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Title: The role of Mg in the crystallisation of monohydrocalcite.

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Abstract: Monohydrocalcite is a member of the carbonate family which forms in Mg-rich environments at a wide range of Mg/Ca ratios ( $Mg^{2+aq}/Ca^{2+aq} \geq 0.17 < 65$ ). Although found in modern sedimentary deposits and as a product of biomineralization, there is a lack of information about its formation mechanisms and about the role of Mg during its crystallization. In this work we have quantitatively assessed the mechanism of crystallization of monohydrocalcite through in situ synchrotron-based small and wide angle X-ray scattering (SAXS/WAXS) and off-line spectroscopic, microscopic and wet chemical analyses. Monohydrocalcite crystallizes via a 4-stage process beginning with highly supersaturated solutions from which a Mg-bearing, amorphous calcium carbonate (ACC) precursor precipitates. This precursor crystallizes to monohydrocalcite via a nucleation-controlled reaction in stage two, while in stage three it is further aged through Ostwald-ripening at a rate of  $1.8 \pm 0.1$  nm/h<sup>1/2</sup>. In stage four, a secondary Ostwald ripening process ( $66.3 \pm 4.3$  nm/h<sup>1/2</sup>) coincides with the release of Mg from the monohydrocalcite structure and the concomitant formation of minor hydromagnesite. Our data reveal that monohydrocalcite can accommodate significant amounts of Mg in its structure ( $\chi_{MgCO_3} = 0.26$ ) and that its Mg content and dehydration temperature are directly proportional to the saturation index for monohydrocalcite (SIMHC) immediately after mixing the stock solutions. However, its crystallite and particle size are inversely proportional to these parameters. At high supersaturations (SIMHC=3.89) nanometer-sized single crystals of monohydrocalcite form, while at low values (SIMHC=2.43) the process leads to low-angle branching spherulites. Many carbonates produced during biomineralization form at similar conditions to most synthetic monohydrocalcites, and thus we hypothesize that some calcite or aragonite deposits found in the geologic record that have formed at high Mg/Ca ratios could be secondary in origin and may have originally formed via a metastable monohydrocalcite intermediate.

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17/02/2013

Dear Executive Editor,

We would like to submit the manuscript listed below for possible publication in *Geochimica et Cosmochimica Acta*.

**Title:** The role of Mg in the formation of monohydrocalcite.

**Authors:** Juan Diego Rodriguez-Blanco, Samuel Shaw, Pieter Bots, Teresa Roncal-Herrero and Liane G. Benning

This manuscript describes an experimental study in which we elucidated the formation mechanism of monohydrocalcite from a poorly-ordered precursor and the role of Mg in its crystallization. Combining in situ synchrotron-based with various off-line laboratory characterizations allowed us to derive complementary quantitative data that explain the monohydrocalcite crystallization via a multiple stage process.

We believe that our paper is of interest to a broad geochemical community and that our results may help explain a number of important biogeochemical processes (including biomineralization and their link to past variations in ocean chemistry).

All authors have read and accepted the manuscript in its current format and we all confirm that this paper represents original work from which no part has been published, nor is being considered for publication, elsewhere.

We suggest **Jacques Schott** as possible Associate Editor.

A few suggested reviewers are included below:

**Prof. Mark Hodson**

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17/02/2013

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Yours sincerely,

Dr. Juan Diego Rodriguez-Blanco.



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Date Oct 15<sup>th</sup> 2013

Dear Frank,

Thank you very much for the comments and suggestions to improve our manuscript. Please find below [in blue](#) the detailed replies to the reviewers' comments. Based on these we have made corrections to the whole manuscript to improve clarity. In our replies we explain in detail the aqueous speciation calculations (done with PHREEQC) and describe how using this approach we calculated the saturation index of monohydrocalcite and other carbonates. We also describe the necessary calculations to determine the Mg content in our monohydrocalcite. These calculations are relatively standard in PHREEQC and thus we did not include them in the Supplementary Information Section. However, if you as the AE consider that this is necessary, we are happy to include these calculations in the Supplementary Information.

In addition we have addressed the comments of Prof Keisuke Fukusjhi regarding (a) the possibility of having metastable nequehonite formed in our experiments, and (b) the speciation of Mg in our monohydrocalcite. All our characterization data clearly indicate an absence of nesquehonite in our experiments and thus our results do not support Prof. Fukushi's hypothesis. However, we have taken his suggestions and ideas into account and discussed them both in the manuscript and in this reply.

We hope now the manuscript is acceptable for publication in GCA.

Yours Sincerely,

Liane G. Benning, Leeds Oct 2013

**Reviewer #1:** I enjoyed reading this clearly written paper that describes an elegant set of experiments monitoring the precipitation of ACC and its subsequent crystallisation to monohydrocalcite. These are techniques I'd love to go and use having read your paper -it is great the way you can follow these reactions in real time.

I have a few trivial queries that it would be good to consider when you have a chance before the final version of the manuscript is produced. I have one substantive point, which is why I've ticked "Substantive modification required" just so the point can be checked by yourselves and the AE, it could be that I'm wrong and it isn't an issue. So, to let suspense build I'll start with a couple of trivial points...

Abstract - SI  $\langle$ subscript $\rangle$  MHC is not defined before it is given - could be worth a mention though an informed reader can work it out.

[This has been now defined in the abstract.](#)

In the Experimental section at the start you describe your experimental protocol for the SAX / WAX. It would be helpful here to say what concentrations you were using - I presume the first line of Table 2 gives this info, but it isn't clear.

Similarly at line 160 when you mention the off line experiments - are these the 350 mM Ca, 150 mM Mg ones.

I think the confusion can be resolved by making clear the concentrations used in this section of text and also altering Table 2 slightly so that it is clear which set of conditions relate to the first experiment (the first line of data) and which to the subsequent off line experiments - the rest of the lines of data.

[We thank the to reviewer for highlighting this issue. We have added all the requested information to the manuscript. Specifically we have text to describe the concentration of ions in the stock solution used in the on-line and off-line experiments, \(lines 136-137 and also 162-163, respectively\), and rewritten the caption for table 2.](#)

This is my big question

Line 175 - I had a bit of a problem here but it might just be me. It's too late now but it would surely have been better to measure the carbonate ion concentrations. You assume that the concentration of  $(\text{CO}_3)_2^-$  is equal to the sum of the concentration of  $\text{Mg}^{2+}$  and  $\text{Ca}^{2+}$ . I wonder 2 things. At the pH you are dealing with, c. 9 won't c. half the carbonate be present as bicarbonate? If that is the case shouldn't you be adjusting the value of  $(\text{CO}_3)_2^-$  you are using. Secondly at pH of 9 won't the concentration of  $\text{OH}^-$  be non-trivial in terms of the mass balance in your system where  $(\text{H}^+) + (\text{Ca}^{2+}) + (\text{Mg}^{2+}) = (\text{OH}^-) + ((\text{CO}_3)_2^-) + ((\text{HCO}_3)_1^-)$ . What difference will this make to all your calculated SIs? I think this is something that you need to address either in the manuscript or to explain to the editor why I'm being stupid and none of this is relevant! This point is the only reason that I've ticked "Substantie modification

required",

The reviewer is correct in her/his assertion that we may not have explained the calculations correctly. In our experiments carbonate was not measured directly in the solutions. The way we determined total carbonate in solution and the carbonate speciation followed 3 steps:

1. After mixing the initial solutions, dissolved [Ca] and [Mg] were measured in the supernatant solution at various time steps as explained in the method section and as shown in Fig 7.
2. Using these values we assumed that all the Ca and Mg removed from our initial solution (prior to mixing) was in the form of Ca/Mg carbonate. Therefore, the number of moles of carbonate removed from solution must be equal to the total moles of Ca + Mg removed from solution.
3. The concentrations of Ca, Mg (measured), total carbonate (calculated) and pH (measured) were then entered into PHREEQC to calculate the speciation of carbonate in solution and the saturation index with respect to MHC and other calcium/magnesium carbonate phases.

We have taken all of these factors into account (as described above) and have updated the manuscript (line 172- 178) to clarify this point.

Lines 198-200 - why the difference in the temperature range used for your amorphous and crystalline solids?

The water content of monohydrocalcite is frequently calculated by TGA above 100 °C because the loss of water from 25 to 100 °C is considered to correspond to adsorbed and not structural water. Other authors we cite in our manuscript also measured the water content of monohydrocalcite using TGA from 100 °C, so we have followed the same procedure to be able to compare all data. See for example:

Hull, H., Turnbull, A. G. (1973). A thermochemical study of monohydrocalcite. *Geoch. Cosmochim. Ac.*, 37, 685-694.

Skinner, H. C. W., Osbaldiston, G. W., and Wilner, A. N. (1977). Monohydrocalcite in a guinea pig bladder stone, a novel occurrence. *Am. Mineral.*, 62, 273-277.

In the case of the amorphous calcium carbonate (ACC) we needed to consider the loss of weight from 25 to 550 °C because this hydrated poorly-ordered precursor is unstable and, depending on its Mg content, it can crystallise at ambient temperatures after a few hours or days. This loss of structural water can occur ambient temperature, hence the reason to use the range 25-550 °C to estimate its water content.

The text in the manuscript has been updated (lines 206 – 211) to clarify these points.

Line 221 - again, perhaps I'm misunderstanding here - you have a formula of  $\text{Ca}_{0.74}\text{Mg}_{0.26}$  etc. surely this indicates that  $X_{\text{MgCO}_3} = 0.26$  but in Table 2 you have  $X_{\text{MgCO}_3} = 0.164$ . is this a typo?

It is not a typo, however we have now clarified it on line 230.

Line 256 - should be "indicating"

Generally actually you need to have "a" in front of words that don't start with a vowel, only use "an" for words that start with a vowel.

Corrected. Thank you.

Line 323 - can you explain the basis for your estimation.

Quantification using X-ray diffraction showed that the final content of hydromagnesite in our experiments was ~4.5 % (i.e. ~ 95.5 % of monohydrocalcite). From these quantitative XRD data and the final solid composition (mixture of monohydrocalcite and hydromagnesite) obtained from chemical analysis, it is possible to calculate the composition the monohydrocalcite.

Line 469 - this section suggests that the previous material related to the experiment described in Table 1 and the first line of Table 2 - I think you need to make this all clearer.

We thank the reviewer for pointing this out. The text has been updated to clarify this point (line 493).

Line 650 -  $\text{PO}_4$  and  $\text{AsO}_4$  don't exist - you need charges on these.

Corrected. Thank you.

Line 694 - big not bug

Corrected. Thank you.

In the figs. could you use different patterning as well as different colours - my b&w print out made it hard to spot the different lines you were talking about.

No details about specific figures are given. However:

1) We have been very careful with colours in the figures and have printed them prior to submission to check that they are understandable and readable in B&W.



2) Furthermore all colour figures also include labels, numeric data, symbols with different sizes and shapes, and even arrows indicating the key information, so the reader can identify the data we are referring to. In some cases we refer to specific stages in the crystallisation of monohydrocalcite, so the data is easier to find.

3) Most of the lines in our pictures (e.g.: Fig 3, 4, 9) need to be continuous lines: it would not make sense to plot fit lines, SAXS patterns or TGA data using different patterning.

table 2 - the column  $SI_{MHC}$  - which solution is this for? Presumably either the initial  $MgCl_2/CaCl_2$  solution or the  $Na_2CO_3$  solution - in either case how can the solution be saturated with respect to MHC - presumably it is the  $MgCl_2/CaCl_2$  solution and the saturation is due to equilibrium with atmospheric  $CO_2$ ? Again, if you could clarify this it would be helpful.

Table 2 of the original manuscript shows calculated  $SI_{MHC}$  in the mixed solutions without any precipitation. This was we could calculate the saturation index of the solution from which the calcium carbonate has formed. This is not “before mixing” and we apologies for this oversight, and the incorrect use of this term. We have updated table 2 and the relevant section in the manuscript to clarify this issue (lines 185-186).

**Reviewer #2:**

This paper describes the role of Mg for MHC formation by using synchrotron based time-resolved XRD and classical off-line approach. The results and implication of the present study are very interesting. However, the discussion is not enough to clarify the role of Mg for MHC formation. The author should be addressed more about the chemical speciation of Mg in MHC.

This paper described that Mg is accommodated in the MHC structure during stage II to III without the sufficient evidences. This implies that Mg does not form the discrete hydrous Mg carbonate during II to III. On the other hand, our recent paper (Nishiyama et al. 2013GCA) showed that the MHC formation requires the paragenesis of hydrous Mg carbonate (most likely fine grained nesquehonite) from the equilibrium calculation. We considered that hydromagnesite is the alteration product from the hydrous Mg carbonate. The authors should discuss more about the chemical speciation of Mg.

On the other hand, I think our conclusion is still valid for the authors experiments and should be considered in the paper. Table 1 shows the SI of MHC and nesquehonite of the solution before the precipitations by using the authors results (Table 2 in the manuscript). The SIs were calculated by using REACT in Geochemists Workbench with "thermo.com.V8.R6+" database. The bicarbonate/carbonate concentrations were calculated from charge balance of the solutions. Although the activity is calculated with extended Debye-Huckel equation but it must be valid for the experiments with lower saturation states. Even for the experiment with highest saturation state, the calculated SI of MHC (3.76) is close to the authors calculation by using Pitzer equation (3.89). This table clearly shows that the formations of MHC correspond to the conditions of equilibrium or superstition with respect to nesquehonite, which is consistent with our conclusion. Table 2 also shows the SIs of relevant minerals from the authors results for the evolution of the aqueous solution compositions during the MHC formation (Table 1 in the manuscript). The results show that the solution before hydromagnesite formation (8h) is almost equilibrium with respect to nesquehonite (SI within  $\pm 0.2$ ). This is also consistent with our results (Nishiyama et al. 2003).

Table 1

Na <sub>2</sub> CO <sub>3</sub> (mM)	Ca (mM)	Mg (mM)	SI MHC before mixing	SI Nesquehonite before mixing	Mineralogy
500	350	150	3.76	1.30	MHC
250	175	75	3.46	1.01	MHC
200	140	60	3.36	0.91	MHC
100	70	30	3.05	0.61	MHC
50	35	15	2.73	0.30	MHC
25	17.5	7.5	2.41	-0.04	MHC
125	8.75	3.75	2.08	-0.39	Mg calcite
5	3.5	1.5	1.62	-0.88	Mg calcite

Table 2

Time (h)	pH	Ca mM	Mg mM	SI MHC	SI hmg	SI nesquehonite	SI aragonite
0.083	9.239	1.25	28.17	0.88	5.54	0.03	1.58
0.5	9.217	0.84	27.75	0.69	5.39	0.00	1.39
2.5	9.214	0.32	28.08	0.27	5.40	0.01	0.97
3	9.203	0.51	31.05	0.49	5.69	0.07	1.20
5	9.167	0.68	34.59	0.63	5.90	0.13	1.33
7	9.171	0.62	35.62	0.60	6.01	0.16	1.30
8	9.111	0.29	25.47	0.16	4.68	-0.12	0.86
9	8.893	0.09	12.64	-0.69	1.28	-0.79	0.01
10	8.646	0.08	12.83	-0.92	0.10	-0.97	-0.22
12	8.609	0.09	13.12	-1.36	-2.99	-1.46	-0.65

We thank Prof Fukusjhi for his review and comments. We have modified our manuscript to take his suggestions into consideration.

We assert, however, that with our new data we have clarified the role of Mg in monohydrocalcite formation because we have shown that:

1. Mg in solution and high supersaturation levels are required to form monohydrocalcite.
2. The crystallisation of monohydrocalcite takes place via the formation of an Mg-bearing poorly-ordered precursor (Mg-ACC). Mg plays an essential role in the stability of this precursor phase controlling the pathway of transformation to monohydrocalcite.
3. Significant amounts of Mg are incorporated into monohydrocalcite structure. The starting concentration of Mg in solution (Mg/Ca/CO<sub>3</sub> ratio) and the initial supersaturation immediately after mixing the stock solutions are key factors that control the content of Mg in monohydrocalcite.
4. The kinetics of monohydrocalcite growth depends on the Mg content in the solid and the concentration of Mg in the aqueous solution.
5. The dehydration process, solubility, unit cell parameters, crystallite size and morphology of monohydrocalcite are dependent on its Mg content.
6. The higher energy required to dehydrate the Mg<sup>2+</sup>-aquo ion compared to Ca<sup>2+</sup>-aquo is essential to understand the longer lifetime of the Mg-ACC precursor and the kinetics of monohydrocalcite growth.
7. Finally, we show a simple methodology to synthesize monohydrocalcite with specific Mg content, particle size and shape by varying the starting Mg/Ca/CO<sub>3</sub> ratio and the initial supersaturation.

Furthermore, Nishiyama et al (2013) GCA state that the formation of monohydrocalcite requires the paragenesis of hydrous Mg carbonate (e.g. nesquehonite). All the different solid characterization techniques applied in our work have however revealed that the only crystalline carbonates that could be unambiguously identified were monohydrocalcite and hydromagnesite. No other phases were detected. In particular:

1. Our XRD data did not reveal the presence of nesquehonite or other crystalline hydrous carbonates at any stage during the replica of the on-line experiment. Also, experiments designed to study the effect of

starting supersaturation in the crystallization of monohydrocalcite did not show any evidence of nesquehonite.

2. An in-depth analysis of the time-resolved synchrotron-based WAXS data did not reveal nesquehonite or other crystalline hydrous carbonates.

3. Our HR-TEM images only showed a poorly-ordered Mg-ACC precursor during the first stage of the experiment, followed by monohydrocalcite (stages II to IV) and hydromagnesite at stage IV. No other solid phases were observed.

However, the calculations of saturation indexes for nesquehonite carried out by Prof. Fukushi show that the aqueous solution was nearly saturated with respect to nesquehonite and therefore suggest the potential formation of this mineral. Our conclusion is that despite the combination of conventional XRD, synchrotron-based WAXS and high-resolution microscopy, which do not show the presence of nesquehonite, we cannot completely discard the fact that small quantities of nanocrystalline nesquehonite or other hydrous carbonate may have been present during the experiment.

We have therefore included this hypothesis in the manuscript (lines 458-467) and also calculated the saturation indexes of nesquehonite in our experiment (now included in Table 1).

Specific comments:

Line 481 and Figure 9a: We observed that Mg content in the solid product decreases by washing with deionized water (probably because of the high solubility of hydrous Mg carbonate coexisting with MHC). The MHC sample used in the experiments by Munemoto and Fukushi (2008) washed by sufficient amount of deionized water. So the Mg content in the sample should not be reflected from the saturation state of initial solution.

This information has been updated in the text. Thank you.

Fig. 8 and Table 1: The SI values of hydromagnesite plotted in the Fig. 8 are different from that in Table 1.

These data is now corrected. Thank you.

