Sources of Particulate Organic Matter across Mangrove Forests and Adjacent Ecosystems in Different Geomorphic Settings

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Abstract
Mangrove forests are among the world’s most productive ecosystems and provide essential ecosystem services such as global climate regulation through the sequestration of carbon. A detailed understanding of the influence of drivers of ecosystem connectivity (in terms of exchange of suspended particulate organic matter), such as geomorphic setting and carbon stocks, among coastal ecosystems is important for being able to depict carbon dynamics. Here, we compared carbon stocks, CO₂ fluxes at the sediment-air interface, concentrations of dissolved organic carbon and suspended particulate organic carbon across a mangrove-seagrass-tidal flat seascape. Using stable isotope signatures of carbon and nitrogen in combination with MixSIAR models, we evaluated the contribution of organic matter from different sources among the different seascape components. Generally, carbon concentration was higher as dissolved organic carbon than as suspended particulate matter. Geomorphic settings of the different locations reflected the contributions to particulate organic matter of the primary producers. For example, the biggest contributors in the riverine location were mangrove trees and terrestrial plants, while in fringing locations oceanic and macroalgal sources dominated. Anthropogenic induced changes at the coastal level (i.e. reduction of mangrove forests area) may affect carbon accumulation dynamics in adjacent coastal ecosystems.

Keywords Connectivity · Blue carbon · Carbon flux · Seagrass beds · Tidal flats · Suspended particulate matter

Introduction
Mangrove forests play an important role of the tropical seascape, as well as for blue carbon accumulation, as they are sites of storage and exchange of carbon (Dittmar et al. 2012; Duarte et al. 2013; Kristensen et al. 2008). Mangrove forests store carbon within their aboveground (AGB) and belowground living biomass (BGB), non-living biomass (e.g. dead wood) and sediments (Alongi 2014). Together with saltmarshes and seagrasses, mangrove forests are known as a “blue carbon” ecosystem, referring to the high rates at which these coastal ecosystems sequester and store carbon (Lovelock and Duarte 2019).

Multiple studies have quantified mangrove carbon stocks (Bhomia et al. 2016; Kauffmann et al. 2011; Murdiyarso et al. 2015), suggesting an important role in climate regulation. Moving beyond stock assessments, studies have now started to show how carbon stocks may differ with climate (Simard et al. 2019) or geomorphic setting (Rovai et al. 2018) at regional to global scales. On a local scale, physical and bio-geographic factors (i.e. climate regime, tidal regime, geomorphology, hydrodynamics, and nutrient dynamics) influence the carbon dynamics in mangrove forests (Alongi 2014).
Understanding the factors that influence carbon storage and estimating carbon emissions and exchange at the seascape level across physical and ecological gradients is important to understand blue carbon dynamics.

Landscape structure includes the spatial arrangement and composition of landscape elements (corridors, patches and matrix) and ecological fluxes (e.g., movements of water, nutrients, organisms and other materials), and is an important landscape ecology research topic (Wu 2012). The fluxes and exchange of organic carbon among ecosystems are particularly important in networks of coastal ecosystems that release and trap carbon and nutrient subsidies from adjacent ecosystems (Gillis et al. 2014a, 2017). Carbon stored in coastal tropical vegetated ecosystems can either be produced autochthonously, or imported from adjacent ecosystems (Adame and Lovelock 2011; Hyndes et al. 2014; Duarte et al. 2013). Carbon flow among ecosystems can occur actively by motile consumers (Bouillon and Connolly 2009) or passively by tides, coastal currents and/or river discharge transporting suspended detrital particulate carbon and dissolved organic and inorganic carbon (Hyndes et al. 2014; Bouillon et al. 2008; Gillis et al. 2014b). Several studies suggest that connectivity is important to the understanding of the ecosystem processes that underlie blue carbon sequestration (Gillis et al. 2014a; Olds et al. 2016; Walton et al. 2014; Twilley et al. 1992). However, only few studies have directly measured carbon stocks (Phang et al. 2015) or nutrient fluxes (Huxham et al. 2018) across adjacent ecosystems such as tidal flats and seagrass beds.

