Influence of changing carbonate chemistry on morphology and weight of coccoliths formed by *Emiliania huxleyi*

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Received: 6 March 2012 – Accepted: 2 May 2012 – Published: 23 May 2012

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Published by Copernicus Publications on behalf of the European Geosciences Union.
Abstract

The coccolithophore *Emiliania huxleyi* is a marine phytoplankton species capable of forming small calcium carbonate scales (coccoliths) which cover the organic part of the cell. Calcification rates of *E. huxleyi* are known to be sensitive to changes in seawater carbonate chemistry. It is, however, not yet understood how these changes are reflected in the morphology of coccoliths. Here, we compare data on coccolith size, weight, and malformation from a set of five experiments with a large diversity of carbonate chemistry conditions. This diversity allows distinguishing the influence of individual carbonate chemistry parameters such as carbon dioxide (CO$_2$), bicarbonate (HCO$_3^-$), carbonate (CO$_3^{2-}$), and protons (H$^+$) on the measured parameters. Measurements of fine-scale morphological structures reveal an increase of coccolith malformation with decreasing pH suggesting that H$^+$ is the major factor causing malformations. Coccolith distal shield area varies from about 5 to 11 µm$^2$. Changes in size seem to be mainly induced by varying [HCO$_3^-$] and [H$^+$] although influence of [CO$_3^{2-}$] cannot be entirely ruled out. Changes in coccolith weight were proportional to changes in size. Increasing CaCO$_3$ production rates are reflected in an increase in coccolith weight and an increase of the number of coccoliths formed per unit time. The combined investigation of morphological features and coccolith production rates presented in this study may help to interpret data derived from sediment cores, where coccolith morphology is used to reconstruct calcification rates in the water column.

1 Introduction

Coccolithophores are unicellular photoautotrophic organisms, able to form blooms in all major ocean basins (Moore et al., 2012). Their unique feature is the intracellular formation of small scales (coccoliths) made of calcium carbonate (CaCO$_3$) covering the organic surface of the cell. Coccolithophores appeared for the first time about 220 million years ago in the fossil record and are found ever since in marine sediments
although their abundance was highly variable (Bown et al., 2004). They are important components in the marine carbon cycle because the CaCO$_3$ in their coccoliths serves as ballasting material accelerating the organic carbon flux from the surface into the deep ocean (e.g., Honjo, 1976; Ploug et al., 2008). In modern oceans, *Emiliania huxleyi* is the most abundant species (Paasche, 2002). It evolved from *Gephyrocapsa oceanica* about 270 ky ago and dominates the coccolithophore community for the last ∼70 ky (Thierstein et al., 1977). *E. huxleyi* frequently forms large blooms that can cover up to a million km$^2$ and can be seen from space (Holligan et al., 1993; Tyrell and Young, 2009). These blooms are typically found in stratified waters during later stages of the phytoplankton spring succession (Tyrell and Merico, 2004), although, recently, high *E. huxleyi* abundance has also been reported in turbulent regimes when cell numbers are integrated over the whole water column (Schiebel et al., 2011).

The invasion of anthropogenic CO$_2$ into the ocean currently changes the marine carbonate chemistry by increasing [CO$_2$] and decreasing seawater pH – a process known as ocean acidification (Raven et al., 2005). Although these changes in carbonate chemistry are known to influence calcification rates of *E. huxleyi* (Riebesell and Tortell, 2011), it is still not understood how changing calcification rates are reflected in coccolith size, weight, and exocytosis rate (i.e. the number of coccoliths formed and egested per day). Such knowledge is, however, urgently needed in case morphometric data originating for example from sediment cores is used to reconstruct calcification rates within the water column. The influence of changing carbonate chemistry on the appearance of coccolith malformations in *E. huxleyi* is better understood (Langer et al., 2010, 2011) but key questions such as for example which carbonate chemistry parameter is actually causing malformations are still unknown.

The study presented here aims to improve our understanding on the following three research questions. (1) Does morphology (size and weight) of *E. huxleyi* coccoliths change in response to changing carbonate chemistry conditions? (2) Are potential changes in morphology reflected in calcium carbonate production rates? (3) Which particular carbonate chemistry parameter(s) drive potential changes in *E. huxleyi* coccolith
morphology? In order to address these questions, we evaluated samples for *E. huxleyi* coccolith size, weight and malformation from five culture experiments with a large diversity of carbonate chemistry conditions. This diversity allowed us to disentangle the carbonate system and assess which of the carbonate system parameters that can directly influence the cell physiology (i.e. CO$_2$, HCO$_3^-$, CO$_3^{2-}$ and H$^+$) are responsible for possible changes in the morphology of coccoliths formed by *E. huxleyi*.

2 Material and methods

2.1 Basic experimental settings

Five experiments were conducted with monospecific cultures of the coccolithophore *Emiliania huxleyi* strain PML B92/11. All experiments are generally similar in their design. Differences between them are exclusively manifested in the carbonate chemistry parameters of the culture medium (see Sect. 2.2).