**Reference:**

Nishiyama, R., Munemoto, T., Fukushi, K. (2013) Formation condition of monohydrocalcite from  $\text{CaCl}_2\text{-MgCl}_2\text{-Na}_2\text{CO}_3$  solutions. *Geochimica et Cosmochimica Acta*, **100**, 217-231.

# The role of Mg in the crystallisation of monohydrocalcite.

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## ABSTRACT

Monohydrocalcite is a member of the carbonate family which forms in Mg-rich environments at a wide range of Mg/Ca ratios ( $\text{Mg}^{2+}_{\text{aq}}/\text{Ca}^{2+}_{\text{aq}} \geq 0.17 < 65$ ). Although found in modern sedimentary deposits and as a product of biomineralization, there is a lack of information about its formation mechanisms and about the role of Mg during its crystallization. In this work we have quantitatively assessed the mechanism of crystallization of monohydrocalcite through in situ synchrotron-based small and wide angle X-ray scattering (SAXS/WAXS) and off-line spectroscopic, microscopic and wet chemical analyses. Monohydrocalcite crystallizes via a 4-stage process beginning with highly supersaturated solutions from which a Mg-bearing, amorphous calcium carbonate (ACC) precursor precipitates. This precursor crystallizes to monohydrocalcite via a nucleation-controlled reaction in stage two, while in stage three it is further aged through Ostwald-ripening at a rate of  $1.8 \pm 0.1 \text{ nm/h}^{1/2}$ . In stage four, a secondary Ostwald ripening process ( $66.3 \pm 4.3 \text{ nm/h}^{1/2}$ ) coincides with the release of Mg from the monohydrocalcite structure and the concomitant formation of minor hydromagnesite. Our data reveal that monohydrocalcite can accommodate significant amounts of Mg in its structure ( $\chi_{\text{MgCO}_3} = 0.26$ ) and that its Mg content and dehydration temperature are directly proportional to the saturation index for monohydrocalcite ( $\text{SI}_{\text{MHC}}$ ) immediately after mixing the stock solutions. However, its crystallite and particle size are inversely proportional to these parameters. At high

35 supersaturations ( $SI_{MHC}=3.89$ ) nanometer-sized single crystals of monohydrocalcite form,  
36 while at low values ( $SI_{MHC}=2.43$ ) the process leads to low-angle branching spherulites. Many  
37 carbonates produced during biomineralization form at similar conditions to most synthetic  
38 monohydrocalcites, and thus we hypothesize that some calcite or aragonite deposits found in  
39 the geologic record that have formed at high Mg/Ca ratios could be secondary in origin and  
40 may have originally formed via a metastable monohydrocalcite intermediate.

41

42 **Keywords:** monohydrocalcite, carbonates, magnesium, synchrotron, SAXS, WAXS,  
43 scattering.

44

45

## 46 1. INTRODUCTION

47

48 A variety of calcium carbonate minerals are stable at Earth surface conditions. These  
49 include the common polymorphs calcite, vaterite and aragonite, and the less common and  
50 hydrated phases monohydrocalcite ( $CaCO_3 \cdot H_2O$ ) and ikaite ( $CaCO_3 \cdot 6H_2O$ ).  
51 Monohydrocalcite forms in a variety of modern natural environments including saline spring  
52 waters (Ito, 1993), marine polar systems (e.g., Antarctic lakes, or Ikka Fjord, Greenland; Bird  
53 et al., 1991; Dahl and Buchardt, 2006), basaltic caves (Broughton, 1972; Onac, 2000;  
54 Léveillé et al., 2000), cold/humid mine galleries and lacustrine deposits. Representative of the  
55 latter environment is for example, the oldest known monohydrocalcite, 800 ka. (Solotchina et  
56 al. (2009), that was found in deep sediments at Lake Hovsgol, NW Mongolia. Other  
57 lacustrine examples include Lake Kivu in Africa, or Lake Fellmongery and Lake Butler in S.  
58 Australia (Stoffers and Fischbeck, 1974 and Taylor, 1975). Monohydrocalcite is usually  
59 found in association with other carbonates like Mg-calcite, aragonite, Ca-rich dolomite  
60 ( $CaMg(CO_3)_2$ ) or Mg carbonates like hydromagnesite ( $Mg_5(CO_3)_4(OH)_2 \cdot 4H_2O$ ) and  
61 nesquehonite ( $MgCO_3 \cdot 3H_2O$ ) (Fischbeck and Mueller, 1971; Broughton, 1972; Nishiyama et  
62 al., 2013). Monohydrocalcite is also formed as a product of biomineralization by certain  
63 molluscs (Lowenstam, 1981), flatworms (calcareous corpuscles of Platyhelminthes; Señorale-  
64 Pose et al., 2008), vertebrates (otoliths; Carlström, 1963), guinea pigs (bladder stones;  
65 Skinner et al., 1977) or is even found as a decay product in Saguaro cacti (Garvie, 2003,  
66 2006). It has also been described associated with algae (Taylor, 1975) or halo bacilli

67 (Rivadeneira et al., 2004). However, despite its occurrence in a wide variety of systems little  
68 is known about the mechanism and kinetics of its formation pathway.

69 At Earth surface temperatures and pressures, monohydrocalcite is thermodynamically  
70 unstable relative to anhydrous calcite and aragonite. In the presence of a Mg-free aqueous  
71 fluid it will slowly (>2 days at 100 °C, several days/weeks at ambient temperature) transform  
72 to calcite (Stoffers and Fischbeck 1974; Taylor, 1975). Yet, even minor concentrations of  
73 aqueous magnesium will lead to its transformation to aragonite over 25 days at ambient  
74 temperatures (Brooks et al., 1950; Kamiya et al., 1977; Dahl and Buchardt, 2006; Munemoto  
75 and Fukushi, 2008; Fukushi et al., 2011). This effect is believed to be due to the inhibiting  
76 effect of  $Mg^{2+}$  on calcite crystallization (Chen et al., 2004; Mucci and Morse, 1983, Davis et  
77 al., 2000; Bots et al., 2011). The instability of monohydrocalcite with respect to calcite and  
78 aragonite explains the relatively low abundance of this phase within modern environmental  
79 systems and the geological record.

80 The mechanism of monohydrocalcite crystallisation in many systems is unknown, but  
81 some studies indicate that it can form from an amorphous precursor (Kamiya et al., 1977;  
82 Loste et al., 2003; Fukushi et al., 2011; Nishiyama et al., 2013). Such a crystallization  
83 pathway has been shown to be common for many Ca-Mg carbonates (e.g., vaterite, aragonite,  
84 dolomite; Bots et al., 2012; Rodriguez-Blanco et al., 2012; Sand et al., 2012; Rodriguez-  
85 Blanco et al., 2013; Ihli et al., 2012), and other carbonate and phosphate systems (Roncal-  
86 Herrero et al., 2009; Roncal-Herrero et al., 2011; Vallina et al., 2013), but this amorphous to  
87 crystalline transition is not ubiquitous (Van Driessche et al., 2012). In the carbonate system,  
88 these studies have shown that the transformation of the amorphous calcium carbonate (ACC)  
89 precursor to stable crystalline phases (i.e., vaterite, dolomite) occurs via a nucleation  
90 dominated (spherulitic) growth mechanism. Despite its presence in a variety of natural  
91 settings and biominerals, a quantitative evaluation of the kinetics and mechanisms of  
92 monohydrocalcite formation from an amorphous precursor is lacking.

93 Monohydrocalcite has a hexagonal structure with space group  $P3_112$  and an atomic  
94 structure consisting of irregular 8-folded Ca-O polyhedra, with a central  $Ca^{2+}$  ion surrounded  
95 by carbonate groups and water molecules (Effenberger, 1981; Neumann and Epple, 2007;  
96 Swainson, 2008). The presence of the water makes its structure more open and less dense  
97 compared to anhydrous  $CaCO_3$  (e.g., calcite or aragonite). Interestingly, monohydrocalcite is  
98 described as a pure 'calcium' carbonate phase in mineralogical databases, but a number of  
99 studies have shown that Mg is ubiquitous in all synthetic and natural monohydrocalcite  
100 samples ( $Mg/Ca = \sim 0.01-0.53$ ; Hull and Turnbull, 1973; Taylor, 1975; Neumann and Epple,

101 2007; Nebel et al., 2008; Fukushi et al., 2011; Nishiyama et al., 2013). The Mg content of  
102 monohydrocalcite is likely to be linked to the overall fluid chemistry and in particular to the  
103 Mg contents of the fluid in which it forms. In both natural and most synthetic  
104 monohydrocalcites the presence of high concentrations of Mg relative to Ca in the  
105 precipitating solution ( $Mg^{2+}_{aq}/Ca^{2+}_{aq} \geq 0.17 < 65$ ) is a prerequisite for its formation (e.g.,  
106 Munemoto and Fukushi, 2008; Neumann and Epple, 2007; Last et al., 2010; Kimura and  
107 Koga, 2011). Recently, Nishiyama et al (2013) have also shown that the crystallinity and  
108 particle size of synthetic monohydrocalcite decrease with Mg content. This clearly shows that  
109 Mg is a key component in monohydrocalcite, and must be present in the fluids in which it  
110 crystallizes. However, the exact mechanisms by which Mg controls the formation and  
111 stability of monohydrocalcite are unknown.

112 This study aims to determine the mechanism of monohydrocalcite crystallisation in  
113 solution via an ACC precursor. We evaluate the role of Mg during this process and test the  
114 hypothesis that a nucleation-dominated process controls the formation of monohydrocalcite  
115 from the Mg-containing precursor ACC. This has been done through a series of experiments  
116 where the nucleation and crystallization of monohydrocalcite from Mg doped-ACC was  
117 followed in situ and in real time with synchrotron-based X-ray scattering combined with  
118 microscopic and wet chemical characterization of the solids and solutions throughout the  
119 reaction. Our results demonstrate that the formation of monohydrocalcite takes place via a 4-  
120 stage process that starts with the precipitation of an Mg-rich ACC precursor. This precursor  
121 crystallizes in stage two to monohydrocalcite through a nucleation-controlled dissolution-  
122 reprecipitation reaction. In stage three Ostwald-ripening dominates, while in stage four, a  
123 secondary monohydrocalcite crystallization, also through Ostwald ripening, occurs  
124 concomitantly with the loss of some of its structural Mg. We also show that, depending on  
125 the aqueous Mg/Ca ratio and supersaturation, monohydrocalcite can accommodate large  
126 amounts of Mg in its structure (up to ~25% Ca replaced by Mg) and that the aqueous  
127 chemistry of the starting solution controls the crystallite size, unit-cell dimensions and  
128 particle size/shape of the forming monohydrocalcite. Based on these findings we discuss the  
129 implications of this monohydrocalcite crystallization pathway, on its occurrence in various  
130 natural settings and as a biomineral.

131

132

## 133 2. EXPERIMENTAL

134



135 Experiments were carried out at 21 °C by rapid addition (under constant and vigorous  
136 stirring) of a CaCl<sub>2</sub>/MgCl<sub>2</sub> solution (700 mM CaCl<sub>2</sub>; 300 mM MgCl<sub>2</sub>; Ca/Mg ratio of 7/3) to  
137 a 1000mM Na<sub>2</sub>CO<sub>3</sub> solution to achieve a (Ca+Mg)/CO<sub>3</sub> ratio of 1/1. In all cases immediately  
138 upon mixing a white gel precipitated. The crystallization of this white gel was followed on-  
139 line using in situ and time resolved small and wide angle X-ray Scattering (SAXS/WAXS) on  
140 beamline I22 (Diamond Light Source Ltd., UK). Solutions were mixed using a stopped-flow  
141 apparatus (Bio-Logic, Paris, France) and the resulting white suspensions were continuously  
142 circulated with a peristaltic pump through a capillary cell in line with the synchrotron beam.  
143 Simultaneous SAXS (RAPID detector; Marchal et al., 2009) and WAXS (HOTWAXS  
144 detector; Bateman et al., 2007) patterns were collected over 12 hours at 1 min/frame. The  
145 time resolved scattering patterns were detector-response corrected, and background  
146 subtracted using a scattering pattern from the starting Na<sub>2</sub>CO<sub>3</sub> solution. Individual WAXS  
147 patterns were fit using XFIT (Cheary and Coelho, 1992) and the areas under the Bragg peaks  
148 normalized to values from 0 to 1 to express the degree of reactions ( $\alpha$ ) as a function of time  
149 (Rodriguez-Blanco et al., 2011). Crystallite size was calculated from the Bragg peak full-  
150 width half-maximum (FWHM) using the Scherrer equation (Scherrer, 1918) and the unit cell  
151 parameters were determined with TOPAS (Coelho, 2006), using the Swainson (2008) model  
152 for the monohydrocalcite structure.

153 The variations in the scattering peak position in the SAXS patterns were used to  
154 derive the changes in the particle sizes of the solids throughout the experiments via the  
155 expression:

$$157 \quad d = 2\pi/q \quad [1]$$

158  
159 where  $d$  (nm) is particle diameter and  $q$  (nm<sup>-1</sup>) is the scattering vector (de Moor et al, 1999;  
160 de Moor et al, 1999b). These particle size results were also cross-validated by whole SAXS  
161 pattern fitting using GNOM (Svergun, 1992).

162 Experiments equivalent to the online SAXS/WAXS experiments (i.e., 1000mM  
163 Na<sub>2</sub>CO<sub>3</sub> solution mixed with a 700 mM CaCl<sub>2</sub> and 300 mM MgCl<sub>2</sub> solution) were performed  
164 to follow the reaction process via time-dependent solid characterization and solution  
165 analyses. At specific time steps aliquots of the reacting suspension were removed and  
166 immediately vacuum filtered (0.2  $\mu$ m polycarbonate membranes). The resulting solids were  
167 washed with water and isopropanol following Rodriguez-Blanco et al (2008). An aliquot of  
168 each solid sample was dissolved in 0.1 M HCl and analysed together with the corresponding

169 solution samples for aqueous Ca and Mg concentrations ( $[Ca^{2+}_{aq}]$  and  $[Mg^{2+}_{aq}]$ ) using ion  
 170 chromatography (Dionex LC 220) following Bots et al (2011). Throughout all experiments  
 171 the pH was recorded at a 10 second time resolution using an Orion pH meter and WinWedge  
 172 v3.4.1 software (TAL technologies) with a precision of 0.001. Finally, the total carbonate  
 173 concentration in solution was determine by assuming that all Ca and Mg precipitated formed  
 174 Ca/Mg carbonate, therefore the moles of Ca + Mg removed form solution must equal the total  
 175 moles of carbonate removed. The total carbonate was then calculated from the difference  
 176 between the initial concentration and the amount removed at each time point. From these  
 177 solution data the saturation indexes (SI) with respect to different Ca and Mg-bearing  
 178 carbonates were calculated using PHREEQC (Parkhurst, 1995) and are presented in Table 1.  
 179 This was done using the Pitzer activity coefficient models (Pitzer, 1979) and two solubility  
 180 products,  $K_{sp(MHC)}$ , for monohydrocalcite ( $10^{-7.60}$ ; Hull and Turnbull, 1973) and ( $10^{-7.05}$ ; Krajl  
 181 and Brečević, 1995). The saturation index of monohydrocalcite is defined as:

$$182 \quad SI_{MHC} = \log \frac{a_{Ca^{2+}} \times a_{CO_3^{2-}}}{K_{sp(MHC)}} \quad [2]$$

183  
 184 To determine the effect of initial supersaturation (i.e. saturation index of the aqueous  
 185 solution for monohydrocalcite immediately after mixing the stock solutions) on the  
 186 crystallization reaction, experiments were also performed following the same procedure but  
 187 starting at different initial supersaturations with respect to monohydrocalcite, yet keeping a  
 188 1/1 (Ca+Mg)/CO<sub>3</sub> ratio (Table 2) in all cases. Using the  $[Ca^{2+}_{aq}]$  and  $[Mg^{2+}_{aq}]$  concentrations  
 189 obtained from the analyses of the dissolved solids, the molar fraction of MgCO<sub>3</sub> in each solid  
 190 sample,  $X_{MgCO_3}$ , was calculated following:

$$191 \quad X_{MgCO_3} = \frac{M_{MgCO_3}}{M_{MgCO_3} + M_{CaCO_3}} \quad [3]$$

192  
 193 where  $M_{MgCO_3}$  and  $M_{CaCO_3}$  correspond to the molar concentration of MgCO<sub>3</sub> and CaCO<sub>3</sub>,  
 194 respectively.

195  
 196 The mineralogy of the solids from off-line experiments was characterized by powder  
 197 X-ray diffraction (PXRD) using a Bruker D8 X-ray Diffractometer (CuK $\alpha$ 1, 5-90° two theta,

198 0.001°/step; 0.1 or 1 sec/step). The PXRD patterns were also used to derive the crystallite  
199 size using the Scherrer equation (Scherrer, 1918). All samples were imaged by high-  
200 resolution transmission electron microscopy (HR-TEM; Philips CM200 field emission gun  
201 TEM equipped with a Gatan Imaging Filter, GIF-200 and a Gatan camera for selected area  
202 electron diffraction, SAED, pattern collection) or scanning electron microscopy (SEM; LEO  
203 1530 Gemini). Finally, the water content of the samples was determined using  
204 thermogravimetric analyses (TGA, Mettler Toledo, 25-800 °C in air, 10°C/min). The total  
205 water content for the initial white precipitate was calculated from the difference in weight  
206 between 25 and 550°C in accordance with the method of Radha et al. (2012) for determining  
207 the water content of ACC. For crystalline samples the difference in weight between 100 and  
208 550°C was used, in accordance with the method of Hull and Turnbull (1973) for determining  
209 the water content of monohydrocalcite.

