How organic carbon derived from multiple sources contributes to carbon sequestration processes in a shallow coastal system?

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Abstract
Carbon captured by marine organisms helps sequester atmospheric CO\(_2\), especially in shallow coastal ecosystems, where rates of primary production and burial of organic carbon (OC) from multiple sources are high. However, linkages between the dynamics of OC derived from multiple sources and carbon sequestration are poorly understood. We investigated the origin (terrestrial, phytobenthos derived, and phytoplankton derived) of particulate OC (POC) and dissolved OC (DOC) in the water column and sedimentary OC using elemental, isotopic, and optical signatures in Furen Lagoon, Japan. Based on these data analysis, we explored how OC from multiple sources contributes to sequestration via storage in sediments, water column sequestration, and air–sea CO\(_2\) exchanges, and analyzed how the contributions vary with salinity in a shallow seagrass meadow as well. The relative contribution of terrestrial POC in the water column decreased with increasing salinity, whereas autochthonous POC increased in the salinity range 10–30. Phytoplankton-derived POC dominated the water column POC (65–95%) within this salinity range; however, it was minor in the sediments (3–29%). In contrast, terrestrial and phytobenthos-derived POC were relatively minor contributors in the water column but were major contributors in the sediments (49–78% and 19–36%, respectively), indicating that terrestrial and phytobenthos-derived POC were selectively stored in the sediments. Autochthonous DOC, part of which can contribute to long-term carbon sequestration in the water column, accounted for >25% of the total water column DOC pool in the salinity range 15–30. Autochthonous OC production decreased the concentration of dissolved inorganic carbon in the water column and thereby contributed to atmospheric CO\(_2\) uptake, except in the low-salinity zone. Our results indicate that shallow coastal ecosystems function not only as transition zones between land and ocean but also as carbon sequestration filters. They function at different timescales, depending on the salinity, and OC sources.

Keywords: blue carbon, carbon sequestration, estuary, organic carbon dynamics, phytoplankton, seagrass meadow, sediment, stable isotope

Introduction
The ocean functions as an important carbon sink by absorbing atmospheric CO\(_2\) at a rate (2.3 ± 0.7 Pg-C yr\(^{-1}\)) comparable to the rate of CO\(_2\) uptake by terrestrial ecosystems (2.6 ± 1.2 Pg-C yr\(^{-1}\)) (IPCC, 2013). One of the contributors to the ocean carbon sink is ‘blue carbon’, which is carbon captured by marine organisms (Nellemann et al., 2009; Duarte et al., 2010, 2013; Macreadie et al., 2014). Processes involved in carbon sequestration include carbon storage in sediments, carbon sequestration in the water column, and air–sea CO\(_2\) exchange.

Blue carbon stored in sediment can be sequestered from atmospheric CO\(_2\) for geological timescales (Mateo et al., 1997; McLeod et al., 2011; Fourqurean et al., 2012; Macreadie et al., 2012). The burial rate of organic carbon (OC) is estimated to be higher in shallow coastal ecosystems (238 Tg-C yr\(^{-1}\)) than in the open ocean (6 Tg-C yr\(^{-1}\)) (Nellemann et al., 2009). Furthermore, because shallow coastal ecosystems receive a substantial input of terrestrial carbon (0.9 Pg-C yr\(^{-1}\)), they store allochthonous carbon captured on land (Regnier et al., 2013). The most productive marine ecosystems are shallow coastal ecosystems (Bianchi, 2007) in which abundant OC is produced via active photosynthesis by aquatic primary producers such as microalgae, macroalgae, and seagrass. Part of the produced OC is stored in the sediments. For example, seagrass-derived OC is estimated to contribute about 50% of sedimentary OC (Kennedy et al., 2010). However, the variability of the origin of the OC stored in sediments and the...
mechanisms mediating the composition of stored OC are largely unknown (Macreadie et al., 2012).

