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Significance of Perylene for Source Allocation of Terrigenous **Organic Matter in Aquatic Sediments**

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Supporting Information

ABSTRACT: Perylene is a frequently abundant, and sometimes the only polycyclic aromatic hydrocarbon (PAH) in aquatic sediments, but its origin has been subject of a longstanding debate in geochemical research and pollutant forensics because its historical record differs markedly from typical anthropogenic PAHs. Here we investigate whether pervlene serves as a source-specific molecular marker of fungal activity in forest soils. We use a well-characterized sedimentary record (1735-1999) from the anoxic-bottom waters of the Pettaquamscutt River basin, RI to examine mass accumulation rates and isotope records of perylene, and compare them with total organic carbon and the anthropogenic PAH fluoranthene. We support our arguments with radiocarbon (14C) data of



higher plant leaf-wax n-alkanoic acids. Isotope-mass balance-calculations of perylene and n-alkanoic acids indicate that ~40% of sedimentary organic matter is of terrestrial origin. Further, both terrestrial markers are pre-aged on millennial time-scales prior to burial in sediments and are insensitive to elevated ¹⁴C concentrations following nuclear weapons testing in the mid-20th Century. Instead, changes coincide with enhanced erosional flux during urban sprawl. These findings suggest that perylene is definitely a product of soil-derived fungi, and a powerful chemical tracer to study the spatial and temporal connectivity between terrestrial and aquatic environments.

■ INTRODUCTION

Perylene is found in marine¹⁻⁵ and lacustrine sediments,⁶⁻⁸ soils,^{9,10} and also petroleum^{11,12} and fuel emissions^{13,14} often associated with other distinctive anthropogenic combustionderived polycyclic aromatic hydrocarbons (PAHs). However, most studies report that sediment records of perylene differ greatly from those of anthropogenic PAHs.¹⁵⁻¹⁹ The latter are typically most abundant in sediments post-dating the Industrial Revolution, particularly those deposited during the latter half of the 20th Century, whereas perylene abundances are lowest near the sediment-water interface and tend to increase with depth, particularly in anoxic sedimentary settings.^{18,20} This depthrelated increase in concentration implies abiotic or biologically mediated in situ production from precursor natural prod $uct(s)^{2,21}$ by either a first- or second-order reaction under anaerobic conditions.²⁰

The origin of perylene has remained unclear for decades, during which has previously been argued for terrestrial^{1,2,22,23} and diagenetic, 6,16,24 as well as petrogenic^{12,25} and pyrogenic^{13,14} sources. It was only recently that Itoh and collaborators²³ determined that the fungal species Cenococcum geophillum produces 4,9-dihydroxyperylene-3,10-quinone, confirming a longstanding hypothesis^{1,22} that there is a naturally produced precursor. This ectomycorrhizal fungus appears almost ubiquitous in boreal, temperate, and subtropical regions, and is present in the rhizosphere of woody-plant roots and more generally in forest soils.²⁶ A fungal origin can explain the widespread abundance of perylene in the environment even over geological time-scales,²⁷ since mycorrhizal associations with vascular plants evolved about 400 million years ago.²⁸

Besides C. geophyllum, other mycorrhiza also produce similar perylene precursor compounds^{29,30} as toxins involved in pathogenesis of their host plant.³¹ Prior stable carbon isotopic $(\delta^{13}C)$ measurements of perylene found values similar to those of terrestrial sources (C₃-vegetation: $\delta^{13}C \approx -27 \%$),

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supporting a wood-degrading origin from the rhizosphere.³² This is further supported by dual-isotope analyses of perylene (δ^{13} C and δ D) in sediments showing similar δ D values as methoxy groups in lignin while δ^{13} C values are consistent with the expected fractionation range in saprophytic fungi.²² In addition, natural abundance radiocarbon (¹⁴C) analyses allows a comparison of the ¹⁴C age of perylene with that of other organic matter constituents,²⁵ including total organic carbon (TOC) and specific markers of terrigenous organic matter co-deposited in aquatic sediments. Such analyses shed light on the origin and provide further evidence on the origin of perylene and its use in environmental forensics.

