SELF-ACCELERATING DOLOMITE-FOR-CALCITE REPLACEMENT:
SELF-ORGANIZED DYNAMICS OF BURIAL DOLOMITIZATION AND
ASSOCIATED MINERALIZATION

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ABSTRACT. A new dynamic model of dolomitization predicts a multitude of
textural, paragenetic, geochemical and other properties of burial dolomites. The
model is based on two postulates, (1) that the dolomitizing brine is Mg-rich but
undersaturated with both calcite and dolomite, and (2) that the dolomite-for-calcite
replacement happens not by dissolution-precipitation as usually assumed, but by
dolomite-growth-driven pressure solution of the calcite host. Crucially, the dolomite-
for-calcite replacement turns out to be self-accelerating via Ca$^{2+}$: the Ca$^{2+}$ released
by each replacement increment accelerates the rate of the next, and so on. As a result,
both pore-fluid Ca$^{2+}$ and replacement rate grow exponentially.

As brine enters and infiltrates a limestone, water/rock disequilibrium plus the
self-accelerating feedback inevitably yield a process that is self-organized, both in time
(as repeated dolomite growth pulses per slice of limestone) and in space (as successive
slices). Self-organization in pulses and slices accounts for several properties of burial
dolomites: (1) generation of dissolution porosity and its spatially periodic distribution;
(2) dolomitization affects only limestones; (3) sharp field contacts between dolo-
mitized and undolomitized limestone; (4) formation of both saddle dolomite and
“late-stage” calcite near the end of each growth pulse, accompanied by Mississippi-
Valley-type ores if the brine also contains Zn, Pb, Ba, sulfate, and other relevant
elements; (5) “sweeping” of ores downflow with accumulation in the last position of
the dolomitization front.

In addition, the combination of the self-accelerating feedback via Ca$^{2+}$ with the
known strain-rate-softening rheology of crystalline carbonates leads to another suite of
predictions that are strikingly confirmed by observation. If the dolomite-for-calcite
replacement becomes fast enough to lower the local rock viscosity sufficiently, then the
dolomite growth will pass spontaneously from replacive to displacive. This is when
thin, self-organized, displacive zebra veins form (Merino and others, 2006), indeed
displaying seamless contacts with their replacive walls and consisting of curved, or
saddle, dolomite crystals. Serendipitously, both the deformation of the dolomite
crystals (produced by Ca-for-Mg substitution driven by the huge pore-fluid Ca$^{2+}$) and
the seamless rheological transition result from the self-accelerating feedback via Ca$^{2+}$
itself; that is why they are always associated. This detail alone strongly suggests that the
new model captures the chemistry, drives, mechanisms, and feedbacks that lend burial
dolomitization and its often associated MVT ore deposits their geological uniqueness.

Key words: Dolomitization dynamics, chemical-rheological feedbacks, self-
organization, replacement physics, petrography of replacement, geochemistry’s blind
spot, dolostone porosity, dolomitizing brines, replacive-to-displacive dolomite transi-
tion, syntaxial zebra veins, MVT lead-zinc ores, saddle dolomite, late-stage calcite,
dedolomitization

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I. INTRODUCTION

Huge volumes of limestone have been transformed into dolostone by replacement of calcite by dolomite in many ancient sedimentary basins. Although the transformation may seem chemically simple—just add Mg to calcite to make dolomite—and despite extensive research by petrologists and geochemists for decades, several aspects of the metasomatic process of dolomitization and several properties of burial dolomites are poorly understood: Why are only limestones dolomitized, and not other rocks? What causes the large porosity that makes burial dolostones excellent petroleum reservoirs? What are the source, nature, and chemistry of the dolomitizing aqueous solutions? How do those Mg-rich brines travel to the site of dolomitization? Is burial dolomitization fast or slow? What are the mechanisms of formation of the two basic types of dolomite, the replacive and the zebra/breccia veins (which are displacive), and why are they associated? Why are the zebra veins thin and equidistant, have no sharp contact with their walls, and consist of dolomite crystals that are curved (thus called saddle dolomite), displaying a consistent assemblage of enigmatic textures? Why is dolomitization of limestones paradoxically accompanied by formation of so-called late-stage calcite and even dedolomitization, or calcite-for-dolomite replacement? Why are the distinctive Mississippi-Valley-type (MVT) ore deposits—consisting of barite, fluorite, sphalerite, galena, and other minerals—typically hosted in burial dolomites? The prevailing theory of dolomitization, gives only ad hoc or unsatisfactory answers to those questions. Furthermore, it contains a flaw that we believe is both fundamental and hidden from view: Whereas according to the prevailing theory replacement, and in particular the dolomite-for-calcite replacement, is produced by “dissolution-precipitation,” alert early investigators (Bastin and others, 1931) long ago showed petrographically that replacement cannot be produced by dissolution-precipitation. Thus, in the section “Replacement Physics” we discuss the evidence against dissolution-precipitation, why both the old petrographic insights and their kinetic implications were forgotten, and how replacement does take place, in light of recent theoretical rheological-kinetic research.

The main purpose of this paper is to propose a new view of dolomitization dynamics that may solve many of the problems of prevailing dolomitization theories; several of these problems are summarized in Section II. The new model is a forward model, one based on two postulates. The first arises in contrast to the widespread assumption that the dolomitizing brine must be supersaturated with dolomite. If so, any rock type could in principle be dolomitized, whereas in fact only limestones are. Therefore, in the new model we assume that the dolomitizing aqueous solution must be Mg-rich but—counterintuitively—undersaturated with dolomite and with calcite.

The second postulate is that replacement, and in particular the dolomite-for-calcite replacement, takes place not by dissolution-precipitation, but by guest-growth-driven pressure solution of the host. This mechanism—as significant for weathering, diagenesis, ore genesis, and metamorphism as it is for dolomitization—is justified in Section III (Replacement Physics).

In the main section of this paper, Section IV, “New Dolomitization Model,” we describe how the two postulates of the model, combined with kinetic and rheological feedbacks inevitable in the transformation of limestone into dolostone, lead to predicting a complex, non-linear, self-organized dolomitization process that yields the two observed dolomite types (replacive and displacive, or dolomite-I and dolomite-II) characteristic of burial dolostones along with a multitude of textural, paragenetic, geological, geochemical, rheological, dynamic, and field properties. All the predictions are confirmed by independent evidence, suggesting the new model is on the right path. Although the new model is conceptual, some of its mechanisms (in particular the self-accelerating kinetics of the dolomite-for-calcite replacement) are
quantified in two Appendices. Implications of the new dynamics are discussed in Section V, including its use as a framework for future reaction-transport calculations of dolomitization and associated mineralization. A Summary of the new paradigm is given in the last section, Section VI.

II. WEAKNESSES OF THE PREVAILING THEORY OF BURIAL DOLOMITIZATION

According to the consensus reached in recent decades (for example, Zenger and Dunham, 1980; Land, 1980; Yao and Demicco, 1997; Nielsen and others, 1998; Gregg and others, 2001; Wilkinson, 2003; Gregg, 2004; Machel, 2004; Gasparrini and others, 2006; Lonee and Machel, 2006; Davies and Smith, 2006) burial dolomitization is thought to take place in two stages separated in time. In the first stage, a huge volume of limestone is replaced by dolomite—the so-called dolomite-I, or replacive dolomite—via dissolution-precipitation driven by a Mg-rich, dolomite-supersaturated brine, possibly evaporative, which in some versions of the theory comes down from the surface environment. An example of replacive dolomite, or dolomite-I, is shown in figure 1A. The replacive dolostone is then assumed to undergo a void-opening process—by fracturing, hydraulic fracturing, karst dissolution, or hydrothermal alteration—and eventually a second phase of dolomite, the so-called dolomite-II, or “void-filling cement,” grows in those voids from a new, hot brine also supposedly supersaturated with dolomite. This dolomite-II often occurs as sets of zebra and “breccia”-like veins (figs. 1B, 1C, 1D, and 1E), textures that are common also in Mississippi-Valley-type (MVT) ore-deposits.

There are several problems with the consensus model.

1. It takes for granted that two phases of dolomite growth (thought to have formed from different brines), the hypothetical void-opening process between them, and MVT ore growth (often hosted in these dolostones) must all coincide in space and in the appropriate temporal sequence, in most burial dolostones.

2. As noted, if the dolomitizing brine is assumed to be supersaturated with dolomite from the start any rock type could in principle be dolomitized. In fact only limestones can be dolomitized. Furthermore, for some investigators the assumption points to seawater, supersaturated with respect to dolomite, as a candidate for the dolomitizing brine. This choice leads to predicting a dolomitization effected by descending, low-temperature solutions (unless one calls for dolomitization by convecting sea-water, for example Machel, 2004). Each of the two implications is contradicted by good evidence. Gasparrini and others (2006) show large ascending fingers of dolomitization into Paleozoic limestones in NW Spain, see figures 2A, 2B; and fluid inclusion data published by many geochemists leave little doubt that dolomitization (dolomite I and dolomite II) happens between 70 and 200 °C, which means dolomitization by solutions that are hot.

3. The prevailing view sheds no light on the origin of the remarkable assemblage of textures systematically displayed by dolomite-II or on its association with dolomite-I. Dolomite-II occurs as thin, white, zebra and breccia-like veins displaying seamless contacts with the replacive dolomite-I of the walls and consisting of coarse saddle-shaped crystals; see figures 1B to 1E. First, if the two dolomites are supposed to be independent and separated in time, why is their contact always gradual, seamless, and optically continuous, instead of sharp under the microscope, as one would expect? Second, while the zebra veins and breccia veins are routinely referred to in the literature as “void-filling cement,” we presented evidence (Merino and others, 2006) indicating that the dolomite-II veins are not cements, but displacive veins. The evidence includes: (a) small transform-fault displacements of veins that could not have taken place if the veins were passive cements; (b) tiny stylolites in the slices of host dolomite-I that formed as the veins grew in order to compensate for the volume added by the veins. And (c), we showed theoretically that the same vein-growth-generated
Fig. 1. Textural properties of replacive and displacive dolomite in burial dolomitization. (A) A dolomite rhombohedron 0.8 mm across replaces parts of three oolites at once. The facts that the dolomite crystal is idiomorphic and that the replacement preserves the oolitic rings undeformed (and thus preserves mineral volume) each separately rules out that it was produced by dissolution-precipitation; see text. (1960 photomicrograph reproduced with permission from its author, Raymond C. Murray.) (B, C) Plane-polarized and cross-polars views under low magnification of one zebra vein consisting of displacive dolomite (or dolomite-II), from the Cretaceous Basque-Cantabrian basin, northern Spain. In (B) note lack of sharp contact between the dolomite II vein (coarse crystalline) and its dolomite I walls (fine crystalline). In (C) note the sweeping extinction, denoting deformation of the structure, of several large saddle dolomite-II crystals. A few of these can be seen to grow in optical continuity with small crystals of dolomite-I; this is visible only while rotating the microscope stage and under higher magnification. (D, E) Plane-polarized and cross-polars views of displacive zebra veins of saddle dolomite in burial dolomites of SW Sardinia. Note the seamless contact between dolomite-I and dolomite-II, and the thin colorless late-stage calcite cement (cc) at center, with low optical relief. In (E) a few instances of saddle dolomite-II crystals in optical continuity with small crystals of dolomite-I are detectable under higher magnification while rotating the stage.
stress, or induced stress, that forcibly displaces the host also triggers a feedback that
tends to pressure-dissolve, and thus “weed out,” incipient veins that happen to be too
close to their neighbors, leaving the surviving veins more equidistant than before, and
explaining the self-organized nature of zebra veins that had been noted by Merino
(1984) and Fontboté (1993). Third, the prevailing dolomitization theory also leaves

