Technical Note: A simple method for vaterite precipitation for isotopic studies: implications for bulk and clumped isotope analysis

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Abstract. Calcium carbonate (CaCO₃) plays an important role in the natural environment as a major constituent of the skeleton and supporting structure of marine life and has high economic importance as an additive in food, chemicals and medical products. Anhydrous CaCO₃ occurs in the three different polymorphs calcite, aragonite and vaterite, whereof calcite is the most abundant and best characterized mineral. In contrast, little is known about the rare polymorph vaterite, in particular with regard to the oxygen isotope fractionation between H₂O and the mineral.

Synthetic precipitation of vaterite in the laboratory typically involves rapid processes and isotopic non-equilibrium, which excludes isotope studies focused on the characterization of vaterite under equilibrium conditions. Here, we used a new experimental approach that enables vaterite mineral formation from an isotopically equilibrated solution. The solution consists of a ∼0.007 mol L⁻¹ CaCO₃ solution that is saturated with NaCl at room temperature (up to 6.4 mol L⁻¹). Vaterite precipitated as single phase or major phase (≥ 94 %) in experiments performed between 23 and 91 °C. Only at 80 °C was vaterite a minor phase with a relative abundance of 27 %. The high mineral yield per experiment of up to 235 mg relative to the initially dissolved CaCO₃ amount of on average 360 mg enables an investigation of the oxygen isotope fractionation between the mineral and water, and the determination of clumped isotope values in vaterite.

1 Introduction

Vaterite is the least known polymorph of calcium carbonate and was first described by Vater (Vater, 1893). In the 1920s and 1930s, the nature and crystallographic structure of vaterite was still questioned and the occurrence of a third polymorph of CaCO₃ was disputed amongst various groups (Johnston et al., 1916; Spangenberg, 1921; Heide, 1924; Rinne, 1924; Gibson et al., 1925). In the following decades, work mainly focused on the structure of vaterite (e.g. McConnell, 1960; Kamhi, 1963; Meyer, 1969; Mann et al., 1991; Wang and Becker, 2009), which continues to be a source of surprises today (Kabalah-Amitai et al., 2013). Vaterite has a major hexagonal structure (von Olshausen, 1925) and appears in different morphologies such as spherulitic aggregates (Han et al., 2006; Nebel and Epple, 2008; Mori et al., 2009; Hu et al., 2012) or hexagonal plates (e.g. Johnston et al., 1916; Kamhi, 1963; Dupont et al., 1997; Xu et al., 2006; Kawano et al., 2009).

Vaterite occurrence in nature is more widespread than generally assumed. It was first observed in gastropods (Mayer and Weineck, 1932), but later studies also discovered vaterite to be related to oil field drilling (Friedman and Schultz, 1994) and to be found in Portland cement (Friedman and Schultz, 1994) and as stones in the urinary system (Prien and Frondel, 1947; Sutor and Wooley, 1968). Vaterite has not been found in the geologic record and is therefore suspected to be metastable. The observation of vaterite in biogenic systems (Mayer and Weineck, 1932; Spann et al., 2010; Nehrke et al., 2012; Kabalah-Amitai et al., 2013) gives some constraints on
its stability, which can be on the order of years (Lowenstamm and Abbott, 1975), but not geological ages.

The natural occurrence of vaterite and its potential economic use due to its large specific surfaces and high porosity (Mori et al., 2009) warrants a precise investigation of this mineral. So far, most laboratory experiments have been designed to precipitate relatively large single crystals of vaterite for X-ray analysis that focused on the crystal structure (e.g. Kamhi, 1963). Vaterite precipitation experiments generally used either mixtures of several solutions such as K$_2$CO$_3$, Na$_2$CO$_3$ and CaCl$_2$ (Kamhi, 1963; Easton and Claugher, 1986; Han et al., 2006; Nebel and Epple, 2008) or CaNO$_3$ (Davies et al., 1978), sometimes with added surfactants (Mann et al., 1991; Dupont et al., 1997; Mori et al., 2009), or additional organic substrates (Falini et al., 1996; Xu et al., 2006; Kirboga and Oner, 2013). These experiments provided crystals with sizes between a few hundred nanometres and a few micrometres and were mostly restricted to the temperature range of 25–60°C.

