Total alkalinity production in a mangrove ecosystem reveals an overlooked Blue Carbon component

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Scientific Significance Statement

Blue Carbon ecosystems (seagrass meadows, mangroves, and saltmarshes) sequester atmospheric CO₂ as organic carbon in their sediments for periods of centuries to millennia. Calcium carbonate (CaCO₃) dissolution is another major permanent sink of atmospheric CO₂ in the coastal area, but has been disregarded in Blue Carbon ecosystems. In the Red Sea, as in many tropical areas, Blue Carbon ecosystems thrive on carbonate sediment generated by the erosion of coral reefs. Our study reveals that the dissolution of CaCO₃ is a major sink of atmospheric CO₂ in carbonate sediment mangroves, representing 23-fold the sink from organic carbon sequestration at our Red Sea mangrove site. The CaCO₃ calcification and dissolution budget should be considered in future Blue Carbon assessments.

Abstract

Mangroves have the capacity to sequester organic carbon (Corg) in their sediments permanently. However, the carbon budget of mangroves is also affected by the total alkalinity (TA) budget. Principally, TA emitted from carbonate sediment dissolution is a perennial sink of atmospheric CO₂. The assessment of the TA budget of mangrove carbonate sediments in the Red Sea revealed a large TA emission of 403 ± 17 mmol m⁻² d⁻¹, independent of light, seasons, or the presence of pneumatophores, compared to -36 ± 10 mmol m⁻² d⁻¹ in lagoon sediment. We estimate the TA emission from carbonate dissolution in Red Sea mangroves supported a CO₂ uptake of 345 ± 15 gC m⁻² yr⁻¹, 23-fold the Corg burial rate of 15 gC m⁻² yr⁻¹. The focus on Corg burial in sediments may substantially underestimate the role of mangroves in CO₂ removal. Quantifying the role of mangroves in climate change mitigation requires carbonate dissolution to be included in assessments.

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Mangrove ecosystems, along with seagrasses beds and saltmarshes, sequester large amounts of organic carbon \((C_{\text{org}})\) in their sediments (Duarte et al. 2005, 2013; Mcleod et al. 2011). Carbon burial in mangrove sediments has been estimated at 21 to 32.4 Tg \(C_{\text{org}}\) yr\(^{-1}\) globally (Breithaupt et al. 2012), contributing about 10% of the \(C_{\text{org}}\) burial in the ocean (Duarte et al. 2005; Breithaupt et al. 2012). Before perennial sequestration, part of the labile fraction of the organic matter (OM) is remineralized through a cascade of redox reactions involving aerobic respiration in surface sediments and hydrolysis and fermentation in the deeper layers (Middelburg 2018).

Degradation of OM in marine sediment can also be a source of total alkalinity (TA) to the marine environment. TA, simplified as the charge balance of conservative ions in seawater (e.g., \(Na^+, Ca^{2+}, K^+, Cl^-\)) and of the redox state of some minor elements (e.g., species of the nitrogen and phosphorus cycle) (Middelburg et al. 2019), is a key parameter affecting the seawater inorganic carbon system and the global carbon cycle (Krumins et al. 2013). Seawater is slightly depleted of conservative anions compared to conservative cations, and the difference (about 2 mmol kg\(^{-1}\)) allows for the uptake of atmospheric \(CO_2\), stored in seawater as \(HCO_3^-\) and \(CO_3^{2-}\). The consequence of this minor imbalance results in the world’s oceans storing about 98% of the \(CO_2\) present in the atmosphere – hydrosphere system.

