Marine carbonate system evolution during the EPOCA Arctic pelagic ecosystem experiment in the context of simulated Arctic ocean acidification

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Abstract

A major, potential stressor of marine systems is the changing water chemistry following increasing seawater carbon dioxide concentration (CO\textsubscript{2}), commonly termed ocean acidification. In order to understand how an Arctic pelagic ecosystem may respond to future CO\textsubscript{2}, a deliberate ocean acidification and nutrient perturbation study was undertaken in an Arctic fjord. The initial setting and evolution of seawater carbonate chemistry were investigated. Additions of carbon dioxide resulted in a wide range of ocean acidification scenarios. This study documents the changes to the CO\textsubscript{2} system throughout the study following net biological consumption and gas exchange with the atmosphere. In light of the common practice of extrapolating results to cover regions away from experimental conditions, a modelling study was also performed to assess the representativeness, in the context of the simulated present and future carbonate system, of the experimental study region to both the near and wider Arctic region. The mesocosm experiment represented the range of simulated marine carbonate system for the coming century and beyond (\textit{pCO}_2 to 1420 µatm) and thus extrapolations may be appropriate to ecosystems exhibiting similar levels of CO\textsubscript{2} system drivers. However, as the regional ocean acidification was very heterogenous and did not follow changes in atmospheric CO\textsubscript{2}, care should be taken in extrapolating the mesocosm response to other regions based on atmospheric CO\textsubscript{2} scenarios.

1 Introduction

Ocean observations of marine carbon dioxide concentrations (e.g. Sabine et al., 2004; Olsen et al., 2006, 2010; Vázquez-Rodríguez et al., 2009) have confirmed the changing ocean chemical state, in response to partial equilibration with increasing atmospheric CO\textsubscript{2} concentrations. This build up of excess carbon results in a redistribution of dissolved carbon speciation and a lowering of pH (e.g. Zeebe and Wolf-Gladrow, 2001). This process has become known as ocean acidification (Caldeira and Wicket, 2003).
Model simulations point to an even greater rate of perturbation of the marine carbon cycle in the future with the earliest changes projected to occur in the Arctic (Orr et al., 2005; Bellerby et al.; 2005, Steinacher et al., 2009; Denman et al., 2011).

This oceanic uptake provides a great climate service by removing CO$_2$ from the atmosphere and thus reducing the greenhouse warming. The flipside of this oceanic uptake is that the shifting chemistry of the oceans may perturb physiological, ecological and biogeochemical processes, thereby changing the way that energy and elements are transported though marine systems. This may alter marine productivity and biodiversity, challenging both the ecological and economic sustainability of some marine systems (Denman et al., 2011). One approach to understand ecological and biogeochemical responses to ocean acidification is to deliberately perturb marine ecosystems and organisms and then propagated through climate driven ocean models (e.g. Oschlies et al., 2007; Cheung et al., 2011). This form of information maybe then be used to inform ocean management frameworks under future climate change (Halpern et al., 2012).

This study explores the response of the marine carbonate system in an Arctic Fjord to deliberate carbonate system and nutrient perturbations and its subsequent evolution in net response to biological processes and air-sea gas exchange. Furthermore, to put the experiment in a broader pan-Arctic perspective, the experimental CO$_2$ system sensitivity and variability are compared to regional scenarios of future ocean acidification around Svalbard from a regional coupled physical-ecosystem-carbon model.

2 Methods

2.1 Experimental

The experiment was performed in Kongsfjorden, Svalbard 78°56’ N 11°54’ E, between 7 June and 7 July 2010. Nine floating mesocosm enclosures were deployed and filled quasi-simultaneously with in situ fjord water prior to additions of CO$_2$ and nutrients.
Concurrent with sampling for other biogeochemical and biological variables, seawater samples for determining the carbon dioxide system were taken daily from the enclosures using an integrated water sampler (IWS, Hydrobios, Kiel) that took a 5 l sample evenly drawn from a 12 m column in the enclosure. Samples for total alkalinity ($A_T$) and total dissolved inorganic carbon ($C_T$) were drawn into 500 ml borosilicate bottles. No filtering of sample prior to analysis was done due to the lack of significant calcifying plankton (Schulz et al., 2012; Brussard et al., 2012; Niehoff et al., 2012). $A_T$ was measured using Gran potentiometric titration (Gran, 1952) on a VINDTA system (Mintrop et al., 2000) with a precision of 2 µmol kg$^{-1}$. $C_T$ was determined using coulometric titration (Johnson et al., 1987) with a precision of $\leq$ 2 µmol kg$^{-1}$. Measurements for both $C_T$ and $A_T$ were calibrated against certified reference material (CRM; Batch 101) (Dickson, 2010).

