Role of oceanic abiotic carbonate precipitation in future atmospheric CO₂ regulation

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The oceans play a major role in the earth's climate by regulating atmospheric CO₂. While oceanic primary productivity and organic carbon burial sequesters CO₂ from the atmosphere, precipitation of CaCO₃ in the sea returns CO₂ to the atmosphere. Abiotic CaCO₃ precipitation in the form of aragonite is potentially an important feedback mechanism for the global carbon cycle, but this process has not been fully quantified. In a sediment-trap study conducted in the southeastern Mediterranean Sea, one of the fastest warming and most oligotrophic regions in the ocean, we quantify for the first time the flux of inorganic aragonite in the water column. We show that this process is kinetically induced by the warming of surface water and prolonged stratification resulting in a high aragonite saturation state (ΩAr ≥ 4). Based on these relations, we estimate that abiotic aragonite calcification may account for 15 ± 3% of the previously reported CO₂ efflux from the sea surface to the atmosphere in the southeastern Mediterranean. Modelled predictions of sea surface temperature and Ω Ar suggest that this process may weaken in the future ocean, resulting in increased alkalinity and buffering capacity of atmospheric CO₂.

The production of CaCO₃ minerals (namely, aragonite and calcite) plays an important role in regulating the ocean's carbon budget¹-². Whereas most of the CaCO₃ production in the oceans is biogenic, a fraction forms in abiotic reactions. To date, abiotic CaCO₃ production in the form of aragonite has been observed in certain localities, such as the Bahamas or the Persian Gulf, where specific conditions are present. However, as ocean acidification intensifies, the production of abiotic CaCO₃ will diminish and may have already diminished significantly, due to the reduction in the saturation state of the carbonate ion. Here we show, for the first time, the occurrence of abiotic aragonite production in the eastern Mediterranean, under different conditions than previously observed and a possible analogue to future ocean conditions under the effects of global warming. According to the proposed mechanism, the abiotic production of CaCO₃ in surface waters may be enhanced by warming and stratification of the oceans, rather than additional seeding or water mass mixing. In contrast, ocean acidification may inhibit the abiotic production of surface waters, countering the positive effect of warming. Based on our observations, we argue that abiotic aragonite production is more ubiquitous than previously considered and delineate the production mechanism. Based on state-of-the-art climate models, we illustrate that the potential that this mechanism is diminishing and discuss its impact.

Background

The exchange of CO₂ between the ocean and the atmosphere has been considered to act as a major feedback mechanism that helps regulate planetary climate¹-². Currently, it is estimated that ca. 25% of anthropogenic CO₂ emitted to the atmosphere annually is absorbed by the ocean³-⁴. The ensuing chemical reaction of the absorbed CO₂ with seawater produces carbonic acid, resulting in a process known as ocean acidification⁵. The production of carbonic acid not only reduces seawater's pH, but also shifts the speciation of dissolved inorganic carbon (DIC = CO₂ + HCO₃⁻ + CO₃²⁻) in seawater away from carbonate (CO₃²⁻) to bi-carbonate (HCO₃⁻), according to the following stoichiometric equation – CO₂ + H₂O + CO₃²⁻→ 2HCO₃⁻.

The speciation of DIC in seawater is also influenced by temperature through its effect on the thermodynamic dissociation constants, where warming shifts the carbonate system towards CO₃²⁻. Most of the transformation of DIC into solid phase, be it organic or inorganic, is biologically-driven⁶. Where, biologically mediated assimilation of DIC into inorganic carbon solid phase (CaCO₃) removes Ca²⁺ from seawater and results also in a reduction in alkalinity and an increase in pH and buffering capacity⁷. Abiotic production of CaCO₃, on the other hand, is a chemical process that can occur without biological intervention. Abiotic precipitation of CaCO₃ may play a role in the carbon cycle, especially under conditions of high CO₂ sequestration by biological processes, such as in the open ocean. Abiotic precipitation of CaCO₃ occurs when CO₂ dissolves in water and reacts with calcium ions (Ca²⁺), forming CaCO₃, according to the following reaction:

\[ \text{Ca}^2+ + \text{CO}_2 + \text{H}_2\text{O} \rightarrow \text{CaCO}_3 + \text{H}^+ \]

This reaction is thermodynamically favourable at low CO₂ levels and high Ca²⁺ concentrations. However, as CO₂ levels increase and Ca²⁺ concentrations decrease, the reaction shifts towards the production of CO₂, leading to acidification of the ocean. The equilibrium constant for this reaction is strongly influenced by temperature and pressure, with increasing temperature shifting the equilibrium towards the production of CO₂.

