

Chapter 9

BLACK CARBON IN COASTAL AND LARGE RIVER SYSTEMS

Siddhartha Mitra¹, Andrew R. Zimmerman², Glendon B. Hunsinger³, William R. Woerner⁴

¹Department of Geological Sciences, East Carolina University, Greenville, NC 27858

²Department of Geological Sciences, University of Florida, Gainesville, FL 32611

³Department of Geology and Geophysics, Yale University, New Haven, CT 06520

⁴Department of Geosciences, Stony Brook University, Stony Brook, NY 11794-2100

FINAL DRAFT. Cite as follows: Mitra, S., Zimmerman, A.R., Hunsinger, G.B.,
Woerner, W.R. In Press - 2013. Black Carbon in Coastal and Large River Systems, in
Biogeochemical Dynamics at Major River-Coastal Interfaces: Linkages with Global
Change, eds. Bianchi T.S., Allison, M.A., Cai.W.J. Oxford Publishing Company

9.1 INTRODUCTION

Black carbon (BC) is defined as the highly condensed carbonaceous products of organic matter combustion (Goldberg, 1985; Novakov, 1984). It can be formed either via the condensation of combustion gases (i.e. soot) or as the residues of incomplete combustion of any type of biomass. Black carbon in the environment has recently received increasing attention with the growing awareness of its potential importance to the short and particularly the long term C cycle, and accompanying climate feedbacks. While BC in the terrestrial environment has been extensively investigated, BC transport to and storage in aquatic environments has received far less attention. In this chapter, the evolution of BC is discussed, from its creation in watersheds to its fate in the oceans, with specific focus on its crossing of the terrestrial-aquatic interface.

The two major possible sources of BC to rivers or the ocean are from fossil fuel and biomass burning. Subsequent to its generation, BC may be transported to the ocean either by atmospheric deposition or after incorporation into soils and sediments, followed by its erosion and transport via fluvial systems (Fig 9-1). While these are important, examination and quantification of fluvial BC in large rivers is likely an important route for BC transfer to the ocean, deserving further research attention for several reasons. First, despite the importance of BC and its fluvial discharge, there are a limited number of studies addressing BC discharge from the world's major rivers and limited measurements of BC in aquatic environments as a whole. Second, when measured, BC has been shown to comprise a substantial portion of sedimentary organic matter across different river systems and their coastal margins globally (Elmquist et al., 2008b; Lohmann et al., 2009; Masiello and Druffel, 2001; Mitra et al., 2002). Third, the quantity of BC identified in riverine particulate organic carbon (POC) is approximately twice the amount of BC calculated to be buried in ocean sediments (Druffel, 2004), suggesting a missing sink, unidentified transformation rates, or uncertainty in the estimates associated with fluvial and

coastal BC sources or oceanic sinks. Finally, as marine sedimentary rock represents, by far, the largest reservoir of organic matter on Earth (Hedges, 1992), it might follow that it is also largest reservoir for BC on the Earth. However, unlike other OM, BC is only created in the terrestrial sphere. Thus, quantifying the efficiency of the terrestrial-to-marine transfer of BC is particularly important towards understanding how biomass combustion is related to climate change and the C cycle. The burial of BC is particularly important to understanding long term climate variation because, as a relatively refractory pool of carbon (e.g. Masiello 2004) , the formation of BC from biomass combustion, followed by its deposition and burial represents a transfer of carbon from the short to the long-term carbon cycling pools (i.e. long term sequestration - Fig 9-2).

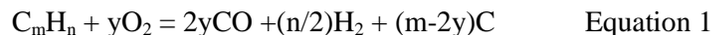
Different aspects of the biogeochemical cycling of BC have been reviewed previously (e.g., Schmidt and Noack, 2000; Forbes et al., 2006; Preston and Schmidt, 2006). The objective of this chapter is to provide a review of studies of BC sources, fluxes and reservoirs with emphasis on its fluvial transport and discharge into the ocean. It will be shown that, to the extent that can be determined, in many ways, the fluvial discharge of BC and its burial in the ocean is similar to that of terrestrial organic matter, which is discussed in other chapters of this book (Chapter 4, Chapter 8), whereas in other respects, its cycling is quite distinct. Indeed, the result of the comparisons made in this chapter suggest that unlike bulk terrigenous organic matter, the sources and sinks of refractory BC, irrespective of isolation method used, may in fact be in balance.

It is now widely recognized that atmospheric BC is an important driver of climate, only second or third in importance behind CO₂ in the warming of the atmosphere (Forster and Raga, 2007) and unlike CO₂ and methane, the reduction of BC emission to the atmosphere may be an easier goal to implement in efforts toward climate change mitigation (Jacobson, 2001). Under future climate change scenarios, the threat of wildfires will increase (IPCC, 2007) and, as society's dependence on fossil fuels continues to grow, concomitant increases in fossil fuel and aerosol BC emissions are expected. In addition, individuals and companies may soon

intentionally produce BC (biochar) from waste biomass in large quantities in efforts to sequester C and improve soil fertility (e.g. Lehmann, 2007). The fate of this increasing BC supply is not known, so knowledge of riverine fluxes and aquatic BC reservoir processes are needed now more than ever.

9.2 SOURCES OF BLACK CARBON

The main sources of BC to the environment are combustion of fossil fuels (coal, oil, gasoline, diesel) and biomass combustion. Except in the case of natural fires, these are driven largely by society's present and historical energy needs. Natural gas combustion is not a significant contributor to BC formation (Goldberg, 1985) thus we will not consider it further in this chapter. Complete combustion of these organic materials results in the production of CO₂, an important greenhouse gas. However, the efficiency of combustion is dependent on several variables including combustion time, temperature, type of fuel, but primarily the presence or absence of oxygen. A sub-stoichiometric reaction of organic matter with oxygen, or what is known as pyrolysis, results in the formation of particulate, carbon rich, and O-H-S-N-poor compounds (Stanmore et al., 2001). For example, in the following equation,



BC will be formed when m exceeds the value of 2y (Goldberg, 1985). It should be noted that kinetics of the reaction will often determine the ratio of m:2y.

Several terms have been used to identify the residuals (or subsets of the residuals) of incomplete combustion of organic matter including char, biochar, charcoal, carbonaceous spherules, soot, fusain, elemental carbon, carbon black, oxidation resistant elemental carbon, and graphitic carbon. It may be best to conceptualize these types of BC as existing along a continuum

of combustion, with the abundance of each of these products dependent on combustion conditions (Fig 9-3) (Hammes et al., 2007; Masiello, 2004). On the left side of this figure are the residues remaining from incomplete combustion of biomass including slightly charred biomass and charcoal; these constitute the larger size class of pyrogenic BC. In contrast, the smallest size class, soot and graphitized C, is a mix of amorphous microscopic particles that contain BC, organic carbon, and smaller amounts of sulfur and other chemicals. During combustion, residual char and charcoal may form simultaneously with soot, depending upon fuel composition and efficiency of combustion. For example, BC from diesel combustion has a higher proportion of soot compared to other incomplete combustion products. In contrast, smoke plumes from wildfires may appear brown, blue, or gray due to the reduced combustion efficiency of wildfires, which produce relatively greater amounts of organic carbon (sometimes called 'brown carbon') and other pyrolysis aerosols in comparison to soot (Bachmann, 2009).

Laboratory charring experiments of individual components of wood (e.g. cellulose, lignin, pectin) as well as bulk wood and grass, offer some clues to understanding the controls on and variations in BC composition. In general, initial heating of organic matter results in dewatering and dehydration to phenols and aromatics, which is then followed by additional chemical transformations that occur at higher temperatures and longer duration of heating. These in turn lead to formation of more condensed refractory organic matter. For example, Czimczik et al (2002) showed that at temperatures below 500 degrees C, BC formed via smoldering hardwood or softwood yielded small clusters of condensed C, lacking a high proportion of graphitic characteristics. Thus, levels of soot-type BC may be low in ecosystems dominated by low-temperature smoldering fires especially when quantified by isolation methods that rely detection of condensed aromatic structures (Czimczik and Schmidt, 2002).

At formation temperatures of 450 °C and beyond, progressive depolymerization, loss of functional groups, aromatization, dehydrogenation reactions and removal of substituents occurs,

resulting in larger sheets of fused aromatic rings (Arranz et al., 2009; Baldock and Smernik, 2002; Keiluweit et al., 2010; Knicker, 2007; Zimmerman and Gao, In press). These condensed structures would likely be very resistant to biotic or abiotic degradation, as would the graphitic ‘turbostratic crystallites’ that have been observed to form at temperatures >600 °C and in soot (Keiluweit et al., 2010; Kercher and Nagle, 2003). In contrast to charcoal, which is irregularly shaped and larger, the basic units of soot are spherical or nearly spherical particles with diameters often in the range 20 -30 nanometers, corresponding to about one million carbon atoms (Haynes and Wagner, 1981). Soot may be formed from the nucleation of gaseous carbonaceous species, such as acetylene and polycyclic aromatic hydrocarbons (PAHs) around uncharged radicals, which occur during intermolecular collisions. Subsequently, condensation reactions of gas phase species lead to the appearance of the first recognizable soot particles during the cooling process (Haynes and Wagner, 1981). For some solid fuels, soot is also formed from the pyrolysis of tarry materials ejected during devolatilization. These gases then subsequently dehydrogenate and partially oxidize into soot spherules (Stanmore et al., 2001).

9.2.1 Atmospheric BC

The atmosphere is an important source of BC influx to rivers and the ocean. Moreover, the presence of BC aerosols in the atmosphere has been shown to affect hydrological cycles. For example, the radiative forcing of most aerosols is negative (i.e. leading to cooling of the Earth’s atmosphere with increased concentrations) (Penner, 2001) but BC aerosol’s radiative forcing is positive as BC absorbs sunlight and warms the atmosphere and may enhance evaporation (Hansen et al., 2000; Jacobson, 2001). These radiative effects have been shown to alter regional atmospheric stability and vertical circulation, while affecting large scale circulation and hydrologic cycles, causing significant regional climate effects such as enhanced regional precipitation (Menon et al., 2002).

The aeolian transport of BC aerosols and their deposition on snow and ice are intimately linked via a positive feedback loop in the climate-carbon cycle. The long-range transport of BC aerosols to snow and ice in remote areas reduces albedo (Warren and Wiscombe, 1979; Warren and Wiscombe, 1981; Warren and Wiscombe, 1985) resulting in increased ice melt in polar and boreal areas (Barry, 1996; Clarke and Noone, 1985; Robock, 1984). For example, a concentration of 15 mg kg^{-1} of BC in snow reduced its albedo by 1% (Warren and Wiscombe, 1980). In some cases, the BC remaining at the surface of glacier snow increased melting by 50% over areas of ice which were not impacted by BC deposition (Conway et al., 1996). Such BC-driven snow melting has been shown to contribute to historical rapid Tibetan glacier retreat, suggesting that BC deposition may significantly affect some of the world's largest freshwater supplies as well as sea level globally (Hansen and Nazarenko, 2004; Xu et al., 2009). Furthermore, some recent studies have alluded to anthropogenic aerosols, rich in fossil carbon, contributing to much of the ancient organic matter in glacial runoff (Stubbins et al., 2012a). As a result, increased BC emission may ultimately enhance melting of ice from polar and boreal areas, leading to increased fluvial transport of BC into the oceans.

There are several factors that affect the concentration of BC aerosols in the atmosphere and their subsequent deposition. Geographically, modern day emissions of BC are highest in the tropics and East Asia (Ramanathan and Carmichael, 2008) suggesting that watersheds and rivers in this area may be enriched in BC. Also, aerosol BC is found above both land and ocean, with concentrations being highest near urbanized areas and lowest above remote areas (Clarke et al., 1984). Present day estimates of annual global BC emission range from 6 to 24 Teragrams ($1 \text{ Tg} = 10^{12} \text{ g}$) BC from fossil fuel emission (Novakov et al., 2003; Penner and Eddleman, 1993) and 50 to 260 Tg BC from biomass burning (Fernandes et al., 2007; Kuhlbusch and Crutzen, 1995), suggesting both processes contribute to aerosol BC loading. However, another recent estimate of

annual fossil fuel and biomass BC emission together ($5 - 20 \text{ Tg y}^{-1}$), suggests that there is still some uncertainty in the scientific community as to how much BC is emitted into the atmosphere.

The residence time of BC aerosols in the atmosphere is determined by initial size distribution, concentration of ambient particles, frequency and duration of precipitation, and wind conditions (Ogren and Charlson, 1983; Ogren et al., 1984). The largest BC particles are not likely to become or stay airborne and as such, will deposit proximally to the source of combustion (Clark, 1988). The majority of aerosol BC is found in the smaller $0.01 \mu\text{m}$ to $1.0 \mu\text{m}$ size fraction (Clarke et al., 2006) and will have atmosphere residence times of days to weeks (Clark, 1988) (Fig. 9-3). Despite its importance locally and regionally, dry deposition does not contribute very much to the total atmospheric deposition of aerosol organic matter (Bidleman, 1988). Additional studies at the regional level have shown that wet deposition accounted for 52 – 99% of aerosol BC deposition and thus is considered to be the most important deposition mechanism by which atmospheric BC enters the ocean (Suman et al., 1997). The most comprehensive estimate for atmospheric BC deposition to the ocean to date, suggests that dry and wet deposition of BC to the ocean are respectively, 2 and 10 Tg C y^{-1} (Jurado et al., 2008). Below, we address how this estimate compares to other BC sources into the ocean, namely that derived through soil erosion and fluvial influx.

9.2.2 Soil BC

Given that humans have been burning biomass since $\sim 9 \text{ kya}$ (Carcaillet et al., 2002), the widespread occurrence of natural fires (when unsuppressed), and the emission products associated with both processes, the occurrence of BC in nearly all soils is not surprising. Although there is still uncertainty over the most appropriate method to measure BC (see later section), calculation of pyrogenic OM contribution to soil organic carbon (SOC) using solid state ^{13}C nuclear magnetic resonance spectroscopy, acid potassium dichromate digestion, and

benzenepolycarboxylic acid concentrations, are becoming most common. Using these methods of determination, concentration of BC in most soils generally ranges 0.2 – 5 weight % of soil and represents between 2 and 15% of the soil TOC with values of about 10% SOC most common (e.g. Bird et al., 1999; Cusack et al., 2012; Hammes et al., 2007). Soils thought to have anthropic inputs of pyrogenic OM such as Amazonian *terra preta* and higher latitude grassland OM-rich soils such as Chernozems and Mollisols have been found to have BC contents as great as 45% SOC (Chasar et al., 2000; e.g. Czimczik et al., 2002; Forster et al., 2007; Forster and Raga, 2007; e.g. Schmidt et al., 2002).

Much as for soil OM as a whole, there is considerable debate as to the primary controls on soil BC concentrations. As with soil OM, the possible controls can be reduced to either input or preservation-related mechanisms. While some studies have found soil BC stocks to be related to fire frequency or intensity, other studies have failed to find such relationships. For example, the BC content of an Andean volcanic agricultural soil that had yearly burns of wheat and barley residues was not different from one that had experienced no burning (Rivas et al., 2012). In contrast, a nearby forest with no recent burning had significantly less BC content (0.5% SOC) than the topsoil of a site affected by wildfire 3 years previously (7% SOC, Drivas, 1982). In studies of Siberian Scots pine forest soils, increases in BC of up to 40% were found in some soils and no lasting effect on others (Czimczik et al., 2003; Czimczik et al., 2005).

Despite all these uncertainties, wildfires have been estimated to produce approximately 0.04 to 0.2 Pg soil BC yr⁻¹ (Preston and Schmidt, 2006). Given these global estimates, as well as those for specific areas, workers have concluded that charcoal soil stocks should be several times greater than actually observed (Baldock et al., 2004; Ohlson and Tryterud, 2000). Part of the difficulty in linking BC production to BC soil stocks is that BC may not be as fully inert as was previously assumed. The observation of strong correlations between SOC and BC levels (Cusack et al., 2012; Hammes et al., 2007)) suggests that similar processes may control their distribution

and perhaps export, including degradation, translocation within soil, dissolution and erosion. Both BC degradation and dissolution (soil as a dissolved BC source) are discussed further below.

