso-Azárate 1999b, Alonso-Azárate 2001a, Alonso-Azárate 2001b; Manilla-Figueroa et al. 1998; Mas et al. 2003). Microthermometry of fluid inclusions, radiometric data, mineral assemblages (chloritoid–chlorite at peak metamorphism), crystallochemical properties of phyllosilicates, chlorite microthermometry, isotopic thermometry, and geochemistry of pyrite deposits indicate that: (1) metamorphism displays clear thermal inversions across areas of rapid deposition; (2) metamorphic grade is more influenced by the changes in rock permeability and composition than by burial depth; (3) metamorphism postdated the filling of the basin (post-rift metamorphic ages range from 106 to 86 Ma (late Albian–Coniacian); and, (4) the metamorphism ranged from low-grade (epizone) to very low-grade (anchizone) conditions, with maximum temperatures and pressures of 350 to 370°C, and 1 kbar, respectively.

This study focuses on the early Kimmeridgian succession, the youngest Jurassic marine unit, which underlies synrift Tithonian to Lower Cretaceous deposits, and is represented by coral-reef complexes of the Torrecilla en Cameros Formation. Specifically, we have examined the Torrecilla Reef Complex, exposed just southwest of Logroño in Northern Spain (Fig. 1B). The reeval succession exposed in this area consists of offlapping reefs, up to 35 m thick, that developed during relative sea-level fall (Alonso et al. 1986–1987). The Torrecilla Reef Complex was buried by up to 4 km of Tithonian to Aptian fluvial and lacustrine deposits, and upper Albian to Upper Cretaceous coastal and marine sandstones, limestones, and dolostones (Alonso et al. 1993; Mas et al. 1993; Gómez-Fernández and Meléndez 1994; García et al. 2004). The contact between the reeval and continental units is a major paleosol-associated erosional unconformity surface.

Fig. 2. Schematic representation of the paragenetic sequence of early (from Benito et al. 2005) and late (this study) diagenetic phases within the Torrecilla Reef Complex. Numbers indicate the order of precipitation of the different diagenetic phases (1, oldest; 11, youngest).

Fig. 3. Cathodoluminescence (CL) photographs showing the early and burial diagenetic phases. A) Coral moldic pore cemented by early diagenetic nonluminescent columnar calcite (CC), followed by brightly luminescent slightly ferroan calcite (IC), a dullly luminescent ferroan calcite with brighter luminescent areas which correspond to regions with abundant inclusions (FC) and nonluminescent ferroan saddle dolomite (FD). B) Primary porosity was first cemented by an early diagenetic generation of zoned nonluminescent and bright luminescent nonferroan drusy calcite cement (NFC). NFC was corroded (blue arrow), and then overlain by a bright to dull slightly ferroan calcite (IC) and dark luminescent ferroan calcite (FC), which occludes the pore. C) Burial diagenesis phases in a coral moldic pore. The corroded surface (white arrow) of a nonluminescent ferroan saddle dolomite (FD) is overlain by a thin generation of zoned red luminescent and dark luminescent nonferroan syntaxial dolomite (SD) followed by a nonferroan, brightly luminescent equant calcite (EC). D) Ferroan calcite cement (FC) in a coral moldic pore. Ferroan calcite is affected by a fracture that is filled by ferroan saddle dolomite (FD). Cloudy areas in ferroan calcite (FC) and early calcite cements (ECC) contain abundant micro-sized dolomite inclusions (MDIs). Note that some of the cloudy areas are preferentially concentrated and aligned parallel to the fracture (red arrows), and cut across the brownish growth zones of the calcite crystals (black arrows). E–F) Stained burial diagenetic phases. E) Ferroan calcite cement (FC) is corroded (red arrows), and then overlain by ferroan saddle dolomite (FD). Note the MDIs inside ferroan calcite cements (black arrows). F) MDIs (black arrows) are preferentially concentrated in the areas next to the corrosional boundary between ferroan calcite and ferroan dolomite cements (red arrow). G) Ferroan calcite cement (FC) containing MDIs (red arrows) is followed by ferroan saddle dolomite (FD) after a corrosion surface (yellow arrows). H) Same photograph as part G under polarized light. Note that most of MDIs (red arrows) are in perfect crystallographic continuity with host ferroan calcite. The largest dolomite inclusion (white arrow), however, is in crystallographic continuity with the surrounding ferroan saddle dolomite.
MICRO-SIZED DOLOMITE INCLUSIONS DEVELOPED DURING BURIAL DIAGENESIS

