Anthropogenic Inputs of Terrestrial Organic Matter Influence Carbon Loading and Methanogenesis in Coastal Baltic Sea Sediments

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Coastal regions globally have experienced widespread anthropogenic eutrophication in recent decades. Loading of autochthonous carbon to coastal sediments enhances the demand for electron acceptors for microbial remineralization, often leading to rearrangement of the sediment diagenetic zonation and potentially enhancing fluxes of methane and hydrogen sulfide from the seafloor. However, the role of anthropogenic inputs of terrestrial organic matter (OMterr.) in modulating diagenesis in coastal sediments is often overlooked, despite being of potential importance in regions of land-use and industrial change. Here we present a dated 4-m sediment and porewater geochemistry record from a eutrophic coastal location in the northern Baltic Sea, to investigate sources of recent carbon loading and their impact on modern diagenetic processes. Based on an end-member mixing model of sediment N/C ratios, we observe that a significant fraction of the late-20th century carbon loading at this location was contributed by OMterr.. Furthermore, analysis of lignin in this material shows depleted ratios of syringyl/vanillyl (S/V) and cinnamyl/vanillyl (C/V) phenols, indicative of enhanced inputs of woody gymnosperm tissue likely from forest industries. The rapid loading of organic matter from combined terrestrial and autochthonous sources during the late 20th century has stimulated methanogenesis in the sediment column, and shoaled the sulfate-methane transition zone (SMTZ) to a depth of 5–20 cm. Optical parameters of colored dissolved organic matter (CDOM) confirm that OMterr. is actively degrading in the methanogenic layer, implying a role for this material in diagenetic processes. Porewater CH4, SO42−, δ13C-DIC, and ∑S2− data suggest that the modern SMTZ is a broad zone in which organoclastic sulfate reduction, methanogenesis and anaerobic oxidation of methane (AOM) co-occur. However, fluxes of CH4 and SO42− show that rates of these processes are similar to other marine locations with a comparably shallow SMTZ. We suggest that the shallow depth of the modern SMTZ is the principal reason for high observed diffusive and ebullitive methane fluxes from sediments in this area. Our results highlight that anthropogenic activities lead to multiple pathways of carbon loading to coastal
INTRODUCTION

Anthropogenic impacts on coastal environments have accelerated in parallel with industrialization during the late 20th century (Diaz and Rosenberg, 2008). Among the most widespread and important impacts is eutrophication, defined as an increased supply of organic matter to a coastal ecosystem (Nixon, 1995), and often primarily driven by enhancement of autochthonous productivity due to loading of nutrients such as nitrogen and phosphorus (Bonsdorff et al., 1997; Anderson et al., 2002). Eutrophication increases the amount of fixed organic carbon in coastal ecosystems and alters the primary producer community structure (Heisler et al., 2008), consequently raising the oxygen demand of microbial remineralization of organic matter (OM) (Middelburg and Levin, 2009; Breitburg et al., 2018). In turn, consumption of oxygen leads to further ecosystem impacts associated with benthic hypoxia (Carstensen et al., 2014; Rabalais et al., 2014). Because a significant fraction of OM remineralization in shallow coastal systems takes place in the underlying sediments, eutrophication can severely impact rates of microbial processes in the sediment column. Ultimately, these processes dictate the rate of organic carbon turnover and burial (Arndt et al., 2013) and hence play a critical role in the global carbon cycle.

One of the key consequences of eutrophication for coastal sediment microbial processes is to enhance rates of anaerobic remineralization after the exhaustion of oxygen (Middelburg and Levin, 2009). Aerobic sulfate reduction is the dominant pathway of anaerobic remineralization in marine sediments (Jorgensen, 1982; Canfield, 1991; Bowles et al., 2014), but in settings with low salinity and high flux of sedimenting OM, sulfate may become completely exhausted within the upper sediment column (e.g., Slomp et al., 2013; Thang et al., 2013). In such cases, a significant fraction of OM remineralization occurs through methanogenesis (Rooze et al., 2016), creating the potential for emissions of this potent greenhouse gas to the water column and ultimately the atmosphere (Bange et al., 1994; Geleš et al., 2016; Humbold et al., 2019; Myllykangas et al., 2020a). Many studies have demonstrated the presence of methane within the uppermost meter of the sediment column in low-salinity coastal settings (e.g., Albert et al., 1998; Martens et al., 1998; Mogollon et al., 2011; Egger et al., 2015), and human-impacted estuaries are considered hotspots for methane emissions (Borges and Abril, 2011). Because such emissions may offset the climate benefits of carbon burial (Dean et al., 2018), it is essential to understand the controls on methanogenesis in human-impacted coastal sediments.

Associated to changes in vertical zonation of primary anaerobic remineralization pathways, eutrophication can also strongly modify the network of secondary microbial processes in the sediment column. For example, a key consequence of enhanced methanogenesis in human-impacted systems is enhanced rates of anaerobic oxidation of methane (AOM). This process proceeds mainly through coupling to sulfate reduction at the so-called sulfate methane transition zone (SMTZ) (Knittel and Boetius, 2009), but can also be coupled to reduction of nitrate (Ettinger et al., 2010) or metal oxides (Beal et al., 2009; Sivan et al., 2011; Egger et al., 2015). Crucially, AOM strongly reduces the fluxes of methane from the sediments to the water column relative to the amount of methane produced during sedimentary methanogenesis, thus acting as a filter for methane emissions (Dean et al., 2018).

In addition to inputs of autochthonous OM, coastal sediments also receive carbon loading from terrestrial organic matter (OMterr.) (Blair and Aller, 2012). Up to 200 Tg particulate organic carbon (POC) are delivered by rivers to coastal environments annually (Schlunz and Schneider, 2000), while a similar flux has been calculated for dissolved organic carbon (DOC) (Dai et al., 2012). The DOC contribution typically dominates OMterr. inputs in boreal systems (e.g., Mattsson et al., 2005). The importance of DOC inputs for coastal sediment carbon loading in the Baltic Sea is not well constrained, but several studies have suggested that salt-mediated flocculation may transfer carbon from the dissolved to particulate phase at the land-sea transition (Sholkovitz et al., 1978; Astma et al., 2014; Jilbert et al., 2018). Typically, the reactivity of OMterr. in microbial remineralization is considered lower than that of autochthonous material (Hedges et al., 2000). This is due to a combination of its refractory molecular composition, being dominated by lignin, cellulose and cutin (de Leeuw and Largeau, 1993) and physical association to protective minerogenic material prior to deposition (Mayer, 1994; Hedges and Keil, 1995). However, riverine fluxes, molecular composition and biological reactivity of OMterr. have been shown to be sensitive to anthropogenic activities such as deforestation, ditching, agriculture and industry (e.g., Lambert et al., 2017; Astma et al., 2019; Deininger and Frigstad, 2019; Landsman-Gerjoi et al., 2020). Rapid inputs of such “anthropogenic” OMterr. to coastal sediments may be expected to confer higher reactivity to OMterr. in the sediment column. Yet, few studies have explicitly investigated the impacts of anthropogenic OMterr. inputs on coastal sediment biogeochemistry.