All experiments were conducted with dilute batch cultures (LaRoche et al., 2010) at 15°C and 150 µmol photons m$^{-2}$ s$^{-1}$ incident photon flux density in a 16/8 light/dark cycle. The growth medium was artificial seawater, prepared as described in Kester et al. (1967) but without the addition of NaHCO$_3$. The artificial seawater medium (free of dissolved inorganic carbon (DIC) and total alkalinity (TA)) was enriched with $\sim$ 64 µmol kg$^{-1}$ nitrate, 4 µmol kg$^{-1}$ phosphate, f/8 concentrations for trace metals and vitamins (Guillard and Ryther, 1962), 2 ml of natural North Sea water and 10 nmol kg$^{-1}$ of SeO$_2$ to avoid nutrient limitation in the course of the experiments. Samples for nitrate and phosphate were 0.7 µm filtered at the beginning and the end of the experiments and measured according to Hansen and Koroleff (1999). The nutrient-enriched medium was sterile-filtered (0.2 µm) into sterile polycarbonate bottles where the carbonate system was adjusted (see following section). Samples for carbonate chemistry measurements ($\sim$ 500 ml) were taken from these bottles after adjustment. The remaining medium was gently transferred into sterile 2 l polycarbonate bottles. The headspace
in the 2l bottles was kept below 5 ml. The culture medium was acclimated to 15°C overnight to avoid a thermal shock when transferring the cells from the pre-cultures to bottles in which the main experiments were performed. Cells were acclimated to the carbonate chemistry conditions of the main experiment for at least 7 generations prior to inoculation.

2.2 Carbonate chemistry manipulations

The experiments only differed with respect to the carbonate chemistry manipulation of the culture medium. In the first experiment, cells were cultured at constant total alkalinity (2320 ± 22 µmol kg⁻¹) and varying fCO₂ levels, ranging from ~ 20 to ~ 5960 µatm. Here, DIC and TA levels were adjusted by adding calculated amounts of Na₂CO₃ and hydrochloric acid (3.571 mol l⁻¹, certified by Merck) (Gattuso et al., 2010). In the second, third and fourth experiment, pH was kept constant at pHᵢ (free scale) 7.74 (±0.004), 8 (±0.01), and 8.34 (±0.008), while fCO₂ was increased from ~ 100 to ~ 3600, from ~ 40 to 3650, and from ~ 21 to ~ 1163 µatm, respectively. Carbonate chemistry in the constant pH approaches was adjusted by adding 2 mmol kg⁻¹ of 2-[-4-(2-hydroxyethyl)-1-piperazinyl]-ethanesulfonic acid (HEPES) to the culture medium which was adjusted to the target pHᵢ levels. DIC was added as NaHCO₃. The small change in pH in the HEPES buffered seawater medium due to NaHCO₃ addition was compensated by adding small amounts of strong NaOH or HCl. In the fifth experiment, fCO₂ was kept constant (430 ± 47 µatm), while DIC ranged from ~ 500 to 4100 µmol kg⁻¹. DIC and fCO₂ were adjusted by adding calculated amounts of Na₂CO₃ and hydrochloric acid (Gattuso et al., 2010). For an overview of the carbonate chemistry in the experiments see Table 1. Note that other data from the constant TA and the constant pHᵢ 8 experiment can be found in Bach et al. (2011).
2.3 Carbonate chemistry sampling and measurements

Samples for TA measurements were filtered (0.7 µm), poisoned with a saturated HgCl$_2$ solution (0.5‰ final concentration) and stored at 4 °C until measurements. TA was measured in duplicate applying a two-stage potentiometric open cell titration (Dickson et al., 2003) and corrected with certified reference material (Prof. A. Dickson, La Jolla, CA). Some TA samples of the constant CO$_2$ experiment were higher than ∼4700 µmol kg$^{-1}$ and had to be diluted in order to get reliable results. Therefore, these TA samples were mixed with double de-ionised water, containing no alkalinity. The ratio of double de-ionised water relative to the TA sample was determined on a balance (Sartorius) with a precision of ±0.01 g.

Samples for DIC were sterile filtered (0.2 µm) with gentle pressure and stored bubble-free at 4 °C in 4 ml borosilicate bottles. All DIC samples were measured according to Stoll et al. (2001). In most treatments of the constant pH experiments, DIC was either too high or too low to be measured according to Stoll et al. (2001). To solve this problem, sample medium was mixed with artificial seawater of known DIC concentration. The ratio of the mixing solvent to the original DIC sample was determined by first weighing the mixing solvent alone and in a second step the mixing solvent plus the original DIC sample on a balance with a precision of ±0.01 mg (Sartorius). The mixture was carefully rotated in a 50 ml tube with ∼1 ml headspace. The ratio of sample to mixing solvent was adjusted to result in a final DIC concentration of approximately 1800–2200 µmol kg$^{-1}$. After the mixing procedure, DIC samples were processed identical to undiluted samples (see above).

Samples for pH$_f$ were measured potentiometrically at 15 °C with separate glass and reference electrodes (Metrohm) which were calibrated with reference material certified for TA and DIC with a salinity of 33.3 (Prof. A. Dickson, La Jolla, CA). pH$_f$ of the reference material was calculated from certified TA and DIC applying the constants of Roy et al. (1993). Measured electromotive force (E) of the samples and standards were
used to calculate the pH$_f$ of the sample as

$$\text{pH}_f = \text{pH}_{f_{\text{ref}}} + \frac{E_s - E_x}{R \times T \times \frac{\ln 10}{F}} \tag{1}$$

where, pH$_{f_{\text{ref}}}$ is the calculated pH$_f$ of the certified reference material, $T$ is the temperature of the sample in Kelvin, $R$ is the universal gas constant, $F$ the Faraday constant and $E_s$ and $E_x$ are the measured electromotive forces in Volts of the standard and the sample, respectively (Dickson et al., 2007).