Our motivation is to reconcile existing hypotheses on the source of perylene and assess the potential of this marker compound as a biogeochemical tracer to follow the trajectories of terrigenous organic matter mobilization and transport within watersheds. Specifically, we investigate whether it serves as a molecular marker for rhizosphere carbon from catchment soils and thus facilitates source approximation of terrigenous organic matter in sediments. Moreover, while researchers currently tend to exclude perylene from forensic investigations involving the apportionment of PAHs due to its incongruent behavior, perylene may have mutagenic effects on organisms.^{33,34} Thus, an improved understanding of the provenance and dynamics of perylene may be pertinent to mapping inventories of natural and anthropogenic pollutants, delineating transport pathways, and reconstructing historical land-use.

In this study, we construct historical records of perylene and TOC abundances and isotope compositions from the anoxic sediments of the suburban Pettaquamscutt River basin, RI. We chose these sediments because they provide an exceptionally detailed chronological record and a wealth of background information on the catchment area, including human influence on local to regional scales.^{15,35-38} In addition, this sedimentary record extends over 260 years, from the preindustrial (\sim 1735) until 1999, including an interval characterized by frequent above-ground nuclear-bomb testing (resulting in elevated atmospheric ¹⁴C concentrations peaking in 1963 when the testing ceased), thus offering the opportunity to study the rate at which specific organic carbon species incorporate atmospheric CO₂ and are sequestered in aquatic sediments. These characteristics of the study site, when coupled with down-core δ^{13} C and ¹⁴C records, provide a novel perspective on perylene biogeochemistry, yielding constraints on its source and on the burial efficiency of terrigenous organic matter in aquatic sediments. We compare mass accumulation rates (MAR) of perylene and TOC to the combustion-derived PAH marker fluoranthene. While several nonalkylated PAHs could be used to trace past-combustion practices, we choose fluoranthene because its quantitative down-core profile resembles the temporal evolution of the sum of 15 parent PAHs in this watershed.¹⁵ We further support our interpretation with downcore data of particle size and leaf wax (C_{30-32} ; *n*-alkanoic acids) ¹⁴C variations. We then carefully assess whether the rhizosphere of wooded land serves as the direct source for perylene in the environment and discuss the process of pre-aging of terrestrial organic matter prior to delivery into the aquatic environment in the context of parallel sedimentary records.

EXPERIMENTAL SECTION

A series of seven freeze cores were collected in the depocenter of the lower basin of the Pettaquamscutt River basin, RI (41.503100; -71.450500) in 1999. The catchment area covers 35 km² of forest, wetlands and open water of which today about 30% is residential land.³⁷ Ocean water flooded the basin about 1700 \pm 300 years ago leading to a stratified water column and sustained anoxic conditions in bottom waters and underlying sediments.³⁶ A detailed description on the sediment chronology and sample processing is presented elsewhere.^{15,39} Sediment chronology was obtained from varve counting of X-ray radiographs as well as from ¹³⁷Cs and ²¹⁰Pb profiles using the model of constant rate of supply that yielded a sequence of about 260 years (1735–1999 AD) and an average sedimentation rate of 0.44 \pm 0.10 cm yr^{-1.39}

Sections from the cores were combined after aligning xradiograph images including hurricane layers in 1938 and 1954 to a reference chronology of varve counting, ²¹⁰Pb, and ¹³⁷Cs. This was necessary to obtain enough sediment for trace molecular isotopic-analyses.³⁵ Hence, we pooled the samples in eight horizons for all isotopic analyses: H1 (1999–1982), H2 (1981–1962), H3 (1960–1931), H4 (1929–1898), and H5 (1896–1873), H6 (1871–1842), H7 (1840–1768), and H8 (1764–1735), respectively, except *n*-alkanoic acids that were analyzed in four individual samples: 1885 \pm 5, 1947 \pm 33, 1972 \pm 2, and 1991 \pm 1. Quantitative measurements for the calculation of the mass accumulation rates as well as grain size analyses were performed on individual samples as well.