As vaterite is a relevant mineral in biogenic systems (Lowenstamm and Abbott, 1975; Pouget et al., 2009; Spann et al., 2010; Nehrke et al., 2012) it may provide new insights into isotope fractionation during biological carbonate formation. Vaterite has been proposed as a potential precursor for high-purity CaCO$_3$ precipitation, later transforming into aragonite or calcite (e.g. Jacob et al., 2008). Depending on conditions prevailing during the vaterite–aragonite or vaterite–calcite transformation, the original isotope signature may be inherited in the final, stable CaCO$_3$ phase. In order to be able to use isotopic tools (for instance, to reconstruct environmental conditions or palaeoclimates) on biogenic carbonate with vaterite precursors or remaining vateritic components, knowledge of the fractionation factor between vaterite and water and the clumped isotope $\Delta^{47}_{\text{C}}$–$T$ relationship of vaterite is required. However, so far little is known about the oxygen isotope fractionation between dissolved inorganic carbon and vaterite and, in particular, the clumped isotope $\Delta^{47}_{\text{C}}$–$T$ relationship of vaterite. Whereas vaterite was reported in a few stable isotopes studies aiming at determining the oxygen isotope fractionation factor in the system CaCO$_3$–H$_2$O (e.g. Kim and O’Neil, 1997), it rarely occurred as a pure phase and thus did not allow for a precise study focused on vaterite. Based on the limited data available, Tarutani et al. (1969) suggested vaterite to be enriched in 18O by +0.5‰ relative to calcite. Kim and O’Neil (1997) obtained a similar value of +0.6‰ with respect to calcite. Both studies were limited to either one (25°C) or two temperatures (25, 40°C), and a more comprehensive study is still lacking. In addition, the clumped isotope $\Delta^{47}_{\text{C}}$–$T$ relationship of vaterite has not been assessed so far, but could give new insights into the effect of polymorphism on isotope ratios or mineral growth-related isotope fractionation.

In this study we present a simple method that allows vaterite precipitation over a wide temperature range (at least between 20 and 91°C) and that provides large quantities of the mineral, enabling for example the investigation of the oxygen isotope fractionation factor between vaterite and H$_2$O, and the $\Delta^{47}_{\text{C}}$–$T$ relationship of vaterite.

### 2 Experimental setup

The precipitating solution was prepared by dissolving high-purity CaCO$_3$ (Merck Suprapur, 99.95 %) in de-ionized water. Light microscopy indicates that this CaCO$_3$ consists of calcite. The water was initially taken from the local water supply, purified with a reverse osmosis technique and finally de-ionized with an Ultrapure™ system with an output quality of 18 MΩ cm. Trace components of the CaCO$_3$ used to prepare the solution are strontium (<100 ppm), rubidium (<20 ppm), sulfate (<20 ppm), and phosphate (<10 ppm).

About 370 mg of CaCO$_3$ was dissolved in ~500 mL de-ionized water at room temperature per experiment. The water was acidified by the purging of CO$_2$ tank gas (normal grade, BOC UK) through the solution. The solution was filtered after 3 h through a double layer of Whatman® filter paper (grade 1, corresponding to 11 µm filtration size) to remove undissolved CaCO$_3$ crystals. Optical inspection of the filtered water via light reflectance confirmed no large crystals to have bypassed the filtration stage. The dissolution step appeared to be quantitative as no CaCO$_3$ was visibly retained on the filter paper.

The filtered CaCO$_3$ solution was then thermally and isothermally equilibrated at a set temperature in a temperature-controlled water bath (a sketch of the experimental setup is depicted in Fig. 1). The experimental temperatures ranged from 23 to 91°C. The solution was enclosed in a 500 mL Erlenmeyer flask with a rubber stopper. The rubber stopper contained two feed-throughs for tubes that were used to maintain a constant gas flow through the solution. Humidified and thermally equilibrated CO$_2$ gas (same temperature as the solution) was passed through the CaCO$_3$ solution (Table 1) at a rate of ~0.03–0.1 mL s$^{-1}$ to prevent carbonate precipitation before complete isotopic equilibrium was achieved. The CO$_2$ gas was humidified and adjusted to the experimental temperature by bubbling it slowly through an Erlenmeyer flask filled with de-ionized water and contained in the temperature-controlled water bath. The equilibration period varied between 3 h at 91°C and 23 h at 23°C. pH values during equilibration are below pH 6 due to the continuous CO$_2$ flux.