### 2.4 DIC estimations

Unfortunately, we lost all DIC measurements of the constant TA experiment and the DIC measurements from the beginning of constant pH$_f$ 8 experiment due to storage problems. The estimation of DIC of these samples is shown in detail in Bach et al. (2011) and shall be outlined only briefly in the following. DIC concentrations from the beginning of the constant pH$_f$ 8 experiment were estimated by adding the total particulate carbon build-up which was produced during the experiment to the final DIC concentrations. Initial DIC concentrations from the constant TA experiment were calculated as:

$$\text{DIC} = \frac{\text{TA}_{\text{measured}} + (\text{Volume}_{\text{acid}} \times 3.571)}{2} \tag{2}$$

where TA$_{\text{measured}}$ is the measured TA in µmol kg$^{-1}$ at the beginning of the experiment, Volume$_{\text{acid}}$ is the volume of acid that was added in µl kg$^{-1}$ and 3.571 is the molarity of the acid (certified by Merck) in mol l$^{-1}$. This estimate has an uncertainty of approximately 40–50 µmol kg$^{-1}$ which is small compared to the large DIC range in this experiment (Bach et al., 2011). DIC concentrations at the end of the constant TA experiment were calculated by subtracting the measured total particulate carbon build-up from the initial DIC concentrations.
2.5 Carbonate chemistry calculations

Carbonate chemistry conditions within experiments were calculated from temperature, salinity, inorganic phosphate concentrations and two measured (or estimated) carbonate system parameters, applying the equilibrium constants of Roy et al. (1993) and the program CO2Sys (Lewis and Wallace, 1998). Measured (or estimated) carbonate system parameters were: TA and DIC in the constant TA experiment; pH\textscript{f} and DIC in the constant pH experiments; TA and pH\textscript{f} in the constant CO\textsubscript{2} experiment. The biological response data are plotted to the mean of initial and final carbonate chemistry conditions.

2.6 Sampling and calculation of coccolith exocytosis rate

Sampling started two hours after the onset of the light period and lasted no longer than two and a half hours. Two samples for particulate organic carbon (POC) and two for total particulate carbon (TPC) were filtered (200 mbar) onto precombusted (5 h, 500 °C) GF/F filters and stored in the dark at −20 °C. POC and TPC samples from constant pH experiments were rinsed with artificial seawater after filtration (supersaturated with respect to calcite) in order to wash off HEPES buffer which otherwise would have contributed ~ 40 pg of carbon to every TPC and POC measurement. POC samples were stored for two hours in a desiccator containing fuming HCl to remove all inorganic carbon and subsequently dried for ~ 6 h at 60 °C. TPC filters were dried in the same way as the POC filters, but in a separate oven and without prior acid treatment. Carbon concentrations of POC and TPC filters were measured using an isotope ratio mass spectrometer (Finnigan) combined with an elemental analyzer (EuroEA, Hekatech GmbH). Particulate inorganic carbon (PIC) was calculated as the difference between TPC and POC. Four POC samples from the constant pH\textscript{f} 8 experiment were lost during measurements.

Cell numbers were measured at the beginning and the end of the experiment with a Coulter Counter (Beckmann). The growth rate (\(\mu\)) was calculated from initial and final
cell numbers as

\[ \mu = \frac{\ln(t_{\text{fin}}) - \ln(t_0)}{d} \]  

where \( t_0 \) and \( t_{\text{fin}} \) is the cell number at beginning and the end of the experiment, respectively and \( d \) is the number of days the cell culture was growing. PIC production rates were calculated by multiplying \( \mu \) and PIC cell\(^{-1} \). CaCO\(_3\) production rates were calculated by multiplying PIC production rates (in \( \mu\text{mol C cell}^{-1} \text{d}^{-1} \)) with the molecular weight of CaCO\(_3\). The number of egested coccoliths per day was subsequently determined as

\[ \text{coccolith exocytosis rate} = \frac{\text{CaCO}_3 \text{ production}}{\text{coccolith weight}} \]  

where coccolith weight was measured as described in Sect. 2.8.

### 2.7 Scanning electron microscopy (SEM)

5–10 ml of sample were filtered by gravity on a polycarbonate filters (0.2 \( \mu \text{m} \) pore size). Samples taken from the constant TA and the constant pH\(_f\) 8 experiment were dehydrated with ethanol and bis(trimethylsilyl)amine solution, to conserve the organic part of the cell, and subsequently dried in a desiccator (Bach et al., 2011). Samples for the other three experiments were not dehydrated and dried for 2 h at 60°C directly after filtration. All samples were kept in the desiccator until they were sputtered with gold-palladium and processed with the scanning electron microscope.

SEM pictures were taken with a CamScan CS 44 scanning electron microscope and evaluated using the software imageJ. Measured lengths or areas on the pictures were calibrated with the size bar given on each SEM picture. Manually measured parameters on coccoliths were the surface area of the distal shield (DSA), the length of the distal shield (DSL), the width of the distal shield (DSW), the length of the central area (CAL) and the width of the central area (CAW) (compare Fig. 1a). CAL and CAW could not
be determined in case the coccolith was lying upside down on the filter. In total 2638 coccoliths were measured manually from SEM pictures. Note that not all treatments could be investigated with SEM due to the extremely elaborate manual evaluation. Evaluated treatments are shown in Table 1.

Measured DSA was compared to an estimated value calculated from DSL and DSW as

\[
\text{calculated DSA} = \pi \times \frac{\text{DSL} \times \text{DSW}}{4}
\]

assuming an elliptical shape of the coccolith. The outer shield length (OSL) was calculated as

\[
\text{OSL} = \frac{\text{DSL} - \text{CAL} + \text{DSW} - \text{CAW}}{4}
\]

2.8 Determination of coccolith weight by birefringence

5–10 ml of sample were filtered with \( \sim 100\) mbar on a cellulose nitrate filter (0.45 µm pore size). Filters were dried for 2 h at 60 °C and subsequently embedded with Acrifix 192 (Roehm) on microscope slides. Acrifix makes cellulose nitrate filters transparent without damaging the coccoliths and has a refraction index of 1.44 so that it does not interfere with the optical analysis.