Although sequestration of OC in the water column has the potential to sequester atmospheric CO$_2$, a mechanism for sequestering CO$_2$ in the water column of shallow coastal ecosystems has not been identified. In the open ocean, water column sequestration is considered to be an important carbon sink, because refractory dissolved organic carbon (DOC) can be sequestered from atmospheric CO$_2$ for geological timescales (Nagata, 2008; Jiao et al., 2010, 2014). In shallow coastal ecosystems, aquatic primary producers, benefitting from the high nutrient concentrations and sufficient light conditions, generate a substantial amount of autochthonous DOC (Carlson, 2002; Wada et al., 2008; Agustí & Duarte, 2013). Thus, one can expect that refractory OC may be sequestered in the water columns of shallow coastal ecosystems.

Air–sea CO$_2$ exchange is the process that directly determines whether a habitat is a sink or source of atmospheric CO$_2$. Shallow coastal ecosystems are considered to be a net source of atmospheric CO$_2$ due to their high rate of mineralization of terrestrial OC (Borges, 2005; Borges et al., 2005; Cai et al., 2006; Chen & Borges, 2009; Cai, 2011; Regnier et al., 2013). However, recent studies have demonstrated that shallow coastal seagrass meadows in estuaries can be sinks for atmospheric CO$_2$ (Maher & Eyre, 2012; Tokoro et al., 2014). Whether the net ecosystem production (NEP) of seagrass meadows is positive or negative is a key determinant of whether they are sinks or sources, respectively, of atmospheric CO$_2$ (Maher & Eyre, 2012; Tokoro et al., 2014), but the linkages between the gas exchange process and OC dynamics such as inflow/outflow and production/consumption remain unclear.

There is increasing interest in understanding the role of OC derived from multiple sources in coastal carbon dynamics in response to global change (Bauer et al., 2013). Unlike sequestration in terrestrial ecosystems, water column OC dynamics intervene and mediate the above-mentioned key carbon sequestration processes in shallow coastal ecosystems. Furthermore, assessment of temporal variations of the carbon sequestration processes was impossible from this single survey. However, our goal was to understand processes by relating the spatial pattern to salinity.

At each station, the salinity of the water was recorded by using a conductivity–temperature sensor (COMPACT-CT, JFE Advantech, Nishinomiya, Japan). Samples for DIC and the fugacity of CO$_2$ (fCO$_2$) were dispensed into 250-mL Schott Duran bottles, which were poisoned with mercuric chloride (200 µL per bottle) to prevent DIC changed due to biological activity. Water samples for the other analyses were collected
in acid-washed polyethylene bottles. Samples for nutrients (NH$_4^+$, NO$_2^-$, and NO$_3^-$), total dissolved nitrogen (TDN), DOC, and optical analyses of chromophoric dissolved organic matter (CDOM) were filtered through 0.2-$\mu$m polytetrafluoroethylene filters (DISMIC–25HP; Advantec, Durham, NC, USA) into precombusted (450 °C for 2 h) 50- or 100-ml glass vials and frozen at −20 °C until analysis. Samples for chlorophyll a (chl a) concentrations and analyses of POC, particulate nitrogen (PN), and stable isotope analysis ($\delta^{13}$C$_{POC}$ and $\delta^{15}$N$_{PN}$) were obtained by filtration (~1 l) onto precombusted (450 °C for 2 h) glass-fiber filters (GF/F, Whatman, Maidstone, Kent, UK) and stored in the dark at −20 °C until analyses.

Sediment cores were collected in October 2012 at stations L7, L8, and L17 and in June 2013 at station C1 along a salinity gradient. The surface sediments (depth: 0–20 mm) of triplicate sediment cores (8.6-cm inner diameter) were used for analysis of total organic carbon (TOC) and total nitrogen (TN) concentrations and stable isotope signatures ($\delta^{13}$C and $\delta^{15}$N).

To evaluate the contribution of phytobenthos to particulate organic matter (POM) in the water column and sedimentary OM, the stable isotope signatures and C/N ratio of the phytobenthos were measured. The dominant seagrass (Z. marina) and periphyton of the seagrass were collected in August 2012 at stations L6, L8, L13, and L17. Samples were kept at −20 °C prior to analysis.