In boreal regions, a key driver of anthropogenic OMterr. inputs to the marine environment over the last century is forestry and the associated paper and pulp industry. Forest industry processes, including debarking, pulping, bleaching and washing (Ali and Sreekrishnan, 2001), have introduced large point-source emissions of raw and processed OMterr. to aquatic systems, in many cases proximal to the coastal environment (Pocklington and McGregor, 1973; Louchouarn et al., 1999; Brandenberger et al., 2011). The molecular composition of solid forest industry sediments, and that forest industry impacts on sedimentation in the northern Baltic Sea may be more widespread than previously acknowledged.

Keywords: terrestrial organic matter, forestry, lignin, diagenesis, methane, blue carbon, baltic sea
wastes in sediments includes the major wood polymers cellulose and lignin (Louchouarn et al., 1997; Dahlberg et al., 2020), together with a wide array of additional compounds from the raw material and its treatment processes, including tannins, resin acids, and polychlorinated biphenyls (PCBs) (Lacorte et al., 2003). Sedimentary contents of such materials have been shown to vary with distance from industrial sources, with the most concentrated deposits (e.g., cellulose-rich fiberbanks) consisting of nearly pure organic material (Dahlberg et al., 2020; Dahlberg et al., 2021).

To date, several studies have used organic biomarkers such as lignin phenols and persistent organic pollutants (POPs) to identify the presence of forest industry-derived anthropogenic OMter. in boreal coastal sediments, and to quantify its local contribution to total OMter. or total sedimentary organic matter (e.g., Louchouaran et al., 1997; 1999; Dahlberg et al., 2020). However, few studies have combined these analyses with a porewater biogeochemistry approach to determine the potential role of such inputs on sediment diagenetic processes, including methanogenesis. This is an important gap in knowledge, since OMter. has been shown to promote methanogenesis in lakes (Tittel et al., 2019) and fiberbank material in particular shows strong methanogenic potential (Kokko et al., 2018). Here, we employ a comprehensive analysis of sediment bulk inorganic and organic chemical parameters, including lignin phenol analysis, coupled to detailed porewater chemical profiling, to determine the impacts of recent anthropogenic OMter. inputs on diagenesis at a site in the northern Baltic Sea. The site is located tens of kilometers from the nearest forest industry point source, allowing investigation of larger spatial-scale impacts of anthropogenic OMter. than previous studies focused on fiberbanks close to industrial point sources.

**SITE DESCRIPTION**

The Baltic Sea is a landlocked brackish marine system in northern Europe with a N-S salinity gradient of 3–15 (Leppäranta and Myrberg, 2009). The whole Baltic Sea was strongly eutrophied by nutrient loading during the 20th century (Gustafsson et al., 2012), leading to the development of widespread deep water hypoxia and anoxia in the central basins (Conley et al., 2009). Many coastal areas have shown localized trends towards oxygen depletion during recent decades, as a consequence of direct nutrient inputs from adjacent land areas leading to coastal eutrophication (Conley et al., 2011). The contribution of OMter. to Baltic Sea sediments as a whole is estimated as ≤30% of total OM, with a N-S compositional gradient from gymnosperm-to angiosperm-dominated material (Miltner and Emeis, 2001). A recent study demonstrated clearly higher OMter. contributions to sedimentary carbon in northern areas such as the Gulf of Bothnia and Gulf of Finland (Nilsson et al., 2021).

This study focuses on the Storfjärden site on the Finnish coast of the Gulf of Finland in the northern Baltic Sea (Figure 1). Storfjärden is a shallow coastal basin forming part of the Tvärminne archipelago system at the mouth of the Pojo Bay estuary, draining the Mustinjoki river catchment in southern Finland. Since the most recent deglaciation at 12250 cal year BP, Storfjärden has been successively covered by late- and post-glacial lacustrine clays and most recently by brackish-water muds (Virtasalo et al., 2014; Virtasalo., 2019). The Gulf of Finland
coastal areas were affected by eutrophication during the 20th century (Weckström, 2006), and the majority of the OM in the modern sediments is derived from autochthonous production (Jilbert et al., 2018). However, Jokinen et al. (2020) measured elevated contents of OM_{terr.} in nearby sediments during the depositional interval corresponding to the late 20th century, suggesting an anthropogenic influence on OM_{terr.} fluxes from the Mustionjoki catchment during this period. The city of Lohja in the southern part of the catchment has been a major center of the Finnish pulp and paper industry since the early 20th century (Katko et al., 2005).

### MATERIALS AND METHODS

#### Sediment Coring

Sediments were collected from the Storfjärden site (33 m water depth, Figure 1) during various sampling campaigns from 2015 to 2020, onboard the vessels R/V Saduria, R/V Electra and R/V Augusta (Table 1). A twin-barrel GEMAX coring device was used to recover the intact sediment surface and approximately the uppermost half meter of the sediment column. A piston corer onboard R/V Electra was used to recover longer sediment cores (max. length 4.5 m in September 2017; 5.5 m in September 2018). Seasonal changes in sediment and porewater chemistry on the decimeter scale are known to be minimal at this site (Myllykangas et al., 2020a), hence the figures in this paper present combined data from multiple sampling campaigns.

#### Porewater Sampling and Treatment

Porewater samples for most parameters were collected with Rhizons\textsuperscript{TM} (pore size 0.15 μm). For each core, a vertical series of holes (diameter 4 mm) was pre-drilled into the liner and taped prior to sampling (resolution 2 cm for GEMAX cores, 10 cm for piston cores). After core recovery, the tape was penetrated and Rhizons\textsuperscript{TM} inserted. Porewater was collected under vacuum in an attached plastic syringe and transferred directly into a 65-ml glass bottle containing supersaturated NaCl solution. The bottles were closed with a butyl rubber stopper and screw cap. A headspace of 10 ml N\textsubscript{2} gas (purity 5.0) was injected through the stopper within 24 h of sampling and bottles were stored inverted until analysis.