The role of abiotic precipitation of CaCO₃ in the carbon cycle is not well understood. While it is known that abiotic precipitation of CaCO₃ occurs in certain localities, such as the Bahamas or the Persian Gulf, where specific conditions are present, its contribution to the global carbon cycle has not been quantified. In this study, we quantify for the first time the flux of inorganic aragonite in the water column of the southeastern Mediterranean Sea, one of the fastest warming and most oligotrophic regions in the ocean. We show that this process is kinetically induced by the warming of surface water and prolonged stratification, resulting in a high aragonite saturation state (ΩAr ≥ 4). Based on these relations, we estimate that abiotic aragonite calcification may account for 15 ± 3% of the previously reported CO₂ efflux from the sea surface to the atmosphere in the southeastern Mediterranean. Modelled predictions of sea surface temperature and Ω Ar suggest that this process may weaken in the future ocean, resulting in increased alkalinity and buffering capacity of atmospheric CO₂.

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of seawater total alkalinity (TA). This process is also non conservative with salinity changes. While the organic sink (“biological pump”) removes carbon from the atmosphere, the inorganic sink (precipitation of carbonate minerals), consumes TA and shifts the carbonate system towards CO₂ and its degassing from the ocean surface to the atmosphere, acting as a positive feedback to global warming. Precipitation of both crystal forms of CaCO₃ (aragonite and calcite) in the ocean is mostly biogenic and is a highly complex process, affected by environmental conditions, such as temperature, dissolved nutrient levels and, most noteworthy, the ratio of the Ca²⁺ and CO₃⁻² ion activity product to the solubility constant of CaCO₃, also known as the CaCO₃ saturation state (Ω). Oceanic surface waters are mostly supersaturated with respect to CaCO₃ (Ω > 1) and therefore, one would rightfully expect the abiotic precipitation of CaCO₃ to be a common occurrence and an important process in the oceanic carbon cycle and climate regulation. However, to date, large spatial scale abiotic CaCO₃ precipitation (usually aragonite) in the marine environment have been observed only during events referred to as “whitening events”, when CaCO₃ is abiotically precipitated in surface waters, turning them milky white in relatively shallow waters. Such events occur almost exclusively in the Persian Gulf and the Bahamas. While the specific mechanism driving whitening events is still a highly contested topic, extremely high levels of Ω seem to be an important factor triggering them.

Abiotic nucleation of CaCO₃ in seawater is strongly inhibited kinetically. Suspended particles, which provide a mineral surface area for nucleation (“seeding”), were also suggested to play an important control for carbonate precipitation from seawater. However, “seeded” experiments have concluded that temperature controls the mineralogy of the precipitating phase and that the precipitation rate is proportional to the saturation state. Hence, vast low latitude oceanic regions, where the saturation state is high and most CO₂ degassing occurs, are suspected to accommodate abiotic precipitation of aragonite. Yet, this process remains largely undetected, as it is often concealed by the larger biological signal.

Abiotic aragonite precipitation. As the ocean warms and stratification intensifies, primary productivity decreases, most notably in oligotrophic regions. This would mean that the carbon cycle in these regions may sway from biological mediation towards greater abiotic (chemical and physical) control. This process is already ongoing in the southeastern Mediterranean, making it a good model system for the warmer stratified future of the oceans. Like most oceanic basins, the region is experiencing acidification, yet, the rate of acidification is relatively low due to increasing alkalinity (supplemental 1, Fig. S1.1). In contrast, the summertime sea surface temperature has risen at an exceptionally high rate of up to 1 °C/decade over the last three decades, substantially higher than warming rates in other low latitude tropical seas. Additionally, the oligotrophic state of the region results in one of the lowest primary productivities of any marine water body in the world.

In a sediment-trap study that we conducted in the southeastern Mediterranean Sea (Fig. 1), we quantify for the first time the flux of inorganic precipitation of aragonite in the water column during the summer (supplemental 2). Unlike calcite, there are only a few sources of biogenic aragonite in the study area, e.g. pteropod blooms and
resuspension of eroded aragonite shells of benthic organisms – none of which were present in the traps based on microscopic examination of the material in the traps and adjacent surface sediments. The Sr/Ca ratios in the traps ranged between 6.8 and 11.4 in the summer months when aragonite was present (Fig. 2; supplemental 1, Table S1.1). The Sr/Ca ratio in the sediment was found to be ~ 5.0 and biogenic sources such as pteropods were found to have ratios ranging between 2.3 and 3.6 (supplement 1, Table S1.2). As a result, neither of these sources can explain the higher Sr/Ca ratio in the traps. However, these values agree well with the Sr/Ca ratio (~ 9.0) of abiotic aragonite precipitated in lab experiments from seawater in the range of summertime temperatures in the study area. This is still lower than the maximum value of Sr/Ca = 11.4, where the excess Sr is likely sourced from Saharan dust, which has a Sr/Ca ratio of ~ 304,35. However, the dust is aragonite-free and its supply rate is low during summer. Thus, neither dust nor resuspension, given the low wave activity during summer, explain the relatively high Sr/Ca ratio in the traps. Thus, it is reasonable to conclude that it is caused by abiotic precipitation of aragonite (supplemental 1, Figs. S1.2–S1.3).

