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DOI: <https://doi.org/10.1002/2016GL069253>

Posted at the Zurich Open Repository and Archive, University of Zurich

ZORA URL: <https://doi.org/10.5167/uzh-124368>

Journal Article

Accepted Version

Originally published at:

Hanke, Ulrich Michael; Eglinton, Timothy I; Braun, Ana L L; Reddy, Christopher M; Wiedemeier, Daniel B; Schmidt, Michael W I (2016). Decoupled sedimentary records of combustion: causes and implications. *Geophysical Research Letters*, 43(10):5098-5108.

DOI: <https://doi.org/10.1002/2016GL069253>

Decoupled Sedimentary Records of Combustion: Causes and Implications

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Key Points:

- Comparison of high temporal-resolution multi-proxy records of particulate pyrogenic carbon (PyC) in aquatic sediments
- Records of combustion are clearly decoupled in the (pre-)industrial era
- Aquatic sediments chronicle local and regional sources of combustion

Abstract

Pyrogenic carbon (PyC) is a collective term for carbon-rich residues comprised of a continuum of products arising from biomass burning and fossil-fuel combustion. PyC is ubiquitous in the environment where it can be transported by wind and water before being deposited in aquatic sediments. We compare results from four different methods used to trace PyC that were applied to a high-temporal resolution sedimentary record in order to constrain changes in PyC concentrations and fluxes over the past ~250 years. We find markedly discordant records for different PyC tracers, particularly during the pre-industrial age, implying different origins and modes of supply of sedimentary PyC. In addition to providing new insights into the composition of sedimentary combustion products, this study reveals that elucidation of past combustion processes, and development of accurate budgets of PyC production and deposition on local to regional scales requires careful consideration of both source characteristics and transport processes.

1 Introduction

Pyrogenic carbon (PyC), including charcoal and soot-like materials, is formed during the incomplete combustion of biomass or fossil fuels and largely consists of condensed polyaromatic structures [Antal and Gronli, 2003; Bird *et al.*, 2015]. The cluster size of PyC increases with temperature, with relative small aromatic domains (≤ 7 rings) at low production temperatures (< 400 °C) and highly condensed domains (> 19 rings) at higher temperatures (700°C) [McBeath *et al.*, 2011] depending on oxygen availability and reaction times [Bockhorn, 1994]. Low molecular weight carbon compounds can be building blocks for formation of soot particles [Pöschl and Shiraiwa, 2015].

PyC is a key component of the global carbon cycle due its slow intrinsic decomposition rate and its ubiquity in the environment [Bird *et al.*, 2015; Goldberg, 1985; Schmidt and Noack, 2000]. PyC originates from wildfires, land clearance, energy production, and from burning of fossil fuel in industry and transportation [Bond *et al.*, 2013; Goldberg *et al.*, 1981; Thevenon *et al.*, 2010]. PyC can subsequently be retained in soils, where its been reported to account for between < 1 -60 % of total organic carbon (TOC) [Preston and Schmidt, 2006], or transported over large distances by wind and water, eventually accumulating in aquatic sediments [Bird *et al.*, 2015]. Riverine transport alone is estimated to transfer 19-80 Tg yr⁻¹ of particulate PyC from terrestrial to marine systems [Bird *et al.*, 2015; Dittmar *et al.*, 2012; Gustafsson and Gschwend, 1998; Jaffé *et al.*, 2013; Sánchez-García *et al.*, 2013]. However, data for lacustrine and estuarine systems remains scarce despite their potential importance as a long-term sink for PyC [Bird *et al.*, 2015; Thevenon *et al.*, 2010]. This information void needs to be filled before modern-day PyC budgets can be calculated and past PyC production and cycling assessed. To address the latter, sedimentary sequences, which integrate signals derived from small catchment areas can yield detailed records of PyC production and deposition that can be linked to local land-use history. Lima *et al.* [2003], for example, developed a detailed record spanning the 19th and 20th Centuries of combustion-derived polycyclic aromatic hydrocarbons (PAH) from Pettaquamscutt River basin sediments, and interpreted PAH fluxes in the context of the history of fuel consumption in the northeastern United States [Lima *et al.*, 2003]. The anoxic depositional setting of this location results in a laminated sediments that are conducive to reconstruction of historical records of combustion at both regional and local scales at high-temporal resolution [Hites *et al.*, 1980; Lima *et al.*, 2003; Lima *et al.*, 2005b].