#### Porewater Analyses

Dissolved S was determined by ICP-OES after dilution (Thermo iCAP 6000 at University of Helsinki), and is assumed to predominantly represent SO\textsubscript{4}^{2-}. Hydrogen sulfide (H\textsubscript{2}S) is lost during acidification hence does not contribute to dissolved S in the ICP-OES subsample (Jilbert and Slomp, 2013). We acknowledge that at low SO\textsubscript{4}^{2-} concentrations, such as in deeper layers of the sediments, non-zero dissolved S values may indicate the presence of dissolved organic sulfur (Jilbert et al., 2020). However, the present study focuses on S gradients in the near-surface sediments, where this fraction is considered negligible, hence we assume dissolved S \(\approx\) SO\textsubscript{4}^{2-}. Reproducibility of the ICP-OES analyses is <5% RSD. Dissolved ammonium (NH\textsubscript{4}+) was determined by the indophenol method (Koroleff, 1976) using an autoanalyzer (Lachat QuikChem 8000, reproducibility <5% RSD). Analysis of δ\textsuperscript{13}C-DIC in headspace gas from the H\textsubscript{3}PO\textsubscript{4}-acidified vials was performed on a Thermo Gasbench II coupled to a MAT 253 mass spectrometer at Stockholm University and is reported in conventional delta notation relative to Vienna PeeDee Belemnite (VPDB). Standard deviation was less than 0.1‰.

| Porewater parameters | Sediment parameters |
|----------------------|---------------------|
| NH\textsubscript{4}\textsuperscript{+} | Alk | δ\textsuperscript{13}C-DIC | S (ICP-OES) | a | CH\textsubscript{4} | DOM (peak) | Pb\textsubscript{206/207Pb} | C, N | Lignin | phenols | Fe, Mn | S (ICP-OES) |
| GEMAX April/June 2015 0–40 cm | | | | | | | | | | | | |
| GEMAX Sept. 2018 0–60 cm | | | | | | | | | | | | |
| GEMAX June 2020 0–60 cm | | | | | | | | | | | | |
| Piston Sept. 2017 40–400 cm | | | | | | | | | | | | |
| Piston Sept. 2018 60–650 cm | | | | | | | | | | | | |

*Data reproduced from Jilbert et al. (2018).
Total alkalinity (AlkT) was determined by HCl titration with a Metrohm Titration 809 (reproducibility <2% RSD). Total dissolved sulfide ($\Sigma S^{2−}$) concentrations in the Zn acetate-treated samples from June 2015 were determined by spectrophotometry (670 nm) after direct addition of an acidic solution of FeCl$_3$ and n,n-dimethyl-p-phenylenediamine (Cline, 1969; Reese et al., 2011) to the sample vials. The procedure dissolves the ZnS precipitate and immediately complexes S as methylene blue for spectrophotometric analysis. Total dissolved sulfide concentrations were calibrated against a series of standard solutions of Na$_2$S·3H$_2$O, fixed in Zn acetate in the same manner as the samples. The exact concentration of S in the Na$_2$S·3H$_2$O stock solution was determined by iodometric titration (Burton et al., 2008).

Dissolved methane (CH$_4$) concentrations were determined by gas chromatography. Subsamples of 1 ml were taken from the headspace of each 65 ml glass bottle with a gas-tight glass syringe and transferred to evacuated 12-ml glass tubes with a butyl rubber septum (LabCo Exetainer™ model 839W). Exetainers were then pressurized with 20 ml N$_2$ (purity 5.0). The mole fraction of methane in headspace of the samples was analyzed with a FID-equipped gas chromatograph (Agilent Technologies 7890B, University of Helsinki) against a standard series of known CH$_4$ concentrations. Porewater concentrations were calculated assuming quantitative evolution of methane into the headspace from the original 10 ml wet sediment sample, using the measured porosity profile from sediment sample processing. Due to use of unpressurized coring apparatus and sampling on deck, partial degassing of CH$_4$ from the cores after recovery cannot be avoided (e.g. Thang et al., 2013; Egger et al., 2016). The saturation concentration range of CH$_4$ at in situ salinity, 1 atm pressure and range of temperatures experienced on deck is therefore given in the plots of dissolved CH$_4$ concentrations to assess the potential impact of degassing.

Absorbance of CDOM was determined using a Shimadzu 2401PC spectrophotometer with 5-cm quartz cuvette (spectral range from 200 to 800 nm with 1 nm resolution). Excitation–emission matrices (EEMs) of FDOM were determined with a Varian Cary Eclipse spectrofluorometer (Agilent). Ultrapure water was used as the blank for all samples, and EEMs were corrected and optical proxies extracted as in Asmala et al. (2018).

### Sediment Sampling, Processing and Analysis

Sediment cores were sliced on deck (GEMAX) or in the laboratory (piston) at intervals of 1 cm (uppermost 10 cm) or 2 cm (>10 cm depth) into plastic bags. Samples were stored frozen until further processing, then freeze-dried and homogenized in an agate mortar. Volumetric porosity was estimated from weight loss on freeze drying, assuming a sediment density of 2.5 g cm$^{-3}$. Dried subsamples were analyzed for total carbon and nitrogen contents by thermal combustion elemental analysis (LECO TruSpec Micro, University of Helsinki, analytical precision and accuracy <10% RSD). A subsample of 3–4 mg dried sediment was weighed into tin cups and loaded into an autosampler rosette. Inorganic carbon and nitrogen forms are considered negligible in this setting, hence measured total carbon and nitrogen concentrations are

$$J = \frac{D \cdot \varphi \cdot dC}{\theta^2 \cdot dz}$$

in which $J$ = flux (initially calculated in $\mu$mol cm$^{-2}$ s$^{-1}$; here negative values indicate downwards fluxes towards the SMTZ, and vice versa), $D$ = ion-specific diffusion coefficient, corrected for temperature, taken from Boudreau (1997), $\varphi$ = porosity and $\theta$ = tortuosity, defined as per Boudreau (1997), and $\Delta C/dz$ is the concentration gradient as given by the linear regression line.

Although fluctuation in temperature in the core profile is expected with depth in the sediments due to propagation of seasonal changes in bottom water temperature (e.g., Mogollon et al., 2011; Mueller et al., 2016), the potential error in calculated fluxes is expected to be <20% for absolute values of each species (based on the climatological range of bottom water temperature at the site (Merkouriad and Leppäranta, 2013) and <3% for the flux ratios, due to the parallel effects of temperature on all species.

For assessing the optical fingerprint of porewater DOM, the magnitude of the C-fluorescence peak (Coble, 1996) and humification index (HX; Zsolnay et al., 1999) were calculated from the measured and corrected EEMs. Absorbance spectra and EEMs were processed using the cdom and eemR packages for R software (Massicotte and Markager 2016; Massicotte 2019, respectively).
considered equivalent to the concentrations of the organic forms (C$_{org}$, N$_{org}$).

Dried subsamples were prepared for further bulk elemental analysis by a triple-acid digestion procedure. From 0.1–0.2 g of sediment was extracted overnight with 5 ml HF (45%) and 5 ml of mixed HClO$_4$ (70%)/HNO$_3$ (65%) (volumetric ratio 3:2) at 90°C in closed teflon vials. The acids were then evaporated at 160°C until samples displayed a gel-like consistency, and 15 ml 1 M HNO$_3$ was added to re-dissolve the material. Where necessary, further dilution was applied prior to analysis. Extracts were analyzed by ICP-OES (Thermo iCAP 6000, Utrecht University) for total lead (Pb$_{tot}$) according to standard procedures (Van der Veer, 2006), was determined by comparison with standard reference material ISE-921 (Van der Veer, 2006), was $\leq$15% for Fe, Mn and S and $\leq$25% for Pb.