Grain Size Analyses. One freeze-core slab was subsectioned using a scalpel blade, transferred into 50 mL round-bottom flasks and treated with a 30% hydrogen peroxide solution at a ratio of 40 mL per 1 g sediment, and heated in a water bath to 70 °C as a means to support mineralization of the organic matter. Subsequently, samples were freeze-dried, and a subsample of about 11 mg was suspended in water and measured on a Beckmann Coulter LS13 320 laser diffraction particle analyzer (Indianapolis, IN) in triplicate.

Total Organic Carbon and Nitrogen. A Fisons 1108 elemental analyzer was used to measure the TOC content of the samples. To remove the inorganic carbon fraction, about 2 mg of dry sample was weighed into a silver capsule and acidified with 20 μ L of 2 N HCl. TOC content was calculated in relation to the whole sediment dry weight while organic carbon/organic nitrogen TOC/TN ratios were calculated on an atomic basis. Samples were run in triplicate and all reported weight percentages represent the mean \pm one standard deviation with an instrumental blank of 0.004 mg for C and smaller than 0.005 mg for N.

Isotope Ratio Monitoring Mass Spectrometry. The stable carbon isotopic composition of bulk sample TOC was determined in triplicate by automated online solid combustion interfaced to a Finnigan Delta Plus isotope ratio mass spectrometer. Isotope ratios were calculated relative to CO_2 reference gas pulses, with standard deviations for replicate measurements were always better than 0.6% and usually within 0.3%.

Extraction, Purification, and Isotope Analyses of Perylene, Fluoranthene, and *n*-Alkanoic Acids. Dry sediment samples (0.5–1.5 g) were extracted by pressurized fluid extraction (Dionex ASE 200) using a mixture of acetone and *n*-hexane (1:1) at 1000 psi at 100 °C. Molecular identification and quantification was achieved using an Agilent 6890 Plus GC System interfaced to a mass selective detector operating at 70 eV in SIM mode using a DB-XLB capillary column (60 m × 0.25 mm × 0.25 μ m).¹⁵



Figure 1. Mass accumulation rates for total organic carbon (TOC), perylene, and fluoranthene as well as carbon to nitrogen (TOC/TN) ratios and down-core shifts in coarse particle abundance (D_{90}) with shaded area A representing the maximum PAH flux in this area, B the period of amplified erosion due to infrastructure development, and C the preindustrial era prior to 1840s.

Compound-specific radiocarbon analyses (CSRA) of perylene and fluoranthene were performed on eight pooled horizons by first using high-pressure liquid-chromatography (HPLC) to separate pure pervlene (98% purity or greater) from the sample extracts. The HPLC procedure isolated PAHs into 2ring and the combined 3 + 4-ring and 5 + 6-ring PAHs. The resulting 16 HPLC fractions (8 horizons \times 2 ring classes) were subjected to two-dimensional preparative capillary gas chromatography for isolation and purification of individual PAHs via HP 7683 autoinjector and a multicolumn switching system (Gerstel MCS 2) connected to a HP 6890 series gas chromatograph, and Gerstel preparative fraction collector (PFC; further details in the Supporting Information (SI)). Purified samples were transferred to precombusted quartz tubes $(7 \text{ mm i.d.} \times 20 \text{ cm})$, dried under nitrogen before adding copper oxide (50 mg). Each tube was then evacuated on a vacuum line while samples were kept at -90 °C to prevent sublimation, sealed, and combusted at 850 °C for 5 h. About 95% of the purified carbon dioxide was reduced to graphite, pressed and analyzed for ¹⁴C at NOSAMS, Woods Hole, MA, and the remaining 5% was used for δ^{13} C measurements.

