The potential of $^{230}$Th for detection of ocean acidification impacts on pelagic carbonate production

Christoph Heinze$^{1,2}$, Tatiana Ilyina$^3$, and Marion Gehlen$^4$

$^1$Geophysical Institute, University of Bergen and Bjerknes Centre for Climate Research, Bergen, 5020, Norway
$^2$Uni Research Climate, Bergen, 5007, Norway
$^3$Max Planck Institute for Meteorology, Hamburg, 20146, Germany
$^4$Laboratoire des Sciences du Climat et de L’Environnement UMR CEA-CNRS-UVSQ, Gif-sur-Yvette, 91191, France

Correspondence to: Christoph Heinze (christoph.heinze@uib.no)

Abstract. Concentrations of dissolved $^{230}$Th in the ocean water column increase with depth due to scavenging and downward particle flux. Due to the $^{230}$Th scavenging process, any change in the calcium carbonate (CaCO$_3$) fraction of the marine particle flux due to changes in biological CaCO$_3$ hard shell production as a consequence of progressing ocean acidification would be reflected in the dissolved $^{230}$Th activity. Our prognostic simulations with a biogeochemical ocean general circulation model using different scenarios for the reduction of CaCO$_3$ production under ocean acidification and different greenhouse gas emission scenarios (RCPs 8.5 to 2.6) reveal the potential for deep $^{230}$Th measurements to detect reduced CaCO$_3$ production at the sea surface. The time of emergence of an acidification induced signal on dissolved $^{230}$Th is of the same order of magnitude as for alkalinity measurements. Yet, deep ocean $^{230}$Th concentrations are less affected by seasonal and multiyearvariability than surface alkalinity. Thus, deep ocean $^{230}$Th observations could be advantageous to guide monitoring and detection campaigns. Furthermore, given that the precision of $^{230}$Th measurements may potentially improve in the near future, earlier detection of ocean acidification impact signals would be possible. Our results indicate that the deep Pacific Ocean and the deep Southern Ocean are the most suitable regions for selected regular reoccupations of deep reaching $^{230}$Th stations.

1 Introduction

Progressing ocean acidification is a fact. It can be directly seen from measurements at Eulerian time series stations (e.g., Bates (2007); Dore et al. (2009); Santana-Casiano et al. (2007)) and also at larger depth from high quality deep hydrography data (Olafsson et al., 2009). Depending on the emission scenario for CO$_2$, the decrease in ocean pH and the decrease in carbonate saturation will continue and become more pressing during this century (e.g., Orr et al. (2005); Steinacher et al. (2009); Bopp et al. (2013)). Temporal and regional patterns of changes in pH and carbonate saturation can be relatively straightforwardly projected by Earth system models including marine inorganic carbon chemistry formulations (e.g. Bopp et al. (2013)). They also can be monitored through long-term high quality measurements of the inorganic carbon system. However, monitoring impacts of ocean acidification on biological processes remains challenging. While for some organisms, especially corals (Kleypas et al., 1999), detrimental effects due to decreasing pH and carbonate saturation have been determined, the various physiological impacts of ocean acidification on specific organisms and ecosystem functioning are still under investigation (e.g., Iglesias-Rodriguez et al. (2008); Kroeker et al. (2013); Meyer and Riebesell (2015); Riebesell et al. (2007)). At present, a series of possible pH-dependencies of governing marine carbon cycle
parameters (such as elemental stoichiometric ratios and nutrient uptake kinetics under biological particle production) are discussed. So far, the potential decrease in calcification due to the lowering of carbonate saturation under high pCO2 is among the key changes which may be expected to occur, especially when it comes to organisms building aragonite shells (the meta-stable form of CaCO3, calcite has a lower solubility than aragonite) (Raven et al., 2005). Two important questions emerge: 1. If changes in biological calcification would indeed occur in the ocean during the coming years – how can they be detected and monitored by observational frameworks (through which methodology and through which variable)? 2. In which oceanic region could one observe early signals of these changes unambiguously at the earliest possible stage?