TA in mangrove sediments is produced through sulfate reduction, calcium carbonate \((CaCO_3)\) dissolution, denitrification, and ammonification (Krumins et al. 2013). In carbonate, iron-depleted sediments characteristics of coral reefs regions (Ku et al., 1999; Hu and Burdige, 2007; Krumins et al., 2013), the TA produced from sulfate reduction is re-uptaken in equal amounts during sulfite oxidation after diffusion into the water column. Hence, much of sediment TA emissions derive from \(CaCO_3\) dissolution along with denitrification and ammonification (Krumins et al. 2013). The emission of TA to the water column by mangrove sediments is a sink of atmospheric \(CO_2\) (Maher et al. 2018). In particular, TA emission from \(CaCO_3\) dissolution is a perennial sink of \(CO_2\) at the geological time scale (Ridgwell and Zeebe 2005), which should be added to the classical assessment of Blue Carbon as \(C_{\text{org}}\) sequestration. Understanding the role of calcification and \(CaCO_3\) dissolution in affecting \(CO_2\) budgets of Blue Carbon ecosystems is a major challenge for Blue Carbon science (Macreadie et al. 2017; Saderne et al. 2019b). Through TA emission, the dissolution of \(CaCO_3\) results in an atmospheric uptake of \(\sim 0.6\) mol of \(CO_2\) per mole of \(CaCO_3\) dissolved under current atmospheric \(CO_2\) partial pressure \((pCO_2)\). This is independent of the pathway involved in the dissolution or metabolic activities occurring in parallel in the ecosystem (e.g., respiration, photosynthesis) (Frankignoulle et al. 1995). The \(CO_2\) produced by the degradation of sediment OM, together with organic acid exuded by mangrove roots (Jiang et al. 2017), results in an acidification of the pore waters and the dissolution of \(CaCO_3\) (Hu and Burdige 2007). Dissolution of \(CaCO_3\) could also occur at the sediment-water column interface in Blue Carbon ecosystems, due to the acidification caused by the metabolic activity of the benthic fauna and flora, principally respiration at night (Rosenreter et al. 2018; Saderne et al. 2019a). Hence, based on a review of carbonate burial and budgets in Blue Carbon sediments, we recently proposed that carbonate dissolution may be an important, but overlooked mechanism of \(CO_2\) removal in Blue Carbon habitats where carbonate is supplied from adjacent calcifying ecosystems, such as coral reefs and lithogenic carbonate deposits (Saderne et al. 2019b). Here, we report a high rate of \(CO_2\) removal from TA net emission by a mangrove stand with carbonate-rich sediments (80% \(CaCO_3\) of dry weight; Saderne et al. 2018) in the central-eastern Red Sea. We do so based on assessments of annual net TA fluxes in a flooded mangrove stand compared to those in bare reef lagoon sediments outside of the mangrove.

**Materials and methods**

**Site**

The incubations were conducted in a mono-specific *Avicennia marina* mangrove swamp of the central Red Sea (22°20′23″N, 39°05′22″E, Fig. 1), in the King Abdullah University of Science and Technology Ibn Sina field research station and conservation area (Saudi Arabia). Coastal expansion of mangrove in the Red Sea is limited by the small tidal range, approximately 20–30 cm in the central part of the Red Sea, combined with seasonal variations of water level, decreasing from May to August of approximately 50 cm and re-increasing thereafter till October, exposing mangroves to increasing period of emersion throughout summer (Pugh and Abualnaja 2015). The water column in our site classically varies between complete emersion and ~30–40 cm depending on the tide and period of the year, which is representative of Red Sea mangroves. Red Sea mangroves thrive in one of the driest areas of the planet, with precipitations ranging from 10 to
25 mm per month on average in 2018 (Global Precipitation Climatology Centre).

**Mangrove incubations**

The incubations were conducted during the last 2 h of the rising tide, around solar noon and at night within the same day, to distinguish between light and dark fluxes of TA and dissolved inorganic carbon (DIC). The incubations were conducted from April 2018 to May 2019. In April/May 2018, day and night incubations were done 15 d apart.

The incubation chambers in the mangrove consisted of transparent PVC pipes, open on both ends, enclosing a sediment surface of 71 cm² (9.5 cm diameter). The mean (± SD) volume of water in the chambers at the start of the incubations was 1.2 ± 0.4 L (height of 16.7 ± 5.4 cm) and increased by 0.20 ± 0.18 L (2.8 ± 2.6 cm) during the incubation, due to the rising tide.

This incubation method, with chambers open on both sides, allows the natural pore-water advective flow of compounds from the sediment to the water column. The problem of reproducing natural advection in benthic incubation systems is a key issue in the estimate of sediment – water fluxes (Tengberg et al., 2004). This is traditionally achieved through the generation of a rotary current within the chamber, creating a central depression pulling the solutes out of the sediment (Tengberg et al., 2004). The fluxes measured are then partly depending on the stirring force imposed in the chamber (Tengberg et al., 2004). In our mangrove incubations, the advection of the solutes from the sediment is ensured by the natural rise of water in the chambers during the rising tide.