Dried subsamples from selected depths were subjected to alkaline CuO oxidation in closed vessels in a furnace for the extraction of lignin phenols (slight modification to Hedges and Ertel, 1982). Briefly, 0.5 g sediment was added to the vessel with 1 g purified CuO and 0.5 mg Fe(NH$_4$)$_2$(SO$_4$)$_2$6H$_2$O (to scavenge O$_2$), 7.0 ml of 8% (wt/wt) aqueous NaOH, and a small stainless steel bar agitator. The oxidation procedure was carried out for 3 h at 155°C, after which the mixture was acidified to pH 1 with HCl. Organic oxidation products were then extracted with purified ethyl acetate and then concentrated, first by rotoevaporation and then to near dryness with a flow of N$_2$ over the final ~1-ml volume, in the presence of anhydrous Na$_2$SO$_4$ to remove moisture. The concentrated extract was re-dissolved in pyridine with trans-1-furoacetamide (BSTFA) as derivitizing agent. Samples were analyzed for concentrations of 8 vanillyl, syringyl and cinnamyl lignin phenols (Table 2) by GC-MS at the University of Edinburgh.

### Sediment Data Processing

A simple two-end-member mixing model was used to investigate bulk OM sources. The calculation uses only the molar N/C ratio of organic matter, and end-member values, N/C$_{EM}$, based on the studies of Goñi et al. (2003) and Jilbert et al. (2018):

$$%OC_{phyt} = \left( \frac{N/C_{sample} - N/C_{EM-terr}}{N/C_{EM-phyt} - N/C_{EM-terr}} \right) \times 100$$

$$%OC_{terr} = 100 - %OC_{phyt}$$

where %OC$_{phyt}$ and %OC$_{terr}$ are the respective contributions of phytoplankton and terrestrial material to total sedimentary organic carbon (OC), with N/C$_{EM-terr}$ = 0.04, and N/C$_{EM-phyt}$ = 0.13. The mixing model assumes that terrestrial plant matter and phytoplankton are the only sources of organic material, that their N/C values are spatially and temporally fixed at the end-member values, and that these values do not alter significantly during sedimentation and burial of organic matter.

Absolute and relative concentrations of lignin phenol oxidation products were computed according to standard notations. These include: $\sum_8$ = sum of 8 measured phenols reported as $\mu$g g$^{-1}$ sediment; $A_8$ = sum of 8 phenols reported as mg/100g C; $A_V$ = sum of 3 vanillyl phenols reported as mg/100g C (Bianchi and Canuel, 2011). Phenol compositions are expressed through ratios of the summed cinnamyl and syringyl phenols to the summed vanillyl phenols (C/V and S/V, respectively) (Table 2).

### Age vs Depth Model

An age vs depth model for the Storfjärd site was produced from the sediment Pb$_{tot}$ data, combining samples from the GEMAX core of June 2015 with the piston core of September 2017 (Figure 2). We used seven tie-points in the combined Pb$_{tot}$ profile, identified as per Brännvall et al. (1999), Zillen et al. (2012) and Jokinen et al. (2018), and assumed to represent known temporal changes in the deposition of anthropogenic Pb over the last two millennia. Age uncertainty of pre-20th century tie-points was set at 25 years, while that of the 1970 tie-point was set at 10 years. Depth uncertainties were set at 4–8 cm. The model was produced from the tie-points using the Undatable software of Lougheed and Obruchta (2019), applying 10$^5$ simulations with an xfactor of 0.1 and 15% bootstrapping. Of these, the xfactor dictates the maximum allowable variation in sediment accumulation rate between all pairs of age-depth constraints, while the bootstrapping percentage determines the number of tie-points that are randomly excluded from the simulations. The approach takes into account uncertainty in both age and depth, with uncertainty allowed to increase with distance from tie-points, and uses a Bayesian approach to estimate a probability-density cloud for the upper 310 cm of the sediment. A subset of tie-points in the Pb$_{tot}$ profile were checked by comparison with $^{206}$Pb/$^{207}$Pb data, but the latter were not used in the construction of the model due to the observed low signal-to-noise ratio in the profile. Linear sedimentation rate (LSR) and mass accumulation rate (MAR) were estimated for each segment between tie-points using the measured porosity profile. Accumulation rate of OC$_{terr}$ was estimated for each sampled interval from the estimated content of OC$_{terr}$ (Eq. 4) and MAR of the corresponding segment.

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### Table 2: Lignin phenol oxidation products analyzed in this study. Codes indicate provenance as described by Hedges and Ertel (1982). G = gymnosperms, woody tissue; g = gymnosperms, non-woody tissue; A = angiosperms, woody tissue; a = angiosperms, non-woody tissue.

| Vanillyl phenols (V) | Syringyl phenols (S) | Cinnamyl phenols (C) |
|---------------------|----------------------|----------------------|
| GgAa Vanillin        | Aa Syringaldhyde     | ga Commaric acid     |
| GgAa Acetovanillione | Aa Acetosyringone    | ga Ferulic acid      |
| GgAa Vanillic acid  |                      |                      |

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RESULTS

Sedimentation Rate and Mass Accumulation Rate

The age vs depth model shows that the 4-m studied interval covers at least the last two millennia (Figure 2), with the earliest dating tie-point identified as 1 AD (as per Zillen et al., 2012). Both LSR and MAR have varied over the studied interval, showing ranges of 0.1–0.5 cm/yr and 0.04–0.15 g/cm²/yr, respectively. The most recently deposited interval (1970–present) is characterized by relatively high values compared to the preceding 500 years (LSR = 0.49 cm/yr, MAR = 0.13 g/cm²/yr), indicating enhanced inputs of sediment at this site in the recent past. One earlier interval of comparably high LSR and MAR values is observed deeper in the sediment column, corresponding to 1200–1350 AD.

Solid-Phase Profiles

Major changes occurred in bulk composition during the deposition of the uppermost meter of the sediments, corresponding to the interval since AD 1600 (Figure 3). Most importantly, this interval is characterized by an increase in organic matter content, as evidenced by elevated Corg values.
up to 4–6% by weight (relative to <4% deeper in the sediments). Changes in the inputs of both terrestrial and phytoplankton-derived OM are responsible for this general increase, although the evolution of the two components since AD 1600 differs markedly (Supplementary Figure S1). While OCphyt. shows fairly stable contents until ~AD 1900 and a steady rise thereafter, OCterr. shows a steady rise from AD 1600 to ~AD 1850, followed by two abrupt increases between the mid-19th century and mid-20th century. Here we note that the absolute ages of events within the interval between the tie-points of AD 1600 and AD 1970 are subject to uncertainty as defined by the age vs depth model (Figure 2). In particular, it is plausible that sedimentation rate began to increase in parallel with changes in sediment composition, whereas the age model assumes linear sedimentation between the two tie-points. This would have the effect of making the estimated dates too old in the interval of most rapid compositional changes.

