Distribution, input pathway and mass inventory of black carbon in sediments of the Gulf of Thailand, SE Asia

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Abstract

The coastal margins around Southeast Asia (SE Asia) may serve as an ideal location to study the source-sink process of sedimentary black carbon (BC) because SE Asia has been identified as one of the major BC emission source regions in the world. This study provides an extensive picture of recent regional-scale sedimentary BC sequestration in the Gulf of Thailand (GOT), a tropical marine system in SE Asia. Generally, the sedimentary BC concentrations (0.07–3.99 mg/g) were in the low to moderate ranges of those obtained in other coastal sediments around the world. Regional variability of the BC and its correlation with the sediment grain size and total organic carbon (TOC) content indicated a general hydrodynamic constraint on BC occurrence in the lower Gulf in contrast to the upper Gulf with a more source dependence due to the direct land-based input. BC/TOC% values and the varied BC components (char and soot), as well as their correlations suggested that char was the predominant constituents of sedimentary BC both in the upper and lower Gulf, which could be mainly derived from biomass burning and entered into the nearshore region through direct fluvial transport and surface run-off. The estimated BC burial flux (~212 μg/cm²/y) and mass inventory (~200 Gg/y) in the GOT on the hundred-year timescale were of the same order of magnitude compared with other oceanic margins, and thus the tropical shelf sediments from SE Asia could serve as an important sink of land-emitted BC.

1. Introduction

Black carbon (BC), including both the char and soot components, is usually defined as the recalcitrant carbonaceous residue produced from the incomplete combustion of biomass and fossil fuels (Goldberg, 1985; Schmidt and Noack, 2000; Baldock and Smernik, 2002; Dickens et al., 2004). The widespread combustion process and refractory nature make BC widely distributed in environmental matrices, such as the atmosphere, soils, fresh/sea water, ice and sediments. The emitted BC can eventually be transferred into the ocean through direct atmospheric deposition and river run-off by soil erosion (e.g., Goldberg, 1985; Suman et al., 1997; Gustafsson and Gschwend, 1998; Kuhlbusch, 1998; Mitra et al., 2002; Elmqquist et al., 2008; Lohmann et al., 2009), which is an important process of the global carbon cycle as transferring the carbon from a rapidly cycling atmosphere-biosphere system into
the long-term geological carbon pool (Seiler and Crutzen, 1980; Suman et al., 1997; Kuhlbusch, 1998; Middelburg et al., 1999). The interaction of BC among the various carbon pools is also relevant in predicting the climate change of the future earth (Bond et al., 2013).

It has been estimated that ~90% of the marine BC deposition occurs on the marginal settings with the area less than 10% of the world ocean (Suman et al., 1997); however, the magnitude of the BC mass inventory in marine sediments on the global scale remains largely unknown (e.g., Gustafsson and Gschwend, 1998; Dickens et al., 2004; Masiello, 2004; Elmquist et al., 2008), which may be partially due to the scarcity of large-scale observations. Therefore, the extensive study of the BC occurrence and mass inventory in the continental margins is essential for a better understanding of the global biogeochemical cycle of BC. However, a vast majority of the existing studies on the sedimentary BC in the coastal margins were geographically within high-latitude regions (e.g., Europe and America), such as the Northern European shelf (Sánchez-García et al., 2012), the Gulf of Cádiz (Sánchez-García et al., 2013), the Gulf of Maine (Gustafsson and Gschwend, 1998; Flores-Cervantes et al., 2009), the pan-arctic estuaries (Guo et al., 2004; Elmquist et al., 2008), the Washington coast (Dickens et al., 2004), etc., however, only little attention has been paid to the continental shelf of Asia (e.g., Wang and Li, 2007; Sun et al., 2008), particularly for the Southeast Asia (SE Asia), a tropical region, which has been identified as a major BC emission source region due to frequent forest fires, biomass burning and escalating fossil fuel utilisation (Streets et al., 2003). Moreover, the tropical coastal margins in SE Asia also serve as an important source emission region for coastal BC export due to the inherent nature of rapid transferring of land-based materials into the aquatic system via strong rain and surface run-off (Nittroer et al., 1995; Zakaria et al., 2002; Saha et al., 2009). Therefore, the coastal sediments from tropical SE Asia may be an important reservoir for the exported BC from the adjacent landmass. Nevertheless, to date, the characterisation of sedimentary BC in this tropical region remains poorly understood.