**Analytical methods**

DIC concentrations and fCO$_2$ values were determined on a batch-sample analyzer (ATT-05; Kimoto Electric, Osaka, Japan). DOC was determined via high-temperature catalytic oxidation with a TOC analyzer (TOC5000A; Shimadzu, Kyoto, Japan). Nutrient concentrations in sample filtrates were measured using an AutoAnalyzer (TRAACS 800; Bran+Luebbe, Norderstedt, Germany). TDN was measured via wet oxidation with persulfate (SWAAT; BL-Tec, Osaka, Japan). Concentrations of dissolved organic nitrogen (DON) were calculated as the differences between TDN and total DIN (the sum of NH$_4^+$, NO$_2^-$, NO$_3^-$).

For the chl a analyses, the GF/F filters were extracted in the dark for 12 h in 90% acetone, and chl a concentrations were measured by using a UV-visible spectrometer (UV-2450; Shimadzu, Kyoto, Japan) (Lorenzen, 1967).

CDOM absorbance spectra were recorded from 250 to 700 nm at 1-nm increments using a UV-visible spectrometer (UV-2450; Shimadzu, Kyoto, Japan) fitted with a 1-cm quartz flow cell and referenced to ultrapure water (Milli-Q water; Millipore, Billerica, MA, USA). The absorbance values at each wavelength ($\lambda$) were transformed into absorption coefficients ($a_{CDOM}$) using:

$$a_{CDOM}(\lambda) = \frac{2.303 \times A_{CDOM}(\lambda)}{m}$$

where $A_{CDOM}$ is the absorbance value per meter. The absorption value at 375 nm, $a_{CDOM}(375)$, was chosen to quantify CDOM because this wavelength has been commonly used as a metric of DOC absorbance in previous studies (e.g., Bricaud et al., 1981; Astoreca et al., 2009; Para et al., 2013). We also measured $a_{CDOM}(254)$ as a metric of the aromaticity of the DOM (e.g., Weishaar et al., 2003; Zurbrügg et al., 2013). We calculated specific UV absorption at 254 nm (SUVA254) using:
SUVA254 = A_{CDOM(254)}/[DOC](1 \text{ mg}^{-1}\text{m}^{-1})

where (DOC) is a DOC concentration. Spectral slopes for the interval of 275-295 nm ($\lambda_{275-295}$) were calculated using linear regression of the log-transformed $a_{CDOM}$ spectra. Slopes are reported as positive numbers to follow the mathematical convention of fitting to an exponential decay. Thus, higher (or steeper) slopes indicate a more rapid decrease in absorption with increasing wavelength (Helms et al., 2008).

Samples for analyses of TOC and TN content and stable isotope signatures ($\delta^{13}C$ and $\delta^{15}N$) were dried in an oven at 60 °C or with a freeze dryer. To remove inorganic carbon, we acidified the samples with 1N HCl and dried them again at 60 °C. TOC and TN concentrations and stable isotope signatures ($\delta^{13}C$ and $\delta^{15}N$) were measured with an isotope-ratio mass spectrometer (Delta Plus Advantage, Thermo Electron, Bremen, Germany) coupled with an elemental analyzer (Flash EA 1112; Thermo Electron, Bremen, Germany). Stable isotope ratios are expressed in $\delta$ notation as the deviation from standards in parts per thousand ($\%$o) according to the following equation:

$$\delta^{13}C, \delta^{15}N = [R_{\text{sample}}/R_{\text{standard}} - 1] \times 10^3$$

where $R$ is $^{13}C/^{12}C$ or $^{15}N/^{14}N$. PeeDee Belemnite and atmospheric nitrogen were used as the isotope standards of carbon and nitrogen, respectively. The analytical precision of the Delta Plus Advantage mass spectrometer system, based on the standard deviation of the internal reference replicates, was <0.2‰ for both $\delta^{13}C$ and $\delta^{15}N$.

Data analysis

To estimate biological and/or physicochemical reduction or addition of carbon in the water column, we calculated the difference between the observed concentration of an element ($X$) and its concentration predicted by conservative mixing ($X_{\text{mix}}$) as $\Delta X$ at each lagoon station: $\Delta X = X - X_{\text{mix}}$. The $\Delta X$ concentrations were determined for DIC, DOC, POC, and total carbon (TC = DIC + DOC + POC). Predicted conservative concentrations ($X_{\text{mix}}$) were estimated using a linear salinity mixing model with two end members. The terrestrial end member was defined as the average concentration in the three rivers weighted by the discharge of the rivers (Furen River: 21.0 m$^3$ s$^{-1}$; Yausubetsu River: 21.0 m$^3$ s$^{-1}$; Pon-Yausubetsu River: 0.7 m$^3$ s$^{-1}$). The concentrations in the bottom water at the lagoon mouth, station L18 with the highest salinity, were used as the marine end member.