### 2.2 Mesocosm CO$_2$ system calculations

The measured $C_T$ and $A_T$, with associated temperatures, salinity and dissolved nutrient data, were applied to the CO2SYS program (Lewis and Wallace, 1998) to calculate additional carbon dioxide system variables. To be consistent with Bellerby et al. (2008), we used the dissociation constants for carbonic acid of Dickson and Millero (1987), boric acid from Dickson (1990a), sulphuric acid following Dickson (1990b) and the CO$_2$ solubility coefficients from Weiss (1974). Values are reported as in situ concentrations. Seawater pH is reported on the total hydrogen scale (pH$_T$) and $p$CO$_2$ in µatm.

### 2.3 Regional model description

This study employed the SINTEF model (SINMOD), a coupled hydrodynamic-ecosystem model encompassed the Nordic Seas, the central Arctic Ocean and the Eurasian shelf (Fig. 1) (Slagstad and Mcclimans, 2005; Wassmann et al., 2006). The hydrodynamic model is based on the primitive Navier-Stokes equations discretised on a z-grid (Slagstad and Mcclimans, 2005), and includes a sea-ice model similar to that of

(Riebesell et al., 2012).
Hibler (1979). The sea ice model includes state variables for properties such as salinity, $A_T$ and $C_T$ allowing for these properties to be transported by ice drift and distributed during sea-ice melt. The structure of the ecological sub model is shown in Fig. 2. The basic model currency is nitrogen (mmol N m$^{-3}$), and the conversion to carbon is performed using a C:N ratio of 7.6, based upon regional data from the Barents Sea (Reigstad et al., 2002). The state variables are: $C_T$, $A_T$, nitrate, ammonium, silicate, diatoms, autotrophic flagellates, bacteria, heterotrophic nanoflagellates, microzooplankton, fast sinking detritus, slow sinking detritus, dissolved organic carbon, bottom sediment and two groups of mesozooplankton, one representing the Atlantic species, *Calanus finmarchicus* and one, *Calanus glacialis*, representing the Arctic species.

3 Set-up and boundary conditions

The model employs a horizontal grid with a uniform resolution of 20 km and 25 vertical levels. Initial values of temperature and salinity were taken from the World Ocean Circulation Experiment Global Data Resource Version 3.0 (http://www.nodc.noaa.gov, Lindstrom, 2001). SINMOD has open ocean boundaries to the Atlantic Ocean and the Bering Sea (Fig. 1). Water fluxes through these open boundaries were kept constant during the simulation; a flux of 0.8 Sv enters the model domain through the Bering Strait (Woodgate et al., 2010); on the Atlantic side the fluxes were derived from a large scale (50 km resolution) model covering the North Atlantic. A total of 8 tidal components were imposed, specifying the various components at the open boundaries of the large-scale model. The model is also forced with freshwater fluxes from both river discharges and diffuse run-off from land. Freshwater run-off along the Norwegian coast and in the Barents Sea is based on data from a simulation with a hydrological model (see Dankers and Middelkoop, 2008 for more details). For Arctic Rivers flows, data were obtained from R-ArcticNet (Vörösmarty et al., 1996, 1998) accessible through http://www.r-arcticnet.sr.unh.edu/v4.0/index.html.
The climatic scenario runs use atmospheric forcing fields derived from a regional model system, configured to cover the model domain of SINMOD, run by the Max Planck Institute, (REMO; Keup-Thiel et al., 2006; Slagstad et al., 2011).

For marine biogeochemistry, monthly mean values of $C_T$, $A_T$, and nutrients from the Bergen Climate Model (BCM) (Assmann et al., 2009), corrected using the CARINA database climatology (Tanhua et al., 2010) and the absolute delta change method (e.g. Hay et al., 2000), were used at the boundaries. From $C_T$ and $A_T$, with associated salinity and temperature, phosphate and silicate concentrations, the other CO$_2$ system variables were determined following Dickson (2007) with the dissociation constants of Mehrbach et al. (1973) modified by Leuker et al. (2000) on the total hydrogen scale. The representation of air-sea exchange of CO$_2$ followed Wanninkhof (1992).