The aim of this study was to evaluate the relative contribution of mangrove trees, seagrass plants, macroalgae, terrestrial plants and oceanic sources (plankton and suspended particulate matter (SPM)), to the particulate organic matter (POM) pool in the coastal water body. We define SPM as all suspended matter including inorganic and organic matter, while POM only includes the organic pool of the suspended matter. In order to explore carbon stocks, CO$_2$ and carbon exchange and the contributions of the main primary producers to POM, we chose three mangrove-seagrass/tidal flat ecotones along the coast of Singapore. Our research questions were: (1) what are the carbon stocks in the mangrove tree biomass, seagrass plants and the upper 15 cm of sediment, and CO$_2$ fluxes from the sediment to the atmosphere of three mangrove forests under similar climatic conditions but different geomorphic settings (riverine vs. fringing, according to (Ewel et al. 1998)) and (2) what is the role of mangrove forests and adjacent ecosystems as donors and recipients of POM across different geomorphic settings? We hypothesized that biomass, CO$_2$ fluxes and connectivity (through POM exchange) with adjacent ecosystems is influenced by landscape structure (i.e. mangrove forest or catchment area spatial arrangement).

Materials and Methods

Sampling Design

Three intertidal continuums in northern Singapore were selected: Sungei Buloh (1°26′47″N, 103°43′25″E), Seletar Island (1°26′36″N, 103°51′43″E) and Chek Jawa (1°24′39″N, 103°59′26″E). Those sites included two mangrove forests-tidal flat (Sungei Buloh and Seletar Island) and one mangrove forests-seagrass bed ecotones (Chek Jawa). In each location, mangrove forests were present alongside other intertidal ecosystems such as tidal flats or seagrass beds (Fig. 1). Sampling was conducted during the late Northeast Monsoon (Dry Phase) from February to March 2017. Daily averages of temperature were between 23 and 32 °C and total daily rainfall between 0 and 31 mm (Meteorological service Singapore, http://www.weather.gov.sg).

 Transects were set across the intertidal zone, starting at the tidal flat or seagrass bed and ending at the mangrove forest-terrestrial ecotone, extending landward. Within each location we established three transects separated by a distance of 50 to 500 m, depending on alongshore mangrove forest extent. The distance between points along each transect was determined by the total cross-shore length along the intertidal gradient of the mangrove forest, but sampling points were generally separated by 20 to 120 m, to gain a representation of the entire forest width.

Sungei Buloh and Chek Jawa had between three to five points on each transect, whilst Seletar Island had two points on each transect; terrestrial sampling points were not possible here due to impenetrable forest cover. Terrestrial points at Sungei Buloh did not follow a line but were taken at the closest point possible to the last mangrove-terrestrial forest ecotone. Terrestrial sampling was not possible at Seletar due to impenetrable forest cover. Transect points inside the mangrove that were closest to the ocean are defined as mangrove forest 1 (MF1), and the three points furthest from the ocean as mangrove forest 3 (MF3). For seagrass beds, the three transect points furthest from the mangrove forest are defined as seagrass bed 1 (SB1), and the three closest to the mangrove forest as seagrass bed 3 (SB3). The tidal flat transect point was classified with (TF), and the terrestrial area with (TE).

Biomass Carbon Estimation

At each transect point inside the mangrove forest, a plot of 100 m$^2$ was established, all mangrove species were identified, and the total number of trees of each species present was recorded. The diameter at breast height (DBH) at 1.3 m from the forest floor, was measured for each tree, except for species of the genus Rhizophora sp. where the diameter at 30 cm above the highest root of the main stem was taken (Kauffman and Donato 2012). From
this, we estimated aboveground and belowground biomass (AGB and BGB respectively), using species-and region-specific allometric equations (Table 1). If no allometric equation had been developed for the species, a genus-level or general equation was used, as commonly conducted in other studies (e.g., Rahman et al. 2015). Species-specific wood densities were used, which is recommended if stand-specific measurements are not available (Chave et al. 2005). Carbon pools were derived from the living biomass measurements using a standard and conservative biomass-to-carbon ratio of 0.464 (Kauffman and Donato 2012).

At each transect point inside the seagrass bed a core of 15 cm diameter, total area of 176 cm$^2$ and 20 cm depth was used to harvest seagrass ABG and BGB. Seagrass parts were cleaned and rinsed in the field with seawater, and again in the laboratory with distilled water. The amount of organic carbon in seagrass structures was calculated by multiplying % carbon content of each species by the biomass present at each point (Howard et al. 2014). Carbon content in the plant biomass and carbon stocks in the ecosystems were calculated as explained below. Carbon stocks were reported in Mg (megagrams) of carbon per hectare, however other units were reported in mg (milligrams) due to the quantities differing by at least ten orders of magnitude.

**Sediment Sampling**

At each point along the transects, two types of suspended sediment samples were evaluated, one from sediment traps on the sediment surface, and a second one from the water column (suspended particulate matter (SPM)).

