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# The interplay between terrestrial organic matter and benthic macrofauna: Framework, synthesis, and perspectives

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#### Abstract

Ecosystems are shaped by physical, chemical, and biological drivers, which affect the quality and quantity of basal energy sources, with impacts that cascade to higher trophic levels. In coastal, shelf, and marine habitats, terrestrial-derived organic matter (ter-OM) can be a key driver of ecosystem structure and function. Climate change is expected to alter land-ocean connectivity in many regions, with a broad range of potential consequences for impacted ecosystems, particularly in the coastal zone. The benthic compartment is an important link between the large organic carbon pools stored on land and the marine environment. At the same time, the macrofauna plays a key role in the processing, biological uptake, and fate of ter-OM in the aquatic environment, with implications for coastal ecosystem functioning, benthic-pelagic coupling, carbon burial, and biogeochemical cycles. However, information about relationships between land-ocean connectivity (including ter-OM loads) and coastal benthic community responses remains spread across disciplines, and a broad perspective on the potential impacts of a changing climate is still missing. Here, we explore the interplay between benthic macrofaunal communities and ter-OM through a paired narrative and research weaving analysis, which combines systematic mapping and bibliometric analysis. The review describes the past development and status of the research field as well as the lack of information in some geographical regions and habitats worldwide. We highlight the role of macrofauna in carbon cycling and the growing evidence that ter-OM plays a key role in the structure and function of benthic communities, not strictly limited to estuarine habitats. Climate change poses challenges for the prediction of future ter-OM fluxes and potential macrofauna responses to this additional stressor, thus requiring new methodological approaches (e.g., multimarker approaches for OM characterization) and long-term monitoring programs across different habitats and spatiotemporal scales.

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## K E Y W O R D S

benthic communities, climate change, land-ocean interactions, macrobenthos, organic carbon, riverine inputs

# INTRODUCTION

The exchange of material and nutrients across ecosystem boundaries can play a key role in shaping habitats as well as ecosystem structure and function (Loreau et al., 2003). This is particularly true for the organisms living at the land-ocean interface (Gounand et al., 2018; Polis et al., 1997; Stoler & Relyea, 2020). Organic matter (OM), with its diverse structure and nature, is central to ecosystem processes and an essential driver of marine pelagic and benthic community structure (Baldock et al., 2004; Fernández-Rodríguez et al., 2019; Lautenschlager et al., 2014). In coastal, shelf, and marine environments, terrestrial-derived organic matter (ter-OM) is an important factor in shaping the relationship and interactions between fauna and the environment as an essential energy source for many taxa as well as a potential stressor to others (Asmala et al., 2013; Bianchi, 2011; Capelle et al., 2020; Cragg et al., 2020). For many years, the origin and fate of ter-OM, and its interaction with other environmental stressors typically associated with river runoff (as a major source of ter-OM to the coastal zone), have been a topic of interest for researchers, policy makers, and civil society (De Wit et al., 2016; Deuser, 1988; Hedges et al., 1997). Thus, the understanding of the role of ter-OM in structuring ecosystems and connectivity between land and sea is a central ecological question (Asmala et al., 2013; Deininger & Frigstad, 2019; Jokinen et al., 2020; Witt et al., 2012).

Natural (i.e., nonanthropogenic) ter-OM is composed of a mix of vascular plants along with (black) carbon from soil and eroded carbonate rocks (Bianchi, 2011 and literature within). Recent and ancient OM stored in land soil, along with vascular plants and other sources, corresponds to the largest storage of carbon, one of the main constituents of ter-OM (Bianchi, 2011). A considerable amount of this material reaches the aquatic benthic compartment, turning it into an important sink for sedimentation and accumulation of land-derived carbon and nutrients (Burdige, 2007; Hedges et al., 1997), where as much as approximately one third of the OM buried in marine sediments is indeed land-derived (Burdige, 2005).

Benthic macrofauna has been included as a reliable indicator in short- and long-term environmental studies and monitoring programs (Jędruch et al., 2019; Trannum et al., 2018). The group is a key component in ecosystem functioning and services by supporting productivity of coastal habitats, nutrient cycling, carbon mineralization, and trophic chains (Griffiths et al., 2017; Janas et al., 2019; Volkenborn et al., 2007). Benthic communities rely on resources coming from within (autochthonous) and/ or outside (allochthonous) their natural habitats. There is growing evidence for the use of less-nutritious ter-OM as food source by macrofaunal groups, varying between habitats and seasons (Attrill et al., 2009; Bianchi et al., 2021; Harris et al., 2018). Thus, the increasing trend of ter-OM inputs in some regions may lead to substantial bottom-up ecosystem changes (Drylie et al., 2020; Fernández-Rodríguez et al., 2019).

Benthic macrofauna is considered a crucial component of climate mitigation and adaptation due to its influence on soil biogeochemistry and carbon mineralization pathways (Bianchi et al., 2021; Solan et al., 2020). To support the understanding of the interplay between estuarine, coastal and shelf macrofauna, and ter-OM dynamics, we have, first, provided a narrative review of the current knowledge on the origin, composition, fate, and impacts of ter-OM on macrofauna functioning and structuring, by exploring the pathways of ter-OM from its source until reaching the benthos, and the outcomes of this interrelationship at species and community levels. Although discharge from farms and municipal wastewater remains the major source of OM from land in some regions worldwide (Mudge & Duce, 2005; Wassmann & Olli, 2005), our review focuses on natural sources of ter-OM. Secondly, we provide a research weaving analysis through a bibliographic review and systematic mapping (Nakagawa et al., 2019). This methodology allows us to synthesize the current knowledge, identify emerging research themes, and highlight critical knowledge gaps (Current status of the research topic: ter-OM and macrofauna communities). Lastly, we provide perspectives and potential future directions on the topic, mainly focusing on contemporary challenges under the prism of climate change (Emerging themes on the research topic).

# THE ORIGIN AND FATE OF ter-OM

# Sources of ter-OM to aquatic habitats

## Composition of ter-OM

Natural OM is delivered in the form of dissolved or particulate OM (DOM and POM, respectively) (He et al., 2016 and literature within). DOM is a major marine carbon reservoir (~662 Gt) (Hansell et al., 2009; He et al., 2016). The molecular structure of marine DOM depends on its source and previous transformation processes, and plays a key role in determining both the bioavailability (quality) as well as the light-attenuating properties of DOM. The light absorbing, or chromophoric, fraction of DOM is referred to as colored DOM (CDOM), and especially present in the surface and coastal ocean influencing both water color and light quality (Deininger & Frigstad, 2019; Nelson & Siegel, 2013).

OM from different terrestrial sources (e.g., plant end-members, primary and secondary producers) and in different degradation stages interact in the coastal zones (Massicotte et al., 2017; McGovern, Pavlov, et al., 2020). Terrestrial and marine OMs have distinct chemical patterns due to the different primary producers (e.g., multicellular plants vs. marine phytoplankton) and biogeochemical processes (Baldock et al., 2004; Burd et al., 2016 and literature within). Although difficult to generalize, ter-OM tends to have macromolecules with more complex structural polymers such as lignin, cellulose, peptidoglycans, and algaenans than marine resources, usually leading to a higher carbon-to-nitrogen ratio (Burd et al., 2016; Burdige, 2007; Dickens et al., 2006), although some marine components (e.g., macroalgae, seagrasses, salt marshes) may show similar values complicating the attribution to various OM sources (see *Emerging themes of the research topic*). Proteins (amino acids) and lipids represent a major fraction of marine OM components, followed by carbohydrates, which are more abundant in vascular plants (~70% of the source) (Burdige, 2007).

The major component of ter-OM is soil-derived material containing POM (Baldock et al., 2004; Bianchi, 2011), but its mixed origin makes it difficult to fit in a simple classification. Along with organic carbon, ter-OM includes nitrogen, phosphorous, and other nutrients and macromolecules (Meybeck, 1982: Wakeham & Canuel, 2006). The varied quality and quantity of compounds mirror the different sources of ter-OM. A considerable fraction of ter-OM derived from land use is composed of fossil refractory matter from erosion of carbonaceous rocks (Galy et al., 2007; Gordon & Goñi, 2004) and black carbon mainly originating from landscape fires and volcano eruptions, also observed in the form of pyrogenic OM (Coppola et al., 2018; Jones et al., 2020; Jones & Gislason, 2008; Li et al., 2021). Vascular plants are a major component of ter-OM not just for being the largest biomass on Earth, but their biochemical composition makes a relevant and eventually an exclusive resource (Bianchi, 2011; Cragg et al., 2020). Originating from diverse sources ranging from plant litter decay to soil erosion, these materials are seasonal or year-around organic

sources across brackish and marine environments (Drenzek et al., 2009; Feng et al., 2013; Massicotte et al., 2017).

### Variation in ter-OM inputs

Geographical regions and habitats have marked variations in ter-OM stocks and dispersion mechanisms. The origin of ter-OM may vary, but natural landing areas are the aquatic compartments (Figure 1), where the three main processes of biodegradation, flocculation, and photodegradation are believed to drive the transformation of ter-OM (Bianchi et al., 2018; Massicotte et al., 2017) (Figure 1). In general, there is a decrease in the reactivity of the organic material (i.e., tendency to participate in chemical reactions, such as oxidation) along the aquatic continuum, indicating that the ter-OM reaching coastal areas is more refractory and potentially less bioavailable. Shallow estuarine regions have a longer retention time and organic material passes through a series of biogeochemical processes until reaching outer areas (Canuel et al., 2012; Middelburg & Herman, 2007). In the temperate northern hemisphere, boreal forests are known for the high contribution to the global terrestrial carbon pool (Bradshaw et al., 2009; Tagesson et al., 2020), with seasonal and continual decay of terrestrial material (mainly senescence leaves). Similarly, Arctic regions are marked by seasonal ter-OM fluxes during spring freshet and late-season melt, where ter-OM from rivers, glaciers, permafrost, and soil erosion flows into fjords and coastal areas (Bring & Destouni, 2011; Clark et al., 2022; McGovern, Pavlov, et al., 2020).

Rivers are responsible for a continuous input of ter-OM, and around 0.35 Pg of carbon are yearly transported by rivers to marine environments worldwide (Hedges et al., 1997). Global estimations predict that more than one tenth of the total carbon exported by major rivers is composed of black carbon, mainly from wildfires and fossil-fuel-derived compounds in its different forms (Jones et al., 2020), with 3%-10% reaching marine sediments (Coppola et al., 2018). However, the estimation of exports could be heavily influenced by intermittent rivers with temporal pulses containing variable ter-OM loads (Datry et al., 2014; del Campo et al., 2021). Regardless of the transportation pathway, the association of ter-OM with minerals supports the dispersion of OM to water compartments (Gordon & Goñi, 2004; Hedges & Keil, 1995; Simoneit, 2006). Muddy finer grained sediments, when compared with coarse grains, usually have a stronger affinity with ter-OM (Hedges & Keil, 1995), thus making it challenging to disassociate OM and sediments, and predict potential impacts to the faunal communities (Lohrer et al., 2006).



**FIGURE 1** Conceptual representation of the land-ocean environmental interface and potential sources of terrestrial organic matter (ter-OM). The material from land has both natural and anthropogenic, or mixed origins. Following the aquatic continuum, a substantial amount of ter-OM reaches the coastal and shelf benthic compartments. When sedimented, the material can follow three paths: (1) remineralized into dissolved inorganic carbon and nutrients, and remixed with the water, (2) stored in the benthic biomass, or (3) buried in the deeper sediments. Thus, ter-OM influences macrobenthic communities in several forms by being a source of nutrients, refuge, physical stressor smoothing epibenthic groups, and also as an important component of benthic-pelagic dynamics, acting as an energy transfer among trophic levels.