Several studies have focused exclusively on PAHs as tracers of condensed PyC by virtue of their ease of extraction and analysis, as well as information embedded in their distributions and isotopic compositions regarding combustion sources [Hites *et al.*, 1980; Lima *et al.*, 2003; Wakeham *et al.*, 2004; Windsor and Hites, 1979]. Other tracers of PyC are now also commonly employed, and some studies have yielded down-core profiles that contrast with those of PAHs [Currie *et al.*, 2002; Elmquist *et al.*, 2007; Louchouart *et al.*, 2007; Muri *et al.*, 2006]. These discrepancies may be related to different modes of formation or transport of combustion products, or possibly to methodological artefacts. For example, PAHs are secondary products derived from condensation of initial low molecular weight combustion species [Masiello, 2004; McBeath and Smernik, 2009], with high-temperature combustion favouring the production of non-alkylated PAH assemblages [Blumer, 1976] that can coagulate to form soot-like structures [Bockhorn, 1994]. In contrast, charcoal comprises primary charred residues of biogenic materials. Compared to PAHs, charcoal and other products of incomplete combustion exhibit heterogeneous chemical characteristics (slightly charred to highly condensed; Figure S1, supporting information) [Bird *et al.*, 2015; Hammes *et al.*, 2007; Wiedemeier *et al.*, 2015]. These two types of PyC represent end-members in a continuum of combustion products that vary in abundance as a function of the sources and conditions of production as well as subsequent fates [Masiello, 2004]. Available methods capture different parts of this PyC continuum, e.g. operationally-defined fractions (Graphitic black carbon (PyC_{GBC}), Chemical thermal oxidation (PyC_{CTO}) and spheroidal carbonaceous particles (SCP) [Gélinas *et al.*, 2001; Gustafsson *et al.*, 2001; Rose, 1994]), chemically-defined (molecular) markers such as PAHs and benzene polycarboxylic acids (PyC_{BPCA}; Figure S1, supporting information) [Glaser *et al.*, 1998], or spectroscopic characteristics [Baldock and Smernik, 2002; Lehmann *et al.*, 2005]. Application of a broad suite of methods is thus favoured in order to constrain the abundance and characteristics of different PyC components (Table 1, supporting information). Operationally-defined fractions and chemically-defined species also offer the potential to derive other information (e.g. carbon isotopic composition) that can be used to constrain the provenance of PyC.

Quantifying the heterogeneous mixtures of PyC in environmental matrices and constraining its sources is a challenging but a necessary step in order to understand this important component in the global carbon cycle. In the present study, we compare sedimentary profiles of PAHs in anoxic and laminated sediments from the Pettaquamscutt River basin, RI, USA [Lima *et al.*, 2003] with those derived from application of three other methods used to quantify PyC at a bulk level aiming at to differentiate between local biomass burning in the catchment and regional-scale inputs from industrial sources of combustion. The goals were to: (1) comprehensively quantify and characterize PyC, (2) compare the information on PyC gleaned from different combustion tracers and proxies, and (3) assess local and regional inputs of PyC.

2 Materials and Methods

2.1 Study area

The Pettaquamscutt River estuary (Figure 1) is located in Washington County (southern Rhode Island, U.S.; 41.503816, -71.450545) and drains a small watershed (35 km²) with a North-South orientation [Orr and Gaines, 1973]. Present-day land cover mainly consists of oak tree forests, wetlands and open waters [Urish, 1991] with approximately 30 % as residential land. The geological setting underlying the Pettaquamscutt River basin is a Pennsylvanian (late

Carboniferous) meta-sedimentary rock formation consisting of sandstone, conglomerate, schist and graphite [*Hermes et al.*, 1994].

Ocean water flooded the Pettaquamscutt estuary and the upper two basins about 1700±300 years ago, leading to the formation of zoned circulation patterns with a vertical salinity gradient (10 ppt) that has caused oxygen depletion in basin bottom waters [*Hubeny et al.*, 2009]. Since then surface sediments have been anoxic, limiting microbial decomposition [*Orr and Gaines*, 1973] and preventing bioturbation by macrofauna [*Hubeny et al.*, 2009]. The water depth at the sampling location in the lower basin was 20m, with the oxic/anoxic transition zone in the water column between 3.5 to 6m and sulphidic waters below 6m [*O'Sullivan et al.*, 1997]. The high-temporal resolution sedimentary record stemming from the confined nature of the basin and the permanently anoxic conditions render it an ideal location for detailed historical reconstruction of PyC deposition.