A

B

C

D

E

F

G

H

JSR
Fibrous calcite and peloidal micrite (Fig. 2) were precipitated syndepositionally in the reefs and were followed by early diagenetic phases during exposure to meteoric waters as detailed by Benito et al. (2005). Early diagenetic processes that affected the reefal complexes were controlled by rising sea level and local tectonism that gave rise to alternations of submergence and reefal exposure; while exposed, reef corals were neomorphosed and dissolved. When submerged, this secondary porosity was then filled by nonferroan columnar calcite cement precipitated from marine-derived waters (Figs. 2, 3A). During the latest stages of reef development, the rate of sea-level rise could no longer keep pace with tectonic uplift, leading to exposure, the development of an extensive paleosol, and the precipitation of nonferroan drusy meteoric calcite cement (Figs. 2, 3B). Subaerial exposure of the reef complex continued until the onset of subsidence and continental sedimentation in the Tithonian.

PETROGRAPHIC OBSERVATIONS AND GEOCHEMISTRY OF LATE DIAGENETIC PHASES

Early diagenetic cements are postdated by a sequence of late carbonate cements that largely filled remaining primary and secondary porosity (Figs. 2, 3). A common feature of these phases is that δ13C values are significantly more negative in samples collected in the upper part of the Torrecilla Reef Complex, compared to those obtained from the lower and middle part (Fig. 4). This pattern of variation of δ13C values is similar to that seen in the composition of pre-burial host-rock carbonates, including reefal carbonates and early marine-derived and meteoric cements, which also record progressive upsection depletion in δ13C values that has been related to early exposure and alteration of the reef complex (Benito et al. 2005). Specifically, columnar marine-derived cement (Figs. 2, 3A), which is more abundant in the lower and middle part of the sequence (first 25–30 m), defines a field of isotopic values that range from −1.2 to −3.3‰ in δ18O and from +2.0 to +3.0‰ in δ13C (Fig. 4). The isotopic values of drusy meteoric cements (Figs. 2, 3B), which preferentially precipitated in the upper part of the sequence (last 5–10 m) in association with the subaerial exposure of the reef complex, define a field centered about −3.5‰ δ18O and 0‰ δ13C (Fig. 4). The isotopic values of drusy meteoric cements (Figs. 2, 3B), which preferentially precipitated in the upper part of the sequence (last 5–10 m) in association with the subaerial exposure of the reef complex, define a field centered about −3.5‰ δ18O and 0‰ δ13C (Fig. 4). These latest phases precipitated in pores and fractures that developed in response to the Tertiary Alpine contraction (Benito et al. 2003; Benito and Mas 2005).

An important accessory component in this sequence of late diagenetic phases is dolomite that occurs as micro-sized inclusions within ferroan calcite and, less commonly, in slightly ferroan and earlier calcite cements (Figs. 2, 3C), fluorate, sphalerite, celestine, barite, nonferroan saddle dolomite, and solid hydrocarbons. These latest phases precipitated in pores and fractures that developed in response to the Tertiary Alpine contraction (Benito et al. 2003; Benito and Mas 2005).