210

211

### 212 3. RESULTS AND DISCUSSION

213

214 Our data revealed that over the 12 hours of our experiments monohydrocalcite formed  
215 through a multi-stage crystallisation pathway. The first stage of the reaction was  
216 characterized by the formation of a Mg-bearing amorphous calcium carbonate phase. In stage  
217 two, this amorphous phase crystallised to nanoparticulate monohydrocalcite. The rate of  
218 crystallization slowed considerably during stage three, coincident with only minor changes  
219 in the solution composition (e.g., pH). Finally, in the fourth stage of the reaction secondary  
220 growth of the nanoparticulate monohydrocalcite was observed. This was accompanied by the  
221 simultaneous decrease in  $[Mg^{2+}_{aq}]$  and the formation of minor amounts of hydromagnesite  
222 ( $Mg_5(CO_3)_4(OH)_2 \cdot 4H_2O$ ). Each of these four stages is described in detail below:

223 In **Stage I** of the reaction a white gel precipitated immediately upon solution mixing.  
224 The PXRD patterns of this gel (Fig. 1a, lower pattern) showed no Bragg peaks and only three  
225 humps located at  $2\theta$  ( $\lambda=1.54 \text{ \AA}$ ) 17, 31 and 45°, indicating the solid is poorly-ordered. These  
226 three broad humps (asterisks in Fig. 1a), are located at the same positions as those observed  
227 for Mg-free amorphous calcium carbonate (ACC; Rodriguez-Blanco et al., 2008). Based on  
228 the chemical analyses of this amorphous phase (Table 1, 0.83 hour sample) and the TGA  
229 results, the formula for this compound was determined to be  $Ca_{0.74}Mg_{0.26}CO_3 \cdot 1.18H_2O$ . In  
230 line with previous studies where the incorporation of Mg into ACC has been quantified

231 (Nebel and Epple, 2008; Wang et al., 2009; Rodriguez-Blanco et al., 2012), we will term this  
232 poorly-ordered phase, Mg-ACC, following the notation of Günther et al (2005). Throughout  
233 this first stage, Mg-ACC was the only solid phase present. In both the on-line and off-line  
234 experiments and regardless of conditions, the Mg-ACC started to crystallize after  
235 approximately 50 minutes to 1 hour, initiating **Stage II**. During this stage the background  
236 intensity in the PXRD patterns decreased with time, and Bragg peaks for monohydrocalcite  
237 formed, grew and sharpened (Fig. 1a). The on-line time resolved WAXS data (Fig. 1b) shows  
238 the simultaneous growth of the monohydrocalcite (110), (111), (300) and (301) peaks, after ~  
239 1 hour. The whole reaction is clearly illustrated when the change in area for a specific Bragg  
240 peak, expressed as the degree of reactions ( $\alpha$ ), is plotted as a function of time (Fig. 2). Stage  
241 II of the reaction lasted for about 1 hour and was characterized by the rapid increase in  
242 monohydrocalcite peak area. After the fast growth in stage II, a much slower growth phase,  
243 between ~ 2 and 8 hours followed (**stage III**). During this stage, only minor changes in Bragg  
244 peak intensities / areas were observed (Fig. 1b and 2). The final stage of the reaction -**stage**  
245 **IV**- started after about 8.3 hours. During this stage a secondary growth in the  
246 monohydrocalcite Bragg peak areas was observed. This growth phase lasted about 2-3 hours  
247 and the peak areas reaching a stable maximum after ~ 11 hours. This same evolution was  
248 observed for all other monohydrocalcite Bragg peaks. An PXRD examination of off-line  
249 experimental solids sampled during stage IV revealed the presence of a second, but minor  
250 phase of hydromagnesite (ICDD PDF 01-070-1177; Fig. S1). Quantitative X-ray refinement  
251 with TOPAS (Coelho, 2006) determined that its abundance was ~4.5%. The on-line time  
252 resolved WAXS patterns (Fig. 3) also revealed small hydromagnesite (110) and (011) Bragg  
253 peaks that began to grow at the onset of stage IV, in parallel with the second stage of  
254 monohydrocalcite crystallization.

255 Analysis of the SAXS data (Fig. 4) indicates that the Mg-ACC that formed in stage I  
256 consisted of particles with a diameter of between 40 and 60 nm in size, a size that remained  
257 constant throughout this first stage. This is consistent with particle sizes derived from the  
258 TEM imaging of the Mg-ACC particles, which indicated an average diameter of ~ 40 nm  
259 ( $n=100$ ; Fig. 5a; Table 1). However, the Mg-ACC particle diameters varied between 15 and  
260 210 nm revealing a high degree of polydispersity. The corresponding SAED patterns revealed  
261 only diffuse rings for the Mg-ACC, confirming its poorly-ordered nature (Fig. 5d). During  
262 stage II, a distinct scattering peak appeared in the SAXS patterns (arrows in Fig. 4a)  
263 indicating that the crystals formed in the experiment were relatively monodispersed. The  
264 position of the scattering peak in the time resolved SAXS patterns moved towards lower  $q$

265 with time indicating particle growth (Fig. 4a). From the position of this peak and using  
266 equation [1] the particle size of the monohydrocalcite crystals was determined (Fig. 4b). At  
267 the beginning of stage II, when the first Bragg peaks for monohydrocalcite appeared in the  
268 WAXS data (Fig. 1b and Fig. 2), the corresponding SAXS data revealed an average diameter  
269 of ~9 nm for the newly forming monohydrocalcite crystals. During the rest of stages II and  
270 III these monohydrocalcite nanocrystals grew only slightly and very slowly reaching a  
271 maximum diameter of ~15 nm after 8 hours. This same trend in average particles sizes was  
272 confirmed (open symbols in Fig. 4b) through selected full pattern evaluation with GNOM  
273 (Svergun, 1992). TEM microphotographs of the monohydrocalcite that formed in stage II and  
274 persisted throughout stage III (Fig. 5b) revealed that the monohydrocalcite consisted of  
275 nanocrystals, which exhibited sub-spherical to slightly elongated morphologies. Their  
276 average dimensions were  $28\pm 7$  (L) x  $20\pm 5$  (W) nm, with the minimum size being around  
277  $14\times 10$  nm (n=100). The corresponding SAED pattern showed diffraction rings with only  
278 poorly developed spots (Fig. 5e), evidencing the presence of very small but crystalline  
279 nanocrystals. At the onset of stage IV, and simultaneously with the secondary growth in  
280 Bragg peak areas for monohydrocalcite, the SAXS data also revealed a steady increase in the  
281 monohydrocalcite nanocrystal diameters, which reached a maximum of ~45 nm at the end of  
282 the reaction. Photomicrographs of a sample obtained after 10 hr (stage IV; Fig. 5c) showed  
283 euhedral or subhedral single monohydrocalcite crystals with sizes of  $77\pm 20$  (L) x  $47\pm 10$  (W)  
284 nm. Again large variations in dimensions were observed, with the largest and smallest  
285 crystals being  $155\times 80$  and  $40\times 30$  nm (n=100). SAED pattern from these crystals exhibited  
286 clearly defined spots within the diffraction rings (Fig. 5f). HR-TEM images of the sample  
287 taken from the end of stage IV (Fig. 5c, white arrow) revealed the presence of  
288 hydromagnesite as a minor phase consisting of ~ $100\times 5$  nm platy crystals, confirming the  
289 WAXS and PXRD observations (Fig. 3 and Fig. S1).

290 Evaluating the corresponding WAXS patterns for the Scherrer crystallite size (Fig. 2)  
291 showed a reasonable agreement with the particle size determined from the SAXS and TEM  
292 data, and also confirmed that the forming monohydrocalcite crystals were anisotropic. For  
293 example, based on the Bragg peak (110) the data reveals that the monohydrocalcite crystals  
294 formed during stage II had a crystallite size of ~15 nm, increasing to ~20 nm during stage  
295 III, which is in agreement with the particle sizes derived from the SAXS and TEM data. Once  
296 stage IV of the crystallization was initiated, the monohydrocalcite nanocrystallite size  
297 steadily increased to > 80 nm, which is almost double the average particle size from SAXS,

298 but closer to that derived from the TEM data. However, although this general trend was the  
299 same in all other Bragg peaks, the crystallite sizes in all stages were dramatically different  
300 between the different Bragg peaks. For example, the final crystallite size from the (141) peak  
301 at the end of stage 3 was 16 nm, half that of the (110) peak, and at the end of the reaction it  
302 only reached a value of 28 nm (Fig. 2), almost three times smaller than the final crystallite  
303 size from the (110) peak, indicating a clear anisotropic growth of the monohydrocalcite  
304 nanocrystals. This is consistent with the elongated particle morphology observed by TEM.

305 Analysis of the WAXS data also revealed changes in unit cell dimensions during the  
306 formation of monohydrocalcite (Fig. 6). Once formed, the unit cell volume ( $721.6 \text{ \AA}^3$ )  
307 remained virtually constant during stages II-III with a slight decrease during stage IV ( $\Delta V \approx -$   
308  $0.2 \text{ \AA}^3$ ; Fig. 6a). However, more substantive changes were observed in the a and c axis  
309 dimensions (Fig. 6b and c, respectively). Their evolution mirrored each other, with the a-axis  
310 decreasing and the c-axis increasing in length ( $\Delta c = -\Delta a \approx 0.02 \text{ \AA}$ ). These changes were small  
311 during stages II and III, but became more significant during the secondary monohydrocalcite  
312 crystallization (stage IV).

313 The changes in solution and solid phase chemistry during the formation and  
314 crystallisation of monohydrocalcite showed a clear link between the evolution of pH,  
315  $[\text{Ca}^{2+}_{\text{aq}}]$ ,  $[\text{Mg}^{2+}_{\text{aq}}]$  and molar fraction of Mg in the solids (Fig. 7) over the four stage of the  
316 reaction. The reaction started with a pH of 11.60 (the pH of the initial  $\text{Na}_2\text{CO}_3$  solution),  
317 which upon mixing with the Ca/Mg solution rapidly dropped to 9.17. During stage I the pH  
318 increased slightly reaching a value of 9.24, while the formation of monohydrocalcite in stage  
319 II did not change the pH. In stage III only a small decrease in pH to a value of 9.16 was  
320 evident. However, at the end of stage III and during stage IV the pH dropped dramatically to  
321 8.65. The  $[\text{Ca}^{2+}_{\text{aq}}]$  was 1.25 mM during stage I and decreased to  $\approx 0.6$  mM by stage III. It  
322 then decreased further reaching a concentration  $< 0.1$  mM at the end of stage IV (Fig. 7).  
323  $[\text{Mg}^{2+}_{\text{aq}}]$  was 28 mM during stages I and II, but increased by  $\sim 20\%$  to 35 mM during stage  
324 III. During stage IV it decreased by 40% to 13 mM. These changes in  $[\text{Mg}^{2+}_{\text{aq}}]$  were mirrored  
325 throughout the whole reaction by the change in  $\chi_{\text{MgCO}_3}$  in the solids (Fig. 7). Furthermore, the  
326 timing of these changes in aqueous composition were all coupled with the increase in  
327 monohydrocalcite particle size (Fig. 4 and 5) and formation of hydromagnesite (Fig. 3).

328 From the  $\chi_{\text{MgCO}_3}$  values and the TGA results, the formula of the stage II  
329 monohydrocalcite was derived to be  $\text{Ca}_{0.74}\text{Mg}_{0.26}\text{CO}_3 \cdot 0.99\text{H}_2\text{O}$ . However, although the total  
330  $\chi_{\text{MgCO}_3}$  increased during stage IV (Fig. 7), this does not represent the Mg in monohydrocalcite

331 but primarily the contribution from the minor (~ 4.5%) hydromagnesite ( $\text{Mg}_5(\text{CO}_3)_4(\text{OH})_2$   
332  $4\text{H}_2\text{O}$ ). Using the quantitative XRD results and the composition of the final solid obtained  
333 from chemical analysis ( $\chi_{\text{MgCO}_3} = 0.26$ , Table 1) we calculated that the molar fraction of Mg  
334 in monohydrocalcite at the end of stage IV was only ~ 0.065 moles, which is a 75% decrease  
335 compared to the Mg content of the stage II monohydrocalcite (0.26 moles). The remaining  
336 Mg was transferred into the newly formed hydromagnesite.

337 One last line of evidence that supports the observations above is the evolution of the  
338 saturation states during the reaction. PHREEQC modelling of the solution during the reaction  
339 showed that immediately after the precipitation of Mg-ACC upon mixing, the aqueous  
340 solution was supersaturated with respect to both monohydrocalcite and hydromagnesite (Fig.  
341 8) and that the SI for both phases varied little during stages I to III ( $\text{SI}_{\text{MHC}} \approx 0.5$ ;  $\text{SI}_{\text{HMgS}} \approx 6$ ).  
342 The biggest change in both SI occurred at the onset of stage IV, where  $\text{SI}_{\text{MHC}}$  became  
343 undersaturated ( $\approx -0.85$ ) while  $\text{SI}_{\text{HMgS}}$  dropped to  $\approx 0.2$ . This change in SI was concomitant  
344 with the removal of Mg from solution and the associated drop in pH, due to the formation of  
345 hydromagnesite. There are however, two factors that need to be taken into account when  
346  $\text{SI}_{\text{MHC}}$  is calculated. Firstly, two solubility products for monohydrocalcite are available in the  
347 literature:  $10^{-7.60}$  (Hull and Turnbull, 1973), and  $10^{-7.05}$  (Krajl and Brečević, 1995) and this  
348 results in a ~0.55 difference in the calculated  $\text{SI}_{\text{MHC}}$  (Fig. 8). Secondly, the total ionic strength  
349 in our experiments was high ( $\leq 1$  M) and thus even an error of 1% in the ionic strength of the  
350 starting solution could also result in variations up to ~0.5 units in the calculated  $\text{SI}_{\text{MHC}}$ .

351

352

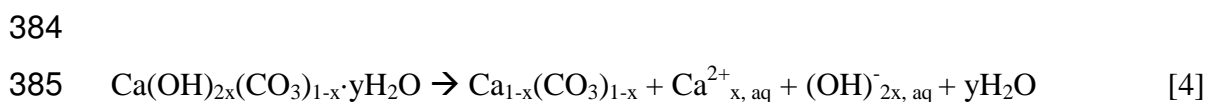
### 353 **3.1. Stability of the precursor Mg-ACC and the mechanism of monohydrocalcite** 354 **nucleation and growth**

355

356 The formation of monohydrocalcite proceeded, in all our experiments, through a  
357 poorly-ordered Mg-ACC precursor, which precipitated from solutions with an initial  
358  $[\text{Ca}^{2+}_{\text{aq}}]:[\text{Mg}^{2+}_{\text{aq}}]$  of 7:3. The resulting phase was considerably more stable (crystallization  
359 started at ~50 minutes) compared to pure ACC (no Mg; crystallization started <2 minutes;  
360 Ogino et al., 1987; Rodriguez-Blanco et al., 2011; Rodriguez-Blanco et al., 2012, Bots et al.,  
361 2012), or Mg-ACC formed in a solution with  $[\text{Ca}^{2+}_{\text{aq}}]:[\text{Mg}^{2+}_{\text{aq}}]$  of 9 to 1 (crystallized to  
362 calcite within ~ 14 minutes; Rodriguez-Blanco et al., 2012). However, the Mg-ACC in the  
363 current study was less stable than Mg-ACC produced from solutions where 50% of the

364 calcium was replaced with magnesium (i.e.,  $[\text{Ca}^{2+}_{\text{aq}}]:[\text{Mg}^{2+}_{\text{aq}}] = 1:1$ ), which did not  
365 crystallize at ambient temperatures even after longer reaction times ( $>1$  day; Rodriguez-  
366 Blanco et al., 2013). The increasing stability of ACC with increasing Mg contents is not  
367 unexpected, and confirms both abiotic (Rodriguez-Blanco et al., 2011; Rodriguez-Blanco et  
368 al., 2013; Ajikumar et al., 2005) and biotic (Politi et al., 2010; Loste et al., 2003; Raz et al.,  
369 2003) data that demonstrate the importance of Mg in stabilizing amorphous precursors and  
370 delaying crystallization. The most likely cause of these effects is the high dehydration energy  
371 of the Mg-aquo ion compared to the Ca ion (di Tommaso and de Leeuw, 2010). Hydrated and  
372 poorly-ordered ACC is less thermodynamically stable than the dehydrated and more-ordered  
373 ACC. This dehydration and local ordering in the amorphous precursor precedes its  
374 crystallization (Radha et al., 2010; Bots et al., 2012). Hydrated Mg located within the  
375 nanoporous structure of ACC (Goodwin et al., 2010) would retard its dehydration and  
376 breakdown, slowing its transformation to crystalline phases.

377 In stage I and prior to the formation of monohydrocalcite, the pH increased by a small  
378 amount (0.07 units) (Fig. 7), which likely corresponds to the release of  $\text{OH}^-$  due to the onset  
379 of Mg-ACC dissolution. Kojima et al (1993) showed that ACC incorporates minor amounts  
380 of  $\text{OH}^-$  ions into its structure when it forms at a basic starting pH. The Mg-ACC in our on-  
381 line experiment started forming at a pH of 11.5 (the pH of the carbonate starting solution;  
382 Fig. 7) and thus minor  $\text{OH}^-$  in our Mg-ACC is not unexpected. Its dissolution would release  
383  $\text{OH}^-$  ions, thus explaining the slight pH increase following:



386  
387 The first monohydrocalcite crystals formed during stage II were significantly smaller  
388 than the initial Mg-ACC nanoparticles (Fig. 4b), yet once monohydrocalcite started forming,  
389 its diameter remained virtually constant throughout stage II and III. Analysis of the WAXS  
390 data shows that the amount of monohydrocalcite crystallising increased at a constant rate  
391 throughout stage II of the reaction (Fig. 2). However, this increase was not due to particle  
392 growth (Fig. 4b) and we suggest that the formation of monohydrocalcite during stage II  
393 proceeds via the nucleation of new crystalline solids, i.e., via a nucleation-controlled reaction.  
394 The constant rate of monohydrocalcite crystallization with time (i.e., linear increase in the  
395 amount of monohydrocalcite) during most of stage II indicates a zeroth order reaction, and a  
396 constant rate of nucleation. Furthermore, the pH remained virtually constant during stage II,



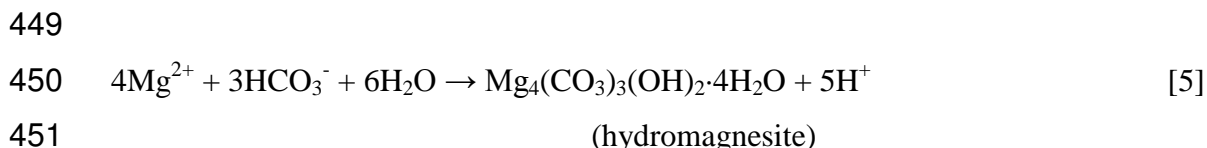
397 indicating a constant dissolution of Mg-ACC coupled with the constant nucleation of  
398 monohydrocalcite. Taking all this into account, we propose that the transformation from Mg-  
399 ACC to monohydrocalcite occurs via a simultaneous dissolution of Mg-ACC and nucleation  
400 of monohydrocalcite at a constant rate. Nucleation controlled growth has been shown for  
401 several other crystalline carbonate phases formed from amorphous Ca-Mg carbonates (e.g.,  
402 vaterite and proto-dolomite; Bots et al., 2012; Rodriguez-Blanco et al., 2013). These studies  
403 suggested that the large difference in solubility between the amorphous precursor and the  
404 crystalline phases as well as the high supersaturations with respect to all crystalline phases,  
405 promotes nucleation. This nucleation process will continue until all of the amorphous  
406 precursor has been consumed, leading to a constant nucleation rate.