2.2 Sampling

A series of freeze cores were collected in 1999 using a rectangular aluminium freeze corer (90 cm x 30 cm x 10 cm), dry ice and methanol [*Lima et al.*, 2003]. Sediment chronology was obtained from varve counting, and ^{137}Cs and ^{210}Pb profiles using with the constant rate of supply model [*Lima et al.*, 2005b]. In short, gamma spectrometry was used to detect ^{210}Pb (46.5 keV), ^{226}Ra (351 keV), ^{137}Cs (661 keV). Excess ^{210}Pb was calculated from total ^{210}Pb and supported ^{210}Pb activity. Limit of detection was 0.040 Bq for excess ^{210}Pb and 0.004 Bq for ^{137}Cs [*Lima et al.*, 2005b]. Sediment chronology (1999±0 to 1855±7) was compared for four key horizons (Chernobyl peak: 1987±1, maximum fallout: 1963±2, and hurricane-sand layers: 1954±3, 1938±4) [*Lima et al.*, 2005b]. Additionally, varve counting with radiograph images allowed extension of the sediment chronology back to 1750. Except for measurements of trace metals and PAHs, all analyses were carried out on four additional cores that were aligned via x-radiograph analysis (to cross-correlate laminae), sliced, freeze-dried, combined and subsequently stored in pre-combusted glass jars.

An elemental analyser (Fisons 1108) was used to determine TOC and nitrogen contents in the samples. In short, 2 mg dry sediment was weighed in silver capsules and acidified with 20 μL 2N HCL under vacuum, dried in the oven (50°C) and analysed [*Lima*, 2004]. Sediment *in situ* densities were calculated from dry densities measured using a 2 mL specific gravity bottle and porosity retrieved from weight percent of water [*Lima*, 2004]. The four PyC records differ in their temporal resolution. The PyC_{PAH} record comprises 77 sample depths to track rapid changes in U.S. fuel combustion, the PyC_{BPCA} record consists of 40 depths and those for PyC_{CTO} , PyC_{GBC} comprise 22 depths. The higher number of PyC_{BPCA} measurements were made to assess the coarser records.

2.3 Polycyclic aromatic hydrocarbons (PyC_{PAH})

PAHs are a group of chemical compounds consisting of fused benzene rings [*Mumtaz and George*, 1995] and in this study we focus on non-alkylated PAHs that represent intermediates in soot-like particle formation [*Blumer*, 1976]. The procedure for acquiring the high-resolution PAH record was previously described [*Lima et al.*, 2003]. Briefly, 0.5-1.5 g dry sediment were spiked with PAH internal standards and extracted by pressurized fluid extraction (1000 psi; 100

°C) with acetone and hexane (1:1). The aromatic hydrocarbon fraction was measured on an Agilent 6890/5970 GC/MSD System operating at 70 eV in selective ion monitoring (SIM) mode using a DB-XLB capillary column to separate compounds [Lima *et al.*, 2003]. Measurement precision is $\pm 5\%$ and the recovery of standards spiked into each sample is $84.9 \pm 9.9\%$ [Lima, 2004]. Total pyrogenic PAH concentrations reported included the following combustion-related species: fluorene, phenanthrene, anthracene, fluoranthene, pyrene, benz[*a*]anthracene, chrysene, benzo[*b*]fluoranthene, benzo[*k*]fluoranthene, benzo[*a*]pyrene, benzo[*e*]pyrene, dibenz[*a,h*]anthracene, indeno[*1,2,3-c,d*]pyrene, benzo[*g,h,i*]perylene, coronene, and together are defined here as PyC_{PAH} .

2.4 Benzene polycarboxylic acids (PyC_{BPCA})

Benzene polycarboxylic acids (BPCA) are molecular markers for PyC that are formed during a laboratory procedure for oxidative degradation of condensed polycyclic aromatic centres, thereby serving as measures for PyC [Glaser *et al.*, 1998]. BPCAs were obtained from the sediments by thermal digestion (170°C, 8 h) with nitric acid and quantified with liquid chromatography [Dittmar, 2008; Glaser *et al.*, 1998; Wiedemeier *et al.*, 2013] using an Agilent 1290 infinity HPLC system and Agilent Poroshell 120 column. Trimellitic acid (Fluka 92119), hemimellitic acid (Aldrich 51520), pyromellitic acid (Fluka 83181), pentacarboxylic acid (Aldrich S437107), mellitic acid (Aldrich M2705) were used as external standards in a seven-point calibration curve (2-200 ng μl^{-1}). The measurement precision of PyC_{BPCA} is, on average, $\pm 5\%$ [Wiedemeier *et al.*, 2013]. For stoichiometric reasons, PyC_{BPCA} accounts for a maximum of about 25 % of the total PyC [Brodowski *et al.*, 2005; Schneider *et al.*, 2011]