Recent climate predictions and ter-OM export estimations have brought a different dimension to the discussion of terrestrial contributions in coastal OM cycles (Cragg et al., 2020; Kandasamy & Nagender Nath, 2016; Qiao et al., 2019; Smeaton et al., 2021). A recent estimation in UK coastal waters showed that benthic subtidal sediments are by far the most important carbon storage area with around 23.9 million tonnes of carbon stored (~0.1–0.5 million tonnes of carbon per annum) (Parker et al., 2021). Fjord systems, for instance, are expected to bury 10 Mt of organic carbon per year (=11% of the global annual budget in marine systems) (Smith et al., 2015), with temperate fjords alone responsible for nearly 12% of global budgets in the last 100,000 years (Nuwer & Keil, 2005). Atmospheric transportation also drives ter-OM deposition in coastal and remote areas, for example, Galletti et al. (2020) have highlighted the importance of delivery of ter-OM associated with Saharan dust to the coast of remote Mediterranean Sea areas via sea spray, where the ter-OM transported by atmospheric inputs is locally higher than deliverables from riverine inputs.

Natural sources of ter-OM can be masked in some regions by anthropogenic-derived inputs. Land-based aquaculture, agriculture, industry, and other activities physically modify the landscape with potential long-lasting impacts on sediment dynamics, bringing up or adding substantial amounts of OM to the environment (Asmala et al., 2013; Bao et al., 2013; Kim et al., 2011; Louchouarn et al., 1999). Deforestation, erosion, and changes in land use can lead to substantial increases in mobilization and downstream transport of sediment OM, often with high molecular weight and aromaticity, with implications for the bioavailability of ter-OM reaching aquatic compartments (Boyer & Groffman, 1996; Coppola et al., 2018; Farella et al., 2001). When evaluating boreal estuaries with different ter-OM sources (forest/ peatland, agricultural, or mixed/urban), Asmala et al. (2013) showed via incubation experiments that ter-OM quality was driving the organic degradation, with higher mineralization rates and bacterial growth efficiency in coastal areas receiving OM inputs from agricultural landscapes.

## Processes in the sediment compartment

Once deposited in the benthic compartment, there are three main "fates" for ter-OM: (1) remineralization into dissolved inorganic carbon and nutrients, and remixed with the water (and therefore returned to the hydrosphere, or even the atmosphere), (2) uptake and short- or medium-term storage in the benthic biomass, or (3) long-term burial in the deeper sediments (Klages et al., 2004) (Figure 1). The different sources of ter-OM and transformation processes to particulate and dissolved fractions make it complex to trace the inputs, mainly when it enters the sediment matrix and food webs (Geraldi et al., 2019). In tidal estuaries, the constant particle mixing of ter-OM and enhanced solid-liquid exchange chemically changes the particles due to ter-OM resuspension into upper and more oxygenated water layers (Komada & Reimers, 2001; Middelburg & Herman, 2007).

The quantity and quality of ter-OM reaching the sediment surface depend on the habitat's geomorphology, physical-chemical characteristics, biological processes on the way down, and the proximity from the land source to the aquatic compartment (source-to-sink) (Bianchi, 2011; Blair & Aller, 2012; Hedges et al., 1997; Hoeinghaus et al., 2011). Nearly half of the total annual budget of ter-OM is indeed degraded before reaching the coastal sediment compartments (Louchouarn et al., 1999), but still a considerable amount is estimated to reach the sediment surface and be consumed or sedimented (Hedges & Keil, 1995; Painter et al., 2018; Sarmiento & Gruber, 2013). The ter-OM burial estimation provided by Burdige (2005) highlighted the inefficient remineralization of ter-OM compared with marine-derived OM. Also, a less explored and associated process in aquatic systems is the priming effect (Aller & Cochran, 2019; Bianchi, 2011), where the interaction between the allochthonous and resident organic material changes the remineralization rates. Experimental assessments exploring the priming effect are scarce (e.g., Turnewitsch et al., 2007; van Nugteren et al., 2009), and required to provide a better understanding of the process, but multiscale priming involving ter-OM seems to vary depending on the geomorphology and physical-chemical characteristics (Bianchi, 2011).

Terrestrial-derived material can also exert influence far from the coast: the massive delivery of ter-OM into the Arctic Ocean Basins from Arctic rivers (~10% of global riverine discharge) is reflected in the prevalence of ter-OM far offshore (Forest et al., 2007; Hwang et al., 2015), with recent models estimating ter-OM POC sources hundreds of kilometers from the shore (e.g., Yukon River delta in Alaska, USA, Clark et al., 2022). Similar observations have been made in large river basins, such as the Amazon shelf, where a seasonal increase in riverine loads drives benthic community changes (Aller & Stupakoff, 1996) with tracers indicating ter-OM up to 700 km from the shelf break (Sun et al., 2017).

# Ter-OM as a source of nutrients and food web link

An ongoing discussion topic is the potential role of ter-OM as a source of energy to coastal and marine food webs. This resource has traditionally been regarded as a low-quality food source for the marine trophic food chains (Cummings et al., 2009; Schell, 1983). Ter-OM has high carbon and nitrogen (C:N) ratio relative to, for example, phytoplankton due to the major presence of macromolecules with low nitrogen (e.g., cellulose, lignin) compared with protein sources. Thus, organisms may require more energy and time to process and assimilate this material, and a longer window to reach the required nutrient levels (Christofoletti et al., 2013; Cragg et al., 2015). Although ter-OM is low in the high-quality polyunsaturated fatty acids characteristic of phytoplankton, it can still have a high content of certain essentials (e.g.,  $\omega$ -3 and  $\omega$ -6 FA families). The material can be separated into nutritional components that are directly (e.g., proteins, sugars) or indirectly (e.g., cellulose, lignin) available to macrofauna (Galloway et al., 2012). Most of the plant-derived compounds, which are a major part of ter-OM, have tissues composed of lignocellulose, a compound that is difficult to break into smaller molecules (i. e., depolymerization) (Cragg et al., 2020). When compared with the long phylogenetic history of major organismal groups, the recent evolution of land plants may explain the sparse appearance of the ability in degrading lignocellulose in the tree of life (Cragg et al., 2015). While several animal groups have cellulose-degrading enzymes, only a few carry modifying molecules able to break lignin. On the other hand, microbial and fungal activities

can play a key role in lignin degradation, depolymerizing the organic material, either as free-living organisms, or living symbiotically with benthic invertebrate taxa (Cragg et al., 2015; Distel et al., 2002). There is a high reliance on bioavailable ter-OM from bacteria depolymerization, and some macrobenthic communities are maintained by the recycled microbial by-products and biomass itself (Harris et al., 2018; McTigue & Dunton, 2014). The macrobenthic wood borers, such as teredinid bivalves, have well-known symbiotic associations with proteobacteria across several genera (Distel et al., 2002). The presence of endogenous cellulase, however, is present in other invertebrate groups (Tanimura et al., 2013 and literature within) but still poorly explored in marine species, with the Limnoriidae isopod crustacean Limnoria quadripunctata being the first group identified with the ability to degrade lignin-derived material unassisted by symbionts (King et al., 2010).

Several estuarine and coastal deposit and suspension feeders utilize ter-OM, in its varied forms, as a continuous or seasonally important food resource (see Table 1). Ter-OM has indeed lower nutritional value when compared with marine-derived organic sources, but the high continuous input of the source in tropical habitats, and pulses during spring and autumn floods in temperate and Arctic regions, turns it into a relevant energy resource, with varied significance (minor to important contribution), to macrobenthic communities' diet (Table 1). As an example, ter-OM was estimated via isotopic mixing models to be responsible for up to 74% of the diet of suspension (bivalve Arthritica helmsi and polychaete Ficopomatus enigmaticus) and deposit (bivalve Soletellina alba) feeders (Lautenschlager et al., 2014), and also the only resource available to some coastal amphipod Gammarus insensibilis populations (Mancinelli & Rossi, 2002).

Ter-OM as a major energy source can be dependent on large-scale events, and higher riverine discharge and ter-OM load may overcome the usual diet contribution of generalist consumers (e.g., oyster Crassostrea gigas, Riera & Richard, 1997; blue crab Callinectes sapidus, Olin et al., 2013). High ter-OM pulses can also potentially reshape the community composition (e.g., positive correlation between macrobenthic abundance and runoff events, Hagberg & Tunberg, 2000) or even trigger life cycle events (e.g., reproduction of lagoon snails due to an increase of ter-OM in rainy seasons, Badano et al., 2016). Notably, high sedimentation accompanies high ter-OM loads and may cause long-lasting burial with potential clogging of filter feeders (Topçu et al., 2019). At the same time, seasonal short-scale events significantly contribute to some species' diets, such as the role of pine pollen for benthic invertebrates

in the Baltic Sea coastal food webs during early summer (Liénart et al., 2022).

Although generalist species can utilize ter-OM resources, the higher intake of less nutrient-rich resources may bring acute to long-term impacts to some groups, particularly to nonselective and filter feeder organisms. For example, the input of lignin alone reduced the food quality and affected the growth of the brittle star Amphiura filiformis, lowering the lipid content and arm regeneration rate in an experimental setup (Gunnarsson et al., 1999). Nevertheless, experimental manipulative studies exploring the effects of ter-OM at the species level are still scarce. At the same time, some taxa rapidly adapt to organic availability changing their feeding strategy, such as A. filiformis, switching from deposit to suspension-feeding (Renz et al., 2018), and the Asian clam Corbicula fluminea, with clear shifts in their food quality intake (marine- or terrestrial-derived) along the estuarine gradient (Dias et al., 2016). Other species even hibernate and enter in aestivation state during unfavorable environmental conditions, a behavior observed across benthic taxa and habitats that potentially is linked to low food availability (Coma et al., 2000). Some taxa simply move to more suitable environments, such as capitellid and nereidid polychaete families (Fernández-Rodríguez et al., 2019 and literature within), a limited behavior for sessile and others living in restricted or fragmented habitats, such as mangroves and tidal salt marshes. Otherwise, several filter feeder species have the ability to sort particles before ingestion supported by morphological and behavior adaptations (Ward & Shumway, 2004 and literature within). Although also scarce in the literature, some species from food-rich areas (e.g., estuarine habitats) are adapted to select ter-OM sources at very fine levels with a better digestion efficiency to specific resources (e.g., fatty acids, chlorophyll-rich sources) (Boon & Duineveld, 2012; Godbold et al., 2009).

At the community level, the reliance on ter-OM is observed across functional macrofauna groups and habitats (Table 1), which also may change along the aquatic continuum and distance from the river outlet (Antonio et al., 2012). McGovern, Poste, et al. (2020) found a higher reliance on ter-OM for subsurface deposit feeders (which feed directly on sediments) than for more selective feeders (such as scavengers/carnivores) in an Arctic fjord, a finding mirroring that of Harris et al. (2018) in the Beaufort Sea, Alaska. However, most studies have focused on general riverine inputs, thus not distinguishing which specific source of ter-OM that had been utilized. Estuarine habitats, such as mangroves, receive a larger input of nondegraded ter-OM directly to the benthic compartment when compared with other

**TABLE 1** Summary of the systematic analysis and additional articles exploring the interplay between macrofauna communities and terrestrial organic material (ter-OM).

