Calcification-driven CO₂ emissions exceed “Blue Carbon” sequestration in a carbonate seagrass meadow

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Long-term “Blue Carbon” burial in seagrass meadows is complicated by other carbon and alkalinity exchanges that shape net carbon sequestration. We measured a suite of such processes, including denitrification, sulfur, and inorganic carbon cycling, and assessed their impact on air-water CO₂ exchange in a typical seagrass meadow underlain by carbonate sediments. Eddy covariance measurements reveal a consistent source of CO₂ to the atmosphere at an average rate of 610 ± 990 μmol m⁻² hour⁻¹ during our study and 700 ± 660 μmol m⁻² hour⁻¹ (6.1 mol m⁻² year⁻¹) over an annual cycle. Net alkalinity consumption by ecosystem calcification explains >95% of the observed CO₂ emissions, far exceeding organic carbon burial and anaerobic alkalinity generation. We argue that the net carbon sequestration potential of seagrass meadows may be overestimated if calcification-induced CO₂ emissions are not accounted for, especially in regions where calcification rates exceed net primary production and burial.

INTRODUCTION

Seagrass ecosystems are some of the most organic carbon (OC)-dense systems on Earth, and it has been argued that OC sequestration here is disproportionately large in comparison to other terrestrial and marine ecosystems, thus constituting an important sink in the global carbon cycle (1, 2). Presently, there exists an apparent consensus that the protection and enhancement of such “Blue Carbon” storage in seagrass meadows is an effective strategy to mitigate increasing atmospheric CO₂ levels (3–7). However, biogeochemical cycling in seagrass ecosystems is complex, and many other processes exist, which may counteract net OC sequestration. These processes collectively regulate local budgets of dissolved inorganic carbon (DIC) and total alkalinity (TA), and include ecosystem calcification (8–10) and anaerobic metabolism (11). When net TA production occurs, the resulting carbonate system reequilibration consumes CO₂, which is then compensated by net CO₂ uptake from the atmosphere. The impact of these TA-generating processes [namely, iron (Fe), sulfate (SO₄²⁻), and nitrate (NO₃⁻) reduction] on net CO₂ uptake depends on the ultimate fate of the metabolic products produced. Net TA generation will only occur if these metabolic products are permanently removed in their reduced form [via long-term burial of reduced Fe or sulfur (S) or release of N₂ gas following denitrification] but will not if the reduced species are reoxidized in place. Competing with these TA sources, a key sink for TA (source of CO₂) in Blue Carbon habitats is the precipitation of carbonate minerals (5, 12, 13). This is especially so in tropical and subtropical seagrasses, which play a disproportionately large role in the global carbonate cycle (14). The impact of these internal reworkings of S, Fe, N, and carbonate minerals on surface water TA/DIC and, ultimately, air-sea CO₂ exchange remains largely unknown. This has been identified as a key gap in our understanding of the role of Blue Carbon ecosystems in the global carbon cycle (10, 15).

Despite the understanding that carbonate precipitation generates CO₂, carbonate seagrass meadows are still considered as important Blue Carbon sinks based largely on the assumption that these carbonate minerals were formed elsewhere (10, 12), or that the CO₂ produced by calcification is rapidly consumed by photosynthesis (16). As a result, policies aimed at restoring or protecting Blue Carbon habitats have gathered momentum as an approach to fight climate change while also achieving co-benefits of habitat improvement (4, 5, 17). In recent Intergovernmental Panel on Climate Change (IPCC) reports, these ocean-based mitigation tools have been assigned a high chance of success (IPCC Chapter 5.5.1.2.2) (18), despite the lack of empirical evidence demonstrating net CO₂ uptake by carbonate seagrass meadows (13). Still, uncertainty in the scientific community regarding the role of calcification in Blue Carbon mitigation is explicitly acknowledged by the IPCC, and its timely resolution is seen as “highly desirable” (18). Blue Carbon mitigation requires public and private buy-in (4, 19), and a general understanding that the actions taken will indeed enhance net carbon sequestration. Many of these Blue Carbon habitats are managed by resource-limited nations, introducing a climate injustice risk, as there exists a perception that they are being asked to carry the climate burden of more industrialized nations (20). It is therefore crucial that these ocean-based management actions only be enacted for Blue Carbon habitats where increased carbon sequestration is plausible.

To address these uncertainties, we used atmospheric eddy covariance (EC) to directly measure air-water CO₂ exchange in Florida Bay, USA, one of the largest seagrass-dominated estuaries in the world, and a known OC sink (21). We combine these EC fluxes with geochemical approaches to attribute the major processes contributing to source or sink behavior. We sampled at three locations in close proximity within this seagrass meadow, representing regions of (i) high...
seagrass aboveground biomass [high density (HD)], (ii) low seagrass density (LD), and (iii) bare sediment (B). The selection of these sites was intended to characterize the seagrass meadow occupying the EC footprint. We combined pore-water and solid-phase analyses with a continuous-flow incubation that either included or excluded living seagrass biomass. This approach lets us ascribe changes in surface water TA and DIC to net process rates in the sediment, which were, in turn, integrated into a biogeochemical budget, assessing their effects on air-water CO₂ uptake or release.