2.5 Chemical thermal oxidation (PyC_{CTO})

This methodological approach targets forms of PyC that exhibit a high degree of condensation, such as soot [Gustafsson *et al.*, 2001; Gustafsson *et al.*, 1997]. In short, dried (60°C) and finely-ground sediments were weighed (about 200 $\mu\text{g C}$) in Ag capsules. Then inorganic carbon was removed *in situ* by microacidification with liquid 1N HCl and dried at 60°C again. Residual sediment was then spread in a thin layer on a pre-combusted watch glass and heated to 375 °C in a programmable oven (Fisher Scientific IsoTemp) for 24 hours in the presence of excess oxygen. The remaining carbon, subsequently measured with an elemental analyser (Fisons 1108) was defined as PyC_{CTO} . Generally, the precision of measurements is reported with a relative standard deviation of 1-3% [Gustafsson *et al.*, 1997]. PyC_{CTO} concentrations may be altered due to additional charring of labile organic matter during the isolation procedure [Accardi-Dey, 2003; Nguyen *et al.*, 2004] and can be overestimated by as much as 3 orders of magnitude [Gélinas *et al.*, 2001].

2.6 Graphitic black carbon (PyC_{GBC})

Measurement of graphitic black carbon (PyC_{GBC}) is based on a modified PyC_{CTO} protocol and aims at reducing the potential formation of condensation products resulting from charring of non-PyC compounds during isolation [Reddy *et al.*, 2002], and thus preventing an overestimation of PyC in environmental matrices. The sediment samples were first treated with 1N HCl and 10% HF, then hydrolysable organic matter was stepwise removed with 2N, 4N and 6N trifluoroacetic acid [Gélinas *et al.*, 2001]. The resulting material was then thermally treated as

for the PyC_{CTO} method, with remaining carbon defined here as PyC_{GBC}. The thermal oxidation including the chemical pre-treatment shows a coefficient of variation of <1% [Gélinas *et al.*, 2001]. However, the chemical pre-treatment enhances the physical loss of sample material of about 15% [Gélinas *et al.*, 2001].

2.7 Data analysis

PyC_{PAH} when absorbed to other materials can be transported through the atmosphere before deposition and temporary storage in intermediate reservoirs [Dachs and Eisenreich, 2000; Koelmans *et al.*, 2006; Wozniak *et al.*, 2011]. We assume that particulate PyC can subsequently be transported from soils in the catchment to the sedimentary archive via surface runoff or soils erosion. Aquatic sediments and associated PyC may also experience short-term storage in sediments on the basin flanks before eventual accumulation in the depocenter. Lima (2004) revised the ²¹⁰Pb chronology for an excess accumulation from the basin periphery and determined a focusing factor of 1.3 based on the atmospheric ²¹⁰Pb inventory measured in New Haven, CT (38.6 dpm cm⁻²) [Turekian *et al.*, 1983] and the vertically integrated unsupported ²¹⁰Pb calculated for Pettaquamscutt River sediments (51.6 dpm cm⁻²) [Lima, 2004]. To account for this phased particle (and associated PyC) deposition, translocation and accumulation, we applied a 20-year moving average on PyC_{PAH}, PyC_{BPCA}, PyC_{CTO}, PyC_{GBC}. This is consistent with a deposition delay of nearly two decades indicated by the high-resolution records of ¹³⁷Cs [Lima *et al.*, 2005b] for the same sediments.

Similarities among the different PyC records were also assessed statistically (Figure 2e) via computing distances for cross-correlations of the four PyC records [Montero and Vilar, 2014]. Using classical multi-dimensional scaling [Cox and Cox, 2001], these distances were represented in two dimensions. Missing data points in the records were replaced with values from linear interpolations prior to these calculations.

3 Results

A comprehensive examination of the PyC_{PAH} record (1750-1999) is described by Lima *et al.* (2003). Individual values of PyC_{BPCA} (1754-1998), PyC_{GBC} (1789-1999) and PyC_{CTO} (1789-1999), as well as data on nitrogen content of residues can be found in Table S3-S4 (supporting information). Corresponding 20-year moving averages of the four PyC methods are shown in Figure 2a-d. We also calculated fluxes, derived from PyC concentrations (g kg⁻¹), deposition rates (g m⁻² yr⁻¹) and *in situ* densities (kg m⁻³). As PyC concentrations are reported more frequently in literature, and since fluxes exhibit very similar patterns, we focus on the former values in discussion of the data.