After equilibration, NaCl was added, reaching concentrations between 4.2 and 6.4 mol L$^{-1}$ (Table 1). The added NaCl (Sigma Aldrich®) has a purity of ≥ 99% and contains minor traces of sulfates (≤ 200 ppm), alkaline earth metals (≤ 100 ppm) and bromides (≤ 100 ppm). Carbonate precipitation was induced by slowly bubbling N$_2$ tank gas (BOC UK, normal grade) through the solution. The N$_2$ gas was humidified and adjusted to the experiment temperature using the same procedure as for the CO$_2$ gas. The bubbling rate
was set to about 1 bubble per second (∼0.03 mL s⁻¹). The gas stream was humidified to prevent water evaporation and a potential change in the solution δ¹⁸O value over time. Minerals always formed on the bottom or the side wall of the Erlenmeyer flask. No crystals were observed on the surface of the solution. After 2–19 days, the solution was passed through a double layer of Whatman® filter paper (grade 1 with a pore size of 11 µm). Crystals on the glass walls were loosened by a thin PVC plastic tube and flushed out with de-ionized water. The precipitated minerals were air-dried at room temperature before microscopic and XRD (X-ray diffraction) analysis was conducted.

### Table 1. Experimental conditions during laboratory precipitation of CaCO₃ (see Sect. 2).

<table>
<thead>
<tr>
<th>Experiment no.</th>
<th>T (°C)</th>
<th>CaCO₃, dissolved (g L⁻¹)</th>
<th>NaCl added (g L⁻¹)</th>
<th>Equilibration (h)</th>
<th>Precipitation (h)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NA-1</td>
<td>23.5 ± 0.5</td>
<td>0.68</td>
<td>250</td>
<td>23</td>
<td>451</td>
</tr>
<tr>
<td>NA-3*</td>
<td>37.5 ± 0.5</td>
<td>0.70</td>
<td>260</td>
<td>21</td>
<td>72</td>
</tr>
<tr>
<td>NA-4</td>
<td>37.5 ± 0.5</td>
<td>0.74</td>
<td>244</td>
<td>14</td>
<td>341</td>
</tr>
<tr>
<td>NA-5</td>
<td>49.6 ± 0.5</td>
<td>0.70</td>
<td>375</td>
<td>16</td>
<td>143</td>
</tr>
<tr>
<td>NA-6</td>
<td>49.6 ± 0.5</td>
<td>0.80</td>
<td>262</td>
<td>17</td>
<td>573</td>
</tr>
<tr>
<td>NA-7</td>
<td>69.9 ± 0.5</td>
<td>0.70</td>
<td>325</td>
<td>3</td>
<td>69</td>
</tr>
<tr>
<td>NA-8</td>
<td>79.9 ± 0.5</td>
<td>0.78</td>
<td>280</td>
<td>3</td>
<td>47</td>
</tr>
<tr>
<td>NA-9</td>
<td>91.0 ± 0.5</td>
<td>0.70</td>
<td>260</td>
<td>3</td>
<td>42</td>
</tr>
</tbody>
</table>

* NA-2 differed in the experimental conditions and is therefore omitted.

### 3 Samples

#### 3.1 Mineral description, microscopy and SEM

Depending on the experiment, temperature and duration, between 5 and 235 mg CaCO₃ was precipitated (Tables 1, 2). Low carbonate recovery < 20 mg is linked to short experiment duration (3–6 days) at lower temperatures (< 70 °C, Table 2). In contrast, experiments with a longer duration of 14–24 days at temperatures ≤ 50 °C yielded on average 135 mg. At 80–91 °C it was sufficient to allow 2 days for mineral precipitation to obtain 80–90 mg of calcite. Note that in all experiments the initially dissolved amount of calcite was similar at about 360 (±20) mg in 500 mL of de-ionized water (Table 1).
Table 2. Mineralogical and isotopic results of the vaterite precipitation experiments. The mineralogy was determined by XRD analysis (Sect. 3.2). The \( n \) values give the number of replicates measured for isotopic analysis. The \( \Delta_{47} \) value is given in the absolute reference frame of Dennis et al. (2011) and is corrected for the acid digestion reaction.

