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Processes determining the marine alkalinity and calcium carbonate saturation state distributions

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Abstract. We introduce a composite tracer for the marine system, Alk*, that has a global distribution primarily determined by CaCO₃ precipitation and dissolution. Alk* is also affected by riverine alkalinity from dissolved terrestrial carbonate minerals. We estimate that the Arctic receives approximately twice the riverine alkalinity per unit area as the Atlantic, and 8 times that of the other oceans. Riverine inputs broadly elevate Alk* in the Arctic surface and particularly near river mouths. Strong net carbonate precipitation results in low Alk* in subtropical gyres, especially in the Indian and Atlantic oceans. Upwelling of dissolved CaCO₃-rich deep water elevates North Pacific and Southern Ocean Alk*. We use the Alk* distribution to estimate the variability of the calcite saturation state resulting from CaCO₃ cycling and other processes. We show that regional differences in surface calcite saturation state are due primarily to the effect of temperature differences on CO₂ solubility and, to a lesser extent, differences in freshwater content and air-sea disequilibria. The variations in net calcium carbonate cycling revealed by Alk* play a comparatively minor role in determining the calcium carbonate saturation state.

1 Introduction

Our goal is to use high-quality total alkalinity (A_T) observations to examine the effects of calcium carbonate cycling on marine A_T and calcium carbonate saturation states. This study is motivated in part by ocean acidification. With marine calcite saturation states decreasing due to anthropogenic carbon uptake (Orr et al., 2005), it is important to understand

the degree to which carbonate cycling impacts the calcium carbonate saturation state.

Carbonate saturation state is a measure of how supersaturated seawater is with respect to a given mineral form of calcium carbonate. It is expressed for calcite as the ratio Ω_C between the product of Ca²⁺ and CO₃²⁻ ion concentrations and the calcite thermodynamic equilibrium solubility product. Values of Ω_C greater than 1 indicate calcite precipitation is favored thermodynamically over calcite dissolution, and the reverse is true for values less than 1.

Marine calcium carbonate cycling includes both internal and external calcium carbonate sources and sinks. Internal cycling refers to net formation of $67-300 \,\mathrm{T}\,\mathrm{mol}\,A_{\mathrm{T}}\,\mathrm{yr}^{-1}$ worth of calcium carbonate in the surface ocean (Berelson et al., 2007) and net dissolution of most of this calcium carbonate at depth. External marine carbonate cycling refers to inputs of carbonate minerals dissolved in rivers, sediment pore waters, hydrothermal vent fluids, and submarine groundwater discharge, as well as to loss due to biogenic carbonate mineral burial and authigenic mineralization in sediments. Rivers add 33 T mol $A_{\rm T}$ yr⁻¹ worth of dissolved bicarbonate to the ocean (Cai et al., 2008). Wolery and Sleep (1988) estimate that hydrothermal vents add an additional 6.6 T mol $A_{\rm T}$ yr⁻¹, though deVilliers (1998) argues the hydrothermal contribution may be as high as $30 \text{ T} \text{ mol } A_{\text{T}} \text{ yr}^{-1}$. Submarine groundwater discharge is poorly constrained, but it is thought to exceed riverine inputs in some areas (Moore, 2010).

We investigate calcium carbonate cycling using the global AT distribution in a dataset we created by merging the Global Data Analysis Project (GLODAP), Carbon in the North Atlantic (CARINA), and Pacific Ocean Interior Carbon (PACIFICA) discrete data products (Key et al., 2004, 2010; Velo et al., 2009; Suzuki et al., 2013). We have combined and gridded these data products using methods detailed in Supplement document SA. We use our gridded data set in our calculations to limit sampling biases and to enable us to make volume-weighted mean property estimates.

Dickson (1981) defines total alkalinity as the concentration excess "of proton acceptors formed from weak acids (pK \leq 4.5) relative to proton donors (weak bases with pK > 4.5)" at a reference temperature, pressure, and ionic strength. $A_{\rm T}$ can be thought of as a measure of how well-buffered seawater is against changes in pH. This operational definition gives $A_{\rm T}$ (expressed in mol kg⁻¹) several properties that make it an especially useful carbonate system parameter for examining carbonate cycling:

- 1. It mixes conservatively
- 2. ... and is therefore diluted and concentrated linearly by evaporation and precipitation.
- It responds in predictable ways to calcium carbonate cycling
- 4. ... as well as organic matter formation and remineralization.
- 5. It is not changed by air-sea exchange of heat or carbon dioxide.
- 6. It is, however, affected by anaerobic redox reactions (Chen, 2002).

We are primarily interested in calcium carbonate cycling, item 3 in our list. In Sect. 2 of this paper we therefore define a tracer we call Alk^{*} that removes the majority of the influences of organic matter cycling (item 4), freshwater cycling (item 2), and non-sedimentary anaerobic redox reactions (item 6) while still mixing conservatively, remaining insensitive to gas exchange, and responding to calcium carbonate cycling. In Sect. 3 we discuss processes that govern the Alk^{*} distribution globally, by ocean basin, and regionally. In Sect. 4 we define a metric to quantify the influence of various processes on the marine calcite saturation state. We use this metric with our gridded data set and Alk^{*} to determine the relative importance of the various controls on calcite saturation state in the ocean and at the ocean surface. We summarize our findings in Sect. 5.

2 The Alk* tracer

In defining Alk^{*}, we take advantage of the potential alkalinity (Brewer et al., 1975) concept to remove the majority of the influence of organic matter cycling and denitrification, and use a specific salinity normalization scheme (Robbins, 2001) to remove the influence of freshwater cycling. We detail the Alk^{*} definition and the reasoning behind it in this section.