The δ^{13} C values of perylene were also determined in triplicate on a Finnigan Delta Plus isotope ratio mass spectrometer with attached Finnigan GC combustion III interface and Hewlett-Packard 6890 GC (irm-GC/MS). Compounds were separated on a CP-Sil 5CB capillary column (50 m × 0.25 mm × 0.25 μ m) and isotope ratios for PAH peaks were calculated relative to CO₂ reference gas pulses. The standard deviation for replicate measurements of perylene was better than 0.6 ‰ and mostly around 0.2 ‰.

The isotopic composition of leaf wax *n*-alkanoic acids (C_{30-32}) were determined for four individual sediment samples deposited in 1885 ± 5, 1947 ± 33, 1972 ± 2, and 1991 ± 1 following the analytical procedure described elsewhere.⁴⁰ In brief, *n*-alkanoic acids were extracted, isolated, and purified using a preparative Hewlett-Packard 5890 series II capillary gas

chromatograph.⁴¹ Following the chromatographic purification of individual compounds, samples were graphitized and measured at NOSAMS, Woods Hole, MA.

Evaluation of Isotope Data. Isotope mass-balance calculations provide quantitative estimates for source apportionment of organic matter. To constrain the sources of TOC and perylene in Pettaquamscutt River sediments, we calculated the relative contribution of possible organic carbon sources in the sediments using the measured ¹⁴C and δ^{13} C values for TOC (Figure 2). We report the ¹⁴C data as F¹⁴C which is the fraction modern independent from the year of measurement.⁴² We assumed a simple mixing model to retrieve an average value for the time lag of perylene and leaf wax *n*-alkanoic acids; the details of which are published elsewhere.³⁵ In brief, these source-specific molecular markers facilitate approximations of the contributions of terrigenous organic matter in aquatic sediments (further details on the concept are included in the SI).

To account for the potential variability under increasing human pressure on the coastal environment, we determined average values of our terrestrial end-members for two time periods: the preindustrial (1735-1840) and the post-"bomb" (1960–1999) period (see SI for more detail). Regarding ¹⁴C data evaluation, the post-1960s period also is a benchmark to test for the resilience and short-term dynamics of catchment processes. In contrast, the aquatic end-member is only affected by the marine reservoir effect, which is about 400 ± 40^{14} C years lower than the atmospheric ¹⁴C concentration at the time of burial.⁴³ Furthermore, we neglect a potential impact of petrogenic carbon⁴⁴ in the Pettaquamscutt River because of the absence of postglacial erosion and due to the absence of carbonaceous metasedimentary rocks in the basin.⁴⁵ We constrain our model by $\delta^{13}C = -21.0\%$ for aquatic production⁴⁶ and use local records of perylene, pyrogenic carbon³⁵ and *n*-alkanoic acids (C_{30-32}) to define terrestrial endmembers. We further obtained the contemporary atmospheric



Figure 2. High-resolution δ^{13} C record for TOC and pooled sample results for perylene (left); molecular ¹⁴C data for the combustion-derived fluoranthene (middle) and results on perylene, leaf wax *n*-alkanoic acids as well as TOC (right).

¹⁴C concentrations from the extended Intcal13 reference chronology.^{43,47}

RESULTS AND DISCUSSION

The exceptionally well-constrained chronology of the Pettaquamscutt River sediments along with detailed reconstructions of the provenance of combustion-derived PAHs^{15,35} and Hg³⁸ provide key constraints that allow us to determine whether perylene signatures are consistent with its production in the rhizosphere of catchment soils.^{23,26} To characterize the export of terrigenous organic matter, including the putative precursors of perylene, and its subsequent burial in aquatic sediments, we report MAR for TOC, perylene, and fluoranthene. Isotopic mass balance calculations for δ^{13} C and ¹⁴C records are employed to further constrain the main source of perylene precursors. With reference to recent research and our results, we then provide a synthesis aiming to merge the existing concepts, reconciling the origin of perylene (i.e., terrestrial, diagenetic or petro-/ pyrogenic).