The mean (± SD) duration of the incubations was 1.48 ± 0.22 h. Incubations of bare mangrove sediment and of bare sediment (without animal burrows) with one pneumatophore, tall enough to emerge from the water, were conducted. Each incubation event (for each day, night, and at different seasons) included 10 mangrove and 10 pneumatophore sediment incubations. Samples (100 mL) were taken at the beginning and end of incubations in each chamber for TA and DIC measurements. In parallel, samples were taken at the beginning and end of the incubation period in surface water outside of the mangrove as reference. Samples were poisoned after collection with HgCl₂ (Dickson et al. 2007).

Two conductivity – temperature – depth (CTD) sensors (Exo1, YSI Inc., Yellow Springs, USA) were installed in the mangrove at the incubation site and outside at the reference sampling point.

**Bare lagoon sediment incubations**

To test whether TA fluxes in mangrove sediments differ from adjacent unvegetated reef lagoon sediments, we carried out incubations with closed benthic chambers in parallel to the mangrove incubations on the 16.01.2019 and 25.05.2019, at daytime and nighttime, in 1–1.5 m water depth. The closed benthic incubation chambers were cylindrical, made of transparent polymethyl methacrylate, with a diameter of 0.5 m, a height of 0.39 m, a volume of 0.68 m³, and covered a sediment surface of 0.2 m². A pump in each chamber was placed to ensure the mixing of water and advection from sediment. A detailed description of the chambers and pumping systems can be found in Roth et al. (2019a). Samples for CO₂ system determinations were taken at the beginning and end of incubations (incubation time of 104–124 min). Samples were transferred to glass bottles and poisoned with HgCl₂ right after collection (Dickson et al. 2007).

**Sample analysis**

TA was measured using a dedicated open-cell titration instrument (AS-ALK2, Apollo SciTech). DIC was measured with an AS-C3 analyzer (Apollo SciTech) by infrared absorption spectrometry in the air phase after complete conversion into CO₂(g) by acidiﬁcation of the sample. The instruments were calibrated on every measurement day using certified CO₂ in seawater reference material (A. Dickson, Scripps Institution of Oceanography, La Jolla, California). Based on a global inter-calibration effort conducted in 2017 by A. Dickson and E. Bockmon (Scripps Institution of Oceanography, La Jolla, California), we estimate the mean ± SD accuracy of our measurements to be 0.21 ± 1.93 μmol TA kg⁻¹ and −3.34 ± 2.2 μmol DIC kg⁻¹. The salinity was measured in each sample using an Exo conductivity sensor (YSI).

**Calculations**

For each mangrove incubation, the TA production rate, in mmol m⁻² h⁻¹, was determined as the change of the total amount of TA (ΔTA_tot, in mmol) in the enclosed water column, from which the variation of TA due to changes in salinity (ΔTA_sal in mmol) was subtracted. The variation in salinity could be due to dilution/concentration with water from incoming tide, from advection of hypersaline pore water or due to evaporation. The relationship between TA and salinity is considered linear in seawater, and therefore any change in salinity causes a proportional change in TA.

\[
ΔTA_{tot} = TA_i V_i ρ_i - TA_f V_f ρ_f
\]

With TAᵢ and TAᵣ are the TA in mmol kg⁻¹ at the beginning and end of incubation, respectively; Vᵢ and Vᵣ are the initial and final volumes of water in the chamber at the beginning and end of incubation (in liter), calculated from the water column height in the chamber using basic cylinder volume formula; ρᵢ is the volumic mass (kg L⁻³) of seawater derived from the samples salinity and CTD temperature.

TA_sal is estimated using a TA to salinity ratio R (mmol kg⁻¹ psu⁻¹) of incoming tidal water, from measurements in the reference sample, taken in surface water outside the mangrove.

\[
R = \frac{TA_{ref}}{S_{ref}}
\]
\[ \Delta T_{\text{sal}} = RV_f S_i \rho_i - RV_i S_i \rho_i \]  \hspace{1cm} (3)

With \( S_{\text{ref}}, S_i, \) and \( S_f \) are the salinities in the reference, initial, and final samples.

\[ T_{\text{prod}} = \frac{\Delta T_{\text{tot}} - \Delta T_{\text{sal}}}{\Delta t \cdot \text{Area}} \]  \hspace{1cm} (4)

With \( \Delta t \) is the duration of the incubation as the difference between the initial and final time \( (t_i \) and \( t_f) \) in hours.