3.1 Polycyclic aromatic hydrocarbons (PyC_{PAH})

Total PyC_{PAH} concentrations vary between 0.00008 g kg⁻¹ (mid-1700s) and 0.0113 g kg⁻¹ (1956±3). Prior to 1842, PyC_{PAH} concentrations remain at background values (≤0.000136 g kg⁻¹), and then increase, concomitant with an increasing consumption of fossil fuels at the onset of industrialization. A first peak in PyC_{PAH} concentrations appears in 1927±4 (0.00676 g kg⁻¹). After a sudden drop during the Great Depression years (1930s), PyC_{PAH} concentrations increase again and reach an overall maximum (0.00919 g kg⁻¹) in 1959±3. Thereafter, the concentrations

tend to decrease, punctuated by another small peak prior to the Organization of Petroleum Exporting Countries (OPEC) oil embargo in 1973±2, and reach a minimum (0.0051 g kg⁻¹) in 1996±1. The slight increase (0.0007 g kg⁻¹) between 1996±1 and 1999±0 may reflect urbanization of the local area [Hubeny *et al.*, 2009].

The PyC_{PAH} record parallels the record of industrialization of the northeastern United States during the 19th and 20th centuries, particularly with respect to the usage of fossil fuels. The relative proportions of 15 different PAHs (and corresponding source diagnostic ratios) indicate minor contributions of petroleum-associated hydrocarbons throughout the entire record, implying that combustion is the primary source of PAHs [Lima *et al.*, 2003]. Moreover, the PAH are considered to be predominantly supplied via atmospheric deposition rather than urban surface run-off as indicated by the absence of an unresolved complex mixture in the sample matrix [Lima *et al.*, 2003].

3.2 Benzene polycarboxylic acids (PyC_{BPCA})

PyC_{BPCA} concentrations vary between 0.25 g kg⁻¹ (1993±1) and 1.43 g kg⁻¹ (mid-1700s). They exhibit a slight decrease from the earliest portion of the record, reaching a broad minimum in the mid-1800s (0.83 g kg⁻¹), before returning to a second maximum in 1920s (1.40 g kg⁻¹). Concentrations then tend to decrease monotonically approaching the end of the 20th Century (0.48 g kg⁻¹).

3.3 Chemical thermal oxidation (PyC_{CTO})

Concentrations of PyC_{CTO} vary between 4.70 g kg⁻¹ (1860±7) and 6.10 g kg⁻¹ (1931±4). In the late 18th Century, concentrations of PyC_{CTO} exhibit intermediate values (5.60 g kg⁻¹) before decreasing slightly until the 1860s (4.70 g kg⁻¹). Concentrations then exhibit a systematic increase to a maximum in 1930s (6.10 g kg⁻¹). Thereafter the PyC_{CTO} record remains relatively constant without any distinct trend until the end of the 20th Century. Nitrogen concentrations (Table S4, supporting information) in the thermal residues are uniform (0.9-1.0 g kg⁻¹), and corresponding C/N ratios of PyC_{CTO} vary between 5.1 and 7.0. These systematically low C/N ratios (Figure S4, supporting information) point towards the formation of artefacts due to charring of nitrogen-containing non-pyrogenic macromolecules during the analysis [Accardi-Dey, 2003]. Thus, we do not discuss the results of PyC_{CTO} further.

3.4 Graphitic black carbon (PyC_{GBC})

PyC_{GBC} concentrations vary from 0.97 g kg⁻¹ (1997±1) to 2.10 g kg⁻¹ (1938±4). PyC_{GBC} concentrations increase from 1.20 g kg⁻¹ in the late 18th Century, to 1.60 g kg⁻¹ in early 19th Century. Concentrations then gradually decline to a minimum of 1.05 g kg⁻¹ at the end of 19th Century before increasing again to a broad maximum in the mid-20th Century (2.10 g kg⁻¹). PyC_{GBC} concentrations then markedly decrease towards the present day (0.99 g kg⁻¹). Nitrogen contents (Table S4, supporting information) in the thermal residues vary between 0.03 and 0.08 g kg⁻¹, and corresponding C/N ratios of PyC_{GBC} vary between 16.3 and 48.8. Compared to PyC_{CTO} these wider and generally higher C/N ratios indicate lesser production of charring artefacts during the isolation of PyC_{GBC}.