Fig. 2. Two views of the sharp contacts between ascending fingers of completely dolomitized (dark gray) and undolomitized Upper Carboniferous limestone (light gray), in the Southwestern Cantabrian Zone, NW Spain (Gasparrini and others, 2006). That the dolomitization rose through the limestone is particularly clear in (A), a photo by M. Gasparrini. (B), photo by A. Canals.
unexplained the facts that the veins are thin and bilateral and consist of crystals of dolomite that are curved and optically continuous with the replacive dolomite-I crystals of the walls.

(4) The prevailing theory leaves unclear what the local reactions of dolomitization really are. On the basis of oilfield-brine compositions from Oklahoma, Weyl (1960) demonstrated that dolomitization happens according to what he called an “approximate mass balance,” \( 2\text{calcite} + Mg^{2+} = \text{dolomite} + Ca^{2+} \), a mass balance involving calcite dissolution and dolomite precipitation, with carbonate being conserved between the two minerals over some mesoscopic scale left unspecified by Weyl. However, despite the fact that the mass balance does not apply to the local, subgrain scale, most investigators since Weyl (1960) have adopted it as the local reaction that brings about dolomitization, disregarding long-known petrographic evidence (see example in figure 1A discussed in the next section) showing that the local replacement does preserve mineral volume and should therefore be \( 1.74 \text{calcite} + Mg^{2+} + 0.26CO_3^{2-} = \text{dolomite} + 0.74Ca^{2+} \), where the 1.74 factor approximately equals the dolomite formula volume, 64.3 cm\(^3\), divided by the calcite formula volume, 36.9 cm\(^3\). Also, use by recent modelers (Wilson and others, 2001; Jones and Xiao, 2005) of Weyl’s mesoscopic mass balance in the continuity equation does not appear to meet the requirement that the reactions whose rates \( R_i \) are inserted in its reaction term must be local—that is, they must take place within a volume element small enough that the difference in flux across it, between entry and exit, can be approximated as linear; only then can quadratic and higher-order terms (in the width of the volume element) in a Taylor expansion of the flux be neglected. (See the derivation of the mass conservation equation in Robinson and Stokes, 1959, p. 47.)

(5) As mentioned above and shown in the next section, “Replacement Physics,” dissolution-precipitation cannot account for the properties of replacement. But dissolution-precipitation also cannot account for the fact that the dolomitization front in the field is often sharp, a feature shown in figures 2A and 2B. Quantitative models of dolomitization based on dissolution-precipitation (Wilson and others, 2001; Jones and Xiao, 2005) predict a transition zone between dolomitized and undolomitized limestone which is tens of meters wide, thus failing to reproduce the sharp contacts observed in the field.

III. REPLACEMENT PHYSICS

We justify here the second postulate of the new dolomitization model, that replacement takes place not by dissolution-precipitation but by guest-growth-driven pressure solution of the adjacent host. Throughout this paper the term replacement refers to the striking phenomenon identified petrographically already by Lindgren (1912, 1925) in many rocks and superbly discussed by Bastin and others (1931), whereby a crystal or crystal aggregate of a new mineral A, the guest, occupies the very space occupied before by the host mineral B, but preserving B’s volume and (some) morphological details of B.

This section may be seen as a digression from our main objective—the dynamics of dolomitization—but it is a necessary digression, because it deals with the fundamental problem of the mechanism of replacement. Our discussion here, a summary from a review in progress, is based on four crucial papers: Bastin and others (1931), Maliva and Siever (1988), Dewers and Ortoleva (1989), and Nahon and Merino (1997). Others also relevant are those by Carmichael (1987), Merino and others (1993), Merino and Dewers (1998), Fletcher and Merino (2001), Merino and Banerjee (2008), and Banerjee and Merino (2011).

In a remarkable paper whose insights would be soon forgotten, Bastin and others (1931) saw that the preservation of volume and morphological details of the host by the guest that are characteristic of replacement (a) requires guest growth and host
dissolution to proceed simultaneously and at the same volumetric rate as each other; and (b) rules out formation by “dissolution-precipitation.” As Bastin and others (1931, p. 603) put it,

“Idiomorphic replacement is . . . illustrated by figures 11 and 12, which show cubic crystals of pyrite transecting the schistosity of a fine-grained schist. Obviously the pyrite [grew] by the simultaneous replacement of the several minerals of the schist. A further implication not generally recognized is that the several minerals of the schist must have been replaced at essentially the same rate, an astonishing fact in view of the diverse solubilities of these several minerals under most circumstances.” (Bastin and others’ italics; text in brackets added.)

Geochemists and modelers, however, still generally assume that replacement results from dissolution-precipitation (see below, Geochemistry’s Blind Spot), not realizing that dissolution-precipitation cannot bring about (except by chance) the equalization of rates that Bastin and others grasped, or that there is solid petrographic evidence that rules out dissolution-precipitation as the mechanism of replacement. We see some of this evidence in figure 1A. The rhombohedral crystal of dolomite in the center has replaced the right end of the round calcium carbonate oölite at left, preserving, undeformed, its ring of black carbonaceous inclusions, thus also preserving mineral volume, since the ring of dust has not been swollen or shrunk upon growth of the dolomite. If the replacement had taken place by dissolution-precipitation, (1) the carbonaceous dust ring of the oölite would have been destroyed before the dolomite rhomb could have grown and preserved it. (2) Also, the calcitic oölite could not have dissolved leaving a hollow space having the characteristic crystal form of the future dolomite idiomorph. (3) And also: even if the oölite had dissolved leaving a hollow space with the form of the future dolomite idiomorph, and if we note that the one idiomorph replaces portions of three separate oölites at once, formation of the replacement by dissolution-precipitation would have required the three oölites to coordinate their respective euhedral dissolutions such that the three voids were oriented making up only one euhedral void that could be later filled by only the one dolomite idiomorphic crystal. This is impossible. Each of the three criteria present in this photomicrograph independently rules out formation of the replacement by dissolution-precipitation. “The criteria of replacement are many . . . The alert observer will discover and use other criteria of relative age and replacement [in other rocks]” (Pettijohn, 1957, p. 112); see more examples in Merino and Dewers (1998).

In any case, it was clear to Bastin and others (1931) that there are genuine cases of dissolution-precipitation, that is, cases in which a mineral grain dissolves and a new one precipitates later in part or all of the space freed, but they were clear that these were not cases of replacement, because neither morphological details nor mineral volume could be preserved, since host dissolution and guest growth could not be simultaneous, nor could their rates be mutually equal (except by chance):

“In other instances, however, minerals are dissolved by one solution and after an interval during which open spaces exist, new minerals are deposited in these spaces . . . This is not replacement.” (Bastin and others, 1931, p. 595; their italics.)

But as noted, Bastin and others’ (1931) insights were soon forgotten, and the next improvement in understanding replacement did not come until decades later, when Maliva and Siever (1988) proposed that replacement happens not by dissolution-precipitation but because of the force of crystallization exerted by the growing guest on the host. Dewers and Ortoleva (1989) applied the Navier-Stokes equation for momentum conservation to demonstrate that growth in a rigid rock must trigger other mineral reaction(s) such that volume is conserved; they thus demonstrated theoretically the existence of the replacement phenomenon. Fletcher and Merino (2001)
showed that the correct coupling factor required for the equalization of volumetric rates in replacement was the crystallization stress or induced stress, which can enter into a feedback with the rates of guest growth and host dissolution, whereas the empirical force of crystallization cannot. Nahon and Merino (1997) showed how the rates are made to become mutually equal by the crystallization stress (fig. 3). The key feedback arises because the crystallization stress generated by guest growth within a (rigid-enough) rock increases the equilibrium constants of both host and guest. Through the appropriate mass-action-law expressions for host and guest, see the caption to figure 3, this causes the guest growth rate to decrease but the host dissolution rate to increase, until the two rates intersect, becoming mutually equal, at a time $t^\ast$. From then on, the two rates remain equal, and mineral volume is thus automatically preserved. The crystallization stress during replacement was calculated by Merino and others (1993) and Fletcher and Merino (2001) for several A,B mineral pairs.

Preservation of morphological details, or “ghosts”.—The preservation or erasure of morphological details by replacement can be understood if we regard the increments
of A growth as small tiles that pressure-dissolve their way into B. If the tiles are smaller than a detail, which is equivalent to the A growth rate being sufficiently small, then the replacement can preserve it. In the Pompeii mosaic in figure 4A the tiles used were so small that it was possible to capture the features of the woman’s face. In contrast, if the increments of A growth, represented by tiles or brush strokes, are as large as the details or larger, these are bound to be erased or half-erased upon replacement. In Cézanne’s “Château Noir,” figure 4B, the brush strokes are so large that we can only barely tell that the château has windows and that the foreground consists of big trees; morphological details start to become erased.

Preservation of ghosts seems to us impossible to explain if replacement is thought to happen by dissolution-precipitation, because precipitation of the guest could not happen simultaneously with, and at the same place as, and much less at the same volumetric rate as, the dissolution of the host, except by chance or by experimental design.