The Gulf of Thailand (GOT), a shallow and semi-enclosed shelf sea in SE Asia, has been subjected to a huge BC emission from biomass burning, forest fires and fossil fuel consumption from the adjacent regions (Saha et al., 2009; Sahu et al., 2011; Huang et al., 2013). Because of the nutrient-rich, shallow waters and their confined nature, the ecosystem of the GOT is especially vulnerable to human activities (Srisukswad et al., 1997), and increased anthropogenic activities around the GOT have induced severe environmental pollution, especially for the upper Gulf with significant river export of land-based contaminants (e.g., the Chao Phraya River) (Wattayakorn et al., 1998). Petroleum hydrocarbon contamination is evident in the nearshore waters of the GOT (Wattayakorn et al., 1998; Wattayakorn, 2012), and sedimentary polycyclic aromatic hydrocarbons (PAHs) exhibit a dominant river influence with mixed pyrogenic and petrogenic origins in the upper Gulf but with a dominant pyrogenic signature in the lower Gulf through atmospheric deposition (Boonyatumanond et al., 2006). This result may imply a different input pathway and source-sink processes for the combustion-derived substances (such as BC and PAHs) in the different GOT compartments (e.g., the upper vs. the lower Gulf).

In the present study, the large-scale occurrence of sedimentary BC and its mass inventory in the GOT was first examined to present a comprehensive study on BC abundance and its distribution pattern, input pathway and mass inventory in this tropical marginal system.

2. Materials and methods

2.1. Study area and sediment sampling

The Gulf of Thailand (GOT) is a shallow, semi-enclosed tropical marine embayment situated in the South China Sea, which is surrounded by the land mass of Malaysia, Thailand, Cambodia and Vietnam (Fig. 1). The GOT is relatively shallow with a mean depth of 45 m and a maximum depth of 80 m. The Gulf can be usually divided into two parts, the upper Gulf and the lower Gulf (Fig. 1). The upper Gulf is the northernmost part of the GOT, covering approximately 10,000 km² and receives a large amount of sewage, runoff and sediments, especially from the Chao Phraya River, which has a catchment area of 162,000 km² and flows through several cities, including the mega-city of Bangkok (Wattayakorn et al., 1998; Boonyatumanond et al., 2006). The regional climate is generally warm and humid, with a typical southwest monsoon that prevails from May to September and a northeast monsoon that prevails from November to February. The induced seasonal circulation in the GOT is generally weak and variable (Wattayakorn et al., 1998; Wattayakorn, 2012), where an apparent counter-clockwise circulation occurs in the upper Gulf during the northeast monsoon, whereas both clockwise and counter-clockwise circulation occurs during the southwest monsoon, depending on the wind conditions and external flow through the open boundary (Sojiesuporn and Putikiatkajorn, 1998; Wattayakorn et al., 1998). The Gulf is poorly flushed on the whole, with little mixing, especially in the upper part (Wattayakorn et al., 1998), resulting in the majority of sediments and associated contaminants from the fluvial inputs primarily depositing in the upper Gulf (Srisukswad et al., 1997). Sediment accumulation in the open GOT is mainly constrained by a combination of wind-driven currents, tides and its bottom topography (Takahashi et al., 1984). Muddy sediments mainly occur in the coastal area near the mouths of the major rivers in the upper Gulf, in addition to the central and western parts (near Samui Island) in the lower Gulf (Windom et al., 1982).

Ninety-three surface sediment samples were strategically collected on a regional-scale through three cruises conducted by R/V Boon-Lerd Pa-Sook and R/V SEAFDEC 2 (SEAFDEC, i.e., Southeast Asian Fisheries Development Center) during 2010–2012. The samples were collected using a stainless steel box corer deployed from the vessels. All sediment samples (0–3 cm) were wrapped in pre-combusted aluminium foil and stored at −20 °C until analysis.

2.2. Analytical procedure

Due to its complex organic and mineral matrices, the separation and quantification of BC in sediments is complicated, and several procedures have been developed for specific studies (Gustafsson et al., 1997, 2001; Song et al., 2002; Hammes et al., 2007; Han et al., 2007a, 2007b; Khan et al., 2009; Poot et al., 2009; Meredith et al., 2012; Wiedemeier et al., 2015). In the present work, the wet-chemical treatment combined with thermal/optical reflectance (TOR) detection was adopted (Han et al., 2007a, 2007b; Fang et al., 2015). Briefly, the thawed, freeze-dried and homogenized (<80 meshes) sediment samples were treated with hydrochloric and hydrofluoric acids to remove inorganic materials. The acid-treated sediment residue was then filtered through pre-combusted quartz fibre filters (Whatman, 450 mm for 4 h and 47 mm in diameter). The filter samples were air dried and subsequently analysed for BC on a Desert Research Institute (DRI) Model 2001 Thermal/Optical Carbon Analyser (Atmoslytic Inc., Calabasas, CA) following the Interagency Monitoring of Protected Visual Environment (IMPROVE) protocols. A 0.544 cm² circular punch drilled from the filter was placed in a quartz boat and then sent into an oven. During the carbon analysis, the oven was ramped from 100 °C to 500 °C at 10 °C/min and held at 500 °C for 120 min. After cooling, the blank and sample filters were treated with 10% HCl and 10% HNO₃ to release the BC mass, which was subsequently measured by TOC analysis.