<table>
<thead>
<tr>
<th>Experiment no.</th>
<th>( T ) (°C)</th>
<th>( \text{CaCO}_3 ) precipitated (mg)</th>
<th>( \delta^{18} \text{O} ) (‰)</th>
<th>( \delta^{13} \text{C} ) (‰)</th>
<th>( \Delta_{47} ) (‰)</th>
<th>( n )</th>
<th>Mineralogy</th>
</tr>
</thead>
<tbody>
<tr>
<td>NA-1</td>
<td>23.5 ± 0.5</td>
<td>120</td>
<td>−8.57 ± 0.16</td>
<td>−18.21 ± 0.06</td>
<td>0.689 ± 0.003</td>
<td>3</td>
<td>vaterite</td>
</tr>
<tr>
<td>NA-3*</td>
<td>37.5 ± 0.5</td>
<td>5</td>
<td>−11.29 ± 0.20</td>
<td>−20.39 ± 0.10</td>
<td>0.639 ± 0.020</td>
<td>1</td>
<td>vaterite (95 %), calcite (5 %)</td>
</tr>
<tr>
<td>NA-4</td>
<td>37.5 ± 0.5</td>
<td>50</td>
<td>−13.30 ± 0.37</td>
<td>−26.06 ± 0.18</td>
<td>0.672 ± 0.027</td>
<td>3</td>
<td>vaterite (&gt; 95 %), rest: calcite</td>
</tr>
<tr>
<td>NA-5</td>
<td>49.6 ± 0.5</td>
<td>15</td>
<td>−13.85 ± 0.26</td>
<td>−21.39 ± 0.03</td>
<td>0.630 ± 0.005</td>
<td>2</td>
<td>vaterite</td>
</tr>
<tr>
<td>NA-6</td>
<td>49.6 ± 0.5</td>
<td>235</td>
<td>−15.06 ± 0.22</td>
<td>−25.26 ± 0.17</td>
<td>0.634 ± 0.008</td>
<td>3</td>
<td>vaterite</td>
</tr>
<tr>
<td>NA-7</td>
<td>69.9 ± 0.5</td>
<td>15</td>
<td>−16.92 ± 0.15</td>
<td>−21.71 ± 0.03</td>
<td>0.577 ± 0.010</td>
<td>3</td>
<td>vaterite</td>
</tr>
<tr>
<td>NA-8</td>
<td>79.9 ± 0.5</td>
<td>80</td>
<td>−17.54 ± 0.03</td>
<td>−25.86 ± 0.10</td>
<td>0.553 ± 0.018</td>
<td>3</td>
<td>calcite (49 %), aragonite (24 %), vaterite (27 %)</td>
</tr>
<tr>
<td>NA-9</td>
<td>91.0 ± 0.5</td>
<td>90</td>
<td>−19.21 ± 0.15</td>
<td>−25.00 ± 0.16</td>
<td>0.545 ± 0.005</td>
<td>5</td>
<td>vaterite (94 %), aragonite (6 %), calcite (&lt; 1 %)</td>
</tr>
</tbody>
</table>

* NA-2 differed in the experimental conditions and is therefore omitted.

Figure 2. Photomicrographs of \( \text{CaCO}_3 \) minerals precipitated in the laboratory experiment. Scale bar is 100 µm in (a)-(c) and 200 µm in (d). Vaterite crystals formed at 50 °C in experiment NA-6 (a), at 70 °C (b) and at 91 °C (d). At 80 °C a mixture of aragonite, calcite and vaterite was precipitated (e).

Figure 3. Photomicrographs of \( \text{CaCO}_3 \) minerals precipitated in the laboratory experiments. Scale bar is 100 µm. Vaterite crystals formed at 23 °C (a), at 37 °C in experiment NA-3 (b) and NA-4 (c), and at 50 °C in NA-5 (d).

Vaterite can be distinguished from other \( \text{CaCO}_3 \) polymorphs by its morphology. Calcite rhombohedra and aragonite needles can be recognized by light microscopy (e.g. Fig. 2c). Vaterite crystals can be similar in size but are more irregular and show a spherulitic shape (Figs. 3, 4). Inspection of large vaterite crystals under normal and polarized light reveals a complex growth history. Various globular segments of 50–100 µm with an internal spherulitic growth pattern coalesce into one larger crystal (Fig. 4). Vaterite crystals showed a typical size of 50 µm (Figs. 2, 3, 4), whereas in a few experiments, crystals of up to 500 µm were observed. Experiments at 70 and 91 °C also resulted in vaterite crystals in the 50 µm size range; however, these are composed of many small (∼10 µm) globular sub-segments. A peculiarity of vaterite crystals precipitated at 23 °C is the combination of rounded, spherical shapes with sharp angular forms (Fig. 3a). Together with the larger crystals sizes observed at this temperature; this points towards slower mineral growth. All minerals of this study were investigated with XRD in ad-
dition to optical microscopy. Note that optical microscopy alone may be ambiguous and should be complemented by additional methods (e.g. XRD).