RESULTS

Net CO₂ emissions associated with carbonate mineral dissolution/reprecipitation

Direct EC measurements reveal this seagrass meadow as a moderate source of CO₂ to the atmosphere (Fig. 1, A and B), with average emissions during the ~1-week study period (10 to 18 November 2019; red vertical line in Fig. 1B) of 610 ± 990 μmol m⁻² hour⁻¹ (green line in Fig. 1B; mean ± 1 SD of 30-min records). This is slightly below the previously reported summer-time CO₂ flux for this site (972 ± 612 μmol m⁻² hour⁻¹) (22) and just below the annual average (700 ± 660 μmol m⁻² hour⁻¹ or 6.1 mol m⁻² year⁻¹; black line in Fig. 1B). While annual hourly climatology demonstrates a clear diel trend in the warmer spring and summer months, with greater CO₂ emissions during the afternoon [Fig. 1A, green (spring) and blue (summer) circles], such a day-night difference was not present during the study period, which occurred in the fall (Wilcoxon, P = 0.61; Fig. 1A, black circles). These seasonal trends are consistent with the hypothesis that factors other than seagrass net ecosystem metabolism control the CO₂ system at this site, contrasting with previous studies describing the importance of water column productivity for seasonal CO₂ uptake in fall (23, 24). In the case of air-water CO₂ flux, this nonmetabolic forcing includes the role of temperature in enhancing convection-mediated air-water gas transfer (22). These CO₂ emissions are likely an underestimate of the global warming potential of this seagrass meadow, which is likely enhanced by the release of other greenhouse gases (25) like CH₄ and N₂O.

In line with strong carbonate dissolution, we observed notable excesses of pore-water DIC coinciding with a moderate (LD) or large (HD) enrichment of ¹³C in DIC such that average pore-water ¹³C_DIC (−1.7 ± 1.4‰) was at least 2‰ “heavier” than surface water (³¹³C_DIC = −3.9 ± 0.02‰) (Fig. 2). For sediment depths outside of the zone of peak DIC accumulation, a Keeling plot points toward an isotopic endmember of ~0‰, characteristic of “heavy” DIC from carbonate mineral dissolution mixed with “light” respiratory DIC from dissolved OC (DOC) degradation (³¹³C_DOC = −18 ± 1.9‰). However,

![Fig. 1. Summary of measured air-water and sediment-water fluxes.](image-url)
within the DIC maximum zone in the HD cores, DIC was enriched in $^{13}$C, with an average $\delta^{13}$CDIC of $0.0 \pm 1.1\%$ ($-2.1 \pm 0.6\%$ in the LD cores). The isotopic endmember indicated for this “DIC-maximum zone” is well above measured particulate IC (PIC) $\delta^{13}$C PIC of $1.9 \pm 0.12\%$. Enrichment in $^{13}$C of this magnitude cannot be explained solely by CaCO$_3$ dissolution. Rather, this enrichment necessitates the consumption of isotopically light pore-water DIC by carbonate precipitation under quasi–closed-system conditions (26). This hypothesis is supported by the modeled DIC production rates (Fig. 2F), where net DIC generated below 10 cm depth is consumed in the upper 10 cm of sediment by combined autotrophic sulfide oxidation and carbonate reprecipitation. Likewise, decreased solid-phase Ca: Sr and Ca: Mg ratios in the upper 10 to 20 cm of sediment support carbonate mineral recrystallization in this region (Fig. 3, C and D). Our findings are consistent with previous work showing this coupled dissolution/reprecipitation to be highest when seagrasses are especially dense (26, 27). This internal carbonate recycling is critically important to the overall C cycle, as it regulates the burial efficiency of CaCO$_3$, ultimately governing CO$_2$ emissions from the sediment related to net carbonate burial.