Reference	Country	Subject	Habitat	Type of ter-OM source	TOM-related variables	Effect/ contribution
Netto and Lana (1999)	Brazil	Macrofauna community structure	Bay, estuary	Spartina alterniflora and detritus (plant material)	OM (LOI) and grain size	Higher density and diversity in salt marshes
Bell et al. (2016)	Alaska, USA	Food source assimilation	Coast	NS	$\delta^{13}$ C, $\delta^{15}$ N, C:N ratio, TOC, and TN	Important food resource
Bongiorni et al. (2016)	Italy	Food source assimilation	Coast	NS	SOM, $\delta^{13}$ C, $\delta^{15}$ N, and POM	Important food source
Bonifácio et al. (2014)	France	Macrofauna community structure	Coast	NS	TOC, TN, chloropigments, amino acids, THAA, EHAA, and grain size	Change in species composition
Forrest et al. (2007)	New Zealand	Macrofauna community structure	Coast	NS	δ <sup>13</sup> C and δ <sup>15</sup> N, C/N, and lipids	Higher density of opportunistic species (capitellidae Heteromastus filiformis)
Hermand et al. (2008)	France	Macrofauna community structure	Coast	NS	TOC, TC, pheopigments, and grain size	Higher density and biomass and lower diversity at river outlet (e.g., polychaeta)
Dunton et al. (2006)	Alaska, USA	Food source assimilation	Coast	NS	$\delta^{13}C$ and $\delta^{15}N$	Important food resource
Lee (1999)	Hong Kong	Macrofauna community structure	Coast	Kandelia candel	OM (LOI), total detritus unit, and tannin	Decreased species richness, diversity, evenness, and abundance with TOM enrichment (*)
Sampaio et al. (2010)	Portugal	Food source assimilation	Coast	NS	$\delta^{13}C$ and $\delta^{15}N$	Important food resource
Sarà et al. (2007)	Iceland	Food source assimilation	Coast	NS	$\delta^{13}C$ and $\delta^{15}N,$ POM, SOM	Food resource
Szczepanek et al. (2021)	Poland	Food source assimilation	Coast	NS	TOC, TN, chl <i>a</i> and pheo concentration, POM, $\delta^{13}$ C, and $\delta^{15}$ N	Important food resource
Darnaude et al. (2004)	France	Food source assimilation	Delta	NS	$\delta^{13}C$ and $\delta^{15}N$	Important food resource (deposit-feeding polychaetes)
Shilla and Routh (2017)	Tanzania	Food source assimilation	Delta	NS	$\delta^{13}C$ and $\delta^{15}N,$ FA	Important food resource (filter feeders and deposit-feeding consumers)

(Continues)

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Reference	Country	Subject	Habitat	Type of ter-OM source	TOM-related variables	Effect/ contribution
Bongiorni et al. (2018)	Italy	Macrofauna community structure and functional traits	Delta lagoons	NS	δ <sup>13</sup> C and δ <sup>15</sup> N, chloroplastic pigments, SOM, suspended POM, and grain size	Important food resource, higher biomass to suspension and surface-deposit feeders at river outlet
Nasi, Auriemma, et al. (2020)	Italy	Macrofauna community structure and functional traits	Delta lagoons	NS	TOC, TN, biopolymeric carbon, pigments, and grain size	Higher functional diversity and redundancy at river outlet
Abrantes et al. (2013)	Kenya	Food source assimilation	Estuary	NS	$\delta^{13}C$ and $\delta^{15}N$	Important food resource
Cummings et al. (2003)	New Zealand	Food source assimilation	Estuary	NS	$\delta^{13}$ C and $\delta^{15}$ N, OM (LOI), POC, TN, TP, chl <i>a</i> , and grain size	Minor food resource
Currin et al. (1995)	USA	Food source assimilation	Estuary	Spartina alterniflora	$\delta^{13}C$ and $\delta^{15}N$	Food resource (e.g., fiddler crabs <i>Minuca pugnax</i> and <i>Leptuca</i> <i>pugilator</i> )
Hughes et al. (2000)	USA	Food source assimilation	Estuary	<sup>15</sup> N-enriched nitrate	$\delta^{13}C$ and $\delta^{15}N,$ and $N$	Important food resource/ assimilation by crustaceans (e.g., sand shrimp <i>Crangon</i> <i>septemspinosa</i> )
Lillebø et al. (1999)	Portugal	Soil/nutrients structure	Estuary	Spartina maritima	OM (LOI), TN, and TP	Macrofauna enhance TOM degradation
Lillebø et al. (2007)	Portugal	Soil/nutrients structure	Estuary	Scirpus maritimus	OM (LOI), TN, and TP	Macrofauna enhance TOM nutrients decomposition and mineralization (*)
Netto and Gallucci (2003)	Brazil	Macrofauna community structure	Estuary	Detritus (plant material)	Detritus (plant material) biomass, TOM, and grain size	Positive correlation between detritus biomass and species no./ abundance (important refuge [?])
Norkko et al. (2002)	New Zealand	Macrofauna community structure	Estuary	Terrigenous clay	OM (LOI) and grain size	High content induces hypoxia and anoxia (?)
Richoux and Froneman (2007)	South Africa	Food source assimilation	Estuary	NS	δ <sup>13</sup> C and δ <sup>15</sup> N and POM	Important food resource, bivalve Solen cylindraceus
Feng et al. (2018)	China	Food source assimilation	Estuary (mangrove, mudflat, bay)	Spartina alterniflora, Kandelia obovata	$\delta^{13}C$ and $\delta^{15}N,$ SOM, and POM	Important food resource

(Continues)

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Reference	Country	Subject	Habitat	Type of ter-OM source	TOM-related variables	Effect/ contribution
Antonio et al. (2012)	Japan	Food source assimilation	Estuary, coast, shelf	NS	$\delta^{13}C$ and $\delta^{15}N$	Important food resource (e.g., bivalve <i>Corbicula</i> <i>japonica</i> )
Gladstone-Gallagher et al. (2020)	New Zealand	Macrofauna community structure and functioning	Estuary and intertidal sandflat	Nitrogen	TOC and grain size	Higher abundance of surface gastropods and deposit-feeding bivalves, and large no. burrows (*)
Cari et al. (2020)	Chile	Food source assimilation	Fjord	NS	Suspended OM, SPOM, POM, $\delta^{13}$ C, and $\delta^{15}$ N	Higher trophic redundancy or overlapping trophic niches
Hagberg and Tunberg (2000)	Sweden	Macrofauna community structure	Fjord	NS		Increased abundance due to runoffs and TOM deposition (?)
Kędra et al. (2012)	Norway	Food source assimilation	Fjord	NS	POM, $\delta^{13}$ C, and $\delta^{15}$ N	Important food resource
Kokarev et al. (2021)	Norway	Macrofauna community structure	Fjord	NS	$\delta^{13}$ C, $\delta^{15}$ N, TOC, TN, and grain size	No effect on community structure (?)
McGovern, Poste, et al. (2020)	Norway	Macrofauna community structure and functional traits	Fjord	NS	TOC, TN, $\delta^{13}$ C, and $\delta^{15}$ N, and grain size	Lower taxonomic and functional diversity, higher community biomass at river outlet
McLeod and Wing (2009)	New Zealand	Food source assimilation	Fjord	NS	δ <sup>13</sup> C, δ <sup>15</sup> N and δ <sup>34</sup> S, FA, TOC, and TN	Important food resource (e.g., sea urchin Echinocardium cordatum and polychate Pectinaria australis)
Quiroga et al. (2016)	Chile	Macrofauna community structure	Fjord	NS	$\delta^{13}$ C and $\delta^{15}$ N, TOM, TOC, C/N ratio, chl <i>a</i> and Phaeo, and grain size	Higher diversity at inner fjord
Sokołowski et al. (2014)	Svalbard, Norway	Food source assimilation	Fjord	NS	$\delta^{13}C$ and $\delta^{15}N$	Food resource
Zapata-Hernández et al. (2014)	Chile	Food source assimilation	Fjord	NS	POM, TOM, $\delta^{13}$ C, and $\delta^{15}$ N	Lower contribution to benthic communities
Zapata-Hernández et al. (2016)	Chile	Macrofauna community structure	Fjord	NS	δ <sup>13</sup> C and δ <sup>15</sup> N, SOM, and POM	Important food resource (wood-boring Xylophagid)
Fanelli et al. (2009)	Italy	Food source assimilation	Gulf	NS	$\delta^{13}C$ and $\delta^{15}N$	Food resource (e.g., amphipods, cumaceans, and decapods)
						(Continues)

Reference	Country	Subject	Habitat	Type of ter-OM source	TOM-related variables	Effect/ contribution
Harmelin-Vivien et al. (2009)	France	Food source assimilation	Gulf	NS	Cisotope, chl <i>b</i> , carbohydrate, lipid, and protein contents	Decreased diversity and increased abundance of specific functional groups (e.g., deposit-feeding polychaetes <i>Laonice cirrata</i> , <i>Mediomastus</i> sp., <i>Cossura</i> sp., <i>Sternaspis</i> <i>scutata</i> , and <i>Polycirrus</i> sp.)
Misic et al. (2016)	Italy	Food source assimilation	Gulf	NS	δ <sup>13</sup> C and δ <sup>15</sup> N, TOC, TN, and carbohydrate	Important food resource (deposit feeders)
Salen-Picard et al. (2002)	France	Macrofauna community structure	Gulf	NS		Higher density and biomass (e.g., polychaeta Mediomastus sp. and Aricidea claudiae)
Dunton et al. (2012)	Alaska, USA	Food source assimilation	Lagoon	NS	$\delta^{13}C$ and $\delta^{15}N,$ POM	Important food resource
Frouin (2000)	French Polynesia	Macrofauna community structure	Lagoon	NS	Grain size, pheop pigment, and organic load	Important food resource
Harris et al. (2018)	Alaska, USA	Food source assimilation	Lagoon	NS	$\delta^{13}C,\delta^{15}N,$ and molar C:N ratios	Important food resource
Jędruch et al. (2019)	Poland	Metal assimilation (Hg)	Lagoon	NS	OM (LOI) and grain size	Increased Hg accumulation on filter feeders
Kanaya et al. (2011)	Japan	Macrofauna community structure	Lagoon	NS	Oxidation-reduction potential, acid-volatile sulfide, TOC, TN, and $\delta^{13}$ C	Lower density of opportunistic species (?)
Marcelina et al. (2018)	Poland	Food source assimilation	Lagoon	NS	$\delta^{13}C$ and $\delta^{15}N$	Food resource
Nasi, Ferrante, et al. (2020)	Italy	Macrofauna bioturbation	Lagoon	NS	TOC, TN, $\delta^{13}$ C and $\delta^{15}$ N, radionuclides $^{137}$ Cs and $^7$ Be, and grain size	Modify the spatial pattern of community traits
Andrade et al. (2014)	Brazil	Macrofauna community structure	Mangrove	Rhizophora mangle		Important refuge
Bouillon et al. (2002)	India	Food source assimilation	Mangrove	NS	$\delta^{13}C$ and $\delta^{15}N,$ and SOM	Minor food resource
Chen et al. (2018)	China	Food source assimilation	Mangrove	Spartina alterniflora	$\delta^{13}C$ and $\delta^{15}N$	Important food resource
De Oliveira et al. (2012)	Brazil	Macrofauna community structure	Mangrove	Rhizophora mangle, Laguncularia racemosa		Food resource and refuge

(Continues)

Reference	Country	Subject	Habitat	Type of ter-OM source	TOM-related variables	Effect/ contribution
Demopoulos et al. (2007)	USA, Hawai	Food source assimilation	Mangrove	Rhizophora mangle	$\delta^{13}C$ and $\delta^{15}N$	Food resource (e.g., polychaetes)
Gladstone-Gallagher et al. (2014)	New Zealand	Macrofauna community structure	Mangrove	Avicennia marina	TOM, chl <i>a</i> , and phaeophytin (phaeo)	Decrease abundance of few taxa (e.g., polychaeta <i>Prionospio</i> <i>aucklandica</i> ), but no effect on composition and density (*)
Churchwell et al. (2016)	Alaska, USA	Macrofauna community structure and food source assimilation	River delta	NS	δ <sup>13</sup> C and δ <sup>15</sup> N, and POM	Important food resource
Bergamino and Richoux (2015)	South Africa	Food source assimilation	Salt marsh	Salt marsh Chenolea diffusa and Sarcocornia perennis, Marsh grass Spartina maritima	δ <sup>13</sup> C and δ <sup>15</sup> N, and FA	Important food resource
Neira et al. (2007)	USA	Macrofauna community structure	Salt marsh	Spartina	Grain size and TOM	Reduced species richness, increase dominance and shift in feeding modes (surface microalgal feeders to subsurface detritus/Spartina feeders, oligochaeta, and polychaeta)
Rezek et al. (2017)	USA	Food source assimilation	Salt marsh	Spartina alterniflora	$\delta^{13}C$ and $\delta^{15}N$	Important food resource
Tang and Kristensen (2010)	Denmark	Macrofauna community structure	Salt marsh	Spartina anglica	OM (LOI)	Low infaunal abundance and diversity; higher abundance of epibenthic grazers and high abundance of <i>Corophium</i> <i>volutator</i> at the mudXat–marsh boundary
Mancinelli and Rossi (2002)	Italy	Macrofauna community structure	Shelf	NS	TOC, OM (LOI), and grain size	Support-adapted species (e.g., Gammarus insensibilis)
McTigue and Dunton (2014)	Alaska, USA	Food source assimilation	Shelf	NS	SPOM, $\delta^{13}$ C, $\delta^{15}$ N, and C:N ratios	Important food source, link with microbial alteration (Continues)