The influence of organic matter cycling on $A_{\rm T}$ is due primarily to the biologically driven marine nitrogen cycle. Nitrate uptake for anaerobic denitrification and the production of amino acids occurs in a $\sim 1:1$ mole ratio with the release of molecules that increase $A_{\rm T}$ (Chen, 2002). Similarly, nitrate from fixation of nitrogen gas and remineralization of amino nitrogen is released in a 1:1 mole ratio with acids that titrate away A_T (Wolf-Gladrow et al., 2007). This observation led Brewer et al. (1975) to propose the idea of "potential alkalinity" as the sum of $A_{\rm T}$ and nitrate, with the aim of creating a tracer that responds to the cycling of calcium carbonates without changing in response to organic matter cycling. Feely et al. (2002) since used a variant that relies on the empirical relationship between dissolved calcium concentrations, A_T, and nitrate determined by Kanamori and Ikegami (1982). This variant has the advantage of implicitly accounting for the $A_{\rm T}$ changes created by the exchange of numerous other components of marine organic matter besides nitrate (e.g., sulfate and phosphate). We thus use the ratio found by Kanamori and Ikegami (1982) to define potential alkalinity $(A_{\rm T})$.

$$A_{\rm P} = A_{\rm T} + 1.26 \times [\rm NO_3^-] \tag{1}$$

While the empirical Kanamori and Ikegami (1982) ratio of 1.26 may be specific to the elemental ratios of the North Pacific, Wolf-Gladrow et al. (2007) provide a theoretical derivation from Redfield ratios and obtain a similar value of 1.36.

The sensitivity of the $A_{\rm T}$ distribution to freshwater cycling is due primarily to the dilution or concentration of the large background $A_{\rm T}$ fraction that does not participate in carbonate cycling on timescales of ocean mixing. This background fraction behaves conservatively, so we call it conservative potential alkalinity ($A_{\rm P}^{\rm C}$) and estimate it directly from salinity as

$$A_{\rm P}^{\rm C} \equiv S \frac{\overline{A_{\rm P}}}{\overline{S}}.$$
 (2)

Here, terms with a bar are reference values chosen as the mean value for those properties in the top 20 m of the ocean. We obtain a volume-weighted surface $\overline{A_P}$ (2305 µmol kg⁻¹) to \overline{S} (34.71) ratio of 66.40 µmol kg⁻¹ from our gridded data set. The mean surface values are chosen in an effort to best capture the impact of freshwater cycling where precipitation and evaporation occur.

Robbins (2001) showed that subtracting an estimate of the conservative portion of a tracer, such as $A_{\rm P}^{\rm C}$, produces a salinity-normalized composite tracer that mixes conservatively. This scheme also retains the 2 : 1 change of $A_{\rm T}$ to dissolved inorganic carbon ($C_{\rm T}$) with carbonate cycling. We follow this approach in our definition of Alk*. In Supplement document SB we estimate this approach removes 97.5% of the influence of freshwater cycling on potential alkalinity and reduces the influence of freshwater cycling on Alk* to less than 1% of the Alk* variability. In Supplement document SC we demonstrate that Alk* mixes conservatively, and we briefly contrast Alk^{*} to traditionally normalized potential alkalinity which does not mix conservatively (Jiang et al., 2014).

In total, we define Alk^{*} as the deviation of potential alkalinity from A_{P}^{C} :

$$Alk^* \equiv A_P - A_P^C \tag{3}$$

$$\equiv A_{\rm P} - \frac{A_{\rm P}}{\overline{S}}S\tag{4}$$

$$\equiv A_{\rm P} - 66.4 \times S,\tag{5}$$

where Alk^{*} has the same units as A_T (µmol kg⁻¹). The Alk^{*} distribution is attributable primarily to carbonate cycling plus the small (in most places) residual variation due to freshwater cycling that is not removed by subtracting A_P^C . However, hydrothermal vent fluid and non-denitrification anaerobic redox chemistry may substantively affect alkalinity distributions in certain marine environments, and Alk^{*} distributions could not be attributed purely to internal and external calcium carbonate cycling in these locations.

Mean global surface Alk* is 0 by definition, and thus Alk* can have negative as well as positive values. For reference, more than 95% of our gridded Alk* data set falls between -35 and $220 \,\mu\text{mol}\,\text{kg}^{-1}$. Comparing gridded Alk* to Alk* from measurements suggests a standard disagreement of order $10 \,\mu\text{mol}\,\text{kg}^{-1}$. We adopt this number as an estimate of standard gridded Alk* error despite noting there are reasons to suspect that this value could be either an underestimate (correlated errors) or an overestimate (we are directly comparing instantaneous point measurements to estimates for annual averages for a grid cell).

3 Alk* distributions

We consider Alk* distributions globally, by ocean basin, and regionally in the context of sources and sinks of the tracer both globally and regionally. We pay special attention to riverine Alk* because it is easily identified where it accumulates near river mouths.

3.1 Global distribution of Alk*

Figure 1 maps surface Alk^{*} (top 50 m) at the measurement stations. We provide this figure to show where we have viable Alk^{*} estimates and to demonstrate that our gridded data product adequately captures the measured Alk^{*} distribution. Figure 2 maps gridded global surface A_T , salinity, Alk^{*}, and phosphate distributions and masks the regions that are lacking data in Fig. 1.

The similarity of the A_T (Fig. 2a) and salinity (Fig. 2b) distributions demonstrates the strong influence of freshwater cycling on the surface marine A_T distribution (see also Millero et al., 1998; Jiang et al., 2014). The dissimilarity between Alk* (Fig. 2c) and salinity (Fig. 2b) suggests Alk* removes the majority of this influence. The phosphate (Fig. 2d) and



Figure 1. A map of station locations at which we use measurements to estimate Alk^{*} (in μ mol kg⁻¹). Dot color indicates surface Alk^{*}. Points with black borders indicate either that AT was measured prior to 1992 (i.e., before reference materials were commonly used) or that no nitrate value was reported (in which case a nitrate concentration of 5 μ mol kg⁻¹ is assumed). Red dots on land indicate the mouth locations and mean annual discharge volumes (indicated by dot size) of 200 large rivers, as given by Dai and Trenberth (2002).

Alk^{*} (Fig. 2c) distributions are similar at the surface. They are also similar at depth: Figs. 3 and 4 show zonally averaged gridded depth sections of Alk^{*} and phosphate. Alk^{*} and phosphate concentrations are low in the deep Arctic Ocean (Figs. 3d, and 4d), intermediate in the deep Atlantic Ocean (Figs. 3a and 4a), and high in the deep North Pacific (Figs. 3b and 4b) and deep northern Indian (Figs. 3c and 4c) oceans. Alk^{*} and phosphate distributions are similar because similar processes shape them: the hard and soft tissue pumps transport A_T and phosphate, respectively, from the surface to depth. The "oldest" water therefore has the highest net phosphate and Alk^{*} accumulation. High surface phosphate and Alk^{*} in the Southern Ocean and North Pacific in Figs. 2, 3, and 4 are due to upwelled old, deep waters.