The method for the calculation of DIC production rates is presented in Supplementary Material and Method. See the DIC production rates in Supplementary Fig. S1.

**Statistical analysis**

To test the differences of our response variable TA, a fully factorial permutational multivariate ANOVA was used, considering our explanatory variable the daytime (two levels: day and night), the Season (three levels: Spring 2018, Winter 2018/2019 and Spring 2019), the type of sediment (two levels, pneumatophores, and bulk) (Supplementary Tables S1–S3). Prior to the analysis, normality of the data was tested. Euclidean distance was used to generate the dissimilarity matrix for the response variable TA. The analysis was performed using the function `adonis2()` in the “R” package “vegan” (Oksanen et al. 2019), and for pairwise comparison, the “R” package pairwiseAdonis (Martinez Arbizu 2019).

**Result and discussion**

**TA production rates in Red Sea mangroves**

The mean (± SE) net TA emission rates during daytime in the bare mangrove sediment and mangrove pneumatophore stands were 16.3 ± 2.7 and 15.4 ± 1.9 mmol m\(^{-2}\) h\(^{-1}\), respectively, similar to the net TA emission rates at nighttime (17.3 ± 1.6 and 18.3 ± 2 mmol m\(^{-2}\) h\(^{-1}\)) in bare sediment and mangrove pneumatophores, Fig. 2a). Neither the time of day nor the presence of a pneumatophore significantly influenced the TA fluxes (all \( p > 0.05 \); Table S1). In contrast, bare sediments in the adjacent lagoon were net sinks of TA at daytime with a mean (± SE) emission rates of \(-2.5 \pm 0.9\) mmol m\(^{-2}\) h\(^{-1}\) (Fig. 2a; Table S2). There was no clear pattern of emission or uptake of TA in the lagoon at nighttime \((0.32 \pm 0.6\) mmol m\(^{-2}\) h\(^{-1}\); Fig. 2a; Table S2). Although we did not find an apparent seasonal pattern in our data (Fig. 2b), we found significant changes over time \((p = 0.001\); Table S1), which differed between sediment types (bare mangrove sediment, sediment with pneumatophore and bare lagoon sand) and the time of the day (both \( p < 0.01 \); Table S1). Pooling together the two mangrove treatments (pneumatophore and bare sediment), all incubation events, and considering 12-h light : 12-h dark daily cycles, the mean (± SE) daily flux of TA from our mangrove site was \(403 \pm 17\) mmol m\(^{-2}\) d\(^{-1}\). This would correspond to an estimated yearly production of \(147 \pm 6\) or \(115 \pm 5\) mol TA m\(^{-2}\) yr\(^{-1}\) considering that the sediments in the mangrove stand are emerged 22% of the year (based on 2016–2017 time series from Saderne et al., 2019a).

The TA emission rates in the Red Sea mangrove reported here are higher than any rates previously reported, all in Australian mangroves (Faber et al. 2014; Sippo et al. 2016; Maher et al. 2018; Santos et al. 2019). Those studies estimated TA fluxes from sediment pore water sampling \((^{222}\text{Rn} \text{mass balance technique;} \) Faber et al. 2014; Santos et al. 2019) and/or at the scale of entire mangrove creeks or inlets using hydrological models coupled with 24 h seawater sampling (Faber et al. 2014; Sippo et al. 2016; Maher et al. 2018; Santos et al. 2019). From pore water, Faber et al. (2014) reported TA fluxes of 310 and 46 mmol m\(^{-2}\) d\(^{-1}\) in two mangrove inlets of Victoria (Australia) and estimated that the contribution of carbonate dissolution to this flux to be rather negligible (about 0.8%). Santos et al. (2019) measured a mean pore water