Reference	Country	Subject	Habitat	Type of ter-OM source	TOM-related variables	Effect/ contribution
Stasko et al. (2018)	Canada	Food source assimilation, Benthic–pelagic coupling	Shelf	NS	δ <sup>13</sup> C, δ <sup>15</sup> N, grain size, chl <i>a</i> and phaeopigment concentrations, LOI, and C:N	Minor and important food source, close link between TOM source and benthic-pelagic trophic coupling strength
Galeron et al. (2009)	Congo	Macrofauna community structure	Shelf (canyon)	NS	TOC, TN, and sulfur	Seasonal density and vertical profile variance
Liao et al. (2018)	China	Macrofauna community structure	Tidal mudflat	Spartina alterniflora, Kandelia candel		Reduced abundance and species no. in <i>Kandelia</i> <i>candel</i> , community ID shifts between <i>Spartina</i> <i>alterniflora</i> and tidal flat habitats
Netto and Lana (1997)	Brazil	Macrofauna community structure	Tidal mudflat	Spartina alterniflora	Grain size and TOM	Correlation between higher density of detritivorous or omnivorous species and higher detritus biomass

*Note*: The content of the articles was organized by country in which the studies were conducted, subject (main content of the article), habitat, type of ter-OM source, ter-OM-related variable, and effect/contribution originated from the ter-OM inputs. Studies with nonspecified type of ter-OM quality (e.g., total riverine input) were specified with NS. Relevance of food resource was split into three levels of magnitude: minor (low relevance to macrofauna diet), ordinary (i.e., food resource), and important food resource (seasonal or permanent relevant resource to macrofauna diet and/or trophic chain). When only specific group/ species are affected by ter-OM, functional groups or species are specified in the effect/contribution column. Effects with vague, uncertain, or hypothesized descriptions were indicated with (?). Manipulative experiments were indicated with (\*) after the effect/contribution description. Table is sorted by habitat type, and the complete list of references is provided in Appendix S1.

Abbreviations: EHAA, enzymatically hydrolysable amino acids; FA, fatty acids; LOI, loss in ignition; POC, particulate organic carbon; SOM, sediment organic matter; TC, total carbon; THAA, total hydrolysable amino acids; TN, total nitrogen; TOC, total organic carbon; TOM, total organic matter; TP, total phosphorus.

coastal and marine environments. In areas with a constant input of fresh and nondegraded ter-OM, macrofauna diet plays a central ecological role in habitat dynamics. As an example, the presence of endogenous cellulase was observed in several leaf eater crabs (e.g., Parasesarma ervthodactyla, Bui & Lee, 2015; Parasesarma bidens, Kawaida et al., 2019) and, by consuming and enhancing the degradation of mangrove litter, they hold an important link in the food web, making nutrients available to both lower and higher trophic levels (Cannicci et al., 2008; Werry & Lee, 2005).

While much research has, quite naturally, focused on macrofaunal utilization of ter-OM in its particulate form, terrestrial DOM can also impact energy sources to macrofaunal communities indirectly by stimulating basal productivity and enhancing vertical carbon export, through plankton and microbial communities. This influence is tightly dependent on light availability in shallow waters and stimulated by riverine inputs of nutrients. Thus, by obtaining energy from ter-OM, macrofauna turns into a crucial link between terrestrial primary production and marine secondary production, through the processes of benthic–pelagic coupling (Antonio et al., 2010; Attrill et al., 2009). Some filter-feeding organisms (e.g., ascidians) are also capable of direct uptake of dissolved and colloidal OM (Riisgård & Larsen, 2010).

Recent studies using multimarker approaches (i.e., approaches combining analysis of multiple geochemical markers) have challenged the paradigm that ter-OM has low importance in marine and coastal food webs (e.g., stable isotopes, lignin phenol). In an Arctic fjord, the large annelid *Scoloplos armiger* was the most abundant species close to the river mouth, and its low  $\delta^{13}$ C value

indicated a considerable terrestrial resource utilization (McGovern, Poste, et al., 2020), thus illustrating how terrestrial inputs may fuel benthic communities. In the Arctic Beaufort Sea, a community trophic characterization highlighted the role of ter-OM as an important subsidy to food webs (Harris et al., 2018) even into higher trophic levels, where microbes efficiently process ter-OM enhancing the availability of high-quality food sources to macrofauna (Bell et al., 2016).

# THE RELATIONSHIP BETWEEN ter-OM AND MACROFAUNA DYNAMICS

# **Diversity and functional traits**

Ter-OM brings either positive, neutral, or negative effects to the community (e.g., structuring and diversity, abundance, recruitment) with both POM and DOM leading

13 of 27 complex multilevel implications to the benthic community (see Figure 2 for a schematical representation of the interrelationship between the effects of ter-OM and macrofauna). Although caution is required while making generalizations about community-level impacts on benthic communities due to the lack of experiments using ter-OM, previous studies suggest that decreased species richness, diversity, evenness, and abundance may be expected under high terrestrial loads (Harmelin-Vivien et al., 2009; Lee, 1999). Similarly to the nonlinear macrofauna responses along the enrichment gradient (Pearson & Rosenberg, 1978), higher richness in the outer estuarine areas (i.e., closer to the marine environment) is expected to have natural ter-OM inputs as an important driver (Giménez et al., 2005; Laurino et al., 2021; McGovern, Poste, et al., 2020).

As components of dynamic environments, benthic estuarine and coastal species are evolutionarily adapted to fluctuating environmental conditions. When ter-OM enrichment is higher than the communities support, it



**FIGURE 2** General schematical interrelationship between the effects of terrestrial organic matter (ter-OM) and macrofauna. Particulate and dissolved organic matter (POM and DOM, respectively) from land, including the dissolved colored form (CDOM), directly affects macrofauna (individuals and community) bringing a cascade of impacts at different levels. In the representation, blue color boxes indicate changes in the biogeochemistry and yellow indicates the effects on the benthic communities structuring and functioning and species fitness. Arrows connecting boxes have uni- or multidirectional impacts. can suppress species behavior and metabolic activity via O<sub>2</sub> depletion, with negative outcomes for both biodiversity and community functions (Drylie et al., 2020). Usually under high ter-OM loadings, opportunistic species tend to dominate the functional groups (Drylie et al., 2020; Fernández-Rodríguez et al., 2019) and higher biomass of suspension, filter- and surface-deposit feeders may be also expected (e.g., Bongiorni et al., 2018; Feng et al., 2018). As an example, Gladstone-Gallagher et al. (2020) showed that similar ter-OM enrichment in a mangrove system affects groups from the same community in different manners: surface gastropods increased and deposit-feeding bivalves decreased in abundance. Community changes, however, vary in intensity and time lags starting several months to even years (e.g., 1–2 years) after the ter-OM input events (e.g., Salen-Picard et al., 2003), depending on the event magnitude and source, as well as the dominant taxa in the impacted communities. Also, ter-OM inputs are often coupled to sediment inputs from land, which can drive community shifts to high abundance of more adapted taxa to high sedimentation rates.

# Links between macrofauna communities, biogeochemical processes, and carbon cycle

Benthic macrofauna is a central player in moderating sediment burial processes and biogeochemical processes within the sediment matrix (Welsh, 2003). Benthic fauna, in association with physical processes, is known to accelerate remineralization by increasing the reoxidation rates and mixing fresh and refractory OM repeatedly into the same reaction zones (Sarmiento & Gruber, 2013; Volkenborn et al., 2012), thereby enhancing ter-OM remineralization. This process may also be facilitated by other faunal groups. For example, macrofauna combined with meiofauna promoted nutrient turnover and supported an increase of up to 90% in the mineralization rate in a Scirpus maritimus salt marsh (Lillebø et al., 2007). While microbial communities are the major drivers of organic remineralization, it is the macrofaunal activity that promotes the sediment mixing through bioturbation, thereby bringing up sediments (including ter-OM) from anoxic to oxic layers and pumping oxygenated water into deeper layers (i.e., biorrigation), which can enhance microbial degradation and heterotrophic metabolism (Wakeham & Canuel, 2006).

Changes in macrofauna community composition limit sediment irrigation and oxygen availability by reducing their remineralization capacity to near-total inefficiency under high anoxic conditions (Levin et al., 2009). The macrofauna also supports the microbial community structure and habitat nutrient cycling by controlling the distribution of key elements (e.g., nitrogen, iron, sulfur) (Kristensen et al., 2012; Sarker et al., 2021; Solan et al., 2019; van de Velde et al., 2020). Although the influence of faunal bioturbation/biorrigation has been better explored in recent decades, the effects of organisms on particle mixing, long-term degradation of ter-OM, and associated processes are still not fully predicted and require a good understanding of the community to provide habitat functional inferences (Kristensen et al., 2014; Sarmiento & Gruber, 2013).

Macrofauna-sediment interactions control and preserve carbon in different ways, and this theme has been extensively reviewed (Bianchi et al., 2021; Kristensen et al., 2008; Snelgrove et al., 2018; Wakeham & Canuel, 2006), but the interplay between habitat dynamics and macrobenthos is sometimes overlooked and neglected in carbon budget assessments (Andreetta et al., 2014; Bianchi et al., 2021; Kristensen et al., 2022; Lillebø et al., 2007). Macrofauna biomass itself is a source and storage of carbon of terrestrial origin, and the ter-OM consumption and sediment mixing behavior make carbon bioavailable to the environment in its different forms (Doering et al., 1986; McLeod & Wing, 2009). There is an increasing interest in the role of particular species able to digest lignocellulose (King et al., 2010) and in macrofauna bioturbation (Bianchi et al., 2021), due to their intrinsic relationship with nutrient regeneration and more specifically carbon cycling.

# CURRENT STATUS OF THE RESEARCH TOPIC: ter-OM AND MACROFAUNA COMMUNITIES

To complement our knowledge synthesis on the interplay between ter-OM and macrofauna communities, we have performed a research weaving systematic review (Nakagawa et al., 2019), using bibliometric and mapping across research networks (see Appendix S1 for our methodological strategy). Studies have been conducted worldwide, however with the majority of knowledge production in European and North American countries (Figure 3). As highlighted by Pearson and Rosenberg (1978) decades ago, most of the studies and collaboration among researchers are still conducted in the northern hemisphere, although some countries from the southern hemisphere are represented in the top 10 countries in number of publications (Figure 4A). Publications included date back to the mid-1990s (Figure 4B) and are primarily published in specialized journals focusing on coastal and marine ecology (Appendix S1: Table S2). These articles explore a wide range of OM-related themes



**FIGURE 3** Global overview of articles exploring the interplay between terrestrial organic material and macrofauna communities. Geographical distribution of publications and number of publications per continent (study sites). The color gradient represents the number of publications based on the corresponding authors' affiliations and lines the collaborations across countries obtained from authors' affiliations.

(e.g., OM-cycling, food web uptake) with a clear shift to carbon-related topics in recent years (Appendix S1: Figure S2, Table S3).