Several qualitative differences between Alk* and phosphate distributions are visible in Figs. 2c, 2d, 3, and 4. Surface phosphate is low in the Bay of Bengal and high in the Arabian Sea (Fig. 2d), while the opposite is true for Alk^{*} (Fig. 2c). Also, Alk* reaches its highest surface concentration in the Arctic (Figs. 2c and 3d) where phosphate is not greatly elevated (Figs. 2d and 4d). These surface differences are due to regional riverine Alk* inputs (Sect. 3.3). Another difference is that Alk* reaches a maximum below 2000 m in all ocean basins except the Arctic, while phosphate maxima are above 2000 m. We attribute the deeper Alk* maxima to deeper dissolution of calcium carbonates than organic matter remineralization. Finally, Alk* values are higher in the deep Indian Ocean than in the deep Pacific. This is likely due to elevated biogenic carbonate export along the coast of Africa and in the Arabian Sea (Sarmiento et al., 2002; Honjo et al., 2008).



Figure 2. Global (a) total alkalinity $A_{\rm T}$, (b) salinity, (c) Alk^{*}, and (d) phosphate distributions at the surface (10 m depth surface) from our gridded CARINA, PACIFICA, and GLODAP bottle data product detailed in Supplement document SA. Areas with exceptionally poor coverage in the data used to produce the gridded product are blacked out.



Figure 3. Zonal mean gridded Alk^{*} (in μ mol kg⁻¹) in the (**a**) Atlantic, (**b**) Pacific, (**c**) Indian, and (**d**) the Arctic oceans plotted against latitude and depth.

3.2 Alk* by ocean basin

In Fig. 5 we provide 2-D color histograms of discrete surface Alk* and salinity measurements for the five major ocean basins. Figure 5 also indicates a single volume-weighted

mean gridded Alk* for each basin (in writing). We attribute the decrease in Alk* as salinity increases – especially visible in the low-salinity bins in the Arctic Ocean (Fig. 5d) to mixing between high-Alk*, low-salinity river water and low-Alk*, high-salinity open-ocean water. Net precipitation in the tropics and net evaporation in the subtropics widens the histograms across a range of salinities and alkalinities without affecting Alk* in Fig. 5a, b, and c. The Alk* elevation associated with upwelled water is most visible in Fig. 5e where Upper Circumpolar Deep Water upwelling near the Polar Front results in high-frequency (i.e., warm colored) histogram bins at high-Alk*. Similarly, the high-frequency Alk* bins in Fig. 5b with salinity between 32.5 and 33.5 are from the North Pacific Subpolar Gyre, and are due to upwelled old, high-Alk* water (cf. the Si* tracer in Sarmiento et al., 2004). River water contributions can be most easily seen in a scattering of low-frequency (cool colored), high-Alk* as well as low-salinity bins in the Arctic Ocean.

The surface Southern Ocean has the highest Alk* followed by the Arctic and the Pacific. The Indian and Atlantic have similar and low mean Alk*. The high mean Southern Ocean Alk* is due to upwelling. The high mean Arctic surface Alk* is due to riverine input. The Atlantic and the Arctic together receive ~ 65 % of all river water (Dai and Trenberth, 2002). We construct a budget for terrestrial $A_{\rm T}$ sources to the various surface ocean basins using the following assumptions:

- 1. the $A_{\rm T}$ of 25 large rivers are as given by Cai et al. (2008);
- 2. the volume discharge rates of 200 large rivers are as given by Dai and Trenberth (2002);
- 3. groundwater and runoff enter each ocean in the same proportion as river water from these 200 rivers;



Figure 4. Zonal mean gridded phosphate (in μ mol kg⁻¹) in the (a) Atlantic, (b) Pacific, (c) Indian, and (d) the Arctic oceans plotted against latitude and depth.

- 4. the $A_{\rm T}$ of all water types that we do not know from assumption 1 is the 1100 µmol kg⁻¹ global mean value estimated by Cai et al. (2008);
- 40° N is the boundary between the Atlantic and the Arctic, and 40° S is the boundary between the Southern and the Atlantic oceans (based upon the region of elevated surface phosphate in Fig. 2d).

Our detailed budget is provided as Supplement file SD. We estimate 40% of continentally derived $A_{\rm T}$ enters the Atlantic, 20% enters the Arctic, and 40% enters all remaining ocean basins. These ocean areas represent 17, 5, and 78 % of the total surface ocean area in our gridded data set, respectively, so the Arctic receives approximately twice as much riverine $A_{\rm T}$ per unit area as the Atlantic, and 8 times the rest of the world ocean. The Atlantic has the lowest openocean surface Alk* value and low basin mean surface Alk* despite the large riverine sources. The large riverine $A_{\rm T}$ input must therefore be more than balanced by strong net calcium carbonate formation. The Indian Ocean has comparably low mean surface Alk* to the Atlantic, but a smaller riverine source. Mean Alk* is higher in the Pacific than the Atlantic and Indian, even when neglecting the region north of 40° N as we do for the Atlantic (Alk^{*} = -16.5 µmol kg⁻¹ when omitted vs. $-22.9 \,\mu mol \, kg^{-1}$ for the Atlantic and $-22.2 \,\mu\text{mol}\,\text{kg}^{-1}$ for the Indian). The difference between the



Figure 5. 2-D histograms indicating the log (base 10) of the number of measurements that fall within bins of Alk* vs. salinity with color. Data are limited to the top 50 m of the (a) Atlantic, (b) Pacific, (c) Indian, (d) Arctic, and (e) Southern oceans. Where basins connect, the boundary between the Atlantic and the Arctic oceans is 40° N, between the Atlantic and the Indian it is 20° E, between the Indian and the Pacific it is 131° E, between the Pacific and the Atlantic it is 70° W, and between the Southern Ocean and the other oceans it is 40° S.

Pacific and the other basins is significant when considering the large number of grid cell Alk^{*} values averaged (> 6000 in the Atlantic), and the small estimated uncertainty for each value ($\sim 10 \,\mu\text{mol}\,\text{kg}^{-1}$). Considering the weak Pacific riverine input, this suggests that, relative to other ocean basins, there are either larger Alk^{*} inputs from exchange with other



Figure 6. Alk^{*} distributions (in μ mol kg⁻¹) (**a**) between 5° and 30° N in the Red and Arabian seas shown against longitude, and (**b**) between 75° and 100° E in the Bay of Bengal plotted against latitude. Small black dots indicate where data are present. The inverted triangle above (**a**) indicates the longitude of the mouth of the Red Sea.

basins and deeper waters or smaller Pacific basin mean net calcium carbonate formation.