4 Discussion

There are relatively few reported studies in which historical records of combustion have been developed based on more than one tracer of PyC. The comparability of different PyC approaches has, up to now, mainly been reported for standard materials, e.g. *Currie et al.* [2002]; *Hammes et al.* [2007]; *Roth et al.* [2012]. In particular, investigations that compare and contrast PyC records in coastal sediments are scarce [*Elmqvist et al.*, 2007; *Louchouart et al.*, 2007; *Muri et al.*, 2006], yet understanding transport and sequestration of PyC in these settings is crucial for developing insights into global-scale biogeochemical cycles of pyrogenic and other forms of reduced (organic) carbon.

Distinctive deposition patterns of PyC in continental shelf sediments of the northeastern United States have been shown previously, e.g. *Gustafsson and Gschwend* [1997; 1998]; *Hites et al.* [1980]; *Windsor and Hites* [1979]. Prevailing (south)westerly winds transport particulate PyC over long distances and link the coastal areas with industrial and urban regions inland.

4.1 Fidelity and representativity of PyC data

While varying sensitivity to different PyC sources and characteristics complicates assessment of the comparability of the four PyC approaches, information derived from these approaches sheds light on dynamics of different components of the PyC continuum, and collectively they offer more comprehensive insights into this important carbon pool. In this regard, the Pettaquamscutt PyC_{PAH} record mirrors the overall history of energy consumption in the northeastern United States, faithfully documenting key episodes and transitions in usage of different types of fuel [*Lima et al.*, 2003]. However, PAHs represent trace PyC constituents, as is evident from their concentrations, and so it is important to compare profiles with other proxies that are more representative of bulk PyC (PyC_{BPCA} , PyC_{GBC}).

The four PyC methodologies used in this study are assumed to capture different portions of the combustion continuum [*Hammes et al.*, 2007]. For the Pettaquamscutt River record, each PyC proxy yields different concentrations (Figure 2). These discrepancies likely reflect the different analytical windows inherent in each measurement approach, modes of PyC formation during burning/combustion (e.g. charring versus condensation reactions), but also methodological artefacts.

4.2 PyC contents in coastal sediments

PyC_{BPCA} concentrations in the Pettaquamscutt River sediments are similar to those found in coastal surface sediments in SW Spain ($0.33\text{-}1.48 \text{ g kg}^{-1}$; $n=4$) [*De la Rosa et al.*, 2011] or ($0.1\text{-}2.3 \text{ g kg}^{-1}$; $n=15$) [*Sánchez-García et al.*, 2013] but higher than those found in the northeast Pacific Ocean sediments ($0.5\text{-}1.1 \text{ g kg}^{-1}$; $n=14$) [*Coppola et al.*, 2014].

For PyC_{GBC} , concentrations are higher compared to those reported for a terrestrial near-shore to marine offshore transect at Washington coast ($0.09\text{-}0.71 \text{ g kg}^{-1}$) [*Dickens et al.*, 2004] but similar to sediments from Lake Washington ($0.22\text{-}0.58 \text{ g kg}^{-1}$; $n=20$) [*Wakeham et al.*, 2004] and from the inner shelf of SW Iberian Peninsula ($0.26\text{-}1.12 \text{ g kg}^{-1}$; $n=4$) [*De la Rosa et al.*, 2011].

The high concentrations of PyC_{PAH} , PyC_{BPCA} and PyC_{GBC} in the Pettaquamscutt River sediments are comparable to other archives that are either down-wind of heavy industry or nearby large industrial areas.

4.3 Decoupled records

PyC_{PAH} concentrations are two orders of magnitude lower than the other methods applied in this study. Nonetheless, PyC_{PAH} have been extensively studied due to the carcinogenic properties of specific PAHs [Mumtaz and George, 1995] and the PyC source information embedded in their distribution [Currie *et al.*, 1997; Rogge *et al.*, 1993]. Thus, although they are trace PyC constituents, it is important to determine temporal variations in their deposition, as well as their origin.