Geochemistry’s blind spot.—Weyl (1959), Garrels (1960), Garrels and Christ (1965), Helgeson (1968), and Helgeson and others (1969) constructed a new, quantitative geochemistry in the 1960s, long after the insights of Bastin and others (1931) had been forgotten. Weyl (1959), in his influential article proposing the idea of modeling water/rock interaction essentially by dissolution and precipitation of minerals enabled by transport in aqueous solution, was unaware of the help petrography may provide:

“In subdividing our field, . . . we are not interested in what [a rock is like] now, but in how it became that way. . . . Instead of looking at a rock and asking for an explanation of its past. . . .” (Weyl, 1959, p. 2001; italics and insert in brackets are the authors’)
The new geochemistry implicitly viewed every geochemical process taking place in rocks, from weathering to metamorphism, as resulting simply from dissolution and precipitation, as though in a fluid medium. Harker’s (1950, p. 29) observation, “. . . the crystallization or recrystallization of minerals in metamorphism proceeds, not in a fluid medium, but in the heart of a solid rock, an environment which cannot fail to modify greatly their manner of growth” (Harker’s italics), also went unnoticed.

Thus the new geochemistry was born with a blind spot: It was unaware of the replacement phenomenon and of its fundamental kinetic implications, which could only be grasped precisely by “looking at rocks,” as Bastin and others (1931) and Bastin (1950) had already done. When the problem of the replacement mechanism arose explicitly later, geochemists and modelers have continued to take for granted that replacement forms by dissolution-precipitation, unaware of the blind spot. To many geochemists (Putnis, 2009, and articles therein) the term “replacement” seems to designate whatever takes place by dissolution-precipitation, in their experiments and/or in nature, and every process from weathering to metamorphism is assumed to result from dissolution-precipitation. As we have seen, however, the term replacement already has since the early 1900s a different, far more specialized meaning: it refers to a unique kinetic-rheological phenomenon which occurs in many sorts of rocks, which is characterized by preserving mineral volume and morphological features of the host, which cannot take place by dissolution-precipitation, and whose existence—only in rigid enough rocks—was predicted theoretically by Dewers and Ortoleva (1989).

IV. NEW DOLomitIZATION MODEL

1. The Dolomitizing Brine

As noted in the Introduction, if the brine were supersaturated with dolomite from the start, as widely assumed in the dolomitization literature, then any kind of rock could in principle become dolomitized, not only limestone as is the case worldwide. Thus in the new model the dolomitizing fluid is hypothesized to be a hot CaCl₂ brine rich in Mg²⁺ but undersaturated with both dolomite and calcite, and thus barren or very poor in carbonate.

Our choice of dolomite-undersaturated dolomitizing brines may seem counterintuitive at first, but it will become clear that it does trigger dolomitization only of limestones (and other rocks rich in CaCO₃). If the dolomitizing brine also contains chemical elements such as Zn, Pb, Fe, Sr, Ba, and sulfate, needed to make typical MVT minerals, then, as will become apparent later, dolomitization may be accompanied by formation of MVT ore deposits—which are well known to be hosted in burial dolostones.

We do not know how our postulated brines would form in nature, but, intriguingly, the deep sedimentary brines from the central Mississippi Salt Dome basin (Kharaka and others, 1987), and also the deep brines of the Smackover Formation in Arkansas (Moldovan and Walters, 1992), which abound in precisely the elements needed to make Mississippi Valley type ore deposits, are according to those two reports Mg-rich and slightly undersaturated with both calcite and dolomite (see table 1). In other words, these brines from the Gulf Coast basin might be natural examples of our postulated dolomitizing brines and at the same time provide a clue to why MVT ore deposits are typically associated with burial dolomites. A word of caution: The slight undersaturation with calcite and dolomite reported by Kharaka and others (1987) and Moldovan and Walters (1992) for the brines of table 1 depends on how the distribution of aqueous species was computed; how the pH at the in situ temperature was computed for each brine; which thermodynamic database was used; and, in
particular, whether the reported pH was calculated taking account of internal consistency issues (Merino, 1979)—a delicate problem whose discussion is beyond the scope of the present paper.

2. The Trigger Reaction: Fast Limestone Dissolution

We adopt a rectangular model system consisting of porous limestone, and assume that the Mg-rich, dolomite- and calcite-undersaturated brine enters through the left side at some velocity (injected by tectonic squeezing of the deep basin or some other cause). Since it is undersaturated with calcite, the brine should first quickly dissolve some calcite according to

$$\text{CaCO}_3 + \text{H}^+ \rightarrow \text{Ca}^{2+} + \text{HCO}_3^-,$$

thereby producing some dissolution porosity at the entry into the current limestone “slice,” between times $a$ and $b$ in figures 5A and B. This is why burial dolostones always

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**Table 1**

|                  | Central Mississippi brine 84-MS-11 | Smackover Fm brine 55 |
|------------------|------------------------------------|-----------------------|
| Depth, m         | 3486                               | 2365                  |
| Temp, °C         | 102                                | 85                    |
| Density, g/cm³   | 1.22                               | 1.225                 |
| Tot salinity, mg/l | 320,000                           | 336,492               |
| Li               | 35                                 | 37                    |
| Na               | 61,700                             | 77,325                |
| K                | 990                                | 893                   |
| Mg               | 3,050                              | 4,265                 |
| Ca               | 48,600                             | 41,225                |
| Sr               | 1,920                              | 2,201                 |
| Ba               | 60                                 | 41                    |
| F                | 1.5                                | 19                    |
| Cl               | 198,000                            | 203,003               |
| Br               | 2,020                              | 6,647                 |
| Alkalinity       | 206                                | ~100                  |
| SO₄              | 64                                 | 128                   |
| Mn               | 212                                | 79                    |
| Fe               | 465                                | 100                   |
| Pb               | 70                                 | 40                    |
| Zn               | 243                                | 143                   |
| SiO₂             | 27.8                               | 41                    |
| pH               | 5.08(3)                            | 5.08(4)               |
| S.I. (dolo)      | -0.37(2)                           | -0.37(2)              |
| S.I. (calc)      | -0.20(2)                           | -0.28(1)              |

All concentrations are in mg/l. S.I. = saturation index.

(1) Calculated with PHREEQC: http://wwwbrr.cr.usgs.gov/projects/GWC_coupled/phreeqc/
(2) From Kharaka and others (1987) and Moldovanyi and Walters (1992).
(3) In situ pH values calculated by Kharaka and others (1987, p. 557).
(4) Measured field pH, Moldovanyi and Walters (1992, table 3), who give also calculated in-situ pHs of 6.26 and 6.19 in their table 5 for this same brine.
have considerable porosity, shown in the figures as a porosity peak. But most importantly, the fast initial dissolution of some calcite at the entry point suddenly releases considerable Ca\(^{2+}\) and carbonate to the Mg-rich brine, which now becomes supersaturated with respect to dolomite. This is the key contribution of any limestone to its own dolomitization; this is why only limestones (and also siliciclastic rocks cemented with abundant CaCO\(_3\)) can be dolomitized. In short, the postulated calcite-undersaturated brine accounts for dolomitization only of limestones, for the high dissolution porosity of dolostones, and for supersaturating itself with respect to dolomite.
3. Crucial Feedback: Self-Accelerating Dolomite-for-Calcite Replacement

At time \( b \) in figure 5A, the Mg-rich solution, having just become dolomite-supersaturated, starts precipitating dolomite according to

\[
Ca^{2+} + Mg^{2+} + 2CO_3^{2-} = CaMg(CO_3)_2(dolo)
\]

as it infiltrates through the limestone. As it grows, the dolomite pressure dissolves (thus replaces, see Section III) an equal volume of adjacent calcite according to

\[
CaCO_3(calc) = Ca^{2+} + CO_3^{2-}.
\]

Note that the dissolution (3) is driven by the induced stress generated by dolomite growth, not by chemical affinity as (1) was. The local mass balance of the two simultaneous, adjacent, induced-stress-coupled reactions (2, 3) is

\[
1.74CaCO_3(calc) + Mg^{2+} + 0.26CO_3^{2-} = CaMg(CO_3)_2(dolo) + 0.74Ca^{2+},
\]

where 1.74 = ratio of dolomite to calcite formula volumes, 64.3 and 36.9 cm\(^3\) respectively.

How and why replacement systematically conserves volume is explained in figure 3 and Section III “Mineral Replacement.”

The aqueous Ca\(^{2+}\) released to the pore fluid by each increment of replacement (4) must increase the local ion-activity product and saturation index for dolomite,

\[
\Omega = Q/K_{eq}^{dolomite} = (a_{Ca^{2+}})(a_{Mg^{2+}})(a_{CO_3^{2-}})^2/K_{eq}^{dolomite},
\]
which in turn must accelerate the growth rate of the next increment of dolomite growth, since

\[ R_{\text{dolomite}} = k_{\text{dolite}}S_\Omega(\Omega - 1). \] (5a)

Note that, although each increment of replacement consumes some carbonate, the \(\text{CO}_3^{2-}\) activity is probably kept roughly constant by bicarbonate dissociation. Likewise, the \(\text{Mg}^{2+}\) concentration also probably remains roughly constant in space during the advance of the brine (say, between times \(t_1\) and \(t_4\) in figs. 5A and 5B), because over that time interval the \(\text{Mg}^{2+}\) advection rate is much larger than the rate at which \(\text{Mg}^{2+}\) is being sequestered by dolomite growth. The strong positive feedback, shown in boxes 2.1–2.3 of figure 5C, forces both the pore-fluid \(\text{Ca}^{2+}_{(aq)}\) concentration and the dolomite-for-calcite replacement rate to increase exponentially with time (see Appendix 1), as shown in figure 5A. Eventually, around time \(t_4\), the dolomite growth rate starts to soar, soon overwhelms the Mg advection flux, and abruptly causes the Mg\(^{2+}\) to plummet between times \(t_5\) and \(T\)—which shuts down the replacement suddenly. \(T\) is thus the time needed for the replacement rate to become so fast that it uses up all the available Mg instantaneously. By time \(T\) the replacement (of only a small fraction of the limestone, see below) has taken place over an interval of length \(L = vT\), where \(v\) is the brine infiltration velocity. This is the reason dolomitization takes place by “pulses” of limited length. A tentative calculation of time \(T\) and length \(L\) are given in the next section. See Glossary, table 2.
4. Complete Dolomitization of a “Slice” and the Jumping Reaction Zone

The first pulse of precipitation just described can replace only a volume fraction of the current limestone slice equal to \( \frac{c}{H_9278} \) (see Appendix 2). Even for a brine with \( 10^4 \) ppm of \( \text{Mg}^2+ \) and a limestone with porosity 0.2, this volume fraction would amount to only 0.005. What happens immediately after the first precipitation pulse suddenly ends, as the brine continues to infiltrate the limestone? How does the unreplaced 99.5 percent limestone in the slice become completely dolomitized—as we know it does from the sharp contacts seen in the field (fig. 2)? We see next that complete dolomitization of a limestone slice takes place by an automatic sequence of many pulses.