407         Stage IV of the reaction is key to understanding the effect of Mg on the formation of  
408 monohydrocalcite. The SAXS data (Fig. 4b) shows a dramatic increase in particle size.  
409 During stage IV, this increase can be fitted to a straight line when plotted as a function of  $t^{1/2}$ ,  
410 indicating a surface-controlled Ostwald-ripening mechanism (Wagner, 1961; Tobler et al.,  
411 2009). TEM images and both WAXS and PXRD data confirm that larger and better  
412 developed monohydrocalcite crystals formed during this stage of the reaction, supporting the  
413 Ostwald-ripening mechanism. The photomicrographs of monohydrocalcites from stages III  
414 and IV (Fig. 5 b and c) show an unambiguous increase in the nanocrystal sizes and a  
415 corresponding change in the SAED patterns from rings with faint spots to discrete spots,  
416 indicating an increase in particle size.

417         Ostwald ripening usually occurs during the later stages of crystallisation reactions and  
418 involves particle growth without the formation of new material (Wagner, 1961). All  
419 nanoparticles formed in stages III are undoubtedly crystalline, but some of them are too small  
420 (<10 nm) to produce a significant amount of diffraction. Therefore scattering from these  
421 crystal is observed in the background of the PXRD/WAXS patterns during stages II and III  
422 (e.g. Fig. 1 after 1 hour of reaction). This behaviour is similar to that observed for ultra-small  
423 iron oxides crystallites which are too small to diffract singly (Machala et al., 2007; Ahmed et  
424 al., 2010). The intensity of the background in the XRD/WAXS patterns decreased during  
425 stages III and IV, while the Bragg peaks increased in intensity and became sharper. These  
426 changes in background and Bragg peak intensity are not due to the formation of more  
427 monohydrocalcite, but reflect the growth of the monohydrocalcite nanoparticles to a size  
428 where they diffract strongly, due to the Ostwald ripening process. The linear fits to the  
429 particle growth data with the surface-controlled growth model Fig. 4b show different rates

430 (slopes) for the Ostwald-ripening process during stages III (slow;  $1.8 \pm 0.1 \text{ nm/h}^{1/2}$ ) and IV  
431 (fast;  $66.3 \pm 4.3 \text{ nm/h}^{1/2}$ ).

432 The rapid change in monohydrocalcite ripening rate is coincident with the formation  
433 of hydromagnesite; its formation lead to the fast removal of Mg from solution and triggered  
434 the rapid increase in monohydrocalcite growth rate. We suggest that structural Mg from  
435 monohydrocalcite during the ripening process was released into the solution leading to the  
436 observed increase in  $[\text{Mg}^{2+}_{\text{aq}}]$  during stage III (Fig. 7). This ultimately drove the  
437 monohydrocalcite composition toward a pure  $\text{CaCO}_3 \cdot \text{H}_2\text{O}$  end-member. The high  
438 concentration of Mg in solution during stage II suppressed the ripening rate, and produced a  
439 feedback effect, which decreased the rate of ripening even more (decrease in the slope of the  
440 degree of reaction plot during stage III; Fig. 2). The presence of Mg in solution is known to  
441 suppress the dissolution and growth of calcium carbonates (e.g., calcite; Chen et al., 2004;  
442 Mucci and Morse, 1983; Davis et al., 2000). Thus, is not unexpected that the high  
443 concentration of  $\{\text{Mg}^{2+}_{\text{aq}}\}$  present during stage III significantly reduced the dissolution and  
444 reprecipitation process during the Ostwald ripening. Although the aqueous solution was  
445 supersaturated with respect to hydromagnesite throughout the reaction, this progressive  
446 increase in  $[\text{Mg}^{2+}_{\text{aq}}]$  eventually triggered the nucleation of hydromagnesite at the end of stage  
447 III (Fig. 3). Its formation removed  $[\text{Mg}^{2+}_{\text{aq}}]$  from solution (Fig. 7) and induced a dramatic  
448 drop in pH following:



453 The main consequence of this Mg removal from the aqueous solution was an acceleration of  
454 the monohydrocalcite ripening reaction, which was translated into a rapid growth in particle  
455 size and the formation of highly crystalline low-Mg monohydrocalcite.

456 PHREEQC calculations (Table 1) also revealed that the aqueous solution was in  
457 equilibrium or slightly supersaturated with respect to nesquehonite ( $\text{Mg}(\text{HCO}_3)(\text{OH}) \cdot 2(\text{H}_2\text{O})$ )  
458 during stages II and II, but became undersaturated in this mineral at stage IV. Nesquehonite  
459 was never detected using conventional XRD, synchrotron-based WAXS or HR-TEM  
460 imaging. However, Nishiyama et al (2013) suggested that the formation of monohydrocalcite  
461 would require the paragenesis of a hydrous Mg-bearing carbonate e.g., nesquehonite, that  
462 would transform to hydromagnesite after longer reaction times. Despite the lack of  
463 experimental evidences, we can not completely discard the possibility that small amounts of

464 nanocrystalline nesquehonite were present during stages II and III, and that these eventually  
465 transformed to hydromagnesite during stage IV.

466 The inhibiting effect of Mg has been described previously for other Ca-Mg carbonates  
467 (Bischoff, 1968; Berner, 1975; Reddy and Wang, 1980; Mucci and Morse, 1983; Davis et al.,  
468 2000). For example, Zhang and Dawe (2000) have suggested that the calcite growth rate is  
469 inversely proportional to the Mg concentration in solution. This effect has been attributed to  
470 the stronger hydration shell of Mg in comparison to Ca (di Tommaso and de Leeuw, 2010;  
471 Moomaw and Maguire, 2008). The higher energy, which Mg needs to dehydrate before  
472 incorporating into a carbonate structure controls the kinetics of crystal growth (Mucci and  
473 Morse, 1983; Nancollas and Purdie, 1964; De Boer, 1977). Therefore, monohydrocalcite  
474 growth would be favoured at lower Mg concentrations.

475

### 476 **3.2. Monohydrocalcite chemical and structural variability**

477

478 The molar fraction of Mg in monohydrocalcite ( $\chi_{\text{MgCO}_3}$ ) was  $\sim 0.25$  during stages II  
479 and III (i.e., before the formation of hydromagnesite). This is close to the maximum values in  
480 the literature for natural or synthetic monohydrocalcites ( $\chi_{\text{MgCO}_3} = \sim 0.01\text{-}0.34$ ; Brooks et al,  
481 1950; Sapozhnikov and Tsvetkov, 1959; Marschner, 1969; Hull and Turnbull, 1973; Taylor,  
482 1975; Skinner, 1977; Neumann and Epple, 2007; Nebel et al., 2008; Munemoto and Fukushi,  
483 2008; Nishiyama et al., 2013). Many of these studies do not include information about the  
484 exact conditions at which monohydrocalcite formed (temperature, solution composition,  
485 synthesis method, etc.), therefore the factors controlling  $\chi_{\text{MgCO}_3}$  in monohydrocalcite are  
486 difficult to evaluate. However, we suggest that the high level of Mg in the monohydrocalcite  
487 crystallised from Mg-ACC in this study (Fig. 7) is likely due to the high supersaturation of  
488 the initial solution with respect to all carbonate phases.

489 To test the effect of supersaturation on the incorporation of Mg into the resulting  
490 monohydrocalcite, a set of batch experiments were performed at different initial  
491 supersaturations (Table 2). Regardless of supersaturation, immediately after mixing the initial  
492 solutions an amorphous precursor formed. Experiments with initial  $SI_{\text{MHC}} > 2.43$  resulted in  
493 the crystallization of monohydrocalcite, while only Mg-calcite was obtained at lower  
494 supersaturations (Table 2 and Fig. S2). No other Ca/Mg-bearing phases were observed.  
495 PXRD analyses of the monohydrocalcite revealed that its crystallite size was inversely  
496 proportional to the starting supersaturation (Table 2), varying between 35 nm at  $SI_{\text{MHC}} = 2.43$   
497 to 16 nm at  $SI_{\text{MHC}} = 3.89$ . Furthermore, the  $\chi_{\text{MgCO}_3}$  of the monohydrocalcite increased with

498 initial supersaturation, from 0.017 ( $SI_{MHC} = 2.43$ ) to 0.164 ( $SI_{MHC} = 3.89$ , Fig. 9a). This data  
499 is consistent with previously published values for  $\chi_{MgCO_3}$  of synthetic monohydrocalcite  
500 produced using different initial Mg content in solution and different supersaturations (Fig. 9a;  
501 Neumann and Epple (2007); Munemoto and Fukushi (2008), however the original  $\chi_{MgCO_3}$  in  
502 the later reference may be higher than reported due to sample treatment). Recently,  
503 Nishiyama et al. (2013) synthesized a series of monohydrocalcites using highly variable  
504 starting  $[Mg^{2+}_{aq}]/[Ca^{2+}_{aq}]/[CO_{3,aq}]$  ratios and most often an excess of carbonate, which lead to  
505 a much higher crystallization pH (9.8-11.4) compared to ours ( $< 9.25$ ). Their data shows that  
506 the  $\chi_{MgCO_3}$  in monohydrocalcite is proportional to their starting  $[Mg^{2+}_{aq}]/[Ca^{2+}_{aq}]$  ratio, but  
507 there is no clear link between supersaturation and Mg contents ( $\chi_{MgCO_3}$ ) (Fig. 9a). In addition,  
508 Nishiyama et al. (2013) used a different solution mixing approach (addition of the  $Na_2CO_3$   
509 solution to the  $CaCl_2/MgCl_2$  solution) compared to ours (addition of the  $CaCl_2/MgCl_2$   
510 solution to the  $Na_2CO_3$  solution). In an earlier study (Rodriguez-Blanco et al., 2012) we  
511 showed that, in the pure carbonate system, different mixing approaches lead to different  
512 initial synthesis pH, which in turn dramatically affected the crystallization pathway. Taken as  
513 a whole, both the initial  $[Mg^{2+}_{aq}]/[Ca^{2+}_{aq}]/[CO_{3,aq}]$  ratios (Nishiyama et al., 2013) and initial  
514 supersaturation (this study) are key factors in controlling the Mg content of monohydrocalcite  
515 (Fig. 9a). This is also consistent with Loste et al. (2003) who showed that in other carbonate  
516 systems, precipitation at high supersaturation levels is so rapid that it results in less  
517 distinction between Ca and Mg ions. Therefore we infer that the Mg content of the solid is  
518 proportional to the initial supersaturation level and the Mg content in the initial amorphous  
519 precursor phase.

520 TGA analysis of the monohydrocalcite samples showed that the  $H_2O$  content (0.93 –  
521 1.03 per formula unit) was similar in all solids and proportional to initial supersaturation.  
522 However, the release of structural water during heating was different depending on the molar  
523 fraction of Mg ( $\chi_{MgCO_3}$ ) (Fig. 9b). In samples with  $\chi_{MgCO_3} < 0.06$  all water was released at low  
524 temperature (150-200°C), while in the Mg-rich monohydrocalcite samples ( $\chi_{MgCO_3} > 0.06$ ) the  
525 retention of water was greater, and it was progressively released during heating to  $T > 450$  °C.  
526 The final weight loss starting at 550 °C was due to carbonate decomposition. The retention of  
527 water to higher temperatures in samples with  $\chi_{MgCO_3} > 0.06$  is related to the greater  
528 dehydration energy (~20% higher) of  $Mg^{2+}$  in comparison to  $Ca^{2+}$  (Nancollas and Purdie,  
529 1964; Lippmann, 1973; De Boer, 1977).  $Ca^{2+}$  and  $Mg^{2+}$  ion dehydration temperature ranges  
530 have been defined from 100 - 300 °C, and from 300 - 550 °C respectively, as the release of  
531 water associated with  $Mg^{2+}$  in the monohydrocalcite structure requires a higher temperature

532 (300-550 °C) compared to water associated with  $\text{Ca}^{2+}$ . A detailed quantitative analysis of the  
533 TGA data (Fig. 9b, inset) shows that the amount of water released at higher temperatures (>  
534 300 °C) is directly proportional to the molar fraction of Mg in the solid.

535 Photomicrographs of high or low Mg-containing monohydrocalcite, reveals two  
536 distinct morphologies and sizes. Monohydrocalcite  $\chi_{\text{MgCO}_3} > 0.06$  consists of individual,  
537 nanometer-sized crystals that are <35 nm in size (Fig. 9c). In contrast, monohydrocalcite with  
538  $\chi_{\text{MgCO}_3} < 0.06$  consists also of nanoparticulate crystallites (<35nm), but these are aggregated to  
539 form elongated particles >5  $\mu\text{m}$  in length. The morphology of these low Mg-  
540 monohydrocalcites (Fig. 9d) is identical to “type 2” spherulites described by Granasy et al.  
541 (2005), who demonstrated that spherulitic growth occurs via growth front nucleation. This  
542 mechanism consists of the continuous nucleation of misaligned equivalent structural units on  
543 the surface of growing spherulites, and proceeds via unidirectional growth with low angle  
544 branching (Fig. 9d). This process is nucleation controlled with very little, if any, growth of  
545 the mineral phase. Spherulites occur in many systems (i.e., organic and inorganic), and  
546 usually require high levels of supersaturation in order drive the nucleation controlled process  
547 (Shtukenberg et al., 2012; Andreassen, 2005; Andreassen et al., 2010; Beck and Andreassen,  
548 2010). These observations are also consistent previous studies who observed  
549 monohydrocalcite crystals with spherulitic morphologies (Kimura and Koga, 2011; Neumann  
550 and Epple, 2007; Dahl and Buchardt, 2006, Dejehet et al., 1999; Ferguson et al., 1978;  
551 Duedall and Buckley, 1971; Nishiyama et al., 2013).

552 Therefore, taking into account its crystal chemistry, crystallite size, thermal behaviour  
553 and morphology, we propose a new categorization for monohydrocalcite:

554 - High-Mg monohydrocalcite ( $\chi_{\text{MgCO}_3} > 0.06$ ) consists of individual nanometer-sized  
555 crystals (<35 nm) (Fig. 9c) with a significant part (6-25%) of its structural  $\text{H}_2\text{O}$  being  
556 associated with the Mg ion, therefore displaying a progressive dehydration during heating to  
557 >500°C (Fig. 9b). Such high-Mg monohydrocalcites are uncommon in nature, but can be  
558 synthesised in the laboratory at high initial supersaturation levels ( $\text{SI} > 3.25$ ).

559 - Low-Mg monohydrocalcite ( $\chi_{\text{MgCO}_3} < 0.06$ ) which forms a “type 2” spherulite  
560 morphology. Less than 6% of the structural water in the low-Mg monohydrocalcite is bonded  
561 to Mg, so it fully dehydrates at low temperatures (150-200°C). They have the same  
562 composition as natural monohydrocalcites reported in the literature, and can be synthesized in  
563 the laboratory at lower supersaturation levels ( $\text{SI} < 3.25$ ).

564 These observations indicate that despite their different morphologies (single  
565 nanometer sized crystals and low-angle branching spherulites, respectively; Fig. 9c and 9d)

566 and levels of supersaturations at which they form, high- and low-Mg monohydrocalcite both  
567 crystallize via a nucleation-dominated growth process. The difference in particle size and  
568 morphology is likely controlled by the aqueous Mg concentration. At high concentrations,  
569 Mg poisons the formation of spherulites but still allows direct nucleation in solution,  
570 producing the non-aggregated, individual high-Mg monohydrocalcite crystals. At low  
571 supersaturations, the Mg concentration is low and monohydrocalcite forms via growth-front  
572 nucleation permitting the development of the low angle branching “type 2” spherulites.

573 Combining the mechanistic results described above with chemical data from our on-  
574 line experiment, and data from other studies (Fig. S3) reveals interesting relationships.  
575 Firstly, our on-line experiment shows an increase in nanocrystal sizes during the secondary  
576 crystallisation of monohydrocalcite, which is coupled with a significant decrease in  $\chi_{\text{MgCO}_3}$   
577 (from  $\sim 0.26$  to  $\sim 0.065$ ). This corresponds to the transition from high- to low-Mg  
578 monohydrocalcite, suggesting that the former would be metastable and rapidly transforming  
579 to the latter, possibly triggered by the removal of Mg from aqueous solution. Secondly, Davis  
580 et al. (2000) determined that the solubility of Mg-calcite ( $\text{Ca}_{1-x}\text{Mg}_x\text{CO}_3$ ;  $x=0-0.20$ ) varies by  
581 approximately half an order of magnitude depending on the Mg content of the solid ( $K_{\text{sp}} = 10^{8.0}$   
582  $- 10^{8.5}$ ). A similar behaviour should be expected for monohydrocalcite. The saturation  
583 indexes calculated for monohydrocalcite using the available solubility products from Hull  
584 and Turnbull (1973) and Krajl and Brečević (1995) are negative and show a difference of  
585  $\sim 0.55$  (Fig. 8). We suggest that may be due to difference in the Mg contents of the  
586 monohydrocalcites used in their respective studies. This hypothesis is supported by the recent  
587 findings of Nishiyama et al. (2013), who reported that the solubility of synthetic  
588 monohydrocalcite increases with higher Mg/Ca ratios in the solid. They showed that the  $K_{\text{sp}}$   
589 of monohydrocalcite can reach maximum values of  $K_{\text{sp}} = 10^{-6.77}$ , which is almost one order of  
590 magnitude higher than the value of Hull and Turnbull (1973). Furthermore, Nishiyama et al.  
591 (2013) also reported an decrease in monohydrocalcite crystallite size (broadening of Bragg  
592 peaks in PXRD) with increasing solid Mg/Ca ratios, which again support our interpretation of  
593 the transition from high- to low-Mg monohydrocalcite.

594 Although an in-depth study of the structural changes in monohydrocalcite as a  
595 function of Mg content are outside of the scope of this study, the changes in unit cell  
596 parameters during crystallisation (Fig. 6) may be better understood when the  $\chi_{\text{MgCO}_3}$  of the  
597 monohydrocalcite is taken into account. Regardless of Mg content, the unit cell volume  
598 remained virtually constant during stages II-III and only decreased slightly during stage IV  
599 ( $\Delta V \approx -0.2 \text{ \AA}^3$ ), in parallel with the decrease in monohydrocalcite  $\chi_{\text{MgCO}_3}$ . This small decrease

600 in volume is a consequence of a mirrored change in a-axis and c-axis dimensions, which may  
601 be explained by the change in the monohydrocalcite internal structure during the loss of Mg  
602 and the transition from the high to low-Mg type (stage III to IV). The structure of  
603 monohydrocalcite is less dense and more open than calcite or aragonite (Effenberger, 1981;  
604 Neumann and Epple, 2007; Swainson, 2008), therefore it may more easily adapt to a  
605 changing Mg content.