The first phase that precipitated after early columnar and drusy calcite cements was a slightly ferroan calcite cement (Fig. 2). This cement exhibits a bright orange luminescence (Fig. 3A, B) and fills fractures that crosscut early diagenetic phases. The boundary between early cements and slightly ferroan calcite is sharp (Fig. 3A) or is defined by a corrosion surface (Fig. 3B). Slightly ferroan calcite is followed by ferroan calcite cement that is darkly dull luminescent (Figs. 2, 3A, B) with a transition that ranges from gradual to sharp. In some cases, slightly ferroan cement is absent, and ferroan calcite is the first burial cement after early diagenetic phases. Both slightly ferroan and ferroan calcites contain little Sr (up to 0.10 and 0.11 mole % SrCO₃, respectively), or Mn (up to 0.11 mole % MnCO₃, respectively) (Table 1). Mg and Fe contents progressively increase from the slightly ferroan to ferroan calcite cements (Table 1). MgCO₃ contents of slightly ferroan calcites range from 0.30 to 1.94 mole %, and FeCO₃ ranges 0.03 to 0.63 mole %. In contrast, ferroan calcite MgCO₃ contents range from 0.28 to 2.2 mole % and FeCO₃ ranges from 0.5 to 3.0 mole %. Isotopic compositions of these cements also show a trend towards progressively more negative δ18O values and slightly more negative δ13C values (Fig. 4). Slightly ferroan calcite cements range from −4.4 to −6.0‰ in δ18O values and from +2.4 to −4.8‰ in δ13C values. The δ18O values of ferroan calcite are more negative and variable than those of the slightly ferroan calcite: δ18O values of the early stages of growth of ferroan calcite, which are next to the slightly ferroan calcite, are more positive (−5.8 to −7.0‰) than the later stages of ferroan calcite cement (−6.9 to −8.4‰). In addition, the δ13C values of the early stages of growth of ferroan calcite are slightly more positive (+1.9 to −2.8‰) than the later stage (+1.2 to −3.7‰). (Fig. 4).

Ferroan calcite cement is followed by unzoned, nonluminescent ferroan saddle dolomite (Figs. 2, 3A, C); an irregular corrosional contact separates these two phases (Fig. 3E–H). Ferroan saddle dolomite, with curved crystal and cleavage faces and sweeping extinction in cross-polarized light, occurs as cement infilling the remnant fabric-specific pores and fractures (Figs. 2, 3A, D), and as replacive patches, up to 10 cm wide, in reeval host rock. The abundance of this dolomite phase is minor, at most constituting 2% of the total reeval rock volume. Although part of the ferroan dolomites contain sufficient Fe to be considered ankerites, they have Fe contents ranging from 3.9 to 20.5 mole % (Table 1). However, Fe content varies by no more than 4 to 5 mole % within any single crystal and exhibits no discernible temporal or spatial trends in concentration. Mg ranges from 27.4 to 44.8 mole %, and Ca from 47.7 to 55.7 mole %. Ferroan saddle dolomite cements contain little Mn (up to 0.2 mole %) and Sr (up to 0.08 mole %) (Table 1). δ18O values are quite constant, ranging from −5.1 to −6.4‰, whereas δ13C values are variable, ranging from +2.1 to −3.7‰ (Fig. 4).

When occurring as cement, the ferroan saddle dolomite can be overlain by a syntaxonal nonferroan saddle dolomite that is zoned with alternating red, dark brown, or no luminescence (Figs. 2, 3C). This nonferroan saddle dolomite overlies ferroan saddle dolomite along an irregular and corrosional boundary, with the red-luminescent dolomite typically penetrating the underlying nonluminescent ferroan dolomite (Fig. 3C). Nonferroan syntaxonal dolomite cement contains little Fe (0.7 to 4.1 mole %), and very low Mn (up to 0.09 mole %) and Sr (up to 0.07 mole %) (Table 1). Mg ranges from 55.3 to 50 mole %, and Ca from 48.1 to 51.4 mole %. The δ18O values range from −4.9 to −5.8‰, and δ13C values range from −0.6 to −1.1‰ (Table 1).