Sediment traps were installed during low tide at a height of 0.05 m above the sediment. The traps were plastic cylinders of 24 cm length and 6 cm diameter, with twenty-four 0.5 cm-diameter holes evenly distributed in the upper 10 cm, with a lid at the bottom that collected the suspended particulate matter that entered the trap. Sediment traps were attached to steel rods that were anchored in the sediment. The traps were emptied after 24 h. To avoid salt contamination, samples were exposed in the oven at 60 °C for 24 h, supernatant water was carefully removed and an additional 150 mL of distilled water was added to dissolve remaining salt. This process was repeated until salt particles were not detected visually.

For SPM, water samples of 1 L from the upper 40 cm of the water column were taken at each point during ebb tide. Admittedly, we acknowledge that SPM being sampled at ebb tide may be biased towards mangroves and terrestrial sources, whereas sediment traps captured POM over two ebb and two flood tides. Samples were kept in freezer bags with freezer blocks and transported within 4 h to the

| No | Species               | Aboveground     | Belowground    | Reference                           |
|----|-----------------------|-----------------|----------------|-------------------------------------|
| 1  | *Avicennia alba*      | 0.251pDBH2.46   | 0.199p0.899DBH2.22 | Komiyama et al. 2005               |
| 2  | *Avicennia officinalis*| 0.251pDBH2.46   | 0.199p0.899DBH2.22 | Komiyama et al. 2005               |
| 3  | *Avicennia rumphiana*  | 0.251pDBH2.46   | 0.199p0.899DBH2.22 | Komiyama et al. 2005               |
| 4  | *Bruguiera cylindrica* | 0.251pDBH2.46   | 0.199p0.899DBH2.22 | Komiyama et al. 2005               |
| 5  | *Rhizophora apiculata*| 0.1709pDBH2.516 | 0.199p0.899DBH2.22 | Putz and Chan 1986; Komiyama et al. 2008 |
| 6  | *Rhizophora stylosa*  | 0.1709pDBH2.516 | 0.199p0.899DBH2.22 | Putz and Chan 1986; Komiyama et al. 2008 |
| 7  | *Sonneratia alba*     | 0.251pDBH2.46   | 0.199p0.899DBH2.22 | Komiyama et al. 2005               |
laboratory, where they were filtered onto pre-combusted (450 °C, 24 h) glass fiber filters (GF/C, 696 grade, 1.2 μm pore diameter). Suspended particulate material, dry filters and sediment trap samples were analyzed for isotopic composition and used later in the mixing model to determine the origin of the POM.

At all transect points, a sediment core of 15 cm depth and 7 cm diameter was taken during low tide. The core was divided into three subsamples of 0–5, 5–10 and 10–15 cm sediment layers. The amount of organic carbon in each layer was calculated by measuring the % OC in the sample and multiplying by sediment density (Howard et al. 2014).

Water and Gas Flux Sampling

Fluxes of CO2 were measured using a portable sampling CO2 data logger (K33-BLG CO2Meter), with an internal CO2 sensor using NDIR (non-dispersive infrared), with an accuracy of ±0.2% volume CO2. The loggers were calibrated using outdoor air as a reference, having 400 ppm as reference value, and the software DAS gas lab®. The sensor was configured with a 15 cm diameter light and dark (covered with aluminum foil) 9 L survey chamber. Light and dark chambers were used to evaluate the influence that photosynthetic microorganism could have on CO2 fluxes. CO2 flux density (mg CO2 ·C m−2 h−1) (F CO2) was calculated following the methods and equation described by Chojnicki et al. (2009) using light and dark chambers.

\[
F_{CO2} = k_{CO2} \cdot (V \cdot A_{-1}) \cdot (dc \cdot dt_{-1})
\]

Where \( k_{CO2} \) is the gas-constant at 273.15 K = 0.536 (μg C μl−1), \( Tair \) represents the air temperature inside the chamber (K), \( V \) is the chamber volume (L), \( A \) is the collar area (m²) and \( dc/dt_{-1} \) is the rate of CO2 concentration change in chamber (ml 1−1 h−1).

At each plot, 20 mL of water was taken from the water column during the ebb tide and filtered (45 μm pore size) into pre-combusted glass vials. Samples were kept in a cooler bag with external freezer blocks and transported to the laboratory. For DOC samples, approximately 200 μL of HCl was added to decrease pH below 2, and preserved samples were transported to ZMT. Dissolved organic carbon was analyzed using an infra-red gas analyzer with a Skalar SAN System.

Samples for Chlorophyll a (Chl-a) were obtained by filtering 500 mL through GF/F filters (1.2 μm). After filtering, samples were stored frozen until measurements were done. All samples were transported in cooler bags with freezer blocks, before being analyzed at ZMT chemical analytical laboratories. For Chl-a, each filter was cut in pieces and left in constant agitation overnight in 8 mL of 96% ethanol in the dark at 20 °C. Later, samples were centrifuged for 20 min at 5000 rpm at 4 °C. Absorbance was measured at 665 nm and 470 nm using a photometer (Shimadzu UV-1700). Calculations of Chl-a were done following the procedure described by Ritchie (2008).