3.3 Riverine Alk* regionally

For river water with negligible salinity, Alk^{*} equals the potential alkalinity. This averages around $1100 \,\mu\text{mol}\,\text{kg}^{-1}$ globally (Cai et al., 2008) but is greater than $3000 \,\mu\text{mol}\,\text{kg}^{-1}$ for some rivers (Beldowski et al., 2010). Evidence suggests that riverine $A_{\rm T}$ is increasing due to human activities (Kaushal et al., 2013).

The most visible riverine Alk* signals are in the Arctic due to the large riverine runoff into this comparatively small basin and the confinement of this low-density riverine water to the surface (Jones et al., 2008; Yamamoto-Kawai et al., 2009; Azetsu-Scott et al., 2010). Figure 3d shows the high Arctic Alk* plume is confined to the top ~ 200 m. Figure 1 shows that these high Alk* values extend along the coast of Greenland and through the Labrador Sea. Alk* decreases with increasing salinity in this region (Fig. 5d) due to mixing between the fresh high-Alk* surface Arctic waters and the salty, lower-Alk* waters of the surface Atlantic. Gascard et al. (2004a, b) suggest that waters along the coast of Norway are part of the Norwegian Coastal Current and originate in the Baltic and North seas where there are also strong riverine inputs (Thomas et al., 2005).

Elevated Alk^{*} can also be seen in the Bay of Bengal, with surface values ~ 100 µmol kg⁻¹ higher than those in the central Indian Ocean. This bay has two high- A_T rivers that join and flow into it, the Brahmaputra ($A_T = 1114 \mu mol kg^{-1}$) and the Ganges ($A_T = 1966 \mu mol kg^{-1}$) (Cai et al., 2008). Figure 6b provides provides an Alk^{*} depth section for this region. The riverine Alk^{*} plume can be clearly seen in the top 50 m. No similar increase is seen in the Arabian Sea (Fig. 6a), where the Indus River (1681 µmol kg⁻¹) discharges only ~ 1/10th of the combined volume of the Brahmaputra and the Ganges.



Figure 7. Alk^{*} (in μ mol kg⁻¹) in top 50 m of the ocean near the Amazon River outflow plotted in color, though with a narrower color scale than is used for all other plots. Panel (**a**) is limited to data collected in November through January, and panel (**b**) is limited to measurements from May through July. Points with black borders indicate either that the AT was measured prior to 1992 (before reference materials were commonly used) or that no nitrate value was reported (in which case a nitrate concentration of 5 μ mol kg⁻¹ is assumed). Red dots on land indicate the mouth locations and mean annual discharge volumes (indicated by dot size) of large rivers, as given by Dai and Trenberth (2002).

The Amazon River is the largest single riverine marine $A_{\rm T}$ source. This river has low $A_{\rm T}$ (369 µmol kg⁻¹; Cai et al., 2008) but has the largest water discharge volume of any river, exceeding the second largest – the Congo – by a factor of \sim 5 (Dai and Trenberth, 2002). Consequently, the Amazon discharges approximately 50% more $A_{\rm T}$ per year than the river with the second-largest $A_{\rm T}$ discharge, the Changjiang (Cai et al., 2008). The Amazon's influence can be seen as a region of abnormally low salinity and A_{T} in Fig. 2a and b. Despite the high discharge volume, the influence is only barely visible as a region of elevated Alk* in Fig. 2c due to the comparatively low Amazon Alk*. However, the influence of the Amazon on Alk* can be seen in the seasonal Alk* cycle in the Amazon plume. Figure 7 provides a map of Alk* for this region scaled to show the influence of this low-Alk* river in the Northern Hemisphere (a) winter and (b) summer months. The higher Alk* found for summer months is consistent with Amazon discharge and AT seasonality (Cooley et al., 2007) and the radium-isotope-based finding of Moore et al. (1986) that Amazon River outflow comprises 20-34 % of surface water in this region in July compared to only 5-9% in December.

3.4 Regional abiotic carbonate cycling

The Red Sea portion of Fig. 6a is strongly depleted in Alk^{*} and contains the lowest single Alk^{*} measurement in our data set, $-247 \,\mu\text{mol}\,\text{kg}^{-1}$. The GEOSECS expedition Red Sea alkalinity measurements (Craig and Turekian, 1980) predate alkalinity reference materials (Dickson et al., 2007) but are supported by more recent measurements (Silverman et al., 2007). Like Jiang et al. (2014), we attribute low Red Sea Alk^{*} to exceptionally active calcium carbonate formation. Brewer and Dyrssen (1985) provide seawater chemistry measurements from the neighboring Persian Gulf that suggest strong calcium carbonate formation results in low Alk^{*} there as well ($< -240 \,\mu\text{mol}\,\text{kg}^{-1}$ along the Trucial Coast).

The Red Sea is one of the only regions where $\Omega_{\rm C}$ is sufficiently high for abiotic carbonate precipitation to significantly contribute to overall carbonate precipitation (Milliman et al., 1969; Silverman et al., 2007). Notably, saturation state remains high at depth in the Red Sea (see Sect. 4.2). In this region, biogenic aragonitic corals and pteropod shells are progressively removed with depth in sediments, and pores left behind are filled in with high-magnesium calcite cement (Gevirtz and Friedman, 1966; Almogi-Labin et al., 1986). We hypothesize biogenic carbonates are dissolved by CO_2 from sedimentary organic matter remineralization, as occurs elsewhere (e.g., Hales and Emerson, 1997; Hales, 2003; Boudreau, 2013), and that high deep Red Sea $\Omega_{\rm C}$ leads to abiotic re-calcification in sediment pores. Morse et al. (2006) find that synthetic high-magnesium calcite - unlike biogenic high-magnesium calcite - is less soluble than aragonite, so this substitution is favored thermodynamically if the abiotic mineral forms similarly to the synthetic mineral.