Down-Core Mass Accumulation Rates of TOC and Perylene. The burial rate of TOC in the sediments of the Pettaquamscutt River ranges from 38 to 61 g m⁻² yr⁻¹ (average = 49 ± 6 g m⁻² yr⁻¹, n = 71). The lowest TOC burial flux is observed in ca. 1938 (Figure 1) and is related to a 16.7% shift in burial flux from 53.0 ± 43 g m⁻² yr⁻¹ prior to ca. 1927 to $44.0 \pm$ 3.7 g m⁻² yr⁻¹ for sediments deposited thereafter. The timing of this change coincides with the construction of the Lacey Bridge on the ocean side of the lower basin in 1934⁴⁵ (Figure 1B) that likely caused a restriction in seawater intrusion, affecting the influx of both marine as well as the terrestrial matter. We argue that this exemplifies the accelerated development of the catchment area beginning early in the last century and the related infrastructure projects that changed the natural flow paths of materials within the watershed.

The N_{org} down-core record reveals only small variations, here illustrated as C_{org}/N_{org} profiles (n = 71; Figure 1) where three trends are delineated: decreasing C/N from 1730s (12.6) to 1840s (9.4), a ~50-year reversal toward higher C/N ratios (13.2 in 1900), and a century-long C/N secular decrease in the upper core (10.0 in 1999). This rather narrow range in C/N ratios suggests that organic matter is predominantly yet not exclusively

composed of labile material with C/N values similar to aquatic organic matter (C/N = 4 to 10).⁴⁸ However, we refrain from further speculations on this bulk-level information due to the absence of a robust reference for C/N values of terrigenous organic matter from this area.

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MARs for perylene (and proto-perylene), which corresponds to the reduced perylenequinone, vary from 29 to 3800 $\mu g m^{-2}$ yr⁻¹, while its down-core profile differs markedly from the combustion-marker fluoranthene (Figure 1). The values were low before 1938, reached a maximum between ca. 1938 and 1973 (1580 $\mu g m^{-2} yr^{-1}$), before diminishing again in most recent decades (~ 100 μ g m⁻² yr⁻¹) (Figure 1). This contrasting accumulation behavior implies a different origin for perylene relative to combustion-derived PAHs, here illustrated by fluoranthene (Figure 1). This nonalkylated PAH appears lower in abundance varying from 1 to 1280 μ g m⁻² yr⁻¹ and also exhibits significant variability,³⁵ however the temporal evolution of MARs of this pyrogenic PAH differs sharply from that of perylene. Specifically, fluoranthene's MAR was low until the mid-19th century (Figure 1C), increased steadily to a maximum in the late-1950s, then declined in the post-1970s due to cleaner burning fuels and as consequence of environmental regulations.^{15,37} Such down-core profiles of combustion-derived PAHs are also observed in other sedimentary records, including the upper basin of the Pettaquamscutt River. 16,19,49

The observed decline in perylene (and proto-perylene) MAR between 1927 and 1938 coincides with the abrupt decrease in TOC accumulation rate (Figure 1B). We attribute the latter to a dilution in organic carbon content resulting from a higher proportion of clastic material due to erosion of soil mineral horizons (Figure 1). Perylene is a trace constituent of TOC, never accounting for more than 0.1% of TOC, and consequently perylene/TOC ratios shift by 500% relative to before or after this time interval. These observations thus support the link between a decrease in perylene's MAR resulting from an increase in erosional flux (Figure 1B).