Carbon Source Analysis

Five sources of POM were considered in this study: oceanic (plankton and SPM), mangrove tree leaves, seagrass leaves, terrestrial plants leaves and macroalgal tissues. Stable isotope signatures, \( \delta^{13}C \) and \( \delta^{15}N \), of the five different POM sources, analyzed through the Bayesian mixing model MixSIAR (Stock et al. 2018), were used for estimating the contribution of the different sources to the POM sampled in each plot. Isotopic signal values for oceanic sources used in this study were taken from studies done in Johor strait (Zhang et al. 2017), in this instance oceanic samples were mostly comprised of plankton and suspended sediment matter. The most abundant species at each mangrove location (Rizophora apiculata, Avicennia alba, Bruguiera cylindrica, Avicennia rumphiana, Nypo fruticans (dominant only in Chek Jawa)), seagrass (Cymodocea rotundata, Halodule uninervis, Halophila ovalis), terrestrial plants (Caryota mitis, Canavalia cathartica, Barringtonia sp., Thespesia populnea, Pinus sp. (dominant only in Seletar Island)) and macroalgae (Ulva sp. and Dictyota sp.) were sampled. Fresh plant leaves (4–5) and algal thalli were taken, placed in separate sample bags and transported to the laboratory. The leaves and thalli were rinsed with distilled water and dried at 60 °C for 48–72 h to constant weight.

All samples were analyzed for POC, and \( \delta^{13}C \) and \( \delta^{15}N \). Sediment traps, SPM, plant, and macroalgal samples were homogenized, then acidified to remove carbonates and analyzed for OC by combustion in an elemental analyzer (EuroVector EA 3000) with a precision of 0.06% for OC and 0.01% for total nitrogen for sediments (organic soil standard), and a precision of 0.36% for OC and 0.05% for total nitrogen for plant materials (Apple leaves standard SRM1515 reference material). Carbon and nitrogen stable isotope ratios were determined using a gas isotope ratio mass spectrometer (Thermo Finnigan Delta Plus) after high temperature combustion in an elemental analyzer (Flash 1112 EA). Isotope ratios were expressed in the delta notation (\( \delta^{13}C \), \( \delta^{15}N \)) relative to Vienna PDB and atmospheric nitrogen. Analytical precision was ±0.10‰ for nitrogen and 0.13‰ for carbon, as estimated from an international standard (Peptone) analyzed together with the samples.

Data Analysis

We compared the water parameters (DOC; Chl-a, and POC), SPM concentration, CO2 fluxes and sediment OC stocks across the three locations and the different ecosystems present in the intertidal zone (i.e. mangrove forest, seagrass bed or
tide tidal flat). A Generalized Linear Mixed-Effects Model was constructed for each variable using location and ecosystem as a fixed effect, and the transect point within the ecosystem (MF1, MF2, SB1 etc.) as a nested random effect within each location. For carbon in AGB and BGB of mangrove trees, only location was used as the fixed effect, and the transect point within the ecosystem (MF1, MF2, etc.) was a random effect, nested within the location.

To test significant differences across the locations, transects points and the different ecosystems in each location, an Analysis of Variance (ANOVA), followed by Tukey’s HSD post hoc comparisons were used. T-tests were used to test for differences between light and dark CO₂ incubations. A type I linear regression was used to evaluate the relationship between Chl-a and SPM, POC, and DOC, to evaluate the influence of phytoplankton on DOC, POC and PN.

Diagnostic plots and visual assessments of normality and homogeneity of variation were used to confirm the data conformed to major statistical assumptions (residual homogeneity, independence and normality). Statistical significance was assessed using α = 0.05. Statistical analyses were completed using R version 3.0.2 (R Core Team, 2013), using the packages ‘lm4’ for GLMM (Bates et al. 2015), ‘CAR’ for ANOVA (Fox and Weisberg 2011) and ‘emmeans’ for Tukey’s HSD post hoc comparisons (Lenth et al. 2018).