When the contributions of OCphyt. and OCterr. are combined, we observe that a significant fraction of total Corg in the late 20th century sediments (up to 30% according to Eq. 3 and Eq. 4) is provided by terrestrial material (Figure 3). This fraction is far higher than in the deeper sediments, indicating a relative increase in OCterr. input over time. Furthermore, the absolute accumulation rate of OMterr. during this interval (up to 0.002 gcm⁻²yr⁻¹) is unprecedented in the record (Figure 3). The accumulation rate of OMterr. has declined since the late 20th century maximum but the surface-sediment C/N ratio, and thus the calculated contribution of OMterr., remains elevated with respect to the deeper sediments.

Similar profiles to that of OCterr. are observed in the ratios of syringyl to vanillyl (S/V) and cinnamyl to vanillyl (C/V) phenols. Both the S/V and C/V phenol ratios show pronounced minima during the interval of maximum accumulation rate of OMterr. (Figure 3). However, the absolute ranges of S/V (0.1–0.6) and C/V (0.3–1.0) are similar to those observed in previous studies of lignin phenols in Baltic Sea sediments (Miltner and Emeis, 2001). Iron (Fe) and manganese (Mn) contents show a parallel evolution throughout the sediment record. Both elements’ contents show a general increase from the core base upwards, towards maximum values of 6.7% (Fe) and 0.1% (Mn) within the interval of maximum OMterr. accumulation. Contents of Fe and Mn decline from this layer towards the sediment surface but the uppermost sample shows elevated values for both elements. Sulfur (S) displays a distinctly different profile, with higher variability throughout the record, and a pronounced maximum at 11 cm depth, clearly shallower than the maxima in OMterr., Fe and Mn.

Porewater Profiles

Porewater NH₄⁺ and Alk₂ show increasing concentrations with increasing depth in the sediment column, towards values of approximately 10 mmol/L and 50 mmol/L, respectively at 400 cm depth (Figure 4). The degree of curvature in both profiles is greatest in the uppermost 50 cm of the sediments, while the gradients below this depth are quite linear. Porewater CH₄ concentrations show a distinct maximum of >4 mmol/L at approximately 25 cm depth, very close to the 1970 horizon at the center of the layer of enhanced OMterr. deposition (Figure 3). A steep upwards gradient in CH₄ concentration is detectable porewater ΣS²⁻ concentrations and encompassing the depth of equal concentrations of CH₄ and S (assumed equivalent to SO₄²⁻). Below the porewater CH₄ maximum, a reverse gradient is observed towards background concentrations of ~1 mmol/L below 3 m depth. We cannot rule out that this gradient is an artefact of degassing effects due to the use of non-pressurized coring apparatus, and that true porewater CH₄ concentrations at depth are much higher as described in e.g. Egger et al. (2016). The increasing NH₄⁺ and Alk₂ concentrations below this horizon would support such a hypothesis, since all three species are
produced during methanogenesis. However, it is also possible that
the profile is genuine, and that the linear NH₄⁺ and Alk₂F
gradients reflect upwards diffusion of these species from a
deep source. We note that measured CH₄ concentrations
decline to values well below the saturation concentration range
during sampling on deck (1.6–2.0 mmol/L, Figure 3), whereas
degassing typically leads to a plateau of values close to this range
(Egger et al., 2016). This observation would support the
hypothesis of a genuine reverse gradient in porewater CH₄.
Above the SMTZ, concentrations of dissolved S (assumed
equivalent to SO₄²⁻) rise steeply towards the bottom water
concentration of approximately 6 mmol/L.

Porewater δ¹³C-DIC values show a minimum of −8.5‰ close
to the SMTZ, while the uppermost measured sample at 1 cm
depth has a value of −2.8‰. Below the SMTZ, δ¹³C-DIC values
show an asymptotic increase towards a stable value of +18–+19‰
in the deeper sediments. In terms of the CDOM optical properties
of the porewaters, the peak C (considered to approximate the
concentration of humic-like DOM; Coble, 1996) shows a
generally concave profile similar to those of NH₄⁺ and Alk₂F.
In contrast, the humification index (HIX; representing the extent
of humification of the CDOM pool, Zsolnay et al., 1999) shows
raised values of up to 14 in the interval of enhanced OMₚterr.
accumulation, superimposed on a trend from lower values at the
surface (<6) to a consistent background of 10–12 at depth.

DISCUSSION
Terrestrial Organic Matter Loading to the
Sediments During the 20th Century
Many coastal areas of the Baltic Sea show evidence for
anthropogenic eutrophication (Conley et al., 2011) which often
leads to enhanced carbon loading to sediments (Jokinen et al.,
2018; Helmond et al., 2020). At Storfjärden, we observe a steady
rise in OCₚhyt contents in the period since AD 1900
(Supplementary Figure S1), consistent with eutrophication.
However, a novel observation of this study is that a significant
proportion of the additional recent carbon at this site is derived
from terrestrial sources. The coincident minima in C/V and S/V
phenol ratios during the period of maximum OMₚterr.
accumulation rates indicate a shift towards more gymnosperm-
dominated material (lower S/V phenol ratio) simultaneously with
a shift towards more woody material (lower C/V phenol ratio, see
Table 2), implying inputs of material originally derived from the
Finnish forests.

Recent changes in C/V and S/V phenol ratios in estuarine
settings have previously been interpreted as evidence for inputs of
lignin-rich waste products from pulp and paper industries in
North America (Louchouarn et al., 1999; Brandenberger et al.,
2011). Such an interpretation appears plausible also for
Storfjärden. Pulp production at Lohja, which peaked in the
catchment of the River Mustionjoki (Figure 1), began in 1907 while paper
production began in 1938 Heikkinen (2000). By 1957, the site
was operating two paper machines with a combined potential
output of 70 kt/y. Considering the uncertainty of the age vs depth
model in this interval, the abrupt increases in OCₚterr. observed in

Supplementary Figure S1 could well relate to these major
changes in industrial activity in the catchment. We also note
that sedimentary contents of OCₚterr. were already increasing for
several centuries prior to this time, indicating additional inputs
potentially from land use change prior to the onset of industrial
activity (e.g. Yang et al., 2021).

Emissions of waste products from forest industries are
expected to be a function of the scale of operations, as well as
the degree of waste water treatment, both of which may change
over time (Katko et al., 2005). Data from the website of Finnish
Forest Industries (https://www.forestindustries.ﬁ/statistics/
environment/), compared with the dated sediment record of total vanillyl
phenols, reported as mg/100 g C (Λᵥ). TSS = total suspended solids,
BOD₇ = 7th day biological oxygen demand.

FIGURE 5 | Time series of production and emissions from forest
industries in Finland since 1950 (https://www.forestindustries.ﬁ/statistics/
environment/), compared with the dated sediment record of total vanillyl
phenols, reported as mg/100 g C (Λᵥ). TSS = total suspended solids,
BOD₇ = 7th day biological oxygen demand.