A striking feature of the perylene profile is that fluxes remain nearly constant ($665 \pm 180 \ \mu g \ m^{-2} \ yr^{-1}$; n = 26) prior to 1865, indicating either steady-state erosional conditions or quantitative conversion of precursor material (Figure 1).²³ Due to the large variations in perylene burial rates in our sedimentary record we refrain from attempting to derive kinetic parameters for conversion of perylenequinone (proto-perylene) precursor to perylene. However, Slater and collaborators⁵⁰ compared two perylene profiles from Lake Siskiwit collected in 1983 and 2005. They found a reaction rate constant (0.048 yr⁻¹), ca. 75% higher than previously reported,²⁰ whereas another study reported much lower rates.¹⁸ All of the above studies suggest that the conversion of perylenequinone to perylene follows first-order kinetics. Nevertheless, the strong (~ 97%) decrease in perylene abundance in Pettaquamscutt River sediments from 1973 to 1999 likely reflects incomplete conversion of precursor compounds (Figure 1).

Isotope Mass Balance Calculations. The δ^{13} C TOC profile (Figure 2) appears relatively invariant throughout the core (-25.4 to -23.0%, average -24.1 ± 0.5%, n = 36), indicating the absence of large variations in organic matter composition. Although the TOC MAR decreased in the 1930s, δ^{13} C values do not suggest this was accompanied by a shift in the nature of organic matter supplied to Pettaquamscutt River basin. Instead, the observed changes in burial flux in this catchment are attributed to a shift in land use stemming from an increase in the local human population and the accompanied development of infrastructure. The δ^{13} C values of perylene (-28.9 to -28.3%); average -28.6 ± 0.6%; Figure 2) were similarly invariant, while significantly lower than TOC and similar to that reported for terrestrial C₃ plants (-29 to -25, average -28%).⁵¹

In contrast, the ¹⁴C record of TOC for pooled samples (n = 8; Figure 2) reveals only small variations in preindustrial time (1735–1840) but then tends to shift toward more ¹⁴C depleted values between 1839 and 1958. This change is thought to integrate the increasing use of fossil fuels slightly diluting the natural atmospheric ¹⁴C concentration (the so-called "Suess effect") as well as ongoing infrastructural development and constructions within the catchment resulting in the mobilization of deeper mineral soils. Thereafter, we observed a drastic (~15%) increase in F¹⁴C TOC values reflecting the shift in atmospheric ¹⁴C derived from thermonuclear weapons testing in the post-1960s. The uppermost sediment layer signals declining ¹⁴C concentrations similar to the atmosphere.⁴⁷

Compound-specific radiocarbon analysis reveals that the ¹⁴C profile of perylene follows that of TOC from the preindustrial period until ~1950. Thereafter, ¹⁴C of TOC increases due to the incorporation of atmospheric bomb ¹⁴C, while perylene tends to more ¹⁴C-depleted values. This could reflect a pyrogenic (fossil fuel) contribution to perylene, mitigating the impact of "bomb" carbon in the Pettaquamscutt River sediments. However, CSRA of leaf wax *n*-alkanoic acids (C_{30-32}) from the same sediments reveals a concomitant ¹⁴C trend with perylene (Figure 2). We infer, therefore, that leaf waxes and perylene share a common source and mode of export, implying that local erosional fluxes and mobilization of soil organic matter are responsible for the decoupled TOC and perylene post-1950s signals rather than global-scale perturbations such as "Suess effect" or above-ground nuclear weapons testing.

In the mainly wooded catchment area of Pettaquamscutt River, organic molecules are produced during photosynthetic activity, translocated and metabolized in plants, as well as released into the rhizosphere. In soil, plants and animals, residues can be assimilated, stabilized and metabolized until they eventually become respired to carbon dioxide,⁵² or they can be eroded and redistributed within the watershed.⁵³ Source-specific molecular markers allow delineation of specific processes associated with organic matter cycling, while molecular ¹⁴C data provide additional information on the time scales of organic matter storage and transport from its source to the "ultimate" site of burial. This "residence time" has been determined for several markers of terrestrial primary productivity, ^{54–56} as well as charred plant biomass, ^{35,57} and range from centuries to several millennia. The underlying processes resulting in organic matter "pre-aging" prior to burial in aquatic depocenters can complicate source apportionment because the ¹⁴C concentration of various organic matter components differs from that of the corresponding atmospheric ¹⁴C reference value.