For the MixSIAR, both SPM and sediment traps isotopic samples were in the range of the 5 potential sources (Fig. 6). Markov Chain Monte Carlo MCM runs with a Chain Length of 1,000,000 were selected, in order to obtain Gelman diagnostics <1.05. Discrimination coefficient was set at 0 (Stock et al. 2018). The relative contribution dimensionless index was calculated by dividing the contribution of each source given by the mixing model (% Contribution) by the percent of surface area (% Area) occupied by the respective ecosystem (terrestrial, mangrove forest, seagrass bed).

\[
\text{Relative Contribution} = \frac{\% \text{Contribution}}{\% \text{Area}} \quad (2)
\]

Percent of surface (% Area) was calculated by dividing the surface area of the ecosystem (i.e. mangrove forest) by the area of the adjacent catchment terrestrial system plus the intertidal systems (mangrove forest, plus tidal flats or seagrass beds).

Results

Carbon Stocks in Biomass and Sediments

Mangrove biomass carbon stocks were significantly different across locations for above ground (GLMM; \( p < 0.01, df = 2, \chi^2 = 10.7 \)) and below ground (GLMM; \( p < 0.01, df = 2, \chi^2 = 9.7 \)) and ranged from 9.5 to 214.8 Mg C ha\(^{-1}\) being lowest at Seletar Island (Fig. 2). Carbon stocks in seagrass plants ranged from 0.02 to 1.4 Mg C ha\(^{-1}\) at Chek Jawa.

Carbon content in the top 15 cm of the sediment ranged from 0.2 to 0.9 Mg C ha\(^{-1}\). Sediment carbon stocks in the 0–15 cm did not differ significantly among mangrove locations (GLMM; \( p = 0.06, df = 2, \chi^2 = 5.5 \)), nor were significant nested random effects detected for sediment carbon stocks for locations or transect points within the ecosystem (GLMM; \( p = 0.08, df = 1, \chi^2 = 2.9 \)) (Fig. 2). No significant differences were found among seagrass beds, mangrove forests and terrestrial soils at Chek Jawa (GLMM; \( p = 0.05, df = 2, \chi^2 = 5.9 \)) or at Sungei Buloh among tidal flats, mangrove forests and terrestrial soils (GLMM; \( p = 0.5, df = 2, \chi^2 = 1.4 \)).

Carbon Content in the Water Column

Mangrove forest organic carbon (OC) was transported towards the sea with the ebb tide as dissolved organic carbon (DOC) and suspended particulate organic carbon (sPOC), the ratio of sPOC/DOC ranging on average across sampling points from 0.3 to 0.8 (Fig. 3b). There was a significant difference among locations for both DOC (GLMM; \( p < 0.01, df = 2, \chi^2 = 45.3 \)) and sPOC (GLMM; \( p = 0.03, df = 2, \chi^2 = 6.9 \)); DOC was lowest at Sungei Buloh (Fig. 3a). No random effects of the transect point within the ecosystem, were detected in any of the water parameters measured, DOC (GLMM; \( p = 0.8, df = 2, \chi^2 = 0.5 \)) and POC (GLMM; \( p = 0.05, df = 2, \chi^2 = 5.9 \)), indicating that neither DOC nor POC concentrations were significantly different inside the mangrove forest and the seagrass bed in Check Jawa. Further, DOC and POC concentrations did not differ significantly among mangrove forests, seagrass beds or tidal flats (Fig. 3).

No significant differences in the quantity of SPM among locations (GLMM; \( p = 0.2, df = 2, \chi^2 = 3.6 \)) or ecosystems (GLMM; \( p = 0.9, df = 2, \chi^2 = 0.1 \)) were observed (Fig. 4a). Chlorophyll-a and DOC showed a significant correlation \( p < 0.01, R^2 = 0.3 \) (Fig. 4b). Location had a significant effect on Chl-a (GLMM; \( p < 0.01, df = 2, \chi^2 = 31.6 \)), with Chl-a concentrations being higher at Seletar Island than the other locations (Fig. 4).

Sediment CO₂ Fluxes

Fluxes of CO₂ from sediment to the atmosphere were variable among transect points. Dark incubations varied between 4 and 79 mmol CO₂ m\(^{-2} \) d\(^{-1}\), and light incubations fluxes between 5 and 61 mmol CO₂ m\(^{-2} \) d\(^{-1}\) (Fig. 5). No significant differences were observed between Seletar Island and Sungei Buloh in dark (GLMM; \( p = 0.6, df = 1, \chi^2 = 0.3 \)) or light incubations (GLMM; \( p = 1, df = 1, \chi^2 < 0.01 \)). Among mangrove forests and tidal flats, only Sungei Buloh presented significantly
different values for light incubation (GLMM; \( p = 0.01 \), \( df = 1 \); \( X^2 = 8.5 \)), being higher in tidal flats.