606  
607

#### 608 **4. SUMMARY AND IMPLICATIONS**

609

610 Results show that under the conditions of our study monohydrocalcite forms via a  
611 Mg-rich ACC precursor. The Mg-ACC forms rapidly then transforms to monohydrocalcite  
612 via dissolution and reprecipitation, with monohydrocalcite forming via a nucleation-  
613 controlled reaction. This crystallization path is consistent with those observed for the  
614 formation of vaterite, calcite and dolomite from (Mg-)ACC. The crystalline phase which  
615 forms from the (Mg-)ACC is controlled by the magnesium content of the precursor: pure  
616 ACC crystallises to vaterite (Bots et al., 2012); 10% Mg-ACC crystallises to calcite  
617 (Rodriguez-Blanco et al., 2012); ~ 30% Mg-ACC crystallises to monohydrocalcite (this study  
618 and Nishiyama et al., 2013), and 50% Mg-ACC crystallises to protodolomite/dolomite at  
619 higher temperatures (Rodriguez-Blanco et al., 2013). The nucleation controlled growth of  
620 these phases from ACC is thought to be due to the high solubility of (Mg-)ACC. Any  
621 solution in equilibrium with the amorphous phase will be highly supersaturated with respect  
622 to the crystalline phase, leading to nucleation controlled growth. This highlights the universal  
623 nature of the nucleation-controlled formation during crystallization from ACC.

624 Our results highlight the importance of Mg in the formation of monohydrocalcite. The  
625 mechanisms by which Mg controls phase stability are complex, but are related to 3 key  
626 factors. Firstly, the presence of Mg in solution is known to inhibit the formation of vaterite  
627 and calcite (Bischoff, 1968; Kitano, 1966; Bischoff and Fyfe, 1968; Berner, 1975; Mucci and  
628 Morse, 1983; Deleuze and Brantley, 1997; Chen et al., 2004; Bots et al., 2011). We suggest  
629 that the Mg levels in the current study (i.e.  $Mg/(Mg + Ca) = 0.3$ ) were high enough to inhibit  
630 the formation of both calcite and vaterite, and favoured the formation of monohydrocalcite.  
631 Secondly, the hydration shell of Mg is more strongly bound than the hydration shell of Ca (di  
632 Tommaso and de Leeuw, 2010; Moomaw and Maguire, 2008). The energy required to  
633 desolvate Mg is higher than that of Ca, a fact which explains the higher temperature required

634 to crystallize anhydrous Ca-Mg carbonates, e.g. dolomite or magnesite (Christ and Hostetler  
635 1970; Lipmann, 1973; Kittrick and Peryea, 1986; Rodriguez-Blanco et al., 2013). The  
636 hydrated nature of monohydrocalcite means that full dehydration of Mg is not required before  
637 incorporation into the crystal lattice, therefore it will more likely form than the anhydrous  
638 calcium carbonate phases. The TGA analysis of the monohydrocalcite samples support this,  
639 as it shows increasing water content with Mg (Fig. 9). Thirdly, monohydrocalcite has a high  
640 capacity to accommodate Mg within its structure favouring its formation at higher Mg/Ca  
641 ratios. Thus although the initial solutions used in our work were more supersaturated with  
642 respect to all anhydrous CaCO<sub>3</sub> polymorphs, the formation of monohydrocalcite was  
643 favoured.

644 We have shown that the crystal chemistry and growth morphology of  
645 monohydrocalcite (Stoffers and Fischbeck, 1974; Taylor, 1975; Munemoto and Fukushi,  
646 2008) can be controlled by the initial supersaturation of the aqueous solution. We showed  
647 that with increasing supersaturation the Mg content in the monohydrocalcite increases up to  
648  $\chi_{\text{MgCO}_3} = \sim 0.26$ , similar to the Mg-contents in Mg-calcites reported previously (Lenders et  
649 al., 2012; Goldschmidt et al., 1955; Goldschmidt and Graf, 1958). This has led to us defining  
650 2 types of monohydrocalcite i.e. high-Mg and low Mg monohydrocalcite, classified  
651 according to their Mg content and morphology. All natural monohydrocalcites, which have  
652 been analysed to date, are low Mg monohydrocalcites (Mg content  $\sim 0.01 > \chi_{\text{MgCO}_3} < 0.075$ ,  
653 Hull and Turnbull, 1973; Taylor, 1975; Fukushi et al., 2011 and references therein). We  
654 suggest that the lack of high-Mg monohydrocalcite in natural environment is due to 2 factors.  
655 Firstly, for high-Mg monohydrocalcite to form, initial solutions must be extremely highly  
656 supersaturated ( $SI_{\text{MHC}} \sim 3.5 - 4$ ; see Table 1 and Table 2). Such extremely high  
657 supersaturation levels are difficult to achieve through natural processes. Secondly, our  
658 analysis of the monohydrocalcite crystallization pathway shows that high-Mg  
659 monohydrocalcite breaks down to low-Mg monohydrocalcite within hours of its formation,  
660 therefore high-Mg monohydrocalcite if it forms at all is likely a transient phase. Furthermore,  
661 despite monohydrocalcite not being as abundant as calcite and or aragonite in the geological  
662 record, it may still have a potential as a palaeoenvironmental indicator (Solotchina et al.,  
663 2009). The isotopic composition of the hydration water in natural low-Mg monohydrocalcites  
664 may be useful as a  $\delta^{18}\text{O}$  record of the palaeoenvironmental conditions of the water from  
665 which it crystallized. Such an approach has been successfully applied to other hydrated  
666 carbonates like ikaite, which is even scarcer in nature and more unstable than  
667 monohydrocalcite (Rickaby et al, 2011; Lu et al., 2012).



668           Nevertheless, our experiments documented a simple methodology to synthesize  
669 monohydrocalcite with specific Mg/Ca ratios, specific particle sizes and shapes (i.e., reactive  
670 surface), which may have applications in a number of fields. For example, synthetic  
671 monohydrocalcite have been reported to have a high sorption capacity for  $\text{PO}_4^{3-}$  and  $\text{AsO}_4^{3-}$   
672 (Yagi and Fukushi, 2012; Yagi and Fukushi, 2011; Fukushi et al., 2011). Thus, by producing  
673 monohydrocalcite with specific compositions, particle sizes and shapes the effectivity of  
674 these remediation materials could be improved.

675           Finally, the formation of monohydrocalcite and other CaMg carbonates as a  
676 consequence of biological activity has been reported in many systems (Lowenstam, 1981;  
677 Sánchez-Román et al., 2011). Microorganisms often promote the formation of highly  
678 supersaturated microenvironments which facilitate the precipitation of carbonates (e.g.,  
679 bacterial cells which absorb Ca, Mg or other metallic cations and act as nucleation sites),  
680 even when the conditions for supersaturation have not been reached in the surrounding  
681 environment (Vasconcelos and Mckenzie, 1997; Párraga et al., 1998). Systems characterized  
682 by high initial aqueous Mg/Mg+Ca values (e.g.,  $\geq 0.3$ ) will lead to the formation of a Mg-rich  
683 ACC precursor, which in turn will crystallize to monohydrocalcite. We suggest that this may  
684 be the primary crystallization pathway during biomineralization of monohydrocalcite.

685           It is interesting to note that monohydrocalcite forms at a broad range conditions in the  
686 laboratory and natural environment (i.e. a wide range of solution compositions and Mg/Ca  
687 ratios) (Nishiyama et al., 2013; Wood, 2012; Fukushi et al., 2011 and references therein).  
688 Yet, monohydrocalcite is scarce in nature, a fact that can be explained by (1) its metastability  
689 with respect to aragonite and calcite, and (2) the high supersaturation required to form it  
690 compared to other calcium carbonates. However, the chemistry of many carbonates formed  
691 during biomineralization processes indicates that they have formed from high Mg/Ca ratios  
692 seawater solutions (Davis et al., 2000; Ries, 2010; Ries et al, 2008; Ries, 2004; Reeder et al.,  
693 1983), and in some cases at high supersaturations (e.g., Señorale-Pose et al., 2008; Hood et  
694 al., 2011; Hood and Wallace, 2012). Furthermore, the paragenesis of some mineral deposits  
695 which now contain aragonite also include a variety of Ca-Mg-bearing carbonates like  
696 dolomite, hydromagnesite or nesquehonite (Fischbeck and Mueller, 1971; Broughton, 1972).  
697 Contrary to monohydrocalcite, aragonite does not necessarily require the presence of Mg to  
698 form (e.g., Ogino et al., 1987; Sand et al., 2012), but it is well known that Mg and  $\text{SO}_4^{2-}$  are  
699 the key ions responsible for the switch from calcite to aragonite seas (Sandberg, 1983; Bots et  
700 al., 2011). During geological timescales the mineralogy of most calcifying organisms was  
701 influenced by the Mg/Ca ratio of the oceans in which they evolved (Stanley, 2006; Porter,

702 2007). For that reason we consider that some aragonite and (Mg)-calcite deposits that are  
703 known to have formed at high Mg/Ca ratios (e.g., coralline red algal beds; rhodoliths, Schäfer  
704 et al., 2011; ooids, Brehm et al., 2006; Davies et al., 1978; Ferguson et al., 1978 or  
705 microbialites, Burne and Moore, 1987) could be secondary in origin and may have originally  
706 been a metastable monohydrocalcite intermediate which recrystallized. This secondary  
707 crystallization via monohydrocalcite could be the reason why the skeletal mineralogy and  
708 chemistry of calcareous organisms that recorded the Mg/Ca ratio of the seawater in which  
709 they grow varied so dramatically during the Phanerozoic (Ries, 2004) and this in turn may  
710 affect Mg/Ca ratio – based paleoenvironmental reconstructions. This may also explain why  
711 some early Cambrian fossils may have preferentially formed low-cost, high-Mg calcite, while  
712 later the increasing physiological cost of biomineralization lead to low Mg-calcite skeletons  
713 (Wood, 2011). Whether monohydrocalcite played a role in these early biomineralization  
714 processes is still an open question, although if monohydrocalcite was an intermediate phase  
715 during these biomineralization reactions it may have big implications for seawater chemistry  
716 reconstructions and other paleoproxies and would require further study.

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## 728 6. REFERENCES

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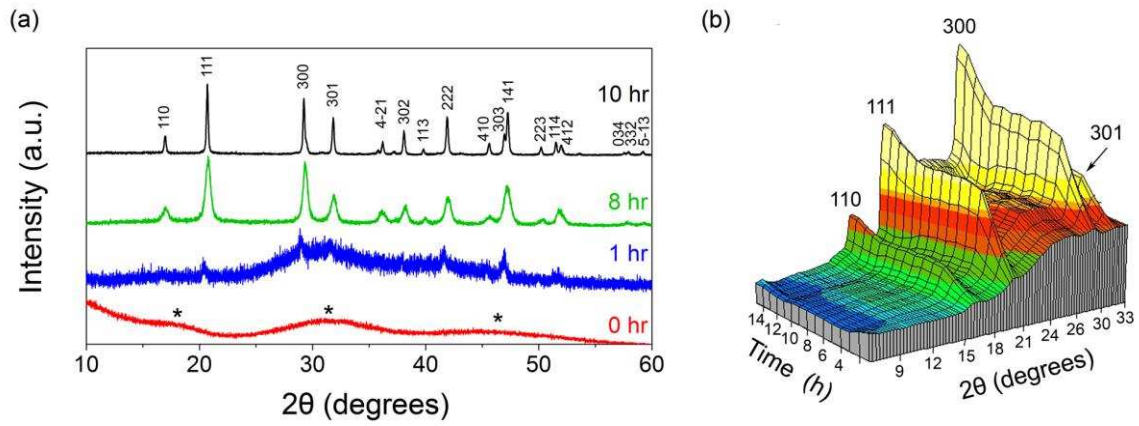
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1032 **Figures**

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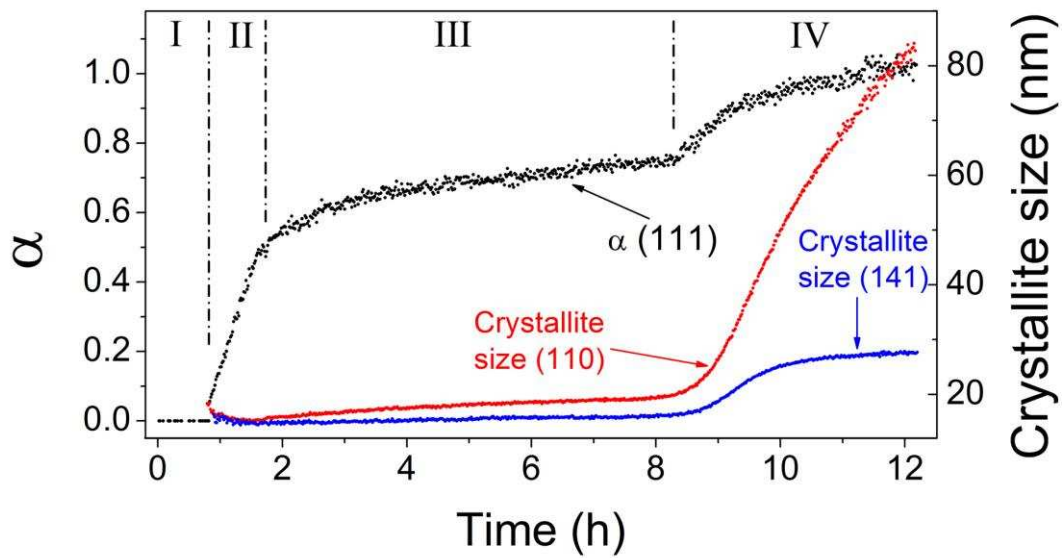
1039 **Fig. 1.** (a) PXR D patterns from off-line crystallization experiments with samples removed

1040 from each run after 0, 1, 8 and 10 hr. (b) 3D WAXS plot showing the time resolved formation

1041 of monohydrocalcite from the on-line experiment with only some major peaks shown.

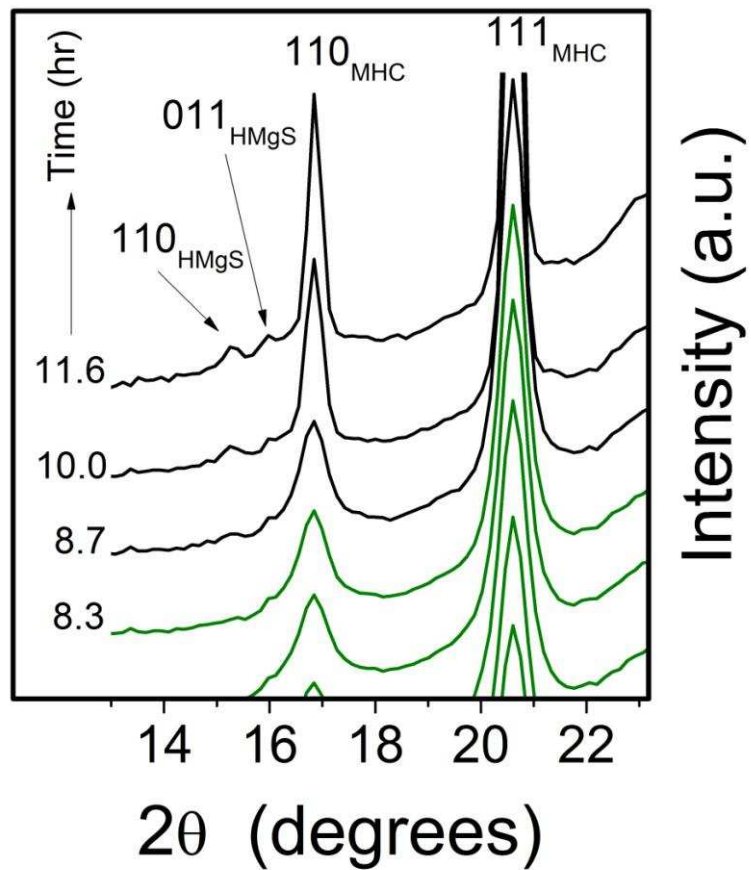
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**Fig. 2.** Evolution of the degree of reaction ( $\alpha$ ) of the monohydrocalcite (111) Bragg peak (black symbols). The same growth profiles are true for all other Bragg peaks. Shown also with the right-hand y-axis are Scherrer crystallite sizes for the (110) and (141) peaks of monohydrocalcite as a function of time as derived from the WAXS data. I to IV correspond to the crystallization stages described in the text.



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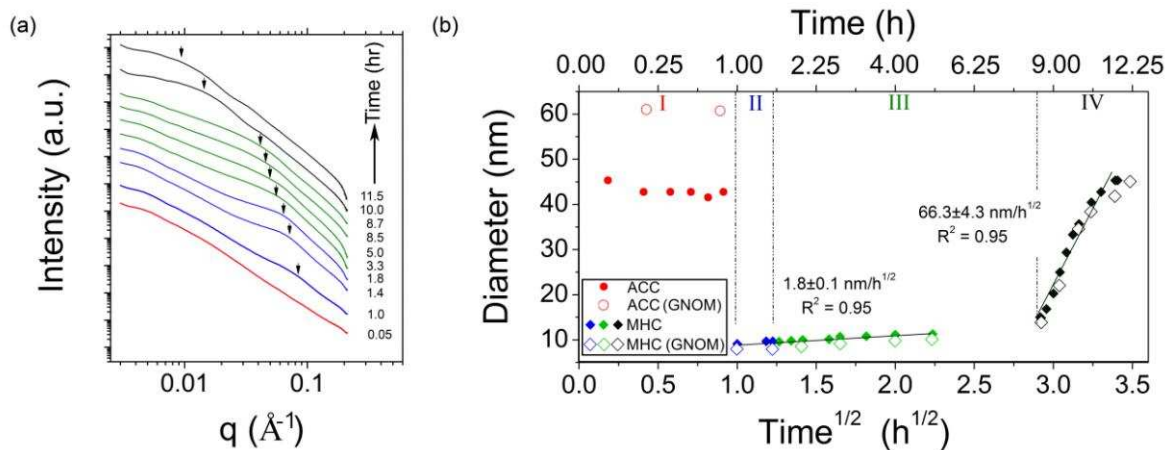
1055 **Fig. 3.** Selected WAXS patterns from the on-line experiment showing the growth of the  
 1056 monohydrocalcite (MHC) 110 and 111 Bragg peaks and the formation of the small  
 1057 hydromagnesite (HMgS) 110 and 011 Bragg peaks during stages III (green patterns) and IV  
 1058 (black patterns) of the reaction.