While horizontal and vertical movement of particulate soil BC by erosion and downward translocation, respectively, are infrequently quantified, there is a growing body of indirect evidence that these processes represent significant losses of soil BC at short timescales. First is the very presence of BC in aquatic environments despite its exclusive production on land. Second, studies examining soil profiles in which the history of fire incidence is known have calculated wildly varying loss rates leading many to conclude that process other than BC degradation, such as erosion, may play a dominant role (e.g. Nguyen et al., 2008; Zimmerman and Gao, In press). Finally, a few studies have attempted to follow all BC losses from a BC-amended soil in the field. Working in Colombia over two years, only a small portion of the BC losses could be accounted for leading the researchers to conclude that 20-53% of the applied BC must have been lost by surface erosion (Major et al., 2010). Another study found that 7–55% of pyrogenic C added to the soil via a surficial grass fire was lost to erosion and another 23–46% was moved vertically to greater soil depths (Rumpel et al., 2009). In addition, observations of low BC concentrations in upper layers of soils subject to frequent burning, but increased BC with depth, suggests mobility of BC within the soil profile (Dai et al., 2005; Dick et al., 2005; Knicker et al., 2012).

Perhaps it is not surprising that BC should make its way from soils into rivers given that natural fires often occur on unstable slopes and may denude the landscape of erosion-detering vegetation. The passage of fire across soil can also lead to an increase in soil hydrophobicity and a concomitant decrease in water infiltration capacity, the net result of which is increased surface runoff and erosion, which are magnified in proportion to the intensity of the fire (Conedera et al., 2009). In addition, BC particles may be preferentially exported compared to other soil OM due to their predominantly small size and low density (Hammes et al., 2007; Skjemstad et al., 1999).

Exported BC was found to be most closely associated with the 20–50 μm soil fraction (Rumpel et al., 2009). Certainly, BC export from soil, versus storage or loss via other mechanisms will vary with such factors as climate, topography, soil type and hydrological regime.

9.2.3 Dissolved BC

Despite the fact that fluvial BC has been typically regarded as particles found in suspended sediments or bedload, the recently proposed presence of BC in the dissolved phase is certainly reasonable. The definition of particulate versus dissolved classes of organic matter is operational, with POC often defined as organic material retained on filters with nominal size cutoffs (e.g. 0.45, 0.7 or 1 μm). Thus, it is conceivable that microparticulate BC escapes the analytical window of what is defined as particulate material. Furthermore, material which passes through such filters may remain suspended in the water column for substantial periods of time, leading to the observed age offset between POC and BC (Abiven et al., 2011; Masiello and Druffel, 1998). Also, desorption from soot BC aerosols at room temperature has yielded condensed aromatic molecules in the water soluble fraction while oxidation of particulate BC has been shown to cause it to become more water soluble (Chughtai et al., 1991; Decesari et al., 2002; Wozniak et al., 2008). Thus, leaching of carbon from the particulate to the dissolved phase, aided by chemical oxidation or biodegradation, may be significant.

Researchers have claimed to identify substantial amounts of pyrogenic substances in the dissolved phase of organic matter in rainwater, rivers, and the ocean (Dittmar, 2008; Hockaday et al., 2006; Kim et al., 2004; Lee, 2005; Mannino and Harvey, 2004). Some of the recent studies that have detected the presence of BC in the sub-particulate phase are summarized in Table 9-1. What is clear from the comparison of these studies is that dissolved BC may comprise trace amounts up to 7 % of the total DOC in an aquatic system. The studies in Table 9-3 suggest that a median value of ~ 5% is appropriate for the fraction of total DOC that is BC. Since DOC in the

oceans is estimated at 7×10^5 Tg C (Hansell and Carlson, 2001), the ocean's reservoir of dissolved BC can be approximated at 4×10^4 Tg. Using a value of $35,000 \text{ km}^3 \text{ y}^{-1}$ for the water discharged from all rivers to the ocean (Hetland and Hsu, Chapter 3), with a median concentration of all dissolved BC concentrations listed in Table 9-1 ($10 \mu\text{g L}^{-1}$), an estimate for the residence time of dissolved BC is calculated to be $\sim 100,000$ y. Since the median age of bulk DOC has been measured to be far younger (~ 6000 y; Williams and Druffel, 1987), our simple calculations suggest that dissolved BC is certainly old enough to bias the age of bulk DOC. That said, several studies have found dissolved BC to be rapidly photooxidized (Stubbins et al., 2012b; Stubbins et al., 2010). It is likely that, as with particulate BC, dissolved BC is composed of materials with a continuum of compositions and labilities. Clearly, additional investigations into the chemical composition, age, and environmental fate of dissolved BC in the ocean, are warranted.

9.3 ISOLATION, QUANTIFICATION, AND SOURCE DISCRIMINATION METHODS FOR BLACK CARBON

There are as many methods to isolate BC as there are BC practitioners. Furthermore, each method of BC isolation is often uniquely suited to the interests of the researcher. For example, since the atmospheric community is concerned about BC's radiative properties, it often quantifies BC by measurement of light transmittance or reflectance on filters (Bond, 2001; Currie et al., 2002; Horvath, 1993; Lioussse et al., 1993; Martins et al., 1998a; Martins et al., 1998b; Moosmuller et al., 2009; Novakov, 1984; Rosen and Novakov, 1977; Smith et al., 1975). But, isolation, quantification, and source discrimination methods for aquatic and sediment-associated BC fall into categories of: visual/optical, oxidation-based (chemical and thermal), spectroscopic approaches, molecular proxy-based, and some combination of these. Specifically, quantification of BC after its isolation involves either direct optical methods (e.g. counting) or quantification of C by high temperature oxidation in the presence of a metal catalyst to CO_2 . These BC isolation

methods are briefly discussed below. The reader is also referred to several additional methodological review studies for further information (Currie et al., 2002; Hammes et al., 2007; Nguyen et al., 2004; Schmidt and Noack, 2000).

The methods used to isolate BC in aquatic systems can result in large differences in the amount of BC quantified. Furthermore, each method for BC isolation has its inherent limitations based on the fact that each BC isolation method may be optimized for a specific portion of the BC continuum (Fig. 9-3). Also, there is no measure of accuracy for BC analysis. Methodological limitations can be attributed to 1) a lack of a common definition for BC, 2) failure to detect a broad range of pyrogenic carbonaceous residue, and 3) false-positive identification of non-BC carbonaceous substances such as soil geopolymers, kerogen, or other petrogenic carbon.

9.3.1 Visual methods for BC quantification

Paleoecological studies have benefited tremendously from quantification of the historical occurrence of fires made by enumeration and quantification on a per unit area basis of macroscopic or microscopic charcoal in the sedimentary record. Several recent studies that have comprehensively quantified historical charcoal deposition globally, have suggested that both geographical and climatological differences contribute to BC loading (Bowman et al., 2009; Carcaillet et al., 2002; Marlon et al., 2009a; Marlon et al., 2009b; Power et al., 2008). Although light microscopy can identify larger particles that are clearly charcoal (i.e. jet black, opaque, angular) some particles can be challenging to identify due to their ambiguous color and/or shape (Patterson et al., 1987). Distinguishing between carbonized plant tissue and much smaller carbonaceous residue of combustion of fossil fuels and other sources is controversial, though important to climate studies. For example, using scanning electron microscopy, Kralovec et al., (2002) propose that larger BC particles (> 20 μm) isolated downcore in Lake Erie sediments, were either 1) porous and spheroidal (oil-derived), 2) elongate with length-to-width ratios greater

than three (wood-derived), or 3) irregular with a length to width ratio between 1 and 2 (coal-derived). An attempt to independently validate the BC isolation methods in that study using historical energy consumption data was only partially successful (Kralovec et al., 2002) suggesting that this approach may be limited in its application.

9.3.2 Chemical and thermal BC quantification methods

Quantification of BC in natural soils and sediments requires the ability to distinguish between inorganic carbon, non-pyrogenic OC and pyrogenic carbon. Chemical and thermal oxidation methods entail the removal of the former two, followed by quantification of residual carbon abundance. Prevalent chemical and thermal methods and their variables currently in use are noted in Table 9-1 of Nguyen et al., (2004). Subtle differences such as isolation temperature (Elmquist et al., 2006; Elmquist et al., 2004) or chemical oxidant reaction time (Masiello et al., 2002) can lead to substantial differences in BC yield for the same natural sample or reference material (Elmquist et al., 2006; Elmquist et al., 2004; Masiello et al., 2002). Thus, it is critical that any BC isolation/quantification method be applied carefully and precisely. To that end, it is advisable to incorporate well-characterized available standard reference materials into any method of BC isolation such as those used in the BC Ring Trial (Hammes et al., 2007).

The thermal and chemical oxidation methods used most often for analysis of sedimentary BC are: 1) acid-dichromate oxidation (AD) and 2) Chemo-Thermal Oxidation at 375 degrees Celsius (CTO-375). The AD technique involves thorough demineralization of a sample via hydrochloric acid (HCl) and hydrofluoric acid (HF), followed by a lengthy potassium dichromate (KCr_2O_7) and sulfuric acid oxidation of the demineralized residue (Wolbach and Anders, 1989). The technique is based on the kinetics of the reduction reaction of an acidic solution of dichromate in conjunction with that solution's simultaneous oxidation of reduced organic matter. The dichromate in acid oxidizes labile and refractory carbon ranging from fresh biomass to

kerogen and even labile BC, depending on the duration of oxidation time (Wolbach and Anders, 1989). It is also assumed that each fraction of OM present in the sample responds differently to the oxidant, which is available in excess. The oxidation of carbon in the residue during the etching procedure is then modeled as a first order exponential loss process, with a “fast” rate constant corresponding to labile non-BC, intermediate rate constants corresponding to kerogens, and a “slow” rate constant corresponding to BC (e.g. elemental carbon) (Wolbach and Anders, 1989) with 400h being an adequate representation of the onset of the slower rate constant (Masiello, 1999). It is commonly and incorrectly assumed that this approach can be approximated as 'BC is what is left.' However, that is not the case as application of this technique relies on knowledge of the chemical kinetics of sample oxidation (Masiello et al., 2002). One criticism of wet-chemical methods is that they may underestimate BC as a result of losses during decantation or other transfers of the samples (Elmqvist et al., 2006; Elmqvist et al., 2004). However, other studies have not observed a difference in BC content of Standard Reference Material 1649a (urban dust) isolated by both the CTO-375 and the AD method (Mitra et al., 2009), suggesting that the BC losses during the washing and transfer steps can be minimized with careful handling. While this method has been used to isolate fluvial BC in numerous studies (see Table 9-2), the toxicity of the reagents and the lengthy duration of the oxidation process, are somewhat disadvantageous.

The CTO-375 technique for isolating BC has probably been the technique most widely applied to soils and sediments, due to its ease of use. This method originally consisted of thermal oxidation of OC in the presence of excess air and was conducted at 375 °C followed by release of carbonates via in situ acidification in 1M HCl (Gustafsson et al., 1997). The residual sedimentary carbon is then quantified by CHN elemental analysis (Gustafsson et al., 1997). Later testing of this technique indicated that this method only quantified soot BC (Nguyen et al., 2004). Several modifications have been made to this technique over the years. For example the temperature as

well as the point at which acidification should be done, have both been modified since the initial study (Elmquist et al., 2006; Elmquist et al., 2004). One problem with thermal techniques is that they can yield false-positives and over-estimation of pyrogenic OM via charring. For example, nitrogen-rich macromolecules seem to be charred during thermal oxidation (Accardi-Dey, 2003; Flores-Cervantes, 2008) leading to quantities of soot BC in coastal planktonic organic matter isolates (Accardi-Dey, 2003; Gelinas et al., 2001), bovine serum albumin (Flores-Cervantes, 2008) and melanoidin (Hammes et al., 2007). It also appears that pollen is refractory to chemical and thermal treatment associated with the CTO-375 (Gustafsson et al., 1997). Thus, the reader is advised to carefully review the evolution of this procedure before proceeding and in particular, evaluate its limitations alongside its simplicity.

Recent advances in spectroscopic methods for identification of BC, which are based on molecular structure such as aromaticity, have yielded intriguing results about chemically and thermally-isolated pools of BC. Spectroscopic methods such as ^{13}C nuclear magnetic resonance (NMR), X-ray absorption near-edge structure (XANES), and near-edge X-ray absorption fine structure (NEXAFS) spectroscopic methods have been used to demonstrate the presence of aromatic, graphitic, as well as amorphous moieties in chemically and thermally-isolated BC (Dickens et al., 2006; Haberstroh et al., 2006; Lehmann et al., 2005). This suggests that material that is chemically or thermally isolated as BC may in fact be extremely heterogeneous in functional groups and composition.

9.3.3 Isolation of BC in the dissolved phase

Researchers have used a variety of techniques to isolate and quantify BC in the dissolved or colloidal phase in aquatic systems. For example, Mannino and Harvey (2004) quantified BC in high molecular weight dissolved organic matter from the Delaware Bay using the CTO-375 method. However, quantities of BC in algal exudates, as well as the modern radiocarbon ages of

the BC isolate (Mannino, 2000) suggest that their results were influenced by some charring of carbon during their sample processing. Indeed, the level of charring in the blanks used in that study created BC that was similar to the concentrations of BC estimated to exist in the open ocean (Mannino and Harvey, 2004). As noted above, selective charring of nitrogen-rich organic matter has been observed to occur via the CTO-375 technique (Accardi-Dey, 2003; Gelinas et al., 2001). Thus, application of thermal oxidation may not be appropriate for isolating BC in coastal or oceanic DOM containing nitrogenous organic matter.

Evidence of BC in the dissolved phase is determined by the presence of condensed polyaromatic moieties in water, which has been revolutionized by Fourier Transform Ion Cyclotron Mass Spectrometry (FT ICR-MS) (Kim et al., 2004; Hockaday et al., 2006; Koch and Dittmar, 2006; Hockaday et al., 2007). This technique, although not quantitative, offers extremely high mass resolving capacity of DOM (~200,000 mass units). In order to quantify dissolved BC, several researchers have relied on the fact that dissolved and un-substituted PAHs, can be oxidized in the lab with nitric acid into various benzenepolycarboxylic acids (BPCA). These BPCAs can then be quantified as a proxy for dissolved BC using liquid or gas chromatography (Brodowski et al., 2005; Glaser et al., 1998). However, it should be understood that the similarity of functionalized water-soluble condensed aromatics in natural waters to the degradative products of BC does not necessarily confirm their common origin. For example, functionalized polyaromatics such as those used as proxies for dissolved BC may originate from oil-weathering (Aeppli et al., 2012) or *in situ* thermogenic processes known to occur in hydrothermal seeps in the ocean (Dittmar and Paeng, 2009). Furthermore, both improper laboratory acidification during the procedure and the presence of darker soil pigments have been shown to yield non-pyrogenic BPCA. Thus, the use of BPCAs as a proxy for dissolved BC should also be done with caution.

One novel method of quantifying colloidal BC suspended in the water column involves using flow injection analysis (FIA) coupled to a single particle (SP) inter-cavity laser induced

incandescence photometer (Bisiaux et al., 2011). The FIA-SP2 method determines BC from the wavelength resolved incandescent light emitted by individual BC particles heated to their boiling point (3700 – 4300 K) via laser. As a result, the method is free of positive interferences from humic acids, charcoal, mineral graphite or PAHs, and is insensitive to particle morphology. However, the method is restricted to a particle size range of ~ 60 – 400 nm and thus, larger colloidal BC would be omitted from being quantified.