**Denitrification and sulfate reduction are only minor alkalinity sources**

Pore-water total dissolved sulfide was elevated relative to surface water (Fig. 2E), especially in the HD cores, and was generated by net sulfate reduction at depths of ~10 cm (LD) to ~15 cm (HD) (Fig. 2G). This excess sulfide corresponded to a large modeled vertical sulfide flux (Fig. 1C) that was presumably reoxidized in the narrow oxic zone of the sediments or the water column. All dissolved sulfide samples from the sediment flux incubation were below the limit of detection, confirming that reduced sulfur diffusing upward is quantitatively lost to either (i) oxidation to sulfate or (ii) burial as FeS$_2$ + S$_0$ (Fig. 3A), or (iii) burial as FeS. We expect that sulfide oxidation is largely limited to depths of 0 to 30 cm, as the relative increase in Mo (a redox-sensitive tracer) below 30 cm depth (Fig. 3B) indicate permanent anoxia below this threshold. While bulk sediments of the upper 0 to 30 cm are also anoxic, the relative Mo depletion in this zone supports localized and sporadic oxidation in the millimeters surrounding seagrass roots. The impact of net sulfate reduction on TA depends on the fate and form of net reduced sulfur burial. To this point, we observe substantial accumulation of...
chromium (II)–reducible sulfur (CRS; FeS$_2$ + S$^0$), with an average content of 94 ± 37 μmol CRS g$^{-1}$ (0.3 ± 0.12% dry weight), and a peak of 200 μmol g$^{-1}$ in the peak sulfate reduction zone of site HD. Lower concentrations of acid-volatile sulfide (AVS) (maximum of 3.6, average of 2.4 ± 1.8 μmol g$^{-1}$) indicate a relatively minor contribution of H$_2$S and FeS to total sulfur burial. Because the total Fe content (44.9 ± 5.3 μmol g$^{-1}$) is half of average CRS, at most ~100% of reduced sulfur may be buried as pyrite (FeS$_2$). However, previous work (28) shows that 50% or more of the sedimentary Fe pool in central Florida Bay is not directly associated with CRS, suggesting substantial burial of elemental sulfur (S$^0$) as a product of partial sulfide re-oxidation (29). This is supported by the CRS peak at 10 cm at the HD site, possibly caused by sulfate oxidation to sulfur at the surface of seagrass roots (30). Assuming that CRS is composed of equal proportions FeS$_2$ and S$^0$, net sulfate reduction and burial will produce TA and DIC in a ratio of 2:1 (Table 2).

Continuous-flow incubations conducted in the dark showed that denitrification was closely balanced by N$_2$ fixation (Fig. 1C), and only cores from the HD site with seagrass biomass included were, on average, net denitrifying (positive N$_2$ flux). In contrast, all other cores were net N$_2$ fixing (negative N$_2$ flux), despite dark conditions favorable to net denitrification (Fig. 1C) (31). Averaged across all cores, net denitrification was not significantly different from zero (± 27 μmol m$^{-2}$ hour$^{-1}$), consistent with the understanding that denitrification in seagrass meadows is spatially heterogeneous, tends to covary with autochthonous organic matter input (32, 33), and is part of a highly conservative N cycle. Our low denitrification measurements are also consistent with the low nitrate concentrations typical of many carbonate seagrasses (34), where all available nitrate appears to be quantitatively denitrified. That net denitrification was only observed in cores from the HD site containing living seagrass biomass (Fig. 1C), where pore-water sulfide was also maximized (Fig. 2E), contrasts with the expected sulfide inhibition of denitrification in Florida Bay (33). Instead, denitrification may have been enhanced by the leaching of labile OM (organic matter) leaching from the seagrass rhizosphere. Likewise, the dissolution of carbonate sediments,
as expected during this dark/heterotrophic incubation, is a potential abiotic source of dissolved OM through the release of carbonate-associated organic matter (35). We posit that these abiotic and biotic OM sources together drove our large measured DOC fluxes. Broadly, the observation of strong internal recycling with respect to N and S supports the notion that alkalinity dynamics in this carbonate seagrass meadow are dominated by CaCO₃ precipitation and dissolution, rather than anaerobic alkalinity generation via nitrate, sulfate, or metal (oxy)hydroxide reduction.

**CO₂ emissions are driven by net CaCO₃ production**

We detected consistent net CO₂ emissions from this seagrass meadow of 610 μmol m⁻² hour⁻¹ over the study period and 700 μmol m⁻² hour⁻¹ (6.1 mol m⁻² year⁻¹) as an annual average (Fig. 1B). This is notable given the consensus impression of seagrasses as net Blue Carbon sinks, and in light of our understanding of these seagrasses as being net autotrophic (36). That CO₂ emissions persisted at this site, despite seagrass ecosystem productivity and known net OC burial (21), suggests that processes besides seagrass net metabolism drive air-water CO₂ exchanges. To find the dominant carbon source for these CO₂ emissions, we constructed a simple TA and DIC budget, considering the impact of calcification (IC accumulation), net Fe³⁺ and SO₄²⁻ reduction (CRS), and net OC burial. Measured denitrification was not significantly different from 0; hence, it was excluded as a putative TA source. CO₂ consumption by net OC accumulation reduced excess CO₂ by 3.4%, while the burial of CRS and Fe (i.e., net SO₄²⁻ and Fe³⁺ reduction) only reduced excess CO₂ by ~1%. When these TA/DIC consuming and producing processes are summed, we find that excess CO₂ available for release to the atmosphere (solid black line in Fig. 4A) increases with sediment accumulation. This excess CO₂ generated by IC accumulation explains the remaining 95.8% of the change in excess CO₂ (Fig. 4A). Using an average gas transfer velocity of 11.7 cm hour⁻¹, our measured CO₂ fluxes can be converted to excess CO₂ values of between 6.0 and 5.2 μmol kg⁻¹ (gray dashed lines in Fig. 4, A to C), within the range of excess CO₂ in the budget. Furthermore, the sediment accumulation rate (SAR) required to sustain measured annual average CO₂ emissions (700 ± 660 μmol m⁻² hour⁻¹ or 6.1 mol m⁻² year⁻¹) is only 460 g sediment m⁻² year⁻¹ (vertical dotted line in Fig. 4A), intermediate of literature values for central Florida Bay (37, 38).