![Fig. 2. Net TA emissions rates. Mean ± SE hourly emission rates of TA (mmol m\(^{-2}\) h\(^{-1}\)); (a) at daytime and nighttime in mangrove bare sediment and pneumatophore incubations, and in lagoon sand incubations summarized by type of sediments; (b) summarized by seasons at day and night.](image-url)
leads to an uptake of 0.6 mol of CO2 per dissolved mole of CaCO3 (Frankignoulle et al. 1995; Abril and Frankignouille 2001). The next highest source of TA emission from the sediment is linked to the transformations of nitrogen in relation to OM degradation, net ammonification (ammonification minus nitrification), and denitrification (Krumins et al. 2013). These emissions can be associated with the Corg cycle. Including them into Blue Carbon assessments would require an estimate of the net TA generated during the cycle of primary production vs. OM degradation. The relationship between TA and DIC emission rates (Fig.3) confirms that the emitted TA does not solely originate from the dissolution of CaCO3 but from a combination of dissolution and redox processes (Fig. 3). However, we cannot determine the proportion of TA resulting from CaCO3 dissolution. The partitioning between the sources of TA emitted from Blue Carbon sediment remains a scientific and analytical challenge to date and requires further investigations.

Hence, considering that 65% of the TA production in reef, bank, and bay sediments is supported by carbonate dissolution (Krumins et al. 2013) and that one mole of CaCO3 dissolved results in the uptake of 0.6 mol of CO2, the mean TA mangrove production rate found in our study (147 ± 6 mol m⁻² yr⁻¹) would be equivalent to an atmospheric uptake of 345 ± 15 g C m⁻² yr⁻¹. This sink is 23-fold larger than the Corg burial rate for central Red Sea mangroves estimated at 15 g C m⁻² yr⁻¹ (Almahasheer et al. 2017). In comparison, in a carbonate poor mangrove of Australia (less than 1% sediment DW), the carbon sink from TA export was estimated 1.7-fold larger than OC burial rates (Sanders et al. 2016; Maher et al. 2018). The central Red Sea mangrove studied here would reach a carbon sink value of 360 g C m⁻² yr⁻¹, which is 2.2-fold above global mangrove mean based on Corg alone, estimated at 163 g C m⁻² yr⁻¹ (Breithaupt et al. 2012). The assumption that only 65% of the

**Fig. 3.** Stochiometry between DIC and TA emission rates in mangrove incubations. Data are the TA to DIC sediment emission rates for all incubations performed (day, night, all seasons). Red line: Linear regression between TA and DIC (F = 66, p < 0.0001, slope and intercept p < 0.0001). Vectors of relative length represent the TA to DIC stoichiometry of the main chemical processes occurring in sediments. Ammo.: ammonification.
TA mangrove production comes from carbonate dissolution, compared to 100% assumed in many studies (e.g., Van Dam et al. 2019 in seagrass meadows), renders our estimate of $345 \pm 15$ g C m$^{-2}$ yr$^{-1}$ of CO$_2$ removed from TA emission in the Red Sea mangroves a conservative one.

The dissolution rates found in our mangrove site can be compared to calcification rates observed in nearby coral reefs, a process that emits large amounts of CO$_2$ to the atmosphere globally (Frankignoulle et al. 1995; Borges et al. 2005). Roth et al. (2019b) found mean ($\pm$ SE) net calcification rates of $4.8 \pm 0.4$ kg CaCO$_3$ m$^{-2}$ yr$^{-1}$ in an adjacent coral reef, equivalent to a CO$_2$ emission of $344 \pm 26$ g C m$^{-2}$ yr$^{-1}$. The CO$_2$ uptake by carbonate dissolution in 1 ha of Red Sea mangrove, therefore, compensates for the CO$_2$ production by calcification of 1 ha of Red Sea coral reef. Mangrove ecosystems are often growing on underlying carbonate platforms, principally due to the presence of extant and fossil coral reefs in the same latitudes (Saderne et al. 2019b). Our results indicate that assessments of the Blue Carbon potential of mangroves need be extended to consider CO$_2$ removal from the TA emission in mangrove stands on carbonate deposits. Likewise, targeting mangroves along carbonate deposits for restoration and expansion increases the potential climate change mitigation of mangrove restoration projects.

**Conflict of interests**

The authors declared no potential conflict of interests.

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