Development of such early warning systems is vital in order to check the validity of parameterisations of pH-dependent processes in ocean models and to take appropriate mitigation/adaptation measures to diminish the consequences of potential considerable ecosystem changes due to ocean acidification. Such changes could affect the marine food chain. In a global modelling study, Ilyina et al. (2009) quantified the detection thresholds for changes in alkalinity due to changes in a series of possible formulations for the reduction of calcification (with the term “calcification” we mean here the production of CaCO3 hard parts by marine biota) with pH decrease. The logic behind the approach of Ilyina et al. (2009) is as follows: If ocean acidification leads to a decrease in calcification, more CO3^2- ions would be retained in the surface water and not become incorporated into CaCO3 shell material. Changes in CO3^2- ion concentration would induce a change in ocean total alkalinity, which could eventually be measured against an earlier baseline. According to that study, the tropical Pacific (with high CaCO3 production rates) would be the region for detecting such alkalinity changes first, as the anticipated change in CaCO3 production rates would be the region for detecting such alkalinity changes first, as the anticipated change in CaCO3 production would be largest there in absolute terms. For intermediate dependencies of CaCO3 production on pH/carbonate saturation, a reduction in bio calcification could unequivocally only be diagnosed from ca, year 2035 on. In the Arctic Ocean, where pH changes are expected to be most pronounced, large scale changes in calcification would be detectable even later on the basis of alkalinity measurements due to the overall lower biological production rates. Likewise, detection may be additionally complicated due to signals from natural seasonal and interannual variations in surface total alkalinity (Carter et al., 2016). There is thus a need for the development of novel detection methods. Heinze et al. (2006) investigated the impact of varying rain ratios CaCO3:Corg on the distribution of the radionuclides thorium (230Th), protactinium (231Pa), as well as beryllium (10Be). As the rain ratio describes the average amount of carbon atoms incorporated into CaCO3 shell material relative to the amount of carbon atoms incorporated into organic matter by plankton, a reduction in calcification would lead to a rain ratio decrease. Indeed, especially for 230Th, due to its affinity for scavenging to CaCO3 particles, a considerable increase in the concentration of dissolved 230Th with depth and time was obtained in a sensitivity experiment with strong rain ratio reduction. In this paper here, we explore the option of using radionuclides for diagnosing changes in calcification and respectively reduced downward marine particle flux in more detail.

2 The concept - radionuclides and particles

We focus here on 230Th, a long lived radionuclide (half-life 7.5x10^4 yr) and a highly particle reactive metal. 230Th is produced in the uranium decay series. As uranium has a very long residence time in the ocean and quasi-
homogenous concentrations in seawater, the present natural marine $^{230}$Th source is constant everywhere in the ocean water column and thus ideally suited for using it as a tracer in modelling studies. $^{230}$Th is strongly particle reactive and is removed from the water column quickly through scavenging by the vertical particle flux in the ocean (for a summary, please, see Henderson et al. (1999)). The majority of $^{230}$Th is removed from the water column and will not re-enter the bottom waters, e.g., through sediment pore water diffusion (though resuspension may potentially cause some re-release). In spite of its strong particle reactivity, concentrations of dissolved $^{230}$Th show horizontal as well as vertical gradients in the water column which are induced by the oceanic current field and differential particle fluxes as well as particle concentrations (e.g., Henderson et al. (1999); Yu et al. (1996)). The distribution of particle bound $^{230}$Th follows the particle concentrations and particle fluxes. Both, the dissolved $^{230}$Th and the particle bound $^{230}$Th show nutrient type vertical profiles with low values at the surface and increasing concentrations with depth. Dissolved $^{230}$Th increases with depth as particles continuously carry $^{230}$Th downwards and new equilibria between the dissolved phase and the particle attached phase establish as illustrated in Figure 1. The equilibrium between the concentration of the dissolved phase $[^{230}\text{Th}_{\text{diss}}]$ and the concentration of the particle attached phase $[^{230}\text{Th}_{\text{part}}]$ can be described in an analog way to a chemical reaction equation:

$$^{230}\text{Th}_{\text{diss}} \leftrightarrow ^{230}\text{Th}_{\text{part}}$$

The respective analog for the mass action law constant describing to which extent the reaction from left to right is carried out is then given through the $k_d$ value governing the equilibrium between the dissolved and particle bound phases for the radionuclide:

$$k_d = \frac{[^{230}\text{Th}_{\text{part}}]}{[^{230}\text{Th}_{\text{diss}}]}$$

Often, $k_d$ values are formulated to account for a shift of the equilibrium towards the particle attached phase at low particle mass concentrations assuming that low particulate concentrations occur in parallel to low particle sizes with correspondingly high reactive surface areas when compared to large particles (e.g. Honeyman et al. (1988)). Respective formulations have been used successfully in $^{236}$Th modelling studies (Heinze et al., 2006; Henderson et al., 1999).