Calcium carbonate has recently been found as metastable ikaite (a hydrated mineral with the formula $CaCO_3 \times 6H_2O$) in natural sea ice (Dieckmann et al., 2008). Ikaite cycling provides a competing explanation for the high Arctic surface Alk* values if high-A_T, low-salinity, ikaite-rich ice melt becomes separated from low-A_T, high-salinity rejected brines. However, riverine A_{T} inputs better explain the magnitude of the feature: the \sim 5 mg ikaite L⁻¹ sea ice that Dieckmann et al. (2008) found in the Antarctic could only enrich $A_{\rm T}$ of the surface 100 m by \sim 1 µmol kg⁻¹ for each meter of ice melted, and Arctic surface 100 m Alk^{*} is elevated by 59 μ mol kg⁻¹ relative to the deeper Arctic in our gridded data set. By contrast, Jones et al. (2008) estimate a \sim 5 % average riverine end-member contribution to the shallowest 100 m of this region, which accounts for $\sim 55 \,\mu mol \, kg^{-1}$ Alk^{*} enrichment. Also, surface Alk* in the Southern Ocean - which has sea ice but lacks major rivers - is not similarly elevated relative to surface phosphate (Fig. 2) or deep Alk* (Fig. 3).

4 Controls on the calcite saturation state

The Alk^{*} tracer provides an opportunity to estimate the impact of carbonate cycling on $\Omega_{\rm C}$. In addition to (1) carbonate cycling, $\Omega_{\rm C}$ is affected by (2) organic matter cycling, (3) freshwater cycling, (4) pressure changes on seawater, (5) heating and cooling, and (6) $A_{\rm T}$ changes from nitrogen fixation and denitrification. For each of these six processes, we estimate the standard deviation of the net influence of the process globally by considering the standard deviation of a "reference" tracer R_i for the process, " σ_{R_i} ", where R_i is Alk^{*} for CaCO₃ cycling, phosphate for organic matter cycling, salinity for freshwater cycling, pressure for pressure changes, temperature for heating and cooling, and N^* (Gruber and Sarmiento, 1997) for nitrogen fixation and denitrification. We use the standard deviation of the reference tracer as a measure of the oceanic range of the net influence of the corresponding process. We measure the impact of this range on $\Omega_{\rm C}$ using a metric *M*, which we define as

$$M_i = \sigma_{R_i} \left| S_{R_i} \right|, \tag{6}$$

where S_{R_i} is the Ω_C sensitivity to a unit process change in R_i , which we estimate in Appendix A. We are interested in the relative importance I of our six processes, so we also calculate the percentage that each metric value estimate contributes to the sum of all six metric value estimates:

$$I_{i} = 100 \% \times \frac{M_{i}}{\sum_{i=1}^{6} M_{i}}.$$
(7)

We derive and estimate our metric and its uncertainty in Appendix A. We carry out our analysis for the full water column assuming it to be isolated from the atmosphere (Sect. 4.1), and also for just the top 50 m of the water column assuming it to be well-equilibrated with the atmosphere (Sect. 4.2). Finally, we consider how equilibration with an atmosphere with a changing pCO_2 alters surface Ω_C .

4.1 Process importance in atmospherically isolated mean seawater from all ocean depths

Our metric M_i is an estimate of the standard deviation of the global distribution of Ω_C resulting from the *i*th process. Our relative process importance metric I_i is an estimate of the percentage of overall variability of the Ω_C distribution that can be attributed to that process. We provide M and I values for mean seawater from the full water column alongside the R_i , S_{R_i} , and σ_{R_i} values used to estimate them in Table 1. These calculations assume that the seawater is isolated from the atmosphere.

Relative process importance estimates *I* indicate organic matter cycling (48%) is the dominant process controlling $\Omega_{\rm C}$ for mean seawater. Changing pressure (28%) is the second-most-important process, followed by calcium carbonate cycling (17%), temperature changes (4%), nitrogen fixation and denitrification (1.21%), and freshwater cycling (0.78%).

4.2 Process importance in well-equilibrated surface seawater

In Table 2 we provide M_i values for well-equilibrated seawater in the top 50 m of the ocean alongside the R_i , σ_{R_i} , and S_{R_i} used to estimate them. These surface seawater M_i values are calculated assuming the water remains equilibrated with an atmosphere with 400 µatm pCO_2 . We test the validity of this assumption by also estimating M for the observed global pCO_2 variability in the Takahashi et al. (2009) global data product. This test reveals transient air–sea disequilibria are indeed important for surface ocean Ω_C , but only as a

Table 1. Metric estimates M_i , relative process importance percentages I_i , calcite saturation sensitivities S_{R_i} to unit changes in the R_i reference properties, and reference property standard deviations σ_{R_i} for the i = 6 processes in atmospherically isolated mean seawater from all ocean depths. We provide details on how these terms are estimated and M_i and I_i uncertainties are obtained.

Process	i	R _i	S_{R_i}	σ_{R_i}	M_i	I _i
Carbonate cycling	1	Alk*	0.0043	$53.5\mu molkg^{-1}$	0.23	17 %
Org. matter cycling	2	Phosphate	-0.0069	$0.60\mu{ m molkg^{-1}}$	0.66	48%
Freshwater cycling	3	Salinity	0.032	0.27	0.011	0.78%
Sinking/shoaling	4	Pressure	-0.00028	1411 db	0.4	28%
Warming/cooling	5	Temp.	0.014	4.20 °C	0.06	4 %
Denit./nit. fix.	6	N^*	-0.010	$1.6\mu molkg^{-1}$	0.017	1.2 %

Table 2. Metric estimates M_i , relative process importance percentages I_i , calcite saturation sensitivities S_{R_i} to unit changes in the R_i reference properties, and reference property standard deviations σ_{I_i} for the i = 6 processes in well-equilibrated surface seawater. We provide details on how these terms are estimated and M_i and I_i uncertainties are obtained.