The determination of BC mass was performed using a Shimadzu TOC-5010C analyzer equipped with an auto sampler and an advance combustor. The digestion of the blank and sample filters was performed at 500 °C for 2 h. The blank reading was subtracted from the sample reading, and the difference represented the BC mass on the filter. The error introduced by this process was kept less than ±10%.

The analytical procedure was conducted twice for each sample, and the results are reported as the mean of the two analyses. The reproducibility of the BC analysis was determined by running the method on the same filters and samples in the same instrument over a period of six months. The reproducibility of the method was found to be ±10% for BC mass. The precision and accuracy of the method were determined using standard reference materials (SRMs) provided by the National Institute of Standards and Technology (NIST) and the International Atomic Energy Agency (IAEA). The recoveries of BC from the SRMs were found to be within the recommended accuracy range of ±10%.

2.3. Characterisation of sedimentary BC

The characterisation of sedimentary BC includes the identification of BC types, BC mass, and the distribution of BC among different sedimentary compartments. The BC types were determined using a combination of thermal-optical analysis and stable carbon isotope analysis. The BC mass was determined using a Shimadzu TOC-5010C analyzer equipped with an auto sampler and an advance combustor. The distribution of BC among different sedimentary compartments was determined using a combination of chemical fractionation and stable carbon isotope analysis.

The stable carbon isotope analysis was performed using a Thermo Fisher Delta V Advantage isotope ratio mass spectrometer. The accuracy of the isotope ratio measurement was ±0.1‰.

The BC mass and the distribution of BC among different sedimentary compartments were determined using a combination of chemical fractionation and stable carbon isotope analysis. The chemical fractionation was performed using a combination of sequential acid digestion and sequential solvent extraction. The sequential acid digestion was performed using a combination of HCl, HNO₃, and HF. The sequential solvent extraction was performed using a combination of chloroform, methanol, and acetone.

The accuracy of the isotope ratio measurement was ±0.1‰.
temperature steps (580, 740, and 840 °C). A portion of the volatile-
zible carbon chars in the absence of oxygen as indicated by the
declined reflectance of the laser, which was termed “pyrolyzed
organic carbon (POC)”. Once in the oxidizing atmosphere, the POC
evolve together with the original EC and leave the filter. The
quantity of POC is defined as the carbon that evolves to the time at
which the laser reflectance achieves its initial value (Han et al.,
2007a). The IMPROVE protocol defined EC as the sum of the three
EC sub-fractions minus the POC (i.e., EC = EC1 + EC2 + EC3 − POC).
Based on the finding that char and soot (two subtype of BC) in
standard reference materials can be stepwise oxidized in the EC1
and EC2 + EC3 steps, respectively, it was proposed that the
IMPROVE method can be used to differentiate between char and
soot, and char was defined as EC1 − POC, and soot as EC2 + EC3

Fig. 1. Locations of the sampling sites (n = 93) in the GOT in SE Asia, the stations located at the upper Gulf (n = 33) with the corresponding labels were marked in the dashed rectangle.
Detailed procedures for the BC separation and quantification are available in our recent work (Fang et al., 2015).

For quality assurance and quality control (QA/QC), whether the residue on the filters has an even distribution is a primary factor that markedly influences the accuracy of the BC concentration determination (Khan et al., 2009; Han et al., 2011; Fang et al., 2015). The typical filters loaded with acid-treated sediment residue are displayed in Fig. S1, and the residue on the filters has an even distribution to the naked eye. To confirm this distribution further, 10% of the total filters (n = 10) were randomly selected, and two punches drilled from different positions within a similar filter were analyzed for BC. The relative standard deviation (RSD, %) of the measured BC concentration was in the range of 2–8%, and averaged 4%, fully demonstrating the even distribution of the residue on the filters.