4 Results and discussion

The seawater for the experiment was isolated from the fjord on 2 June (t-5). The initial characterization of the CO$_2$ system in the mesocosm and the fjord was performed on t-3 prior to the CO$_2$ addition (Riebesell et al., 2012). The mesocosm values agreed to $\pm 1.2 \mu$mol kg$^{-1}$, i.e. within the measurement precision, for both $C_T$ and $A_T$ (not shown). This confirmed that the closing of the bags isolated water of very similar biogeochemical properties in each mesocosm; a significant feat due to the typical small scale heterogeneity of the fjord (Svendsen et al., 2002). Following the final carbon dioxide perturbations on t4 (Schulz et al., 2012; Czerny et al., 2012a) it took a further four days for the CO$_2$ system to settle down in the mesocosms due to exchange with “dead volume” in the base of the bags and thus, all changes to the CO$_2$ fields were referenced to t-8.

$A_T$ increased steadily in all the bags from 2242 $\mu$mol kg$^{-1}$ on t8 to 2247 $\mu$mol kg$^{-1}$ on t25 falling back to the original 2242 $\mu$mol kg$^{-1}$ by t27. The increase was due to freshwater losses, following evaporation, and nutrient uptake (Silyakova et al., 2012). However, in the absence of significant numbers of calcifiers (Schulz et al., 2012; Brussard et al., 2012; Niehoff et al., 2012), there were no significant alkalinity changes following

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The effect of nutrient addition on t13 could not be seen in $A_T$ as the addition was alkalinity neutral due to the concomitant addition of acid (Riebesell et al., 2012). As there were no other changes in other associated biogeochemical variables and salinity, it is likely that the drop in $A_T$ on T27 was a calibration offset.

$C_T$ concentrations showed high variability between the mesocosms in response to the deliberate additions of CO$_2$. From an original fjord value of about 1982 µmol kg$^{-1}$, the perturbations spanned a range from 1982 to 2270 µmol kg$^{-1}$. In the high CO$_2$ scenarios, $C_T$ drops rapidly and consistently throughout the experiment with net $C_T$ changes between 52 and 63 µmol kg$^{-1}$. In the intermediate CO$_2$ scenarios, $C_T$ concentrations change much more slowly until about t23 when there is a much faster reduction. Total reductions in the intermediate scenario were between 54 and 58 µmol kg$^{-1}$. In the low CO$_2$ scenario mesocosms, $C_T$ increases until t19 before exhibiting the fastest decline of all the scenarios towards the end of the experiment resulting in a net change of between 31 and 40 µmol kg$^{-1}$.

The initial mesocosm pCO$_2$ concentrations were chosen to represent a range of atmospheric values corresponding to anticipated carbon fossil fuel release scenarios. pCO$_2$ showed very large inter- and intra-mesocosm variability, particularly in the high CO$_2$ scenarios. This is due to the poor buffer capacity of the seawater that results in increasing sensitivity in pCO$_2$ to even small changes in $C_T$ and $A_T$ that result from both net ecosystem perturbations and from measurement sensitivity. The higher CO$_2$ scenario mesocosms also exhibited the largest reductions in pCO$_2$ enhanced by rapid gas exchange with the atmosphere.

Initial pH$_T$ levels ranged from 7.5 to 8.3 and, in all bags, increased through the experiments according to the relative amounts of CO$_2$ exchange with the overlying atmosphere and biological net carbon production. The high CO$_2$ mesocosm exhibited the greatest pH changes.

The aragonite saturation state ($\Omega_{ar}$) had the highest values (2.6) in the control mesocosms. The seawater was under-saturated with respect to aragonite in the four highest CO$_2$ mesocosms with the lowest $\Omega_{ar}$ of the experiment being 0.5. Seawater was
under-saturated with respect to aragonite for the entire experimental period under the highest CO₂ scenario.

Carbonate system variability can be high in the surface ocean due to seasonal perturbations from biological CO₂ uptake following photosynthesis and release through respiration. The system also responds to the changes in solubility of CO₂ that follow seasonal and episodic heating events. Furthermore, in the Arctic freshwater melt and river inputs can play a large role in determining the seasonal range and speciation of total inorganic carbon. Understanding the complex interplay of carbon cycle drivers requires a modeling approach to develop optimized simulations of future ocean acidification.