At time \( T \) the brine in the pores of interval \((0, L)\) in figure 5B is left completely depleted in \( \text{Mg}^2+ \) but calcite-supersaturated. Fresh Mg-rich calcite-undersaturated brine continues to enter through the limestone cross section at \( x = 0 \), see figure 6. Because there is still 99.5 percent by volume of limestone left, the fresh brine immediately dissolves some calcite, supersaturates itself with dolomite, and proceeds to deposit a second pulse of dolomite precipitation similar to the first. This repetitive process continues for as long as there is some undissolved calcite left in the \((0, L)\) interval. \( N = \frac{1}{\phi c_{Mg}} b \) pulses are needed to dolomitize one slice completely, taking roughly a total time \( NT \). It will be seen later that each pulse consists mainly of replacive dolomite, but also of very minor calcite and ores (see Section IV-7); the many thin layers of which bothryoidal aggregates of sphalerite typically consist (fig. 7A) might be seen as a record of the repeated pulses of precipitation. Tentative values of \( T \) (<100 years), \( L \) (several meters), and \( N \) (a few hundred) are calculated in Appendix 2, table A1.

This is how each limestone slice becomes completely dolomitized—in a time \( NT < 20,000 \) years—before the replacement zone jumps to the next slice. In figure 6 each pulse has been represented by its exponential replacement rate versus time; only four (should be a few hundred) pulses are shown, versus time, for each of the two slices that have been already completely dolomitized—and only one for the slice currently under dolomitization. All these exponential curves and their abrupt stops are similar versions of the rate curve in figure 5A. Dolomitization of the current slice of limestone starts only after the second slice became completely dolomitized, and after a dead time \( T \).
during which fresh Mg-rich brine flows through the slice just dolomitized—without reacting with it—and starts entering the next slice at $x = 2L$. The reaction zone now jumps forward by a distance $L$ to its next position, $(2L, 3L)$ in figure 6, where it will remain for the next $NT$ years.

The final phase of each pulse (especially for the latest several pulses of each slice), when the dolomite growth reaches its greatest rate, is a moment of particular activity and complexity. There may occur a self-induced change in the rheology of the local dolostone; this is discussed in Sections IV-5 and IV-6. Several minerals grow in this final phase, during and after the dolomite shutdown, that constitute a characteristic paragenesis in burial dolostones; this mineralization is described in Section IV-7.

5. From Replacive to Displacive Growth: Self-Induced Change in Dolostone Rheology

The self-accelerating dolomite-for-calcite replacement ends up—if aqueous Mg$^{2+}$ is not used up first—modifying the rheological response of the crystalline dolostone to the crystallization stress, or induced stress, generated by dolomite growth. The property that makes this possible is that crystalline carbonates are non-newtonian materials of the strain-rate-softening kind (Verhoogen and others, 1970, p. 508), that is, their viscosity decreases as the carbonate is deformed at an increasing strain rate. This behavior was demonstrated experimentally by Heard and Raleigh (1972) for Yule marble at temperatures between 300 and 800 °C; their results, fitted by them to the Weertman equation, yielded

$$\dot{\varepsilon} = H\sigma^N,$$

with a stress exponent $N \approx 8$. The stress versus strain rate curve, graphed schematically in figure 8A, is knee-shaped. The viscosity of the marble, $\eta (= d\sigma/d\dot{\varepsilon} = G\dot{\varepsilon}^{-7/8})$, equal to the slope of the curve at each point, obviously decreases with increasing strain rate, expressing that marble is indeed strain-rate-softening in the temperature ranges studied. [$\varepsilon$ is strain, $\dot{\varepsilon}$ is strain rate ($= d\varepsilon/dt$), $\sigma$ is stress, and $H$ and $G$ ($= H^{-1/8}$) are temperature-dependent factors.] We assume that this strain-rate-softening behavior
applies also to crystalline dolostones, and down to the 300 to 100 °C range of relevance to burial dolomitization. The mechanism of dolostone deformation at this low temperature range is non-pervasive volume loss by pressure solution at stylolites, as shown by Merino and others (2006).

Fig. 7. Replacements among dolomite and ores in burial dolostones. (A) A curved dolomite rhombohedron replaces sphalerite at the Galmoy ore deposit, Ireland, preserving the layering of the disappeared sphalerite. (B) Dolomite replaces botryoidal sphalerite at Riopar, Albacete (Spain). (C) Sphalerite partly replaces dolomite leaving unreplaced the central portion of each zebra vein, at San Vicente, Peru. (D) Plane-polarized view under low magnification of the area marked in (C). Note that the medium gray sphalerite preserves rhombohedral terminations and cleavages of displacive dolomite-II, one of them curved; the sphalerite preserves both the small crystal size of the replacive dolomite-I, and the large crystal size of the displacive dolomite-II. (E) Barite at center replaces a curved dolomite-II crystal at the Hammam Zriba deposit (Zaghouan province, north-eastern Tunisia). The dark gray areas at top right and left are fluorite replacing displacive dolomite.

applies also to crystalline dolostones, and down to the 300 to 100 °C range of relevance to burial dolomitization. The mechanism of dolostone deformation at this low temperature range is non-pervasive volume loss by pressure solution at stylolites, as shown by Merino and others (2006).
The combination of self-accelerating replacement and strain-rate softening dolostone has a surprising consequence. Since the self-accelerating replacement imposes—via the crystallization stress exerted by the growing dolomite—an ever-increasing (elastic) strain rate on the host dolostone itself, it follows (fig. 8A) that the dolostone’s bulk viscosity becomes low enough to let the until-then replacive growth pass continuously into displacive, naturally causing the contact between replacive and displacive dolomite to be gradual and seamless—as sketched in figure 8B and as observed petrographically (figs. 1C,D and E,F).

Thus, when the slice is sufficiently crystalline (in late pulses) and its viscosity is lowered sufficiently—that is, when the dolomite growth rate and thus the strain rate reach a high enough value $R_{\text{critical}}$, figures 5A and 8A—the dolomite growth should pass continuously from replacive to displacive, as sketched in figure 8B. The replacive crystals 1, 2, 3, 4 (fig. 8B) that happened to be growing at time $t_5$ (see fig. 5A)—when the rheological transition takes place—would each consist of a replacive portion...
(formed just before \(t_5\)) and of a displacive portion \((1',2',3',4')\) grown between times \(t_5\) and \(T\) with no sharp boundary between them, and with the displacive portion in optical continuity with the replacive portion, as shown in figure 8B. The forced displacement of 1,2 from 3,4 will propagate sideways, resulting in a bilateral, syntaxial vein. Many such displacive veins would be simultaneously generated over tens of centimeters or more; their orientation and interactions were studied by Merino and others (2006)—see next section.

6. Displacive Veins of Saddle Dolomite

Gradual, seamless contact between dolomites I and II.—The predicted seamless, optically continuous displacive crystals and vein growth just described exactly match the two examples shown in figures 1B, 1C and 1D, 1E as well as those reported by Tona (ms, 1973), Fontboté and Amstutz (1980), Horton (1989), Wallace and others (1994), Gasparrini (ms, 2003, plates 8, 9), Vandeginste and others (2005), Gasparrini and others (2006), and Merino and others (2006), who pointed out that the margins of the dolomitic zebra and breccia veins are, under the microscope, gradual (over a length of a few micrometers) and seamless.

It now becomes clear that the observed gradual transition from replacive dolomite to white displacive veins means that there was no growth interruption between them; that the displacive dolomite-II is not filling a prior void (but making room for itself by pushing the host replacive dolomite aside); and that the two dolomites form at the same temperature and from the same (though quickly evolving) aqueous solution.

Genesis of saddle-shaped dolomite.—Saddle-shaped dolomite crystals are known to be extra-calcic. Radke and Mathis (1980) hypothesized that the extra Ca substitutes for Mg in dolomite and deforms its structure, because \(Ca^{2+} > Mg^{2+}\) in size. By transmission electronic microscopy Barber and others (1985) found submicroscopic bifurcating calcitic ribbons, each a few to 100 nm thick, coherently incorporated in crystals of saddle dolomite, and attributed the deformation to such microstructures. Either or both microstructures would account for the characteristic deformation of the structure that results in saddle-shaped dolomite-II crystals and for their concomitant extra-calcic composition. But what neither article solved was why those microstructures were systematically present in the dolomite-II of a burial dolostone and even in some dolomite-I crystals. Radke and Mathis (1980) thought that the proposed cation substitution might result from high temperature.

Now we realize, however, that the driving force that pushes \(Ca^{2+}\) to substitute for \(Mg^{2+}\) in dolomite crystals, and/or that forces the precipitation of submicroscopic calcitic slivers within the dolomite crystal structure, must be none other than the exponentially rising pore-fluid \(Ca^{2+}\) concentration, which is forced to reach very high values by the same feedback that forces the replacive dolomite growth rate to become huge and—if the \(Mg^{2+}\) lasts—to seamlessly transition into displacive growth. This is why the fastest growing dolomite crystals, regardless of whether they have already become displacive, are also the ones that are systematically saddle-shaped. Examples of saddle dolomites are in figures 1B, 1C and 1D, 1E.

Syntaxial, self-organized, thin displacive veins.—The displacive dolomite crystals should make up syntaxial (or bilateral) veins, as shown in figure 8B. (A syntaxial vein consists of crystals that grow inwards from each side and that are in optical continuity with crystals of the same mineral from the walls; the vein ends up having a center seam.) At time \(t_5\) in figure 5A many such incipient displacive veins would start to grow within the reaction zone. The orientation of the veins was predicted by Merino and others (2006): If the principal stresses are mutually equal the incipient veins tend to be oriented randomly, and upon further growth they intersect each other and generate a so-called “breccia” texture. If the principal stresses are slightly unequal the veins tend to orient themselves normal to the least stress, generating a zebra texture, or rhythmite.
In this case the induced stress exerted by the veins as they thicken (1) should generate stylolites (marked “S” in fig. 8B) in the slice of the host dolomite-I caught between adjacent veins, and (2) should trigger a feedback that by pressure-solution “weeds out” veins that are too close to their neighbors, with the survivors of this triage being left more equidistant than before (Merino and others, 2006).

But this displacive dolomite growth phase cannot last long. The transition of replacive to displacive only happens if and when the rate of dolomite-for-calcite replacement reaches a high enough value, $R_{\text{critical}}$, that corresponds to the strain rate for which the dolostone’s viscosity becomes low enough for the growth to be accommodated displacively instead of replacively (fig. 5A). But as soon as the replacement transitions into displacement, pore fluid $\text{Ca}^{2+}$ stops being raised, which shuts off the self-accelerating feedback. The displacive dolomite continues to grow at the very high rate $R_{\text{critical}}$ very fast scavenging all the aqueous Mg from the brine, whereupon the growth rate of dolomite itself comes down to zero abruptly. The displacive veins only have between times $t_5$ and $T$ to grow, figure 5A, a very short interval because the drop in Mg and in rate is practically instantaneous relative to the infiltration velocity. Thus the veins are predicted to be thin, as observed (Merino and others, 2006).