Articles, with few exceptions, cover two main themes: macrofauna community structure and food source assimilation (Table 1). The former explores how functional and diversity traits respond to ter-OM, and how communities (re)arrange to the potential stressors, with some exploring the functional traits of the communities. The latter explores how communities utilize ter-OM as a potential energy source. The use of functional analysis and trait-based approaches has been more recently incorporated into the ter-OM and benthic macrofauna studies (2016-present) (Table 2). Despite the small number of studies available limiting the potential to make broad exploring studies inferences, sedimentand feeding-related effect traits are relatively well-represented and show that suspension and sediment surface feeders tend to be more abundant and/or cope relatively well compared with other taxa where ter-OM inputs are high. Long, thin, and threadlike shape of macrofauna was indicated as a response trait to habitats receiving larger inputs of ter-OM inputs (inner fjord stations); however, other drivers (e.g., grain size) may also exert a significant influence in the group (McGovern, Poste, et al., 2020). Trait-based approaches complement species composition measurements to predict and determine ter-OM as a stressor. Feeding-mode traits support the understanding of OM pathways and how it is incorporated and passed across trophic levels (McTigue & Dunton, 2014).

Additional articles exploring changing sediment biogeochemistry, bioturbation potential, and metal assimilation as a consequence of benthic macrofauna and ter-OM interactions were also observed, although less in number.

Most of the studies were conducted in estuarine habitats (51 out of 65), followed by coastal (10) and shelf (4) environments, as well as one study that covered all the gradients (Antonio et al., 2012). Fjords, followed by lagoons and mangroves, are the most explored estuarine habitat. The majority of the studies are field-based evaluating the effect of total riverine input on the communities. However, just a few have explored the net effects of specific plant material (e.g., *Spartina alterniflora, Scirpus maritimus*) or organic compounds (e.g., nitrogen) on macrofauna, even less under manipulative laboratory or field experiments (Table 1).

# Emerging themes on the research topic

Bridging the traditionally separate fields of terrestrial and marine research is a challenging task (Munguia & Ojanguren, 2015), and investment in interdisciplinary studies targeting key knowledge gaps related to terrestrial-marine coupling is still required. Climate change and other anthropogenic-derived stressors impact ecosystems and connectivity between land and sea, making the need for a better understanding of these complex multifaceted changes. Despite the potential bias from the languages used in the literature search (i.e., excluding







**FIGURE 4** (A) Top 10 countries in the number of publications exploring the interplay between terrestrial organic material and macrofauna communities based on the country of the authors' affiliations. MCP, multiple country publication; SCP, single country publication. (B) Number of publications on the topic per year.

**TABLE 2** Effect and response functional traits used in studies exploring the interplay between terrestrial organic material (ter-OM) and macrofauna communities on estuarine, coastal, and shelf habitats. Specific functional trait levels used for each study can be traceable in each reference.

	Func	tional traits	ter-OM-related traits		
Study	Response	Effect	Response	Effect	
McGovern, Poste, et al. (2020)	Diversity, larval type, adult habitat, degree of attachment, adult mobility, and body form	Normal size, life duration, sediment depth, sediment reworking, fecal deposition, and feeding habit	Body form: long, thin, and threadlike shape <sup>a</sup>	Sediment dwelling depth: deep dwellers (5–15 cm); Sediment reworking: upward conveyor, biodiffusors; fecal deposition: subsurface (0–5 cm); feeding habit: subsurface deposit feeders <sup>a</sup>	
Nasi, Auriemma, et al. (2020)	Sediment position, adult mobility, and adult movement method	Maximum size, adult longevity, reproductive frequency, adult feeding habit, and bioturbation			
Cari et al. (2020)		Feeding habitat			
Nasi, Ferrante, et al. (2020)		Feeding strategy, mobility, sediment reworking, and burrow ventilation		Sediment reworking: semi-motile and conveyors	
Jędruch et al. (2019)		Feeding strategy		Suspension feeders	
Bongiorni et al. (2018)		Feeding strategy		Suspension and surface-deposit feeders	
Shilla and Routh (2017)		Feeding strategy			
Stasko et al. (2018)		Feeding habit and feeding strategy			

<sup>a</sup>Functional traits observed in the riverine stations.

other non-English language publications than Portuguese and Spanish), our bibliographic analysis highlighted fewer studies and lower levels of multicountry collaboration in the Global South, which is consistent with previously reported trends in ecological science (Massicotte et al., 2017; Nuñez et al., 2021; Pearson & Rosenberg, 1978). The lack of knowledge in certain regions or collaboration across disciplines and countries can lead to a bias toward generalizations based on studies with limited global coverage, particularly when dealing with climate-change-related topics where impacts on north temperate systems often dominate the scientific discourse (Nuñez et al., 2021).

Climate change alters the fluxes of ter-OM (and other terrestrial material) into aquatic systems, with an increase due to heavy precipitation and erosion (Panagos et al., 2017) mainly in Europe, Asia, and southeastern South America (IPCC, 2021). For example, several studies have pointed to an ongoing increase in runoff of freshwater and terrigenous material from land to the Arctic Ocean (Bring & Destouni, 2011; Saito et al., 2021), where the land-ocean connectivity is particularly strong (McClelland et al., 2012). The region has been warming at unprecedented rates and contains large natural organic carbon stocks, such as permafrost regions, which store the largest pool of organic carbon in the world along with a high nitrogen reservoir (Abbott, 2022; Turetsky et al., 2019). Since macrofauna is recognized as a strong indicator of environmental conditions, and as a potential tool to manage protected areas under climate change scenarios (Laurino et al., 2021), investments in poorly explored areas are crucial to support multiple-scale mitigation measures.

Ter-OM is a good indicator of environmental quality due to its influence on carbon budget and nutrient composition; however, it is usually degraded, consumed, or stored quickly, making it challenging to trace when entering aquatic systems. Even when considering OM derived only from vascular plant material, the polysaccharides, lignin, and other compounds differ in their degradation rates and bioavailability to organisms (Wakeham & Canuel, 2006). Tracing ter-OM utilization by benthic macrofauna can also be complex, where geochemical signatures of ter-OM (e.g.,  $\delta^{13}$ C, fatty acid composition) are often seasonally and spatially heterogeneous and can overlap with marine OM sources. Specific biomarkers have been used to trace ter-OM along the land-ocean gradient, performing better when integrated into multimarker approaches (Forrest et al., 2007; Rodil et al., 2020). As previously mentioned, stable isotopes or carbon and nitrogen have been broadly utilized, but their use as single tracers should be carefully applied, for example, in Arctic systems, where the high inorganic nitrogen may lead to an underestimation of ter-OM if not properly removed (Kumar et al., 2016). Thus, apart from habitat-specific approaches, the use of multi and complementary ter-OM tracers is needed for more reliable estimations, such as using radiocarbon (Feng et al., 2013; Guo et al., 2004) and lignin phenols (Rezende et al., 2010; Sobrinho et al., 2021), and a strong need to focus on the integrative use of land-derived tracers, especially in environmental monitoring programs.

Ter-OM influences all living coastal sediment taxa, including the more marine-restricted species (Birchenough et al., 2015). Looking to the past, paleoclimatic events with ter-OM overloading have shaped benthic community composition and functioning (Barash, 2012; Bianchi et al., 2021). Apart from the rapid temperature increase, the Paleocene-Eocene Thermal Maximum (~56 Mya) period experienced a larger input of sediments with high carbon content and  $\delta^{13}$ C-depleted carbon entering the coastal and marine systems; these changes resulted in high benthic fauna extinction rates due to changes in food sources and high productivity and low-oxygen conditions (McInerney & Wing, 2011). Thus, predicted environmental changes (e.g., an increase in flood events) may bring complex impacts to the macrofauna, highlighting the need for long-term, seasonal ecological studies and experimental manipulative approaches to clarify the role of ter-OM as a driver for the structure and function of macrofauna communities. Presently, tube-building infauna group support sediment stability and thereby represent an ecosystem engineering role for the surrounding and associated taxa (Bailey-Brock, 1984; Van Hoey et al., 2008). However, builder species tend to be more sensitive to organic enrichment than free-living species, such as the tube-builder polychaete Diopatra neapolitana, (Carregosa et al., 2014; Harkantra & Rodrigues, 2004). A potential negative effect on tube builders' density may modify the whole community structure including the environmental dynamics. Additionally, the increasing input of CDOM into aquatic systems resulting in the darkening of coastal and shelf waters (Aksnes et al., 2009; Deininger & Frigstad, 2019) have more obvious implications for light-dependent organisms living in the pelagic compartment, but changes in the food supply for the benthic compartment and cascading trophic level changes may also be expected (e.g., Frigstad et al., 2023).

Biogeochemical models have focused efforts on disentangling processes between benthic and pelagic compartments (Griffiths et al., 2017). Recent efforts support the observation and improvement of models to address the lack of baseline information and predictions aquatic processes (Kandasamy & Nagender on Nath, 2016; Rühl et al., 2020). Global models are limited (e.g., faunal biomass, Jones et al., 2014; Yool et al., 2017) and estimations of the direct impacts on faunal activity are inexistent (Bianchi et al., 2021). Recent advances in modeling approaches have included e.g. improved representation of the role of benthic substrate in driving biogeochemical cycling (e.g., European Regional Seas Ecosystem Model [ERSEM], Aldridge et al., 2017) and the inclusion of benthic bioturbation and bioirrigation (e. g., BAltic sea Long-Term large Scale Eutrophication Model [BALTSEM], Ehrnsten et al., 2022; Bottom RedOx Model [BROM v.1.1], Yakushev et al., 2017). However, all of the studies listed above point to the need for improved modeling tools for capturing the complexity of the interactions between benthic biota and biogeochemical cycling.

# **CONCLUSION AND PERSPECTIVES**

Our review highlights the current knowledge of the interplay between coastal, shelf and marine macrofauna communities and ter-OM. A large amount of ter-OM is brought to the coast from land, however there are still many knowledge gaps related to the transformation and fate of this material, especially in the context of a changing climate (Bianchi, 2011). Although marine systems are shaped by a complex set of abiotic and biotic drivers, understanding the impacts of ter-OM as an individual stressor can provide important insight into potential combined and cumulative impacts (Crain et al., 2008). In general, the direct or indirect effect of ter-OM depends on the benthic community structure and functional groups and the characteristics and amount of the OM itself.

Most studies explore the ultimate impacts of ter-OM stressors on macrofauna communities (e.g., species richness, biomass), but physiological and behavior impairments may bring ecological consequences yet are still not well explored. Changes in the source, quality and quantity of ter-OM inputs may lead to unimodal and more complex multimodal cascade responses. As the evidence of ter-OM contribution to coastal and marine food webs grows, novel relationships with ter-OM sources and macrofauna are observed. Both the interplay between terrestrial versus marine and anthropogenic versus natural organic sources, have different chemical compositions and residence time, and thus generalizing their impacts on benthic communities may lead to inaccurate conclusions.

The role played by macrofauna in carbon cycling and sequestration is not fully understood due to the lack of experimental information, but its importance in controlling these processes is unquestionable (Snelgrove et al., 2018; Solan et al., 2020). It seems also clear from recent studies and our compiled information that classifying ter-OM as a nonrelevant energy resource to estuarine, coastal, and shelf regions is no longer valid. As highlighted by Attrill et al. (2009), this consensus may origin from not considering estuarine studies and the closer evolutionary affinity of estuarine species with freshwater groups. The use of biogeochemical tracers for understanding OM sources and cycling, including food web utilization, has also contributed to a changing perspective on the topic.

When it comes to global ter-OM fluxes and characterization, the lack of in situ observations mainly in (sub) tropical areas and small riverine systems and the general lack of benthic-pelagic biogeochemical and integrative approaches contribute as missing pieces to this environmental puzzle. Monitoring macrofauna under a changing climate scenario is challenging with the traditionally used community indices, which are the tools mainly environmental supporting assessments worldwide. Community traits, for example, inform on evolutionary adaptations to stressors and species shifts to buffer areas. Such approach could be coupled with newly developed community indices safeguarding decision makers to protect the benthic ecosystem goods and services in coastal, shelf, and marine environments.

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# **CONFLICT OF INTEREST STATEMENT**

The authors declare no conflicts of interest.

#### DATA AVAILABILITY STATEMENT

Data (Pardo, 2023) are available from Zenodo: https:// doi.org/10.5281/zenodo.7702058.