Overall no significant difference was observed between dark and light incubation chambers at each transect point (T-test; \( p = 0.4 \), \( df = 12 \); \( t = 1 \)) with the exception of transect point MF1 at Seletar Island (Fig. 5), where CO2 fluxes out of the sediment were significantly higher in dark chamber than in light chamber incubations (T-test; \( p = 0.01 \), \( df = 1 \); \( t = 63.7 \)). We did not complete a statistical test for the transect points at Sungei Buloh due to insufficient replication.

**Carbon Sources in SPM and Trap Samples**

The isotopic signatures of SPM grouped close to oceanic and macroalgae in Seletar Island and Chek Jawa (Fig. 6a and c), whereas Sungei Buloh samples grouped closer to the terrestrial and mangrove sources (Fig. 6a). Whilst most of the isotopic signatures of sediment trap samples grouped closer to mangrove and terrestrial sources in Sungei Buloh and Seletar Island, in Chek Jawa the signatures were spread mainly among mangrove plants, terrestrial plants, macroalgae and oceanic sources (Fig. 6b and d).

Contributions of the different sources to the POM analyzed at the different points varied among locations and between type of sample (SPM or trap). Oceanic sources were the highest contributor of POM for SPM at all locations, while sediment trap samples were dominated by mangrove POM (Fig. 7a and b). Seagrass beds were only present at Chek Jawa, and covered a smaller area than the terrestrial zone (Fig. 1), their relative contribution was similar in both SPM...
and trap samples (Fig. 7c and d). The relative contributions showed that mangrove forests were the greatest contributor of the three ecosystems with fixed area (mangrove, seagrass, terrestrial) at all locations and for all samples except for the SPM samples in Sungei Buloh.

Suspended particulate matter samples exhibited greater contribution of terrestrial sources than sediment trap samples at all locations. Seagrass beds were not a strong contributor of POM in SPM or sediment traps samples (Fig. 7a and b). However, at Chek Jawa the contribution of macroalgae was higher than that of seagrass in both SPM and sediment traps samples (Fig. 7).

Discussion
Carbon Stocks, Carbon in the Ebb Tide and CO₂ Fluxes

Despite similar climatic conditions, mangrove forests in Northern Singapore exhibited differing standing biomass stocks, but carbon stocks in, and CO₂ fluxes from, the upper sediment layers 0–15 cm showed no difference. Mangrove forests at Sungei Buloh presented the highest carbon stocks (Fig. 2), and Sungei Buloh also had the largest mangrove forest cover with a higher riverine/fresh water influence and terrestrial contribution than Chek Jawa and Seletar Island (Figs. 1 and 2). Carbon stocks in the upper sediment layer (0–15 cm) did not significantly differ among mangrove forests, terrestrial forests, tidal flats and seagrass beds at each location. This could mainly be due to high variability of organic carbon content in the sediment across the sampled transect points especially at the mangrove forests. Mangroves at Seletar Island were smaller, patchier, and without riverine influence (Fig. 1), and as such showed lower aboveground carbon stocks (Fig. 2). Results of this study, compared to a previous study of mangrove above- and belowground biomass measurements in northern Singapore, were lower at Chek Jawa (Phang et al. 2015) but similar or higher at Sungei Buloh (Friess et al. 2016). However, variation may be due to differences in specific sampling sites, temporal variation, or calculations of carbon biomass methods between the studies. Mangrove forests in Singapore, as well as other mangrove forests, show spatial variation in biomass density, and therefore in carbon stocks observed between the locations, corroborating a common pattern in mangrove forests, due to variation in natural factors such as geomorphic setting and previous
Fig. 6 Scatter plots with isotopic of carbon ($\delta^{13}$C o/oo) and nitrogen ($\delta^{15}$N o/oo) composition of suspended particulate matter (SPM) (Panel A and C) and sediment trap samples (Panel B and D). Sources are expressed as mean ± SE in different shapes without fill, in Isolated locations only 4 sources (Panel C and D), see legend for details. Check Jawa was the only connected location therefore the only location where seagrass were used as a source (Panel A and B).

Fig. 7 Stacked columns with percentage contributions of each source determined from the mixing model (a and b) and relative contribution calculated from Eq. 2(c and d). a-b show total percentage contribution of each source for suspended particulate matter (SPM) and sediment trap samples. c-d represents a dimensionless index indicating the relative contribution of terrestrial zone, mangrove forest and seagrass bed sources for SPM and sediment trap samples. X-axis shows the transect points inside the ecosystem where samples were taken; mangrove forests (MF), seagrass bed (SB), tidal flat (TF). The lowest the number is closer to the oceanic fringe of the ecosystem. Different colors indicate different sources indicated in the legend.
higher concentration than sPOC (Fig. 3). To measure the contribution of different ecosystem sources, elucidating the origin of DOC is necessary in order to determine which process is occurring since we did not remove the biofilms of the sediments to measure CO₂ fluxes and we could not measure CO₂ fluxes in Chek Jawa. Additional studies including different tidal regimes and removal of biofilms will help to determine what controls CO₂ fluxes in Singapore mangrove forest. As we found no differences across locations and only in one location light chambers show between mangrove forest and seagrass beds.