We determined an average ¹⁴C residence time for perylene of 1300 ± 300 ¹⁴C years for samples pre-dating the industrialization (1735-1840, n = 3) and 2000 ± 500^{14} C years for the post-"bomb" era (1960–1999; n = 3). These values agree well with leaf wax *n*-alkanoic acids (C_{30-32}) that showed a 2700 ± 700 ¹⁴C year residence time for the post-"bomb" era (Figure 2), and a record of biomass-derived pyrogenic carbon from the same core revealed an average age of 1460 \pm 490 14 C years for the preindustrial era.³⁵ The apparent average age or residence time over the preindustrial and the post-"bomb" era for perylene and higher plant n-alkanoic acids of ca. 2000 ¹⁴C years is slightly higher than it has been reported for (micro) charcoal in the same catchment³⁵ as well as for leaf wax *n*-alkanoic acids in the Bengal fan⁵⁵ yet similar to that of Cariaco basin.⁵⁶ However, in the post"bomb" era we observed a decoupling of ¹⁴C concentrations among molecular markers and TOC. Specifically, the increase in atmospheric ¹⁴C concentrations post-1960s is not apparent in perylene or *n*-alkanoic acids, suggesting that soil carbon storage on millennial time scales may conceal or delay the legacy.

Isotope mass balance calculations based on TOC ¹³C and ¹⁴C data yield similar values for the proportion of organic matter of terrestrial origin (δ^{13} C: -43 ± 10% and ¹⁴C: -37 ± 10%), with the remaining carbon derived from aquatic productivity. These results for the Pettaquamscutt River basin are slightly higher than an estimated global average of one-third of sedimentary organic is of terrestrial origin.⁵⁸ The similar estimates obtained from dual-carbon isotope evaluations based on perylene reinforce its value in constraining the terrestrial end-member in source apportionment.

New Constraints on the Source of Perylene. The origin and widespread distribution of perylene in the environment has been a subject of scientific debate for decades, although its conversion from the precursor molecule, 4,9-dihydroxyperylene-3,10-quinone, has long been suspected.^{1,2,23,29⁴} In particular, it has remained unclear whether perylene is formed during in situ diagenesis^{6,16} or is of petro-/pyrogenic origin.^{11,14,25,59} The former hypothesis was stimulated by the presence of perylene in Antarctic marine sediments⁶⁰ and by large similarities between perylene and total organic matter down-core profiles, fueling the notion of in situ synthesis from TOC.¹⁶ Along these lines, Gschwend and collaborators²⁰ calculated kinetic parameters necessary to yield perylene from biogenic precursor concentrations that are consistent with results from a recent study comparing down-core profiles 20 years apart in Lake Siskiwit,⁵⁰ and both are in line with the observed reduction of perylenequinones.²³ The second alternative hypothesis is based on the abundance of perylene in ancient sediments,^{22,61} fossil fuels^{11,12} or combustion emissions.^{13,14,59} Regarding the latter sources, perylene concentrations remain below 1.4% of the total PAH and thus are negligible for the overall budget. Its presence in these matrices, however, raises the question of whether perylene is exclusively of fungal origin. A possible explanation for the