One of the main purposes of this modelling exercise was to examine how the mesocosm carbonate system changes scaled in the context of broader regional ocean acidification changes simulated for the coming century. This study chose two approaches: the first was to study regions in juxtaposition with the Svalbard archipelago and; the second to widen the view to encompass the waters of the northern Nordic Seas and the shelf regions and western Eurasian Basin of the Arctic Ocean. As the IPCC scenario SRESA1B was used in the model simulations, comparisons with the more wider ranging mesocosm experiment are constrained to those with atmospheric CO₂ simulations between present and year 2100. These correspond most closely to mesocosm M1 and M4.

Most simulations of the future Arctic Ocean CO₂ system have documented yearly average (Orr et al., 2005; Steinacher et al., 2009; Denman et al., 2011) or wintertime (Bellerby et al., 2005) ocean acidification. Of more relevance to biological systems and biogeochemical cycles is the seasonal cycle and overlying trends. For the regions around the Svalbard archipelago: the northern Barents Sea opening; the West Spitsbergen Current (WSC); and the northern Svalbard shelf (Fig. 3), the seasonal variability in the bi-weekly, mixed layer means for \( pCO₂ \), \( pH_T \) and \( \Omega_{ar} \) are shown in Fig. 6. The periods of study are the decades 2006–2015 and 2090–2099.
The modern model $pCO_2$ cycle for the WSC is somewhat higher than the summertime values of the fjord and the control mesocosms. Further, due to the unseasonal forcing through a significant nutrient addition to the mesocosms, it is difficult to determine which stage of the year the mesocosms were simulating and thus allow a complete comparison with rate of change of the spring bloom. The comparison does, however, enable a regional scaling of the experiment to anticipated changes in the coupled Arctic system and thus can inform on the limits to representation of ocean-acidification-ecosystem responses founded from the mesocosm results.

This experiment pushed the envelope of $pCO_2$ ranges beyond that which was studied in the PeECE experiments (Bellerby et al., 2008). The large seasonal variability in the CO$_2$ system exhibited in the models (Fig. 6) challenges the use of fixed CO$_2$ system approaches. This study highlights the necessity to operate with mesocosms closely simulating natural variability the CO$_2$ system following biological activity to allow more realistic seasonality. However, as the mesocosms were open to the contemporary atmosphere, and not one simulating the correct scenario atmospheric $pCO_2$, as was done in the semi-closed systems in the PeECE study (Riebesell et al., 2008), air-sea exchange was a prominent driver of the net CO$_2$ changes, especially at the lowest and highest scenarios (Czerny et al., 2012b). The rates of CO$_2$ system change seen in the experiment can then only approximate the anticipated ocean acidification response expected from the initial CO$_2$ levels. Accordingly, any results from the experiment should not be interpreted as corresponding to the representative year corresponding to a particular IPCC scenario. The experiment does allow for interpretation of results to the carbonate system chemistry. In summary, organisms, ecosystems and biogeochemistry are blind to operationally defined CO$_2$ scenarios and thus it is not the atmospheric CO$_2$ values that should guide interpretation of a response to CO$_2$ but the absolute ocean acidification of the system at the phase of interest.

A larger sub-section of the full model domain was chosen to represent the variability in the marine carbonate system in the waters of the Nordic Seas and Arctic shelves north of Svalbard for decades at the beginning and the end of this century (Fig. 7).
This region is driven by water masses and currents that incur a dominant control on the marine carbonate systems around Svalbard. These are predominantly the North Atlantic-, Norwegian Atlantic-, and West Spitsbergen Current chain, the East Greenland Current and the waters in the northern Barents Sea. The current chain can be considered to be the “North Atlantic acidification highway” to the Arctic as it presently brings with it most of the anthropogenic carbon to the region (Olsen et al., 2006).

For the contemporary period 2006–2016, there is a $p\text{CO}_2$ undersaturation for the entire region, with the highest values associated with the waters of the Iceland and Norwegian Seas. Concentrations increase towards the north until a rapid reduction in $p\text{CO}_2$ is found at the Western Greenland Sea running north through the Fram Strait and rounding Svalbard towards the shelf north of the Barents Sea. Further north of this “front”, $p\text{CO}_2$ increases again to values similar to those in the northern Greenland Sea.