**Mangrove Forests and Adjacent Ecosystems: Donors and Recipients**

While climatically similar, the mangrove forests in this study had different local geomorphic settings and landscape structures, such as higher riverine influence in Sungei Buloh, fringing patches of mangrove forests in Seletar Island, and fringing mangrove forests with an adjacent seagrass bed in Chek Jawa (Fig. 1). Sampling was completed during late Northeast Monsoon, which is a season of low rainfall and high wind speeds, suggesting that POM from runoff within the catchment area contributed little during this time of the year. We hypothesize that the influence of terrestrial plants will be higher during highly rainy seasons due to the higher input of terrestrially derived POM. However, further studies are needed to test this hypothesis.

The transport of POC from mangroves to adjacent ecosystems depends on different factors, such as river discharge, surface area of ecosystems, estuarine geomorphology and hydrodynamics (Hyndes et al. 2014). Geomorphic settings corresponded to the isotopic signal of SPM samples. For example, Sungei Buloh (riverine location) had isotopic signals closer to mangrove trees and terrestrial plants, while fringing locations had signals closer to oceanic and macroalgal sources (Figs. 1 and 6). The geomorphic setting of Sungei Buloh facilitates the transport of detrital material from terrestrial plants and mangrove trees into coastal water bodies due to channel-associated water movement. Samples of SPM at Sungei Buloh showed a higher terrestrial contribution than the other locations (Fig. 7a), as Sungei Buloh is influenced by upstream freshwater sources containing terrestrial plant detritus (Fig. 1). In general, the biggest contribution of all sediment trap samples were mangrove vegetation (Fig. 7b and d), as indicated by the closer grouping of all samples to the isotopic signatures of mangrove trees (Fig. 6b and d). While fringing mangrove forests, such as Chek jawa and Seletar island, are less likely to have terrestrial inputs to the POM, due to the lack of water inflow from the catchment area. Therefore, we hypothesize that exchange of POM between terrestrial and coastal waters would be mainly driven by wind action or runoff. At Seletar Island, the terrestrial plant sources are located closer to the oceanic fringe, hence, leaves move easily (i.e. via wind action) and therefore are transported out with the wave currents faster. In Chek Java, the terrestrial plant area is isolated...
from the ocean by the mangrove forest (Fig. 1) showing a higher contribution of mangrove trees to the POM than Sungei Buloh (Fig. 7).

According to the relative contribution dimensionless index (Eq. 2), catchment areas contributed relatively (as per unit area) more to POM than seagrass in Chek Jawa, whereas the mangrove forest contributed relatively more to POM than the catchment area in Seletar Island. In contrast, the relative POM contribution of the catchment area was higher than that of the mangrove forest in Sungei Buloh (Fig. 7c). The relative contribution was not calculated for oceanic and macroalgal sources, because no representative surface area for the ‘ocean’ could be calculated. Additionally, we suspect that the macroalgae found in the mangrove forest were mainly allochthonous and transported by local currents, since collected algae were not attached to any hard substrate. Local geomorphological setting together with local differences in hydrodynamics will influence the relative contribution of vegetated coastal ecosystems and catchment area. However, more studies with more with different landscape structure mangrove forests are necessary to understand those processes, since patch size and seascape configuration are rarely included in connectivity studies, and if so mainly focus on fish (Boström et al. 2011). In addition to landscape structure, information on litter production and decay will be necessary for understanding the contributions of each primary producer, since it has been shown that decay rates are slower for mangrove trees and terrestrial plants, intermediate for seagrass plants and fast for macroalgae (Kristensen, 1994; reviewed in Zimmer 2019).