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occurrence of perylene in fossil fuels lies in the evolutionary development of fungi. The symbiotic interaction between fungi and vascular plants evolved about 400 Ma ago, and ectomycorrhiza have existed for at least 56 Ma,⁶² while the divergence and evolution of the mycorrhiza gene pool remains largely unexplored.²⁸ However, Blumer proposed the conversion of pigments to hydrocarbons through a geochemically irreversible deoxygenation of functional groups and hydrogenation under sustained reducing conditions.⁶³ Even though this was exemplified for fossil crinoids in a pigment-rich Triassic oil shale, these reactions likely also apply to the precursors of biogenic PAH. We deduce, therefore, that fungal-derived perylenequinone is deoxygenated during sediment diagenesis and, at least partly, survives coalification. This could explain the occurrence of perylene in fossil deposits³² and could explain its abundance in combustion residues. One evolutionary rationale for the occurrence of perylene in scletoria, the resting structures of C. geophyllum,⁶⁴ is the extreme mutagenic activity of the unsaturated nucleus of perylenequinones on Gram-negative bacteria,³³ protecting the reproductive function of fungal spores. Overall, the preponderance of evidence,^{17,22,32} reinforced by our new data, indicates that perylene is a remnant of wood degradation or fungal activity in the rhizosphere, a hotspot of biological activity on land, and source of organic matter to aquatic systems.

Parallels between Perylene and TOC. Sedimentary organic matter is a highly complex mixture composed of aquatic and terrigenous organic carbon, and this complexity complicates source apportionment calculations, as well as determination of carbon fluxes and thus the carbon burial efficiencies.^{65,66} Measurement of the abundance and isotopic composition of source-specific molecular markers can provide constraints on terrigenous organic matter in sediments.⁶⁵ In addition to source constraints from δ^{13} C and 14 C signatures, the latter also yields information on whether organic matter is of modern or fossil origin,⁴⁴ as well as on time scales of terrigenous organic matter between formation on land and burial in aquatic depocenters (i.e., "average residence times").³⁵ The process of "pre-aging" implies a time lag between biosynthesis and export of organic matter from land to depocenter, with natural as well as anthropogenic molecules being retained in catchment soils from annual to millennial time-scales.35,55,56 Here, we find that this pre-aging process occurs on similar time-scales for leaf wax fatty acids (*n*-alkanoic acids), biomass-derived pyrogenic carbon and the rhizosphere tracer perylene-all together terrestrial markers with different modes of formation, chemical structures and functionalities. This agreement between these different source-specific markers suggests that retention within soils regulates the export of most terrigenous organic matter and associated carbon-based pollutants that enter the soil column.

TOC and perylene MARs covary in the preindustrial era until about 1850 despite some variability in the TOC flux (Figure 1). We attribute the latter to the increasing pressure by extensive farming, as well as potential changes in aquatic productivity. The dual carbon isotope records for TOC and perylene (Figure 2) support the absence of drastic compositional shifts throughout the record, despite the influence of the "Suess effect" and nuclear weapons testing on the latter part of the TOC ¹⁴C record. Notably, perylene shows large variation in MAR after 1850, which we suspect is due to the conversion of the catchment from a rural into a suburban landscape that was accompanied by an enhanced erosional flux of carbon from bare soils. Similarities between TOC and perylene records have been observed previously,^{6,16,22,61} yet in these cases it was suspected to reflect direct in situ microbial production of perylene¹⁶ rather than the reduction and conversion of a precursor into perylene. This depletion of the precursor pool may continue over geological time scales,⁶³ thus limiting comprehensive quantitative assessment of perylene in recent sediments.⁵⁰ In contrast, the isotopic composition of perylene is expected to be insensitive to conversion efficiency, and thus can serve as a robust tracer of soil-derived terrestrial organic matter.

ASSOCIATED CONTENT

S Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.est.9b02344.

Details on the preparative chromatographic isolation of PAHs for compound-specific isotope analysis, the concept of two modes of terrigenous organic matter export from land to oceans, the impact of the nuclear weapon testing on the atmospheric and marine ¹⁴C partitioning over time and results for grain size analyses, respectively (PDF)

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Notes

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