To find the rate of OC burial required to balance the CO₂ produced by calcification, we repeatedly ran the model, increasing the fraction of sediment composed of OC until total excess CO₂ was equal to 0. For these processes to balance out, the sediment OC fraction would need to increase by a factor of nearly 20, from 2.1 to 39% (compensated by IC decrease from 97.5 to 60.6%; Fig. 4B). OC contents of this order are much greater than those that can be found in carbonate seagrass meadows elsewhere in Florida Bay (39) or globally (40, 41). In a final model, we increased OC burial by only one order of magnitude (to 21%), but increased CRS and Fe fractions by a factor of 45, simulating an artificial Fe amendment scenario. In this admittedly unrealistic scenario, CO₂ emissions due to calcification are balanced approximately equally by OC burial and anaerobic TA generation by net sulfate and Fe reduction. In these additional scenarios (Fig. 4, B and C), some unknown additional CO₂ source would be required to bring modeled excess CO₂ (thick black line) in line with the excess CO₂ associated with measured CO₂ emissions (gray dashed lines). As nearly one-third of seagrass meadows (12) overlie sediments with CaCO₃ contents exceeding 80% dry weight, it is likely that our observation of calcification-driven CO₂ emissions from this seagrass meadow applies to carbonate systems globally.

**DISCUSSION**

**Progress toward realistic Blue Carbon accounting**

Previous Blue Carbon syntheses have suggested carbonate seagrass meadows as strong carbon sinks, based largely on the assumption that much of the CaCO₃ was allochthonous (10, 12). The present study site in central Florida Bay is isolated from coral reefs and other calcifying ecosystems, meaning that all the CaCO₃ buried here is produced in situ (38, 42). Measured rates of lime mud production in Florida Bay suggest that Florida Bay exports carbonates rather than importing them (42). In addition, extensive mud banks throughout Florida Bay restrict horizontal exchange (43), further decreasing the chance of internal CaCO₃ redistribution, as previously suggested (12).

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**Fig. 4. Modeled excess CO₂ across a range in literature sediment accumulation rates.** In (A), sediment fractions of CRS, Fe, IC, and OC were set according to direct measurements from this study. In (B), the OC fraction was artificially increased until excess CO₂ remained at 0 across the range in sediment accumulation such that CO₂ released by calcification was offset by CO₂ consumed by OC burial and Fe and SO₄ reduction. In (C), net CO₂ balance was achieved by increasing OC, Fe, and CRS fractions by factors of 10, 45, and 45, respectively.

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Therefore, our direct EC measurements, which reveal net annual emissions of 700 ± 660 μmol CO₂ m⁻² hour⁻¹ (6.1 mol m⁻² year⁻¹), place this seagrass meadow as a clear net CO₂ source to the atmosphere. Calcification explained essentially all the measured CO₂ emissions and exceeded the OC sink by more than 300%, rather than offsetting 30 to 40% of OC burial as previous work indicated (12). This net CaCO₃ burial is sustained by carbonate dissolution and reprecipitation, enhancing CaCO₃ burial efficiency and ultimately shaping CO₂ emissions. While OC burial does offset a small fraction (3.4%) of these emissions, an unrealistic 20-fold increase would be required for the site to become a net CO₂ sink. The impact of TA productivity, denitrification, etc., on net CO₂ emissions vary with seasonal variations in temperature and salinity. The seasonal increase in afternoon CO₂ emissions during warmer months (Fig. 1A) is consistent with the role of temperature in enhancing net CO₂ release. This is likely a combined result of increased ecosystem calcification rates, along with the role that temperature plays in physically enhancing air-water gas transfer through convective forcing as documented previously at this site (22).

Our empirical approach, supported by micrometeorological and isotope geochemical tools, contributes fundamental evidence toward resolving one of the key uncertainties in Blue Carbon accounting (13, 18). This approach is especially suitable for other ecosystems considered for Blue Carbon storage where hydrodynamic forcing is minimal, allowing measured air-water CO₂ exchanges to be more reliably linked to benthic biogeochemical processing. However, a decoupling of benthic processes from air-water exchanges may occur when tidal exchanges are dominant, challenging this approach in more complex estuarine environments. Furthermore, our study considered a seagrass meadow with relatively low biomass and P limitation, but in seagrass meadows with increased P availability, greater standing biomass, enhanced primary production, or net carbonate dissolution (35), CO₂ emissions could be decreased facilitating net C storage. Nevertheless, as more than a quarter of all Blue Carbon habitats also have high carbonate contents (>80% dry weight) (12), reliable carbon balancing such as this study will help identify cases of “surprise” CO₂ emissions (44). Our study motivates a more rigorous (re-)assessment of the coastal carbon balance, which is essential to protect public and private sectors’ initiatives and investments in ocean-based mitigation of climate change as an accountable contribution to the COP21 Paris agreement.