Note that the displacive growth phase can be bypassed altogether. If the $\text{Mg}^{2+}$ concentration in the initial dolomitizing brine is not sufficiently high, then $\text{Mg}^{2+}$ gets used up completely—and the replacement rate is suddenly brought down to zero—before the strain-rate can reach the appropriate $R_{\text{critical}}$ value needed for the rheological transition to displacive growth discussed above. This is the case for the “Mg2” profile in figure 5A, which may explain why the thin displacive vein sets may not occur at all in some burial dolomites, or may occur only at a few, meter-sized regions within huge volumes of dolomite-replaced limestone.

7. Mineral Paragenesis of Each “Pulse”

We have discussed the dynamics of the replacive (± minor displacive) dolomite growth and have seen how the self-accelerating replacement automatically leads to the complete dolomitization of each $(0, L)$ slice by hundreds of pulses, before the reaction zone jumps to the next slice, $(L, 2L)$. But we left out that when the self-accelerating dolomitization shuts itself down by sudden $\text{Mg}^{2+}$ “sequestration” in each pulse, and even before, other minerals must precipitate as well. We look at these in this section. Since pore fluid $\text{Ca}^{2+}$ concentration is necessarily very high when the $\text{Mg}^{2+}$ concentration plummets (fig. 5A), we expect $\text{Ca}^{2+}$ minerals to start forming immediately.

The first is calcite, which should cement the small spaces in the center seam of displacive zebra veins (see fig. 1D-E), and which should back-replace, or “dedolomitize,” some of the replacive and especially the displacive dolomite just made (see figures 169, 170 in Adams and others, 1984). The new model thus naturally explains a feature—growth of so-called late-stage calcite—that had long been puzzling (Spirakis and Heyl, 1995) from the perspective of the prevailing theory of burial dolomitization. Benito and others (2006) have found what we take to be good indirect evidence of dedolomitization in the form of minute dolomite inclusions within late-stage calcite yielding an isotopic composition identical to that of nearby ferroan saddle dolomite crystals. Note that the submicroscopic calcitic slivers that form within the late, fastest-growing dolomite crystals, deforming them into saddle shapes, are driven by the same huge $\text{Ca}^{2+}$ that drives the late-stage calcite; see section on saddle dolomites above.

Fluorite, anhydrite, barite.—If the original brine contains significant fluoride (as the brine from the Smackover Formation in table 1 does), the huge $\text{Ca}^{2+}$ concentration reached should form fluorite, an ore commonly found in MVT deposits associated with burial dolomites, also back-replacing the host dolomite. Similarly, if the interstitial
solution contained sulfate and barium, and if this sulfate has not been appreciably reduced—see next section—then minor anhydrite and barite (fig. 7E) would also be expected to form at some place(s) within the current \((0,L)\) pulse, at around or right after time \(T\) in each pulse, the moment when \(\text{Mg}^{2+}\) becomes abruptly zeroed out in the pore fluid.

**Sulfates versus sulfides.**—If the brine contains base metals and sulfate (as those in table 1 do), and if the limestone undergoing replacement by dolomite contained appreciable organic matter, several sulfides—galena, sphalerite, pyrite, marcassite—should precipitate during and after the late-replacive and displacive dolomite phases. This is because the organic carbon that is released to the pore fluid by the dolomite-for-limestone replacement would reduce the aqueous sulfate to sulfide, as suggested by Thom and Anderson (2008) and Anderson and Thom (2008). The probable concentrations of sulfate, organic carbon, and sulfide versus time within one pulse of growth are shown qualitatively in figure 5A.

8. **Self-Cannibalization: MVT Ores Are “Swept” Downflow**

When much of the limestone slice currently under dolomitization is dolomitized, the dolomite and ores of late pulses may partly replace ores and dolomite, respectively, formed in earlier growth pulses of the slice. Several examples are shown in figure 7. Dolomite replaces ores at Galmoy (Ireland) and Riopar (Albacete, Spain), figures 7A and 7B. Sphalerite replaces dolomite I and II at San Vicente, Peru, figures 7C and 7D. Barite and fluorite replace dolomite-II at the Hammam-Zriba Mine (Zaghouan district, Tunisia), figure 7E. Fluorite replaces dolomite at Sierra de Gádor (Almería, Spain), and at the Dau Range (Carinthia, Austria), see Zeeh (1995). At the Polaris deposit (Canadian Arctic Archipelago) dolomite was dissolved or replaced by sulfide-precipitating fluids (Savard and others, 2000). For the Buick mine of southeast Missouri Sverjensky (1981) gave and cited petrographic evidence that both galena and sphalerite have precipitated and dissolved repeatedly during ore formation, and suggested that repetitive precipitation and dissolution of galena was probably an important process during the formation of the entire Viburnum Trend. We have seen above that some late-stage calcite produced at the end of late pulses may “dedolomitize” some dolomite.

As a result of all these replacements, the aqueous components released to the interstitial brine—mainly \(\text{CO}_3^{2-}, \text{Mg}^{2+}, \text{Zn}^{2+}\) and \(\text{S}^{2-}\)—are advected forward and promote reprecipitation of more dolomite and/or sphalerite and other ores downflow, generating a larger and larger ore deposit always associated with the current position of the advancing dolomitization front. This progressive accumulation of a scarce mineral at a moving reaction front was modeled quantitatively for a generic, simplified case by Ortoleva and others (1986). We have loosely adapted their dynamic model to dolomitization in figure 9, which shows that at any given time most or all of the ore formed previously (both in previous pulses of the same slice and in previous slices) accumulates at the current front. In fact Harper and Borrok (2007) have reported that the largest zinc and lead deposits of the midcontinent of the U.S. do occur at the last position of the dolomitization front, against the undolomitized limestone.

9. **Geochemical Features of the Dolomite Growth**

Since the displacive dolomite grows continuously on replacive dolomite, it is not surprising that they have been found to have the same or very similar geochemistry. Previous investigators supporting the “two-dolomites” model have been understandably intrigued by the observed geochemical similarity, and have endeavored to explain it. For example, Lonnie and Machel (2006) proposed that the dolomite-I is altered hydrothermally by the same solution that deposits the dolomite-II. However, the fact
that the dolomite-for-calcite replacement is self-accelerating can be expected to have a
differential kinetic effect on the contents of trace elements and O, C isotopes
incorporated by the growing dolomite. If heavy carbonate ions, $^{13}\text{C}^{18}\text{O}_3^{2-}$, move
more sluggishly than light ones, $^{12}\text{C}^{16}\text{O}_3^{2-}$ (O’Neil, 1986, p. 2), then we can expect
that a dolomite crystal that grew at the high growth rate corresponding to times
$t_4$ and $T$ in figure 5A should have lower $\delta^{18}\text{O}$ and $\delta^{13}\text{C}$ values than another that had grown at
a lower rate, say, that between times $t_1$ and $t_3$. $\delta^{18}\text{O}$ and $\delta^{13}\text{C}$ values for four pairs of
replacive and displacive dolomite crystals from the dolomitized Cretaceous limestones
of the Basque-Cantabrian basin are given in table 3 (Simón and others, 1999), with the
replacive and displacive crystals of each pair only a few centimeters from each other, do
indeed show the predicted trend: the replacive dolomite of each pair is a few permil
higher than the nearby displacive dolomite crystal, which according to the new

Table 3

Comparison of $\delta^{18}\text{O}$ (% VSMOW) and $\delta^{13}\text{C}$ (% PDB) values for four pairs of nearby replacive (R, or dolomite-I) and displacive (D, or dolomite-II) dolomite crystals from dolostones in the Cretaceous Basque-Cantabrian basin, northern Spain

| Sample#    | $\delta^{18}\text{O}$ | $\delta^{13}\text{C}$ |
|------------|----------------------|----------------------|
| C98-15b-R  | 19.2                 | 0.2                  |
| C98-15a-D  | 15.5                 | -0.8                 |
| C98-53b-R  | 19.9                 | 1.5                  |
| C98-53a-D  | 14.5                 | 0.6                  |
| C98-59b-R  | 18.3                 | 0.5                  |
| C98-59c-D  | 14.8                 | 0.4                  |
| S98-23b-R  | 20.3                 | 2.8                  |
| S98-23a-D  | 15.7                 | -2.4                 |

Data from Simón and others (1999).
In each pair the displacive (D) crystal is isotopically lighter than the nearby replacive (R) crystal.
dynamic model grew at a much higher rate. The close proximity of the two crystals of
each pair makes it unlikely that the Δ18O and Δ13C differences could be due to a
temperature difference between their locations. The same trend—isotopically slightly
lighter dolomite II—was found also by Fontboté and Gorzawski (1990) for the burial
dolostones at San Vicente, Peru, by Gasparrini and others (2006) for burial dolostones
of the Cantabrian Zone, northern Spain, and by Mattes and Mountjoy (1980).

Since the Ca2+ concentration in the pore fluid increases exponentially along with
the growth rate, very-fast-growing dolomite crystals (those growing between times \( t_4 \)
and \( T \) in fig. 5A) must be more Ca-rich (and thus deformed) than those that grew
earlier, between times \( t_1 \) and \( t_4 \).

Similarly, the degree of trace element incorporation should differ considerably
from fast-growing to slow-growing dolomite crystals that are near each other, warrant-
ing caution in dealing with trace element and fluid inclusion analyses, which should
not be interpreted as reflecting equilibrium. In particular, the ratio Mn/Fe, which is
thought to be one of the controls of cathodoluminescence in dolomite crystals, may be
expected to vary wildly especially for the higher rates of dolomite growth at times \( t_4 \) to
\( T \) (fig. 5A), potentially leading to sudden changes in cathodoluminescence in the
saddle dolomites but not in nearby replacive dolomite crystals formed from the same
solution. The observation that the dolomite-I and dolomite-II have different cathodo-
luminescence (for example, Gregg and others, 2001; Chen and others, 2004; Gaspar-
rini and others, 2006; Lonnee and Machel, 2006; Choquette and Hiatt, 2008), could be
explained by the differential effect of self-accelerating kinetics on the incorporation of
Fe and Mn as traces.

The dolomite- and calcite-undersaturated, and thus very carbonate-poor, property
of the dolomitizing brine—see probable analogues in table 1—suggests that Δ13C
values of the dolostone should be determined by those of the host limestone, since the
brine can contribute only negligible carbonate to the dolostone. This is borne out by
the analyses of Land (1980), Montañez and Read (1992), Hitzman and others (1998),
Gasparrini (ms, 2003), and Carmichael and others (2008), among others.