The Bayesian isotopic modeling package, Stable Isotope Analysis in R (SIAR) (Parnell et al., 2010), was used to partition the proportional contributions of potential OM sources to the bulk POM and sedimentary OM based on their N/C, $\delta^{13}C$, and $\delta^{15}N$ signatures. We chose N/C rather than C/N ratios in the model because the former were statistically more robust; the higher number (TOC concentration) is the denominator and behaves linearly in end-member mixtures (Goñi et al., 2003; Perdue & Koprivnjak, 2007). The SIAR model works by determining probability distributions of the sources that contribute to the observed mixed signal while accounting for the uncertainty in the signatures of the sources and isotopic fractionation. We assumed an isotopic fractionation of 0 and ran the model through 1 $\times$ 10$^6$ iterations. For each potential source, we report the median and the 95% confidence interval (CI) of the estimate of the proportional contribution of each source to the observed value.

We defined four sources (terrestrial OM, coastal OM, lagoon OM, and phytobenthos-derived OM) as end members for the isotopic and elemental mixing model. To estimate the OM components in sediments, we pooled coastal OM and lagoon OM as phytoplankton-derived OM (PhytoOM). The signature values of terrestrial OM (TerrOM) were determined as the average values of samples taken at the three river mouths (salinity = 0). The values in the bottom layer at the lagoon mouth station (L18) with the highest salinity were used as the signature values of coastal OM (CoastOM). Lagoon OM (LagOM) was defined as autochthonous phytoplankton growing in the brackish area of the lagoon. Because the C/N ratio of fresh phytoplankton is close to 7.0 on a molar basis (Redfield et al., 1963) and a low POC/chl $a$ ratio (<100 on a mass basis) is characteristic of living microalgae (Zeitzschel, 1970; Maksymovska et al., 2000), LagOM from samples collected in the brackish area (salinity range: 5–25), which was characterized by a low POC/chl $a$ ratio (<50) and low C/N ratio (<7.0), was considered to be representative of phytoplankton. Phytobenthos-derived OM (BenthOM) was defined as a mixture of the dominant seagrass (Z. marina) and the epiphyte. The signature values for BenthOM were calculated using a linear interpolation between the values characteristic of seagrass (seagrass; 100%) and of epiphytes (seagrass; 0%). The total range of the calculated signature values revealed that the mixing ratio had only about a 5% effect on the estimation. We therefore determined the BenthOM to be a mixture of 50% seagrass and 50% epiphyte. Because the N/C ratio generally declines while OM is decomposing (Van Mooy et al., 2002), it should be noted that the contribution of OM with a high N/C ratio (i.e., LagOM and CoastOM; Table S1) to POM and sedimentary OM was probably underestimated. In contrast, isotopic fractionation does not occur during the decomposition of organic compounds with a high molecular weight (Fry, 2006).

Results

Characteristics of organic matter in the water column

Concentrations of POM (POC, PN) were higher than predicted by conservative mixing, with the highest values around a salinity of 20 in the lagoon (Fig. 2a, b). The distribution of POM concentrations corresponded approximately to the distribution of chl $a$ concentration (Fig. 2f). The POC/PN ratio was lower than predicted by the conservative mixing, and there were several high values in the river mouths (Fig. 2c). $\delta^{13}C_{\text{POC}}$ plotted along the predicted conservative mixing line, but the $\delta^{15}N_{\text{PN}}$ values were above the predicted line at many stations (Fig. 2d, e).
Dissolved OC and DON concentrations were higher than predicted by the conservative mixing, and the differences were large in the salinity range 5–25 (Fig. 3a, b). The DOC/DON ratios were lower than predicted by the conservative mixing (Fig. 3c). Although \( \delta^{13}C_{POC} \) values plotted along the conservative mixing line, \( \delta^{15}N_{PN} \) values were higher than predicted by the conservative mixing (Fig. 3d, e). SUVA254 values plotted along the conservative mixing line (Fig. 3f). \( S_{275-295} \) values were higher than predicted by the conservative mixing, and the highest values occurred at high salinity (Fig. 3g).