Additionally, blank and replicate samples as well as standard reference material (NIST 1941b) were analyzed simultaneously at a frequency of one per ten samples. The blank samples (n = 10) yielded 0.00 ± 0.00 μg/cm² for BC, and the RSD of 10 pairs of replicate analyses was in the range of 0–11%, and averaged within 5%. The BC concentration measured in SRM 1941b was 10.56 ± 1.28 mg/g (n = 10), which was well in accordance with the values reported by recent studies (Hammes et al., 2007; Han et al., 2007a; Cong et al., 2013; Fang et al., 2015). These QA/QC results suggested that the BC analytical method used here was reliable and repeatable (see Supplementary materials).

Detailed analytical methods for the TOC and sediment grain size were provided elsewhere (Hu et al., 2009, 2011). In brief, the grain size composition of each sample was determined using a Laser Particle Size Analyser (Mastersizer 2000, Malvern Instruments Ltd., UK). Portions of the sediment samples were treated with 4 M HCl to remove carbonates. The resulting carbonate-free samples were analysed for TOC in duplicate with a Vario EL-III Elemental Analyser.

### 2.3. Estimation of the burial flux and mass inventory of the BC in the GOT

The burial flux and mass inventory of BC in this work were estimated according to the procedures from Tolosa et al. (1996); Hu et al. (2014). Briefly, the study area was divided into 93 homogenous sectors, with the sampling sites located in the centre of their respective sectors by coupling with the recent regional-scale varied sediment mass accumulation rates for each sample site (Fig. 7a, Srisuksawad et al., 1997). The BC burial flux (μg/cm²/yr) among each homogenous sector was estimated by the following equation:

$$ F_{\text{burial}} = C_i \cdot M_i $$

where $C_i$ is the BC concentration in the sediment sample of each sector, and $M_i$ is the reported mass sediment accumulation rate among each divided sector (Srisuksawad et al., 1997). The calculated $F_{\text{burial}}$ is then applied to the corresponding areal extension of each divided sector to construct the BC mass inventory (l) in the study area according to the following equation:

$$ I = \sum_{i=1}^{93} C_i \cdot M_i \cdot A_i $$

where $C_i$ is the BC concentration in the sediment sample of each sector, $M_i$ is the reported mass sediment accumulation rate among each divided sector (Srisuksawad et al., 1997) and $A_i$ is the area of each divided sector.

### 3. Results and discussion

#### 3.1. Sediment grain-sizes and background depositional settings

Surface sediments in the GOT primarily consisted of silt and sand silt with mean grain size (Mz) ranging from 11.5 to 547.6 μm (mean, 52.9 μm) (Fig. 2a). Fine deposits predominantly occurred nearshore around the mouth of the major rivers from the upper Gulf and the central basin (Fig. 2b). The south of the Chao Phraya delta, the transition zone between the upper and lower Gulf, was mainly characterised as silty sand with coarser particle size (>60 μm). The upper Gulf was mainly influenced by four rivers (the Chao Phraya, the Mae Klong, the Ta Chin and the Bang Pakong rivers) (Fig. 1), of which the total suspended sediments into the upper Gulf were estimated to be 6.32 million tons per year; but they were not efficiently transported southward into the central basin (Emery and Niino, 1963; Windom et al., 1982; Srisuksawad et al., 1997), resulting in relatively higher sediment accumulation rates in the upper Gulf (Windom et al., 1982; Srisuksawad et al., 1997). Water circulation and the depositional mechanism in the open GOT is complex and dominated by the combination of wind-driven currents, tides, density gradients and bottom topography (Takahashi et al., 1984). Even the sources and formation mechanism of the fine deposits in the central basin are not clear; the water circulation, inflowing and/or exchange with the open sea could be responsible for the depositional settings in the lower Gulf (Srisuksawad et al., 1997).