Images of coccoliths were taken with a Leica DM6000B light microscope equipped with a SPOT Insight b/w camera. Under cross-polarized light only the birefringent calcite of the coccoliths is illuminated. 200 images were randomly taken per sample and analyzed with the software SYRACO (Beaufort and Dollfus, 2004). The software identifies \( E. \) huxleyi coccoliths and measures the grey level for each pixel. Coccolith weight was subsequently calculated from measured grey level following Beaufort et al. (2008). On average, \( \sim 500 \) coccoliths were evaluated for each treatment.
2.9 Calculation of malformation

Coccolith malformation has been defined as “irregular coccolith formation as a result of departure from the normal growth process” and is commonly expressed in reduced symmetry or altered shape of individual elements (Young and Westbroek, 1991). In order to meet the demands given in this definition, fine-scale morphological structures of individual coccoliths were measured and subsequently used in an algorithm to quantify the degree of malformation. The measured morphological quantities comprised vectors associated to the openings between distal shield elements (slits). These were: (1) the distances between the distal ends of two adjacent slits (dl). (2) the distances between the proximal ends of two adjacent slits (pl). (3) The length of each slit (sl) (compare Fig. 1b). Incomplete coccoliths were not measured.

In general, regular and repetitive structures like the individual elements composing a coccolith appear to be malformed in case these adjacent structures differ in an irregular manner. Malformations are therefore characterized in the evaluation procedure as the degree of asymmetry of adjacent slits.

The algorithm to calculate the malformation index makes use of the average deviation. It is defined as:

\[
\text{Average deviation} = \frac{1}{n} \sum_{i=1}^{n} |x_i - \text{mean}(x)_n| \tag{7}
\]

where \( n \) is the number of all measured elements, \( x_i \) is a measured element and \( \text{mean}(x)_n \) is the mean value of all measured elements. Using the average deviation has the advantage that it is not influenced by the number of measured elements. Hence, the malformation index is not sensitive to the number of distal shield elements of the investigated coccolith. Applying the average deviation the degree of malformation is
calculated as:

\[ \text{Malformation} = \frac{1}{n} \sum_{i=1}^{n} \left| \frac{d l_i - d l_{i+1}}{1/2(d l_i + d l_{i+1})} - \text{mean} \left( \frac{d l_i - d l_{i+1}}{1/2(d l_i + d l_{i+1})} \right) \right| \]

\[ + \frac{1}{n} \sum_{i=1}^{n} \left| \frac{s l_i - s l_{i+1}}{1/2(s l_i + s l_{i+1})} - \text{mean} \left( \frac{s l_i - s l_{i+1}}{1/2(s l_i + s l_{i+1})} \right) \right| \]

\[ + \frac{1}{n} \sum_{i=1}^{n} \left| \frac{p l_i - p l_{i+1}}{1/2(p l_i + p l_{i+1})} - \text{mean} \left( \frac{p l_i - p l_{i+1}}{1/2(p l_i + p l_{i+1})} \right) \right| \]

\[ \text{(8)} \]

where \( d l \), \( s l \) and \( p l \) are the measured quantities of the distal shield elements (see above) and \( n \) is the total number of slits. In this way, higher values calculated from Eq. (8) reflect increased malformation (Fig. 1c).

3 Results

3.1 Carbonate chemistry

A large diversity of carbonate chemistry conditions was set up in the five experiments presented in this study. In each one of the five experiments, one particular carbonate system parameter was kept constant while all the others changed with increasing DIC (Fig. 2). In the constant TA experiment, \( fCO_2 \) and \([HCO_3^-] \) increased, while pH and \([CO_3^{2-}] \) decreased with increasing DIC. The carbonate system manipulation of this experiment is similar to the way seawater carbonate chemistry is currently changing due to anthropogenic CO\(_2\) invasion. In the three constant pH experiments, all carbonate system parameters except for pH\(_f\) were increasing linearly with increasing DIC. pH\(_f\) remained constant in all of these experiments but at different levels. In the constant CO\(_2\) experiment, all carbonate system parameters except for CO\(_2\) were increasing with DIC.
3.2 General morphological features

Measured and calculated DSA are in excellent agreement to each other. The slope of the linear regression is close to one which shows that DSA can reliably derived from DSL and DSW (Fig. 3a). The aspect ratio of the coccolith (i.e. DSL : DSW) gets closer to one with increasing coccolith length, indicating that larger coccoliths are rounder than smaller ones (Fig. 3b). The same trend was found for the aspect ratio of the central area. The larger the central area became, the rounder it was (Fig. 3c) which is in good agreement with results obtained by Young and Westbroek (1991). The relative contribution of OSL to total DSL increased with increasing DSL (Fig. 3d).

3.3 Malformation

Malformations of coccoliths correlated best to seawater $\mathrm{pH}_f$, indicating a key influence of $\mathrm{H}^+$. Malformations remained relatively stable above a $\mathrm{pH}_f$ of about 8 whereas they increased with decreasing $\mathrm{pH}_f$ in the range from $\sim 8$ down to 7.1 (Fig. 4). $\mathrm{CaCO}_3$ production rates did not correlate with malformations (data not shown) suggesting that the appearance of malformations is not coupled to calcification rates in *E. huxleyi*.

3.4 Coccolith size

Changes in DSA, DSL and DSW in response to varying carbonate chemistry conditions were largely identical to each other. All three parameters increased most pronounced in the range from low to intermediate $\mathrm{[HCO}_3^\text{-}]$ or $f\mathrm{CO}_2$ whereas changes were minor above this threshold in all except the constant TA experiment where a decreasing trend above $\sim 1000 \mu\text{atm}$ was observed (Fig. 5a–d; data for DSW not shown).