Scanning electron microscope (SEM) images were made at the Institute of Earth Sciences at Heidelberg University to investigate the morphology in more detail. The scanning electron microscope LEO 440 was used for imaging. It has a tungsten cathode, was operated at an accelerating voltage of 20 kV, and enables a minimal resolution of ca. 5 nm. Samples were sputtered with a thin gold layer for imaging and with carbon for elemental analysis. A summary with characteristic vaterite aggregates is shown in Fig. 5. The size of individual grains that make up the vaterite aggregates was investigated using SEM images (Figs. 5, 6). The grain size decreases with increasing temperature, from about 100 µm at 23°C, 10 to 20 µm at 50°C, to < 10 µm at 91°C. The relatively large grains detected at 23°C are internally composed of ~2 µm long elongated fibrous crystallites (Fig. 6a). The smallest grains observed at 91°C are of a similar size of a few micrometres. Sub-micrometre-sized crystallites and frambooidal aggregates (e.g. Nehrke and Van Cappellen, 2006) were not observed. The minerals show a radial growth pattern from a central nucleus leading to spherical conglomerate particles (Fig. 5a). The spherical shape is still dominant at 50°C (Fig. 5b); however, it changes to the growth of flat platelets at 91°C (Fig. 5c). The radiating growth pattern at 91°C is restricted to two dimensions with a tree-like branching structure characteristic for diffusion-controlled dendritic crystalization (Fig. 5d).

Additional elemental analyses on carbon-sputtered vaterite grains using energy-dispersive X-ray spectroscopy (EDS) at the SEM system at Heidelberg University revealed minor traces of sodium and chloride to be occasionally incorporated in the vaterite mineral. Quantitative analysis was not attempted due to the very sporadic and dispersed occurrence of the sodium chloride crystals.

3.2 XRD analysis

The carbonate samples were analysed at the National History Museum, London, using an Enraf Nonius FR 590 powder diffractometer with Cu-Kα radiation (40 kV, 35 mA). In brief, the sample powder was placed in a thin layer on a sapphire substrate and measured by fixed beam-sample-detector geometry. Analysis times were adjusted to the counting statistics and varied between 10 and 90 min. Signals and phase fractions were evaluated by comparing measured spectra with a mineral database using the program X’Pert Highscore (PANalytical B.V., 2009). Peak positions were calibrated with two standards (silver behenate and quartz). For phase quantification, a pure calcite standard and an aragonite standard were additionally measured.

For most samples, the dominant XRD peaks were found at 20.98(±0.04), 24.86(±0.02), 27.03(±0.03), 32.74(±0.03), 43.79(±0.09), and 50.0(±0.04)° (2θ, Fig. 7). In contrast, the characteristic and dominant calcite peak of the calcite standard is observed at 29.46°; those of the aragonite calibration standard are at 26.36, 27.35, 33.25 and 46.01° (2θ, Fig. 8). The aragonite peaks at 26.36 and 27.35° were resolved for the pure aragonite standard, but the peak at 27.35° overlapped with the vaterite peak at ~27.03° for aragonite–vaterite mixtures. Our laboratory CaCO₃ samples are clearly different from aragonite and calcite, but coincide with the XRD data and d-spacing of vaterite. Kabalah-Amitai et al. (2013) measured vaterite d-spacing of 2.07 and 3.63 Å, corresponding to 43.69 and 24.50° (2θ at Cu-Kα radiation). Earlier work of Dupont et al. (1997) determined similar d-spacing values of 4.254, 3.591, 3.307, 2.741, 2.07 and 1.826 Å, corresponding to 20.86, 24.77, 26.94, 32.64, 43.69 and 49.90° (2θ at Cu-Kα radiation). Our own results are
close to these values confirming the precipitates to be composed of vaterite. In the case of the 37 °C experiment, calcite is additionally present as a minor phase (about 5 %, visible in the peak at 29.46°, Fig. 7), whereas only traces of calcite are found in the 91 °C experiment. An exact quantification of this small calcite fraction was not possible due to a general uncertainty of about 3 % in the phase quantification. Half of the mineral phase at 80 °C consists of calcite, whereas the other half is made up of equal proportions of aragonite and vaterite (Table 2, Fig. 8). Aragonite is also a minor phase in the 91 °C experiment (6 %, Fig. 7).