MATERIALS AND METHODS
Site description and sample collection
All collection of discrete samples was carried out in November 2019 at a well-characterized and shallow site (~1 m depth, 25°1.718’ N, 80°40.736’ W) near Bob Allen Keys in central Florida Bay, one of the largest seagrass-dominated estuaries in the world. Sediments consist of 70 to 90% of carbonate minerals (>90% for Bob Allen Keys) (21), mostly aragonite and high Mg calcite (26), with relatively low iron content (28, 45), and only trace amounts of silicate minerals (46). The site is a sparse-medium density seagrass meadow dominated by Thalassia testudinum (turtle grass) with occasional occurrences of Halodule wrightii and sparsely distributed macroalgae. Low phosphorus availability limits the development of dense seagrass meadows in this part of Florida Bay (47). We characterized the species composition and relative abundance with surveys occurring six times per year for the period 2000–2021 along a permanently marked 50-m transect. During each survey, all conspicuous taxa in 10 randomly placed 0.25 m² sampling quadrats along the transect were identified and assigned a Braun-Blanquet cover-abundance score (48, 49). We transformed the cover and abundance scores to percent cover (defined as the fraction of the bottom obscured by the taxon of interest when viewed from above) assuming the midpoint of each cover range for each cover-abundance score. Over the 112 surveys conducted over the 22-year period of record, T. testudinum was the dominant species, averaging 18.6 ± 1.0% cover (Table 1). The only other seagrass species encountered was H. wrightii, which was detected in 18 of the surveys with a low period-of-record average abundance of 0.05 ± 0.01% cover (Table 1). Calcifying and non-calcifying macroalgae were also present, but at lower percent cover than the dominant T. testudinum. Of these macroalgae, Penicillus sp. was the most often encountered and was observed on 100 of the 112 total transects (at an average of 0.8 ± 0.1% cover). The macroalga Batophora sp. had the highest average percent cover of 2.8 ± 0.7% but was present less frequently.

Both the aboveground biomass and the aboveground net primary productivity of T. testudinum were seasonal at the site and showed no long-term monotonic trends. We measured productivity using the leaf-mark technique (50) in six 10 cm–by–20 cm quadrats six times per year from the period 2000–2021. Aboveground biomass was assessed as the mass of green leaves in the productivity quadrats and averaged 22.2 ± 1.8 g (dry weight) m⁻² (±1 SE, n = 96; Fig. 5), and aboveground net primary productivity averaged 0.40 ± 0.04 g (dry weight) m⁻² day⁻¹. For each biomass sample, we dried the compounded leaves and ground them to a fine powder for analysis of C content using an elemental analyzer; T. testudinum leaves averaged 38.3 ± 0.2% C (n = 125) over the 22 years of sampling. The C stocks in the aboveground biomass of T. testudinum (which comprised the vast majority of the total aboveground biomass of macrophytes at the site) averaged 0.085 Mg C ha⁻¹, falling below the global median C stock (1) of 0.264 Mg C ha⁻¹. This low seagrass biomass is typical of this region of Florida Bay (51) and representative of the seagrass meadows of the footprint of our EC tower. Our average percent cover of T. testudinum of 18.6% was similar to the percent cover detected in the Florida Fish and Wildlife Institute’s Fisheries Habitat Assessment Program (FHAP) for Florida Bay that has been assessing percent cover using the Braun-Blanquet technique twice per year since 1995 (52). The FHAP program uses a stratified-random procedure to distribute 30 observation points evenly across basins of Florida Bay for each survey. In the 36-km² basin immediately to the north of our study site, T. testudinum averaged 12.4 ± 0.4% cover for 1011 observations over the period of record. In the 20-km² basin immediately to the east of our study site, T. testudinum cover averaged 14.3 ± 0.4%. No FHAP data exist for the basin to the west of our study site, but it has a visually similar characteristic seagrass meadow as the site and the other two basins.

We assessed net benthic metabolic rates at three locations in close proximity within this seagrass meadow, representing regions of (i) high seagrass aboveground biomass (HD), (ii) low seagrass density (LD), and (iii) bare sediment (B). We collected samples at these three sites to characterize as fully as possible the seagrass meadow that occupies the EC footprint. Quadruplicate cores containing approximately 10 cm of sediment were collected for a continuous-flow incubation
in which net sediment-water fluxes were directly measured using cores that either included or excluded living seagrass biomass. This steady-state incubation does not represent net ecosystem metabolism because it was conducted in the dark. Rather, the intent was to create net conditions representative of night-time heterotrophy when denitrification is expected to be greatest (31), thereby providing a conservative estimate for the maximum anaerobic alkalinity that could be expected, across a realistic range in seagrass density. Separately, duplicate 30- to 50-cm sediment cores were collected at HD and LD sites for pore-water and solid-phase analysis.