Carbonate cycling 1 Alk* 0.0034 $36.9 \mu\text{mol}\text{kg}^{-1}$ 0.13 7.8% Org. matter cycling 2 Phosphate -0.0045 $0.51 \mu\text{mol}\text{kg}^{-1}$ 0.037 2.3% Freshwater cycling 3 Salinity 0.20 0.86 0.22 13.2 Sinking/shoaling 4 Pressure -0.00083 15db 0.011 0.70 Warming/cooling 5 Temp. 0.14 $8.8 ^{\circ}\text{C}$ 1.2 76% Denit /nit fix 6 N^* -0.0043 $15 \mu\text{mol}\text{kg}^{-1}$ 0.006 0.40	Process	i R _i	S_{R_i}	σ_{R_i}	M_i	I _i
Denit /nit fix $6 N^* = -0.0043 + 5.00 k g^{-1} = 0.006 - 0.40$	Carbonate cycling Org. matter cycling Freshwater cycling Sinking/shoaling Warming/cooling	1 Alk [*] 2 Phosp 3 Salinit 4 Pressu 5 Temp	$\begin{array}{r} 0.0034 \\ hate & -0.0045 \\ ty & 0.20 \\ tre & -0.00083 \\ 0.14 \end{array}$	36.9 μmol kg 0.51 μmol kg 0.86 15 db 8.8 °C	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	7.8 % 2.3 % 13.2 % 0.70 % 76 %
Dent./mt. nx. 0 1/ 0.0045 1.5 µmorkg 0.000 0.40	Denit./nit. fix.	$6 N^*$	-0.0043	$1.5 \mu mol kg^{-1}$	0.006	0.40 %
Denit /nit fix 6 N^{*} = 0.0043 1.5 µmol kg ⁻¹ 0.006 0.40	Warming/cooling	5 Temp.	0.14	8.8 °C	1.2	76%

^a standard deviation of the Takahashi et al. (2009) revised global monthly pCO_2 climatology. ^b the *M* value for disequilibria is only calculated to test our assumption of surface seawater air–sea equilibration and is omitted from calculations of I_i for comparison with Table 1.

secondary factor when considered globally. Despite this, it is important to recognize that air–sea equilibration following a process is not instantaneous, and that the S_{R_i} value estimates in Sect. 4.1 may be better for estimating short-term changes following fast-acting processes such as spring blooms (e.g., Tynan et al., 2014) or upwelling events (e.g., Feely et al., 1988). We omit the disequilibrium M value estimate from the denominator of Eq. (7) to allow I values for surface seawater to be compared to I values from mean seawater globally.

Warming and cooling are the dominant processes controlling Ω_{C} for well-equilibrated surface seawater (76%). The large increase in M for warming and cooling relative to the value calculated for mean seawater is due to lower equilibrium C_T at higher temperatures. Freshwater cycling is the second-most-important process (13%), followed by carbonate cycling (8%), organic matter cycling (2%), pressure changes (1%), and denitrification and nitrogen fixation (0.4%). The increased importance of freshwater cycling compared to Sect. 4.1 is because freshwater dilutes $C_{\rm T}$ by more than the equilibrium $C_{\rm T}$ decreases from $A_{\rm T}$ dilution, so carbon uptake tends to follow freshwater precipitation and carbon outgassing follows evaporation. Carbonate cycling is less important because $A_{\rm T}$ decreases with carbonate precipitation lead to lower $C_{\rm T}$ at equilibrium. Organic matter cycling is much less important because atmospheric reequilibration mostly negates the large changes in $C_{\rm T}$. Pressure changes are negligible because we only consider water in the surface 50 m. Our air–sea disequilibrium M estimate suggests surface disequilibria are comparably important to freshwater cycling for surface $\Omega_{\rm C}$ but substantially less important than temperature changes (this would correspond to an I value of ~ 14%).

The dominance of warming and cooling and freshwater cycling over carbonate cycling is most evident in the Red Sea where high temperatures (> 25 °C) and high salinities (> 40) lead to surface $\Omega_{\rm C}$ exceeding 6 despite extremely low Alk* (< -200 µmol kg⁻¹). The deep Red Sea is also unusual for having deep water that was warm when it last left contact with the atmosphere (the Red Sea is > 20 °C at > 1000 m depth). This provides high initial deep $\Omega_{\rm C}$ that – combined with decreased influence of pressure changes at higher temperatures – keeps deep Red Sea $\Omega_{\rm C}$ > 3. Similarly, the lowest surface $\Omega_{\rm C}$ values are in the Arctic where there are low temperatures, low salinity, and high Alk* values from riverine inputs. The importance of warming and cooling is also suggested by the correlation between $\Omega_{\rm C}$ and the surface temperature ($R^2 = 0.96$). These properties are plotted in Fig. 8.



Figure 8. Gridded global (a) calcite saturation state $\Omega_{\rm C}$ and (b) temperature at the surface (10 m depth surface) of our gridded CA-RINA, PACIFICA, and GLODAP bottle data products. Areas with exceptionally poor coverage in the data used to produce the gridded product are blacked out.

5 Conclusions

Alk^{*} isolates the portion of the A_T signal that varies in response to calcium carbonate cycling and exchanges with terrestrial and sedimentary environments from the portion that varies in response to freshwater and organic matter cycling. The salinity normalization we use has the advantage over previous salinity normalizations that it allows our tracer to mix linearly and to change in a 2 : 1 ratio with C_T in response to carbonate cycling. We highlight the following insights from Alk^{*}:

- 1. Alk* distribution: the Alk* distribution clearly shows the influence of biological cycling, including such features as the very low Alk* in the Red Sea due to the high calcium carbonate precipitation there. We also find evidence of strong riverine A_T sources in the Bay of Bengal and in the Arctic. We show river inputs likely dominate over the small influences of ikaite cycling on the Arctic alkalinity distribution.
- Influence of calcium carbonate cycling on marine calcite saturation state: Alk* allows us to quantify the net influence of calcium carbonate cycling on marine Ω_C. For well-equilibrated surface waters, carbonate cycling is less influential for Ω_C than gas exchange driven by warming and cooling and freshwater cycling. At depth, the carbonate cycling signal is smaller than the signal

from organic matter cycling and from pressure changes. Temperature is the dominant control on Ω_C of surface waters in equilibrium with the atmosphere. This accounts for the low calcite saturation states in the cold surface of the Arctic and Southern oceans despite high regional Alk^{*}, and high Ω_C in the warm subtropics despite low regional Alk^{*}.