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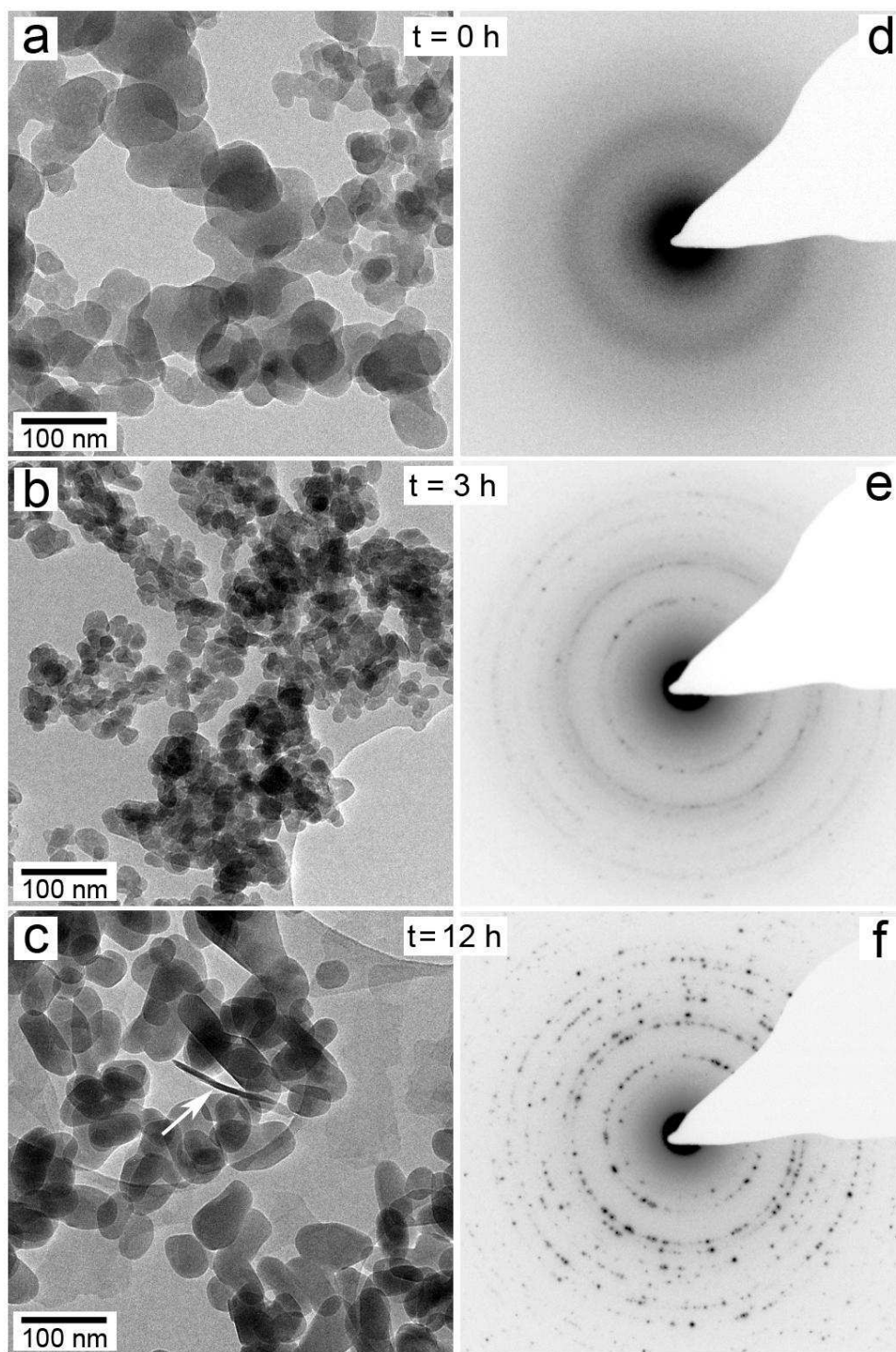
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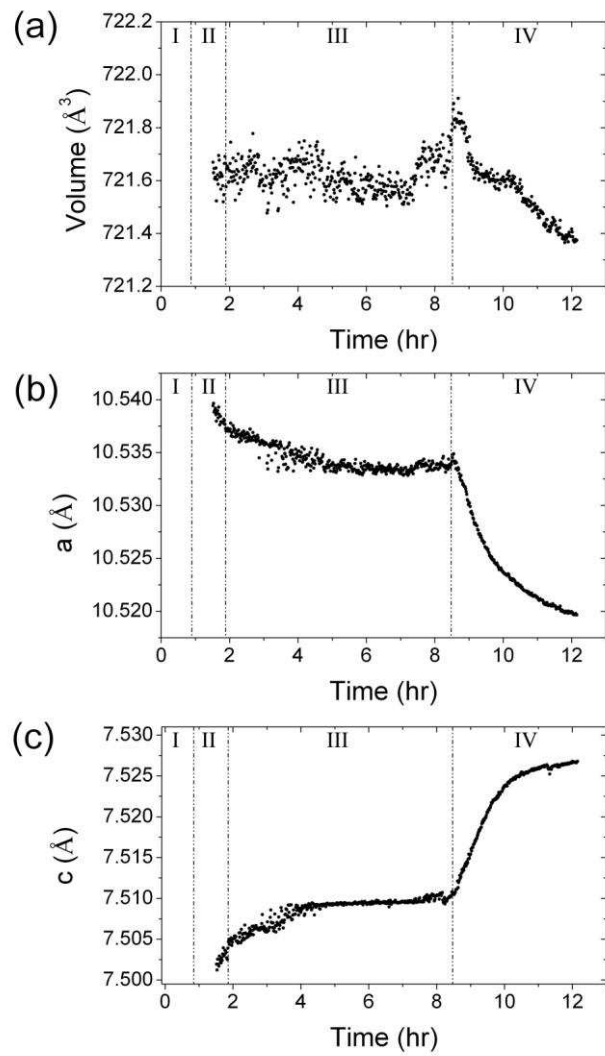
**Fig. 4.** (a) Double logarithmic plot showing selected SAXS patterns from the on-line in situ experiment with arrows indicating a changing position of the scattering peak, reflecting the growth of monohydrocalcite particles with time; colours of the patterns matches the colour code of the four stages of reaction shown in (b) Time evolution of the particle diameters for the initial amorphous phase (Mg-ACC stage I, red symbols) and for monohydrocalcite (MHC stages II to IV, blue, green and black symbols respectively) as derived from the SAXS data. Empty symbols correspond to GNOM code (Svergun, 1992). The differences in growth rates of the monohydrocalcite particles are evidenced by the fit in stages II-III and IV.





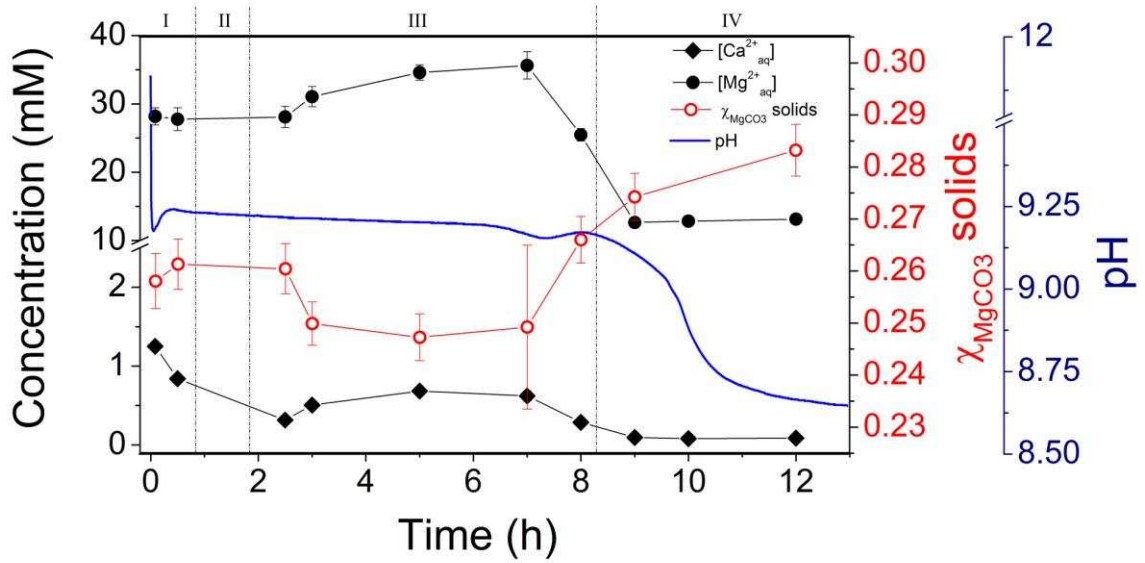
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**Fig. 5.** HR-TEM microphotographs and corresponding SAED patterns of solids during the crystallization reaction. (a, d) Mg-ACC precursor; stage I); (b, e) Monohydrocalcite nanocrystals from stage III; (c, f) Bigger monohydrocalcite particles and a platy hydromagnesite crystal (white arrow) from the end of stage IV.



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**Fig. 6.** Evolution of the unit cell volume (a) and of the a and c parameters (panels b and c) for monohydrocalcite over time as derived from the on-line WAXS data for stages II to IV.



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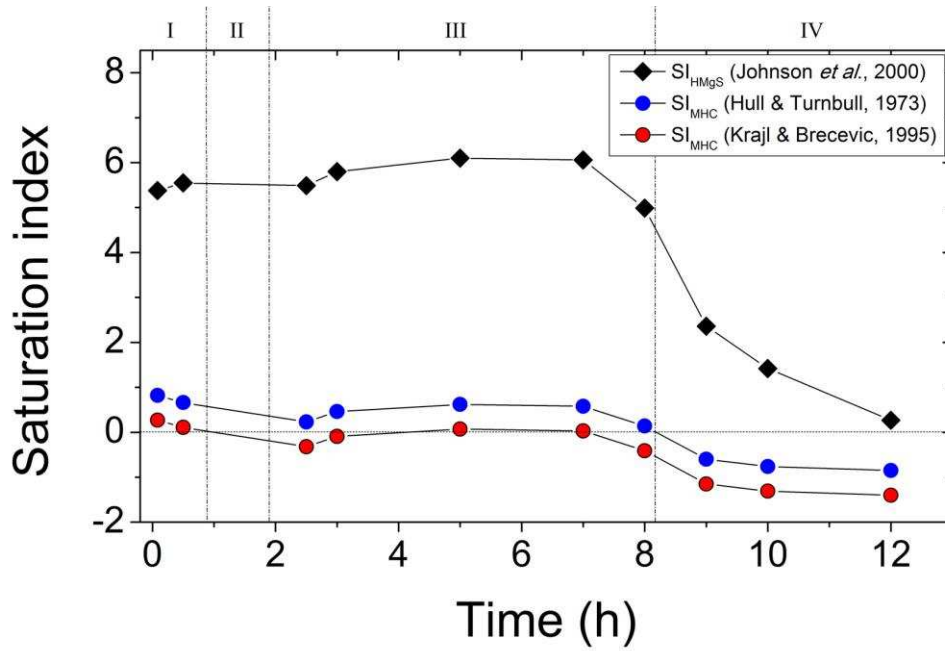
1092 **Fig. 7.** Evolution of the pH, the concentrations of  $[Ca^{2+}]_{aq}$ ,  $[Mg^{2+}]_{aq}$  and of the Mg molar  
 1093 fraction ( $\chi_{MgCO_3}$ ) in the solids over the course of the four stages of the crystallization reaction.

1094 Error bars correspond to the standard deviation of three measurements each.

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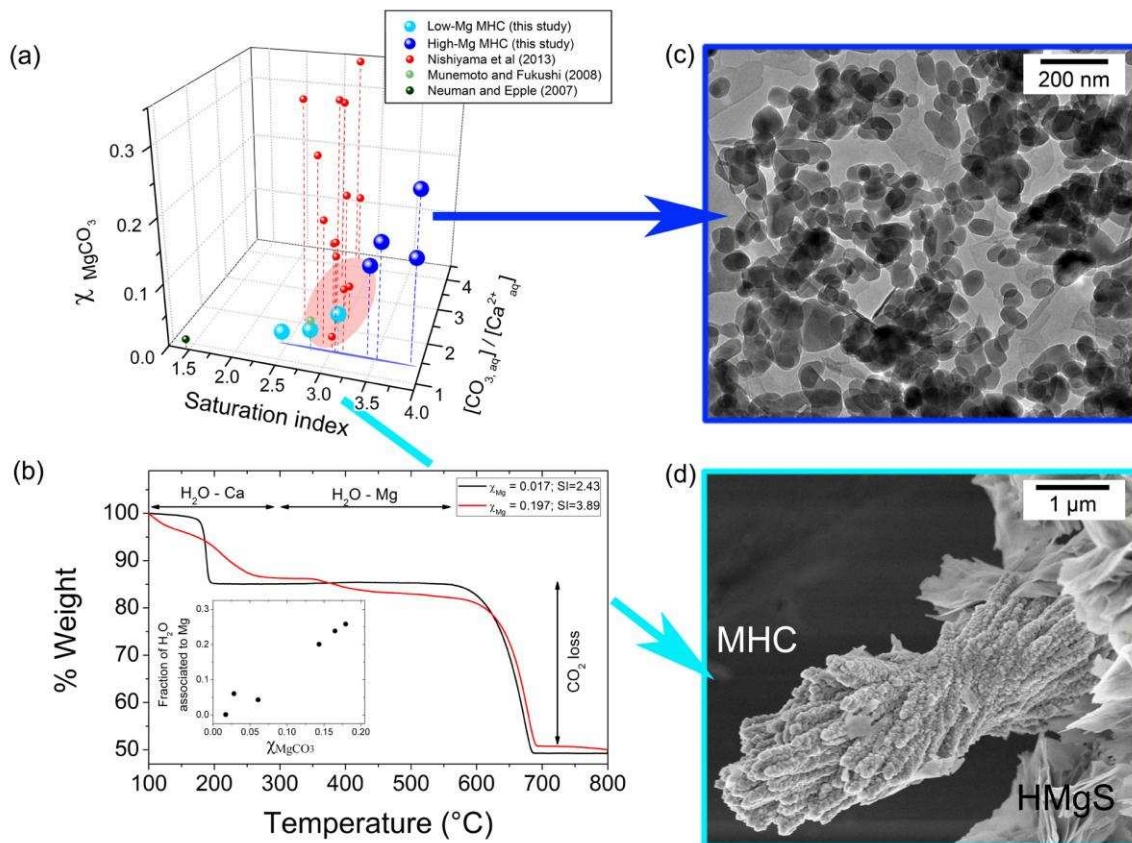
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1100 **Fig. 8.** Evolution of the saturation indexes for monohydrocalcite ( $SI_{MHC}$ ) and hydromagnesite

1101 ( $SI_{HMgS}$ ) calculated from the data shown in Fig. 7 and Table 1.

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1107 **Fig. 9.** (a) Three dimensional plot showing the relationship between the molar fraction of Mg  
 1108 ( $\chi_{\text{MgCO}_3}$ ) in monohydrocalcite as a function of its initial saturation index ( $\text{SI}_{\text{MHC}}$ ) and starting  
 1109  $[\text{CO}_{3,\text{aq}}]/[\text{Ca}^{2+}_{\text{aq}}]$  ratios. Blue and turquoise larger spheres are our off-line experimental data  
 1110 (Table 2) that were all obtained at the same starting  $[\text{CO}_{3,\text{aq}}]/[\text{Ca}^{2+}_{\text{aq}}]$  ratio; the smaller, red,  
 1111 green and black spheres correspond to literature data. (b) Comparative TGA graphs for  
 1112 monohydrocalcite synthesised at high (SI=3.89) and low (SI=2.43) initial supersaturations  
 1113 (Table 2) with the inset showing the fraction of Mg-bonded structural water in  
 1114 monohydrocalcite vs.  $\chi_{\text{MgCO}_3}$ . Images (c) and (d) illustrate the differences in morphology,  
 1115 particle sizes and formation modes for our high- and low-Mg monohydrocalcite (MHC),  
 1116 respectively. Hydromagnesite (HMgS) crystals are also present in image (d).

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**Table 1.**

Evolution of the aqueous solution composition, the magnesium content in the solids ( $\chi_{\text{MgCO}_3}$ ) and the solid mineralogy during the course of a typical experiment. Saturation indexes for monohydrocalcite, hydromagnesite and nesquehonite ( $\text{SI}_{\text{MHC}}$  and  $\text{SI}_{\text{HMgS}}$ ,  $\text{SI}_{\text{nesq}}$ ) were calculated from the measured pH and  $[\text{Ca}^{2+}_{\text{aq}}]$  and  $[\text{Mg}^{2+}_{\text{aq}}]$  values combined with the solubility products for monohydrocalcite reported by (1) Hull and Turnbull (1973), and (2) Krajl and Brečević (1995) and those for hydromagnesite and nesquehonite from Johnson et al. (2000). Average particle sizes were evaluated from the SAXS data or from photomicrographs.

Stage	Time (h)	pH	$[\text{Ca}^{2+}_{\text{aq}}]$ (mM)	$[\text{Mg}^{2+}_{\text{aq}}]$ (mM)	$\chi_{\text{MgCO}_3}$ solid	Mineralogy	Particle sizes from SAXS / Crystallite sizes from Scherrer (nm)	Particle sizes from TEM imaging (nm)	$\text{SI}_{\text{MHC}}$ (1)	$\text{SI}_{\text{MHC}}$ (2)	$\text{SI}_{\text{HMgS}}$	$\text{SI}_{\text{nesq}}$
Pre-mixing	0	-	350	150	-	-	40-60 / -	-	3.89	3.34	13.06	1.17
I	0.083	9.239	$1.25 \pm 0.05$	$28.17 \pm 1.26$	$0.258 \pm 0.005$	Mg-ACC	40-60 / -	~ 40	0.82	0.27	5.38	0.03
	0.5	9.217	$0.84 \pm 0.04$	$27.75 \pm 1.68$	$0.261 \pm 0.005$		43 / -		0.66	0.11	5.55	0.04
II	-	-	-	-	-	-	-	-	-	-	-	-
III	2.5	9.214	$0.32 \pm 0.02$	$28.08 \pm 1.57$	$0.260 \pm 0.005$	MHC	10 / 15	28±7 (L) x 20±5 (W)	0.23	-0.32	5.49	0.04
	3	9.203	$0.51 \pm 0.03$	$31.05 \pm 1.48$	$0.250 \pm 0.004$		11 / 15		0.46	-0.09	5.8	0.11
	5	9.167	$0.68 \pm 0.03$	$34.59 \pm 1.14$	$0.247 \pm 0.005$		11 / 16		0.62	0.07	6.1	0.18
	7	9.171	$0.62 \pm 0.06$	$35.62 \pm 2.00$	$0.249 \pm 0.016$		- / 17		0.58	0.03	6.06	0.19
	8	9.111	$0.29 \pm 0.01$	$25.47 \pm 0.93$	$0.266 \pm 0.004$		- / 17.5		0.14	-0.41	4.99	-0.05
IV	9	8.893	$0.09 \pm 0.01$	$12.64 \pm 0.75$	$0.274 \pm 0.005$	MHC+HMgS	15 / 22.5		-0.6	-1.15	2.36	-0.60
	10	8.646	$0.08 \pm 0.00$	$12.83 \pm 0.39$	-		35 / 37.5	155±80 (L) x 40±30 (W)	-0.76	-1.31	1.42	-0.70
	12	8.609	$0.09 \pm 0.00$	$13.12 \pm 0.54$	$0.283 \pm 0.005$		45 / 45		-0.85	-1.4	0.27	-0.85

**Table 2.**

Initial stock solution concentrations, and corresponding calculated monohydrocalcite saturation index ( $SI_{MHC}$ ) for the solution immediately after mixing for each experiment. The first line represents the conditions used in the on-line SAXS/WAXS experiments. The mineralogy, Scherrer crystallite size and Mg ( $\chi_{MgCO_3}$ ), and H<sub>2</sub>O content of the solid products after 2 h of reaction are also presented.. The saturation index of monohydrocalcite was calculated using the solubility product from Hull and Turnbull (1973).