**Sediment and pore-water sampling and analysis**

For solid-phase characterization, we collected and froze 50-cm-long acrylic cores at both HD and LD sites, which were thawed and subsampled at 5-cm intervals within 1 week of sampling. Sediment compaction due to inserting the tubes was not visible or indicated by the vertical bulk density profiles. Following transport on dry ice back to the laboratory, sediment was freeze-dried, ground (PM 400, Retsch), and microwave-digested (MARS 5 Xpress, CEM Corp.) in a solution of 5-ml concentrated HNO$_3$ and 2-ml concentrated HCl before trace-metal analysis described below. Frozen sediment was freeze-dried and analyzed for $\delta^{13}$C of the inorganic carbonate fraction ($\delta^{13}$C$_{PIC}$) or preserved with 25% Zn(oAc)$_2$ for the analysis of CRS (a measure for $S^0$ and FeS$_2$) and AVS (a proxy for H$_2$S and FeS). The AVS and CRS fractions were extracted from the sediments according to a two-step distillation procedure (53), as described in (54, 55). Duplicate acrylic cores of 25 to 35 cm length were further collected from HD and LD sites, and pore water was retrieved at 5-cm-depth resolution using Rhizons (0.12-μm-diameter filtration) on the same day as sampling. Pore water was preserved with HgCl$_2$, stored cool and dark, and analyzed for the concentration and stable carbon isotope composition of DIC ($\delta^{13}$C$_{DIC}$) (56) using a Thermo Finnigan MAT 253 gas mass spectrometer coupled to GasBench II. Pore-water aliquots preserved with 5% Zn(oAc)$_2$ were used for spectrophotometric measurement of the total dissolved sulfide concentration (57) using an Analytik Jena AG, Specord 40 photometer (56). $\delta^{13}$C measurements of DIC and CaCO$_3$ were carried out via isotope ratio monitoring mass spectrometry as described by (58) and (59), respectively. The resulting per mil values are equivalent to mUr [milli-Urey (60)]. Isotope ratio measurements are given with respect to the international Vienna Pee Dee Belemnite (VPDB) standard.

**Sediment trace metal analysis**

Sediment solids were sampled and frozen, as described above, before trace metal analysis. Type I reagent-grade water (>18.2 megohm-cm) for trace metal analysis was obtained from an ultrapure water system consisting of an Elix 3 module (Merck Millipore), a Milli-Q element module (Merck Millipore), and a Q-POD element (Merck Millipore). Analytical grade HNO$_3$ (65%, w/w; Fisher Scientific GmbH) and analytical grade HCl (30%, w/w, Carl Roth GmbH + Co. KG) were further purified by double subboiling in perfluoroalkoxy-polymer (PFA)–subboiling stills (DST-4000 and DST-1000, Savillex). Sediment was freeze-dried (Beta 1-8, Christ Gefriertrocknungsanlagen), ground (PM 400, Retsh), and microwave-digested (MARS 5 Xpress, CEM Corp.) in a solution of 5-ml concentrated HNO$_3$ and 2-ml

| Taxon               | Mean ± 1 SE | No. of occurrences | Minimum | Maximum |
|---------------------|-------------|--------------------|---------|---------|
| **Seagrasses**      |             |                    |         |         |
| *T. testudinum*     | 18.6 ± 1.0  | 112                | 2.2     | 57.5    |
| *Halodule wrightii* | 0.05 ± 0.01 | 18                 | 0       | 1       |
| **Macroalgae**      |             |                    |         |         |
| **Chlorophyta**     |             |                    |         |         |
| Penicillus sp.      | 0.8 ± 0.1   | 100                | 0       | 3.8     |
| Rhipocephalus sp.   | 0.002 ± 0.001 | 3                | 0       | 0.125   |
| Acetabularia sp.    | 0.002 ± 0.001 | 2                | 0       | 0.125   |
| Neomeris sp.        | 0.05 ± 0.04 | 3                  | 0       | 3.75    |
| Cymoporia sp.       | 0.07 ± 0.01 | 54                 | 0       | 0.5     |
| Batophora sp.       | 2.8 ± 0.7   | 76                 | 0       | 57.5    |
| Dasycladus sp.      | 0.2 ± 0.1   | 11                 | 0       | 15      |
| Other Chlorophyta   | 1.4 ± 0.3   | 49                 | 0       | 15      |
| **Rhodophyta**      |             |                    |         |         |
| Drift algae         | 0.3 ± 0.2   | 10                 | 0       | 11.25   |
| Other Rhodophyta    | 0.2 ± 0.1   | 9                  | 0       | 11.25   |
| **Phaeophyta**      |             |                    |         |         |
| Sargassum sp.       | 0.014 ± 0.007 | 5                | 0       | 0.5     |
| Other Phaeophyta    | 0.4 ± 0.2   | 7                  | 0       | 21.75   |
| **Sponges**         | 0.12 ± 0.02 | 46                 | 0       | 1.1     |
concentrated HCl before trace-metal analysis (61) by inductively coupled plasma–tandem mass spectrometry (ICP-MS/MS) (Agilent 8800, Agilent Technologies). Digestion took place at 180°C for 300 min using 35-ml precleaned TFM digestion vessels. ICP-MS/MS was coupled to an ESI SC-4 DX FAST autosampler (Elemental Scientific) equipped with a discrete sampling system with a loop volume of 1.5 ml. Selection of the analyzed isotope, as well as cell gas mode, was based on the achieved sensitivity and occurrence of spectral interferences (isobaric and polyatomic interferences): 24 Mg (H2 mode), 40 Ca (H2 mode), 56 Fe (H2 mode), 88 Sr (He mode), and 95 Mo (O2 mass shift mode). The instrument was optimized on a daily basis using a tune solution containing Li, Y, Ce, and TI (10 μg liter−1). Quantification was performed on the basis of external calibration, covering a concentration range from 0.1 to 100 μg liter−1 for all analytes and from 10 to 10,000 μg liter−1 for Ca. Solutions were prepared on a daily basis from custom-made multielement standards (Inorganic Ventures). Multielement data were processed using MassHunter versions 4.4 (Agilent Technologies) and a custom-written Excel spreadsheet. The significant number of digits of mass fractions is given according to GUM (Guide to the Expression of Uncertainty in Measurement) and EURACHEM guidelines, whereby the uncertainty determines the significant number of digits to be presented with the value. A detailed description of the trace metal determination can be found in the literature (61).