The smallest coccoliths were measured at very low $\mathrm{HCO}_3^\text{-}$ of $\sim 500 \mu\text{mol kg}^{-1}$ (Table 1). These carbonate chemistry conditions are unrealistically low and most likely rarely existed in the natural habitat of *E. huxleyi* since its appearance about 270 ky ago. Hence, sizes determined in these particular treatments should be considered as
physiological potentials rather than realistic representations of naturally occurring *E. huxleyi* coccolith sizes. Variations in size were minor within a realistic DIC and $fCO_2$ range of the last 270 kyr (i.e. from present conditions down to about 1800 µmol kg$^{-1}$ and 180 µatm, respectively). DSA for example, varied from about 8 to 9 µm$^2$ and showed no clear trend within that range if all experiments are considered. Note, however, that DSA increases in the constant TA experiment which simulates ocean acidification, from ~180 to 650 µatm by about 10% and starts to decrease slightly above this threshold.

CAL and CAW remained largely unaffected by changing carbonate chemistry except for the very lowest $fCO_2$ levels in the constant TA experiment (below ~100 µatm) where they showed a decreasing tendency (Fig. 5e, f).

### 3.5 Coccolith weight and production

The mean weight of coccoliths increased by approximately 100% from lowest to highest CaCO$_3$ production rates (estimated from linear fit). Measured mean weight ranged from ~1 to 4 pg (Fig. 6a) which is in reasonable agreement with previous estimates of ~2 pg for the same *E. huxleyi* morphotype (Fagerbakke et al., 1994; Young and Ziveri, 2000). The coccolith exocytosis rate also increased with CaCO$_3$ production by an estimated 100%, similar as for coccolith weight. Minimum and maximum calculated coccolith exocytosis rates were ~12 and 45 coccoliths cell$^{-1}$ d$^{-1}$, respectively (Fig. 6b). Changes in coccolith weight correlate with changes in coccolith size (Fig. 6c).

In the three highest DIC treatments of the constant pH 8 experiment, mean coccolith weight was up to 8 pg which seems unrealistically high. After careful re-evaluation of SEM samples we occasionally found coccoliths that were associated with cubic crystals of unknown material. This might have caused interference with the coccolith weight estimation. Since we could not find a concomitant increase in the CaCO$_3$ content per cell we expect these values to be the result of non-biological processes.
4 Discussion

4.1 Comparison of different evaluation methods of malformations

Malformations of coccoliths in response to changing carbonate chemistry conditions have been observed in several coccolithophore species (e.g., Riebesell et al., 2000; Langer et al., 2006; Müller et al., 2010). In case these malformations were quantified, it was done by visual comparisons of individual coccoliths and subsequent classifications to fixed categories like for example “normal”, “slightly malformed”, “strongly malformed”, and “incomplete” (Langer et al., 2006, 2011; Kaffes et al., 2010; Langer et al., 2011; Bach et al., 2011). Here, we propose an alternative method to approximate malformations of *E. huxleyi* which aims to quantify malformations by direct measurements of fine-scale morphological structures (see Sect. 2.9). A direct comparison of both methods shows that they both lead to similar conclusions. Bach et al. (2011) have visually evaluated the same samples of the constant TA and the constant pH_8 experiment as used in this study and concluded that malformations are mainly induced by seawater pH below ∼ 8. This is largely confirmed by the results from the evaluation of malformation by Eq. (8) (Fig. 4) indicating that both methods seem to be equally appropriate to evaluate malformations of *E. huxleyi*.

The application of these two methods reveals distinct advantages of each one of them. Visual evaluations can be done “online” during scanning of the sample by electron microscopy whereas measurements of morphological structures require a time-intensive analysis of pictures taken by the microscope after having scanned the sample. Hence, visual evaluations facilitate analysis of high number of coccoliths per treatment and therefore usually lead to an investigation of a more representative sample size. Furthermore, visual evaluations are easy to adapt to all coccolithophore species whereas quantification of malformations by direct measurements of morphological structures can so far only be applied to complete coccoliths from *E. huxleyi* morphotype A, B, C and O (for morphotype taxonomy see Young et al., 2003; Hagino et al., 2011). Although adaption of Eq. (8) to other species or morphotypes is generally possible, it
would require measurements of other fine scale structures than the ones used in \textit{E. huxleyi}. The major advantage of the new method is the reduction of subjectivity. A direct measurement of morphological structures reduces human influence on the measurement and makes it easier to compare with results of other studies. Furthermore, Eq. (8) could be implemented in an evaluation software which would analyze malformations automatically. This would be the most efficient and the most reproducible way to quantify malformations.