$Na_2CO_3$ <sub>(ini)</sub> (mM)	$[Ca^{2+}]$ <sub>(ini)</sub> (mM)	$[Mg^{2+}]$ <sub>(ini)</sub> (mM)	$SI_{MHC}$ (t = 0 sec)	Mineralogy from PXRD	Scherrer crystallite size (average) (nm)	$\chi_{MgCO_3}$ <sup>*</sup> solid	H <sub>2</sub> O content
500	350	150	3.89	MHC	16	0.164	0.99
250	175	75	3.52	MHC	17	0.180	1.03
200	140	60	3.41	MHC	-	0.143	1.00
100	70	30	3.08	MHC	-	0.061	0.97
50	35	15	2.76	MHC	35	0.028	0.95
25	17.5	7.5	2.43	MHC	35	0.017	0.93
12.5	8.75	3.75	2.10	Mg-Calcite	43	0.073	0
5	3.5	1.5	1.63	Mg-Calcite	131	0.032	0

\* Errors for  $\chi_{MgCO_3}$  of solids are in all cases lower than 0.002

Fig 1.jpg

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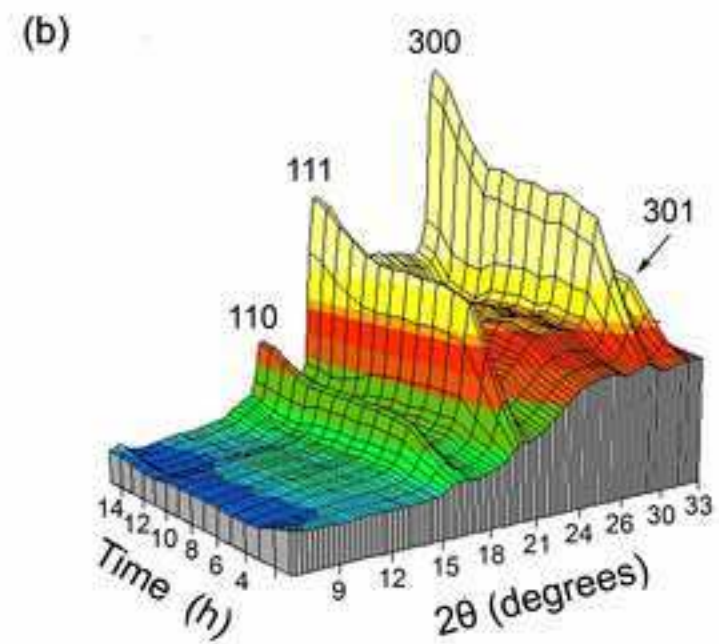
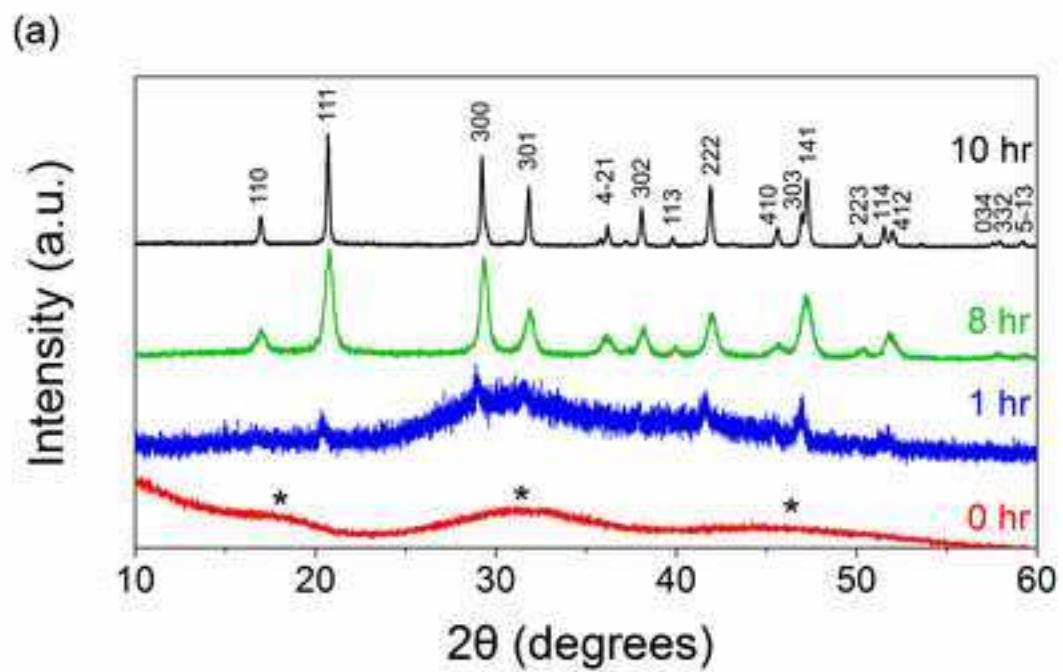




Fig 2.jpg

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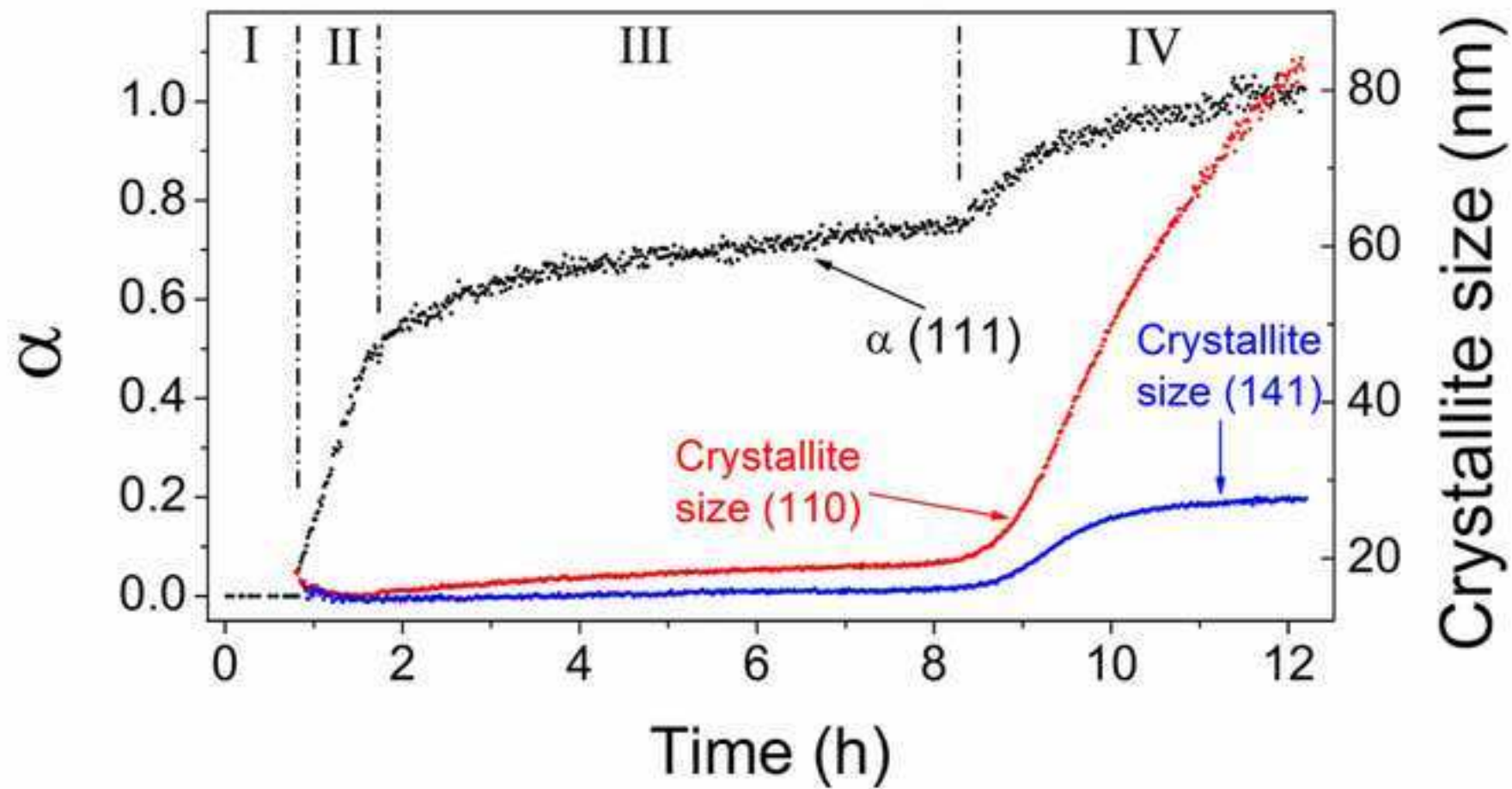
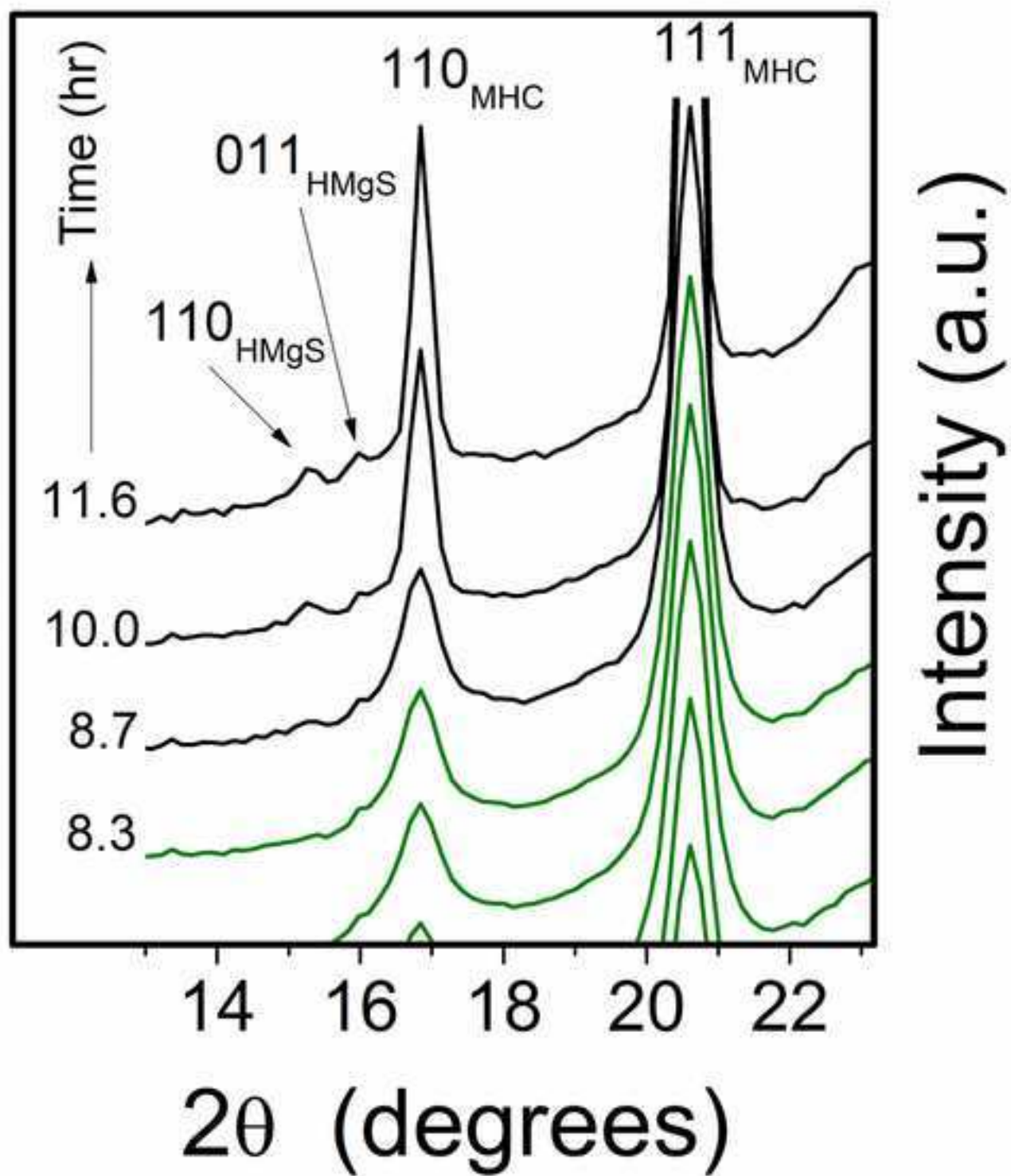


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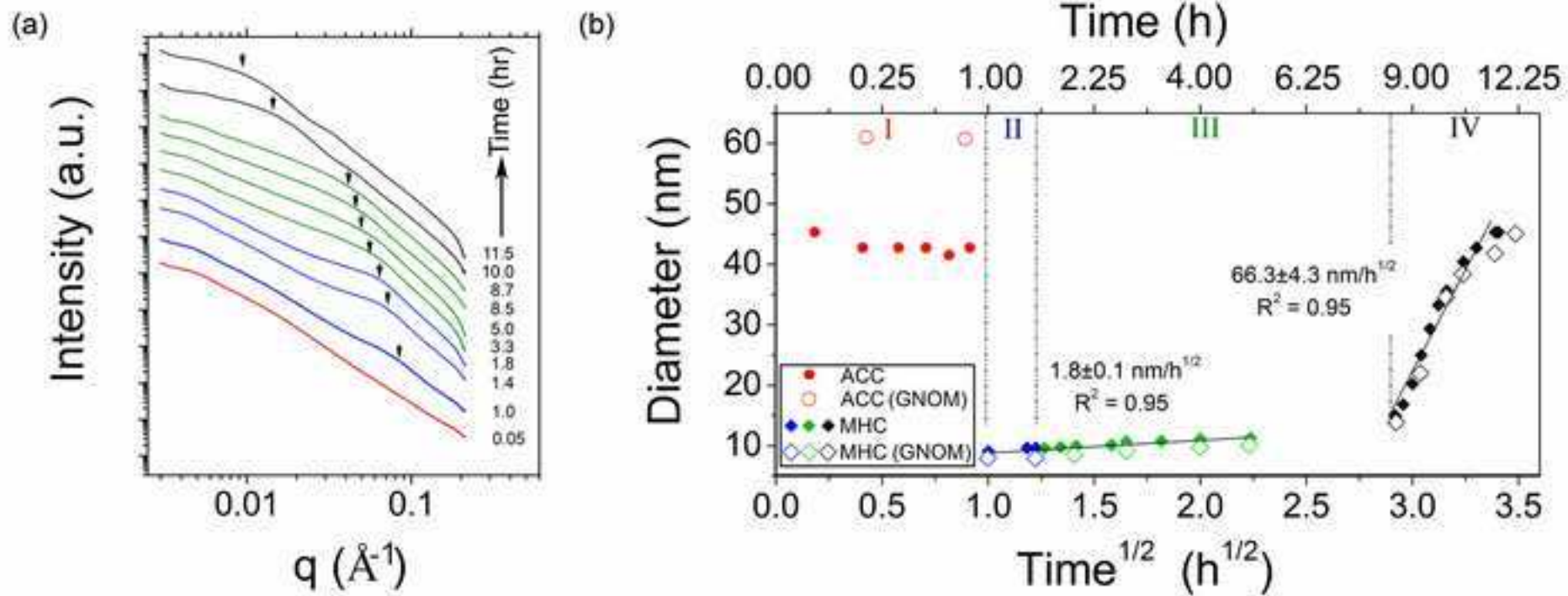
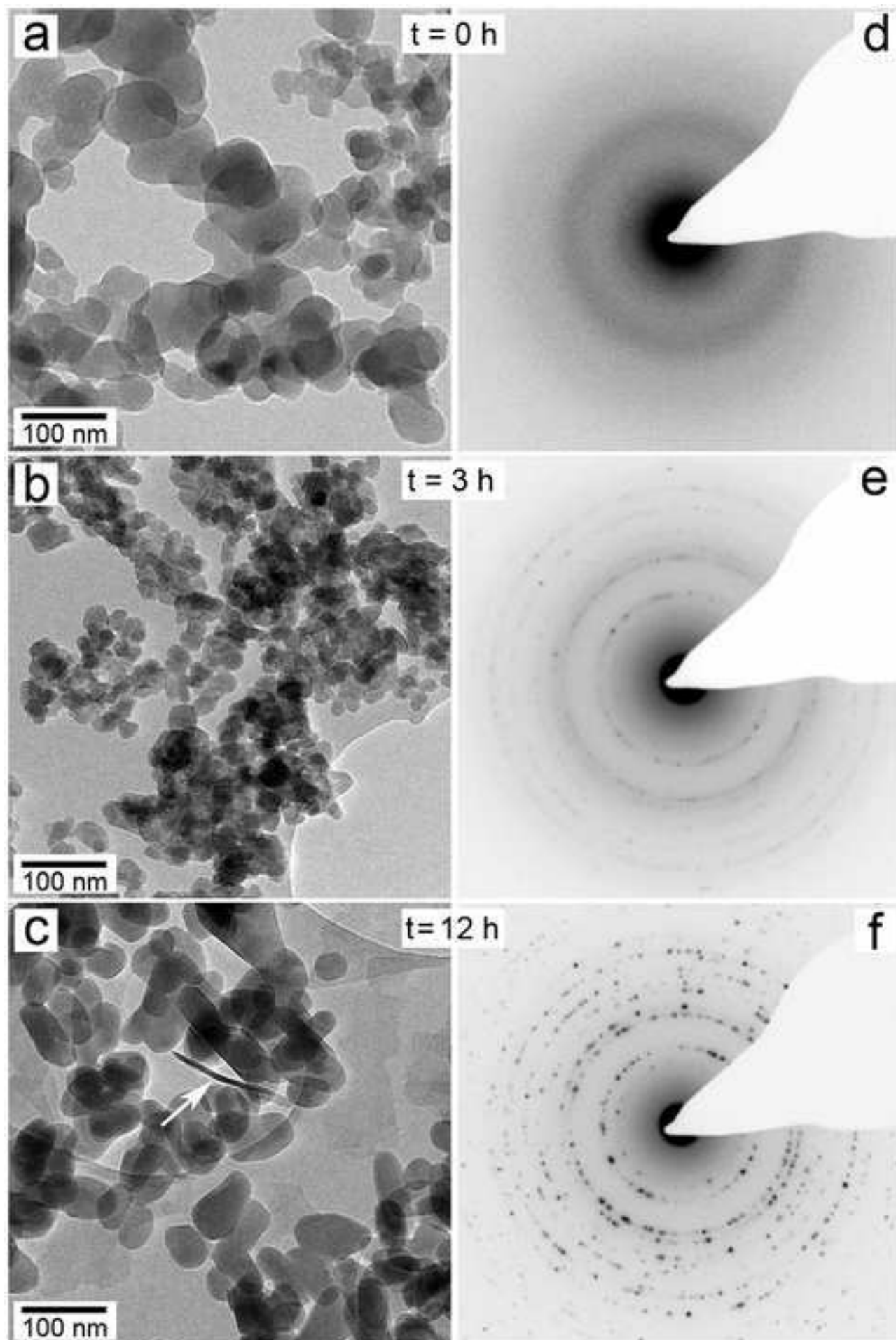