3.3 Isotope analysis

Oxygen, carbon and clumped isotopes were analysed at the Qatar Stable Isotope Laboratory at Imperial College. Details of the sample preparation and mass spectrometric procedures are given in Kluge et al. (2015). In brief, per analysis ~5 mg sample was dissolved in orthophosphoric acid at 70 °C to produce CO₂ for the mass spectrometric measurement. The CO₂ is cleaned manually, comprising a step for cryogenic water separation and one for contaminant removal via porous polymers (Porapak™ Q). Analyses were done using two dual-inlet isotope ratio mass spectrometers (Thermo Scientific MAT 253) that measure sample and reference gas alternately. Individual analyses have a precision of 0.2 ‰ for δ¹⁸O, 0.1 ‰ for δ¹³C, and 0.03 ‰ for Δ₄⁷, based on replicate analyses of standards. Samples were measured repeatedly (typically three times) to reduce the uncertainty.

Carbonate δ¹⁸O values follow the trend determined by the temperature-dependent isotopic fractionation between water and calcite (Fig. 9). For calculation of the expected carbonate δ¹⁸O values, we used the fractionation factors of Kim and O’Neil (1997) and a water value of −6.4 ± 0.7 ‰ (−7.8 ± 0.5 ‰ for the later repeat experiments NA-4 and NA-6). The water δ¹⁸O value was not directly measured but is based on back-calculation of aragonite and calcite samples that were precipitated in close temporal connection (within 3–12 days) of the vaterite experiments using the same water source and the same precipitation technique (see Kluge et al., 2015). The back-calculated water values from the pure Ca(HCO₃)₂ solution, used as a reference, correspond to the observed range of surface and ground water values of the London metropolitan area (Darling, 2003). For all experiments, the water was taken from the local water supply. Although we have a defined water reference value based on the pure Ca(HCO₃)₂ solution, we cannot exclude short-term fluctuations of the tap water δ¹⁸O. However, it is unlikely that it exceeds the observed long-term variability of ±0.7 ‰ (June–October 2012) and ±0.5 ‰ (March–April 2013) for the back-calculated solution δ¹⁸O value.

The δ¹³C values vary between −18 and −26 ‰ and reflect the negative signature of the CO₂ tank gas used during the equilibration phase (Table 2). The δ¹³C values do not show a temperature dependence. Clumped isotope Δ₄⁷ values of vaterite samples decrease with increasing temperature and are similar to calcite or aragonite–calcite mixtures precipitated at the same temperature (Fig. 10). Δ₄⁷ values of calcite and calcite–aragonite mixtures were taken from Kluge et al. (2015) for comparison.
4 Discussion

Vaterite was obtained over the entire experimental temperature range of 23–91 °C. It is detected either as the only phase (23, 50, 70 °C) or as the major phase (≥ 94 %) with minor contributions from calcite or aragonite (37, 91 °C). An exception is the experiment at 80 °C where all anhydrous CaCO₃ polymorphs were precipitated simultaneously. On average 80 mg of vaterite was formed per experiment. This amount may be increased by longer experiment runs or by up-scaling of the setup using larger beakers with the same solution concentrations. A longer experiment duration appears to be the most effective approach. Considering the experiments from 23 to 70 °C only, the yield increases exponentially with the duration, reaching a recovery rate (relative to the initially dissolved CaCO₃) of 70 % after ∼ 570 h (Fig. 11). An extrapolation of this trend is not easily possible as the precipitation rate will eventually decrease due to the continuously decreasing supersaturation. Another option of increasing the CaCO₃ supersaturation in the initial solution was not tested, but has to be treated carefully. A higher initial chemical potential may produce a higher yield but also lead to the precipitation of other forms of CaCO₃ such as ikaite (calcium carbonate hexahydrate, CaCO₃(H₂O)₆) or amorphous calcium carbonate (Kawano et al., 2009).

This study shows that vaterite precipitation is not limited to a certain temperature range, e.g. to room temperature or from 10 to 48 °C (Gussone et al., 2011), but can be performed at least up to 91 °C. A pressurized reaction vessel that prevents boiling of the solution could be used to extend vaterite mineral formation to much higher temperatures (e.g. Kluge et al., 2015). A thermally and isotopically equilibrated CaCO₃ supersaturated solution could be injected into the thermally equilibrated and saturated NaCl solution of a pressurized reaction vessel.