\[\text{Figure 1: Schematic illustration of the equilibration process between the dissolved and particle-attached phases of }^{230}\text{Th and the increasing concentrations downward in the water column.}\]
How would then the distribution of dissolved $^{230}\text{Th}$ in the ocean reflect changes in marine calcification at the sea surface? There are indications, that $^{230}\text{Th}$ is attached first of all to CaCO$_3$ and clay particles in the water column as preferential carrier phases (Chase et al., 2002, 2003; Luo and Ku, 1999). Indeed, Heinze et al. (2006) reproduced the large scale $^{230}\text{Th}$ distribution in the world ocean with a coarse resolution biogeochemical ocean general circulation model based on a formulation allowing $^{230}\text{Th}$ scavenging only by CaCO$_3$ and clay particles. Other studies have indicated that preferential carrier phases may vary regionally in the ocean (Scholten et al., 2005). However, Heinze et al. (2006) could demonstrate that rain ratio changes (CaCO$_3$:Corg) in marine biological particle export production could also be recorded, if $^{230}\text{Th}$ became in addition attached to particulate organic carbon (POC). A weakening of CaCO$_3$ particle production would result in a less efficient $^{230}\text{Th}$ scavenging as less particles (in terms of mass) would be available to carry $^{230}\text{Th}$ to larger depths and finally the sediment. Changes in the strength of CaCO$_3$ production and the respective downward particle flux are reflected increasingly better in the $^{230}\text{Th}$ distribution with increasing depth (see Heinze et al. (2006)) due to two reasons. First, CaCO$_3$ particles get less well degraded as POC (which is remineralised mostly in the upper 1000 m of the water column) and thus reach larger depth; this is also corroborated from sediment trap measurements (e.g., Honjo, 1996). Second, due to the scavenging of $^{230}\text{Th}$ by particles, the vertical downward particle flux, and the equilibration between dissolved and particle bound phases, $^{230}\text{Th}$ is continuously transferred from shallower layers to larger depths. Thus temporal changes in $^{230}\text{Th}$ scavenging in the upper ocean become enlarged in the deep $^{230}\text{Th}$ distribution as in a kind of “magnifying glass” (see Figure 1). We investigate here, whether this feature can be exploited for an early detection method of large scale reductions in calcification and correspondingly reduced rates in CaCO$_3$ particle export and CaCO$_3$ particle fluxes through the water column.

3 Model description

In this study we use the Hamburg ocean carbon cycle circulation model HAMOCC (Maier-Reimer, 1993) in its annually averaged version (time step 1 yr, Heince and Maier-Reimer (1999); Heince et al. (2009); Heince et al. (2016)) with a horizontal resolution of 3.5°x3.5°. This coarse resolution model is computationally very efficient and useful when multiple integrations are needed such as for the testing and adjusting of scavenging codes. An advantage of this fast model is, that it includes a fully equilibrated early diagenesis module (10 layers) under each grid point and thus can account for alkalinity changes induced by dissolution of CaCO$_3$ from the seafloor. The model version employed here corresponds to the version as used in Heince et al. (2009) and Heince et al. (2016) with the scavenging module of Heince et al. (2006). For details, please, see these publications. The model uses a fixed ocean velocity field (and thus provides no dynamical computation of the ocean currents; velocities, temperature, salinity and ice cover are rather read from an input file). The model includes an atmospheric compartment (“slab atmosphere”) which allows for prognostic computation of the atmospheric CO$_2$ concentration as well as meridional atmospheric CO$_2$ transport. We describe here only briefly features of specific relevance for this study. The water column is structured into 11 layers (centred at 25, 75, 150, 250, 450, 700, 1000, 2000, 3000, 4000, and 5000 m). The bioturbated top sediment zone of the early diagenesis module is divided into 10 layers which are separated by interfaces at 0, 0.3, 0.6, 1.1, 1.6, 2.1, 3.1, 4.1, 5.1, 7.55, and 10 cm “downcore.” We make the simplifying assumption that no pore water reactions take place below 10 cm depth in the sediment (see, e.g., Smith and Rabouille (2002); Boudreau (1997)). The biogeochemical model includes the
processes of air-sea gas exchange, biogenic particle export production out of the ocean surface layer, particle
flux through the water column and particle degradation by dissolution as well as remineralization, transport of
dissolved substances with the ocean currents, deposition of particulate constituents on the ocean floor, pore water
chemistry and diffusion, advection of solid sediment weight fractions (organic carbon, organic phosphorus,
CaCO₃, opal, and clay), bioturbation, and sediment accumulation (export out of the sediment mixed layer). The
model predicts the following tracer concentrations in the atmosphere, the ocean water column and in the
sediments. Atmospheric tracers include the concentrations of $^{12}$CO₂ (carbon dioxide), $\delta^{13}$CO₂, and O₂. In the
water column, concentrations of DIC (dissolved inorganic carbon), POC (particulate organic carbon), POP
(particulate organic phosphorus), DOC (dissolved organic carbon), CaCO₃ (calcium carbonate or particulate
inorganic carbon) – all for $^{12}$C as well as $^{13}$C, dissolved oxygen O₂, dissolved PO₄³⁻ as biolimiting nutrient,
silicic acid Si(OH)₄ and opal (biogenic particulate silica BSi) are calculated. In the sediment pore waters, the
same dissolved substances as in the water column, as well as solid sediment constituents such as clay, CaCO₃,
opal, and organic carbon are simulated. The inorganic carbon chemistry is computed following Dickson et al.
(2007). In the advection scheme and for the other chemical reactions, DIC and TAlk are used as “master
tracers” form which derived quantities such as the CO₃²⁻ concentration and the pH value are computed through a
Newton-Raphson algorithm. In the annually averaged model as employed in this study, only export production
of biogenic particles is modelled (and no explicit phytoplankton and zooplankton concentrations). Particle
production takes place in the model surface layer representing the euphotic zone. Phosphate serves as biolimiting
nutrient. POC and opal export productions are simulated following Michaelis Menten kinetics for nutrient uptake
(e.g., Sarmiento and Gruber (2006)) (where the phytoplankton concentration is replaced by the phosphate
concentration as ecosystem processes as such are not explicitly modelled):