4.2 Cause of malformations

Malformations of \textit{E. huxleyi} coccoliths are most likely induced by high concentrations of $H^+$ (Fig. 4). In order to understand how excess seawater $[H^+]$ could interfere with coccolith formation, it may be helpful to consider the development of a coccolith on a cellular basis. The formation takes place in a Golgi-derived vesicle (coccolith vesicle or “CV”) which is closely associated with a labyrinthine membrane system (reticular body). Coccolith formation is initiated inside the CV with the production of an organic base plate which serves as template (van der Wal et al., 1983; Westbroek et al., 1984, 1989; Young et al., 1999). Nucleation of calcite occurs subsequently on the rim of the organic base plate from where the initial crystals start to grow in a radial direction until coccolith formation is completed (Westbroek et al., 1984; 1989; Young et al., 1992). Crystal growth is tightly controlled by the cellular machinery. The inner side of the CV membrane always remains in close contact with the coccolith and is actively expanded from the outside by the cytoskeleton located within the cytosol so that the growing calcite crystals fill the space defined by the expanding vesicle (Westbroek et al., 1984, 1989; Didymus et al., 1994, Marsh et al., 1994; Young et al., 2009). Inside the CV, coccolith-associated polysaccharides (CAPs) bound the inner side of the membrane, have a crucial role in controlling CaCO$_3$ precipitation due to their potential to bind Ca$^{2+}$ (De Jong et al., 1976) and inhibit precipitation at places where they cover the calcite (Borman et al., 1982; Henriksen et al., 2004).
Considering the pathway described above, the cytoskeleton and CAPs seem to be two major factors controlling the correct growth of calcite crystals within *E. huxleyi* (Young et al., 1999; Langer et al., 2006). Langer et al. (2010) examined in detail the consequences of a malfunctioning of the cytoskeleton on coccolith formation by applying chemical inhibitors for microtubules and actin filaments. They found an increasing degree of coccolith malformation, the more these cytoskeleton structures and therefore the active expansion of the CV was disturbed by these inhibitors. Possibly, malformations found in our study are also resulting from a malfunctioning of the cytoskeleton, in our case with the chemical driving force being H\(^+\). Such an explanation is feasible because H\(^+\) is known to easily enter into the cytosol of *E. huxleyi* (Suffrian et al., 2011). Here, a change in [H\(^+\)] could disturb the correct functioning of cytoskeleton elements or the enzymes associated with them so that the controlled expansion of the CV is handicapped.

The second possible option of a negative effect of H\(^+\) on crystal growth is a disturbance of CAPs inside the CV (Langer et al., 2006, 2011). Henriksen and Stipp (2009) demonstrated that the capability of CAPs to bind onto calcite crystals depends on the ion composition of the solution in which CAPs are dissolved. A change of the ion composition inside the CV may therefore disturb controlled crystal growth. Such a change could be the direct consequence of a change in [H\(^+\)] inside the CV in case H\(^+\) can somehow enter this compartment. Ion composition inside the CV could, however, also indirectly be altered by changing [H\(^+\)] in the cytosol assuming that H\(^+\) gradients between the cytosol and the CV potentially drive numerous transport processes of major ions such as Ca\(^{2+}\) or Mg\(^{2+}\) (Mackinder et al., 2010, 2011).

The prominent influence of [H\(^+\)] on malformations observed in the investigated *E. huxleyi* strain raises the question whether this particular carbonate chemistry parameter is also responsible for observed malformations in other coccolithophore species and *E. huxleyi* strains. Langer and Bode (2011) examined coccolith malformation of *Calcidiscus leptopus* in response to various carbonate system parameters. In contrast to our findings, they identified CO\(_2\) as the key carbonate chemistry parameter causing
malformations which suggests that the control mechanisms of coccolith formation are affected by different carbonate system parameters on a species level. A comparison on the strain level is not possible because there is no such data on another *E. huxleyi* strain available so far. The only information available at the moment is that the carbonate chemistry conditions at which malformations start to appear differ between different strains of *E. huxleyi* (Langer et al., 2011). Clearly, this does not mean that the key carbonate chemistry parameter causing malformations differs between *E. huxleyi* strains but it shows that sensitivities to changes in carbonate chemistry do.

### 4.3 Influence of individual carbonate chemistry parameters on coccolith size

There is experimental evidence that HCO$_3^-$ is the principal inorganic carbon source utilised for calcification (Sikes et al., 1980; Buitenhuis et al., 1999). The dependence of calcification on HCO$_3^-$ seems to be reflected in DSA, DSL, and DSW which increased comparably in all experiments from low to high [HCO$_3^-$] up to ~ 2000 µmol HCO$_3^-$ kg$^{-1}$. The close correlation to [HCO$_3^-$] makes this ion a key candidate responsible for at least some of the pronounced increase in DSA, DSL, and DSW observed within that range (Fig. 5a–d; data for DSW not shown). Next to HCO$_3^-$, H$^+$ is another factor with potential influence. H$^+$ might be particularly important in the high fCO$_2$ range of the constant TA experiment. Here, the observed decrease in size cannot be explained by [HCO$_3^-$] since the concentration of this ion is still increasing whereas DSA, DSL, and DSW are already decreasing (Fig. 5d). A potential negative effect of very high [H$^+$] seems possible since H$^+$ has already been shown to be detrimental to calcification rates above certain thresholds (Bach et al., 2011). In contrast to HCO$_3^-$ and H$^+$, the influence of CO$_2$ is of minor importance (Fig. 5a–d; data for DSW not shown). DSA, DSL, and DSW clearly decreased in the constant CO$_2$ experiment. If CO$_2$ was of primary importance in determining these morphological parameters, then DSA, DSL, and DSW would have remained constant in the constant CO$_2$ experiment. The fourth parameter with possible influence is CO$_3^{2-}$. CO$_3^{2-}$ is of high importance for the dissolution of calcite due to
its influence on the calcium carbonate saturation state of seawater. Whether it is of
direct physiological influence on coccolith formation is more difficult to assess because
relatively little is known about the possibilities of cells to transport CO$_2^-$ across mem-
branes. Such transporters have so far not been identified (Mackinder et al., 2010). In
our study, a differentiation between HCO$_3^-$ and CO$_3^{2-}$ is only possible in the constant
TA experiment because they co-correlate in all others (Fig. 1). In the constant TA ex-
periment, DSA, DSL and DSW correlate with [HCO$_3^-$] and not [CO$_3^{2-}$] in the low $f$CO$_2$
range. This indicates that increasing [CO$_3^{2-}$] is not likely to be responsible for increasing
DSA, DSL and DSW.