We intend to use Alk* for two future projects. First, Alk* is superior to $A_{\rm T}$ for monitoring and modeling changes in marine chemistry resulting from changes in carbonate cycling with ocean acidification. $A_{\rm T}$ varies substantially in response to freshwater cycling, so Alk* trends may be able to be detected sooner and more confidently attributed to changes in calcium carbonate cycling than trends in $A_{\rm T}$ (Ilyina et al., 2009). Secondly, we will estimate global steady-state Alk* distributions using Alk* sources and sinks from varied biogeochemical ocean circulation models alongside independent water mixing and transport estimates (e.g., Khatiwala et al., 2005; Khatiwala, 2007). We will interpret findings in the context of two hypotheses proposed to explain evidence for calcium carbonate dissolution above the aragonite saturation horizon: (1) that organic matter remineralization creates undersaturated microenvironments that promote carbonate dissolution in portions of the water column which are chemically supersaturated in bulk, and (2) that high-magnesium calcite and other impure minerals allow chemical dissolution above the saturation horizon.

Appendix A: Definition of the process importance metric *M*

In simplest terms, our metric is the product of the Ω_C sensitivity to a process and the variability of the net influence of the process globally. The difficulty in this calculation lies in quantifying the "net influence of a process." We first show how we change coordinates so we can use reference tracers as a proxy measurement for these net influences.

Our metric for $\Omega_{\rm C}$ variability resulting from the *i*th process is expressed as M_i :

$$M_i = \sigma_{P_i} \left| \frac{\partial \Omega_{\rm C}}{\partial P_i} \right|,\tag{A1}$$

where P_i is an abstract variable representing the net process influence (that we will later factor out), and $\frac{\partial \Omega_C}{\partial P_i}$ is the Ω_C sensitivity to the process. We expand $\frac{\partial \Omega_C}{\partial P_i}$ using the chain rule to include a term for Ω_C sensitivity to changes in the reference tracer R_i (see Sect. 4) and a term $\frac{\partial R_i}{\partial P_i}$ representing changes in R_i resulting from the *i*th process:

$$\frac{\partial \Omega_{\rm C}}{\partial P_i} = \frac{\partial \Omega_{\rm C}}{\partial R_i} \frac{\partial R_i}{\partial P_i}.$$
 (A2)

In practice, we calculate $\Omega_{\rm C}$ as a function of j = 7 properties: (1) pressure, (2) temperature, (3) salinity, (4) phosphate, (5) silicate, (6) $A_{\rm T}$, and (7) $C_{\rm T}$ for mean seawater and $p{\rm CO}_2$ for surface seawater, so we use the chain rule again to expand the $\frac{\partial \Omega_{\rm C}}{\partial R_{\rm c}}$ terms as follows:

$$\frac{\partial \Omega_{\rm C}}{\partial R_i} = \sum_{j=1}^7 \frac{\partial \Omega_{\rm C}}{\partial X_j} \frac{\partial X_{j,i}}{\partial R_i}.$$
 (A3)

Here, the $\frac{\partial X_{j,i}}{\partial R_i}$ are assumed terms (assumptions detailed shortly) that relate the effect of the *i*th process on the *j*th property to the effect of the process on R_i , and the $\frac{\partial \Omega}{\partial X_j}$ terms reflect $\Omega_{\rm C}$ sensitivity to changes in the *j* properties used to calculate it.

We make assumptions regarding the $\frac{\partial X_{j,i}}{\partial X_R}$ terms: we relate changes in temperature from sinking or shoaling to changes in pressure using the potential temperature (θ) routines of Fofonoff and Millard (1983); we assume freshwater cycling linearly concentrates $A_{\rm T}$, $C_{\rm T}$, phosphate, and silicate by the same ratio that it changes salinity; we relate $C_{\rm T}$, phosphate, and AT changes from organic matter formation to changes in phosphate using the remineralization ratios found by Anderson and Sarmiento (1994) and the empirical relationship of Kanamori and Ikegami (1982); we also use Kanamori and Ikegami's (1982) constant to relate changes in $A_{\rm T}$ from nitrogen fixation and denitrification to changes in N* from these processes; and we assume that an increase in $A_{\rm T}$ from calcium carbonate dissolution equals the Alk* increase, and that the corresponding increase in $C_{\rm T}$ equals half of this Alk* increase. We neglect any changes in $C_{\rm T}$ from denitrification and nitrogen fixation because these changes are better thought of as organic matter cycling occurring alongside nitrogen cycling.

We estimate $\frac{\partial \Omega}{\partial X_j}$ property sensitivity terms as the differences between $\Omega_{\rm C}$ calculated before and after augmenting *j*th property by one unit. $\Omega_{\rm C}$ is calculated with the MATLAB CO2SYS routines written by van Heuven et al. (2009) using the carbonate system equilibrium constants of Mehrbach et al. (1973), as refit by Dickson and Millero (1987). Seawater $p{\rm CO}_2$ is used in place of $C_{\rm T}$ for the surface seawater calculations (when j = 7) to calculate the change in $\Omega_{\rm C}$ that remains after the surface seawater is allowed to equilibrate with the atmosphere.

We assume that the distributions of our R_i reference properties are linearly related to the P_i net activities of their associated processes. This assumption implies

$$\sigma_{P_i} = \sigma_{R_i} \left| \frac{\partial P_i}{\partial R_i} \right|. \tag{A4}$$

We can then substitute Eq. (A3) into Eq. (A2), and substitute this combined equation for $\frac{\partial \Omega_C}{\partial P_i}$ and Eq. (A4) into Eq. (A1). We then cancel the $\frac{\partial P_i}{\partial R_i}$ and $\frac{\partial R_i}{\partial P_i}$ terms to obtain

$$M_{i} = \sigma_{R_{i}} \left| \sum_{j=1}^{7} \frac{\partial \Omega_{C}}{\partial X_{j}} \frac{\partial X_{j,i}}{\partial R_{i}} \right|.$$
(A5)

We then define $\Omega_{\rm C}$ sensitivity S_{R_i} as

$$S_{R_i} = \left| \sum_{j=1}^{7} \frac{\partial \Omega_{\rm C}}{\partial X_j} \frac{\partial X_{j,i}}{\partial R_i} \right|,\tag{A6}$$

where S_{R_i} is the $\Omega_{\rm C}$ sensitivity to a change in the *i*th process scaled to a unit change in the reference variable for that process. We can then substitute Eq. (A6) into Eq. (A5) to obtain Eq. (6). We use Eq. (A6) to define S_{R_i} and Eq. (6) to calculate M. We provide the $\frac{\partial \Omega_{\rm C}}{\partial X_j}$ and $\frac{\partial X_{j,i}}{\partial R_i}$ values we use to estimate S_{R_i} for atmospherically isolated seawater from all depths in Table A1 and for well-equilibrated surface seawater in Table A3. We perform a sample I and M calculation in Supplement document SE.