9.3.4 Source Discrimination of BC

Most BC quantification methods do not provide information about the BC parent material or mode of combustion. Nonetheless, this source information is useful in order to understand BC's role in the global carbon cycle or effect on climate. Stable and radiogenic isotopes of BC as well as molecular proxies of combustion have been used to determine its sources. If minimal isotopic fractionation of the natural isotopes of carbon occurred during 1) combustion of fuel, 2) post combustion preservation, and 3) while carrying out a specific BC isolation technique, stable carbon isotopic signatures ($\delta^{13}\text{C}$) of residual BC might be used to distinguish BC from different sources. However, evidence suggests there is carbon isotopic fractionation of both fossil fuels and biomass, both during combustion as well as diagenesis. For example, the isotopic signature of fossil fuels is a function of their physical state (i.e. solid, liquid or gas) and fractionation during combustion leads to isotopic depletion during the production of combustion gases, but is generally positive during the process of particle formation (Widory, 2006). Similarly, isotopic shifts during the heating process of biomass resulted in fractionation of C-13 by as much as 2-4 ‰ (Gilhooly et al., 1996; Turekian et al., 1996). Also, up to 8 ‰ depletion in $\delta^{13}\text{C}$ was observed in C4-derived chars from natural burning without a similar isotopic change in chars from wood or C3 grasses, due to the presence of occluded phytoliths in C4 derived chars (Krull et al., 2003). Residual BC may also contain some nitrogen, oxygen, hydrogen, and sulfur, in which case their

isotopic signature may be used fingerprint the source of BC. Since these elements comprise such a minor percentage of its composition, they are often not considered practical in BC source discrimination (Bird and Ascough, 2012).

Radiocarbon (C-14) abundance can be used to elucidate the extent to which BC was produced from the combustion of fossil sources or modern biomass (Gustafsson et al., 2009; Reddy et al., 2002). While living biomass incorporates C-14 from the atmosphere, it is deficient in ancient organic material. This method was applied in a most elegant manner to the infamous “brown cloud” or annual haze of combustion-derived aerosols over the Indian Ocean (Gustafsson et al., 2009). In that study, the C-14 value of aerosol BC was quantitatively used to conclude that 50% of the BC was derived from biomass combustion. Unfortunately, ancient petrogenic and paleosol carbon, which have been found in stream and river waters (Raymond and Bauer, 2001), are also likely to be depleted in radiocarbon. Thus, improper accounting for such recycled carbon may lead to overestimates of carbon sequestration in marine sediments. While physical gravimetric separation conducted prior to oxidation may be effective in separating out some petrogenic material (Dickens et al., 2004), more robust confirmation of BC source is generally necessary as each source of carbon (fossil fuel or biomass) will have different net effects on the global carbon cycle.

Some of the uncertainties associated with chemical and thermal BC isolation methods of aquatic sediments have been overcome by quantifying trace organic chemical molecules co-generated during combustion such as selected polycyclic aromatic hydrocarbons (PAHs) (Gustafsson and Gschwend, 1997; Louchouart et al., 2007; Mitra et al., 2009) and levoglucosan, a byproduct of cellulose combustion (Elias et al., 2001; Kuo et al., 2011). Isomer ratios of high molecular weight and particle reactive PAHs are often used to discriminate between sources of combustion (Yunker et al., 2002), but have also been used to corroborate the abundance and provenance of sedimentary BC (Mitra et al., 2002c; Mitra et al., 2009). Another similar technique

along these lines employs pyrene fluorescence quenching as a proxy for BC concentrations in natural sediments (Flores-Cervantes et al., 2009b). This method relies on the concept that BC has a high sorption affinity for PAHs such as pyrene, and has yielded promising results for its quantification in coastal aquatic ecosystems (Flores-Cervantes et al., 2009). In summary, it is recommended that any attempt at BC quantification should be coupled to multiple methods of its source discrimination.

9.4 BLACK CARBON IN LARGE RIVER SYSTEMS

Despite the importance of the fluvial delivery of BC in the global carbon cycle as shown above, only a few studies have characterized or quantified BC in the suspended river and coastal bed sediments associated with major rivers. The results of those studies are summarized in Table 9-2. Quantification of BC discharge from measurements of BC in riverine suspended sediments (e.g. Masiello and Druffel, 1999; Mitra et al., 2002) are associated with uncertainties in that BC in suspended sediments can vary dramatically within and among river watersheds and likely vary with river size and discharge (Masiello and Druffel, 2001). The amount of BC transported as bedload is also unknown. Additionally, erosion of soil from fire-affected areas can introduce non-BC soil-derived geopolymers which may produce false-positive measurements of BC suspended in the water column. Other researchers have attempted to estimate BC discharge by quantifying BC in the seabed adjacent to major river systems (Elmqvist et al., 2008; Lohmann et al., 2009) but this approach also has inherent limitations. For example, BC may be preferentially transported away from river deltas. And bed sediments adjacent to the mouth of large rivers have been termed “fluidized bed reactors”, which are extremely efficient hot spots for remineralization of carbon (Aller, 1998). As a result, quantities of BC in these areas may be disproportionately low and unrepresentative of global fluvial quantities entering the oceans.

9.4.1 Fluvial BC

Existing fluvial sedimentary BC concentration data were collected and parsed into estimates for BC in riverine suspended sediments as well as bed sediments (Table 9-2). In comparing these numbers a few salient trends are apparent. First, irrespective of the isolation method used or geographic location, BC concentrations per mass of sediment are generally greater in suspended sediments than in bed sediments. Second, the concentration of POC, when quantified concurrently with BC, is significantly higher in suspended sediments than bed sediments. As a result, BC-to-TOC concentration ratios appear similar in all the fluvial systems in Table 9-2. This suggests that non-BC sedimentary organic carbon is transported, sorted or degraded by similar processes as BC across coastal systems. For these reasons, we suggest that the average value of 1.6 mg BC per gram dry weight (gdw) of sediment, integrated across suspended and bed sediments, be used as a convenient representative concentration of fluvial BC entering the oceans. In order to calculate global particulate BC exported from large rivers for which BC has not yet been quantified, the value of 1.6 mg gdw⁻¹ was multiplied by suspended sediment discharged from each large river listed in Table 9-3. A similar approach has been used to calculate global fluvial discharge of sedimentary POC (Burdige, 2005). Estimated values of BC were then compared to known values of BC in a subset of the large river systems listed in Table 9-3, as determined by others. This comparison shows that estimated values of fluvial BC export are similar to values of soot BC for the Congo, the Mississippi, and selected Pan-Arctic rivers (Fig 9-4). Part of the reason that this estimation method agrees well with existing values of soot BC may be that averaging BC/TSS across different methods and across a larger number of rivers may eliminate differences due to methodological artifacts. Alternatively, the comparison may be biased by the fact that most of the values and the 1.6 mg gdw⁻¹ in Table 9-2 are derived from refractory BC, isolated using the AD and CTO-375 procedures.

The estimates for each river's BC discharge noted in Table 9-3 are graphically depicted in Figure 9-5, where the area of each arrow is proportional to the estimated annual BC discharge. The rivers in Table 9-3 are organized in order of decreasing sediment discharge. It is apparent from Figure 9-5 that not only is the Amazon the dominant large river for BC discharge, but the BC discharge of the four major South American rivers (Amazon, Magdalena, Orinoco, and Paraná) nearly equal to the BC discharge of every other major river in the study combined (2.64 vs 2.72 Tg y⁻¹, respectively). Perhaps this is due to the compound effects of 1) residual widespread biomass burning (Dittmar et al., 2012) exacerbated by the lengthy watershed residence time of the basin, coupled with the sheer enormity of that river system's discharge. Nonetheless, the fluvial BC discharged from South America seems to supersede that of any other continent. But, as we note in another section below, the quantities of BC measured in the seabed adjacent to the Amazon River, are surprisingly low.

While quantifying fluvial BC discharge is not a straightforward process, the BC discharge of some large river systems as determined with our simple BC discharge model agree well with the measured values (Fig 9-3). As noted in other chapters of this book, the large rivers listed in Table 9-3 are viewed as some of the most influential ones globally in terms of either water or sediment discharge (e.g. see Chapter 3). Thus, we assume their collective contribution to BC discharge to the ocean to be the most important, globally.

Although collectively, small mountainous high yield river systems (SMR) provide 40-70% of the global sedimentary delivery to the ocean (Leithold et al., 2006; Milliman and Syvitski, 1992), they are not included in the global BC discharge estimates in Table 9-3. This may be justified given that their watersheds are smaller. Nonetheless, we do discuss their potential role in fluvial BC discharge below.

9.4.2 Global pyrogenic PAH discharge and comparison with BC

As noted earlier, PAHs co-evolve with BC during combustion (Lima et al., 2005). During biomass and fuel combustion, the liberation of hydrogen associated with fuel combustion enroute to its formation of BC produces aromatic molecules (Chakraborty and Long, 1967). Several researchers have isolated PAHs concurrently with BC in an attempt to confirm the pyrogenic origin of BC quantified (Flores-Cervantes et al., 2009b; Kuo et al., 2008; Louchouart et al., 2007; Mitra et al., 2002a). Here we attempt to validate our approach to calculating global BC discharge for the river systems listed in Table 9-3 by making a comparison between the discharge of benzo[*a*]pyrene (a high molecular weight particle-reactive PAH) and the estimated values of large river BC discharge depicted in Fig 9-5. The significant correlation between the two values ($r^2 = 0.47$; $p < 0.005$ - Fig 9-6) suggests that the fluvial BC estimated using our approach is pyrogenic and not substantially influenced by non-pyrogenic refractory material (e.g. petrogenic C or soil-derived polymers). In contrast, river discharge of total PAH and BC was not significantly correlated ($r^2 < 0.01$), a reasonable result, since PAHs are not all pyrogenic. For example, low molecular weight and alkylated PAHs would not be expected to be coeval with combustion-derived BC (Lima et al., 2005).

The global sources and sinks of BC to the ocean can be summarized as follows. First, the most recent estimate of atmospheric BC deposition to the ocean was combined with our estimated fluvial discharge summary (Table 9-3). It was assumed that the most recent value for atmospheric deposition of BC into the world's oceans (12 Tg y^{-1}) is also the most reliable (Jurardo et al., 2008), since it is similar in magnitude to the most updated and comprehensive estimate of the amount of fossil fuel and biomass BC emitted annually throughout the world ($5 - 20 \text{ Tg y}^{-1}$; Bond et al., 2004). This value for annual BC emission globally is, however, much lower than the often cited $50 - 270 \text{ Tg y}^{-1}$ for atmospheric BC emission strictly attributed to biomass burning (Kuhlbusch et al., 1996). Use of the larger range for BC emission calculated by Kuhlbusch et al., (1996) has yielded higher and perhaps erroneous estimates for the magnitude of BC introduced

into the ocean compared to the lower amounts found in ocean sediments (Druffel, 2004; Masiello, 2004).

The value for the global estimate for large river delivery of BC into the ocean calculated above is 5.4 Tg y^{-1} (Table 9-3). If this fluvial estimate is increased by 40% to incorporate SMR contributions, the total influx of particulate BC to the oceans from rivers may be as high as 8 Tg y^{-1} . This value is comparable to the 8 Tg y^{-1} supplied by atmospheric deposition and implies a total flux of $17 - 20 \text{ Tg y}^{-1}$ BC to the ocean. It should be noted, however, that these estimates pertain mainly to more refractory BC (e.g. soot) and may not include other more labile portions of the BC continuum such as charcoal.

9.5 FATE OF BLACK CARBON

Despite efforts to understand the short and long term cycling of BC derived from both natural and anthropogenic sources, the role of the coastal zone in sequestering fluvial BC is still uncertain. As noted earlier, long-term burial of biomass-derived BC in sediments is a net long-term sink for atmospheric CO_2 . In contrast, deposition of fossil fuel BC amounts to reburial of ancient carbon and does not promote net carbon sequestration. It is important to note that understanding of the role of BC in the carbon cycle is confounded by the timescales over which it is quantified in various reservoirs. For example, variations in aeolian BC production and deposition have been observed at seasonal-to-annual time scales whereas the marine sedimentary record can only be reconstructed at decadal or longer resolution. Consequently, the bulk of the sedimentary record of BC in many areas of the open ocean where sedimentation is low ($\text{mm } 1000\text{y}^{-1}$) will be dominated by BC derived from natural forest fire sources as opposed to modern anthropogenic sources. In contrast, coastal areas with higher sedimentation rates are more likely to record BC inputs from both anthropic fossil fuel and biomass combustion. Below, we discuss

sedimentary BC after it leaves a river, and consider its fate both in margins as well as the deep ocean.

9.5.1 Storage BC in the ocean

Globally, continental margins (shelf, slope and rise), estimated to be responsible for about 80-90% of the ocean's OC storage (Berner, 1982; Hedges and Keil, 1995), are the most significant reservoirs of marine OC. This is likely to be true for the majority of sedimentary BC, as well, which is thought to be deposited in continental shelf regions where water depths are less than 200 m (Suman et al., 1997). The amount and ultimate fate of sedimentary BC on continental margins will vary as a function of the concentration and flux of delivered BC, climate, tectonics, biological and physical mixing, and geomorphology of the margin. Additionally, throughout the Quaternary, there is evidence that stochastic events such as large storms may be drivers of the geochemical record in coastal environments (Goni et al., 2007; Rogers and Goodbred, 2010). Most of these drivers have not been comprehensively addressed in existing studies of bulk sedimentary carbon, let alone in BC studies.

We address this data gap by compiling existing estimates of BC deposition in coastal and open ocean sediments (Table 9-4) to determine if any salient trends can be attributed to any of the state variables noted above. Further, this compendium may help determine whether there is imbalance in BC production and deposition in the ocean. The data in Table 9-4 are listed geographically from high-to-low latitude. What is particularly striking is that the highest concentrations and largest fluxes of BC to coastal sediments are neither driven by latitude nor by proximity to large river mouths. And, once again, there does not appear to be a trend in BC concentrations or flux that occurs as a function of its quantification method. Rather, the highest fluxes of sedimentary BC listed in Table 9-4 (Swedish Continental Shelf, Chesapeake Bay, coastal East China Sea, and the Palos Verdes Shelf) are adjacent to urbanized fluvial systems,

suggesting that perhaps fossil fuel combustion primarily drives the bulk of sedimentary BC influx into coastal margins. Without higher temporal or spatially-resolved data, little more can be said about some of additional controls on the BC in the coastal sedimentary record.

The global flux of BC to the ocean is estimated at $\sim 20 \text{ Tg y}^{-1}$, with the reasonable assumption that approximately half of this is contributed by rivers and another half by the atmosphere. Though there exists the methodological issue of whether this estimate is mainly for soot only or includes all BC, we have shown that estimates don't vary greatly across detection methods that employ chemical and thermal oxidation (Tables 9-2 and 9-4). Assuming a figure of about 90% of total seaward flux of particulate BC to be deposited on continental margins (Suman et al., 1997), BC storage in these regions can be estimated to be 18 Tg y^{-1} . This leads to a supply-side estimate of $\sim 2 \text{ Tg of BC y}^{-1}$ deposited annually in the deep ocean at water depths greater than 200 m. Assuming an area of the deep ocean of $3.34 \times 10^8 \text{ km}^2$ (Suman et al., 1997), we calculate an expected BC flux of $0.60 \mu\text{g cm}^{-2} \text{ y}^{-1}$. This is similar to the range of measured flux values ranging from 0.001 to $3.6 \mu\text{g cm}^{-2} \text{ y}^{-1}$ (Table 9-4). Using a similar approach, 18 Tg y^{-1} delivered to continental shelves globally can be scaled to the total area of continental shelves ($2.75 \times 10^7 \text{ km}^2$), to yield $65 \mu\text{g cm}^{-2} \text{ y}^{-1}$, a comprehensive estimate for BC burial flux in coastal regions globally. In summary, our compilation of data for the annual budget of refractory BC suggests that its sources and sinks in the coastal and deep ocean are in balance. What our estimates are not able to provide however, is an evaluation of the role of labile pools of BC in the global carbon cycle. In light of recent studies quantifying BC degradation in the environment, data on the production, transport and fate of these labile pools such as char and charcoal are sorely needed.

9.5.2 Degradation of BC

There is a growing awareness that, while BC is a relatively refractory OM form, it is not completely inert in the environment. There are a number of lines of evidence that some pools of BC degrade via abiotic and biotic processes. Physical processes that may enhance BC degradation include UV exposure, leaching/solubilization, volatilization and consumption by later fires, erosion, resuspension and transport in the seabed and biological processes that include bioturbation and ingestion by macrofauna or remineralization by microbes.