Frontiers in Earth Science | www.frontiersin.org
October 2021 | Volume 9 | Article 716416
9
20th century lignin accumulation with maximum emissions from forest industries in Finland (Figure 5, note that age vs depth uncertainties are lower in this interval due to the tie-point at 1970). Furthermore, Heikkinen (2000) reported high contents of sedimentary resin acids during the same period in the southern part of Lake Lohjanjärvi, adjacent to the industrial site (Figure 1), while high contents of OMterr. in sediments close to Storfjärden were reported by Jokinen et al. (2020). Combined, these studies strongly support the interpretation that emissions from industries in Lohja were the main reason for the high inputs of OMterr. to Storfjärden during the late 20th century. This observation is remarkable, as it implies that significant amounts of waterborne waste materials were transported through Lake Lohjanjärvi, the Mustionjoki river and Pojo Bay as far as the archipelago area where the study site is situated, a linear distance of over 60 km (Figure 1).

An ongoing mapping exercise in the Swedish coastal zone has identified 29 locations of fiberbank deposits close to point source industrial sites (Norrlin and Josefsson, 2017). In terms of surface area at any given location, true fiberbanks (sediments dominated by coarse waste material fibers) typically occupy up to 10 hectares, while fiber-rich sediments (sediments with visible fibers) may extend over up to 1 square kilometer. Our data raises the possibility that the true spatial extent of anthropogenic OMterr. inputs to sediments of the northern Baltic Sea may be larger, due to the transport of finer particulate and dissolved material away from point sources. Using the Swedish coastal area as an example due to the good data coverage, we performed a speculative mass balance calculation to investigate the scale of carbon accumulation in mapped fiberbank locations in relation to total inputs from forest industry sources (Table 3). From the reported sediment carbon concentrations in Dahlberg et al. (2020), we estimate that 0.34 million tonnes of carbon currently reside in the fiberbanks and fiber-rich sediments at the 29 mapped locations of Norrlin and Josefsson (2017). We subsequently derived a first-order estimate for the magnitude of forest industry carbon inputs to the Baltic Sea from Sweden. This estimate is based on an assumed Gaussian distribution of total COD emissions from Swedish forest industries over the period 1930–2000, fitted to the existing data from the Swedish Forest Industries Federation (Supplementary Figure S2) and corrected downwards assuming that only 5–15% of Swedish forest industry point source emissions enter the Baltic Sea, based on the locations of point sources in the maps of Norrlin and Josefsson (2017). The conversion from COD to total organic carbon (TOC) is based on the regression for high-latitude systems presented by Jiao et al. (2021). This exercise yields a total carbon input of 3.2–9.5 million tonnes, up to an order of magnitude greater than the estimated carbon stock in the 29 mapped locations (Table 4). Approximately half of the expected Swedish coastal point sources to the Baltic Sea have been mapped, thus its seems likely that the total fiberbank and fiber-rich sediment carbon stock in the Swedish coastal zone, once known, will be significantly less than the total input of carbon from anthropogenic OMterr. during 1930–2000. In this context, a diffuse contribution of anthropogenic OMterr. to a larger coastal sediment area is plausible, although part of the difference may be explained by mineralization and other transformations in the coastal environment.

Additionally, we estimated the quantitative significance of anthropogenic OMterr. inputs in the context of total terrestrial carbon loading to the Baltic Sea from Finland and Sweden. During the period 1993–2012, when standardized TOC analysis protocols were routinely used and therefore data are most reliable (Asmala et al., 2019), annual TOC loading to the Baltic Sea from the two countries combined varied from approximately 1.0–2.5 million tonnes per year (Table 4). Using the same Gaussian approach described above to provide a first-order estimate of the historical point source anthropogenic OMterr. inputs from Finland (Supplementary Figure S2) we calculate a total loading during the period 1930–2000 of approximately 4–13 million tonnes of TOC for the two countries combined. During the loading maximum of the mid-1960s, we estimate approximately 0.18–0.53 million tonnes TOC per year. In conclusion, during the period of maximum emissions, carbon inputs from industrial sources may have been a significant component (up to >10%) of total annual carbon loading from Finland and Sweden, although the contribution likely declined steeply during the last decades of the 20th century.

**Overall Impact of Carbon Loading on Sediment Diagenesis**

The additional carbon loading from both aquatic primary production and terrestrial sources has contributed to higher LSR and MAR at Storfjärden in the recent past, and driven high rates of remineralization in the upper sediments, as
shown by the curvature of the NH$_4^+$ and Alk$_T$ profiles in the upper 50 cm. The porewater data suggest that enhanced demand for electron acceptors has driven a rearrangement of the diageneric zones in the upper sediment column (e.g., Middelburg and Leen, 2009). Most notably, carbon loading has enhanced methanogenesis and consequently shoaled both the interval of organoclastic sulfate reduction and the SMTZ. This sequence of eutrophication-driven changes in diageneric zonation in the northern Baltic Sea has been described in several previous studies (e.g., Slomp et al., 2013; Egger et al., 2016; Rooze et al., 2016; Jilbert et al., 2018). A key piece of evidence for non-steady state conditions – the solid-phase S maximum indicating enhanced formation of sulfide minerals in the modern SMTZ – is observed in our profiles at 11 cm depth (Figure 3). Our data suggests that the methanogenic horizon is focused in the recently deposited sediments, with a gradient towards lower CH$_4$ concentrations observed in deeper layers. Even in the case of degassing impacting on the methane profile, the strong curvature of the NH$_4^+$ and Alk$_T$ profiles in the upper 50 cm supports the theory that this interval is characterized by enhanced rates of methanogenesis relative to the rest of the sediment column. This confirms that enhanced methane production, and associated effluxes to the water column, are truly a legacy effect of recent carbon loading as postulated by Myllykangas et al. (2020a) from short-core data. Indeed, promotion of methanogenesis through rapid accumulation of organic material in sediments has been described previously in both freshwater and estuarine systems (e.g., Egger et al., 2016; Steinsberger et al., 2017). In the northern Baltic Sea, naturally low sulfate concentrations and modern high sedimentation rates allow degradable organic material to pass through the zone of sulfate reduction within the uppermost decimeters of the sediment column, thereby facilitating high rates of methanogenesis (e.g., Thang et al., 2013; Sawicka and Bruechert, 2017).