The regional, annually averaged, surface carbonate chemistry exhibits even greater heterogeneity in all $\text{CO}_2$ system variables (Fig. 7) than the seasonal study local to the Svalbard region (Fig. 6). The largest centennial changes are seen in the waters associated with the Arctic (Fig. 8) concurring with another independent model study of the region (Bellerby et al., 2005). There is a very clear demarcation where the largest changes in sea ice cover (not shown, Slagstad et al., 2011). It is this region that the greatest warming and freshening are simulated and these drive changes in ocean acidification in addition to those caused solely by increased $\text{CO}_2$. Warming attenuates the drop in the saturation state and yet increases the pH reduction. Freshening causes a lowering of $A_T$ and $C_T$ that together result in a lowering of pH and of saturation state (e.g. Yamamoto-Kawai et al., 2009).

5 Conclusions

This study has documented the potential for significant ocean acidification perturbations in a future Arctic fjord environment. It reports on the the most northerly, large scale, mesocosm experiment to date and the most detailed to study an Arctic pelagic
ecosystem. The model analysis is a first attempt to put mesocosm perturbations into a regional perspective. It recognizes the limitations of downscaling scenarios at the local scale with the coarse model resolution but it highlights the requirement for detailed local studies of ocean acidification relevant to individual organisms and ecosystems. The modeling study highlights large heterogeneity in the regional and seasonal response of the marine carbonate system in the waters of the Arctic, with a focus on the waters around Svalbard. Climate change feedbacks on the CO$_2$ system mainly through the very significant processes of warming and freshening. Incongruities with the absolute seasonal timing of the mesocosm with the model results in no way lessens the importance in this experiment and its relevance for understanding ecosystem response. However, for inclusion of these results in management plans for the sustainability of this important region, caution should be taken in extrapolating the mesocosm results to represent thresholds or responses relevant to a particular time period. Whilst the results of organisms, ecosystems and biogeochemistry can be linked to absolute carbonate system properties, if information is required on the timing of these responses or the atmospheric CO$_2$ concentrations under which these will occur, then a better understanding of the chemical setting is required. This highlights the need for more dedicated studies relevant to key species and biogeochemical provinces. For more realistic representations on future Arctic ecosystem responses it is required that multi-stressor experiments be performed. It is also apparent that under the same atmospheric forcing, the regional heterogeneity of future ocean acidification requires dedicated organismal and ecosystem investigations if informed management plans are to be developed.

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Fig. 1. Regional domain of the SINMOD model.
Fig. 2. Ecological and biogeochemical components of the SINMOD model and their interactions.
Fig. 3. Sub-regional domains for the seasonal modelling study. (1) Barents Sea Opening; (2) West Spitsbergen current and (3) Arctic Shelf.
Fig. 4. Absolute values for the marine carbonate system variables. Measured values are (a) total inorganic carbon ($C_T$) and (b) total alkalinity ($A_T$). Calculated values are (c) partial pressure of carbon dioxide ($pCO_2$), (d) pH$_T$ on the total hydrogen scale and (e) aragonite saturation state ($\Omega_{ar}$). Red symbols: high $pCO_2$ mesocosms (M5, M6, M9); grey symbols: medium $pCO_2$ mesocosms (M1, M4, M8); blue symbols: low $pCO_2$ mesocosms (M2, M3, M7).
Fig. 5. Cumulative changes relative to the start of the post CO₂ perturbation (T-8). (a) total inorganic carbon (Cₜ), (b) total alkalinity (Aₜ), (c) partial pressure of carbon dioxide (pCO₂), (d) pHₜ on the total hydrogen scale and (e) aragonite saturation state (Ωₘ). Red symbols: high pCO₂ mesocosms (M5, M6, M9), grey symbols: medium pCO₂ mesocosms (M1, M4, M8), blue symbols: low pCO₂ mesocosms (M2, M3, M7).
Fig. 6. Simulated mixed layer seasonal marine carbonate system dynamics for the regional sub-domains: $pCO_2$, pH and $\Omega_{ar}$. The decadal means are shown as blue (period 2006–2015) and red (period 2090–2099) for the regions of the Barents Sea Opening ($\bigcirc$); West Spitsbergen current ($\Box$) and the Arctic Shelf ($\triangledown$).
Fig. 7. Simulated mixed layer marine carbonate system values for the northern Nordic Seas and Eurasian Arctic Ocean. Decadal means are presented for the time-slices 2006–2015 and 2090–2099 for $p\text{CO}_2$, pH$_T$ and $\Omega_{ar}$.
Fig. 8. Centennial changes in mixed layer marine carbonate system values for the northern Nordic Seas and Eurasian Arctic Ocean. The figures represent the difference between the decadal means in Fig. 6 for (a) $p$CO$_2$; (b) pH$_T$ and (c) $\Omega_{ar}$. 