High oceanic contribution was found in SPM samples (Fig. 7a), corroborating former studies (Almahasheer et al. 2017; Gao et al. 2012; Gillis et al. 2014b) and stressing the importance of ocean-derived input for coastal ecosystems (Orr et al. 2005). Inputs of detritus from adjacent ecosystems (i.e. seagrass and macroalgal beds) can enhance sediment respiration (Lovelock 2008). Even though the catchment area was smaller than the seagrass bed in Chek Jawa, terrestrial plants had a greater contribution to the POM. Isotopic signatures of SPM grouped closer to oceanic and macroalgae, while sediment trap samples grouped closer to terrestrial and mangrove sources (Fig. 6). According to these findings, as well as the results of the MixSIAR models (Fig. 7), seagrass did not contribute markedly. Seagrass beds have been reported significant donors of POM in other studies (Almahasheer et al. 2017; Gillis et al. 2014b). However, these studies reported larger seagrass beds dominated by larger species, such as *Enhalus acoroides*. Seagrass beds in the study area, by contrast, are relatively small and dominated by smaller species like *Cymodocea rotundata*, *Halodule uninervis*, and *Halophila* spp. (Phang et al. 2015). The contribution of seagrass beds could be more important than macroalgal contributions in long-term sediment carbon stocks at the coastal and seascape level, as they are formed by recalcitrant compounds, which could be buried for a longer period within sediments (Trevathan-Tackett et al. 2017).

While we did not measure the standing biomass of all primary producers, we assumed that macroalgal biomass was less than mangrove trees, seagrass and terrestrial plants biomass. However, macroalgal contributions were larger than, or almost equal to, those of mangrove trees or terrestrial plants in fringing landscape structure (i.e. Chek Jawa) (Fig. 7). Particulate organic matter had a higher percentage of material derived from macroalgae than from seagrass plants in Chek Jawa. (Fig. 7a and b). Contrary, in Sungei Buloh (riverine geomorphic setting), even macroalgae contributed more than terrestrial plants to sediment trap samples, they contributed equal to, or less than, mangrove trees and oceanic sources, and they were the smallest contributors of SPM samples. Those results highlight that despite their low biomass, macroalgae are an important source of POM for fringing mangroves along the northern coast of Singapore. The role of macroalgae has been discussed as a potential carbon donor to adjacent ecosystems, due to algal material being exported and stored into adjacent habitats (Hill et al. 2015; Krause-Jensen et al. 2018). Macroalgal carbon compounds degrade faster, being released in form of DOC or CO₂ (Hill et al. 2015; Krause-Jensen et al. 2018; Mews et al. 2006). The presence of refractory macroalgal compounds (carbonates, long-chain lipids, alginates, xylans, and sulfated polysaccharides), combined with factors such as retention time in the coastal habitat, fast burial, and anaerobic decay, could render macroalgae a significant contributor to long-term carbon storage (Trevathan-Tackett et al. 2015). However, further research is needed to quantify these contributions (Hill et al. 2015). To our knowledge, no research has focused on the exchange of POM derived from macroalgae in the tropics; our results suggest that the contribution of macroalgae could be significant in understanding carbon dynamics in tropical coastal areas.

Other studies have shown that mangrove forests that extend inland may retain much of their POM and nutrients of allochthonous origin (Adame and Lovelock 2011; Alongi et al. 2016). Interactions between productivity, geomorphology and hydrology influence the connectivity of ecosystems, changes in connectivity could alter the exchange and storage of organic matter provided by mangrove forests (Alongi 2014; Feller et al. 2010). Recently, a global study showed the influence of coastal morphology on carbon budgets (Twilley et al. 2018). They found that mangrove stocks in the sediment varied markedly across different types of coastal environmental settings, increasing from river-dominated to tide/wave-dominated to carbonate coastlines, but independently of forest
architecture. Our results also showed that local landscape structure were important for carbon stocks and for connectivity in the sense of the exchange of organic matter. In our study location, mangrove forests with riverine influence (Sungei Buloh) and with seagrass beds (Chek Jawa) tend to have higher carbon stocks i.e. and compared to Seletar Island. Nevertheless, further studies evaluating these and additionally landscape structure interactions will help to further understand carbon dynamics.

Conclusions

In order to sustain carbon storage provided by coastal ecosystems, geomorphological settings and ecosystem connectivity should be taken into account in conservation and restoration plans. Mangrove forests clearly contributed the most to POM across the intertidal landscape, with differences in contributions from different primary producers across different landscape spatial arrangements. Therefore, anthropogenic induced modifications of local geomorphic settings at the coastal level (i.e. reduction of mangrove forests area or changing fresh water fluxes) may affect carbon accumulation in mangrove forests and adjacent coastal ecosystems. Conversely, active human modifications of local geomorphic settings may also increase carbon sequestration in mangrove forests and adjacent coastal ecosystems, such as seagrass and tidal flats. Further studies determining the origin of DOC together with tidal fluxes will help to elucidate the role of hydrodynamics, driven by the geomorphic setting, in the interchange and sequestration of dissolved and particulate organic carbon at the coastal landscape level.

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Compliance with ethical standards

Data Repository Data used in this study will be deposited at scientific data repository of ZMT.

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