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# REFERENCES

- Abbott, B. W. 2022. "Permafrost Climate Feedbacks." In Global Arctic: Permafrost Climate Feedbacks, edited by M. Finger and G. Rekvig, 189-209. Cham: Springer International Publishing.
- Abrantes, K. G., A. Barnett, T. R. Marwick, and S. Bouillon. 2013. "Importance of Terrestrial Subsidies for Estuarine Food Webs in Contrasting East African Catchments." Ecosphere 4: 1-33.
- Aksnes, D. L., N. Dupont, A. Staby, Ø. Fiksen, S. Kaartvedt, and J. Aure. 2009. "Coastal Water Darkening and Implications for Mesopelagic Regime Shifts in Norwegian Fjords." Marine Ecology Progress Series 387: 39-49.
- Aldridge, J. N., G. Lessin, L. O. Amoudry, N. Hicks, T. Hull, J. K. Klar, V. Kitidis, et al. 2017. "Comparing Benthic Biogeochemistry at a Sandy and a Muddy Site in the Celtic Sea Using a Model and Observations." Biogeochemistry 135: 155-82.
- Aller, R. C., and J. K. Cochran. 2019. "The Critical Role of Bioturbation for Particle Dynamics, Priming Potential, and Organic C Remineralization in Marine Sediments: Local and Basin Scales." Frontiers in Earth Science 7: 157.
- Aller, J. Y., and I. Stupakoff. 1996. "The Distribution and Seasonal Characteristics of Benthic Communities on the Amazon Shelf as Indicators of Physical Processes." Continental Shelf Research 16:717-51.
- Andrade, J. T. M., N. B. Palhano, C. H. Tagliaro, and C. R. Beasley. 2014. "Spatial and Temporal Variation in the Abundance and Taxonomic Composition of Estuarine and Terrestrial Macrofauna Associated with Mangrove Logs." Journal of the Marine Biological Association of the United Kingdom 94: 35-42.
- Andreetta, A., M. Fusi, I. Cameldi, F. Cimò, S. Carnicelli, and S. Cannicci. 2014. "Mangrove Carbon Sink. Do Burrowing Crabs Contribute to Sediment Carbon Storage? Evidence from a Kenyan Mangrove System." Journal of Sea Research 85: 524-33.
- Antonio, E. S., A. Kasai, M. Ueno, N. il Won, Y. Ishihi, H. Yokoyama, and Y. Yamashita. 2010. "Spatial Variation in Organic Matter Utilization by Benthic Communities from Yura River-Estuary to Offshore of Tango Sea, Japan." Estuarine, Coastal and Shelf Science 86: 107-17.
- Antonio, E. S., A. Kasai, M. Ueno, Y. Ishihi, H. Yokoyama, and Y. Yamashita. 2012. "Spatial-Temporal Feeding Dynamics of Benthic Communities in an Estuary-Marine Gradient." Estuarine, Coastal and Shelf Science 112: 86-97.
- Asmala, E., R. Autio, H. Kaartokallio, L. Pitkänen, C. A. Stedmon, and D. N. Thomas. 2013. "Bioavailability of Riverine Dissolved Organic Matter in Three Baltic Sea Estuaries and the Effect of Catchment Land Use." Biogeosciences 10: 6969-86.
- Attrill, M. J., S. D. Rundle, A. Fraser, and M. Power. 2009. "Oligochaetes as a Possible Entry Route for Terrigenous Organic Carbon into Estuarine Benthic Food Webs." Marine Ecology Progress Series 384: 147-57.
- Badano, E. I., F. A. Labra, C. G. Martínez-Pérez, and C. H. Vergara. 2016. "Changes in Body Size Spectra of Benthic Caridean Shrimps (Decapoda: Caridea) and Snails (Gastropoda) as Response to Seasonal Variability." Revista de Biologia Tropical 64: 33-44.
- Bailey-Brock, J. H. 1984. "Ecology of the Tube-Building Polychaete Diopatra leuckarti Kinberg, 1865 (Onuphidae) in Hawaii: Community Structure, and Sediment Stabilizing Properties." Zoological Journal of the Linnean Society 80: 191-9.

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- Baldock, J. A., C. A. Masiello, Y. Gélinas, and J. I. Hedges. 2004. "Cycling and Composition of Organic Matter in Terrestrial and Marine Ecosystems." *Marine Chemistry* 92: 39–64.
- Bao, H., Y. Wu, D. Unger, J. Du, L. S. Herbeck, and J. Zhang. 2013. "Impact of the Conversion of Mangroves into Aquaculture Ponds on the Sedimentary Organic Matter Composition in a Tidal Flat Estuary (Hainan Island, China)." Continental Shelf Research 57: 82–91.
- Barash, M. S. 2012. "Mass Extinction of Ocean Organisms at the Paleozoic-Mesozoic Boundary: Effects and Causes." Oceanology 52: 238–48.
- Bell, L., B. Bluhm, and K. Iken. 2016. "Influence of Terrestrial Organic Matter in Marine Food Webs of the Beaufort Sea Shelf and Slope." *Marine Ecology Progress Series* 550: 1–24.
- Bergamino, L., and N. B. Richoux. 2015. "Spatial and Temporal Changes in Estuarine Food Web Structure: Differential Contributions of Marsh Grass Detritus." *Estuaries and Coasts* 38: 367–82.
- Bianchi, T. S. 2011. "The Role of Terrestrially Derived Organic Carbon in the Coastal Ocean: A Changing Paradigm and the Priming Effect." Proceedings of the National Academy of Sciences of the United States of America 108: 19473–81.
- Bianchi, T. S., R. C. Aller, T. B. Atwood, C. J. Brown, L. A. Buatois, L. A. Levin, J. S. Levinton, et al. 2021. "What Global Biogeochemical Consequences Will Marine Animal–Sediment Interactions Have during Climate Change?" *Elementa: Science* of the Anthropocene 9: 180.
- Bianchi, T. S., X. Cui, N. E. Blair, D. J. Burdige, T. I. Eglinton, and V. Galy. 2018. "Centers of Organic Carbon Burial and Oxidation at the Land-Ocean Interface." Organic Geochemistry 115: 138–55.
- Birchenough, S. N. R., H. Reiss, S. Degraer, N. Mieszkowska, Á. Borja, L. Buhl-Mortensen, U. Braeckman, et al. 2015. "Climate Change and Marine Benthos: A Review of Existing Research and Future Directions in the North Atlantic." Wiley Interdisciplinary Reviews: Climate Change 6: 203–23.
- Blair, N. E., and R. C. Aller. 2012. "The Fate of Terrestrial Organic Carbon in the Marine Environment." *Annual Review of Marine Science* 4: 401–23.
- Bongiorni, L., F. Fiorentino, R. Auriemma, F. B. Aubry, E. Camatti, F. Camin, F. Nasi, M. Pansera, L. Ziller, and J. Grall. 2016. "Food Web of a Confined and Anthropogenically Affected Coastal Basin (the Mar Piccolo of Taranto) Revealed by Carbon and Nitrogen Stable Isotopes Analyses." *Environmental Science and Pollution Research International* 23: 12725–38.
- Bongiorni, L., F. Nasi, F. Fiorentino, R. Auriemma, F. Rampazzo, M. C. Nordström, and D. Berto. 2018. "Contribution of Deltaic Wetland Food Sources to Coastal Macrobenthic Consumers (Po River Delta, North Adriatic Sea)." Science of the Total Environment 643: 1373–86.
- Bonifácio, P., S. Bourgeois, C. Labrune, J. M. Amouroux, K. Escoubeyrou, R. Buscail, A. Romero-Ramirez, et al. 2014. "Spatiotemporal Changes in Surface Sediment Characteristics and Benthic Macrofauna Composition off the Rhône River in Relation to Its Hydrological Regime." *Estuarine, Coastal and Shelf Science* 151: 196–209.
- Boon, A. R., and G. C. A. Duineveld. 2012. "Phytopigments and Fatty Acids in the Gut of the Deposit-Feeding Heart Urchin *Echinocardium cordatum* in the Southern North Sea: Selective

Feeding and its Contribution to the Benthic Carbon Budget." Journal of Sea Research 67: 77–84.

- Bouillon, S., A. V. Raman, P. Dauby, and F. Dehairs. 2002. "Carbon and Nitrogen Stable Isotope Ratios of Subtidal Benthic Invertebrates in An Estuarine Mangrove Ecosystem (Andhra Pradesh, India)." *Estuarine, Coastal and Shelf Science* 54: 901–13.
- Boyer, J. N., and P. M. Groffman. 1996. "Bioavailability of Water Extractable Organic Carbon Fractions in Forest and Agricultural Soil Profiles." Soil Biology and Biochemistry 28: 783–90.
- Bradshaw, C. J. A., I. G. Warkentin, and N. S. Sodhi. 2009. "Urgent Preservation of Boreal Carbon Stocks and Biodiversity." *Trends in Ecology & Evolution* 24: 541–8.
- Bring, A., and G. Destouni. 2011. "Relevance of Hydro-Climatic Change Projection and Monitoring for Assessment of Water Cycle Changes in the Arctic." *Ambio* 40: 361–9.
- Bui, T. H., and S. Y. Lee. 2015. "Endogenous Cellulase Production in the Leaf Litter Foraging Mangrove Crab Parasesarma erythodactyla." Comparative Biochemistry and Physiology Part B: Biochemistry and Molecular Biology 179: 27–36.
- Burd, A. B., S. Frey, A. Cabre, T. Ito, N. M. Levine, C. Lønborg, M. Long, et al. 2016. "Terrestrial and Marine Perspectives on Modeling Organic Matter Degradation Pathways." *Global Change Biology* 22: 121–36.
- Burdige, D. J. 2005. "Burial of Terrestrial Organic Matter in Marine Sediments: A Re-assessment." *Global Biogeochemical Cycles* 19: 4011.
- Burdige, D. J. 2007. "Preservation of Organic Matter in Marine Sediments: Controls, Mechanisms, and an Imbalance in Sediment Organic Carbon Budgets?" *Chemical Reviews* 107: 467–85.
- Cannicci, S., D. Burrows, S. Fratini, T. J. Smith, J. Offenberg, and F. Dahdouh-Guebas. 2008. "Faunal Impact on Vegetation Structure and Ecosystem Function in Mangrove Forests: A Review." Aquatic Botany 89: 186–200.
- Canuel, E. A., S. S. Cammer, H. A. McIntosh, and C. R. Pondell. 2012. "Climate Change Impacts on the Organic Carbon Cycle at the Land-Ocean Interface." *Annual Review of Earth and Planetary Sciences* 40: 685–711.
- Capelle, D. W., Z. Z. A. Kuzyk, T. Papakyriakou, C. Guéguen, L. A. Miller, and R. W. Macdonald. 2020. "Effect of Terrestrial Organic Matter on Ocean Acidification and CO<sub>2</sub> Flux in an Arctic Shelf Sea." *Progress in Oceanography* 185: 102319.
- Cari, I., C. Andrade, E. Quiroga, and E. Mutschke. 2020. "Benthic Trophic Structure of a Patagonian Fjord (47° S): The Role of Hydrographic Conditions in the Food Supply in a Glaciofluvial System." *Estuarine, Coastal and Shelf Science* 233: 106536.
- Carregosa, V., C. Velez, A. Pires, A. M. V. M. Soares, E. Figueira, and R. Freitas. 2014. "Physiological and Biochemical Responses of the Polychaete *Diopatra neapolitana* to Organic Matter Enrichment." *Aquatic Toxicology* 155: 32–42.
- Chen, Q., G. Xu, S. Zhang, and K. Ma. 2018. "Consumption of an Exotic Plant (*Spartina alterniflora*) by the Macrobenthic Fauna in a Mangrove Wetland at Zhanjiang, China." Wetlands 38: 327–35.
- Christofoletti, R. A., G. Y. Hattori, and M. A. A. Pinheiro. 2013. "Food Selection by a Mangrove Crab: Temporal Changes in Fasted Animals." *Hydrobiologia* 702: 63–72.