Fig. 5. Time series of aboveground biomass and aboveground net primary productivity of the dominant seagrass T. testudinum at the EC tower site for the period of record demonstrating seasonality but no long-term monotonic trends.

**Measured sediment-water flux**

Continuous flow experiments with intact sediment cores were used to determine fluxes of dissolved gases across the sediment-water interface (33). Quadruplicate cores of 10 cm deep, 6.4 cm diameter with overlying water were collected at site B, and at two separate subsites for HD and LD, one set containing aboveground seagrass biomass, and the other omitting biomass for 20 cores total. These cores were incubated under continuous flowing water at a flow rate of 1.5 ml/min with aerated site water at room temperature (21.6°C) and in the dark. Cores were preincubated for 24 hours to allow the system to reach steady state (33). This setup prevented changes between net photosynthesis and respiration driven by light–to–dark transitions, a necessary step to prevent the formation of bubbles that can affect dissolved gas measurements and ensure that calculated N2 and DOC fluxes were representative of steady-state conditions (62). Inflow and outflow samples were collected five times over the next 48 hours. Sediment-water fluxes of N2 (positive flux is net denitrification in excess of nitrogen fixation) and DOC (Fig. 1) were determined from the difference between each outflow and inflow of site reference water, flow rate, and surface area (62). Samples for N2 were preserved with 100 μl of saturated ZnCl2. N2 was measured by membrane inlet mass spectrometry. Samples for DOC analysis were filtered and acidified with HCl and frozen before analysis. DOC concentration was measured on a modified OI Analytical Aurora Model 1030 TOC analyzer using a high-temperature combustion method, coupled to a Delta V Plus isotope ratio mass spectrometer for δ13C measurement (63). Samples were acidified to PH 2 with 85% phosphoric acid and sparged using ultrahigh-purity argon. DOC values were blank-corrected with ultrapure Milli-Q water (18.2-megohm resistivity) and calibrated with caffeine standards, ranging from 83 to 3320 μmol C liter−1. Both caffeine and sucrose standards of known carbon isotopic composition obtained from the International Atomic Energy Agency (IAEA C6) were used to express stable isotope values in the delta notation “δ,” where the ratio (R) of 13C to 12C for an unknown sample is calibrated against a known standard

\[
\delta^{13}C (\text{‰}) = [(R_{\text{sample}} / R_{\text{standard}} - 1) \times 1000]
\]

The values are reported as per mil relative to VPDB. The precision of the analysis for [DOC] was ±16 μmol C liter−1 and ±0.2‰ for δ13C-DOC. By convention, positive fluxes represent a release from
the sediment to the overlying water and negative fluxes are movement from the water column to the sediment.

**Modeled sediment-water flux**

Direct measurements of sediment-water fluxes were augmented by advective-diffusive modeling in PROFILE (64) for H$_2$S and DIC. We assumed no irrigation and estimated biodiffusivity in the upper 15 cm of sediment at 1 and 3 cm$^2$ hour$^{-1}$ (0.00028 and 0.00083 cm$^2$ s$^{-1}$) at the LD and HD sites, respectively, representing seagrass O$_2$ pumping into the rhizosphere. We arrived at these parameterizations through iterative calculation runs attempting to match model output DIC concentration profiles and vertical flux with measured DIC profiles and fluxes. Molecular diffusivity was calculated at in situ temperature and salinity using the “marelac” package (65) in R. Boundary conditions at the top were set by the surface water concentration and at the bottom of the model domain by a diffusive flux calculated from the gradient across the bottom two concentration measurements.