**Composition of particulate organic matter**

The isotopic and elemental mixing model (Fig. 4a, b and Table S1) showed that each OM source made a different contribution to water column POM along the salinity gradient in the Furen Lagoon. Dashed lines indicate the predicted conservative mixing relationships.

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salinity gradient (Fig. 5a–d). The relative contribution of TerrOM to POM was high (median value: 89%) at the river mouths and low (<10%) around the lagoon mouth (Fig. 5a). LagOM was the largest source of POM in the lagoon (~83%) in the salinity range 15–25 (Fig. 5b). The relative contribution of CoastOM to POM was high (~92%) around the lagoon mouth and low (<5%) in low-salinity areas (Fig. 5c). The relative contribution of BenthOM was low (~19%) in the lagoon, with the highest contribution (~20%) around a salinity of 20 (Fig. 5d).

We calculated the concentrations of water column POC derived from each source along the salinity gradient (Fig. 6a–d) by multiplying the relative contribution of each source (Fig. 5a–d) by the bulk POC concentration (Fig. 2a). The terrestrial POC (TerrPOC) concentration was high (median value: 39.1 μmol-C l⁻¹) at the river mouths and low (0.2 μmol-C l⁻¹) around the lagoon mouth (Fig. 6a). The lagoon POC (LagPOC) concentration was high in the salinity range 15–25, and the maximum median concentration was 69.6 μmol-C l⁻¹ at station L5 (Fig. 6b). The ranges of salinity associated with the highest concentrations were similar for chl a and LagPOC (Figs 2f and 6b). Coastal POC (CoastPOC) and Phytobenthos-derived POC (BenthPOC) concentrations were relatively low in the lagoon (Fig. 6c, d). The maximum median concentration of CoastPOC was 18.5 μmol-C l⁻¹ at station L18. The BenthPOC concentration was the highest (median value: 17.3 μmol-C l⁻¹) at station L5.

Composition of sedimentary organic matter

The isotopic and elemental mixing model (Fig. 4a, b and Table S1) showed that each OM source made a
different contribution to sedimentary OM along the salinity gradient (Fig. 7a–c). TerrOM made the greatest contribution to OM (49–78%) in the lagoon sediments but contributed only 10% to the sedimentary OM at the coastal station (Fig. 7a). The relative contribution of TerrOM was high near the river mouths, where the annual mean salinity at station L7, for example, was 15. PhytoOM was a minor component (3–29%) of the OM in the lagoon sediments (Fig. 7b). The relative contributions of PhytoOM were high at the lagoon mouth (station L17) and the coastal station. BenthOM was the second largest OM pool in the lagoon sediments (19–36%; Fig. 7c). The relative contribution of BenthOM was lowest (17%) at the coastal station and highest in low-salinity areas, where the annual mean salinity at station L8, for example, was 13.

Carbon budget in the water column

ΔTC, ΔDIC, ΔDOC, and ΔPOC values were positive at salinities <15 (Fig. 8a, b). At salinities >15, a decrease in DIC corresponded to an increase of OC (e.g., ΔTC near 0; Fig. 8a, b). The surface-water fCO2 values were high (~7332 μatm) at salinities <15 and were close to atmospheric values at salinities >20 (Fig. 8c). Surface-water fCO2 values at several stations where salinity was >15 were actually lower than the atmosphere value of 397 μatm (Fig. 8c).

Discussion

Selective carbon storage in sediments

To our best knowledge, this study first revealed that the efficiency of OC storage in sediments is dependent on OC derived from multiple sources (Fig. 9c, d). The fact that terrestrial OC and phytobenthos-derived OC were more efficiently stored than phytoplankton-derived OC in seagrass meadow sediments suggests that the variability of the composition of OC affects the carbon burial rate.