The first type of evidence showing BC oxidation is studies measuring CO₂ efflux during *in vitro* laboratory incubations of laboratory-produced pyrogenic OM. These experiments have recorded pyrogenic C losses ranging about 0.5 to 5% over a few months to a few years, with some larger losses for materials produced at lower temperatures (<400 C, Baldock and Smernik, 2002; Bruun et al., 2012; Cross and Sohi, 2011; Hamer et al., 2004; Nguyen and Lehmann, 2009; Nguyen et al., 2010; Zimmerman, 2010). About half of these losses can be attributed to abiotic mechanisms (Zimmerman, 2010) and greater losses have been observed in the presence of labile sources of OM (so called positive priming, Cross and Sohi, 2011; Zimmerman et al., 2011). These observed losses translate to half-lives of 10 to several hundred years. However, exponential decreases in rates of C loss over time suggest that the so-called ‘power model’ of OM decomposition (Middleburg, 1989) might be more appropriately applied, leading to the calculation of BC half-lives in the thousand to million year range.

Second, measurements of BC distributions in soil profiles with known pyrogenic OM input histories lead to calculations of apparent BC field loss rates ranging from zero loss (completely stable) to BC half-lives of 3 y. Discrepancy among these studies are likely due to uncertainties that include, 1) the likelihood that BC particles are mobile in the soil column and lost via erosion, 2) differing BC degradation rates in different climates and soil types, 3) poorly known BC input rates, 4) differing BC quantification methods. Only a few studies have examined BC in marine sediment columns and none of these show progressive BC losses with time of burial (Masiello

and Druffel, 1998; Masiello and Druffel, 2003b; Middleburg et al., 1999). However, observations of 64% greater BC contents in abyssal plain sediments exposed to oxygen over 10-20 thousand years versus anoxic sediments suggest BC half-lives of ~10 ky (Middleburg et al., 1999). In addition, smaller BC-nonBC OM ^{14}C -age offsets in oxygenated versus anoxic Santa Monica Basin sediments (Masiello and Druffel, 2003b) also indicates that sedimentary remineralization of BC does occur, at least in the presence of oxygen.

Next, mass balance considerations suggest the occurrence of large-scale BC remineralization at some location between its production and final marine burial. For example, given estimated BC production rates, over the past 7000 y, soil OC would become 25 – 125% BC in just 20,000 years (Masiello and Druffel, 2003b). Calculations presented in this chapter also suggest losses between soils and coastal deposition. For example, globally averaged BC concentration in TSS of 2.9 mg/g versus 1.2 mg/g in deltaic sediment beds suggest about half the BC was lost during transport (though this could also result from preferential transport processes). This is despite the fact that many of these estimates are based on BC measurement by chemical and thermal oxidation methods which isolate relatively refractory BC, perhaps only soot. However, at this time, estimates of BC production, river export, and sediment sink fluxes are too few and too widely ranging (see Sanchez-Garcia et al., 2012) to use for predicting global BC mineralization rates.

One site that may be responsible for much of the BC degradation is on broad continental shelves in physically dynamic environments that, due to periodic resuspension, subjects sediments to redox oscillations which have been shown to enhance BC degradation (Nguyen and Lehmann, 2009). These areas have been termed ‘fluidized bed reactors’ (Aller, 1998) and have been shown to be hot spots of terrestrial OM of all types. In addition, continuous addition of labile planktonic debris may co-metabolize the degradation of OM, which has also been shown to enhance BC degradation (i.e. positive priming, Hamer et al., 2004; Zimmerman et al., 2011). This

may explain the seemingly contradictory observations of large amounts of BC export by the Amazon and other South American rivers (calculated in this chapter) and relatively low concentrations of BC in Amazon shelf sediments (Lohmann et al., 2009).

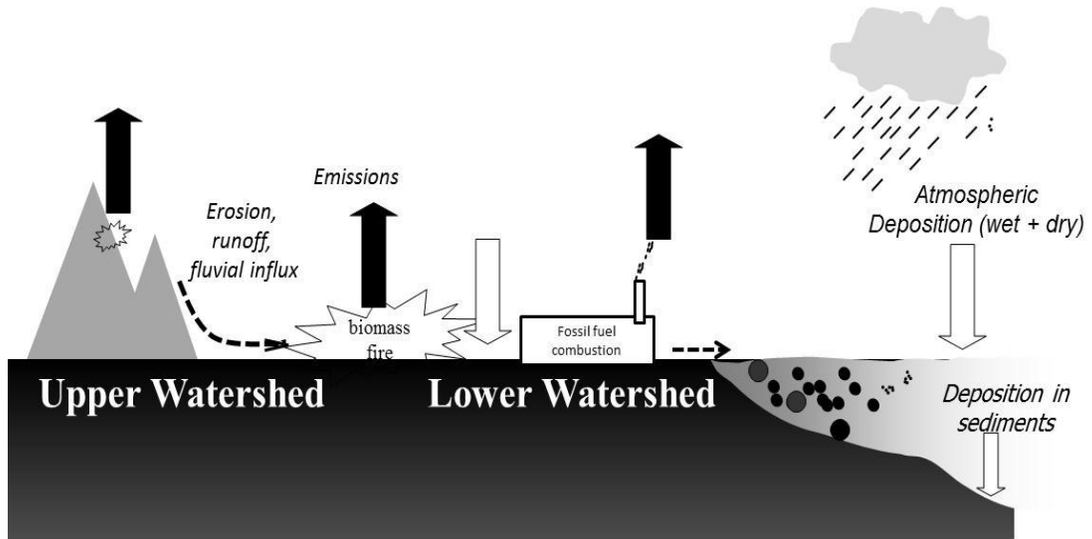
Finally, losses to BC may occur by solubilization or leaching either in the soil, or as part of the suspended or deposited sediment. While laboratory leaching experiments have recorded only small losses of BC ranging 0.04 to 1% (Abiven et al., 2011; Hammes et al., 2007; Katsoyiannis et al., 2011; Zimmerman and Gao, in press), aged charcoals seem to exhibit leach C rates similar to fresh ones, and detection of mobilized BC in rivers draining Brazil's Atlantic forest even 40 years after widespread burning (Dittmar et al., 2012), suggests that C losses may occur by this method may be continuous. And there is evidence from both laboratory microbial incubations showing up to 70% C loss over 1 year (Zimmerman and Gao, in press) and photo-degradation experiments showing a 20-fold decrease in coastal and North Atlantic deepwater dissolved BC (Stubbins et al., 2012b; Stubbins et al., 2010) that losses via leaching and then mineralization may be a primary sink for BC.

9.6 CONCLUSIONS

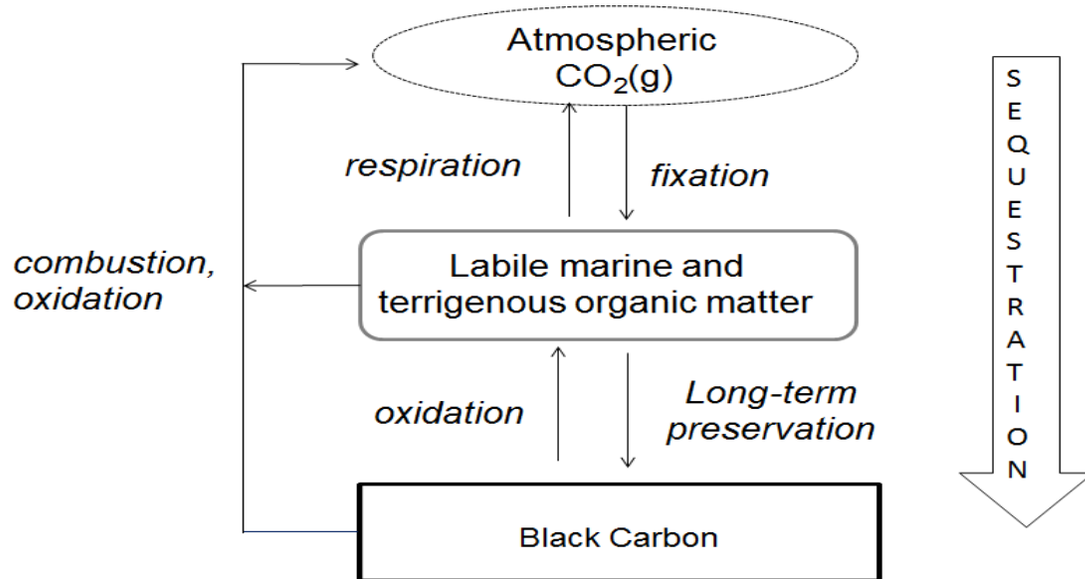
Many recently developed analytical techniques show great promise toward identification and quantification of pyrogenic carbon. Yet, even with the advent of these techniques, much more study of BC in the environment is warranted in order to fully understand it from a source-to-sink perspective. This is because BC generated on land and eventually deposited to the ocean, follows a complex and yet dynamic pathway enroute to its final burial. Consequently, several important questions about the global cycling of BC still remain. For example, how long does BC remain in soil and what processes move it to river systems? Does BC spend very little or a substantial amount of time in transit through river systems? To what extent does BC comprise the bedload versus suspended load of a river system? As there are few studies of the transport and

fate of BC within river systems, these questions cannot yet be answered. Since few studies concurrently measure char, charcoal, and refractory BC, questions also remain about the fate and transport of labile pools of BC shown in the left side of Figure 9-3. Since quantification of carbon within these more labile pools of BC is not typically done, we do not know their mobility through the short term carbon cycle. Also, the presence of dissolved BC in rainwater as well as in fluvial and ocean systems begs the question, “what are the mechanisms by which dissolved BC cycles in the ocean?” Lastly, while it seems clear that continental margins are repositories for the bulk of sedimentary BC, a comprehensive understanding of BC fate and degradation in these areas is lacking. With ever-increasing changes in land use and anthropic impacts to climate, now more than ever, it is necessary to address these data gaps in order to understand the role of BC in the global carbon cycle and its possible use for climate mitigation.