#### 3.2. Occurrence and spatial distribution of BC and the potential controlling factors

3.2.1. Occurrence and distribution of BC and TOC in the surface sediments

The BC concentrations in the GOT ranged from 0.07 to 3.99 mg/g with a mean of 1.0 ± 0.92 mg/g (Table 1). The BC content values in this work (measured by the IMPROVE_A TOR method; from Han et al., 2007a, 2007b) were in the low to moderate ranges of the BC concentrations obtained by different versions of the thermal methods in other coastal sediments around the world, such as the East China Seas (<0.1–2.45 mg/g, CTO method; Wang and Li, 2007; Kang et al., 2009), the Laizhou Bay in China (0.02–1.29 mg/g, TOR method; Fang et al., 2014), the Mexican margin (0.9–2.6 mg/g, modified CTO method; Gelin et al., 2001), the Gulf of Maine (0.11–1.73 mg/g, CTO method; Gustafsson and Gschwend, 1998) and coastal Pakistan (0.6–2.5 mg/g, CTO method; Ali et al., 2014). The BC values were much lower than that from the Swedish continental shelf (0.58–17.66 mg/g, CTO method; Sánchez-García et al., 2012) (Table 1). The TOC content in the GOT showed a relatively wide range, varying between 1.8 and 29.3 mg/g (dry weight), and the BC accounted for 3.2–26.7% of the TOC with a mean of 10.8 ± 4.9%. The obtained BC/TOC values in this work were relatively lower than those in other coastal sediments (up to 50 ± 40%), but comparable to those in abyssal sediments (15 ± 2% to 21 ± 6%) (Verardo and Ruddiman, 1996; Masiello and Druiff, 1998; Gustafsson and Gschwend, 1998; Lohmann et al., 2009; Coppola et al., 2014), revealing that the BC represents an important fraction of the sedimentary OC pool in the tropical marine system of SE Asia.

Spatially, higher BC values (>1 mg/g) were found in the nearshore of upper Gulf, the offshore regions over the central basin and the western part around the Sami Island (Fig. 3a). Pronounced low BC concentrations (<0.3 mg/g) were mainly found at the south of the Chao Phraya delta and the central and southern part of the lower Gulf. This spatial variability of BC generally displayed a similar pattern with that of sediment grain size (Fig. 2), suggesting that the spatially higher BC value area is mainly controlled by the grain size of the sediments.
indicating a potential influence of hydrodynamic condition on the BC accumulation. The TOC distribution in the GOT was also consistent with that of BC, and the highest values were determined in the nearshore of the upper Gulf (Fig. 3b), where the mean TOC concentration was $11.8 \pm 7.0$ mg/g compared to that of $7.2 \pm 4.5$ mg/g in the lower Gulf, suggesting a more source-dependent characteristic in the upper Gulf due to the direct fluvial inputs, in contrary to the more hydrodynamic constrains in the lower Gulf. Relatively higher BC/TOC% values were mainly observed in the nearshore with proximity to the coast, which occurred not only in the upper Gulf, but also in the southern-most part of the lower Gulf (Fig. 3c). This finding could be related to the direct land-based input of BC in these areas through fluvial discharge and/or surface runoff. It has been suggested that terrigenous sediments and associated materials discharged by the coastal rivers in the upper Gulf were mainly constrained within the nearshore with limited offshore dispersion (Srisuksawad et al., 1997; Boonyatumanond et al., 2007); while the high values of BC/TOC% in the southern part of the nearshore in lower Gulf could be more determined by the lower TOC abundance (TOC < 3 mg/g) with a steady concentration of BC in the sediments. The abrupt decrease of TOC concentration in these typical nearshore samples were more related to the intermittent extensive surface run-off owing to the heavy and frequent rain (Boonyatumanond et al., 2006), which could result in a large influx of heavier and/or large particles with limited TOC loading.
3.2.2. Relationships between BC, TOC and sediment grain size and potential implications

BC in the majority of the samples correlated with the sediment fine fraction (Fig. 4a), suggesting a general sorption control of the BC enrichment in fine-grained sediments; however, a separated grouping of the fine samples with relatively higher BC contents was observed and marked with the corresponding sites (Fig. 4a), and they were mainly from the river mouth in the coastal upper Gulf (e.g., sites T003, T004, T030) and from the eastern offshore area in central basin (e.g., sites T26, T34, T43). BC showed a significant and comparable correlation with TOC in the samples both from the upper and lower Gulf (Fig. 4b), reflecting a general hydrodynamic controls and/or their similar initial input pathways (e.g., land-based provenance). For the upper Gulf, the positive correlation between BC and TOC could be more related to the similar direct land-based input pathways, i.e., the extensive fluvial input and surface run-off from fired-impacted soils; while for the lower Gulf, this co-variance should be more explained by their coupled sedimentary dynamics caused by the high sorption and affinity of BC and TOC on the finer particles (e.g., Muri et al., 2003; Hu et al., 2014). This suggests that the hydrodynamic transport and depositional mechanism could

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**Fig. 4.** The scatter plots of the BC abundance between sediment grain size (a) and TOC (b).