$$P_{\text{POC}} = \frac{V_{\text{max}}^{\text{POC}} \cdot \left[\text{PO}_4^{3-}\right]^2}{K_{s}^{\text{POC}} + \left[\text{PO}_4^{3-}\right]^2};$$

and

$$P_{\text{opal}} = \frac{V_{\text{max}}^{\text{opal}} \cdot \left[\text{Si(OH)}_4\right]^2}{K_{s}^{\text{opal}} + \left[\text{Si(OH)}_4\right]^2};$$

where $P_{\text{POC}}$ and $P_{\text{opal}}$ are the POC and opal export production rates (mol l⁻¹ yr⁻¹), Red(C:P) is the Redfield ratio
C:P, $V_{\text{max}}^{\text{POC}}$ and $V_{\text{max}}^{\text{opal}}$ are the maximum uptake rate of phosphate and silicic acid from the water column (yr⁻¹), and $K_{s}^{\text{POC}}$ as well as $K_{s}^{\text{opal}}$ are the respective half saturation constants. $V_{\text{max}}^{\text{POC}}, V_{\text{max}}^{\text{opal}}, K_{s}^{\text{POC}},$ and $K_{s}^{\text{opal}}$ are
simulated as a function of sea surface temperature as described by Heinze et al. (2003). POP production follows
POC production with a constant stoichiometry here. The export production of CaCO₃ is coupled to the local
production ratio $P_{\text{opal}}/P_{\text{POC}}$. It starts to increase gradually (parameter R see below) if $P_{\text{opal}}/P_{\text{POC}}$ sinks below a
threshold value $S_{\text{opal}}$, i.e., when not enough silicic acid is available in the ocean surface layer to fuel full diatom
growth:

$$P_{\text{CaCO}_3} = P_{\text{POC}} \cdot R \cdot a \cdot \left(1 - \frac{P_{\text{opal}}}{P_{\text{POC}}} \right) \quad \text{for} \quad \frac{P_{\text{opal}}}{P_{\text{POC}}} < S_{\text{opal}}; \quad (1)$$
Parameter R is the maximum possible rain ratio C(CaCO₃)/C(POC), a is the CaCO₃ saturation dependent factor to account for an ocean acidification impact (see Figure 2, following Ilyina et al. (2009)), and S_{opal} is the threshold value of P_{opal}/P_{POC} for gradual onset of CaCO₃ production. Particle fluxes and particle degradation are simulated through balance equations for sinking particulate matter as in Heinze et al. (2009) and Heinze et al. (2016) taking the saturation state for CaCO₃ and biogenic silica into account.

Figure 2: Assumed sensitivities of CaCO₃ export production in response to changes in calcite saturation. The rain ratio coefficient is used as factor “a” applied to the right hand side of the equation for CaCO₃ production (see eq. 1). For the control run, “a” was fixed at each grid point according to the prevailing pre-industrial carbonate saturation (following the moderate sensitivity). Blue: Moderate sensitivity. Green: Linear sensitivity. Red: Extreme sensitivity. The shape of the curves would look similar at each grid point; only the pre-industrial cross-over point would be different.

Scavenging of $^{230}$Th is simulated according the reversible first-order scavenging reaction (Gehlen et al., 2003; Heinze et al., 2006):

$$\frac{d c_{\text{part}}}{dt} = K \cdot (c_{\text{eq}}^\text{part} - c_{\text{part}})$$

$c_{\text{eq}}^\text{part}$ is the concentration of particle bound $^{230}$Th, $c_{\text{dis}}$ the concentration of dissolved $^{230}$Th, $c_{\text{part}}$ the equilibrium concentration of particle bound $^{230}$Th. The first-order rate constant $K$ (in [yr⁻¹]) is set here to $10^4$ yr⁻¹ thus assuming a quasi-instantaneous equilibration. Concentration of suspended particulate material is represented by $M$. For the partitioning coefficient $k_d$ between the dissolved and particle attached phases of $^{230}$Th we follow the formulation of (Honeyman et al., 1988), which accounts implicitly for the changing reactive surfaces of particles with particle size:

$$\log_{10} k_d = A + B \cdot \log_{10} M$$

where $M$ is the particle concentration (here taken in mg particles per liter) and $A$ as well as $B$ are tunable parameters. In addition to $^{230}$Th we carry also $^{231}$Pa as well as $^{10}$Be in our model (see Heinze et al. (2006)), but focus here on $^{230}$Th only. For each of these radionuclides, the preferential carrier phase can be selected in a dedicated switchboard (the particle masses of the different particle species are then cumulatively summed up for computing the equilibrium between particle attached and dissolved $^{230}$Th phases using the respective $k_d$ value). For $^{230}$Th scavenging, we used here CaCO₃, POC, and clay as carrier phases. The clay flux is computed according to the modern dust deposition from Mahowald et al. (1999) assuming that respective clay particles are chemically quasi-inert in seawater. Particle bound $^{230}$Th phases never get to zero, even if all biogenic particles
may get degraded, because at each grid point there is – at least a tiny – dust flux consisting of inert clay. Therefore, particle concentrations as such never go to zero (a minimum concentration could be prescribed, but this was not necessary in our case).

### 4 Control run and scenario experiments

The model was spun-up re-starting from an earlier integration and computed into quasi-equilibrium including the sediment over 40,000 years. The equilibrium coefficient $k_d$ was determined through a fit to observations of dissolved $^{230}$Th taken from the literature (Bacon and Anderson, 1982; Bacon et al., 1989; Chase et al., 2002; Cochran et al., 1995; Cochran et al., 1987; Colley et al., 1995; Guo et al., 1995; Huh and Beasley, 1987; Moore, 1981; Moran et al., 1997; Moran et al., 1995; Nozaki and Horibe, 1983; Nozaki and Yang, 1987; Nozaki et al., 1987; RoyBarman et al., 1996; Scholten et al., 1995; Vanderloeff and Berger, 1993; Vogler et al., 1998) and combined with the data as given in the GEOTRACES Intermediate Data Product Version 3 (Mawji et al. (2015), where especially the southern hemisphere data by Hein de Baar and Edward Boyle have been used). Important global bulk numbers and the parameters for the partitioning coefficient $k_d$ resulting from an optimal linear fit to observed data are listed in Table 1.

<table>
<thead>
<tr>
<th>Key variable as simulated</th>
<th>Control run value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Atmospheric CO$_2$ mixing ratio</td>
<td>281.9 ppm</td>
</tr>
<tr>
<td>POC export production (pre-industrial)</td>
<td>9.76 Gt C yr$^{-1}$</td>
</tr>
<tr>
<td>CaCO$_3$ export production (pre-industrial)</td>
<td>1.16 Gt C yr$^{-1}$</td>
</tr>
<tr>
<td>Coefficient A for $k_d$</td>
<td>- 6.39</td>
</tr>
<tr>
<td>Coefficient B for $k_d$</td>
<td>- 0.4</td>
</tr>
</tbody>
</table>

Table 1: Summary of control run results.

Meridional sections of dissolved $^{230}$Th concentrations for the Atlantic and Pacific Oceans are given in Figure 3. The vertical distribution clearly shows the increase of concentrations with depth due to the downward transfer of $^{230}$Th with the marine particle flux. A comparison of simulated and observed dissolved $^{230}$Th values in the water column (Figure 4) indicates that the model is in a good agreement with observations regarding vertical distributions of $^{230}$Th in selected locations. The standard-run export production rates for POC and CaCO$_3$ are given as maps in Figure 5.

We carried out a total of 12 different model projections under three different scenarios for changes in calcification (deviating from the control run with constant calcification) and four different scenarios of the future development of the atmospheric CO$_2$ concentration. All scenarios were restarted from the same previously performed spin-up model run reflecting a preindustrial biogeochemical state of the ocean. Throughout the experiments, the ocean circulation field was not changed. The model was integrated during the calendar years 1700-2300 spanning the time period of 600 years. In the model spin-up simulation the atmospheric CO$_2$ concentration was a prognostic variable. For the computations from 1700 onwards, we prescribed the atmospheric CO$_2$ concentration according to the Representative Concentration Pathways (RCPs) including their extension to year 2300 (van Vuuren et al., 2011) as used in CMIP5 (Coupled Model Intercomparison Project Phase 5). Our simulations follow scenarios RCP 2.6, RCP 4.5, RCP 6.0, and RCP8.5 designed to spanning a
range of radiative forcing between 2.6 and 8.5 W/m² by the year 2100. The respective atmospheric CO₂ concentrations are shown in Figure 6. For the decrease in calcification with decreasing pH as well as carbonate saturation, functional relationships between CaCO₃ export production and carbonate saturation were chosen which correspond to the respective experiments in Ilyina et al. (2009). For each RCP, we carried out a control simulation with constant calcification, a moderate decrease of CaCO₃ production with decreasing saturation, a linear dependency simulation with respectively stronger decrease in CaCO₃ production) and an extreme scenario (Figure 2). To date, no clear bulk formulation for the dependency of CaCO₃ export on CaCO₃ saturation exists. Therefore, the scenarios carried out here are only sensitivity experiments and not overall exhaustive simulations in order to reproduce the entire range of possible changes in ocean biogeochemistry.