The youngest diagenetic phases in this reeval unit include trace amounts of nonferroan equant calcite cement (Figs. 2, 3C), fluorite, sphalerite, celestine, barite, nonferroan saddle dolomite, and solid hydrocarbons.
MDIs have elemental and isotopic compositions very similar to those of ferroan saddle dolomite cement (Fig. 4; Table 1). Fe contents are moderate to high, ranging from 3.1 to 18.2 mole %; Mg ranges from 30.7 to 46 mole %, and Ca from 48.1 to 57 mole %. Mn contents are low, ranging from 0.03 to 0.2 mole %, and Sr contents are very low, ranging up to 0.11 mole %. Isotopic compositions are relatively invariant in $\delta^{18}O$, and range from $-6$ to $-6.2\%$. The $\delta^{13}C$ values range from $+0.4$ to $-1.2\%$. Importantly, isotopic values of MDIs are quite
similar to those of ferroan saddle dolomite occurring in the same samples (Table 2).

**DISCUSSION: EVOLUTION OF THE DIAGENETIC SYSTEM DURING BURIAL**

**Slightly Ferroan and Ferroan Calcite Cements**

Petrographic, elemental, and isotopic compositions of the slightly ferroan and ferroan calcite cements suggest that they precipitated during progressive burial of the reefal unit. This is supported by their elevated Mg and Fe contents and by δ¹⁸O compositions that gradually decrease from starting δ¹⁸O values of early marine-derived and meteoric cements (mean values = −2.3%, and −3.5%, respectively) to the progressively more negative δ¹⁸O values of slightly ferroan calcite, early ferroan calcite, and late ferroan calcite (mean values = −5.3%, −6.2%, and −7.8%, respectively) (Fig. 4). These elemental and isotopic changes can be compatible with a progressive decrease in the Eh of burial pore waters and a concomitant increase in temperature, respectively (e.g., Dickson and Coleman 1980; Meyers and Lohmann 1985; Choquette and James 1990).

Carbon isotope compositions of slightly ferroan and ferroan calcites also show a general trend toward slightly more negative values. For example, the heaviest δ¹³C values of slightly ferroan calcite, early ferroan, and late ferroan calcite progressively decrease from +2.4‰, to +1.9‰ and +1.2‰, respectively (Fig. 4). This change is also compatible with cementation during progressive burial (e.g., Dickson and Coleman 1980; Choquette and James 1990). More importantly, the δ¹³C values of ferroan calcite cements are significantly more negative in the upper part of the reefal unit compared to cements occurring in the middle and lower parts (Fig. 4). We interpret this stratigraphic trend to be inherited from the δ¹³C values of host reefal carbonate whose composition reflects earlier stages of alteration. In response to late Kimmeridgian subaerial exposure and localized input of highly ¹³C-depleted soil–gas CO₂, meteoric alteration gave rise to δ¹³C compositions in the upper part of the reef, which are significantly more negative than host-calcite compositions of the middle and lower reefs (Benito et al. 2005) (Fig. 4). Progressive increase in δ¹³C values with stratigraphic distance from the surface of meteoric recharge is a trend predicted for increased rock–water exchange reactions (Allan and Mathews 1982; Lohmann 1987). Inheritance of these host-rock δ¹³C values by later burial ferroan cements requires dissolution of some of the preexisting host rock and equilibration of fluid δ¹³C values to local host rock values (i.e., in a rock-buffered system with respect to carbon), while a pattern of progressive decrease of δ¹⁸O values within the burial cements requires a water-dominated diagenetic system with respect to oxygen.

**Formation of Micro-Sized Dolomite Inclusions**

The occurrence of MDIs as a phase developed within the burial paragenetic sequence provides an opportunity to determine the conditions for their formation. Furthermore, examination of their geochemical and petrographic characteristics allows the elucidation of the microscale process associated with their emplacement. Prior models for the creation of MDIs within preexisting carbonates include the diagenetic stabilization of a HMC precursor in a relatively closed intracrystalline system (e.g., Lohmann and Meyers 1977; Meyers and Lohmann 1978) and the preservation of dolomite relics during dedolomitization (e.g., Goldberg 1967; Frank 1981; Jones et al. 1989). The MDIs in ferroan calcite cements cannot be accounted for by either of these processes.