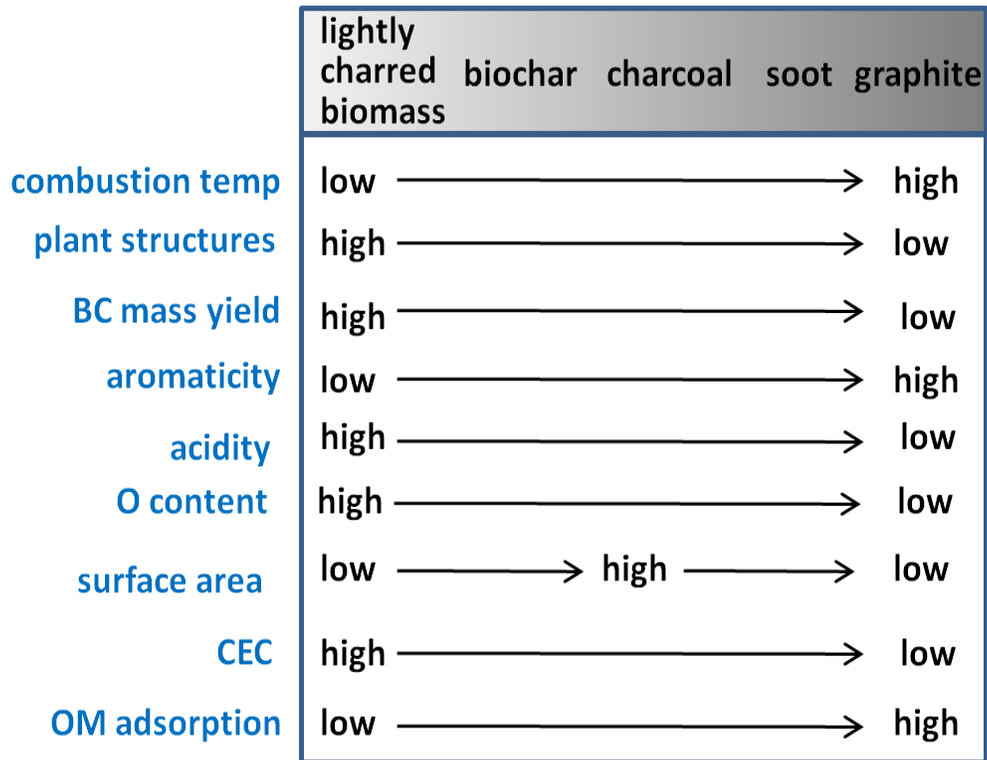
FIGURES



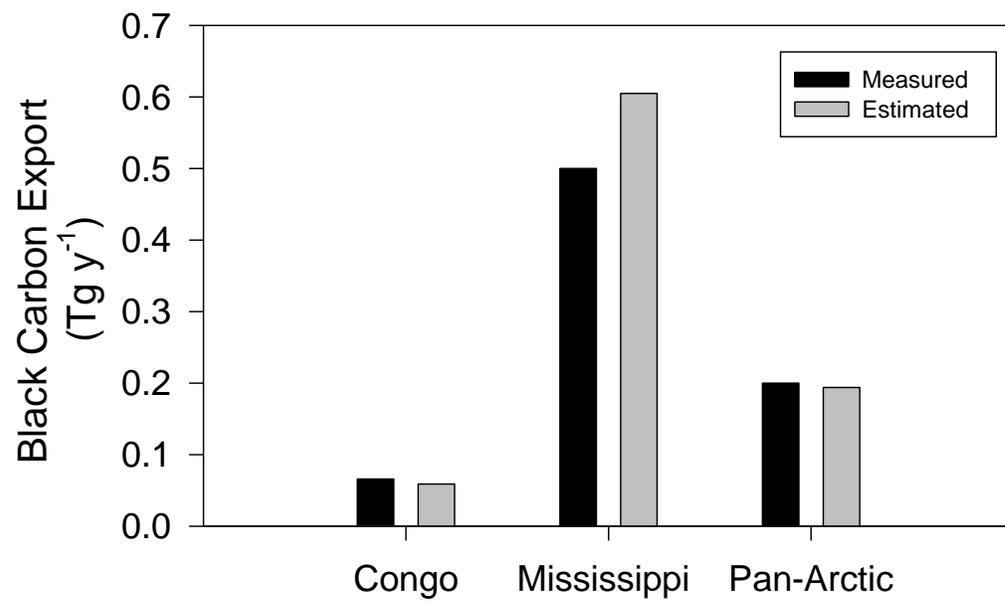
Mitra et al., Figure 1



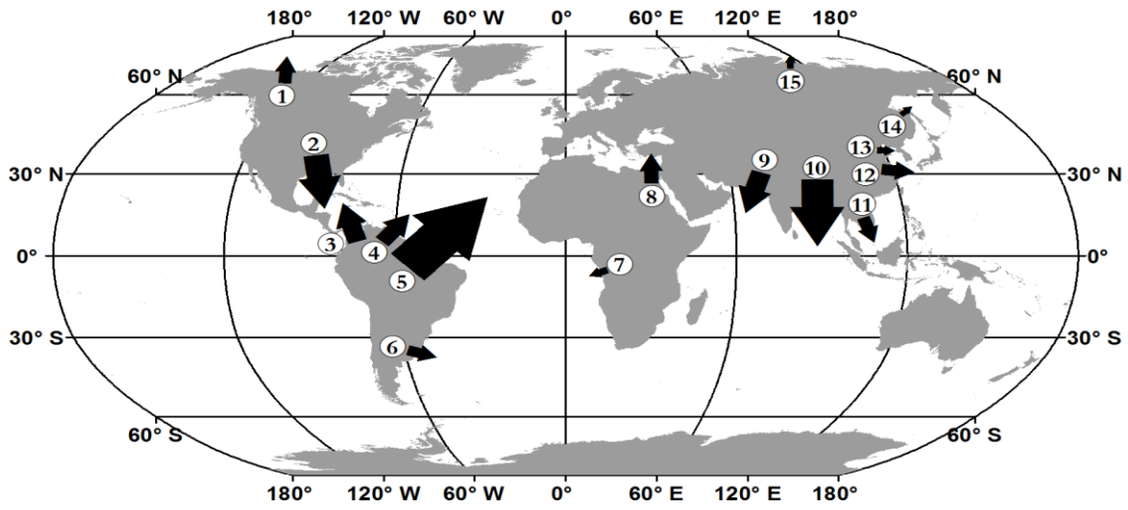
Mitra et al Figure 9-2



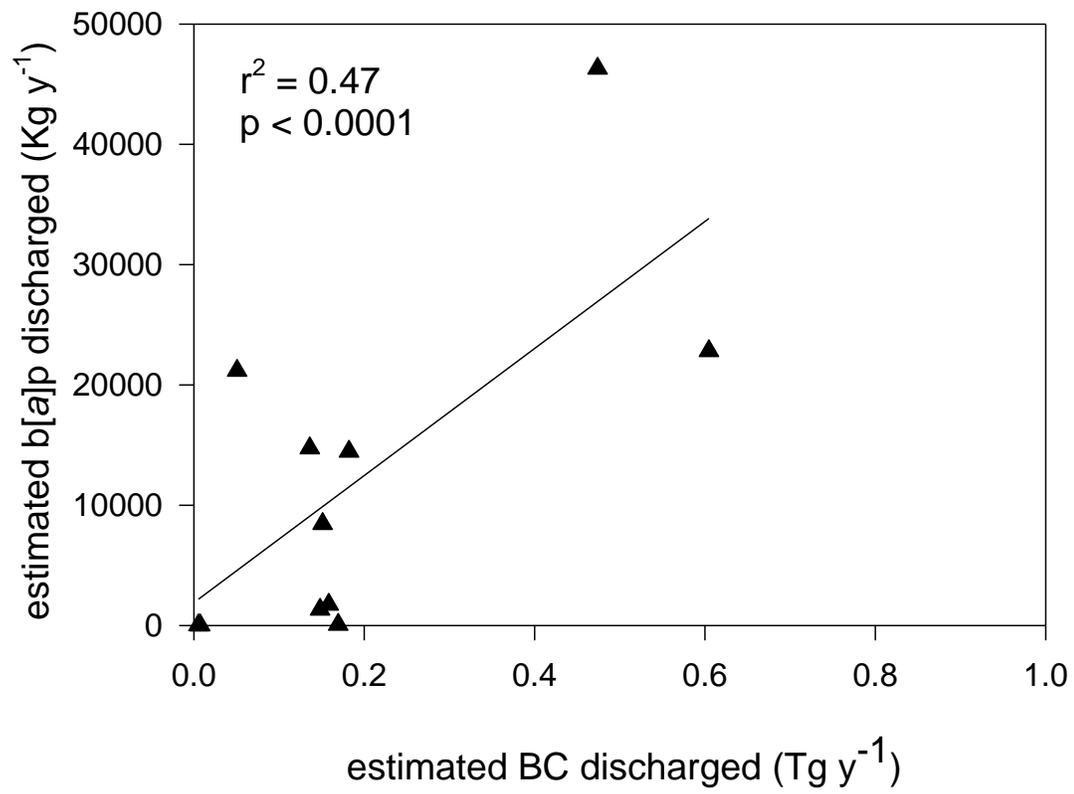
Mitra et al., Figure 9-3



Mitra et al., Figure 9-4



Mitra et al., Figure 9-5



Mitra et al., Figure 9-6

FIGURE CAPTIONS

Figure 9-1. Conceptual model of reservoirs and fluxes of particulate black carbon in the terrestrial and aquatic environments. Large black arrows depict BC emissions processes, white arrows depict deposition processes. Small dashed arrow depicts erosion, runoff, and fluvial influx which may occur throughout a watershed.

Figure 9-2. The relationship between biomass-derived black carbon and the global carbon cycle.

Figure 9-3. The combustion continuum. Modified from Masiello (2004) and Hedges et al. (2000).

Figure 9-4. Measured and estimated black carbon exported from a selected subset of major river systems from Table 9-3. Measured values for the Congo, Mississippi, and Pan-Arctic Rivers are from Lohmann et al., 2009, Mitra et al., 2002, and Elmquist et al., 2008, respectively. Pan-Arctic rivers BC export is calculated as the sum of the well-studied Yenisey, Ob, Lena, and MacKenzie rivers.

Figure 9-5. World map with arrows depicting fluvial BC discharge from major rivers noted in Table 9-3. Names of rivers are: 1) Mackenzie, 2) Mississippi, 3) Magdalena, 4) Orinoco, 5) Amazon, 6) Parana, 7) Congo, 8) Nile, 9) Indus, 10) Ganges - Brahmaputra, Irrawaddy, and Godavari, 11) Mekong, 12) Yangtze, 13) Huanghe, 14) Amur, and 15) Lena, Ob, and Yenisey.

Figure 9-6. Relationship between estimated values of benzo[*a*]pyrene and black carbon discharged in rivers listed in Table 9-3. Estimated values are the product of benzo[*a*]pyrene and black carbon concentrations in sediment multiplied by total suspended sediment discharged for each river system.

TABLES

Table 9-1. Compilation of dissolved or colloidal black carbon data measured in aquatic systems.

Geographical Region	Isolation method	Comments on isolation method	Amount of BC in Dissolved or Colloidal Phase ($\mu\text{g BC L}^{-1}$)	Dissolved or Colloidal BC/DOC (%)	Reference
Delaware Bay, Chesapeake Bay, USA	CTO-375 on HMW DOM	some charring observed but not quantified	45 - 1,038	4 - 7	(Mannino and Harvey, 2004)
Stream in New Jersey, USA	FT ICR-MS of aromatic compounds in DOC	non-quantitative	NA	NA	(Kim et al., 2004)
Rio Negro, Amazon River, Brazil	FT ICR-MS on HMW DOM	non-quantitative	NA	NA	(Kim et al., 2004)
River-to-ocean transect, north eastern Gulf of Mexico	Benzenepolycarboxylic acid quantification on cartridge isolated DOM via HPLC		9,600 – 44,000	0.9 -2.6	(Dittmar, 2008)
Southern Ocean	Benzenepolycarboxylic acid quantification on cartridge isolated DOM via HPLC	functionality confirmed using FT ICR-MS	7.3 - 9.6	~2	(Dittmar and Paeng, 2009)
Open Ocean (Sargasso Sea, North Central Pacific)	Radiocarbon age of benzenepolycarboxylic acid on cartridge-isolated DOM	ultrafiltered material	0.96 – 6	0.5 - 3.5	(Ziolkowski and Druffel, 2010)
North Atlantic Deep Water	Benzenepolycarboxylic acid quantification on cartridge isolated DOM	functionality confirmed using FT ICR-MS	0.66 – 13	0.2 – 2.1	(Stubbins et al., 2012a)
Lake Tahoe, USA	SP2-FIA ¹ on whole water	restricted to ~ 60 - 400 nm particle size range	0.02 - 0.45	0.01 - 0.02	(Bisiaux et al., 2011)
Amazon River	Benzenepolycarboxylic acid quantification on cartridge isolated DOM	aromaticity confirmed using FT ICR-MS	0 – 240	≤ 7	(Dittmar et al., 2012)

Notes:

¹flow injection, ultrasonic nebulization, laser induced incandescence photometer.

NA - not available.

Table 9-2. Values of particulate black carbon and organic carbon concurrently measured in river systems globally.

River	BC (mg/g)	TOC ¹ (mg/g)	BC/TOC (%)	Type of BC ²	Sample Type	Reference
Santa Clara ³	3.0	35.6	8.5	AD	surface water TSS	(Masiello and Druffel, 2001)
	7.1	16.0	44.3	AD		
	1.0	11.5	8.8	AD		
Ganges-Brahmaputra	2.9	5.7	50.9	AD	surface water TSS	Mitra et al., unpublished
Mississippi	3.8	28.0	16.0	Soot	surface water TSS	(Mitra et al., 2002)
	4.2	25.0	16.8	Soot	bottom water TSS	
	0.5	32.0	1.5	Soot	surface water TSS	
	0.8	25.0	3.0	Soot	bottom water TSS	
AVERAGE BC IN TSS	2.9 ± 2.2	22.4 ± 10.3	18.7 ± 18.7			
Changjiang River	0.6	2.6	24.6	Soot	surface sed (0-10cm)	(Kang et al., 2009)
	1.4	5.4	25.6	Soot		
	0.6	2.7	21.9	Soot		
	1.2	5.4	21.9	Soot		
	0.2	2.5	7.2	Soot		
	0.7	6.2	11.9	Soot		
	1.9	6.9	27.1	AD	surface sed (0-2 cm)	(Aeppli et al., 2012)
	2.2	4.1	52.9	AD		
	2.2	4.7	46.2	AD		
	1.7	5.9	28.8	AD		
	2.2	6.3	34.1	AD		
	1.5	5.2	29.2	AD		
	2.3	6.7	34.0	AD		
Congo Fan	0.8	33.0	2.4	Soot	surface sed (0-1 cm)	(Lohmann et al., 2009)
	0.9	20.2	4.5	Soot		
	0.9	13.3	6.8	Soot		
Amazon fan	1.2	3.7	32.4	Soot	surface sed (0-1 cm)	(Lohmann et al., 2009)
	1.1	7.1	15.5	Soot		
	1.1	5.7	19.3	Soot		
	1.1	5.6	19.6	Soot		

Ob	0.5	9.2	5.1	Soot	surface sed (0-2 cm)	(Elmqvist et al., 2008)
Yenisey	1.2	19.4	6.2	Soot		
Lena	0.2	5.7	3.0	Soot		
Indigirka	1.4	18.1	7.7	Soot		
Kolyma	1.5	19.2	7.8	Soot		
Yukon	1.2	12.4	9.7	Soot		
Mackenzie	1.0	20.2	5.0	Soot		
<hr/>						
AVERAGE BC IN SEABED SEDIMENTS ⁴	1.2 ± 0.6	9.5 ± 7.5	18.9 ± 13.7			
<hr/>						
OVERALL AVERAGE	1.6 ± 1.3	12.5 ± 9.7	18.9 ± 14.7			
<hr/>						

Notes:

¹TOC values correspond to the same time period as that of BC collection.

²Type of BC "AD" refers to isolation by acid-dichromate, "soot" refers to that isolated using CTO-375 or a similar chemothermal oxidation method. See original references for additional method details.

³Although not a large river, BC concentrations in the Santa Clara were included due to the overall paucity of information on BC in suspended sediments for large rivers globally.

⁴While many stations were sampled within each of these studies, values of BC in seabed were only included if they were obtained directly at the mouth of that river system.

Table 9-3. Sediment, particulate organic carbon, black carbon and polycyclic aromatic hydrocarbon discharge from large rivers.

River	Qs (T yr ⁻¹)	Qav (m ⁻³ s)	Yield (T km ⁻² yr ⁻¹)	TSS (kg m ³)	TSS (MT yr ⁻¹)	POC (mg L ⁻¹)	POC reference	BC discharged ¹ (tg y ⁻¹)	tPAH (ng g ⁻¹)	b[a]p (ng g ⁻¹)	b[a]p discharged (kg y ⁻¹)	PAH reference
Amazon	1.2x10 ⁹	2.0x10 ⁵	193	0.19	1190.4	2.83	(Richey et al., 1990)	1.91	701	NA		(Budzinski et al., 1997; Lohmann et al., 2009)
Mississippi	4.0x10 ⁸	1.7x10 ⁴	124	0.11	61.3	16.90	(Bianchi et al., 2007)	0.60	5300	372	22817	(Maruya et al., 1997; Overton et al., 2004; Santschi et al., 2001)
Ganges- Brahmaputra	3.1x10 ⁸	3.1x10 ⁴	701	0.32	312.8	2.50	(Aucour et al., 2006)	0.47	2938	148	46300	(Ahmad et al., 1996; Domínguez et al., 2010)
Indus	2.5x10 ⁸	3.2x10 ³	261	1.05	105.0	8.82	(Ludwig et al., 1996a)	0.38	NA	NA		
Magdalena	2.2x10 ⁸	7.2x10 ³	868	0.63	143.3	NA		0.33	NA	NA		
Orinoco	1.7x10 ⁸	3.4 x10 ⁴	180	0.06	65.7	1.50	(Paolini, 1995)	0.26	NA	NA		
Godavari	1.7x10 ⁸	3.3x10 ³	611	2.03	213.2	7.20	(Balakrishna and Probst, 2005)	0.26	NA	NA		
Nile	1.2x10 ⁸	3.5x10 ³	43	1.4	153.8	3.85	(Ludwig et al., 1996b)	0.18	1229	94	14459	(Badawy and Emababy, 2010)
Irrawaddy	1.1x10 ⁸	1.4x10 ⁴	614	0.26	111.2	3.59		0.17	133	0.6	67	(Chernova et al., 2001)
Yangtze	1.0x10 ⁸	2.8x10 ⁴	251	0.13	119.3	1.62	(Wang et al., 2012)	0.16	502	14.6	1741	(Bouloubassi et al., 2001; Li et al., 2012a)
MacKenzie	1.0x10 ⁸	1.0x10 ⁴	58	0.39	128.0	6.36	(Macdonald et al., 1998)	0.15	1200	66	8450	(Yunker et al., 1993; Yunker et al., 2011)
Mekong	9.8x10 ⁷	1.7x10 ⁴	123	0.08	44.2	2.01	(Ellis et al., 2012)	0.15	1379	30	1326	(Keenan et al., 2010)
Paraná	9.0x10 ⁷	1.5x10 ⁴	31	0.18	82.3	1.69	(Ludwig et al., 1996a)	0.14	2120	179	14739	(Cataldo et al., 2001; Colombo et al., 2006)
Congo ²	3.9x10 ⁷	3.7x10 ⁴	8.8	0.03	35.0	1.70	(Coynel et al., 2005)	0.06	227	NA		(Lohmann et al., 2009)
Huanghe	3.3x10 ⁷	1.5x10 ³	1122	2.52	117.7	15.26	(Wang et al., 2012)	0.05	5734	180	21189	(Jiao et al., 2012; Ma et al., 2008; Wu et al., 2001; Xu et al., 2007)
Amur	2.4x10 ⁷	1.1x10 ⁴	28 ³	0.07	25.3	0.24	(Meybeck and Ragu, 1997)	0.04	1100			(Nemirovskaya, 2007)
Lena	2.0x10 ⁷	1.7x10 ⁴	8	0.02	10.8	0.99	(Lobbés et al., 2000)	0.03	97			(Elmquist et al., 2008b)
Yenisey	4.9x10 ⁶	1.8x10 ⁴	5 ³	0.008	5.0	0.98	(Unger et al.,	0.01	636	4.3	22	(Fernandes and Sicre, Mitra et al – p.41

Ob	3.5x10 ⁶	1.3x10 ⁴	6.4 ³	0.009	3.8	0.66	2005) (Unger et al., 2005)	0.01	376	6.9	26	1999;Dahle et al., 2003) (Fernandes and Sicre, 1999;Dahle et al., 2003)
----	---------------------	---------------------	------------------	-------	-----	------	----------------------------------	------	-----	-----	----	---

Notes:

¹A value of 5.4 Tg y⁻¹ is estimated as the sum of BC discharged.

²Although the Congo's POC load is low, its expansive watershed is in an area subject to high biomass burning (Cachier and Ducret, 1991; Delmas et al., 1991) and 2 – 7% of its POC is made up of soot BC.

³Milliman and Meade, 1983.

Unless otherwise noted, physical variables for rivers are from Syvitski and Saito (2007). Qs: suspended load, Qav : average discharge Yield, TSS: total suspended sediments, POC: particulate organic carbon, BC: black carbon, b[a]p: benzo[a]pyrene. NA - not available.

Table 9-4. Deposition of BC to the coastal and open ocean.