Figure 3: Meridional ²³⁹Th cross sections in [dpm/1000l] (dpm = disintegration per minute) for the model control run. (a) Dissolved, Atlantic. (b) Dissolved, Pacific. (c) Particle attached, Atlantic. (d) Particle attached, Pacific.
Figure 4: Comparison of dissolved $^{230}$Th model results with observations (for references of observations see text). (a) Correlation between observations and modelled values. (b) Location of all vertical profiles as used in this work. (c)-(f) Selected single profiles (locations marked orange in the map (b); observations in blue, model results in red): (c) Western North Atlantic (obs. data from Colley et al., 1995). (d) Western North Pacific (obs. data from Nozaki and Yang, 1987). (e) South Pacific, Southern Ocean (obs. data from Chase et al., 2002). (f) Sargasso Sea (obs. data from Cochran et al., 1995). These profiles and locations were chosen to illustrate the model performance for typical profiles at a variety of different oceanic regions.

Figure 5: Model control run results for (a) biological export production of particulate organic carbon and (b) CaCO$_3$ export.
5 Results and discussion

According to the various scenarios applied considerable changes in surface alkalinity (not shown) and corresponding changes in CaCO$_3$ export production evolve at high CO$_2$ (Figure 7). The effect is on the average larger in the Pacific than in the Atlantic due to the overall higher modelled biological production rates of CaCO$_3$ in the Pacific Ocean. The extreme calcification scenario would lead to vastly reduced CaCO$_3$ export production after year 2100 in all RCPs except the most moderate RCP2.6 where even a partial recovery occurs.

For dissolved $^{230}$Th, we show here time series for depth levels 700 m, 2000 m, and 4000 m (Figures 8, 9, and 10, respectively). According to recent intercalibration experiments, still sizable discrepancies exist between the absolute values in $^{230}$Th measurements from different laboratories, though these measurements show a smaller scatter for replicates within a single laboratory. Intercalibration experiments reveal a standard deviation for measurements on one sample by different laboratories of about 0.07-0.08 dpm/1000l (Anderson et al., 2012). We take 0.075 dpm/1000l as an approximate indicative value for the detection level for changes in dissolved $^{230}$Th concentrations in seawater in (Figures 8-10, see orange line therein). This may be a somewhat optimistic estimate for samples of past decades, but future developments of measuring techniques could possibly reduce measurement errors and the spread across analyses from different laboratories and application of different measurement methods. The detection level is shown here relative to the modelled preindustrial values. The time of emergence as indicated by the orange line in Figures 8-10 is, therefore, the earliest possible time of detection if preindustrial $^{230}$Th would be known. Figures 8-10 show the increase of dissolved $^{230}$Th activities with depth from 700 m (Figure 8), over 2000 m (Figure 9) to finally 4000 m (Figure 10). At 700 m the increase in $^{230}$Th activities due to the assumed ocean acidification effect is too small to unambiguously show an effect (Figure 8). This picture changes when going down to 2000 m, where an effect would be detectable within this century at least for the stronger forcing scenarios TCP8.5 and RCP6.0 (Figure 9). At 4000 m depth finally, the effect would be measurable quite soon for all RCPs and a moderate impact of ocean acidification on the calcification (Figure 7).
10. For constant CaCO$_3$ production, the intermediate and deep water $^{230}$Th activities start to rise around year 2100 as well (see black curves in Figures 8-10). This effect is due to the increasing dissolution of CaCO$_3$ particles in the water column in parallel with downward mixing of waters that carry anthropogenic loads of dissolved organic carbon and hence subsurface and deep acidification.

In order to look for most suitable regions for detection of ocean acidification effects on CaCO$_3$ particle fluxes using modern $^{230}$Th data, we plotted the year of emergence for different reference years and different assumed detection levels (Figures 11 and 12). The year of emergence is defined here as the earliest possible year of a potential detection in case that the $^{230}$Th signal (rate of change over time) is only influenced by a decrease in calcification and not any other processes (such as potential shifts in circulation and associated changes in biological particle production). We choose two different pairs of reference years, to which the $^{230}$Th activity at a later stage would be compared to, and two different limits for the detection of a signal: (1) Year 2000 and 0.075 dpm/1000l and (2) year 2020 and 0.0325 dpm/1000l. Case (1) should reflect high quality observations of the first decade of the 21st century while case (2) should reflect hopefully further increased precision in the measurements in the few years to come next. We first looked at these two cases for the moderate calcification scenario (Figure 11). The deeper one goes down in the water column the younger the year of emergence becomes. For case (1) with reference year 2000 and detection limit 0.075 dpm/1000l earliest years would be around 20 years from now at 4000 m depth in the North Pacific (Figure 11 a,b). For case (2) even some earlier thresholds would occur where the shorter difference to the reference year would be more than made up by a higher precision of the measurements (Figure 11 c,d).