- Churchwell, R. T., S. J. Kendall, A. L. Blanchard, K. H. Dunton, and A. N. Powell. 2016. "Natural Disturbance Shapes Benthic Intertidal Macroinvertebrate Communities of High Latitude River Deltas." *Estuaries and Coasts* 39: 798–814.
- Clark, J. B., A. Mannino, M. Tzortziou, R. G. M. Spencer, and P. Hernes. 2022. "The Transformation and Export of Organic Carbon across an Arctic River-Delta-Ocean Continuum." *Journal of Geophysical Research: Biogeosciences* 4: 88–100.
- Coma, R., M. Ribes, J. M. Gili, and M. Zabala. 2000. "Seasonality in Coastal Benthic Ecosystems." *Trends in Ecology & Evolution* 15 (11): 448–53.
- Coppola, A. I., D. B. Wiedemeier, V. Galy, N. Haghipour, U. M. Hanke, G. S. Nascimento, M. Usman, et al. 2018. "Global-Scale Evidence for the Refractory Nature of Riverine Black Carbon." *Nature Geoscience* 11: 584–8.
- Cragg, S. M., G. T. Beckham, N. C. Bruce, T. D. H. Bugg, D. L. Distel, P. Dupree, A. G. Etxabe, et al. 2015. "Lignocellulose Degradation Mechanisms across the Tree of Life." *Current Opinion in Chemical Biology* 29: 108–19.
- Cragg, S. M., D. A. Friess, L. G. Gillis, S. M. Trevathan-Tackett, O. M. Terrett, J. E. M. Watts, D. L. Distel, and P. Dupree. 2020. "Vascular Plants Are Globally Significant Contributors to Marine Carbon Fluxes and Sinks." *Annual Review of Marine Science* 12: 469–97.
- Crain, C. M., K. Kroeker, and B. S. Halpern. 2008. "Interactive and Cumulative Effects of Multiple Human Stressors in Marine Systems." *Ecology Letters* 11: 1304–15.
- Cummings, V., S. Thrush, J. Hewitt, A. Norkko, and S. Pickmere. 2003. "Terrestrial Deposits on Intertidal Sandflats: Sediment Characteristics As Indicators of Habitat Suitability for Recolonising Macrofauna." *Marine Ecology Progress Series* 253: 39–54.
- Cummings, V., K. Vopel, and S. Thrush. 2009. "Terrigenous Deposits in Coastal Marine Habitats: Influences on Sediment Geochemistry and Behaviour of Post-Settlement Bivalves." *Marine Ecology Progress Series* 383: 173–85.
- Currin, C., S. Newell, and H. Paerl. 1995. "The Role of Standing Dead Spartina alterniflora and Benthic Microalgae in Salt Marsh Food Webs:Considerations Based on Multiple Stable Isotope Analysis." Marine Ecology Progress Series 121: 99–116.
- Darnaude, A. M., C. Salen-Picard, and M. L. Harmelin-Vivien. 2004. "Depth Variation in Terrestrial Particulate Organic Matter Exploitation by Marine Coastal Benthic Communities off the Rhone River Delta (NW Mediterranean)." *Marine Ecology Progress Series* 275: 47–57.
- Datry, T., S. T. Larned, and K. Tockner. 2014. "Intermittent Rivers: A Challenge for Freshwater Ecology." *BioScience* 64: 229–35.
- De Wit, H. A., S. Valinia, G. A. Weyhenmeyer, M. N. Futter, P. Kortelainen, K. Austnes, D. O. Hessen, A. Räike, H. Laudon, and J. Vuorenmaa. 2016. "Current Browning of Surface Waters Will Be further Promoted by Wetter Climate." *Environmental Science and Technology Letters* 3: 430–5.
- Deininger, A., and H. Frigstad. 2019. "Reevaluating the Role of Organic Matter Sources for Coastal Eutrophication, Oligotrophication and Ecosystem Health." *Frontiers in Marine Science* 6: 210.
- del Campo, R., R. Corti, and G. Singer. 2021. "Flow Intermittence Alters Carbon Processing in Rivers through Chemical Diversification of Leaf Litter." *Limnology and Oceanography Letters* 6(5): 232–42.

- Demopoulos, A. W. J., B. Fry, and C. R. Smith. 2007. "Food Web Structure in Exotic and Native Mangroves: A Hawaii-Puerto Rico Comparison." *Oecologia* 153: 675–86.
- De Oliveira, A. B., A. E. Rizzo, E. Da Conceicao, and G. Couto. 2012. "Benthic Macrofauna Associated with Decomposition of Leaves in a Mangrove Forest in Ilhéus, State of Bahia, Brazil." *Journal of the Marine Biological Association of the United Kingdom* 92(7): 1479–87.
- Deuser, W. G. 1988. "Whither Organic Carbon?" Nature 332: 396-7.
- Dias, E., P. Morais, A. M. Cotter, C. Antunes, and J. C. Hoffman. 2016. "Estuarine Consumers Utilize Marine, Estuarine and Terrestrial Organic Matter and Provide Connectivity among These Food Webs." *Marine Ecology Progress Series* 554: 21–34.
- Dickens, A. F., J. A. Baldock, R. J. Smernik, S. G. Wakeham, T. S. Arnarson, Y. Gélinas, and J. I. Hedges. 2006. "Solid-State 13C NMR Analysis of Size and Density Fractions of Marine Sediments: Insight into Organic Carbon Sources and Preservation Mechanisms." *Geochimica et Cosmochimica Acta* 70: 666–86.
- Distel, D. L., W. Morrill, N. MacLaren-Toussaint, D. Franks, and J. Waterbury. 2002. "Teredinibacter turnerae Gen. Nov., Sp. Nov., a Dinitrogen-Fixing, Cellulolytic, Endosymbiotic Gamma-Proteobacterium Isolated from the Gills of Wood-Boring Molluscs (Bivalvia: Teredinidae)." International Journal of Systematic and Evolutionary Microbiology 52: 2261–9.
- Doering, P. H., C. A. Oviatt, and J. R. Kelly. 1986. "The Effects of the Filter-Feeding Clam Mercenaria mercenaria on Carbon Cycling in Experimental Marine Mesocosms." Journal of Marine Research 44: 839–61.
- Drenzek, N. J., K. A. Hughen, D. B. Montlucecon, J. R. Southon, G. M. dos Santos, E. R. M. Druffel, L. Giosan, and T. I. Eglinton. 2009. "A New Look at Old Carbon in Active Margin Sediments." *Geology* 37: 239–42.
- Drylie, T. P., A. M. Lohrer, H. R. Needham, and C. A. Pilditch. 2020. "Taxonomic and Functional Response of Estuarine Benthic Communities to Experimental Organic Enrichment: Consequences for Ecosystem Function." Journal of Experimental Marine Biology and Ecology 532: 151455.
- Dunton, K. H., S. V. Schonberg, and L. W. Cooper. 2012. "Food Web Structure of the Alaskan Nearshore Shelf and Estuarine Lagoons of the Beaufort Sea." *Estuaries and Coasts* 35: 416–35.
- Dunton, K. H., T. Weingartner, and E. C. Carmack. 2006. "The Nearshore Western Beaufort Sea Ecosystem: Circulation and Importance of Terrestrial Carbon in Arctic Coastal Food Webs." *Progress in Oceanography* 71(2–4): 362–78.
- Ehrnsten, E., O. P. Savchuk, and B. G. Gustafsson. 2022. "Modelling the Effects of Benthic Fauna on Carbon, Nitrogen and Phosphorus Dynamics in the Baltic Sea." *Biogeosciences* 19: 3337–67.
- Fanelli, E., J. E. Cartes, F. Badalamenti, P. Rumolo, and M. Sprovieri. 2009. "Trophodynamics of Suprabenthic Fauna on Coastal Muddy Bottoms of the Southern Tyrrhenian Sea (Western Mediterranean)." *Journal of Sea Research* 61: 174–87.
- Farella, N., M. Lucotte, P. Louchouarn, and M. Roulet. 2001. "Deforestation Modifying Terrestrial Organic Transport in the Rio Tapajós, Brazilian Amazon." Organic Geochemistry 32: 1443–58.
- Feng, J., Q. Huang, H. Chen, J. Guo, and G. Lin. 2018. "Restoration of Native Mangrove Wetlands Can Reverse Diet Shifts of

Benthic Macrofauna Caused by Invasive Cordgrass." *Journal* of Applied Ecology 55: 905–16.

- Feng, X., J. E. Vonk, B. E. Van Dongen, Ö. Gustafsson, I. P. Semiletov, O. V. Dudarev, Z. Wang, D. B. Montluçon, L. Wacker, and T. I. Eglinton. 2013. "Differential Mobilization of Terrestrial Carbon Pools in Eurasian Arctic River Basins." *Proceedings of the National Academy of Sciences of the United States of America* 110: 14168–73.
- Fernández-Rodríguez, V., C. S. G. Santos, and A. P. F. Pires. 2019. "Meta-Analysis of the Effects of Organic Matter on Polychaetes of the East Coast of South America." *Marine Environmental Research* 149: 148–56.
- Forest, A., M. Sampei, H. Hattori, R. Makabe, H. Sasaki, M. Fukuchi, P. Wassmann, and L. Fortier. 2007. "Particulate Organic Carbon Fluxes on the Slope of the Mackenzie Shelf (Beaufort Sea): Physical and Biological Forcing of Shelf-Basin Exchanges." Journal of Marine Systems 68: 39–54.
- Forrest, B. M., P. A. Gillespie, C. D. Cornelisen, and K. M. Rogers. 2007. "Multiple Indicators Reveal River Plume Influence on Sediments and Benthos in a New Zealand Coastal Embayment." New Zealand Journal of Marine and Freshwater Research 41: 13–24.
- Frigstad, H., G. S. Andersen, H. C. Trannum, M. McGovern, L.-J. Naustvoll, Ø. Kaste, A. Deininger, and D. Ø. Hjermann. 2023.
  "Three Decades of Change in the Skagerrak Coastal Ecosystem, Shaped by Eutrophication and Coastal Darkening." *Estuarine, Coastal and Shelf Science* 283: 108193.
- Frouin, P. 2000. "Effects of Anthropogenic Disturbances of Tropical Soft-Bottom Benthic Communities." *Marine Ecology Progress Series* 194: 39–53.
- Galeron, J., L. Menot, N. Renaud, P. Crassous, A. Khripounoff, C. Treignier, and M. Sibuet. 2009. "Spatial and Temporal Patterns of Benthic Macrofaunal Communities on the Deep Continental Margin in the Gulf of Guinea." *Deep-Sea Research II* 56: 12.
- Galletti, Y., S. Becagli, A. Di Sarra, M. Gonnelli, E. Pulido-Villena, D. M. Sferlazzo, R. Traversi, S. Vestri, and C. Santinelli. 2020.
  "Atmospheric Deposition of Organic Matter at a Remote Site in the Central Mediterranean Sea: Implications for the Marine Ecosystem." *Biogeosciences* 17: 3669–84.
- Galloway, A. W. E., K. H. Britton-Simmons, D. O. Duggins, P. W. Gabrielson, and M. T. Brett. 2012. "Fatty Acid Signatures Differentiate Marine Macrophytes at Ordinal and Family Ranks." *Journal of Phycology* 48: 956–65.
- Galy, V., C. France-Lanord, O. Beyssac, P. Faure, H. Kudrass, and F. Palhol. 2007. "Efficient Organic Carbon Burial in the Bengal Fan Sustained by the Himalayan Erosional System." *Nature* 450: 407–10.
- Geraldi, N. R., A. Ortega, O. Serrano, P. I. Macreadie, C. E. Lovelock, D. Krause-Jensen, H. Kennedy, et al. 2019.
  "Fingerprinting Blue Carbon: Rationale and Tools to Determine the Source of Organic Carbon in Marine Depositional Environments." *Frontiers in Marine Science* 6: 1–9.
- Giménez, L., A. I. Borthagaray, M. Rodríguez, A. Brazeiro, C. Dimitriadis, L. Gimínez, A. I. Borthagaray, M. Rodríguez, A. Brazeiro, and C. Dimitriadis. 2005. "Scale-Dependent Patterns of Macrofaunal Distribution in Soft-Sediment Intertidal Habitats along a Large-Scale Estuarine Gradient." *Helgoland Marine Research* 59: 224–36.