10. Lobed Dolomitization Front and the Reactive Infiltration Instability

The large porosity created by limestone dissolution via reaction (1) in each slice
should trigger the reactive infiltration instability, first identified and modeled by Chadam
and others (1986) and Ortoleva and others (1987b) and proposed recently to generate
the funnels and sinkholes characteristic of karst (Merino and Banerjee, 2008). The
dissolution porosity created in each position of the reaction front attracts additional
brine flux, which in turn carries out faster the dolomitization reactions described in
this chapter, increasing faster the creation of further new dissolution porosity, thus
permeability. The basic consequence of this instability (for example, Wei and Ortoleva,
1990; and Aharonov and others, 1997) is that the reaction zone, even if initially planar,
tends to become fingered in the general direction of the brine's flow. Then the
competition for reactive flux continues among the fingers themselves, leading to
successive jumps in the size and spacing of the lobes (Szymczak and Ladd, 2006).
Actual fingering of the dolomitization front has been described by Mattes and
Mountjoy (1980) for the Miette buildup of Alberta; Wilson and others (1990) for the
Latemar buildup of northern Italy; Gasparrini and others (2006) for Upper Carbonifer-
erous carbonates of the Cantabrian Zone of northwestern Spain (fig. 2A); and many
others.

11. Spatial Distribution of Dissolution Porosity

Because for every pulse the initial fast dissolution of limestone takes place
immediately upon entry of the calcite-undersaturated, Mg-rich brine into the current
reaction zone, we expect that there would be a porosity maximum at the entry to each
limestone slice dolomitized. The small circles in the entry points of the slices already
dolomitized and of the “current slice,” figure 6, represent that dissolution porosity
maximum. If each slice turns out to be several meters thick (see table A1, Appendix 2)
then we would expect a spatial dissolution porosity periodicity of several meters.
Interestingly, this prediction may be confirmed by Budd and others (2006), who have
reported porosity periodicities on several scales, up to several meters, in fair agreement
with tentative calculations of \( L \) in table A1.

**V. Discussion and Future Work**

*A serendipitous discovery.*—The new dolomitization model is a forward, nonlinear
model that predicts a string of linked loops and cyclicities (figs. 5C and 6) and
self-cannibalization (fig. 9), the whole driven by disequilibrium between the dolomitiz-
ing brine and a platform limestone. Based on two postulates and on the operation of
several feedbacks, the model generates a self-organized dynamics many of whose
predictions and consequences are confirmed by observation. The first postulate—that
the dolomitizing brine must be Mg-rich and *undersaturated* with both calcite and
dolomite—was made because, if the brine were dolomite-supersaturated as assumed in
the literature, *any* rock type could in principle be dolomitized, not just limestones. The
undersaturation with calcite, combined with the fast kinetics of calcite dissolution,
provides the crucial “kick” to the system that starts the process (but only in limestones)
resulting right away in dissolution porosity and in supersaturation with dolomite,
which in turn, when combined with the self-accelerating feedback via \( \text{Ca}^{2+} \) inevitably
involved in the dolomite-for-calcite replacement (Section IV-3), triggers self-organized
dolomitization in both time (in the form of many repeated pulses per limestone slice)
and space (in the form of successive slices). The second postulate—that replacement
happens by guest-growth-driven pressure solution of the host mineral—has been
justified in detail in Section III.

Remarkably, the assumed chemistry of the dolomitizing brine and the replace-
ment physics of the second postulate, combined with the self-accelerating feedback via
\( \text{Ca}^{2+} \) involved in the dolomite-for-calcite replacement (Section IV-3), and with the
non-newtonian rheology of crystalline carbonates (Section IV-5), also accounts for a
suite of heretofore enigmatic textures, microstructures, parageneses, and lithological
association with MVT ores that are characteristic of burial dolostones. That suite of
properties includes: the *displacive* nature of dolomitic zebra and breccia veins; a
replacive/displacive contact that is systematically *seamless*; the *saddle shape* and extra-
calcic composition of the displacive dolomite crystals and of the replacive crystals
formed late in each pulse; the displacive veins are always thin and equidistant (by a
“triage” or “weeding” feedback described by Merino and others, 2006); that the growth
of dolomite must periodically shut itself down and cause immediate growth of
“late-stage” calcite and ores; that the late dolomite of each pulse has lower \( \delta^{18} \text{O} \) and
\( \delta^{13} \text{C} \) than the early dolomite; that the MVT ores tend to be concentrated in the last
position of the dolomitization front; that dissolution porosity should tend to be
spatially periodic. The feedback causes dolomite to grow even though Mg/Ca *decreases*
during any one pulse (see fig. 5A), illustrating how disequilibrium plus feedback
trumps the conventional dolomitization condition, that Mg/Ca > \( K_{sp} \). The growth only
stops when Mg\(^{2+}\) is abruptly and totally scavenged from the current packet, or “filling”
(table 2), of aqueous solution.

All of the pieces of the new dynamics work together to form a consistent model. If
any of the postulates were wrong, or if the dolomite-for-calcite replacement were not
self-accelerating, or if it were not self-accelerating *via Ca\(^{2+}\)*, or if calcite dissolution
were not inherently very fast, or dolostones not strain-rate-softening—the model would
not work.
Self-organization.—Since the new model involves disequilibrium (between the dolomitizing brine and a limestone) and feedbacks, the two necessary conditions for geochemical self-organization (Merino, 1984; Ortoleva and others, 1987a; Merino and Wang, 2001), it is not surprising that the model predicts that dolomitization should be self-organized, and self-organized in several ways at once. First, as noted, the self-accelerating-replacement feedback leads to a temporal self-organization in the form of many successive “pulses,” or spurts, of mineral growth in each “slice,” even under steady infiltration. The feedback, because it couples the replacement rate and the pore fluid Ca$^{2+}$ (Section IV.3), causes each pulse to consist of the same paragenesis, namely replacive dolomite with small amounts of ± displacive dolomite + calcite ± ores. Each pulse however replaces less than one percent of the limestone in the slice. Each pulse is triggered by the supersaturation with dolomite caused by dissolution of some unplaced limestone by a new packet of brine. The pulses repeat themselves (probably in the hundreds, see table A1) until the current limestone slice is completely replaced, providing a straightforward way to account for the precipitation of the layered sphalerite aggregates common in MVT deposits. The predicted self-organization in the form of pulses thus makes it unnecessary to postulate repetitive seismic pumping of brine (for example, Sibson, 1994) or episodic fault-related brine flow (McLimans and others, 1980) to account for the sphalerite layering. There is also a spatial self-organization in the form of sequential “slices” with spatially periodic dissolution-porosity highs concentrated preferentially at the entry region of each slice, as shown in figure 6 and Section IV-11, explaining the porosity periodicities detected by Budd and others (2006).

Second, the rheologic-kinetic feedback described in sections IV-5 and -6, causes the replacive dolomite growth to become displacive (though only for a very short time; see fig. 5A), and this leads to the formation of sets of thin veins that if parallel become roughly equidistant through a “weeding” or “triage” feedback described by Merino and others (2006), forming the so-called zebra veins found in most burial dolomites. The zebra veins of a set are simultaneous, and do not result from repeated episodic flows.

Finally, the reactive-infiltration instability causes dolomitization to proceed in self-organized large fingers and lobes (Section IV-10).

Burial dolomitization is fast.—We deduced that both the coupled pore-fluid Ca$^{2+}$ and dolomite-for-calcite replacement rate grow exponentially on time as the dolomitizing brine infiltrates the limestone, during each pulse (see Appendices 1 and 2). The time, $T$, at which the replacement rate becomes so high as to suddenly use up all the aqueous Mg$^{2+}$ contained in the current packet of interstitial brine, denotes the duration of one pulse. It is calculated with equation A7 and given in table A1 (Appendix 1) for each of two estimates of the 100 °C dolomite growth rate constant, and for a brine with $10^4$ mg Mg/l infiltrating a 0.2 porosity limestone at 10 cm/year. The calculated $T$ is 5.5 and 55 years, respectively, which multiplied by the assumed infiltration velocity yield lengths $L$ of 0.5 and 5.5 meters, respectively. ($L$ is the predicted thickness of a slice.) Since each pulse affects only 0.5 percent of the slice, about 200 pulses (each lasting $\ll$100 years) will take place automatically until the dolomitization of the slice is completed—in only $<20,000$ years. A faster advection velocity would proportionally increase the predicted thickness of a slice, $L$, without changing $T$. Lower dolomite rate constant and/or lower surface area increase both $T$ and $L$. The new model poses no restrictions as to the type and timing of the advection

1 This time $T$ for simplicity neglects the time $\Delta T$ (fig. 5A) required for the growth of the late-stage calcite + ores towards the end of each pulse.

2 Recall that each pulse is triggered by the dissolution of some limestone by fresh brine at the entry into the slice.
that brings the deep hot dolomitizing brine to a platform limestone. The predicted fast
dolomitization implies that the dynamic process is probably essentially isothermal.

Syntaxial veins.—The new model explains physicochemically why zebra veins are
displacive and how they form in burial dolomites. The zebra veins are of the kind called
syntaxial by structural geologists. Wiltschko and Morse (2001) and Fletcher and Merino
(2001) independently showed that many bilateral veins grow displacively, pushing
aside their walls through the crystallization stress generated by the inward growth itself.
Merino and others (2006) gave evidence that dolomitic zebra and breccia veins are
indeed displacive. What the new dolomitization dynamics now shows is that the
replacive dolomite-for-calcite growth, because it is self-accelerating, becomes eventual-
ly fast enough to lower the dolostone viscosity sufficiently to permit the replacive
growth to seamlessly become displacive—and generating a vein oriented at right
angles to its growth direction, as shown in figure 8B. The conventional view (for
example, Bons, 2000)—that a syntaxial vein grows by repeated cycles of cracking and
cementation, as per the so-called crack-seal mechanism—rests on the assumption that
there is an exterior force that, conveniently, repeatedly cracks the rock at the place
where, and at the time when, the vein is ready to grow by precipitation of cement, and
does not explain why the vein consists of the same mineral as its walls, or why the
contact between wall and vein is seamless.