Recently, it was suggested that abiotic aragonite precipitation may be caused by seeding, where aragonite crystallizes on suspended nucleation particles. In this model, a high load of nucleation seeds/surface area promotes increased precipitation. The saturation state, while elevated (≥ 4), does not appear to be sufficient for spontaneous nucleation of aragonite from seawater. Therefore, our results support utilization of preexisting nuclei in the water mass by a sustained high aragonite saturation state during summer. Acoustic backscatter data from the ADCP adjacent to the sediment trap (Fig. 2), a proxy for turbidity in the water, indicated highest turbidity between January and May, and much lower turbidity during the summer months. This trend was mirrored in the total mass and CaCO₃ fluxes in the sediment traps (supplement 1; Table S1.1) decreased from 82 and 15.8 gr m⁻² d⁻¹ in January 2016 to 2.9 and 0.7 gr m⁻² d⁻¹ in August 2016, respectively. As such, we would argue that even in stratified, low-turbidity environments, precipitation nuclei are not a limiting factor for inorganic precipitation of aragonite.

The other major factor for the precipitation of aragonite is the seawater CaCO₃ saturation state. Abiotic nucleation of aragonite has been demonstrated to be possible with ΩAr = 4, given induction time (the period of time that is necessary to initiate a reaction) of around two months. To maintain stable ΩAr conditions, a prolonged period of water column stability is required which is consistent with the stratification of the surface water. Our observations show that aragonite formation started in late-spring and continued throughout the summer months, between May and October 2016 (Fig. 2), corresponding to the time of year when warming and weaker mechanical mixing causes intense stratification of the surface layer (Fig. 3). At its apex, the gradient across the seasonal thermocline was on the order of 10 °C (supplement 1; Fig. S1.4). As this layer concentrates heat at the surface, pCO₂ levels reach super saturation in excess of 450 µatm (Fig. 2). During winter, the upper water column is mixed (Fig. 3), temperature and ΩAr are lower (Fig. 2), while biogenic production and sediment load are higher, suppressing abiotic formation of aragonite (Fig. 4a). The absence of abiotic aragonite in the sediment traps during the autumn of 2015 when ΩAr was at times > 4, is likely the result of two notable (4–6 °C) cooling events that correspond to a CO₂ efflux of 0.12 ± 0.01 mol CO₂ m⁻² year⁻¹ (supplemental 4). The total annual net CO₂ flux to the atmosphere in the southeastern Mediterranean Sea was previously estimated at 0.85 ± 0.27 mol m⁻². This suggests that the abiotic aragonite precipitation may account for ~ 15 ± 3% of the annual net CO₂ flux from the sea surface to the atmosphere in the southeastern Mediterranean. Relatively stable and consistent sea surface temperature and salinity conditions in the region allow for the assumption that this represents at least the region of the Levant Basin and possibly extends beyond it in the southeastern Mediterranean. This net annual calcification flux per unit area is only ~ 0.5% of CaCO₃ flux estimated for coral reefs, but the potential area of production is orders of magnitude larger, making it a significant source of atmospheric CO₂.

This is the first time that the relation between warming, stratification, abiotic aragonite precipitation and CO₂ release is demonstrated in any marine system. The observed link between surface warming and CO₂ efflux due to abiotic aragonite precipitation provide a new mechanism by which warming reduces the southeastern Mediterranean Sea buffering capacity for sequestering atmospheric CO₂ and possibly even becoming a CO₂ source. This is a novel feedback mechanism of the carbonate system under global warming.

Implications. More broadly, it was recently shown that much of the lower latitude surface ocean is characterized by ΩAr ≥ 4, and that ΩAr is positively correlated with temperature. As the lower latitude ocean warms, much of that heat is stored in the surface waters, resulting in enhanced stratification and oligotrophy. These three properties (warming, stratification and oligotrophy) will likely modify the balance of the carbon system, thereby affecting the ocean's capacity to store CO₂. Warming, in particular, is of significant importance as it affects the apparent rate of CaCO₃ precipitation through the kinetic rate coefficient Kₜ and order of reaction nₜ (which are temperature dependent; supplement 3; Fig. S3.1). Thus, if the temperature increases Kₜ and nₜ faster than ocean acidification decreases ΩAr, the net apparent CaCO₃ precipitation rate may increase. However, atmospheric buildup of CO₂ is faster than warming in terms of their combined and individual effects on Kₜ and
Ω_{Ar} (supplement 3). Under the influence of warming alone, global potential aragonite production rate would...
increase, but due to the effect of CO2 induced acidification, they have been and are expected to continue to drop (Fig. 5). As a result, the abiotic aragonite sink may be weakening, resulting in reduced TA uptake and non-conservative build-up of TA in the oceans as suggested by the Hawaiian Ocean Time Series salinity normalized TA time series (supplement 1, Fig. S1.8). In turn, this TA build-up could potentially increase the ocean's buffering capacity for increasing atmospheric CO2, increasing oceanic uptake. However, it should be noted that decadal changes in terrestrial runoff, deep mixing and evaporation regimes in and around the Pacific could also explain the non-conservative long-term trend in salinity45. Regardless, the exact contribution of abiotic aragonite precipitation in the upper water column cannot be fully quantified at this time—but it is a feedback mechanism that should be further investigated and incorporated into planetary models to move towards a more complete description of the ocean/atmosphere system.