Geographical Region	Isolation method	BC Concentration in Sediments (mg g ⁻¹)	BC Flux (µg cm ⁻² y ⁻¹)	Reference ¹
Coastal Areas				
Saanich Inlet, BC, Canada	Chemical treatment, IR spectrometry	0.55	74	(Griffin and Goldberg, 1975)
Pan Arctic Shelf	CTO - 375	0.5 – 1.5	0.13 -5.6	(Elmquist et al., 2008)
Swedish Continental Shelf	CTO - 375	2 – 2.8	270 – 8100	(Sanchez-Garcia et al., 2012)
Gulf of Maine, USA	CTO-375, pyrene fluorescence quenching	0.11 – 0.17; 0.1 – 15 ²	100 – 200; 80 – 170	(Flores-Cervantes et al., 2009a; Gustafsson and Gschwend, 1998)
Puget Sound, Washington, USA	GBC ³	0.1 – 0.3	30 – 1,040	(Kuo et al., 2011)
Washington Coast, USA	GBC ³	0.14 – 0.71	1.2 – 3.1	(Dickens et al., 2004)
Chesapeake Bay, USA	AD	1.1 – 7.5	760 – 5,400	(Mitra et al., 2009)
Coastal East China Sea	AD	1.5 – 2.2	210 – 1300	(Li et al., 2012)
Santa Barbara Basin	Chemical treatment, IR spectrometry ; AD	0.29; 2.2	26; 35	(Griffin and Goldberg, 1975) (Masiello and Druffel, 2003)
Gulf of Panama	Chemical treatment, IR spectrometry	NQ	109 -354	(Suman, 1986)
Continental rise adjacent to Nicaragua	Chemical treatment, IR spectrometry	NQ	23 - 135	(Suman, 1986)
Eastern South Atlantic Shelf	CTO-375	0.8 – 1.6	0.5 – 7.8	(Lohmann et al., 2009)

Western South Atlantic Shelf	CTO -375	0.4 – 1.7	0.6 – 2.6	(Lohmann et al., 2009)
Palos Verdes Shelf	CTO -375	1.2 - 1.4	700 – 1000	(Gustafsson and Gschwend, 1998)
<hr/>				
Open Ocean				
<hr/>				
South China Sea	AD	0.01 – 1.9	NQ	(Jia et al., 2003)
Pelagic Pacific and Atlantic	Chemical treatment, IR spectrometry	0.01 - 1	0.002 – 0.2	(Smith et al., 1973)
Pelagic Pacific	Chemical treatment, IR spectrometry	0.44	0.057	(Griffin and Goldberg, 1975)
Pelagic Pacific	Chemical treatment, IR spectrometry	0.001 - 3	< 0.0014 – 3.6	(Herring, 1985)
Equatorial Pacific	GBC	0.092	0.05	(Dickens et al., 2004)
<hr/>				

Notes:

¹The following studies of BC in the deep ocean sediments were reviewed but were not included in the table because neither concentration nor sediment mass flux were provided: (Bird and Cali, 1998; Masiello and Druffel, 1998; Thevenon et al., 2004).

²Concentrations in $\mu\text{g L}^{-1}$.

³The GBC (graphitic black carbon) technique is a thermal oxidation technique with additional organic acid pre-treatment specifically designed to minimize charring of occluded organic matter (Gelinis et al., 2001).

NQ – unable to be quantified.

REFERENCES

- Abiven, S., Hengartner, P., Schneider, M.P.W., Singh, N., Schmidt, M.W.I., 2011. Pyrogenic carbon soluble fraction is larger and more aromatic in aged charcoal than in fresh charcoal. *Soil Biology & Biochemistry*, 43(7): 1615-1617.
- Accardi-Dey, A., 2003. Black carbon in marine sediments: quantification and implications for the sorption of polycyclic aromatic hydrocarbons, Massachusetts Institute of Technology, Cambridge, 282 pp.
- Aeppli, C. et al., 2012. Oil Weathering after the Deepwater Horizon Disaster Led to the Formation of Oxygenated Residues. *Environmental Science and Technology*, 46: 8799-8807.
- Ahmad, S., Ajmal, M., Nomani, A.A., 1996. Organochlorines and Polycyclic Aromatic Hydrocarbons in the Sediments of Ganges River (India). *Bulletin of Environmental Contamination and Toxicology*, 57(5): 794.
- Aller, R., 1998. Mobile deltaic and continental shelf muds as suboxic, fluidized bed reactors. *Marine Chemistry*, 61: 143-155.
- Arranz, J.M.D. et al., 2009. Structural properties of non-combustion-derived refractory organic matter which interfere with BC quantification. *Journal of Analytical and Applied Pyrolysis*, 85(1-2): 399-407.
- Aucour, A.M., France-Lanord, C., Pedoja, K., Pierson-Wickmann, A.C., Sheppard, S.M.F., 2006. Fluxes and sources of particulate organic carbon in the Ganga-Brahmaputra river system. *Global Biogeochemical Cycles*, 20(2).
- Bachmann, J., 2009. *Black Carbon: A Science/Policy Primer*.
- Badawy, M.I., Emababy, M.A., 2010. Distribution of polycyclic aromatic hydrocarbons in drinking water in Egypt. *Desalination*, 251(1): 34-40.
- Balakrishna, K., Probst, J.L., 2005. Organic carbon transport and C/N ratio variations in a large tropical river: Godavari as a case study, India. *Biogeochemistry*, 73(3): 457-473.
- Baldock, J.A., Masiello, C.A., Gelin, Y., Hedges, J.I., 2004. Cycling and composition of organic matter in terrestrial and marine ecosystems. *Marine Chemistry*, 92: 39-64.
- Baldock, J.A., Smernik, R.J., 2002. Chemical composition and bioavailability of thermally altered *Pinus resinosa* (Red Pine) wood. *Organic Geochemistry*, 33(9): 1093-1109.
- Barry, R.G., 1996. The parameterization of surface albedo for sea ice and its snow cover. *Progress in Physical Geography*, 20(1): 63-79.
- Berner, R., 1982. Burial of organic carbon and pyrite sulfur in the modern ocean and its geochemical and environmental significance. *American Journal of Science*, 282: 451-473.
- Bianchi, T.S., Wysocki, L.A., Stewart, M., Filley, T.R., McKee, B.A., 2007. Temporal variability in terrestrially-derived sources of particulate organic carbon in the lower Mississippi River and its upper tributaries. *Geochimica Et Cosmochimica Acta*, 71(18): 4425-4437.

- Bidleman, T.F., 1988. Atmospheric processes - wet and dry deposition of organic compounds are controlled by their vapor particle partitioning. *Environmental Science & Technology*, 22(4): 361-367.
- Bird, M.I., Ascough, P.L., 2012. Isotopes in pyrogenic carbon: A review. *Organic Geochemistry*, 42(12): 1529-1539.
- Bird, M.I., Cali, J.A., 1998. A million-year record of fire in sub-Saharan Africa. *Nature*, 394(6695): 767-769.
- Bird, M.I., Moyo, C., Veenendaal, E.M., Lloyd, J., Frost, P., 1999. Stability of elemental carbon in a savanna soil. *Global Biogeochemical Cycles*, 13(4): 923-932.
- Bisiaux, M.M. et al., 2011. Stormwater and Fire as Sources of Black Carbon Nanoparticles to Lake Tahoe. *Environmental Science & Technology*, 45(6): 2065-2071.
- Bond, T.C., 2001. Spectral dependence of visible light absorption by carbonaceous particles emitted from coal combustion. *Geophysical Research Letters*, 28(21): 4075-4078.
- Bouloubassi, I., Fillaux, J., Saliot, A., 2001. Hydrocarbons in surface Sediments from the Changjiang (Yangtze River) Estuary, East China Sea. *Marine Pollution Bulletin*, 42(12): 1335-1346.
- Bowman, D.M.J.S. et al., 2009. Fire in the Earth System. *Science*, 324: 481-484.
- Brodowski, S., Rodionov, A., Haumaier, L., Glaser, B., Amelung, W., 2005. Revised black carbon assessment using benzene polycarboxylic acids. *Organic Geochemistry*, 36(9): 1299-1310.
- Bruun, E.W., Ambus, P., Egsgaard, H., Hauggaard-Nielsen, H., 2012. Effects of slow and fast pyrolysis biochar on soil C and N turnover dynamics. *Soil Biology & Biochemistry*, 46: 73-79.
- Budzinski, H. et al., 1997. Identification of polycyclic aromatic hydrocarbons in sediments from the Amazon Fan: occurrence and diagenetic evolution, *Proceedings- Ocean Drilling Program Scientific Results*. NSF, pp. 555-564.
- Burdige, D.J., 2005. Burial of terrestrial organic matter in marine sediments: A re-assessment. *Global Biogeochemical Cycles*, 19: doi:10.1029/2004GB002368, 2005.
- Cachier, H., Ducret, J., 1991. Influence of biomass burning on equatorial African rains. *Nature*, 352(6332): 228-230.
- Carcaillet, C. et al., 2002. Holocene biomass burning and global dynamics of the carbon cycle. *Chemosphere*, 49: 845-863.
- Cataldo, D., Colombo, J., Boltovskoy, D., Bilos, C., Landoni, P., 2001. Environmental toxicity assessment in the Paran· river delta (Argentina): simultaneous evaluation of selected pollutants and mortality rates of *Corbicula fluminea* (Bivalvia) early juveniles. *Environmental Pollution*, 112(3): 379-389.
- Chakraborty, B.B., Long, R., 1967. Gas chromatographic analysis of polycyclic aromatic hydrocarbons in soot samples. *Environmental Science & Technology*, 1(10): 828-834.
- Chasar, L.S., Chanton, J.P., Glaser, P.H., Siegel, D.I., Rivers, J.S., 2000. Radiocarbon and stable carbon isotopic evidence for transport and transformation of dissolved organic carbon, dissolved inorganic carbon, and CH₄ in a northern Minnesota peatland. *Global Biogeochemical Cycles*, 14(4): 1095-1108.

- Chernova, T. et al., 2001. The composition and the source of hydrocarbons in sediments taken from the tectonically active Andaman Backarc Basin, Indian Ocean. *Marine chemistry*, 75(1): 1-15.
- Chughtai, A.R., Jassim, J.A., Peterson, J.H., Stedman, D.H., Smith, D.M., 1991. Spectroscopic and solubility characteristics of oxidized soots. *Aerosol Science and Technology*, 15: 112 - 126.
- Clark, J.S., 1988. Particle motion and the theory of charcoal analysis: source area, transport, deposition, and sampling. *Quaternary Research*, 30: 67-80.
- Clarke, A.D., Noone, K.J., 1985. Soot in the Arctic snowpack - a cause for perturbations in radiative-transfer. *Atmospheric Environment*, 19(12): 2045-2053.
- Clarke, A.D., Owens, S.R., Zhou, J.C., 2006. An ultrafine sea-salt flux from breaking waves: Implications for cloud condensation nuclei in the remote marine atmosphere. *Journal of Geophysical Research-Atmospheres*, 111(D6).
- Clarke, A.D., Weiss, R.E., Charlson, R.J., 1984. Elemental carbon aerosols in the urban, rural, and remote-marine troposphere and in the stratosphere - inferences from high-absorption data and consequences regarding radiative transfer. *Science of the Total Environment*, 36(JUN): 97-102.
- Colombo, J.C. et al., 2006. Sources, vertical fluxes, and equivalent toxicity of aromatic hydrocarbons in coastal sediments of the Rio de la Plata Estuary, Argentina. *Environmental Science & Technology*, 40(3): 734-740.
- Conedera, M. et al., 2009. Reconstructing past fire regimes: methods, applications, and relevance to fire management and conservation. *Quaternary Science Reviews*, 28(5-6): 555-576.
- Conway, H., Gades, A., Raymond, C.F., 1996. Albedo of dirty snow during conditions of melt. *Water Resources Research*, 32(6): 1713-1718.
- Coyne, A., Seyler, P., Etcheber, H., Meybeck, M., Orange, D., 2005. Spatial and seasonal dynamics of total suspended sediment and organic carbon species in the Congo River. *Global Biogeochemical Cycles*, 19(4).
- Cross, A., Sohi, S.P., 2011. The priming potential of biochar products in relation to labile carbon contents and soil organic matter status. *Soil Biology & Biochemistry*, 43(10): 2127-2134.
- Currie, L.A. et al., 2002. A critical evaluation of interlaboratory data on total, elemental, and isotopic carbon in the carbonaceous particle reference material, NIST SRM 1649a. *Journal of Research of the National Institute of Standards and Technology*, 107(3): 279-298.
- Cusack, D.F., Chadwick, O.A., Hockaday, W.C., Vitousek, P.M., 2012. Mineralogical controls on soil black carbon preservation. *Global Biogeochemical Cycles*, 26.
- Czimczik, C.I., Preston, C.M., Schmidt, M.W.I., Schulze, E.D., 2003. How surface fire in Siberian Scots pine forests affects soil organic carbon in the forest floor: Stocks, molecular structure, and conversion to black carbon (charcoal). *Global Biogeochemical Cycles*, 17(1).
- Czimczik, C.I., Preston, C.M., Schmidt, M.W.I., Werner, R.A., Schulze, E.-D., 2002. Effects of charring on mass, organic carbon, and stable carbon isotopic composition of wood. *Organic Geochemistry*, 33: 1207-1223.

- Czimczik, C.I., Schmidt, M.W.I., 2002. Effects of charring on mass, organic carbon, and stable carbon isotope composition of wood. *Organic Geochemistry*, 33: 1207-1223.
- Czimczik, C.I., Schmidt, M.W.I., Schulze, E.D., 2005. Effects of increasing fire frequency on black carbon and organic matter in Podzols of Siberian Scots pine forests. *European Journal of Soil Science*, 56(3): 417-428.
- Dahle, S., Savinov, V.M., Matishov, G.G., Evenset, A., Naes, K., 2003. Polycyclic aromatic hydrocarbons (PAHs) in bottom sediments of the Kara Sea shelf, Gulf of Ob and Yenisei Bay. *The Science of The Total Environment*, 306(1-3): 57.
- Dai, X., Boutton, T.W., Glaser, B., Ansley, R.J., Zech, W., 2005. Black carbon in a temperate mixed-grass savanna. *Soil Biology & Biochemistry*, 37(10): 1879-1881.
- Decesari, S. et al., 2002. Water soluble organic compounds formed by oxidation of soot. *Atmospheric Environment*, 36(11): 1827-1832.
- Delmas, R.A., Loudjani, P., Podaire, A., Menaut, J., 1991. Biomass burning in Africa: an assessment of annually burned biomass. In: Levine, J. (Ed.), *Global Biomass Burning*. MIT Press, Cambridge.
- Dick, D.P. et al., 2005. Characteristics of soil organic matter of different Brazilian Ferralsols under native vegetation as a function of soil depth. *Geoderma*, 124(3-4): 319-333.
- Dickens, A., Gelinas, Y., Masiello, C.A., Wakeham, S.G., Hedges, J.I., 2004a. Reburial of fossil organic carbon in marine sediment. *Nature*, 427: 336-339.
- Dickens, A.F. et al., 2006. Solid-state C-13 NMR analysis of size and density fractions of marine sediments: Insight into organic carbon sources and preservation mechanisms. *Geochimica Et Cosmochimica Acta*, 70(3): 666-686.
- Dickens, A.F., Gelinas, Y., Hedges, J.I., 2004b. Physical separation of combustion and rock sources of graphitic black carbon in sediments. *Marine Chemistry*, 92: 215-223.
- Dittmar, T., 2008. The molecular level determination of black carbon in marine dissolved organic matter. *Organic Geochemistry*, 39(4): 396-407.
- Dittmar, T. et al., 2012. Continuous flux of dissolved black carbon from a vanished tropical forest biome. *Nature Geoscience*, 5(9): 618-622.
- Dittmar, T., Paeng, J., 2009. A heat-induced molecular signature in marine dissolved organic matter. *Nature Geoscience*, 2: 175-179.
- Domínguez, C. et al., 2010. Quantification and source identification of polycyclic aromatic hydrocarbons in core sediments from Sundarban Mangrove Wetland, India. *Archives of environmental contamination and toxicology*, 59(1): 49-61.
- Drivas, P.J., 1982. Calculation of evaporative emissions from multicomponent liquid spills. *Environmental Science & Technology*, 16(10): 726-728.
- Druffel, E.R.M., 2004. Comments on the importance of black carbon in the global carbon cycle. *Marine Chemistry*, 92(1-4): 197-200.
- Elias, V.O., Simoneit, B.R.T., Cordeiro, R.C., Turcq, B., 2001. Evaluating levoglucosan as an indicator of biomass burning in Carajas, Amazonia: a comparison to the charcoal record. *Geochimica et Cosmochimica Acta*, 65: 267-272.
- Ellis, E.E., Keil, R.G., Ingalls, A.E., Richey, J.E., Alin, S.R., 2012. Seasonal variability in the sources of particulate organic matter of the Mekong River as discerned by