For the extreme calcification impact scenario, earlier strong changes in calcification and $^{230}$Th activity occur and suitable detection limits would be available over large parts of the Pacific Ocean and in general at southern high latitudes latest by year 2025 for case (1) (Figure 12 a,b). For newer measurements to come (reference year 2020 and detection limit 0.0325 dpm/1000l, case (2)), the earliest years of emergence are a bit later than for case (1) in contrast to the situation for the moderate calcification scenario. This behaviour can be explained by the lower baseline level for CaCO$_3$ production at year 2020 where it is already strongly reduced as compared to the preindustrial values. But still large parts of the Pacific Ocean would have an earlier detection level than 2025. As compared to the detection approach for ocean acidification impacts through total alkalinity measurements as pursued by Ilyina et al. (2009) the $^{230}$Th method presented here results in similar time of emergence of a signal reflecting changes in biocalcification. While for alkalinity, surface measurements would primarily be involved, for $^{230}$Th one would use deep ocean measurements. For very deep depth levels below 4000 m and very high precision measurements of $^{230}$Th one could presumably shorten the time of emergence. Possibly, for the deep measurements some averaging effect over the water column would occur and results from a few stations may be representative for a larger area. The deep Pacific Ocean and to some degree the deep Southern Ocean look as the best areas for detecting ocean acidification impacts through $^{230}$Th. The dust deposition onto the Atlantic Ocean surface originating from mobilisation in the Saharan desert – and the hence strong clay particle flux – blur the imprint of ocean acidification induced changes in CaCO$_3$ flux onto $^{230}$Th.
Figure 7: Temporal evolution of Atlantic (left column) and Pacific (right column) annual mean CaCO$_3$ export production under the different scenarios for the sensitivity of calcification under high CO$_2$ (unit: GtC yr$^{-1}$). From top to bottom for greenhouse gas scenarios RCP8.5, RCP 6.0, RCP4.5, and RCP2.6.
Figure 8: Time series for the evolution of mean Atlantic (left column) and Pacific (right column) dissolved $^{230}\text{Th}$ concentrations at 700 m under the different scenarios for reduction of calcification under high CO$_2$ (unit: dpm/1000l). From top to bottom for greenhouse gas scenarios RCP8.5, RCP 6.0, RCP4.5, and RCP2.6. The orange line indicates the theoretical detection limit for changes with respect to the pre-industrial.
Figure 9: Time series for the evolution of mean Atlantic (left column) and Pacific (right column) dissolved $^{230}$Th concentrations at 2000 m under the different scenarios for reduction of calcification under high CO$_2$ (unit: dpm/1000l). From top to bottom for greenhouse gas scenarios RCP8.5, RCP 6.0, RCP4.5, and RCP2.6. The orange line indicates the theoretical detection limit for changes with respect to the pre-industrial.
Figure 10: Time series for the evolution of mean Atlantic (left column) and Pacific (right column) dissolved $^{230}$Th concentrations at 4000 m under the different scenarios for reduction of calcification under high CO$_2$ (unit: dpm/1000l). From top to bottom for greenhouse gas scenarios RCP8.5, RCP 6.0, RCP4.5, and RCP2.6. The orange line indicates the theoretical detection limit for changes with respect to the pre-industrial.
Figure 11: Prospective year of emergence for an ocean acidification induced signal in dissolved $^{230}$Th activity as derived from the model runs. Shown are the calendar years of emergence for the depth levels 2000m and 4000m. All figures are shown for the strong RCP8.5 scenario concerning atmospheric CO$_2$ concentration and the moderate scenario of calcification decrease with saturation state. (a) For year 2000 as reference year for $^{230}$Th activity and 0.075 dpm/1000l as analytical threshold between different samples, depth level 2000 m. (b) Same as (a) but for depth level 4000 m. (c) For 2020 as reference year for $^{230}$Th activity and 0.0325 dpm/1000l as analytical threshold between different samples, depth level 2000 m. (d) The same as (c) but for depth level 4000 m.