CAL and CAW displayed no clear change in response to changing carbonate chem-
istry except for the very low $f$CO$_2$ range in the constant TA experiment where they
tended to decrease (Fig. 5f, data for CAW not shown). As discussed in Sect. 4.2, coc-
colith formation starts with the construction of an organic base plate on which calcite
crystals start to grow in a radial direction. The dimensions of the central area of the coc-
colith largely reflect the dimensions of the organic base plate (Westbroek et al., 1984;
Young, 1994). Hence, if the dimensions of the central area are affected to a lesser ex-
tent by changing carbonate chemistry, so are the dimensions of the organic base plate.
Accordingly, we hypothesize that changing carbonate chemistry primarily influences
crystal nucleation and growth and not so much formation and size of the organic base
plate.

4.4 Correlation between calcification rates and coccolith weight

Changes in cellular calcification rates can be expressed in three different ways. (1)
A change in coccolith weight at constant coccolith exocytosis rate. (2) A change in coc-
colith exocytosis rate at constant coccolith weight. (3) A simultaneous change in coccol-
ith weight and exocytosis rate. Results presented in Fig. 6a, b support the third option,
indicating that a correlation between calcification rates and coccolith weight exists in
the investigated E. huxleyi strain. This suggests that measurements of coccolith weight
could potentially be useful to reconstruct calcification rates. Nevertheless, this correlation bears uncertainties which should be considered before extrapolating these results to the field. There is the high genetic variability between different coccolithophore species and even strains of the same species (e.g., Brand et al., 1982; Westbroek and Young, 1991; Iglesias-Rodriguez et al., 2006). It has been demonstrated that this variability can cause species- and strain-specific responses in calcification rates within a narrow range of carbonate chemistry conditions (Langer et al., 2006, 2009). However, over a broader range the general response pattern was always similar (an optimum curve type response) where this was tested (Langer et al., 2006; Bach et al., 2011). Therefore, one would expect a similar coupling between CaCO$_3$ production and coccolith weight as shown in Fig. 6a to be present in other species and strains as well, at least over a wider range of carbonate chemistry conditions.

Another factor to consider is the driving force that causes changes in CaCO$_3$ production rates and coccolith weight. Aside from carbonate chemistry, temperature can influence CaCO$_3$ production. A temperature rise from 10 to 20$^\circ$C increases CaCO$_3$ production rates in *E. huxleyi* by more than 40% under ambient carbonate chemistry conditions (unpublished data) whereas it influences coccolith size (and therefore most likely also coccolith weight) only marginally (Watabe and Wilbur, 1966; Fielding et al., 2009). Under this consideration it seems possible that the physiological coupling between CaCO$_3$ production and coccolith weight is not universal but rather specific for changes induced by the carbonate chemistry conditions.

Furthermore, it is important to keep in mind that the correlation between coccolith weight and CaCO$_3$ production rates given in Fig. 6a is derived from monoclonal culture experiments which exclude ecological processes. This is a limitation of the correlation because in a natural *E. huxleyi* assemblage, changing carbonate chemistry could not only directly affect the cell physiology but also induce a shift in the dominant strain. A strain shift in a natural assemblage can change both, mean calcification rate and mean coccolith weight but these two factors do not necessarily have to be correlated to each other as implied in Fig. 6a. The unknown role of ecological processes should...
therefore clearly be considered before using the correlation between calcification rates and coccolith weight to interpret field data.

4.5 Environmental control of coccolith size and weight

There are two different mechanisms how an environmental change in the habitat of an *E. huxleyi* assemblage can induce a change of mean coccolith size and/or weight:

1. The changing environmental factor (e.g. temperature) induces a dominance shift in the assemblage towards an *E. huxleyi* strain or morphotype which forms coccoliths of different size and weight than the one dominant initially. Here, the influence is indirect and in the following termed “ecophysiologically” driven change in coccolith size and/or weight. Morphotype-specific size and weight variations range from 2.5–5 µm and 0.6–4.6 pg, respectively (Young and Ziveri, 2000). Coccoliths of morphotype *R* or over-calcified coccoliths of morphotype *A* are usually relatively heavy, whereas the delicate coccoliths of morphotype *B* (pujosiae) are particularly large (Young and Ziveri, 2000; Young et al., 2003).

2. A change in some environmental factor directly affects the physiology of the dominant *E. huxleyi* strain or morphotype present in the assemblage thereby directly causing a change in mean size and weight (in the following termed “physiologically” driven change in coccolith size and/or weight). Environmental factors known to modify size and/or weight are salinity (Green et al., 1998; Bollmann and Herrle, 2007; Fielding et al., 2009), temperature (Watabe and Wilbur, 1966), nutrient availability (Batvik et al., 1997; Paasche, 1998), growth stage (Westbroek and Young, 1991), seasonality (Triantaphyllou et al., 2010) and carbonate chemistry (Iglesias-Rodriguez et al., 2008; Halloran et al., 2008; Beaufort et al., 2011; this study). In the following we discuss the potential of some of these environmental factors (salinity, temperature and carbonate chemistry) to induce either ecophysiologically or physiologically driven change in coccolith size and/or weight of *E. huxleyi*.

Increasing salinity was shown to positively influence the size of *E. huxleyi* coccoliths (e.g., Green et al., 1998). Fielding et al. (2009) reported a 30% increase in DSW in...
a salinity gradient ranging from 26 to 41 under constant culture conditions indicating that salinity has a relatively high physiological influence on coccolith size. However, whether changing salinity also has the potential to cause a shift in the dominant morphotype in a natural *E. huxleyi* assemblage is unknown. An ecophysiological influence seems conceivable in coastal environments with comparatively large salinity variations while it is less likely in the more stable conditions found in the open ocean.