It is not reasonable to invoke a HMC precursor for slightly ferroan and ferroan calcite cements within which the MDIs are developed. HMC calcite cements typically precipitate in the marine environment, and it is through their transformation during early diagenesis that they are converted to stable mineralogical phases of LMC and microdolomite. The microdolomites formed by this mechanism are typically 10–30 microns in size and occur within host cements that exhibit fabrics characteristic of neomorphic replacement. In the Cameros Basin, MDIs are highly variable in size (1–300 µm) and develop both as irregular anhedral patches and as fine euhedral inclusions (Fig. 5). Moreover, MDIs occur in the ferroan calcite host cements that show no petrographic evidence of recrystallization and are clearly not former marine precipitates.

Six lines of evidence indicate that it is also unreasonable to invoke a process of incomplete calcitization (dedolomitization) of an earlier dolomite cement: (1) During dolomite calcitization, the dolomite...
precursor is dissolved and replaced by calcite that fills the available space; thus dolomite relics are irregular and anhedral. In contrast, individual crystals of MDIs of this study commonly show euhedral to subeuhedral faces (Fig. 5). (2) Luminescent zonation and paragenetic relations clearly indicate that slightly ferroan and ferroan calcite precipitated as primary cements filling free space, not as a replacement of a dolomite precursor. Ferroan calcite crystals display euhedral growth zones that nucleated at the margins of primary and secondary pores, which, in turn, were mostly created during the early stages of meteoric diagenesis (Fig. 3A, B). Moreover, the external surfaces of ferroan calcite crystals exhibit evidence of corrosion in contact with ferroan dolomite (Fig. 3D–H), suggesting a process of calcite alteration prior to or during ferroan dolomite precipitation, not the opposite. (3) MDIs do not occur in samples where ferroan saddle dolomite is absent and ferroan calcite is the latest cement-filling porosity (Fig. 3B). This suggests that the MDIs appearance is directly related to occurrence of ferroan dolomite. (4) In most cases, the...
extinction of MDIs is the same as that of the host calcite crystals, suggesting that MDIs grew in optical continuity with the host calcite crystals, and thus, they were formed after precipitation of ferroan calcite. (5) MDIs have been observed along cleavage planes within the ferroan calcite (Fig. 5E, F) and along shear planes that cut across growth zones of the ferroan calcite crystals (Fig. 3D). Therefore, MDIs necessarily precipitated after the host calcite. (6) Significantly, the elemental and isotopic compositions of most MDIs are quite similar to those of the ferroan saddle dolomite (Fig. 4, Table 2). In this sense, it is difficult to argue that an earlier dolomite, with a specific geochemical composition, precipitated after the early diagenetic cements and was subsequently partially to completely replaced by the ferroan calcite. This, in turn, was followed by precipitation of the ferroan saddle dolomite in the remaining porosity, which coincidently has the identical geochemical elemental and isotopic composition of the calcitized dolomite.

Thus, rather than a dolomite precursor undergoing partial replacement by ferroan calcite, it is more plausible that it was the ferroan calcite that was altered and replaced by the MDIs. On this basis, we propose that MDIs formed through the partial dissolution of the preexisting ferroan calcites during the emplacement of the ferroan saddle dolomite. The nature of this replacement varied. Larger and irregular patches of dolomite occur in areas affected by extensive corrosion, such as adjacent to the exterior of the calcite crystals and/or in shear planes (Figs. 3D–H, 5). These larger crystals probably fill irregular microporosity. It is more difficult to explain the development of the intracrystalline MDIs by such a simple dissolution process, because these occur as small subhedral to euhedral crystallites isolated from the margins of the host crystals. The pathways by which dolomitizing fluids penetrated into the interiors of the crystals would presumably be controlled by the network of connected micropores, cleavage, minute fractures, and shear planes in the calcite crystals. At sites where dolomite nucleated, continued growth in response to very small differences in solubility of ferroan calcite and ferroan dolomite could explain their euhedral shapes and their isolation as discrete inclusions. This process, as described by Martin et al. (1986) for the progressive microscale replacement of aragonite by calcite, would proceed across a thin fluid film separating the two carbonate phases, and calcite dissolution and dolomite precipitation would occur at such a small scale that substantial voids were not produced.