Notably, PyC_{PAH} concentrations remain low (natural background levels) during the pre-industrial era and then reach their maximum during the second half of the 20th Century. The overall trend in PyC_{PAH} concentrations parallels the use of fossil fuels in the northeastern United States. Variations in PyC_{PAH} concentration reflect changes in the type of fuel (coal, oil) and combustion processes (e.g. use of catalytic converters) [Lima *et al.*, 2003] and are superimposed on a constant increase in the consumption of fossil fuels during the industrial era [Lima *et al.*, 2005a; Lima *et al.*, 2003]. In contrast, the results of the “bulk” PyC approaches (PyC_{BPCA} and PyC_{GBC}) vary only slightly in terms of concentration and show broadly parallel down-core variations, implying that they track an important yet distinct component of PyC. Determining the causes of this apparent decoupling of records for different PyC tracers is crucial for a comprehensive quantification of this key carbon pool, as well as for informed interpretation of combustion records.

Within the combustion continuum, PyC can also be comprised of residues produced at low temperature (wood burning) and thus at a lower combustion efficiency. Such residues typically consists of larger PyC particles characterized by a low to intermediate degree of condensation [Masiello, 2004]. The other end of the PyC spectrum derives from condensation reactions in the gas phase during high-temperature combustion of fossil fuels and mainly yields PAH and other fine-carbon particles with a high degree of condensation, including soot.

We examined correlations among the four PyC records in order to further assess their degree of similarity/dissimilarity (Figure 2e). This revealed coherence between PyC_{BPCA} and PyC_{GBC} records, but not in that of PyC_{CTO} , while all three are clearly decoupled from the PyC_{PAH} record. Moreover, different modes PyC productions are evident in the data. The regional-scale history of fossil-fuel combustion since industrialization is echoed in the PyC_{PAH} record. In contrast, the local wood burning practices appear to manifest themselves in PyC_{BPCA} , PyC_{GBC} records, particularly during pre-industrial times when such practices were most pronounced and combustion of fossil fuels had yet to begin.

4.4 Origin and transport pathways

There are several lines of evidence to suggest that the Pettaquamscutt River sediments integrate emissions and deposition on a regional scale as chronicled in the PyC_{PAH} record and in a Pb isotope-based record of combustion sources [Lima *et al.*, 2005a; Lima *et al.*, 2003; Lima *et*

al., 2005b]. Mining and industrial activities in the Upper Mississippi Valley (about 1600 km westward of our study area) and Pennsylvania coal region (about 450 km west-southwest of our study area) [*Census*, 1975] were considered primary sources of the signals observed in the Pettaquamscutt River basin. Wind vector data (Figure 1) lend support for this interpretation of PyC_{PAH} provenance in the northeastern United States [*Golomb et al.*, 2001; *Lima et al.*, 2005a].

Given that the PyC_{PAH} record appears to reflect long-range atmospheric transport, the sharply contrasting PyC_{BPCA} , PyC_{GBC} down-core profiles must trace different PyC source(s) and/or modes of supply to Pettaquamscutt River basin sediments. In particular, the high abundance of PyC_{BPCA} , PyC_{GBC} in sediments deposited prior to the industrial era, as well as their relatively modest variations throughout the record, indicate different source inputs. We infer that the latter derive from inputs stemming from run-off and soil erosion in the Pettaquamscutt watershed, and that the corresponding down-core record therefore reflects the local land-use history. The Pettaquamscutt River catchment was deforested during the late 17th Century and then used as agricultural land until the beginning of 18th Century [*Hubeny et al.*, 2009]. Land under cultivation gradually decreased by 88 % from 1850 to 1969 [*Census*, 1975], eventually accounting for only 4.1 % land usage at the end of the 20th Century [*Widmann*, 2002]. Widespread deforestation has been accompanied by extensive burning, promoting accumulation of charcoal residues in soils [*Santín et al.*, 2015]. Mobilization of burning residues via run-off and bank erosion, potentially also exacerbated by agricultural practices such as ploughing, would eventually result in delivery of particulate PyC to river basin sediments. This process of mobilization and lateral transfer could have continued long after the land use change occurred [*Whitlock and Larsen*, 2001]. Formerly deforested areas underwent regeneration until mid-20th Century. Since then, timber stock has further increased by a factor of three even though the forested area declined by ~5 % due to increasing population density [*Widmann*, 2002]. Local erosion of particulate PyC is probably on going, albeit at a lower level.

In pre-industrial times, PyC must have been supplied predominantly via local surface run-off and erosion processes. With the onset of industrialization, and particular with extensive use of coal for fuel, local PyC inputs have been augmented by atmospheric supply of PyC residues from upwind sources (e.g. Pennsylvania). This additional source of PyC may be responsible for the relatively (compared to changes in PyC_{PAH} concentrations) modest increase in PyC_{BPCA} and PyC_{GBC} concentrations from mid-19th Century to the first half of the 20th Century. During this time period all four PyC records reach their maxima, implying that atmospheric deposition must have influenced overall PyC sedimentary burdens. Subsequent to the 1970s, with the exception of a brief maximum in PyC_{BPCA} in the 1980s that may reflect a surface run-off (storm) event (Table S2, supporting information), all records show a decline in concentration that parallels the implementation of liquid fuels (more efficient conversion to energy and carbon dioxide) and improved controls on particle emissions.