**Fig. 5.** Spatial distribution of the char (a) and soot (b) and their corresponding ratios (char/soot) (c).

**Fig. 6.** The scatter plots of the correlation between char and soot components.

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also be an important factor affecting the occurrence of BC and TOC and their relationships, especially in the open GOT. Actually, this good correlation between BC and bulk OM was not only noticed in the particle phase (e.g., in soil, Glaser and Amelung, 2003; Cusack et al., 2012) but also in the dissolved phase and has been reported worldwide (e.g., Dittmar et al., 2012; Jaffé et al., 2013; Ding et al., 2013, 2015). The close relationship between BC and bulk OM in the dissolved phase could indicate that the release and subsequent sorption and desorption processes of BC are a complex function of environmental factors and particle properties, just as for bulk dissolved OM (Jaffé et al., 2013), which suggests that the mobilization of BC and bulk OM in various matrices is also mechanistically coupled.

It was however, also noticed, that the corresponding slope between TOC and BC in the upper Gulf were different from that of the lower part (Fig. 4b) as reflected by the lower percentage of BC in TOC in the upper Gulf (~12%, p < 0.01) compared to that in the lower GOT (~17%, p < 0.01). This difference could be explained as the following points: (1) the lower GOT has more significant input of soot-BC compared to the upper GOT (see below), which is served as the additional BC source and largely uncoupled from land-based sedimentary OM, resulting in the higher slope therein; (2) the different sedimentary OM pool in the upper and lower Gulf. Since phytoplankton production was reported to be a major contributor to the sedimentary OM pools in the lower Gulf (e.g., Meksumpun et al., 2005), the less refractory nature of marine OM

Table 1

| Study area            | C_{BC} (mg/g) | C_{TOC} (mg/g) | F_{burial of BC} (μg/cm²/yr) | References                |
|-----------------------|---------------|----------------|------------------------------|---------------------------|
| This study            |               |                |                              |                           |
| Gulf of Thailand      | 0.07–3.99     | 1.01 ± 0.93    | 1.8–29.3                    | 8.8 ± 5.9                 | 15–980 212 ± 228          | This study |
| The upper Gulf        | 0.07–3.63     | 1.23 ± 0.97    | 2.3–29.3                    | 11.8 ± 7.0                | 36–980 336 ± 259          | This study |
| The lower Gulf        | 0.13–3.99     | 0.89 ± 0.88    | 1.8–21.2                    | 7.2 ± 4.5                 | 15–923 144 ± 178          | This study |
| Coastal margin settings |             |                |                              |                           |                           |           |
| Coastal of Pakistan   | 0.60–2.50     | 0.65 ± 0.46    | 2.8–28.6                    | 11.7 ± 6.4                | 183–766 330 ± 155        | Ali et al., 2014 |
| Gulf of Cadiz, SW Spain | 0.10–1.10    | 0.40 ± 0.30    | 5.0–14.0                    | 8.8 ± 2.6                 | 42–285 120 ± 76          | Sánchez-García et al., 2013 |
| Swedish Continental Shelf | 0.58–17.66   | 2.41 ± 2.18    | 4.8–168.0                   | 49.1 ± 28.2               | 267–8067 1103            | Sánchez-García et al., 2012 |
| Gulf of Maine, USA    | 0.11–1.73     | 0.65 ± 0.46    | 1.5–22.5                    | 12.7 ± 6.6                | 86–194 131 ± 38          | Gustafsson and Gschwend, 1998 |
| Mexican margin        | 0.9–2.6       | –              | 17.6–79.9                   | –                         | –                         | Gélinas et al., 2001 |
| East China Sea        | <0.1–2.45     | 0.86 ± 0.88    | 3.0–44.4                    | 7.6 ± 3.2                 | –                         | Kang et al., 2009 |
| Pelagic regimes       |               |                |                              |                           |                           |           |
| Pacific Ocean         | 0.01–0.99     | 0.45 ± 0.31    | –                            | –                         | 0.002–0.2 0.09 ± 0.06    | Smith et al., 1973 |
| South Atlantic Ocean  | 0.40–1.70     | 1.02 ± 0.26    | 2.5–44.7                    | 14.1 ± 12.3               | 0.50–7.80 2.27 ± 1.73    | Lohmann et al., 2009 |
| Madeira Abyssal Plain | 0.74–2.85     | 1.62 ± 0.83    | 1.8–11.4                    | 6.1 ± 4.4                 | –                         | Middelburg et al., 1999 |

Fig. 7. Spatial distribution of the regional-scale recent sediment mass accumulation (from Srisuksawad et al., 1997) (a) and the BC burial flux (b) in the GOT.
thus may also be partly related to the varying slope observed in the lower GOT.