**Saddle Dolomites**

Ferroan saddle dolomite formed after partial corrosion of the ferroan calcite (Figs. 2, 3), in fractures, or as a phase that replaced the host rock. The development of these features suggests that the dolomitizing fluid was undersaturated with respect to calcite, and therefore may have derived some carbonate necessary for dolomite precipitation from the dissolution of calcite within the reefal succession. Moreover, ferroan saddle dolomite shows a wide range of carbon and oxygen isotope values in a pattern similar to that observed for the ferroan calcites. The most positive values are for dolomite from the lower part of the section, and the most negative values are from the upper part and top of the unit (Fig. 4). Whereas δ13C values are variable, ferroan saddle dolomite δ18O values are very constant, even in samples where several successive analyses were made within a single crystal. This suggests that both temperature and fluid δ18O composition did not change significantly during precipitation.

Nonferroan syntaxial saddle dolomite precipitated after corrosion of the ferroan dolomite (Figs. 2, 3C). However, the isotopic compositions of nonferroan syntaxial dolomites are similar to those of the ferroan dolomites (Fig. 4), suggesting that they precipitated from fluids with similar oxygen isotope compositions and/or at similar temperatures, but with either a higher Eh or no source for Fe.

**Source of Mg and Fe for Dolomitization**

In order to estimate the extent of the water–rock interaction during the emplacement of the saddle dolomite and MDIs, a necessary constraint is the elemental mass balance for Mg and Fe needed for precipitation of the volume of dolomite (around 2% of the total volume of rock). Specifically, we might ask: sufficient Mg and Fe could be sourced locally through the dissolution of preexisting host rock, or is an extrinsic source necessary?

In order to answer this question, we have taken into consideration the concentration of Fe and Mg in the ferroan saddle dolomite (Table 1) and the average chemical composition of the host-rock components (Benito et al. 2005), which include: fossil fragments and micrite (60%); neomorphosed coral (15%); marine-derived cements (15%); meteoric cements (7%); and ferroan calcite cements (3%). On this basis, in order to precipitate 1 cc of ferroan dolomite, it would be necessary to dissolve 32 cc of the host rock to derive the requisite Fe, and 44 cc to obtain sufficient Mg. Thus, the newly formed dolomite would only fill about 2.5% of the porosity that was created by dissolution of the host rock. This is not consistent with the petrographic and field observations because only minor amounts of dissolution associated with dolomite emplacement are evident petrographically, and ferroan saddle dolomite occurs either as a replacement (on a one-to-one volume basis) or as cement filling primary and secondary porosity that was created during the early stages of meteoric diagenesis. Therefore, it is more likely that the diagenetic system was open during dolomitization, and the Mg and Fe needed for precipitation of dolomite were derived from an external source.

We might then ask what fluid Mg/Ca ratios were needed for dolomite precipitation. It has been shown that, at temperatures above 60–70°C, most Ca-rich groundwaters could be dolomitizing fluids (Hardie 1987). Specifically, dolomite is stable in high-salinity brines with a Mg/Ca ratio of around 0.7 at 80°C; this ratio decreases to 0.15 at 200°C (Hardie 1987). These Mg/Ca ratios are typical in groundwaters from sedimentary rocks and basinal brines (Hardie 1987; Hanor 1994). Moreover, prior studies have suggested that saddle dolomite generally precipitates from basinal brines with salinities 5 to 7 times greater than seawater, at temperatures ranging between 80 and 200°C (Spötl and Pittman 1998). Thus, it is possible that ferroan saddle dolomite precipitated from a fluid with Mg/Ca ratio lower than 1.