The detection of vaterite minerals over the large temperature interval of this study and its predominating character is surprising, given that many other studies emphasized the low stability of vaterite (e.g. McConnel, 1959). McConnel (1959) states that vaterite dissolves at room temperature at contact with water. However, our precipitates were air-dried at room temperature on Whatman® filter paper and stayed wet for a few hours, but did not transform into calcite. Furthermore, vaterite minerals were stored between several weeks and a year before being analysed by XRD and SEM. Despite long storage periods vaterite did not transform into other CaCO₃ polymorphs; this implies that vaterite can be precipitated and stored for periods that are long enough to enable precise and detailed experimental analyses. Independent evidence for the stability of vaterite over years comes from biogenic samples such as bivalves, mollusks and other marine organisms (Lowenstamm and Abbott, 1975; Spann et al., 2010; Nehrke et al., 2012).

Isotopic analysis of vaterite

Our study presents a well-defined method for the isotopic study of vaterite. The technique for vaterite precipitation was modified from the procedures of McCrea (1950), O’Neil et al. (1969) and Kim and O’Neil (1997) that were developed for isotopic studies on calcite and aragonite minerals. The
values of Kim and O’Neil (1997) are commonly used as a reference for the oxygen isotope fractionation between calcite and water. Furthermore, for the temperature calibration of clumped isotopes, an analogous method was applied to precipitate calcite (Ghosh et al., 2006; Zaarur et al., 2013). The closeness of the methodology for synthetic vaterite and calcite precipitation ensures good comparability of the obtained results with widely used calibration and fractionation data.

The long equilibration procedure used in our experimental approach (Table 1) enables isotopic equilibration between the dissolved inorganic carbon (DIC) and water and among the DIC species. A 99% equilibrium between oxygen isotopes in water and DIC takes about 9 h at 25°C and a pH of ~8, whereas it is less than 2 h at temperatures above 40°C (Beck et al., 2005). For comparison, the equilibration duration was 23 h at 23°C, 14–21 h at 37.5°C, and 3–17 h above 40°C (Table 1). This provides the necessary basis for a meaningful isotopic analysis of the precipitated vaterite which has not been attempted in a systematic manner so far.

Beyond the isotopic equilibration of the DIC with water, the precipitation rate and the ionic concentration of the solutions can affect isotope values. The growth rate has to be considered in the interpretation of isotope values as differences can be substantial between slow and rapidly grown minerals (up to 2‰ for Δ18O; Coplen, 2007; Dietzel et al., 2009; Gabitov et al., 2012). The growth rate in our study was not precisely monitored, but can be estimated from the derivative of the general relationship between experiment duration and mineral yield (Fig. 11). The mineral growth for temperatures below 80°C started very slowly with a value of ~1.4 × 10^-10 mol s^-1 and increased exponentially (e.g. to 1.1 × 10^-9 mol s^-1 after 350 h). Converting it relative to the growth surface (using the walls of the Erlenmeyer flask as a first-order estimate), the growth rate was about 3.7 × 10^-9 mol (m² s)^-1 at the beginning of the mineral formation and increased to 2.9 × 10^-8 mol (m² s)^-1 at 350 h. These values only give an impression of the order of magnitude due to the limitation of the assumptions, but already demonstrate that the mineral formation was unlikely affected by rapid growth disequilibrium that is typically encountered at values above 10^-7–10^-8 mol (m² s)^-1 (compare with, for example, Watkins et al., 2013). In contrast, in other vaterite precipitation techniques, two solutions were mixed, leading to almost instantaneous precipitation (e.g. Nebel and Eppe, 2008). As rapid mineral growth may induce disequilibrium fractionation related to a mineral surface effect (Watson, 2004; Dietzel et al., 2009; Watson and Müller, 2009; DePaolo, 2011; Reynard et al., 2011; Gabitov et al., 2012; Gabitov, 2013), experiments with quasi-instantaneous mineral growth are not suitable for isotope studies. Consequently, we neither rapidly grew vaterite nor conducted isotope measurements on potentially fast growth phases.

Traditionally, vaterite was synthesized from mixtures of CaCl₂, K₂CO₃ (Kamhi, 1963) and admixtures of calgon (McConnell, 1960), or included other surfactants (Mori et al., 2009). In other experiments, a CaCO₃ supersaturated solution was treated with surfactants (Dupont et al., 1997) or polymeric substances (Kirboga and Oner, 2008). In a few experiments, Na₂CO₃ replaced K₂CO₃ as the solution containing the carbonate ion (Nebel and Eppe, 2008). The use of CaCl₂ and especially K₂CO₃ could impact on the isotopic values of the forming minerals via preferential fractionation related to the hydration sphere of the Ca²⁺ and K⁺ ions (Taube, 1954; Sofer and Gat, 1972; O’Neil and Truesdell, 1972; Taube, 1954).
and thus should either be restricted to low concentrations or avoided. Our method uses only NaCl as an additive that has been confirmed not to affect the isotope values of the DIC (e.g. O’Neil and Truesdell, 1991).