We use a Monte Carlo analysis to estimate variability and uncertainty in our metric M and our percent relative process importance I calculations. We calculate the standard deviations, σ_M and σ_I , of pools of 1000 M and I estimates calculated after adjusting the seawater properties X_i with a normally distributed perturbation with a standard deviation equal to the property standard deviation from the gridded data set. We find $\frac{\sigma_I}{I}$ is typically much smaller than $\frac{\sigma_M}{M}$. This is because $\Omega_{\rm C}$ sensitivity is typically proportional to the $\Omega_{\rm C}$ itself, so individual Monte Carlo M estimates vary with the initial $\Omega_{\rm C}$ and one another. Our σ_M estimates are therefore better thought of as measures of the ranges of sensitivities found in the modern ocean, while σ_I represent variability in the relative importance of processes. We provide σ_M and σ_I for atmospherically isolated seawater globally in Table A2, and for well-equilibrated surface seawater in Table A4.

Table A1. $\frac{\partial \Omega_{\rm C}}{\partial X_j}$ (bold text) and $\frac{\partial X_{j,i}}{\partial R_i}$ (italic text) terms used in Eq. (A5) for atmospherically isolated mean seawater from all ocean depths. These terms are specific to the j = 7 (columns) properties we use to calculate $\Omega_{\rm C}$ and i = 6 (rows) processes we consider. Units for $\frac{\partial \Omega_{\rm C}}{\partial X_j}$ are the inverse of the listed X_j units. Units for $\frac{\partial X_{j,i}}{\partial R_i}$ are the X_j units divided by the R_i units given in Table 1.

Properties		Pressure	Temp	Salinity	Phos. -1	Silicate	$A_{\rm T}$	$C_{\rm T}$
Units		db	-0	2	µmol kg	µmol kg	µmol kg	µmorkg -
J		1	2	3	4	5	6	1
Mean seawater value	s	2235	3.7	34.71	2.15	49.0	2362	2254
$\frac{\partial \Omega_{\rm C}}{\partial X_j}$		-0.00028	0.014	-0.011	-0.0085	-0.00012	0.0082	-0.0079
Process	i	$\frac{\partial X_{1,i}}{\partial R_i}$	$\frac{\partial X_{2,i}}{\partial R_i}$	$\frac{\partial X_{3,i}}{\partial R_i}$	$\frac{\partial X_{4,i}}{\partial R_i}$	$\frac{\partial X_{5,i}}{\partial R_i}$	$\frac{\partial X_{6,i}}{\partial R_i}$	$\frac{\partial X_{7,i}}{\partial R_i}$
Carbonate cycling	1	-	-	_	-	-	1	0.5
Org. matter cycling	2	_	-	_	1	-	-20.16	117
Freshwater cycling	3	_	-	1	0.062	1.4	68	65
Sinking/shoaling	4	1	0.00010	-	_	_	_	_
Warming/cooling	5	_	1	-	_	_	_	_
Denit./nit. fix.	6	_	_	_	_	_	-1.26	_

Table A2. Monte Carlo-derived estimates for M_i variability (σ_{M_i}) and I_i variability (σ_{I_i}) for atmospherically isolated mean seawater from all ocean depths.

Process	i	σ_{M_i}	σ_{I_i}
Carbonate cycling	1	0.09	1 %
Org. matter cycling	2	0.2	3%
Freshwater cycling	3	0.006	0.08~%
Sinking/shoaling	4	0.2	5 %
Warming/cooling	5	0.02	2 %
Denit./nit. fix.	6	0.006	0.1 %

Table A3. $\frac{\partial \Omega_C}{\partial X_j}$ (bold text) and $\frac{\partial X_{j,i}}{\partial R_i}$ (italic text) terms used in Eq. (A5) for well-equilibrated surface seawater. These terms are specific to the j = 7 (columns) properties we use to calculate Ω_C and i = 6 (rows) processes we consider. Units for $\frac{\partial \Omega_C}{\partial X_j}$ are the inverse of the listed X_j units. Units for $\frac{\partial X_{j,i}}{\partial R_i}$ are the X_j units divided by the R_i units given in Table 2.

Properties		Pressure	Temp	Salinity	Phos.	Silicate	A_{T}	pCO_2
Units		db	°C		μ mol kg $^{-1}$	μ mol kg $^{-1}$	μ mol kg ⁻¹	µatm
j		1	2	3	4	5	6	7
Mean seawater value	S	25	18.3	34.82	0.51	2.5	2305	350
$\frac{\partial \Omega_{\rm C}}{\partial X_j}$		-0.00084	0.14	-0.022	-0.0038	-0.00013	0.0034	-0.0086
Process	i	$\frac{\partial X_{1,i}}{\partial R_i}$	$\frac{\partial X_{2,i}}{\partial R_i}$	$\frac{\partial X_{3,i}}{\partial R_i}$	$\frac{\partial X_{4,i}}{\partial R_i}$	$\frac{\partial X_{5,i}}{\partial R_i}$	$\frac{\partial X_{6,i}}{\partial R_i}$	$\frac{\partial X_{7,i}}{\partial R_i}$
Carbonate cycling	1	_	_	_	-	-	1	_
Org. matter cycling	2	_	_	_	1	_	-20.16	-
Freshwater cycling	3	_	_	1	0.015	0.072	65.9	-
Sinking/shoaling	4	1	0.00010	_	_	_	_	-
Warming/cooling	5	_	1	_	_	_	_	-
Denit./nit. fix.	6	_	_	_	_	_	-1.26	_

Table A4. Monte Carlo-derived estimates for M_{i}	variability (σ_{M_i}) and I_i	I_i variability (σ_{I_i}) for well-equilibrated	surface seawater.

Process	i	σ_{M_i}	σ_{I_i}
Carbonate cycling	1	0.03	0.8 %
Org. matter cycling	2	0.01	0.2 %
Freshwater cycling	3	0.04	0.5 %
Sinking/shoaling	4	0.001	0.03 %
Warming/cooling	5	0.2	1 %
Denit./nit. fix	6	0.002	0.04 %
<i>p</i> CO ₂ disequilibria	а	0.05	a

^a disequilibria are included only as a test of our assumption of surface seawater air–sea equilibration, so these M_i values are omitted from calculations of *I*.

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