- elemental and lignin analyses. *Journal of Geophysical Research*, 117(G1): G01038.
- Elmquist, M., Cornelissen, G., Kukulska, Z., Gustafsson, O., 2006. Distinct oxidative stabilities of char versus soot black carbon: Implications for quantification and environmental recalcitrance. *Global Biogeochemical Cycles*, 20(2).
- Elmquist, M., Gustafsson, O., Andersson, P., 2004. Quantification of sedimentary black carbon using the chemothermal oxidation method: an evaluation of ex situ pretreatments and standard addition approaches. *Limnology and Oceanography: Methods*, 2: 417-427.
- Elmquist, M., Semiletov, I., Guo, L., Gustafsson, 2008a. Pan-Arctic patterns in black carbon sources and fluvial discharges deduced from radiocarbon and PAH source apportionment markers in estuarine surface sediments. *Global biogeochemical cycles*, 22(2): GB2018.
- Elmquist, M., Semiletov, I., Guo, L.D., Gustafsson, O., 2008b. Pan-Arctic patterns in black carbon sources and fluvial discharges deduced from radiocarbon and PAH source apportionment markers in estuarine surface sediments. *Global Biogeochemical Cycles*, 22(2).
- Fernandes, M., Sicre, M.A., 1999. Polycyclic aromatic hydrocarbons in the Arctic: Ob and Yenisei estuaries and Kara Sea shelf. *Estuarine, Coastal and Shelf Science*, 48(6): 725-737.
- Fernandes, S.D., Trautmann, N.M., Streets, D.G., Roden, C.A., Bond, T.C., 2007. Global biofuel use, 1850-2000. *Global Biogeochemical Cycles*, 21: doi:10.1029/2006GB002836.
- Flores-Cervantes, D.X., 2008. Black carbon in the Gulf of Maine: new insights into inputs and cycling of combustion-derived organic carbon, Massachusetts Institute of Technology, Boston, 289pp.
- Flores-Cervantes, D.X., Plata, D.L., MacFarlane, J.K., Reddy, C.M., Gschwend, P.M., 2009a. Black carbon in marine particulate organic carbon: Inputs and cycling of highly recalcitrant organic carbon in the Gulf of Maine. *Marine chemistry*, 113(3): 172-181.
- Flores-Cervantes, D.X., Reddy, C.M., Gschwend, P.M., 2009b. Inferring black carbon concentrations in particulate organic matter by observing pyrene fluorescence losses. *Environmental Science and Technology*, 43: 4864-4870.
- Forbes, M.S., Raison, R.J., Skjemstad, J.O., 2006. Formation, transformation, and transport of black carbon (charcoal) in terrestrial and aquatic ecosystems. *Science of the Total Environment*, 370: 190-206.
- Forster, P. et al., 2007. *Changes in Atmospheric Constituents and in Radiative Forcing*, Cambridge.
- Forster, P., V. Ramaswamy, P. Artaxo, T. Berntsen, R. Betts, D.W. Fahey, J. Haywood, J. Lean, D.C. Lowe, G. Myhre, J. Nganga, R. Prinn, G. Raga, M.S.a.R.V.D., 2007. *Changes in Atmospheric Constituents and in Radiative Forcing*, Cambridge, UK and New York, NY, USA.
- Gelinas, Y., Prentice, K.M., Baldock, J.A., Hedges, J.I., 2001. An improved thermal oxidation method for the quantification of soot/graphitic carbon in sediments and soils. *Environmental Science and Technology*, 35: 3519-3525.

- Gilhooly, W.P. et al., 1996. Stable carbon isotopic analysis of charcoal from single plant sources. In: Levine, J. (Ed.), *Biomass Burning and Global Change*. MIT Press, Cambridge, pp. 466-471.
- Glaser, B., Haumaier, L., Guggenberger, G., Zech, W., 1998. Black carbon in soils: the use of benzenecarboxylic acid as specific markers. *Organic Geochemistry*, 29(4): 811-819.
- Goldberg, E., 1985. *Black carbon in the environment: properties and distribution*. John Wiley & Sons, New York.
- Goni, M., Alleau, Y., Crobett, R., Walsh, J., 2007. The Effects of Hurricanes Katrina and Rita on the Seabed of the Louisiana Shelf. *Sedimentary Record*, 5: 1.
- Griffin, J., Goldberg, E.D., 1975. The Fluxes of Elemental Carbon in Coastal Marine Sediments. *Limnology and Oceanography*, 20(3): 456-463.
- Gustafsson, Gschwend, P.M., 1998. The flux of black carbon to surface sediments on the New England continental shelf. *Geochimica et Cosmochimica Acta*, 62(3): 465-472.
- Gustafsson, O., Gschwend, P.M., 1997. Soot as a strong partitioning medium for polycyclic aromatic hydrocarbons in aquatic systems. Molecular markers in environmental geochemistry, ACS Symposium Series Vol. 671. American Chemical Society, Washington DC, 365-381 pp.
- Gustafsson, O., Haghseta, F., Chan, C., Macfarlane, J., Gschwend, P., 1997. Quantification of the dilute sedimentary soot phase: implications for PAH speciation and bioavailability. *Environmental Science and Technology*, 31: 203-209.
- Gustafsson, O. et al., 2009. Brown Clouds over South Asia: Biomass or Fossil Fuel Combustion? *Science*, 323(5913): 495-498.
- Haberstroh, P.R. et al., 2006. Chemical composition of the graphitic black carbon fraction in riverine and marine sediments at sub-micron scales using carbon X-ray spectromicroscopy. *Geochimica Et Cosmochimica Acta*, 70(6): 1483-1494.
- Hamer, U., Marschner, B., Brodowski, S., Amelung, W., 2004. Interactive priming of black carbon and glucose mineralisation. *Organic Geochemistry*, 35: 823-830.
- Hammes, K. et al., 2007. Comparison of quantification methods to measure fire-derived (black/elemental) carbon in soils and sediments using reference materials from soil, water, sediment and the atmosphere. *Global Biogeochemical Cycles*, 21(3).
- Hansen, J., Nazarenko, L., 2004. Soot climate forcing via snow and ice albedos. *Proceedings of the National Academy of Sciences of the United States of America*, 101(2): 423-428.
- Hansen, J., Sato, M., Ruedy, R., Lacis, A., Oinas, V., 2000. Global warming in the twenty-first century: An alternative scenario. *Proceedings of the National Academy of Sciences of the United States of America*, 97(18): 9875-9880.
- Haynes, B.S., Wagner, H.G., 1981. Soot formation. *Progress in Energy and Combustion Science*, 7(4): 229-273.
- Hedges, J.I., 1992. Global biogeochemical cycles: progress and problems. *Marine Chemistry*, 39: 67-93.
- Hedges, J.I., Keil, R.G., 1995. Sedimentary organic matter preservation: an assessment and speculative synthesis. *Marine Chemistry*, 49(2-3): 81-115.

- Herring, J.R., 1985. Charcoal fluxes into sediments of the North Pacific Ocean: The Cenozoic record of burning. In: Sundquist, E.T., Broecker, W.S. [eds.] (Ed.), *The Carbon Cycle and Atmospheric CO₂: Natural Variations Archean to Present*. American Geophysical Union, Washington, D.C.
- Hockaday, W.C., Grannas, A.M., Kim, S., Hatcher, P.G., 2006. Direct molecular evidence for the degradation and mobility of black carbon in soils from ultrahigh-resolution mass spectral analysis of dissolved organic matter from a fire-impacted forest soil. *Organic Geochemistry*, 37(4): 501-510.
- Hockaday, W.C., Grannas, A.M., Kim, S., Hatcher, P.G., 2007. The transformation and mobility of charcoal in a fire-impacted watershed. *Geochimica Et Cosmochimica Acta*, 71(14): 3432-3445.
- Horvath, H., 1993. Atmospheric light absorption - a review. *Atmospheric Environment Part a-General Topics*, 27(3): 293-317.
- IPCC, 2007. *Fourth Assessment Report (AR4)*, Cambridge University Press.
- Jacobson, M.Z., 2001. Strong radiative heating due to the mixing state of black carbon in atmospheric aerosols. *Nature*, 409: 695-698.
- Jia, G., Peng, P.A., Zhao, Q., Jian, Z., 2003. Changes in terrestrial ecosystem since 30Ma in East Asia: Stable isotope evidence from black carbon in the South China Sea. *Geology*, 31(12): 1093-1096.
- Jiao, W. et al., 2012. PAHs in surface sediments from coastal and estuarine areas of the northern Bohai and Yellow Seas, China. *Environmental geochemistry and health*: 1-12.
- Jurado, E., Dachs, J., Duarte, C.M., Simo, R., 2008. Atmospheric deposition of organic and black carbon to the global oceans. *Atmospheric Environment*, 42(34): 7931-7939.
- Kang, Y. et al., 2009. Black carbon and polycyclic aromatic hydrocarbons (PAHs) in surface sediments of China's marginal seas. *Chinese Journal of Oceanology and Limnology*, 27(2): 297-308.
- Katsoyiannis, A., Sweetman, A.J., Jones, K.C., 2011. PAH Molecular Diagnostic Ratios Applied to Atmospheric Sources: A Critical Evaluation Using Two Decades of Source Inventory and Air Concentration Data from the UK. *Environmental Science & Technology*, 45(20): 8897-8906.
- Keenan, H.E., Bangkedphol, S., Sakultantimetha, A., Songsasen, A., 2010. The ecological complexity of the Thai-Laos Mekong River: II. Metals and polyaromatic hydrocarbons (PAHs) monitoring, modelling and environmental fate. *Journal of Environmental Science and Health Part A*, 45(13): 1674-1680.
- Keiluweit, M., Nico, P.S., Johnson, M.G., Kleber, M., 2010. Dynamic Molecular Structure of Plant Biomass-Derived Black Carbon (Biochar). *Environmental Science & Technology*, 44(4): 1247-1253.
- Kercher, A.K., Nagle, D.C., 2003. Microstructural evolution during charcoal carbonization by X-ray diffraction analysis. *Carbon*, 41(1): 15-27.
- Kim, S., Kaplan, L.A., Benner, R., Hatcher, P.G., 2004. Hydrogen-deficient molecules in natural riverine water samples-evidence for the existence of black carbon in DOM. *Marine Chemistry*, 92(1-4): 225-234.
- Knicker, H., 2007. How does fire affect the nature and stability of soil organic nitrogen and carbon? A review. *Biogeochemistry*, 85(1): 91-118.

- Knicker, H., Nikolova, R., Dick, D.P., Dalmolin, R.S.D., 2012. Alteration of quality and stability of organic matter in grassland soils of Southern Brazil highlands after ceasing biannual burning. *Geoderma*, 181: 11-21.
- Koch, B.P., Dittmar, T., 2006. From mass to structure: an aromaticity index for high-resolution mass data of natural organic matter. *Rapid Communications in Mass Spectrometry*, 20: 926-932.
- Kralovec, A.C., Christensen, E.R., Van Camp, R.P., 2002. Fossil Fuel and Wood Combustion As Recorded by Carbon Particles in Lake Erie Sediments 1850-1998. *Environmental Science and Technology*, 36: 1405-1413.
- Krull, E.S. et al., 2003. ¹³C-depleted charcoal from C4 grasses and the role of occluded carbon in phytoliths. *Organic Geochemistry*, 34: 1337-1352.
- Kuhlbusch, T.A., Crutzen, P.J., 1995. Toward a global estimate of black carbon in residues of vegetation fires representing a sink of atmospheric CO₂ and a source of O₂. *Global Biogeochemical Cycles*, 9(4): 491 - 501.
- Kuhlbusch, T.A.J. et al., 1996. Black carbon formation by savanna fires: measurements and implications for the global carbon cycle. *Journal of Geophysical Research*, 101: 23,651 - 23,665.
- Kuo, L.J., Herbert, B.E., Louchouart, P., 2008. Can levoglucosan be used to characterize and quantify char/charcoal black carbon in environmental media? *Organic Geochemistry*, 39(10): 1466-1478.
- Kuo, L.J. et al., 2011. Combustion-derived substances in deep basins of Puget Sound: Historical inputs from fossil fuel and biomass combustion. *Environmental Pollution*, 159(4): 983-990.
- Lee, E.J., 2005. Studies of dissolved black carbon and organic matter utilizing electrospray ionization mass spectrometry MS Thesis, The Ohio State University, Columbus, OH, 1-117 pp.
- Lehmann, J., 2007. A handful of carbon. *Nature*, 447(7141): 143-144.
- Lehmann, J. et al., 2005. Near-edge X-ray absorption fine structure (NEXAFS) spectroscopy for mapping nano-scale distribution of organic carbon forms in soil: Application to black carbon particles. *Global Biogeochemical Cycles*, 19(1).
- Leithold, E., Blair, N., Perkey, D., 2006. Geomorphologic controls on the age of particulate organic carbon from small mountainous and upland rivers. *Global Biogeochemical Cycles*, 20: Art. No. GB3022 SEP 9 2006.
- Li, B. et al., 2012a. Spatial distribution and source apportionment of PAHs in surficial sediments of the Yangtze Estuary, China. *Marine Pollution Bulletin*.
- Li, X. et al., 2012b. Composition, Abundance and Age of Total Organic Carbon in Surface Sediments from the Inner Shelf of the East China Sea. *Marine chemistry*.
- Lima, A.L.C., Farrington, J.W., Reddy, C.M., 2005. Combustion-derived polycyclic aromatic hydrocarbons in the environment - A review. *Environmental Forensics*, 6(2): 109-131.
- Lioussé, C., Cachier, H., Jennings, S.G., 1993. Optical and thermal measurements of black carbon aerosol content in different environments - variation of the specific attenuation cross-section (σ). *Atmospheric Environment Part a-General Topics*, 27(8): 1203-1211.

- Lobbés, J.M., Fitznar, H.P., Kattner, G., 2000. Biogeochemical characteristics of dissolved and particulate organic matter in Russian rivers entering the Arctic Ocean. *Geochimica et Cosmochimica Acta*, 64(17): 2973-2983.
- Lohmann, R. et al., 2009. Fluxes of soot black carbon to South Atlantic sediments. *Global Biogeochemical Cycles*, 23, doi:10.1029/2008GB003253.
- Louchouart, P. et al., 2007. Elemental and Molecular Evidence of Soot- and Char-Derived Black Carbon Inputs to New York City's Atmosphere during the 20th Century. *Environmental Science and Technology*, 41: 82-87.
- Ludwig, W., Amiotte Suchet, P., Probst, J.L., 1996a. River discharges of carbon to the world's oceans: determining local inputs of alkalinity and of dissolved and particulate organic carbon. *Sciences de la terre et des planÈtes (Comptes rendus de l'AcadÈmie des sciences)*, 323: 1007-1014.
- Ludwig, W., AmiotteSuchet, P., Probst, J.L., 1996b. River discharges of carbon to the world's oceans: Determining local inputs of alkalinity and of dissolved and particulate organic carbon. *Comptes Rendus De L Academie Des Sciences Serie Ii Fascicule a-Sciences De La Terre Et Des Planetes*, 323(12): 1007-1014.
- Ma, Y. et al., 2008. Distribution, sources, and potential risk of polycyclic aromatic hydrocarbons (PAHs) in drinking water resources from Henan Province in middle of China. *Environmental monitoring and assessment*, 146(1): 127-138.
- Macdonald, R. et al., 1998. A sediment and organic carbon budget for the Canadian Beaufort Shelf. *Marine geology*, 144(4): 255-273.
- Major, J., Lehmann, J., Rondon, M., Goodale, C., 2010. Fate of soil-applied black carbon: downward migration, leaching and soil respiration. *Global Change Biology*, 16(4): 1366-1379.
- Mannino, A., 2000. Discussion at USGS Reston on Radiocarbon Ages of BC in Delaware River HMW DOM.
- Mannino, A., Harvey, H.R., 2004. Black carbon in estuarine and coastal ocean dissolved organic matter. *Limnology and Oceanography*, 49(1): 735-740.
- Marlon, J.R. et al., 2009a. Climate and human influences on global biomass burning over the past two millennia (vol 1, pg 697, 2008). *Nature Geoscience*, 2(4): 307-307.
- Marlon, J.R. et al., 2009b. Wildfire responses to abrupt climate change in North America. *Proceedings of the National Academy of Sciences of the United States of America*, 106(8): 2519-2524.
- Martins, J.V. et al., 1998a. Effects of black carbon content, particle size, and mixing on light absorption by aerosols from biomass burning in Brazil. *Journal of Geophysical Research-Atmospheres*, 103(D24): 32041-32050.
- Martins, J.V., Hobbs, P.V., Weiss, R.E., Artaxo, P., 1998b. Sphericity and morphology of smoke particles from biomass burning in Brazil. *Journal of Geophysical Research-Atmospheres*, 103(D24): 32051-32057.
- Maruya, K., Loganathan, B., Kannan, K., McCumber-Kahn, S., Lee, R., 1997. Organic and organometallic compounds in estuarine sediments from the Gulf of Mexico (1993ñ1994). *Estuaries and Coasts*, 20(4): 700-709.
- Masiello, C., Druffel, E., 2003a. Organic and black carbon ¹³C and ¹⁴C through the Santa Monica Basin sediment oxic-anoxic transition. *Geophysical research letters*, 30(4): 1185.