**Budget creation**

Last, we constructed a carbon and alkalinity budget, combining literature SARs (37, 38) with our solid-phase results to estimate net accumulation of CaCO$_3$, and reduced sulfur and iron species. Using the stoichiometry outlined in Table 2, these burial rates were converted into net fluxes of DIC and TA, which were, in turn, used to calculate the CO$_2$ production or consumption due to each biogeochemical process. The ultimate goal of this exercise was to compare budgeted excess CO$_2$ with the excess CO$_2$ that would be required to sustain the CO$_2$ fluxes that were directly observed in this study.

We assumed a range in literature carbonate accumulation rates (37,38) of 1.9 to 1042 g m$^{-2}$ year$^{-1}$ or 0.019 to 10.4 mol CaCO$_3$ m$^{-2}$ year$^{-1}$. According to our solid-phase analysis, this sediment is 97.5% IC, 2.1% OC, 0.3% CRS (representing FeS$_2$ and S$^0$), and approximately 0.5% “free” Fe (calculated as the difference between CRS and total Fe). The net impact of production and burial of IC and OC on TA and DIC is shown in Table 2. The role of sulfate reduction is rather more complicated, as the sulfide produced by sulfate reduction ($\Delta$TA: DIC = 1:1) may or may not react with various phases of Fe, with various effects (66) on TA. Therefore, we take a simplifying approach of assuming that CRS burial occurs in equal proportions as FeS$_2$ ($\Delta$TA = 0) and S$^0$ ($\Delta$TA = 2) such that the net burial of reduced sulfide results in the production of TA and DIC in a ratio of 2:1 (Table 2). The net formation of free Fe produces TA and DIC in a ratio (66) of 8:2. Our modeling and experimental results demonstrate that N cycling is tightly internally recycled such that the net rates are indistinguishable from zero, meaning that the impact of net denitrification on sediment TA release can be neglected from this model. Similarly, because NO$_x$ flux was balanced very closely by NH$_4$ flux (not shown in figures), we can assume that the net effect of net ecosystem production on TA was small (as indicated by the asterisk in Table 2). These molar equivalents were combined with literature SARs to estimate the change in water-column TA and DIC due to each of the abovementioned processes [IC burial, OC burial, net SO$_4$ reduction (CRS), and Fe reduction].

We calculated the change in surface water CO$_2$ concentration ($\Delta$CO$_2$/CO$_2$) for each process using the above TA and DIC anomalies (X and Y; Table 2) and buffer factors (67) for TA ($\gamma_{\text{TA}}$) and DIC ($\gamma_{\text{DIC}}$): $\Delta$CO$_2$/CO$_2$ = $\gamma_{\text{TA}}$X + $\gamma_{\text{DIC}}$Y. Subsequently, the CO$_2$ concentration (μmol kg$^{-1}$) resulting from this CO$_2$ was then calculated using the “seacarb” package (68) in RStudio. Initial conditions were set according to direct measurements in the water column (TA, 2596 μmol kg$^{-1}$; DIC, 2295 μmol kg$^{-1}$; salinity, 38; temperature, 26°C). The excess CO$_2$ (μmol kg$^{-1}$) shown in Fig. 4 is the difference between initial CO$_2$ and the CO$_2$ concentration after sediment TA/DIC exchange. Water currents at this site are very minor (22), allowing us to exclude lateral exchanges from this simple model. However, an important implication of this model is that any net TA and DIC production/consumption is compensated by import/export with the coastal ocean, albeit over longer time scales.

### Table 2. Selection of generalized biogeochemical reaction pathways affecting TA and DIC.

| Process | Generalized reaction | $\Delta$TA | $\Delta$DIC | Inferred from |
|---------|----------------------|-----------|-----------|--------------|
| Primary production | CO$_2$ + H$_3$PO$_4$ + HNO$_3$ → OM + O$_2$ | 0* | -1 | O$_2$, nutrient flux |
| Sulfate reduction + burial as 50% FeS$_2$ and 50% S$^0$ | 4OM + 2SO$_4^{2-}$ + 2H$^+$ → 4CO$_2$ + 2H$_2$S | +2 | +1 | CRS |
| | 0.5[FeS + H$_2$S → FeS$_2$ + H$_2$] | | | |
| | 0.5[X + 2H$^+$ + H$_2$S → S$^0$ + 2H$_2$O] | $[1 + 0.8\times 2] \over 2$ | $[1 + 0.8\times 2] \over 2$ | |
| Fe reduction | OM + 2FeO$_2$ + 8H$^+$ → 2CO$_2$ + 4Fe$^{2+}$ | +8 | +2 | Fe—CRS |
| Canonical denitrification | OM + 0.8HNO$_3$ → CO$_2$ + 0.4N$_2$ + NH$_3$ + H$_3$PO$_4$ | +0.8 | +1 | N$_2$ flux |
| CaCO$_3$ precipitation | Ca$^{2+}$ + 2HCO$_3^-$ → CaCO$_3$ + CO$_2$ + H$_2$O | -2 | -1 | Range of literature CaCO$_3$ accumulation rates |

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