The source of saline basinal fluids can be discerned from the regional occurrence of late burial dolomite. Ferroan saddle dolomite cements are also observed in coeval Jurassic reefal units in other parts of the basin, as in the southern Bigornia and Soria sectors (Benito et al. 2001; Benito and Mas 2002; Benito et al. 2003) (Fig. 1A). In these places, ferroan dolomite cement occurs at the same position in theparagenetic sequence and is very similar in its petrographical and geochemical character to that described here in the Torrecilla Reef Complex. Based on prior studies (Radke and Mathis 1980; Spötl and Pittman 1998), ferroan saddle dolomite generally precipitates at temperature above 80°C; this requires a minimum burial depth of 2 km given a conservative geothermal gradient of 30°C per km.
Although the sequence examined here achieved sufficient depths to reach this temperature, ferroan dolomite is also present in the southern Bigornia area, where maximum burial depth never exceeded 1 km (Benito et al. 2001). It is therefore apparent that the emplacement of the late ferroan saddle dolomites must record a regional event that is independent of depth of burial. The requirement for elevated temperatures and a source for elemental constituents suggests that high temperature basinal fluids migrated during a single event and led to burial dolomitization in all sectors of the basin.

Precipitation of ferroan saddle dolomite, MDIs, and also nonferroan syntaxial dolomite was likely associated with the hydrothermal metamorphic event that affected the Cameros Basin during the late Albian–Coniacian (Benito et al. 2001; Benito and Mas 2002) (Fig. 1A). Upper Jurassic–Lower Cretaceous deposits of the Cameros Basin are composed mainly of Fe-rich sandstones and Fe- and Mg-rich clay minerals. This metamorphic event led to extensive transformation of clays and associated minerals in the northwestern parts of the basin (Alonso-Azcárate et al. 1995, 2005; Alonso-Azcárate et al. 1999b; Barrenechea et al. 1995; Barrenechea et al. 2000; Barrenechea et al. 2001) (Fig. 1A). Additionally, the entire 600-meter-thick Jurassic sequence in the Cameros Basin directly overlies the Triassic Buntsandstein and Keuper facies, which comprise Fe-rich red beds, Mg-rich clays, and evaporites (Mas et al. 1993; Mas et al. 2003; Alonso-Azcárate et al. 2005). Thus, it is probable that Fe and Mg necessary for saddle dolomite precipitation were sourced by the hydrothermal fluids from either Triassic and/or Upper Jurassic–Lower Cretaceous units.

CONCLUSIONS

This study describes a new type of micro-sized dolomite inclusions (MDIs) observed within ferroan calcite and suggests a process that can explain their formation during late-stage burial diagenesis. The paragenetic relationship of MDIs to their host ferroan calcites combined with measures of isotopic and elemental chemistry suggests a burial origin for MDIs associated with the emplacement of ferroan saddle dolomite, which postdates the ferroan calcite. The pathways by which dolomitizing fluids penetrated into the interiors of the calcite crystals may have been controlled by the network of interconnected micropores, cleavage, minute fractures, and shear planes in the calcite cements.

Carbon and oxygen isotope data suggest that during precipitation of the burial phases, fluids reacted and exchanged with the host rock such that \( \delta^{13}C \) values are inherited in both the ferroan calcites and the saddle dolomites. On the basis of petrographic features and the elemental composition of the host rock and ferroan dolomite and MDIs, the origin of these fluids can be constrained. It is unlikely that Fe and Mg necessary for precipitation of the ferroan dolomite and MDIs were sourced from the dissolution of host-rock carbonate, but rather must have been derived from an extrinsic source. Such fluids likely were associated with the hydrothermal metamorphic event that affected the Cameros Basin during the middle to Late Cretaceous.

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