3.3. Occurrence and input pathways of char and soot in the GOT

Although BC is produced by incomplete combustion of fossil fuels and vegetation (Gustafsson and Gschwend, 1998; Schmidt and Noack, 2000), it is not a well-defined constituent, but a collective term, which refers to a combustion continuum and can be further divided into two subtypes of BC (i.e., char and soot) based on the temperature control in the analytical procedures (Masiello, 2004; Han et al., 2007b, 2015a). In this study, char and soot were measured using the IMPROVE thermal optical method according to Han et al. (2007b), which has been validated and successfully applied in aerosol studies (Han et al., 2010; Lim et al., 2012) and in soil and sediment studies (Han et al., 2009, 2011, 2015a, 2015b; Cong et al., 2013), in order to yield a better understanding of the distribution, sources and potential input pathways of char and soot within the tropical coastal system in SE Asia.

As shown in Fig. 5, the spatial variations of char and soot and their ratio values (char/soot) in the study area were presented and compared. The char/soot varied from 3.11 to 24.75, with a mean value of 9.57 ± 5.02, indicating that char was a predominating constituent of BC in the GOT sediment samples. Char is produced mainly in a low-temperature combustion and has a large particle size (generally 1–100 μm), whereas the soot is mainly composed of submicrometer particles formed from the condensation of hydrocarbon radicals at high temperature combustion (>600 °C) (Masiello, 2004; Han et al., 2011). Therefore, char is usually more related to the local biomass burning and industrial activities and is more easily incorporated into local soils and discharged into the adjacent aquatic sediments through rivers and surface run-off (Knicker et al., 2008). In the present study, char showed a more heterogeneous spatial pattern with several extreme high concentration values occurred in the nearshore area along the GOT (Fig. 5a); whereas soot concentrations and the corresponding char/soot ratios showed offshore decreased pattern (Fig. 5b and c).

Furthermore, the relationship between char and soot was in general less correlated in the upper Gulf with corresponding higher slopes (i.e., char/soot ratios) as compared to the lower Gulf (Fig. 6), which could be partially explained by a decoupling of the char and soot sources as noted above. Biomass burning and forest fires have been suggested to serve as the predominant sources for BC in aerosol samples in Thailand, especially in the hot season (Sahu et al., 2011), and the larger particles of char mainly from the biomass burning residues tend to first accumulate in soils in close proximity to their site of production and then be primarily transported by surface run-off and local river inputs to the aquatic sediments (Gélinas et al., 2001; Yunker et al., 2002; Pietzsch et al., 2010; Han et al., 2015a). Actually, in addition to the sediment-associated BC, Ding et al. (2015) also demonstrated that there existed different input sources for the dissolved BC in the boreal forest streams through active removal process in soils from the local biomass burning as compared to those in the remote glacial rivers with a more atmospheric delivered fossil fuel-derived BC. Above all, the occurrence of the composition of sedimentary BC in the GOT in this work suggested that char as the predominant constituents of sedimentary BC both in the upper and lower Gulf could be mainly derived from biomass burning and it entered into the nearshore region mainly through fluvial transport and surface run-off from fire-impacted soils, as compared to the fossil fuel combustion-derived soot with a more ubiquitous atmospheric loading, especially in the lower Gulf. This point could also be supported by the high abundance of perylone in the coastal GOT that was originated from the direct terrestrial soil-derived material contribution (Boonyatumamond et al., 2006).

3.4. Mass inventory of BC in the GOT sediments and its comparison with other coastal regions around the world

The BC burial flux over the Got was estimated based on the measured BC abundances and reported mass sediment accumulation rates (Fig. 7a; from Srisuksawad et al., 1997). The spatial distribution of the BC burial flux showed an apparent regional viability (Fig. 7b), ranging from 15 to 980 μg/cm²/y with a mean of 212 ± 228 μg/cm²·y. The highest BC burial flux was found in the upper Gulf with the average of 336 μg/cm²·y, significantly higher than that of the lower Gulf (144 μg/cm²·y). In addition to the upper Gulf, the other two higher BC burial fluxes (>300 μg/cm²·y) were concurrent within the finer deposits in the lower Gulf, indicating a potential depository for the land-based exported BC. The estimated BC burial fluxes in this study were comparable with those of the coastal regimes around the world (Table 1), such as the Gulf of Cádiz (120 μg/cm²·y) and the Gulf of Maine (131 μg/cm²·y) (Gustafsson and Gschwend, 1998; Sánchez-García et al., 2013), but they were lower than those areas with strong human impact, such as the Swedish Continental Shelf (1103 μg/cm²·y) (Sánchez-García et al., 2012). In contrast, this result was still 2–4 orders higher than those from the pelagic and deep sea sediment due to the relatively lower sedimentation rates, such as the Pacific deep sea (0.01–0.1 μg/cm²·y) (Smith et al., 1973) and the South Atlantic Ocean (2.27 μg/cm²·y) (Lohmann et al., 2009).