In the geological record, shifts between periods of aridity and strong oceanic stratification with high aragonite/calcite ratios and periods of high humidity, cooler mixed water column conditions with low aragonite/calcite ratios, are evident in low latitude sediment cores42. Taking into account that surface seawater pCO2 is mostly temperature-dependent, and that ocean temperatures will continue to increase in the near future46, a reduction in the ocean's capacity to absorb CO2 is to be expected under prolonged stratification. In both Earth's past and near future, warming due to the earth orbital changes could enhance the abiotic carbonate formation, potentially resulting in a greater release of CO2 from the surface waters. The switch from cooler waters (glacial/icehouse periods), where carbon is removed by the biological pump to warm (interglacial/greenhouse period), stratified and oligotrophic surface water, where abiotic aragonite is precipitated may have altered the oceanic buffering capacity. This means that the surface water will become a much less effective sink and under extreme scenarios, larger areas of the surface ocean may become a CO2 source. However, our proposed feedback mechanism suggests a future reduction in abiotic aragonite precipitation under the increasing effect of ocean acidification. The

**Figure 3.** Time series of temperature in the upper 100 m of the water column at THEMO-1 station (early 2018 to early 2019) showing the summertime stratification of the water column.

**Figure 4.** Schematic model for aragonite formation in the Eastern Mediterranean. During cooling events and vertical mixing of the water column (a), the cooler water with increased productivity acts as a CO2 sink. During prolonged periods of warming (b), the water column becomes thermally stratified, CO2 degasses due to super saturation, CO3^2− and ΩAr increase, and abiotic aragonite precipitation occurs and intensify CO2 supersaturation and degassing.
unprecedented rate of anthropogenic climate change is not paired with a comparable increase in continental weathering. In the geological past, weathering may have provided TA to sustain this mechanism, at least in epicontinental seas which could decouple with respect to $\text{Ca}^{2+}$ and $\text{Mg}^{2+}$ from the ocean$^{47}$.

We therefore suggest that the feedback between warming, acidification, and induced $\text{CO}_2$ release due to abiotic aragonite precipitation can be regarded as a potential feedback to global warming, adding to a growing list of feedbacks such as reduced primary productivity in the ocean, desertification, and melting of permafrost$^{46}$, and therefore should be accounted for in future estimation of ocean evolution in response to climate change.

**Methods.** Samples were collected from sediment traps deployed 2 km offshore of Hadera, Israel at 25 m bottom depth. Analysis of sediment samples was carried out using X-ray fluorescence (XRF) and X-ray diffraction (XRD). Seawater carbonate chemistry parameters were estimated from in situ measurements of temperature and salinity from the ISRAM R data base, which were converted to $\text{pCO}_2$ and total alkalinity respectively. See supplement 2 for more information on methods. Global state of the carbon system was estimated from CMIP6 products, see supplement 3 for more information.
Data availability

Scripts, results and additional material for model simulations are available via a the Figshare repository at https://doi.org/10.6084/m9.figshare.15121131; all other material in the supplement are available at https://doi.org/10.6084/m9.figshare.1972181. Oceanographic data used in this manuscript is available through the ISRAMAR (https://isramar.ocean.org.il/isramar2009/) databases, see https://themo.haifa.ac.il/ for more information on the THEMO project. Additional notes on data are listed in supplements 2 and 3.

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Author contributions
O.M.B.—Conceptualization, contributed data and analysis tools, carried out analysis and took part in writing of the manuscript. G.S.V.—Conceptualization, contributed data and analysis tools, carried out analysis and took part in writing of the manuscript. N.V.V.—Carried out analysis and contributed to the writing of this manuscript. J.S.—Contributed analysis tools and contributed to the writing of this manuscript. T.K.—Conceptualization, conceived the design of the data collection, obtained funds, contributed data and analysis tools, carried out analysis and took part in writing of the manuscript.

Competing interests
The authors declare no competing interests.

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