**Methanogenesis, Organoclastic Sulfate Reduction and Anaerobic Oxidation of Methane**

The shoaling of the diageneric zones has triggered a cascade of secondary redox reactions in the sediment column at Storfjärden. Most importantly, the presence of methane in the shallow porewaters has likely led to the initiation of anaerobic oxidation of methane (AOM). Elevated rates of methane oxidation in the SMTZ at this site have been previously suggested from porewater data (Jilbert et al., 2018) and later confirmed by $^{14}$CH$_4$ incubations (Myllykangas et al., 2020b). The present study shows the enrichment of $\Sigma S^{\text{aq}}$ at the SMTZ reported in Jilbert et al. (2018) in the context of the wider porewater data (Figure 4). The enrichment is considered to indicate sulfide production via sulfate-mediated AOM (S-AOM):

$$\text{CH}_4(aq) + \text{SO}_4^{2-}(aq) + \text{CO}_2(aq) \rightarrow \text{H}_2\text{S}(aq) + 2\text{HCO}_3^-(aq) + \text{H}_2\text{O}(l)$$

To further investigate the dynamics of methanogenesis, organoclastic sulfate reduction and S-AOM at Storfjärden, we determined the $\delta^{13}$DIC of porewaters throughout the core profile. Porewater $\delta^{13}$DIC is influenced by each of these processes as described by Whiticar (1999) and Meister and Reyes (2019). Namely, organoclastic sulfate reduction depletes $\delta^{13}$DIC from the bottom water value of ~0‰ towards the value of decaying organic matter (~−20‰). If S-AOM is active at the SMTZ, this process further depletes $\delta^{13}$DIC due to consumption of isotopically light methane. In the underlying methanogenic zone, in contrast, $\delta^{13}$DIC becomes enriched due to preferential consumption of

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**TABLE 4 | Estimate of total carbon emissions to the Baltic Sea from forest industry point sources in Finland and Sweden during 1930–2000.**

|                      | Finland     | Sweden      | Total       |
|----------------------|-------------|-------------|-------------|
| **Forest industry emissions 1930–2000** |             |             |             |
| Modeled integrated BOD$^a$ or COD$^b$ 1930–2000 (kilotonnes) | 9496        | 69973       |             |
| Modeled integrated TOC$^c$ 1930–2000 (kilotonnes) | 24350       | 63612       |             |
| Modeled integrated TOC to Baltic Sea 1930–2000 (kilotonnes, 5%)$^d$ | 1217        | 3181        | 4398        |
| Modeled integrated TOC to Baltic Sea 1930–2000 (kilotonnes, 15%)$^e$ | 3652        | 9542        | 13194       |
| Modeled max. annual TOC to Baltic Sea 1965 (kilotonnes, 5%) | 49          | 127         | 176         |
| Modeled max. annual TOC to Baltic Sea 1965 (kilotonnes, 15%) | 146         | 380         | 527         |

**Total riverine carbon inputs to Baltic Sea**

|                      | High        | Low         | Total       |
|----------------------|-------------|-------------|-------------|
| TOC to Baltic Sea 1993–2012 (kilotonnes/yr)$^f$ | 1148        | 404         | 2547        |
|                      | 1399        | 547         | 961         |

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$^a$Total emissions in Finland, modeled using data for BOD$_7$ from Finnish Forest Industries from 1950 to 2000 (Figure 5) with hindcasting to 1930 assuming a Gaussian distribution (Supplementary Figure S2).

$^b$Total emissions in Sweden, modeled using data for COD from Swedish Forest Industries Federation from 1978 to 2000 and hindcasted to 1930 assuming a parallel Gaussian distribution to that observed in Finland (Supplementary Figure S2).

$^c$Estimated using the regressions for BOD and COD vs DOC in high-latitude systems presented in Jiao et al. (2021), assuming DOC = TOC.

$^d$5% of modeled integrated TOC emissions 1930–2000.

$^e$15% of modeled integrated TOC emissions 1930–2000.

$^f$Data from Asmala et al. (2019)

$^g$Data from Baltic Environmental Database.
The relationship between $\Delta$[DIC]/$\Delta$[SO$_4^{2-}$] and [DIC] * $\delta^{13}$C-DIC strongly varies with depth in the upper sediment column (Figure 6). Samples from the SMTZ plot in the lower-right corner of the diagram, with $\Delta$[DIC]/$\Delta$[SO$_4^{2-}$] values < 2 coincident with negative [DIC] * $\delta^{13}$C-DIC. Although the [DIC] * $\delta^{13}$C-DIC values are less negative than observed in Miller et al. (2017) due to the diluting effect of methanogenesis in this system, the progression towards $\Delta$[DIC]/$\Delta$[SO$_4^{2-}$] values < 2 supports the hypothesis that sulfate-AOM occurs in the SMTZ. However, the minimum detected value for $\Delta$[DIC]/$\Delta$[SO$_4^{2-}$] at 13 cm depth is 1.65, still significantly elevated with respect to the 1:1 value predicted from sulfate-AOM alone. This result suggests that an important fraction of total sulfate consumption is in the SMTZ is likely contributed by organoclastic sulfate reduction (e.g., Jorgensen et al., 2019). This is further supported by the relative diffusive fluxes of SO$_4^{2-}$ and CH$_4$ into the SMTZ, which show a ratio of 2.66 (Table 5), clearly in excess of the 1:1 requirement of the reactants of sulfate-AOM (Eq. 5). Combined, the results imply that the SMTZ at our study site should be considered as a broad zone of overlapping diagenetic processes, in which methanogenesis, organoclastic sulfate reduction and S-AOM are all active simultaneously.

Consequences of the Shallow Sulfate-Methane Transition Zone at Storfjärden

The SMTZ at Storfjärden, similarly to much of the northern Baltic Sea today, is among the shallowest in the marine realm. Due to the strong correlation observed globally between SMTZ depth and associated diffusive fluxes of SO$_4^{2-}$ and CH$_4$ (Egger et al., 2018), the values of these fluxes at Storfjärden are also high in global terms. Our calculated values for $J$CH$_4$ and $J$SO$_4^{2-}$, as well as the flux ratio of 2.66, fall within the ranges of compiled data for 40 marine coring sites with SMTZ depth < 1 m (Table 5), suggesting that rates of methane-related processes at this site are typical for similar coastal locations of high carbon loading in the modern ocean.

While consumption of upwards-diffusing methane by AOM in marine sediments is considered near-quantitative on a global scale (Reeburgh, 2007; Saunois et al., 2016), sites with a shallow SMTZ often show inefficient AOM and significant fluxes of CH$_4$ to the water column and hence potentially to the atmosphere (e.g. Thang et al., 2013; Egger et al., 2016). Indeed, the Storfjärden area today is characterized by active fluxes of methane from sediments through both diffusion (Myllykangas et al., 2020a) and ebullition (Humborg et al., 2019). Our data confirms that the ultimate source of these emissions is the methanogenic layer in the late 20th century sediments, and hence that these emissions are a direct consequence of anthropogenic carbon loading through both eutrophication and inputs of OM$_{terrestrial}$.
Table 5: Comparison of fluxes of methane (JCH4) and sulfate (JSO4^2-) into the SMTZ at Storfjärden with 40 other locations in the marine realm with SMTZ depth < 1 m (data from Egger et al. (2018)). Only sites with data for all parameters were used in the analysis.

| SMTZ depth (m) | JCH4 (mmol/m^2/d) | JSO4^2- (mmol/m^2/d) | JSO4^2-/JCH4 |
|----------------|-------------------|----------------------|--------------|
| Storfjärden, Baltic Sea (this study) 40 sites of SMTZ < 1 m depth Egger et al. (2018) | 0.12 | 1.73 | 4.59 | 2.66 |
| mean | 0.44 | 1.81 | 3.11 | 1.38 |
| min | 0.04 | 0.07 | 0.06 | 0.60 |
| max | 1.00 | 15.93 | 13.07 | 3.69 |