Implications for reaction-transport modeling.—Existing quantitative models of dolomiti-
zation (for example, Wilson and others 2001; Caspard and others, 2004; Whittaker and
others, 2004; Jones and Xiao, 2005; and references therein) have explored the ability
of different large-scale brine flow regimes—reflux, gravity, thermal convection, tec-
tonic squeezing, compaction—to drive dolomitization, incorporating premises of the
prevailing model that we have discussed in Section II, such as that dolomitization is
effected by dolomite-supersaturated brines through the mass balance $2\text{calcite} + \text{Mg}^{2+} = \text{dolomite} + \text{Ca}^{2+}$ or that the dolomite-I and the dolomite-II are produced by
separate brines or at different times. Similarly, there are models quantifying the
formation (by cooling of the ore fluid; by mixing of hydrothermal fluids; by regional
brine flows) of Mississippi-Valley type ore deposits but without accounting for their
association with burial dolostones (see Bethke and Marshak, 1990; Garven and others,
1993; Plumlee and others, 1995; Corbella and others, 2004; among others). We do not
discuss those models here. We suggest instead that the dynamics presented here
provides a blueprint of the brine chemistries, driving forces, mineral reactions, and
feedbacks that should be included in future quantitative reaction-transport models of
dolomitization and MVT mineralization. Crucial in such models will be to monitor the
varying aqueous speciation of the interstitial brine during the precipitation of the
characteristic paragenesis that constitutes each “pulse” (namely, replacive dolomite ±
minor displacive dolomite + calcite ± ores, Section IV.7). The aqueous speciation
should vary during each pulse because of the exponential increase in pore-fluid $\text{Ca}^{2+}$,
the consumption of $\text{Mg}^{2+}$ and carbonate by the dolomite-for-calcite replacement
expressed by equation (4), its effect on pH, and the increasing reduction of aqueous
sulfate by the organic matter released from the limestone by the replacement itself.
The varying aqueous speciation during each pulse could help determine (1) what
minimum Mg concentration the dolomitizing brine should contain for the rheological
transition from replacive to displacive dolomite to be reached; (2) whether aqueous
carbonate could become a limiting factor in the formation of dolomite during each
pulse even before $\text{Mg}^{2+}$ does, since the original brine is carbonate-poor and the
replacement mass balance (equation 4) consumes carbonate; (3) how early or late
within the course of a pulse will the release of organic matter from the limestone start,
as well as the reduction of brine sulfate to sulfide by it, and the growth of Zn and Pb
sulfides.
An interesting possibility for future quantitative reaction-transport models of dolomitization based on the dynamics presented here will be to try to calculate the amount of ore precipitation per pulse and the degree of ore “sweeping” needed to form a large deposit downflow.

Finally, there is the matter of modeling replacement itself as part of the reaction-transport modeling of dolomitization or any other process where replacements are known to take place. Since during any replacement the volumetric rates of guest growth and host dissolution are forced by the crystallization stress to be equal to each other at every moment (see Section III and fig. 3), reaction-transport modeling of a replacement should ensure that equality—an equality that the standard kinetic laws for guest and host would be unable to bring about by themselves if replacement is viewed as resulting by dissolution-precipitation. In modeling by reaction-transport the replacement of limestone by kaolinite that brings about the formation of terra rossa, Banerjee and Merino (2011) used rate laws from Fletcher and Merino (2001) to implement the desired rate equality. In the case of dolomitization, the equality of rates should take place even as they grow exponentially by the self-accelerating feedback.

Predicted periodic dissolution porosity.—Another interesting aim of future quantitative reaction-transport models might be to calculate the amount and perhaps spatial distribution of dissolution porosity. As noted, a basic feature of the dynamic model is that the considerable dissolution porosity carried out by the calcite-undersaturated dolomitizing brine when the brine enters the current limestone slice tends to be concentrated, for each pulse, at the entry into the slice (Section IV-11), and therefore would tend to be accumulated in the entry region of each slice. The predicted dissolution porosity should therefore be periodically distributed in space with a period roughly equal to \( L \), the thickness of the slices, as shown schematically in figure 6. The predicted periodic porosity is confirmed by the interesting data presented by Budd and others (2006), but only in part, since they tease from their data several porosity periodicities, not just one. Therefore, future quantitative models could aim at improving the calculation of \( L \) that we present in table A1, Appendix 2. This would necessitate reliable estimates for the specific surface area and high-temperature growth rate constant of dolomite, and for the brine velocity through the limestone.

VI. SUMMARY OF THE MODEL

We propose a new dynamic model of burial dolomitization which is based on two postulates. The first is that the dolomitizing solution must be Mg-rich but undersaturated with respect to both dolomite and calcite. The second is that the replacement of limestone by dolomite takes place not by dissolution-precipitation as usually assumed, but by dolomite-growth-driven pressure solution of the calcite host.

The postulated brine dissolves some calcite as soon as it enters a limestone, generating dissolution porosity in the entry region and—only then—supersaturating itself with dolomite. Dolomite then grows as the brine infiltrates the limestone, pressure-dissolving, and thus replacing, more calcite. But the dolomite-for-calcite replacement turns out to be self-accelerating, via \( \text{Ca}^{2+} \): the \( \text{Ca}^{2+} \) released by each increment of replacement must increase the ion-activity product for dolomite and thus rate of the next replacement increment, which increases \( \text{Ca}^{2+} \) further, and so on. As a result both the \( \text{Ca}^{2+} \) pore fluid concentration and the dolomite-for-calcite replacement rate increase exponentially with time, as the brine infiltrates the limestone.

The combination of chemical water/rock disequilibrium and positive feedback brings about temporally self-organized dynamics whereby the self-accelerating replacement takes place in repeated growth-and-replacement pulses, each of which shuts itself down when it becomes so fast that all available aqueous Mg is scavenged abruptly.
Towards the end of each pulse, as the pore fluid gets to contain a huge concentration of Ca$^{2+}$, calcite and other Ca-minerals such as fluorite and anhydrite, are also predicted to crystallize. In addition, if the dolomitizing brine contains Zn, Pb, Fe, Ba, F, sulfate, and other appropriate elements (as do brines from the Smackover Formation in Arkansas and the Central Mississippi Salt Dome basin, which seem to satisfy the first postulate of the model too), Mississippi-Valley-type (MVT) minerals will also precipitate at the end of each pulse. The replacement pulses repeat themselves automatically—each triggered by calcite dissolution by a new packet of brine—until the current slice of limestone under dolomitization is completely replaced. The thickness of the slice is determined by the velocity of brine infiltration and the time it takes for each pulse of replacement to become so fast that it shuts itself off abruptly. The dolomitization front thus remains fixed in space until the slice is completely replaced; only then does it jump to the next. Dolomite formed in late pulses may partly replace ore minerals formed in earlier pulses, partly “sweeping” the ores downflow. For a brine with $10^4$ mg/l of Mg infiltrating a 0.2 porosity limestone at 10 cm/year, each replacement pulse is calculated (table A1) to take place in $<100$ years, and about 200 pulses take place one after another until a limestone slice 0.5 to 5 meters thick is wholly replaced, in $<20,000$ years. (These calculations in table A1 were carried out for two values of the high-temperature rate constant for dolomite adopted loosely from Gautelier and others, 1999.)

Self-organization in pulses and slices thus accounts for: the generation of dissolution porosity, and its periodic distribution of maxima; the complete replacement of each limestone slice by dolomite, developing a sharp field contact between dolostone and unreplaced limestone; formation of late-stage calcite and MVT ores in each pulse; “sweeping” of ores downflow with ore accumulation in the last position of the dolomitization front; formation of fingers of dolomitization via the reactive infiltration instability.

In addition, for each pulse, the combination of the self-accelerating feedback via Ca$^{2+}$ with the known strain-rate-softening, non-newtonian rheology of crystalline carbonates leads inevitably to an unlikely suite of associated textural, paragenetic, and geochemical predictions that are however strikingly confirmed by observation. If toward the end of a pulse the dolomite-for-calcite replacement rate gets to be fast enough to lower the viscosity of the local carbonate rock sufficiently (this needs a threshold concentration of Mg in the brine), then the dolomite growth, which has been replacive until that moment, is predicted to transition seamlessly to displacive. This is when displacive, self-organized zebra veins should form, making themselves equidistant by a secondary, “weeding” feedback described previously (Merino and others, 2006). Simultaneously with the fast dolomite growth near the end of each pulse, and regardless of whether the rheological transition has been reached, the Ca$^{2+}$ pore fluid concentration becomes also so high that it drives substitution of Ca$^{2+}$ for Mg$^{2+}$, or of submicroscopic calcitic slivers, on the structure of the growing dolomite crystals, replacive or displacive, forcing them to grow curved because of the larger size of Ca$^{2+}$. Also, the calcite may grow in interstices of the dolostone, and/or may replace some dolomite, a replacement known as dedolomitization. The predicted association of: displacive, equidistant, thin dolomitic veins displaying seamless contacts with their replacive walls; dolomite crystals that are extra-calcic, curved, and slightly lower in $\delta^{18}O$; and minor amounts of “late-stage” calcite, “dedolomite,” and other Ca-bearing minerals (fluorite, anhydrite) and other MVT ores (galena, sphalerite) agrees very well with observations, and suggests that the new model captures the basic mechanisms, drives, and interactions that lend burial dolomitization and its often associated MVT mineralization its geological uniqueness.
where the equilibrium constant is consistent with molality units but dimensionless, the activities of Mg$^{2+}$ and CO$_3^{2-}$ are also dimensionless and the activity of Ca$^{2+}$ is written as (molarity of Ca$^{2+}$ in moles/ℓ) × (Ca$^{2+}$ activity coefficient, in ℓ/mole) assuming for simplicity that molality is approximately equal to molarity. The quantities ($k_{dolo}$ and $R_{dolomite}$) have units of moles/ℓ second. Since the Mg$^{2+}$ and CO$_3^{2-}$ activities remain roughly constant during, say, the first two-thirds of each pulse of replacement (see eqs 5, 5a and text, and fig. 5B), and since the supersaturation $\Omega$ becomes ≫1, the “−1” can be neglected in (5a):

$$R_{dolomite} \approx k_{dolo} S_0 (\Omega - 1),$$  (5a)

where

$$\Omega = Q / K_{eq}^{dol} = (m_{Ca^{2+}} \cdot Y_{Ca^{2+}})(a_{Mg^{2+}})(a_{CO_3^{2-}})^2 / K_{eq}^{dol},$$  (5)

By equations (5, 5a) the growth rate of dolomite is

$$R_{dolomite} = k_{dolo} S_0 (\frac{d m_{Ca^{2+}}}{d t})_{Ca^{2+}}.$$  (A1)