4.5 Implications – Combining complementary analytical tools

The observations resulting from the present study suggest that complementary approaches are required for comprehensive reconstruction past combustion practices and fire history of a watershed from sedimentary records. In particular, it is important to use methods that capture both charcoal and soot PyC , as well as other components of the combustion continuum. The PyC_{PAH} mainly trace fossil-fuel combustion whereas the other tools applied here appear to

primarily reflect biomass-burning residues. Molecular markers can provide qualitative information on PyC source (PyC_{PAH}) and the degree of condensation (PyC_{BPCA}), however quantitative assessments of PyC for the purpose of constraining budgets require use of proxies such as PyC_{BPCA} and PyC_{GBC} that reflect a larger part of the PyC continuum (slightly charred to highly condensed PyC) in combination with further approaches to constrain PyC sources (e.g., radiocarbon analysis).

5 Conclusions

This study sheds new light on the provenance and mode of supply of PyC accumulating in aquatic sediments. Comparison of down-core concentration profiles of four different proxies measured at high-temporal resolution shows that PyC_{PAH} are strongly decoupled from those of the other PyC methods, while PyC_{BPCA} concentration profiles parallel those from the PyC_{GBC} method.

The PyC_{PAH} record mirrors the general increase in production and consumption of fossil fuels brought on by the industrial revolution, with atmospheric transport serving as the primary vector of PyC_{PAH} supply from upwind source regions. In contrast, the PyC_{BPCA} , PyC_{GBC} records likely reflect local land use changes (deforestation, agriculture, urbanization) that resulted in PyC accumulation in, and mobilization from soils within the Pettaquamscutt river watershed. Regional-scale supply and atmospheric deposition of PyC combustion residues appears to be much less important in terms of overall PyC burdens, with only a minor increase in overall PyC quantities accompanying the peak in PyC_{PAH} deposition in the mid-20th Century. All PyC records exhibit a marked decline during the latter half of the 20th Century, concomitant with the use of cleaner fuels and the advent of more stringent emissions regulations.

This study highlights the need for careful consideration of both PyC source characteristics and transport processes in deciphering past records of combustion, and in development of accurate budgets of PyC production and deposition on local to regional scales. Our findings clearly show that different methodological approaches track different portions of the PyC continuum. Further approaches, such as compound-specific radiocarbon analysis, are needed to resolve and quantitatively apportion PyC signatures accumulating in environmental matrices such as sediments.

Acknowledgments

We are grateful to J.W. King for helping us collect such amazing cores; to J. Andrews and P. G Appleby for their help establishing the sediment chronology; to D. Montluçon, S. Sylva, C. Johnson, B. Bergquist, M. Reuer, and A. Dickens for all their help and advice with the lab work; to the late J. Hedges for opening the doors of his home and lab to A.L.L.B.; as well as to M. Hilf for technical support at University of Zurich (UZH). We acknowledge the funding of UZH as well as NSF grants to T.I.E. (OCE-9708478 and CHE-0089172). We also thank two anonymous reviewers for their constructive comments on an earlier version of this manuscript. The data for this study is included in Supporting Information Figure S1-S4 and Table S1-S4 and can be made available upon request from the authors.

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Figure 1: *left:* North-eastern North America with major coal-producing states (light grey) and year (*) when coal production has started (U.S. Geological Survey (<http://pubs.usgs.gov/of/1997/of97-447/text.htm>) and vector wind composite mean from NOAA/ESRL (1871-1998; <http://www.esrl.noaa.gov/psd/cgi-bin/data/composites/plot20thc.v2.pl>); *right:* Pettaquamscutt River catchment (35 km²), RI, U.S. showing boundaries of the drainage basin (dotted line) and the sampling location.

Figure 2: (a-d) Four records on PyC concentrations (closed symbols) and fluxes (open symbols) of the Pettaquamscutt River sediments, RI, U.S. Data are displayed as 20-year moving averages (dark grey) and single data points (light grey). (e) Similarities between the four PyC records, expressed as distances in a two-dimensional coordinate system. Entries in proximity to each other are more similar than distant ones.