Temperature seems to have a small physiological influence on *E. huxleyi* coccolith size. Watabe and Wilbur (1966) observed no change in DSL from 7 to 18°C and only a minor decrease of about 10% from 18 to 27°C. This is largely in line with results by Fielding et al. (2009) who found no detectable influence between 10 and 20°C. In contrast to that, the ecophysiological influence of temperature on coccolith size could be considerably larger. It is likely that coccolithophores are adapted to the mean temperature of their natural habitat (Buitenhuis et al., 2008). In case the mean temperature in a given area changes, another strain or morphotype (potentially having a different coccolith size and/or weight) could take over. A possible example where this might have been observed is given by Triantaphyllou et al. (2010) who investigated changes in *E. huxleyi* coccolith size in a seasonal cycle in the Aegean Sea (Eastern Mediterranean Sea). They reported a shift towards larger coccoliths during cooler winter/spring periods with one possible explanation being the dominance of another *E. huxleyi* strain during that time of the year.

The results presented in our study demonstrate a negligible physiological influence of carbonate chemistry on *E. huxleyi* coccolith size and weight within a realistic range (DIC ∼ 1800–2400 and $fCO_2$ ∼ 180–1000). However, there seems to be a high potential of changing carbonate chemistry to cause ecophysiological driven change in coccolith size and/or weight. In a recent investigation, Beaufort et al. (2011) concluded that carbonate chemistry conditions regulate the relative abundance of different species and morphotypes in the oceans and that species and morphotypes which form heavier coccoliths are predominantly found at sites with supposedly more favourable carbonate chemistry conditions. According to the interpretations by Beaufort et al. (2011), the
carbonate chemistry has a particularly large ecophysiological influence on coccolith weight.

Currently, the physiological influence of the environmental factors mentioned above is understood better than the ecophysiological influence. This is probably due to the fact that physiological experiments are in most cases easier to perform and easier to evaluate than ecological data sets. However, in order to improve our understanding of what drives changes in coccolith size and weight in the oceans, it is essential to focus particularly on the ecophysiological component since this seems to be of larger influence.

Acknowledgement. We thank Andrea Ludwig for her support on DIC measurements, and Ute Schuldt and Arno Lettmann from the SEM laboratory at the Institute of Geosciences for their support during scanning electron microscopy. This research was funded by the Federal Ministry of Education and Research (Bundesministerium für Bildung und Forschung; 03F0608A) in the framework of the Biological Impacts of Ocean Acidification (BIOACID) project (subproject 3.1.1 in collaboration with subproject 3.5.3).

References


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Table 1a. Overview on investigated samples. DIC in µmol kg\(^{-1}\); \(fCO_2\) in µatm; weight (of coccoliths) in pg CaCO\(_3\); DSA in µm\(^2\); malformation (mal) dimensionless. Empty fields indicate that data was not collected for these treatments.

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* No coccoliths found by scanning electron microscopy.
** Coccoliths not detectable in cross-polarized light.
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Table 1b. (Continued.)

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* No coccoliths found by scanning electron microscopy.
** Coccoliths not detectable in cross-polarized light.
*** Coccoliths incomplete.
**** No slit present in between adjacent distal shield elements so that malformation could not be determined with the applied method.
Fig. 1. Morphological quantities measured by SEM. (A) Measured size attributes, distal shield area (DSA), distal shield length (DSL), distal shield width (DSW), central area length (CAL), central area width (CAW), and outer shield length (OSL). (B) Measurements for calculation of malformation index, proximal distance between two adjacent slits (pl), length of slit (sl), and distal distance between two adjacent slits (dl). pl, sl, and dl were measured for each slit of the investigated coccolith and then processed with Eq. (8). (C) Examples of coccoliths with corresponding malformation calculated from Eq. (8). The arrow indicates increasing malformation.
Fig. 2. Carbonate chemistry speciation in relation to DIC. Error bars denote the change in carbonate chemistry from the beginning to the end of the experiment. Note that error bars are in most cases masked by symbol size. (A) fCO$_2$ (B) HCO$_3^-$ (C) CO$_3^{2-}$ (D) pH$_f$. Symbol and colour coding is shown in (B).
Fig. 3. Morphology of coccoliths. Each data point represents an individual coccolith. (A) Correlation between measured DSA and DSA calculated from DSL and DSW using Eq. (4). (B) Aspect ratio of distal shield with increasing DSL. (C) Aspect ratio of central area with increasing CAL. (D) Percentage of OSL that contributes to the total DSL. Symbol size and colour coding is shown in (A).
**Fig. 4.** Malformation of coccoliths calculated with Eq. (8).
**Fig. 5.** Coccolith size (DSA, DSL, and DSW) as a function of carbonate chemistry speciation. (A, C, E) in relation to \( \text{HCO}_3^- \). (B, D, F) in relation to \( f\text{CO}_2 \). Symbol and color coding shown in (E).
Fig. 6. Coccolith weight, production and size. (A) Correlation between cellular CaCO₃ production rate and mean coccolith weight. (B) Correlation between cellular CaCO₃ production rate and the number of coccoliths formed per day. (C) Correlation between coccolith size and weight. Error bars denote the standard deviation from measured mean weight or size of all coccoliths of a treatment. Regression line shows a fit through data from all experiments with fit equation and significance given in the figure. Note that the correlation does not apply equally well for individual experiments.