Specific Impacts of OM_{terr}. Inputs on Diagenetic Processes

Our data show indications that the additional loading of OM_{terr} during the 20th century has had specific impacts on diagenetic processes at Storfjärden. Terrestrial OM is traditionally considered to be relatively inert to remineralization in sediments (Hedges et al., 2000; Arndt et al., 2013) due to its refractory macromolecular composition (de Leeuw and Largeau, 1993) as well as protective associations formed with other materials prior to sedimentation (e.g., Hedges and Keil, 1995; Huguet et al., 2009). Indeed, lignin in sediments has been studied extensively as a biomarker for OM_{terr} largely due to its high preservation potential (Bianchi et al., 2018). However, several studies have reported that sites influenced by large direct inputs from forest industry display evidence for relatively high reactivity of this material in the sediment column. For example, high potential rates of methanogenesis were recently observed in wood fiber-rich sediments from the Finnish Lake Nasijärvi (Kokko et al., 2018). Similarly, fiber-rich sediments in Swedish coastal areas of the Baltic Sea support diffusive fluxes of organic contaminants to overlying waters, implying release from the solid-phase during diagenesis (Dahlberg et al., 2021).

At Storfjärden, we observe a maximum in the humification index (HIX) of porewater CDOM (values up to 14) coincident with the layer of enhanced OM_{terr} accumulation (Figure 4). This maximum is superimposed onto a trend from lower values at the surface (<6) to a consistent background of 10–12 at depth in the profile. Previous studies of HIX in sediment porewaters have typically interpreted this index as a proxy for CDOM sources in the degrading sedimentary OM, with low values (e.g. <3) indicative of autochthonous or microbial material and high values (e.g. >6) indicative of terrestrial humic matter (Chen et al., 2016; Li et al., 2021). Our values are generally higher than those reported in these studies, implying an overall more humic composition of the source material. Furthermore, maximum values are observed in the layer of maximum OM_{terr} accumulation, suggesting solubilization of lignin and other terrestrial macromolecules, which are important components of humic-like CDOM (Del Vecchio & Blough, 2004). The underlying trend in HIX is also of interest. The observation of low-HIX CDOM in the shallowest layers and a stable background of high-HIX CDOM at depth implies a diagenetic shift in CDOM composition after sedimentation of degrading OM. Specifically, low-HIX CDOM produced during degradation in the shallowest layers appears to be consumed after release into the porewaters, leaving a residual accumulation of high-HIX CDOM at depth. We suggest that such consumption may occur through utilization of low-HIX DOM, which is considered more labile, by the microbial community. We note that the HIX profile contrasts with that of peak C, which shows a concave-down profile throughout the sediment column (Figure 4). This suggests that overall production of humic-like CDOM occurs similarly to that of inorganic degradation products such as NH4^+ and Alk_T.

The additional flux of OM_{terr} to the coring location during the 20th century was likely accompanied by other terrestrial materials transported in association with organic matter. As described by Jokinen et al. (2020), metals such as Fe, Mn, Co, Cd, Pb, Sn and Zn are all enriched in the same depth interval as OM_{terr} in sediment cores from the Storfjärden area. Our long core data confirm that contents of Fe and Mn are higher in this layer than at any other depth in the sediment column (Figure 3). These metals derive from the terrestrial environment (including from anthropogenic sources), and are sedimented both as oxide minerals physically associated with organic matter, and as metal-OM complexes that undergo flocculation and aggregation at the estuarine salinity gradient (Widerlund and Ingri, 1996; Jilbert et al., 2018). Temporal changes in the inputs of oxides and metal-OM complexes may impact on diagenetic processes in several ways. For example, higher inputs of oxides may affect rates of oxide-mediated AOM (Lenstra et al., 2018) and thus the potential for phosphorus retention through vivianite formation (Slomp et al., 2013). Furthermore, complexation between OM and Fe in sediments has been shown to generally protect OM from remineralization and thus promote long-term carbon burial (Lalonde et al., 2012; Shields et al., 2016). Hence, not only the direct OM_{terr} inputs from forest industry themselves, but also the associated materials, may have altered diagenetic processes at Storfjärden during the 20th century.

Conclusion

This study shows that recent carbon loading to coastal sediments of the northern Baltic Sea has occurred not only as a consequence of increased aquatic primary production due to increased nutrient loading, but also of enhanced inputs of terrestrial organic matter from forest industry activities. This additional material has contributed to high sedimentation rates and demand for electron acceptors for organic matter remineralization, leading to a shoaling of the diagenetic zones in the sediment column. At the Storfjärden site, a distinct layer of elevated OM_{terr}-contents in the late 20th century sediments are characterized by lignin phenol signatures typical of woody gymnosperm material, implying inputs from forest industries in the catchment. This layer coincides with high porewater methane concentrations and
high curvature in the profiles of NH₄⁺ and AlkT, indicating that the carbon loading has enhanced methanogenesis in the recently deposited sediments. Moreover, optical characteristics of porewater CDOM indicate active degradation of OMterr in the same layer. The SMTZ is observed directly above the methanogenic layer, and is characterized by sulfate-mediated AOM occurring alongside organoclastic sulfate reduction and methanogenesis. Fluxes of methane and sulfate into the SMTZ are high, but typical for similar eutrophic systems throughout the coastal oceans.

The broader spatial impact of anthropogenic OMterr inputs on coastal sediment biogeochemistry in the northern Baltic Sea remains to be established. Our data from Storfjärden shows that the signatures of such inputs may be identifiable several tens of kilometers from the industrial sources, confirming previous observations from the St. Lawrence estuary (Louchouarn et al., 1999). Moreover, our estimates of carbon storage in fiberbank locations in the Baltic Sea show that these proximal deposits are unlikely to account for the entire carbon loading from forest industry sources to the Baltic Sea during the 20th century. Determining the wider extent of anthropogenic OMterr-rich deposits in boreal coastal zones is essential for constraining carbon budgets in these areas, both in terms of carbon burial and greenhouse gas emissions.

DATA AVAILABILITY STATEMENT

All new data presented in the paper are accessible via Zenodo from 1 December 2021 (10.5281/zenodo.5570261) or directly from the corresponding author without undue reservation.

AUTHOR CONTRIBUTIONS

TJ devised the study, led the field and lab work and wrote the manuscript. GC and LL performed lignin phenol analyses and contributed to the interpretations and writing. SJ produced the age vs depth model and contributed to the interpretations and writing. EA performed the porewater CDOM analyses and contributed to the interpretations and writing. XS and CM performed the porewater δ¹³C and Alkalinity analyses and contributed to the interpretations and writing. CH and AN co-ordinated the field campaigns with R/V Electra and contributed to the interpretations and writing.

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SUPPLEMENTARY MATERIAL

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