Before discussing the measured vaterite $\delta^{18}O$ values and its implication for the oxygen isotope fractionation factor $\alpha_{\text{CaCO}_3\text{-H}_2\text{O}}$, we note that we did not analyse the oxygen isotope composition of the solution per se. However, as the solution water is ultimately taken from the local water supply that reflects the London Metropolitan ground- and surface water $\delta^{18}O$ of $-6$ to $-8\permil$ (Darling, 2003) and which is cross-examined using values from independent NaCl-free experiments carried out in parallel, we have a defined reference water $\delta^{18}O$ value ($-6.4 \pm 0.7, -7.8 \pm 0.5\permil$ for the later repeat experiments NA-4 and NA-6; see Sect. 3.3). Thus, our results give a first-order guideline with respect to the temperature dependence of $\alpha_{\text{CaCO}_3\text{-H}_2\text{O}}$ and can provide an upper limit for the deviation of the fractionation factor of vaterite compared to aragonite and calcite. We compare our values to data of Kim and O’Neil (1997) due to similar experimental procedures and their universal use. Note that the exact value of the equilibrium oxygen isotope fractionation factor for calcite is under debate (Coplen, 2007; Dietzel et al., 2009; Gabitov et al., 2012; Kluge et al., 2014).

The fractionation $1000 \ln \alpha_{\text{CaCO}_3\text{-H}_2\text{O}}$ of vaterite closely follows the values of Kim and O’Neil (1997) for calcite (Fig. 9). Over the entire experimental temperature range, vaterite values agree within the calcite errors determined by analysing multiple samples and calibration standards. Vaterite was successfully synthesized from a NaCl-saturated CaCO$_3$ solution. Vaterite formed between 23 and 91°C and was generally pure or the major phase. The precipitation technique allows thermal and isotopic equilibration of the oxygen and carbon isotopes in the solution. The precipitation technique fosters slow mineral formation, which enables a meaningful isotopic analysis of the precipitated vaterite. Recovered vaterite amounts of up to 235 mg also permit the precise determination of the clumped isotope value at each experiment temperature, which requires a relatively large sample aliquot of 5 mg per replicate measurement.

The oxygen isotope fractionation between water and the vaterite mineral agrees within uncertainty with calcite (average difference $0.0 \pm 0.4\permil$). Clumped isotope $\Delta_{47}$ values that are indistinguishable from calibration data (difference of $-0.003 \pm 0.013\permil$). The presented precipitation technique for vaterite, in particular the possibility for thermal and isotopic equilibration, opens research opportunities for precise investigation of isotope ratios on this unexplored CaCO$_3$ polymorph.

Equation (1) is given in the absolute reference frame of Dennis et al. (2011). Vaterite $\Delta_{47}$ values scatter around the $\Delta_{47}-T$ line of Eq. (1) with an average difference of $-0.003 \pm 0.013\permil$ and are thus indistinguishable from the calibration line (Fig. 10). Subtle differences in the mineral structure of the CaCO$_3$ polymorph vaterite appear to be irrelevant for the $^{13}$C$-^{18}$O clumping.

### 5 Conclusions

Vaterite was successfully synthesized from a NaCl-saturated CaCO$_3$ solution. Vaterite formed between 23 and 91°C and was generally pure or the major phase. The precipitation technique allows thermal and isotopic equilibration of the oxygen and clumped isotopes in the solution. The precipitation technique fosters slow mineral formation, which enables a meaningful isotopic analysis of the precipitated vaterite. Recovered vaterite amounts of up to 235 mg also permit the precise determination of the clumped isotope value at each experiment temperature, which requires a relatively large sample aliquot of 5 mg per replicate measurement.

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### Appendix A

$\delta^{18}O$, $\delta^{13}C$ and $\Delta_{47}$ values of samples and calibration standards are provided in the supplementary data file, together with raw data and transfer functions.

The Supplement related to this article is available online at doi:10.5194/bg-12-3289-2015-supplement.

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References


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