PhytoOM, derived mainly from autochthonous production, was dominant in the water column POM pool (~95%; Figs 5b, c and 9b, c), as has been found in other shallow coastal systems worldwide (Sato et al., 2006; Harmelin-Vivien et al., 2010; Dubois et al., 2012; Savoye et al., 2012; Guerra et al., 2013). Because the composition of POM in the water column fluctuates seasonally, our estimation from a single survey is subject to considerable uncertainty. However, we believe that PhytoOM is the primary component of the water column POM throughout the year, because the chl a concentration during this study (mean value: 10 μg L⁻¹; Fig. 2f) was lower than the annual mean value (14 μg L⁻¹; Montani et al., 2011). In addition, the POC/PN ratio in this study (mean value: 7.7 mol mol⁻¹; Fig. 2c) was lower than TerrOM (12.5 mol mol⁻¹; Table S1) and similar to the
annual mean value (7.7 mol mol\(^{-1}\); S. Montani, S. Shibano\-nana and Y. Tsuji, unpublished data). Thus, our assessment that PhytoOM accounted for ~95% of the POM in the water column is basically conservative. Although differences in sinking rates of POM sources affect carbon accumulation rates in sediments, such differences may be of minor importance in very shallow systems like Furen Lagoon. We thus hypothesize that the carbon accumulation rate reflects the composition (concentration) of water column POM and that PhytoOM accounts for the largest flux of POM into the sediments. However, PhytoOM was a relatively minor component (3–29%), and TerrOM and BenthOM were major components (49–78% and 19–36%, respectively) of the sedimentary OM (Figs 7a–c and 9d), that is, TerrOM and BenthOM were more efficiently preserved than PhytoOM in the sediments. The ratios of TerrOM to PhytoOM in the sediments were at least 10 times the analogous ratios in the water column of the lagoon (Fig. 9c, d). Because TerrOM is derived from terrestrial plants with hard cell walls, it contains refractory OM such as aliphatic (cutan, cutin, suberan, and suberin) and aromatic ( lignin, sporopollenin, and dinosporin) cell wall biopolymers (Zonneveld et al., 2010). In addition, the terrestrial OC transported to the ocean has been reported to be relatively refractory and selectively preserved in marine sediments, because much terrestrial OC is highly degraded during the process of being transported through soils (Hedges et al., 1994; Zonneveld et al., 2010).

The contributions of BenthOM in the lagoon sediments were 0.8–6 times the contributions of PhytoOM (Fig. 9d). The efficient storage of phytobenthos-derived OM may be partly explained by the fact that seagrass...
detritus contains a high proportion of refractory OM such as a lignin, especially detritus derived from rhizomes, and roots (Kennedy & Björk, 2009; Kennedy et al., 2010). Like other OM, the flux of BenthOM from the water column to the surface sediment is accounted for by fine particles (<1 mm) derived from plant thalli, because these fine particles are readily resuspended in the water column at our site due to the very shallow, windy conditions. However, it is noteworthy that BenthOM is also supplied from belowground biomass;
thus, the actual total flux of BenthOM to the sediments may have been underestimated.

**Carbon sequestration in the water column**

The dynamics of DOC tend to be overlooked, but they can play a key role in the sequestration of carbon in the water columns of shallow coastal ecosystems, because DOC is generally the dominant fraction of the water column OC pool and includes a large proportion of refractory OC (Nagata, 2008; Jiao et al., 2014). However, the production rates and distribution of refractory DOC in shallow coastal ecosystems are largely unknown (e.g., Wada et al., 2008; Lønborg & Søndergaard, 2009). The release of DOC from seagrass meadows have been reported (Opsahl & Benner, 1993; Barrón & Duarte, 2009); however, using optical analysis, we first showed the possibility of DOC released from seagrass is indeed remained within the whole system.