The probably most important limitation of the approach here is the fixed velocity field which does not vary with seasons, interannual variability, climatic variability modes (such as El Niño Southern Oscillation or North Atlantic Oscillation). In reality, the distribution of $^{230}$Th depends also on variations in the flow field and resulting changes in nutrient availability, and consequently changes in particle production as well as flux. Also the effect of boundary scavenging (Anderson et al., 1983; Roy-Barman et al., 2009), i.e. the transport of dissolved $^{230}$Th from areas of low particle concentrations to those with high particle concentrations (especially at ocean margins at shelf seas) is not spatially resolved in our model. Our results concerning the time of emergence may, therefore, be somewhat optimistic. On the other hand, given a larger amount of vertical observed $^{230}$Th profiles, still a common effect due to reduced CaCO$_3$ production should be detectable through statistical analysis even in the presence of shorter term natural variability in circulation and particle fluxes. Further uncertainties are associated with the choice of the $k_d$ values and particle specific reactivity. Concerning the latter, another study did not reveal a substantial change in results, if different combinations of particle specific scavenging were applied in the model (Heinze et al., 2006).
Figure 12: Prospective year of emergence for an ocean acidification induced signal in dissolved $^{230}$Th activity as derived from the model runs. Shown are the calendar years of emergence for the depth levels 2000m and 4000m. All figures are shown for the strong RCP8.5 scenario concerning atmospheric CO$_2$ concentration and the extreme scenario on calcification.

(a) For year 2000 as reference year for $^{230}$Th activity and 0.075 dpm/1000l as analytical threshold between different samples, depth level 2000 m. (b) Same as (a) but for depth level 4000 m. (c) For 2020 as reference year for $^{230}$Th activity and 0.0325 dpm/1000l as analytical threshold between different samples, depth level 2000 m. (d) The same as (c) but for depth level 4000 m.

The extreme scenario on pH-dependent CaCO$_3$ production is likely to be an overestimation. According to the results from this study and Ilyina et al. (2009) a respective large change in the real world would probably have been detected through the alkalinity signal and other methods such as remote sensing and sediment trap measurements already. Also, the reaction of CaCO$_3$ shell producing organisms to sinking carbonate saturation levels is not a simple function and can vary between taxa (Kroeker et al., 2013).

The detection method for ocean acidification impacts through $^{230}$Th would work best, if the preindustrial values for dissolved oceanic $^{230}$Th activities could reliably be reconstructed. From core top samples and corals one could possibly determine whether the particle attached $^{230}$Th activities and hence those of the dissolved fraction would have undergone any variations over the last centuries and millennia. However, the overall uncertainties associated with the formation and analysis of the paleo-record values may be too high to provide an accurate enough baseline for comparisons with modern water column measurements.
6 Conclusions

In this study, we investigated the potential of the particle reactive radionuclide $^{230}$Th for detection of reduced calcification by biota due to progressing ocean acidification. The time of emergence of a dissolved $^{230}$Th activity signal with respect to a reference year and reference activity is about the same as for detection approaches employing alkalinity. Nevertheless, regular selected reoccupations of a series of deep stations in the Pacific Ocean and Southern Ocean with highest quality $^{230}$Th measurements would be helpful to accompany alkalinity measurements that are easier to do in order to see whether both tracers give consistent results. Surface alkalinity measurements include signals of natural variability on seasonal and multiyear scales. Likewise, changes in ocean circulation and changes in biological particle production due to other processes than ocean acidification may lead to changes in the marine $^{230}$Th distribution (and alkalinity). As the deep ocean is less prone to effects of natural variability and the quality of observations does not change with depth, deep ocean observations of $^{230}$Th could be advantageous for monitoring and detecting ocean acidification effects on calcification. Furthermore, substantial improvements in the overall precision of water column $^{230}$Th measurements could potentially lead to earlier detection years than using alkalinity measurements. In any case, $^{230}$Th represents a fascinating magnifying glass for changes in ocean surface processes seen through the deep ocean signal. Its potential has not yet been fully exploited.

Acknowledgment: This work was partially supported through the “European Project on Ocean Acidification” EPOCA and project “Changes in Carbon Uptake and Emissions by Oceans in a Changing Climate” CARBOCHANGE which received funding from the European Community’s Seventh Framework Programme (FP7) under grant agreements no. 211384 and no. 264879 respectively. The Research Council of Norway supported this study through project “Overturining circulation and its implications for global carbon cycle in coupled models” (ORGANIC; grant no. 239965) and the nationally coordinated project “Earth system modelling of climate variations in the Anthropocene” (EVA; grant no. 229771). This is a contribution to the Bjerknes Centre for Climate Research (Bergen, Norway). A part of the computations were carried out under project NN2980K at the Norwegian Metacentre for Computational Science (NOTUR) and its dedicated storage and archiving project NorStore (NS2980k). This is an in-kind contribution to the GEOTRACES project. We would like to thank the GEOTRACES scientists for the interesting discussions during several workshops and for the fabulous new observational data.

References:


Heinze, C., Maier-Reimer, E.: Age offsets among different biogenic and lithogenic components of sediment cores revealed by numerical modeling, Paleoceanography, 24, 2009.