Based on the estimated BC burial flux and accumulation areas of each of the divided homogenous sectors, the total mass inventory of depositional BC ~200 Gg/y can be finally obtained for the ~150,000 km² of the study area. Considering the areal extension in the current work represents only half of the entire GOT (~320,000 km² in total), the estimated BC mass inventory in the GOT was on the same order of magnitude as other oceanic margins in the world, such as the New England continental shelf off northeastern USA (400–800 Gg/y, ~380,000 km²), the total South Atlantic Ocean (480–700 Gg/y, deep sea sediments) and the Swedish Continental Shelf (~300 Gg/y, ~155,342 km²). For the sub-regions, the estimated sedimentary BC mass inventory for the upper Gulf (~7000 km²) was ~15 Gg/y. The total suspended solids (TSS) delivered into the upper Gulf by the major four rivers have been estimated at 6.32 million tons per year (Srisuksawad et al., 1997); therefore, it could result in a riverine BC flux delivery of ~12 Gg/y into the upper Gulf (if BC constituted 0.2% of TSS). This result partially supports that the riverine export is a predominant pathway for BC input in the coastal upper Gulf. For the lower Gulf, in spite of the influx of fluvial TSS was much less due to the relatively low water discharge (Simpson and Snidvongs, 1998), the significant land-based BC input through the extensive surface runoff could also be important, especially for the nearshore regions with obvious higher values of BC/TOC% and char/soot ratios as noted above. Besides, the atmospheric delivery of combustion-derived soot-BC could also serve an important role for the buried BC in the open GOT.

A major problem in quantifying BC is that it is not a well-defined chemical component, but rather exists as a complex concept of combustion/temperature continuum, with widely varying physical and chemical characteristics (Hedges et al., 2000; Masiello, 2004; Hammes et al., 2007; Conedera et al., 2009; Wiedemeier et al., 2015). Therefore, it is worth noting that the direct comparison as noted above was complicated considering the used different BC quantification methods with regard to the BC sources and sinks. Anyway, based on the same BC measurement procedures, the mass inventory of BC in the GOT in the present study was comparable
with our recent work in the Bohai Sea (~147 Gg/y) and the Yellow Sea (~171 Gg/y) off the western Pacific margins (Fang et al., 2015). The semi-enclosed Bohai Sea and Yellow Sea, from the moderate-high latitude region, are mainly situated at the downwind area of the East Asian Monsoon, receiving large amounts of combustion-derived substances (e.g., PAHs) through atmospheric deposition, especially in the winter season due to the extensive regional heating activities in China (Wang et al., 2014). The surrounding region of the GOT from the tropical regime is more subjected to the extensive biomass burning, especially during the hot season (Sahu et al., 2011). Although there is no extensive coal consumption around the GOT, such as that in North China, the inherent biomass burning in tropical SE Asia is also significant (Bond et al., 2004), and atmospheric deposition is expected to be an important BC vector for the Indian Ocean (Lohmann et al., 2009). Therefore, the atmospheric delivery of BC could play an important role in the regional BC budget in the open GOT, and the shelf sediments could serve as a significant depository of BC in the large-scale fate within SE Asia.

4. Conclusions

BC concentration in the GOT ranged from 0.07 to 3.99 mg/g and was comparable with those in other shelf settings around the world. Spatial variability of BC closely corresponded to sediment grain size and TOC content, suggesting a general hydrodynamic forcing on the BC occurrence in the open GOT in addition to the source-dependent control in the coastal region. Positive correlation between BC and TOC in the upper Gulf could be more related to their similar land-based input pathways; while this co-variation in the lower Gulf should be more explained by their coupled sedimentary dynamics. Char as the predominant constituents of sedimentary BC both in the upper and lower Gulf samples could be mainly derived from biomass burning and entered into the nearshore through fluvial transport and surface run-off, while the lower Gulf had a more significant input of soot compared to the upper Gulf. The estimated BC mass inventory in the GOT was ~200 Gg/y and comparable with other shelf regimes, implying that the coastal margins in tropical SE Asia can serve as a significant reservoir for the exported BC from the adjacent landmass.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.ecss.2015.12.019.

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