The production of autochthonous DOC and the dilution of terrestrial DOC regulate the dynamics of DOC in the water column (Figs 3a and 9a). To date, \( a_{\text{CDOM}}^{(254)} \) values have been used as a metric of the concentration of terrestrial aromatic compounds (Weishaar et al., 2003; Zurbrüggh et al., 2013) and as a proxy for potential refractory OM (Saadi et al., 2006; Hur et al., 2009). This study revealed that aromatic compounds have an autochthonous origin at our site (Fig. 3e), that is, refractory compounds are produced in situ in shallow coastal ecosystems. In a mangrove-dominated estuary located within the Everglades, aromatic DOM is exuded from the mangroves (Bergamaschi et al., 2012). In seagrass meadows, the potential sources of aromatic DOM would be microalgae and seagrasses. Phytoplankton release CDOM with a protein-like fluorescence (Romera-Castillo et al., 2010) that is related to freshly produced aromatic amino acids, such as tyrosine, tryptophan, and phenylalanine (Yamashita et al., 2008). The decrease of DOC/DON ratios and the increase of \( S_{275–295} \) values in the lagoon (Fig. 3c, g) support the hypothesis that phytoplankton release labile, protein-rich DOM (Biddanda & Benner, 1997; Fichot & Benner, 2012). In contrast, aquatic vascular plants such as seagrasses contain lignin, which is a refractory biopolymer. The subtropical seagrass *Halodule wrightii* exudes dissolved, lignin-derived phenols as it decomposes (Opsahl & Benner, 1993). Barrón & Duarte (2009) reported that meadows of the Mediterranean seagrass *Posidonia oceanica* release DOC that represents ~71% of the net community production. These findings suggest that seagrasses directly produce refractory DOM that may be preserved in the water column.

The linear decrease with salinity of \( a_{\text{CDOM}}(375) \), which is also a proxy of potentially terrestrial OM (Astoreca et al., 2009; Para et al., 2013), suggests that conservative mixing of terrestrial DOC with seawater accounted for much of the variability of DOC concentrations in Furen Lagoon (Fig. 9a), as has been observed in other estuaries (Chen et al., 2007; Astoreca et al., 2009).

**Air–sea CO\(_2\) exchange**

The uptake of DIC in the surface-water column by aquatic primary producers stimulates an influx of atmospheric CO\(_2\), although mixing of high-fCO\(_2\) water from land runoff with low-fCO\(_2\) ocean water is the major process that mediates air–sea CO\(_2\) exchange in shallow coastal ecosystems. The low ΔTC values in the high-salinity zone of Furen Lagoon suggest that the benthic release/adsorption of carbon and biological metabolism (photosynthesis and mineralization) were more or less in balance in the water column. Relevant to this point is the fact that OC increases coincided with DIC decreases (Fig. 8a, b), indicating that the aquatic primary producers were converting DIC to OC and thereby decreasing the fCO\(_2\) of the water column (Figs 8c and 9d). Biological metabolism determines whether a body of water is a sink or a source of atmospheric CO\(_2\) (Maher & Eyre, 2012; Tokoro et al., 2014). Our results show that OC sequestration by aquatic primary producers was directly linked to atmospheric CO\(_2\) uptake in the high-salinity zone of Furen Lagoon, as expected by previous studies (Maher & Eyre, 2012; Tokoro et al., 2014).

In contrast, the positive values of ΔTC in the low-salinity zone suggest that DIC and/or OC were being released from the benthic system (Fig. 8a, b). This fact, combined with the general high-fCO\(_2\) of inflowing freshwater (Chen et al., 2012), caused the low-salinity zone to be a source of atmospheric CO\(_2\) (Figs 8c and 9d).

How does organic carbon derived from multiple sources contribute to carbon sequestration?

Shallow coastal ecosystems function not only as transition zones between the land and ocean but also as carbon sequestration filters that mitigate atmospheric CO\(_2\) increases, where planktonic and benthic primary producers sequester OC from atmospheric CO\(_2\) and thereby mitigate atmospheric CO\(_2\) increases (Fig. 9d). We found that (i) phytobenthos-derives and terrestrial OC are stored more efficiently than phytoplankton-derives OC in sediments, (ii) DOC production by aquatic primary producers sequesters carbon in the water column, and (iii) OC sequestration in the water column contributes to the influx of atmospheric CO\(_2\) at biological timescales. These three findings are dependent on the salinity gradient. Our findings reveal that the
dynamics of OC derived from multiple sources link with carbon sequestration processes at multiple timescales. This discovery suggests that criteria for evaluating the effectiveness of processes for sequestering carbon should be related to the salinity and OC sources. Finally, our methodology inferring the multiple sources and rates of carbon and the results provide a step toward better understanding coastal carbon dynamics in response to global change, in particular, continued human pressures (Bauer et al., 2013).

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Supporting Information

Additional Supporting Information may be found in the online version of this article:

Table S1. Isotopic and elemental signatures of OM sources.