Note that the local mass conservation equation for Ca$^{2+}$ probably involves little or no transport because, as the incipient dolomite growth is starting to replace calcite everywhere, both the Ca$^{2+}$ and Mg$^{2+}$ profiles remain horizontal and preclude diffusion of Ca$^{2+}$ or Mg$^{2+}$ forward or backwards. Thus,

$$\frac{d m_{Ca^{2+}}}{d t} = h R_{dolo},$$  (A2)

where $h$ is the stoichiometric coefficient of Ca$^{2+}$ in the replacement mass balance (4), 0.74 moles of Ca$^{2+}$ released to pore fluid/dolomite formula, and molality $m$ is again expressed in molarity units. Putting (A1) into (A2), and moving $m_{Ca^{2+}}$ to the left side and $d$ to the right, one obtains

$$\frac{d m_{Ca^{2+}}}{m_{Ca^{2+}}} = h k_{dolo} S_0 (\frac{d Y_{Ca^{2+}}}{d t})(a_{Mg^{2+}})(a_{CO_3^{2-}})^2 / K_{eq}^{dol}) dt$$  (A3)

$$= h k_{dolo} S_0 H dt$$

where

$$H = (\frac{Y_{Ca^{2+}}}{a_{Mg^{2+}}})(a_{CO_3^{2-}})^2 / K_{eq}^{dol}.$$  (A4)
By integration of (A3) from \( m_{\text{Ca}^2+/\text{H}_11001} \), initial to \( m_{\text{Ca}^2+/\text{H}_11001} \) and from \( t/\text{H}_11005 \) to \( t/\text{H}_11005 \), one gets
\[
\frac{m_{\text{Ca}^2+/\text{H}_11001}}{\text{exp}(t/\text{H}_20849)} = \frac{m_{\text{Ca}^2+/\text{H}_11001}}{\text{exp}(t/\text{H}_9270/\text{H}_20850)}.
\]
where \( \phi \) has units of time and the \( 1000 \) factor converts liters in the \( \text{H}_9270/\text{H}_11005/\text{H}_20849 \) units to cm\(^3\) in the \( \phi \) units. From (A1, A5),
\[
\text{R}_{\text{dolo}}/\text{H}_11015/\text{H}_20849 \text{kdolo}S_0 \text{H}_{\text{mCa}^2+/\text{H}_11001} \text{exp}(t/\text{H}_9270/\text{H}_20850).
\]
In short, because of the feedback between the two, both the pore-fluid \( \text{Ca}^2^+ \) concentration and dolomite growth rate increase exponentially with time, eqs (A5, A7), as shown in figure 5A.

### Appendix 2

#### Replacement of One Limestone Slice By Many Pulses

With reference to figure 6, replacement of a limestone slice takes place completely by many sequential pulses of dolomite growth (plus minor calcite and ores). Each pulse of dolomite precipitation grows—at a self-accelerating rate—from the Mg-rich pore brine contained in the slice’s pores, lasts a time \( T \) calculated in Appendix 1, and has an approximate length \( L = \nu T \). The slice becomes Mg-deficient after a time \( T \) when its aqueous Mg is used up abruptly; \( L \) is the thickness of limestone slice affected by each pulse in the time \( T \); \( N \) is the number of pulses needed to replace each slice of thickness \( L \) completely.

| Parameters Used | Parameters Used |
|-----------------|-----------------|
| \( h \) | 0.75 mol(Ca)/mol(do) | \( h \) | 0.75 mol(Ca)/mol(do) |
| \( b \) | 64 cm\(^3\)(do)/mol Mg | \( b \) | 64 cm\(^3\)(do)/mol Mg |
| \( S_0 \) | 100 cm\(^3\)/cm\(^3\) | \( S_0 \) | 100 cm\(^3\)/cm\(^3\) |
| \( \phi(dolo) \) | 0.0156 mol/cm\(^3\) | \( \phi(dolo) \) | 0.0156 mol/cm\(^3\) |
| \( \text{veloc, } \nu \) | 10 cm/yr | \( \text{veloc, } \nu \) | 10 cm/yr |
| \( m(Mg)_{\text{initial}} \) | 0.4 mol/l = \( 10^4 \) mg/l | \( m(Mg)_{\text{initial}} \) | 0.4 mol/l = \( 10^4 \) mg/l |
| \( \gamma(Ca, Mg) \) | 0.3 (mol/l) | \( \gamma(Ca, Mg) \) | 0.3 (mol/l) |
| \( \text{activ(CO3)} \) | 3.16E-09 | \( \text{activ(CO3)} \) | 3.16E-09 |
| \( \text{activ(Mg)} \) | 1.20E-01 | \( \text{activ(Mg)} \) | 1.20E-01 |

| Calculated Quantities | Calculated Quantities |
|-----------------------|-----------------------|
| \( H/\text{eq A4} \) | 5.32E+01 (mol/l) | \( H/\text{eq A4} \) | 5.32E+01 (mol/l) |
| \( \tau/\text{eq A6} \) | 2.50E+07 s = 0.8 yrs | \( \tau/\text{eq A6} \) | 2.50E+08 s = 8 yrs |
| \( T/\text{eq A7} \) | 1.73E+08 s = 5.5 yrs | \( T/\text{eq A7} \) | 1.73E+09 s = 55 yrs |
| \( L = \nu T \) | 55 cm | \( L = \nu T \) | 5.5 m |
| \( R(dolo), \text{eq A7} \) | 5.32E-08 mol/cm\(^3\) s | \( R(dolo), \text{eq A7} \) | 5.32E-09 mol/cm\(^3\) sec |
| \( R'(dolo) \) | 0.088 cm/month | \( R'(dolo) \) | 0.009 cm/month |
| \( N = 1/\phi_{\text{cum}b} \) | 195 pulses | \( N = 1/\phi_{\text{cum}b} \) | 195 pulses |

* Bowers and others (1984).
** Rate-constant values were loosely extrapolated from values graphed by Gautelier and others, 1999. \( T \) is the time it takes for the replacement rate \( R \) to become 1000 times greater than it was at the start of each growth pulse, by which time aqueous Mg is used up abruptly; \( L \) is the thickness of limestone slice affected by each pulse in the time \( T \); \( N \) is the number of pulses needed to replace each slice of thickness \( L \) completely.

By integration of (A3) from \( m_{\text{Ca}^2+/\text{initial}} \) to \( m_{\text{Ca}^2+} \) and from \( t = 0 \) to \( t = t_1 \), one gets
\[
m_{\text{Ca}^2+} = m_{\text{Ca}^2+/\text{initial}} \exp(t/\tau),
\]
where
\[
\tau = (1000h_{dolo}S_0H)^{-1}
\]
has units of time and the 1000 factor converts liters in the \( H \) units to cm\(^3\) in the \( (kdoloS_0) \) units. From (A1, A5),
\[
R_{dolo} = (k_{dolo}S_0H m_{\text{Ca}^2+/\text{initial}}) \exp(t/\tau).
\]
In short, because of the feedback between the two, both the pore-fluid \( \text{Ca}^2^+ \) concentration and dolomite growth rate increase exponentially with time, eqs (A5, A7), as shown in figure 5A.
\[
\epsilon_{\text{Mg}} = 10,000 \text{ mg/\ell} = (10/24) \times 10^{-3} \text{ mol Mg/cm}^3 \text{ pore} = 0.4 \times 10^{-3} \text{ mol Mg/cm}^3;
\]

\[
\phi = 0.2 \text{ cc pore/cc bulk rock;}
\]

and
\[
b = 64 \text{ cm}^3 \text{ of dolo/mol Mg}^{2+},
\]
a fraction \( f = 0.2 \times (0.4 \times 10^{-3} \text{ mol Mg/cm}^3 \text{ pore}) \times (64 \text{ cm}^3 \text{ dolo/mol Mg}) \approx 0.005 = 0.5 \text{ percent is}
dolomitized by one pulse (in time \( T \)), and \( N = 1/f \approx 200 \) fillings, lasting roughly a time \( 200T \), are needed to
completely dolomitize that one slice, or reaction zone, of limestone. With parameter values adopted in table
A1 and equations A4, A6 we calculate estimates of \( H, \tau \), and the time \( T \) needed for \( R(\text{dolo}) \) to become 1000
times greater than it was at \( t = 0 \). The value of the dolomite growth rate at \( t = T \) is given in mole/cm\(^3\) sa s as
\( R(\text{dolo}) \) and in cm/month as \( R'(\text{dolo}) \). All the calculations are performed for two estimates, \( 10^{-14} \) and \( 10^{-15}
\text{ mol/cm}^2 \text{ s} \), of the high-temperature dolomite-growth rate constant in equation 5a. The dolomite rate-

constant values chosen were loosely extrapolated from rates graphed vs pH by Gautelier and others (1999,
their fig. 7). The time \( T \) for one growth pulse is 5.5 and 55 years, respectively, yielding, for an arbitrary brine
velocity of 10 cm/year, a slice thickness, or thickness of the reaction zone, of 55 cm and 5.5 meters,
respectively.

Since dolomitization of limestone self-accelerates exponentially on time, brine with concentration \( \epsilon_{\text{Mg}} \)
(mol Mg/cm\(^3\) of pore fluid) infiltrates limestone of porosity \( \phi \) over a thickness \( \nu T \), where \( T \) is the time it takes
for the dolomitization to become so fast that all the aqueous Mg in the segment is used up instantaneously,
and \( v \) is the brine velocity through the limestone pores. That length, \( \nu T \), is the thickness \( L \) of the reaction
zone. The reaction zone remains stationary until all the limestone in it is completely dolomitized. Each
successive packet of Mg brine that fills the pores of the reaction zone drops a pulse of dolomite which
replaces only a fraction \( f \) of the current limestone slice. \( N = 1/f \) “fillings” are needed to dolomitize the slice
completely.

The volume of dolomite made by one “filling” of the limestone volume \( \nu TA \) with brine is:
\[
V_{\text{dolo},T} = (\text{flux}) \times (\text{time } T) \times (\text{cross section } A) \times (b) = (\phi \epsilon_{\text{Mg}}) TAb,
\]

where \( b = 64 \text{ cm}^3 \text{ dolo/mol Mg}^{2+} \).

The initial bulk volume of limestone containing \( V_{\text{dolo},T} \) is:
\[
V_{\text{limest}} = \nu TA.
\]

Thus the volume fraction dolomitized in each pulse is:
\[
f = (\phi \epsilon_{\text{Mg}}) TAb/\nu TA = \phi \epsilon_{\text{Mg}} b.
\]

(Note that this is independent of \( T \), the duration of one pulse, which can be calculated from Appendix 1.
Once \( T \) is known we can calculate the length of one “pulse,” or thickness of the “reaction zone,” equal to \( \nu T \).)
The number of fillings needed to dolomitize one reaction zone completely is
\[
N = 1/\phi \epsilon_{\text{Mg}} b.
\]
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