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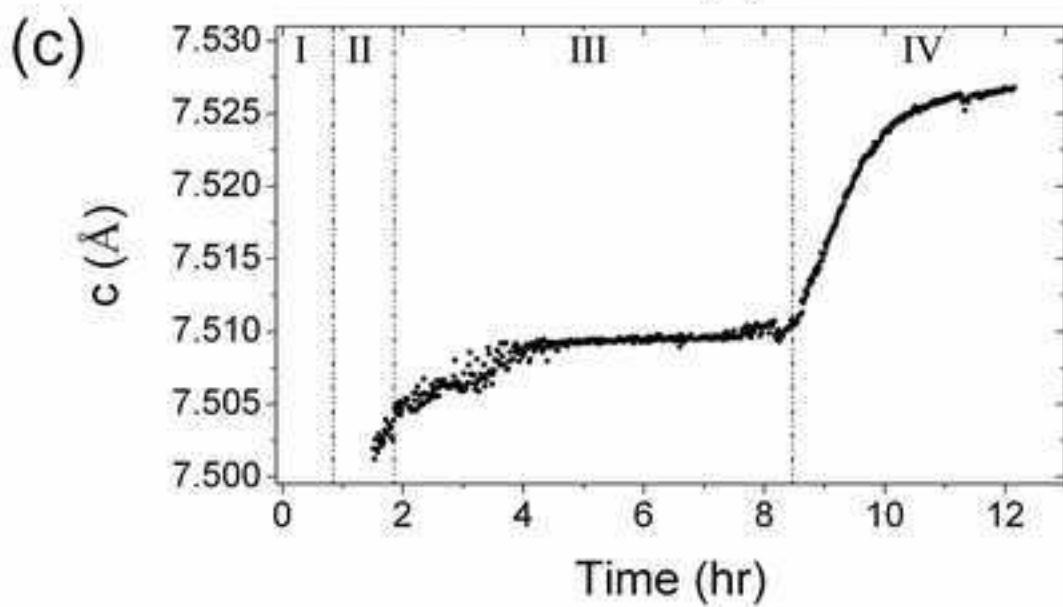
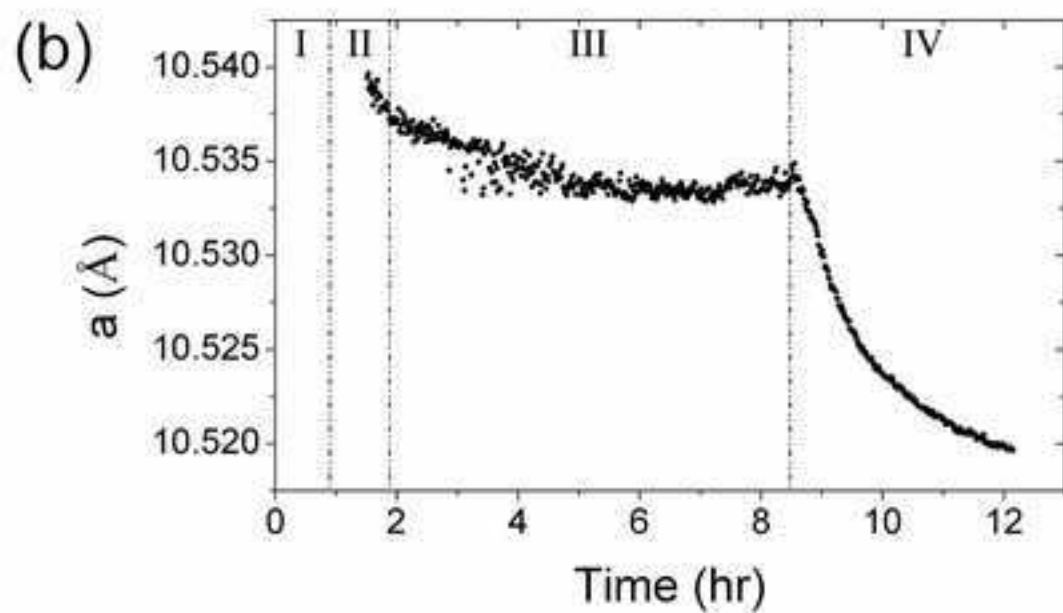
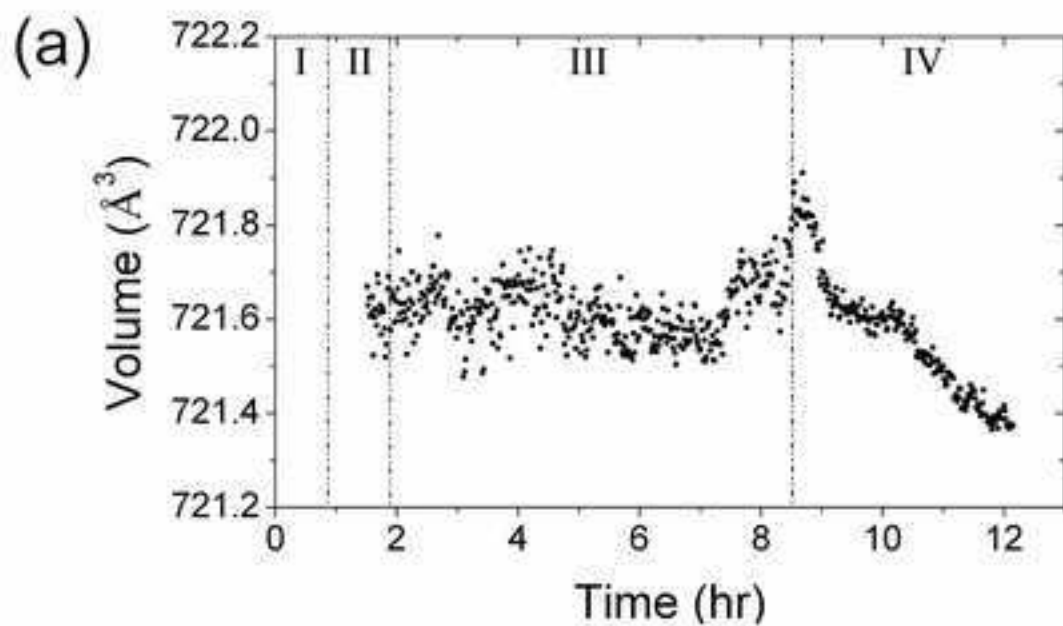




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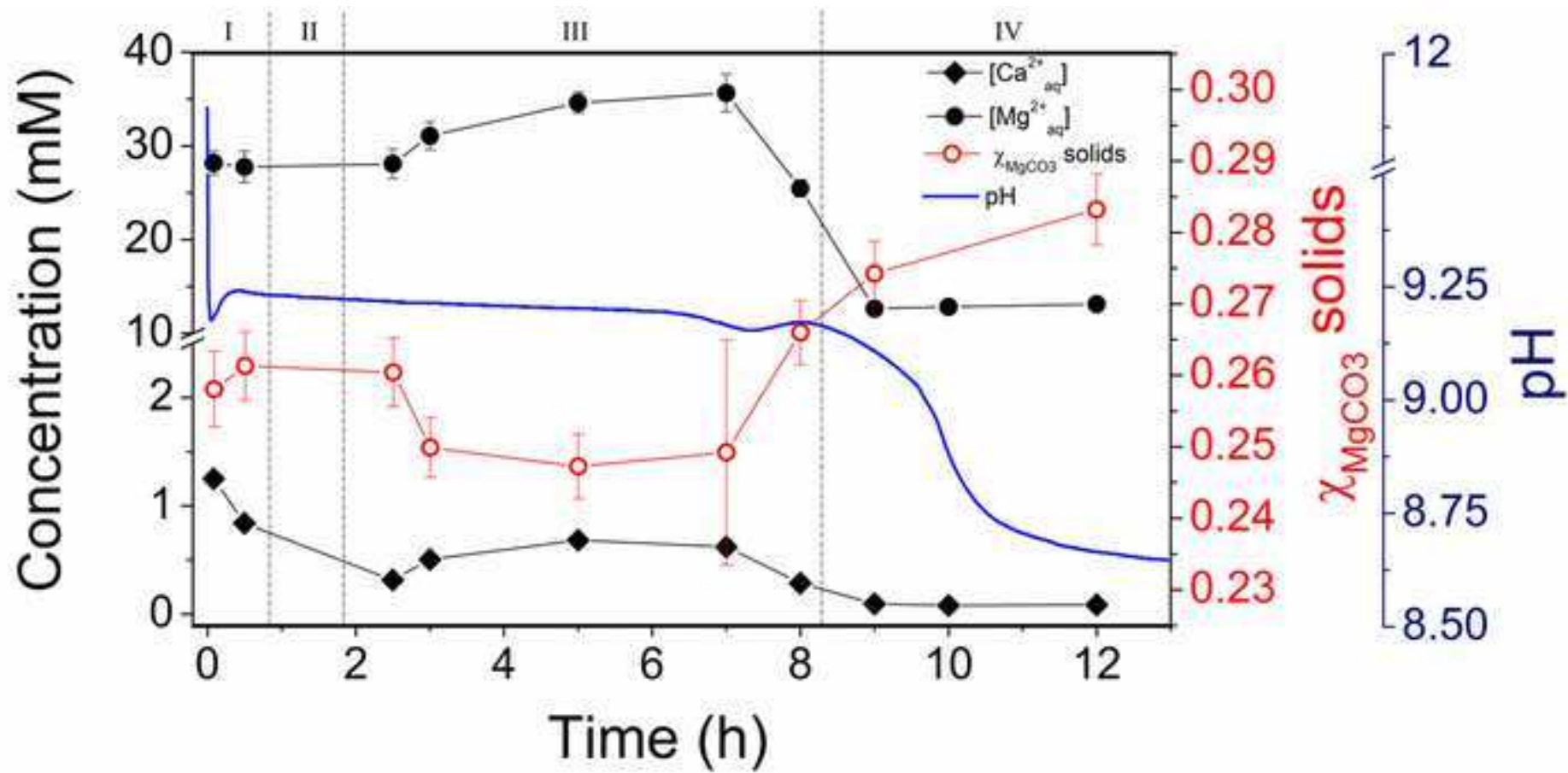


Fig 8.jpg

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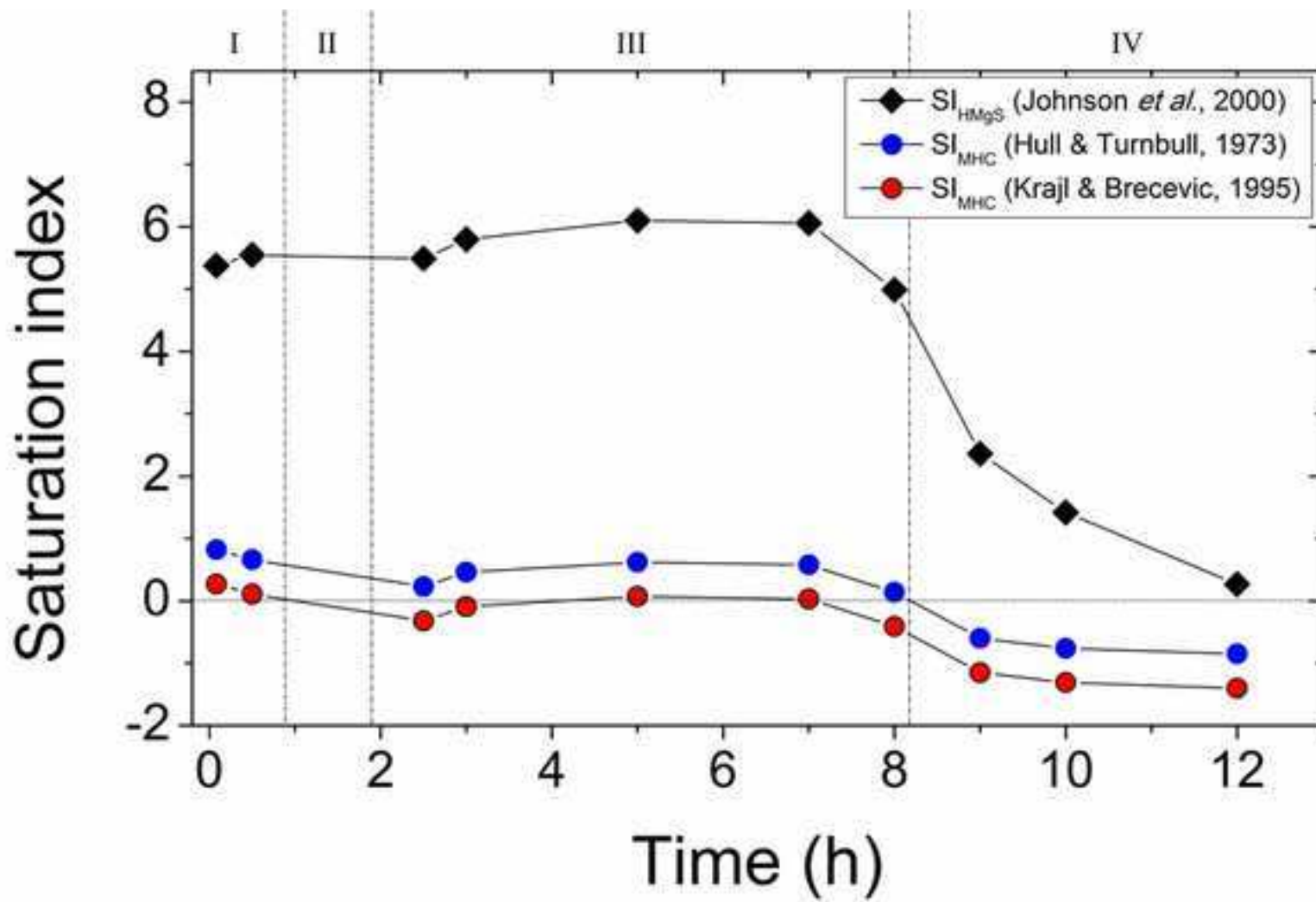
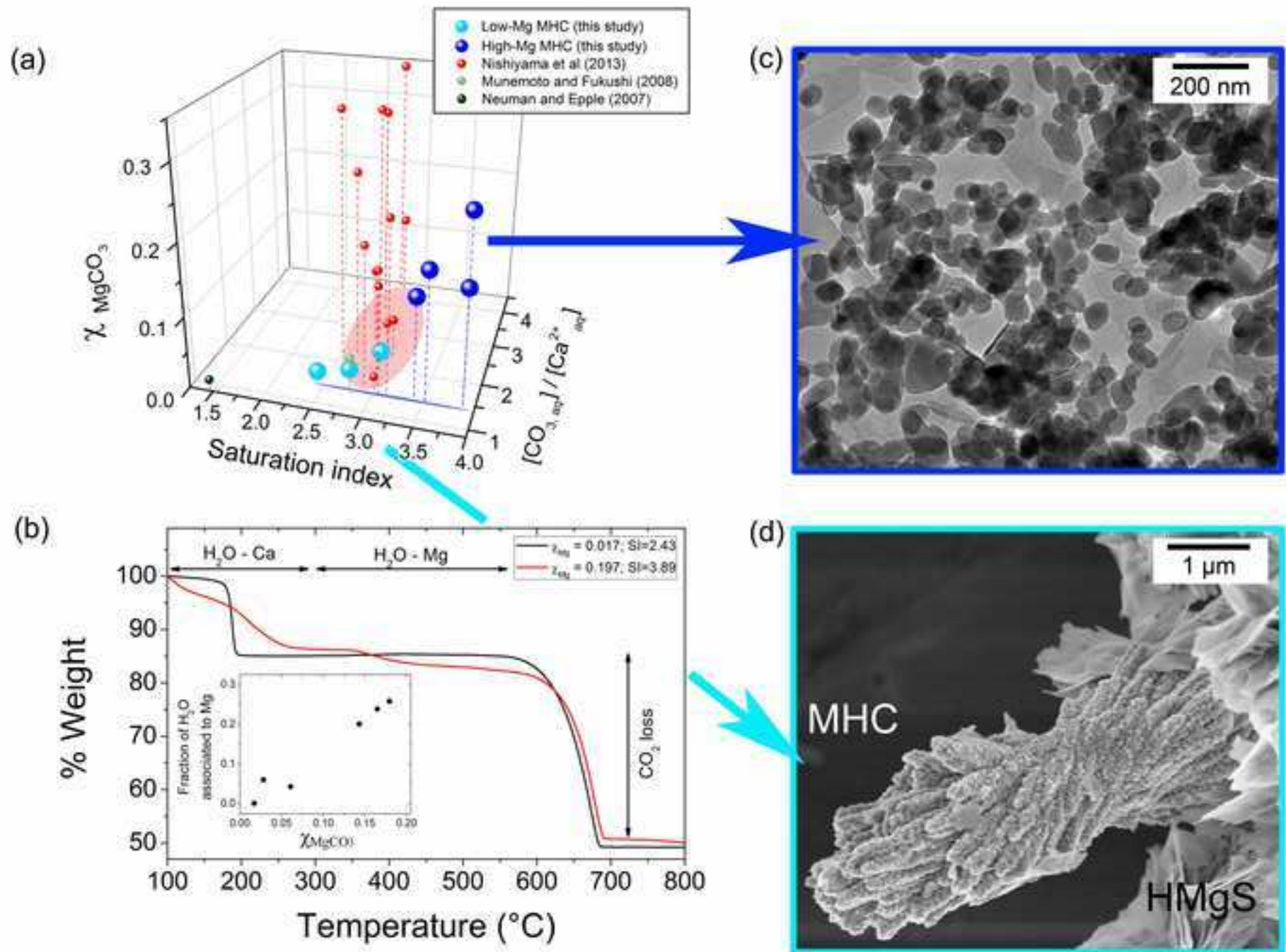


Fig 9.jpg

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