- Masiello, C.A., 1999. Radiocarbon measurements of black carbon in sediments and a small river. PhD Thesis, University of California, Irvine, 150 pp.
- Masiello, C.A., 2004. New directions in black carbon organic geochemistry. *Marine Chemistry*, 92(1-4): 201-213.
- Masiello, C.A., Druffel, E.R.M., 1998. Black carbon in deep sea sediments. *Science*, 280: 1911-1913.
- Masiello, C.A., Druffel, E.R.M., 2001. Carbon isotope geochemistry of the Santa Clara River. *Global Biogeochemical Cycles*, 15: 407 - 416.
- Masiello, C.A., Druffel, E.R.M., 2003b. Organic and black carbon C-13 and C-14 through the Santa Monica Basin sediment oxic-anoxic transition. *Geophysical Research Letters*, 30(4).
- Masiello, C.A., Druffel, E.R.M., Currie, L.A., 2002. Radiocarbon measurements of black carbon in aerosols and ocean sediments. *Geochimica et Cosmochimica Acta*, 66(6): 1025-1036.
- Menon, S., Hansen, J.E., Nazarenki, L., Luo, Y., 2002. Climate effects of black carbon aerosols in China and India. *Science*, 297: 2250-2253.
- Meybeck, M., Ragu, A., 1997. Presenting the GEMS-GLORI, a compendium of world river discharge to the oceans, Freshwater Contamination (Proceedings of Rabat Symposium S4, Aprii-May 1997) IAHS Publ. no. 243., PANGAEA.
- Middleburg, J.J., 1989. A simple rate model for organic matter decomposition in marine sediments. *Geochimica et Cosmochimica Acta*, 53: 1577-1581.
- Middleburg, J.J., Nieuwenhuize, J., van Breugel, P., 1999. Black carbon in marine sediments. *Marine Chemistry*, 65: 245 - 252.
- Milliman, J.D., Syvitski, J.P.M., 1992. Geomorphic-tectonic control of sediment discharge to the ocean: the importance of small mountainous rivers. *Journal of Geology*, 100: 525-544.
- Mitra, S., Bianchi, T.S., McKee, B.A., Sutula, M., 2002a. Black carbon from the Mississippi River: Quantities, sources, and potential implications for the global carbon cycle. *Environmental Science and Technology*, 36(11): 2296-2302.
- Mitra, S., Zimmerman, A.R., Hunsinger, G.B., Willard, D., Dunn, J.C., 2009. A Holocene record of climate-driven shifts in coastal carbon sequestration. *Geophysical Research Letters*, 36: doi:10.1029/2008GL036875.
- Moosmuller, H., Chakrabarty, R.K., Arnott, W.P., 2009. Aerosol light absorption and its measurement: A review. *Journal of Quantitative Spectroscopy & Radiative Transfer*, 110(11): 844-878.
- Nemirovskaya, I., 2007. Hydrocarbons in the water and bottom sediments of a region with continuous petroleum contamination. *Geochemistry International*, 45(7): 638-651.
- Nguyen, B.T., Lehmann, J., 2009. Black carbon decomposition under varying water regimes. *Organic Geochemistry*, 40(8): 846-853.
- Nguyen, B.T., Lehmann, J., Hockaday, W.C., Joseph, S., Masiello, C.A., 2010. Temperature sensitivity of black carbon decomposition and oxidation. *Environmental Science & Technology*, 44(9): 3324-3331.
- Nguyen, B.T. et al., 2008. Long-term black carbon dynamics in cultivated soil. *Biogeochemistry*, 89(3): 295-308.

- Nguyen, T.H., Brown, R.A., Ball, W.P., 2004. An evaluation of thermal resistance as a measure of black carbon content in diesel soot, wood char, and sediment. *Organic Geochemistry*, 35: 217-234.
- Novakov, T., 1984. The role of soot and primary oxidants in atmospheric chemistry. *Science of the Total Environment*, 36(JUN): 1-10.
- Novakov, T. et al., 2003. Large historical changes of fossil-fuel black carbon aerosols. *Geophysical Research Letters*, 30(6): 57-61.
- Ogren, J.A., Charlson, R.J., 1983. Elemental carbon in the atmosphere: cycle and lifetime. *Tellus B*, 35B(4): 241-254.
- Ogren, J.A., Groblicki, P.J., Charlson, R.J., 1984. Measurement of the removal rate of elemental carbon from the atmosphere. *Science of the Total Environment*, 36(JUN): 329-338.
- Ohlson, M., Tryterud, E., 2000. Interpretation of the charcoal record in forest soils: forest fires and their production and deposition of macroscopic charcoal. *Holocene*, 10(4): 519-525.
- Overton, E., Ashton, B., Miles, M., 2004. Historical polycyclic aromatic and petrogenic hydrocarbon loading in Northern Central Gulf of Mexico shelf sediments. *Marine Pollution Bulletin*, 49(7): 557-563.
- Paolini, J., 1995. Particulate organic carbon and nitrogen in the Orinoco River (Venezuela). *Biogeochemistry*, 29(1): 59-70.
- Patterson, W.A., Edwards, K.J., Maguire, D.J., 1987. Microscopic charcoal as a fossil indicator of fire. *Quaternary Science Reviews*, 6(1): 3-23.
- Penner, J.E., Eddleman, H., 1993. Towards the development of a global inventory for black carbon emissions. *Atmospheric Environment*, 27A(8): 1277-1295.
- Penner, J.E.e.a., 2001. *Climate Change 2001: The Scientific Basis*, (Cambridge Univ. Press, Cambridge, 2001), pp. 289–348.). Cambridge University Press, Cambridge, 289-348 pp.
- Power, M.J. et al., 2008. Changes in fire regimes since the Last Glacial Maximum: an assessment based on a global synthesis and analysis of charcoal data. *Climate Dynamics*, 30: 887-907.
- Preston, C.M., Schmidt, M.W.I., 2006. Black (pyrogenic) carbon: a synthesis of current knowledge and uncertainties with special consideration of boreal regions. *Biogeosciences*, 3(4): 397-420.
- Ramanathan, V., Carmichael, G., 2008. Global and regional climate changes due to black carbon. *Nature Geoscience*, 1(4): 221-227.
- Raymond, P.A., Bauer, J.E., 2001. Riverine export of aged terrestrial organic matter to the North Atlantic Ocean. *Nature*, 409: 497-500.
- Reddy, C.M. et al., 2002. Radiocarbon as a tool to apportion the sources of polycyclic aromatic hydrocarbons and black carbon in environmental samples. *Environmental Science and Technology*, 36: 1774 - 1782.
- Richey, J.E. et al., 1990. Biogeochemistry of carbon in the Amazon River. *Limnology and Oceanography*, 35(2): 352-371.
- Rivas, Y., Matus, F., Rumpel, C., Knicker, H., Garrido, E., 2012. Black carbon contribution in volcanic soils affected by wildfire or stubble burning. *Organic Geochemistry*, 47: 41-50.

- Robock, A., 1984. Snow and ice feedbacks prolong effects of nuclear winter. *Nature*, 310(5979): 667-670.
- Rogers, K.G., Goodbred, S.L., 2010. Mass failures associated with the passage of a large tropical cyclone over the Swath of No Ground submarine canyon (Bay of Bengal). *Geology*, 38(11): 1051-U123.
- Rosen, H., Novakov, T., 1977. Raman scattering and the characterisation of atmospheric aerosol particles. *Nature*, 266: 708-710.
- Rumpel, C., Ba, A., Darboux, F., Chaplot, V., Planchon, O., 2009. Erosion budget and process selectivity of black carbon at meter scale. *Geoderma*, 154(1-2): 131-137.
- Sanchez-Garcia, L., Cato, I., Gustafsson, O., 2012. The sequestration sink of soot black carbon in the Northern European Shelf sediments. *Global Biogeochemical Cycles*, 26.
- Santschi, P., Presley, B., Wade, T., Garcia-Romero, B., Baskaran, M., 2001. Historical contamination of PAHs, PCBs, DDTs, and heavy metals in Mississippi River Delta, Galveston Bay and Tampa Bay sediment cores. *Marine Environmental Research*, 52(1): 51-79.
- Schmidt, M.W.I., Noack, A.G., 2000. Black carbon in soils and sediments: analysis, distribution, implications, and current challenges. *Global Biogeochemical Cycles*, 14(3): 777-793.
- Schmidt, M.W.I., Skjemstad, J.O., Jager, C., 2002. Carbon isotope geochemistry and nanomorphology of soil black carbon: Black chernozemic soils in central Europe originate from ancient biomass burning. *Global Biogeochemical Cycles*, 16(4).
- Skjemstad, J.O., Taylor, J.A., Smernik, R.J., 1999. Estimation of charcoal (Char) in soils. *Communications In Soil Science and Plant Analysis*, 30(15, 16): 2283-2298.
- Smith, D.M., Griffin, J.J., Goldberg, E.D., 1973. Elemental carbon in marine sediments: a baseline for burning. *Nature*, 241: 268-270.
- Smith, D.M., Griffin, J.J., Goldberg, E.D., 1975. Spectrometric method for quantitative-determination of elemental carbon. *Analytical Chemistry*, 47(2): 233-238.
- Stanmore, B.R., Brilhac, J.F., Gilot, P., 2001. The oxidation of soot: a review of experiments, mechanisms and models. *Carbon*, 39(15): 2247-2268.
- Stubbins, A. et al., 2012a. Anthropogenic aerosols as a source of ancient dissolved organic matter in glaciers. *Nature Geoscience*, 5: 198-201.
- Stubbins, A., Niggemann, J., Dittmar, T., 2012b. Photo-lability of deep ocean dissolved black carbon. *Biogeosciences*, 9(5): 1661-1670.
- Stubbins, A. et al., 2010. Illuminated darkness: Molecular signatures of Congo River dissolved organic matter and its photochemical alteration as revealed by ultrahigh precision mass spectrometry. *Limnology and Oceanography*, 55(4): 1467-1477.
- Suman, D.O., 1986. Charcoal production from agricultural burning in Central Panama and its deposition in the sediments of the Gulf of Panama. *Environmental Conservation*, 13(1): 51-60.
- Suman, D.O., Kuhlbusch, T.A.J., Lim, B., 1997. Marine sediments: a reservoir for black carbon and their use as spatial and temporal records of combustion. In: Clark, J., Cachier, H., Goldammer, J.G., Stocks, B. (Eds.), *Sediment records of biomass burning and global change*. NATO ASI Series. Springer-Verlag, Berlin, pp. 271-293.

- Thevenon, F., Bard, E., Williamson, D., Beaufort, L., 2004. A biomass burning record from the West Equatorial Pacific over the last 360 ky: methodological, climatic and anthropic implications. *Palaeogeography Palaeoclimatology Palaeoecology*, 213(1-2): 83-99.
- Turekian, V.C. et al., 1996. Bulk and compound-specific isotopic characterization of the products of biomass burning: laboratory studies. In: Levine, J. (Ed.), *Biomass Burning and Global Change*. MIT Press, Cambridge, pp. 422-427.
- Unger, D., Gaye-Haake, B., Neumann, K., Gebhardt, A.C., Ittekkot, V., 2005. Biogeochemistry of suspended and sedimentary material in the Ob and Yenisei rivers and Kara Sea: amino acids and amino sugars. *Continental Shelf Research*, 25(4): 437-460.
- Wang, X.C., Ma, H.Q., Li, R.H., Song, Z.S., Wu, J.P., 2012. Seasonal fluxes and source variation of organic carbon transported by two major Chinese Rivers: The Yellow River and Changjiang (Yangtze) River. *Global Biogeochemical Cycles*, 26.
- Warren, S.G., Wiscombe, W.J., 1979. New model for the spectral albedo of snow. *Bulletin of the American Meteorological Society*, 60(7): 846-846.
- Warren, S.G., Wiscombe, W.J., 1980. A model for the spectral albedo of snow .2. Snow containing atmospheric aerosols. *Journal of the Atmospheric Sciences*, 37(12): 2734-2745.
- Warren, S.G., Wiscombe, W.J., 1981. Comment on radiative properties of snow for clear sky solar-radiation. *Cold Regions Science and Technology*, 5(2): 177-180.
- Warren, S.G., Wiscombe, W.J., 1985. Dirty snow after nuclear war. *Nature*, 313(6002): 467-470.
- Widory, D., 2006. Combustibles, fuels and their combustion products: A view through carbon isotopes. *Combustion Theory and Modelling*, 10(5): 831-841.
- Williams, P.M., Druffel, E.R.M., 1987. Radiocarbon in dissolved organic matter in the Central North Pacific Ocean. *Nature*, 330(6145): 246-248.
- Wolbach, W.S., Anders, E., 1989. Elemental carbon in sediments: Determination and isotopic analysis in presence of kerogen. *Geochimica et Cosmochimica Acta*, 53: 1637-1647.
- Wozniak, A.S., Bauer, J.E., Sleighter, R.L., Dickhut, R.M., Hatcher, P.G. 2008. Technical Note: Molecular characterization of aerosol-derived water soluble organic carbon using ultrahigh resolution electrospray ionization Fourier transform ion cyclotron resonance mass spectrometry. *Atmos. Chem. Phys.*, 8: 5099–5111.
- Wu, Y., Zhang, J., Mi, T., Li, B., 2001. Occurrence of *n*-alkanes and polycyclic aromatic hydrocarbons in the core sediments of the Yellow Sea. *Marine chemistry*, 76(1): 1-15.
- Xu, B.Q. et al., 2009. Black soot and the survival of Tibetan glaciers. *Proceedings of the National Academy of Sciences of the United States of America*, 106(52): 22114-22118.
- Xu, J. et al., 2007. Polycyclic aromatic hydrocarbons in the surface sediments from Yellow River, China. *Chemosphere*, 67(7): 1408-1414.
- Yunker, M.B., Macdonald, R.W., Cretney, W.J., Fowler, B.R., McLaughlin, F.A., 1993. Alkane, terpene and polycyclic aromatic hydrocarbon geochemistry of the

- Mackenzie River and Mackenzie Shelf: riverine contributions to Beaufort Sea coastal sediment. *Geochimica et Cosmochimica Acta*, 57(13): 3041-3061.
- Yunker, M.B., Macdonald, R.W., Snowdon, L.R., Fowler, B.R., 2011. Alkane and PAH biomarkers as tracers of terrigenous organic carbon in Arctic Ocean sediments. *Organic Geochemistry*, 42(9): 1109-1146.
- Yunker, M.B. et al., 2002. PAHs in the Fraser River basin: a critical appraisal of PAH ratios as indicators of PAH source and composition. *Organic Geochemistry*, 33(4): 489-515.
- Zimmerman, A., 2010. Abiotic and microbial oxidation of laboratory-produced black carbon (biochar). *Environmental Science and Technology*, 44: 1295-1301.
- Zimmerman, A.R., Gao, B., in press. The Stability of Biochar in the Environment. In: Ladygina, N., Rineau, F. (Eds.), *Biochar and Soil Biota*. CRC Press, (A Taylor & Francis Group Co.), Boca Raton, FL.
- Zimmerman, A.R., Gao, B., Ahn, M.-Y., 2011. Positive and negative carbon mineralization priming effects among a variety of biochar-amended soils. *Soil Biology & Biochemistry*, 43: 1169– 1179.
- Ziolkowski, L., Druffel, E., 2010. Aged black carbon identified in marine dissolved organic carbon. *Geophys. Res. Lett.*, 37: L16601.