- Gladstone-Gallagher, R. V., C. J. Lundquist, and C. A. Pilditch. 2014. "Response of Temperate Intertidal Benthic Assemblages to Mangrove Detrital Inputs." *Journal of Experimental Marine Biology and Ecology* 460: 80–8.
- Gladstone-Gallagher, R. V., S. Mangan, S. F. Thrush, and C. A. Pilditch. 2020. "Porewater Nutrient Enrichment Alters Benthic-Pelagic Coupling on Intertidal Sandflats." *Journal of Sea Research* 159: 101876.
- Godbold, J. A., R. Rosenberg, and M. Solan. 2009. "Species-Specific Traits Rather than Resource Partitioning Mediate Diversity Effects on Resource Use." *PLoS One* 4: e7423.
- Gordon, E. S., and M. A. Goñi. 2004. "Controls on the Distribution and Accumulation of Terrigenous Organic Matter in Sediments from the Mississippi and Atchafalaya River Margin." *Marine Chemistry* 92: 331–52.
- Gounand, I., E. Harvey, C. J. Little, and F. Altermatt. 2018. "Meta-Ecosystems 2.0: Rooting the Theory into the Field." *Trends in Ecology and Evolution* 33: 36–46.
- Griffiths, J. R., M. Kadin, F. J. A. Nascimento, T. Tamelander, A. Törnroos, S. Bonaglia, E. Bonsdorff, et al. 2017. "The Importance of Benthic-Pelagic Coupling for Marine Ecosystem Functioning in a Changing World." *Global Change Biology* 23: 2179–96.
- Gunnarsson, J. S., M. E. Granberg, H. C. Nilsson, R. Rosenberg, and B. Hellman. 1999. "Influence of Sediment-Organic Matter Quality on Growth and Polychlorobiphenyl Bioavailability in Echinodermata (*Amphiura filiformis*)." Environmental Toxicology and Chemistry 18: 1534–43.
- Guo, L., I. Semiletov, Ö. Gustafsson, J. Ingri, P. Andersson, O. Dudarev, and D. White. 2004. "Characterization of Siberian Arctic Coastal Sediments: Implications for Terrestrial Organic Carbon Export." *Global Biogeochemical Cycles* 18(1).
- Hagberg, J., and B. G. Tunberg. 2000. "Studies on the Covariation between Physical Factors and the Long-Term Variation of the Marine Soft Bottom Macrofauna in Western Sweden." *Estuarine, Coastal and Shelf Science* 50: 373–85.
- Hansell, D. A., C. A. Carlson, D. J. Repeta, and R. Schlitzer. 2009."Dissolved Organic Matter in the Ocean a Controversy Stimulates New Insights." *Oceanography* 22: 202–11.
- Harkantra, S. N., and N. R. Rodrigues. 2004. "Numerical Analyses of Soft Bottom Macroinvertebrates to Diagnose the Pollution in Tropical Coastal Waters." *Environmental Monitoring and Assessment* 93: 251–75.
- Harmelin-Vivien, M. L., D. Bănaru, J. Dierking, R. Hermand, Y. Letourneur, and C. Salen-Picard. 2009. "Linking Benthic Biodiversity to the Functioning of Coastal Ecosystems Subjected to River Runoff (NW Mediterranean)." Animal Biodiversity and Conservation 32: 135–45.
- Harris, C. M., N. D. McTigue, J. W. McClelland, and K. H. Dunton.2018. "Do High Arctic Coastal Food Webs Rely on a Terrestrial Carbon Subsidy?" *Food Webs* 15: e00081.
- He, W., M. Chen, M. A. Schlautman, and J. Hur. 2016. "Dynamic Exchanges between DOM and POM Pools in Coastal and Inland Aquatic Ecosystems: A Review." Science of the Total Environment 551: 415–28.
- Hedges, J. I., and R. G. Keil. 1995. "Sedimentary Organic Matter Preservation: An Assessment and Speculative Synthesis." *Marine Chemistry* 49: 81–115.
- Hedges, J. I., R. G. Keil, and R. Benner. 1997. "What Happens to Terrestrial Organic Matter in the Ocean?" Organic Geochemistry 27: 195–212.

- Hermand, R., C. Salen-Picard, E. Alliot, and C. Degiovanni. 2008. "Macrofaunal Density, Biomass and Composition of Estuarine Sediments and Their Relationship to the River Plume of the Rhone River (NW Mediterranean)." *Estuarine, Coastal and Shelf Science* 79: 367–76.
- Hoeinghaus, D. J., J. P. Vieira, C. S. Costa, C. E. Bemvenuti, K. O. Winemiller, and A. M. Garcia. 2011. "Estuary Hydrogeomorphology Affects Carbon Sources Supporting Aquatic Consumers within and among Ecological Guilds." *Hydrobiologia* 673: 79–92.
- Hughes, J. E., L. A. Deegan, B. J. Peterson, R. M. Holmes, and B. Fry. 2000. "Nitrogen Flow through the Food Web in the Oligohaline Zone of a New England Estuary." *Ecology* 81: 433–52.
- Hwang, J., M. Kim, S. J. Manganini, C. P. McIntyre, N. Haghipour, J. Park, R. A. Krishfield, R. W. MacDonald, F. A. McLaughlin, and T. I. Eglinton. 2015. "Temporal and Spatial Variability of Particle Transport in the Deep Arctic Canada Basin." *Journal* of Geophysical Research: Oceans 120: 2784–99.
- IPCC. 2021. "Climate Change 2021: The Physical Science Basis." In Contribution of Working Group I to the Sixth Assessment Report of the Intergovernmental Panel on Climate Change, edited by V. Masson-Delmotte, P. Zhai, A. Pirani, S. L. Connors, C. Péan, S. Berger, N. Caud, Y. Chen, L. Goldfarb, M. I. Gomis, M. Huang, K. Leitzell, E. Lonnoy, J. B. R. Matthews, T. K. Maycock, T. Waterfield, O. Yelekçi, R. Yu, and B. Zhou. Cambridge: Cambridge University Press.
- Janas, U., D. Burska, H. Kendzierska, D. Pryputniewicz-Flis, and K. Łukawska-Matuszewska. 2019. "Importance of Benthic Macrofauna and Coastal Biotopes for Ecosystem Functioning – Oxygen and Nutrient Fluxes in the Coastal Zone." *Estuarine, Coastal and Shelf Science* 225: 106238.
- Jędruch, A., M. Bełdowska, and M. Ziółkowska. 2019. "The Role of Benthic Macrofauna in the Trophic Transfer of Mercury in a Low-Diversity Temperate Coastal Ecosystem (Puck Lagoon, Southern Baltic Sea)." Environmental Monitoring and Assessment 191: 137–7.
- Jokinen, S. A., T. Jilbert, R. Tiihonen-Filppula, and K. Koho. 2020. "Terrestrial Organic Matter Input Drives Sedimentary Trace Metal Sequestration in a Human-Impacted Boreal Estuary." *Science of the Total Environment* 717: 137047.
- Jones, M. W., A. I. Coppola, C. Santín, T. Dittmar, R. Jaffé, S. H. Doerr, and T. A. Quine. 2020. "Fires Prime Terrestrial Organic Carbon for Riverine Export to the Global Oceans." *Nature Communications* 11: 1–8.
- Jones, M. T., and S. R. Gislason. 2008. "Rapid Releases of Metal Salts and Nutrients Following the Deposition of Volcanic Ash into Aqueous Environments." *Geochimica et Cosmochimica Acta* 72: 3661–80.
- Jones, D. O. B., A. Yool, C. L. Wei, S. A. Henson, H. A. Ruhl, R. A. Watson, and M. Gehlen. 2014. "Global Reductions in Seafloor Biomass in Response to Climate Change." *Global Change Biology* 20: 1861–72.
- Kanaya, G., T. Suzuki, and E. Kikuchi. 2011. "Spatio-Temporal Variations in Macrozoobenthic Assemblage Structures in a River-Affected Lagoon (Idoura Lagoon, Sendai Bay, Japan): Influences of Freshwater Inflow." *Estuarine, Coastal and Shelf Science* 92: 169–79.
- Kandasamy, S., and B. Nagender Nath. 2016. "Perspectives on the Terrestrial Organic Matter Transport and Burial along the

Land-Deep Sea Continuum: Caveats in our Understanding of Biogeochemical Processes and Future Needs." *Frontiers in Marine Science* 3: 259.

- Kawaida, S., K. Nanjo, N. Ohtsuchi, H. Kohno, and M. Sano. 2019. "Cellulose Digestion Abilities Determine the Food Utilization of Mangrove Estuarine Crabs." *Estuarine, Coastal and Shelf Science* 222: 43–52.
- Kędra, M., K. Kuliński, W. Walkusz, and J. Legeżyńska. 2012. "The Shallow Benthic Food Web Structure in the High Arctic Does Not Follow Seasonal Changes in the Surrounding Environment." *Estuarine, Coastal and Shelf Science* 114: 183–91.
- Kim, J. H., F. Peterse, V. Willmott, D. Klitgaard Kristensen, M. Baas, S. Schouten, and J. S. Sinninghe Damstéa. 2011. "Large Ancient Organic Matter Contributions to Arctic Marine Sediments (Svalbard)." *Limnology and Oceanography* 56: 1463–74.
- King, A. J., S. M. Cragg, Y. Li, J. Dymond, M. J. Guille, D. J. Bowles, N. C. Bruce, I. A. Graham, and S. J. McQueen-Mason. 2010. "Molecular Insight into Lignocellulose Digestion by a Marine Isopod in the Absence of Gut Microbes." *Proceedings* of the National Academy of Sciences of the United States of America 107: 5345–50.
- Klages, M., A. Boetius, J. P. Christensen, H. Deubel, D. Piepenburg, I. Schewe, and T. Soltwedel. 2004. "The Benthos of Arctic Seas and Its Role for the Organic Carbon Cycle at the Seafloor." In *The Organic Carbon Cycle in the Arctic Ocean*, edited by R. Stein and R. W. Macdonald, 139–67. Berlin: Springer.
- Kokarev, V., M. Tachon, M. Austad, M. McGovern, and H. Reiss. 2021. "Strong Macrobenthic Community Differentiation Among Sub-Arctic Deep Fjords on Small Spatial Scales." *Estuarine, Coastal and Shelf Science* 252: 107271.
- Komada, T., and C. E. Reimers. 2001. "Resuspension-Induced Partitioning of Organic Carbon between Solid and Solution Phases from a River–Ocean Transition." *Marine Chemistry* 76: 155–74.
- Kristensen, E., S. Bouillon, T. Dittmar, and C. Marchand. 2008."Organic Carbon Dynamics in Mangrove Ecosystems: A Review." *Aquatic Botany* 89: 201–19.
- Kristensen, E., M. Delefosse, C. O. Quintana, M. R. Flindt, and T. Valdemarsen. 2014. "Influence of Benthic Macrofauna Community Shifts on Ecosystem Functioning in Shallow Estuaries." Frontiers in Marine Science 1: 1–14.
- Kristensen, E., G. Penha-Lopes, M. Delefosse, T. Valdemarsen, C. O. Quintana, and G. T. Banta. 2012. "What Is Bioturbation? The Need for a Precise Definition for Fauna in Aquatic Sciences." *Marine Ecology Progress Series* 446: 285–302.
- Kristensen, E., C. O. Quintana, and S. G. G. Petersen. 2022. "The Role of Biogenic Structures for Greenhouse Gas Balance in Vegetated Intertidal Wetlands." In *Carbon Mineralization in Coastal Wetlands*, edited by X. Ouyang, S. Y. Lee, Y. F. D. Lai, and C. Marchand, 233–67. Amsterdam: Elsevier.
- Kumar, V., M. Tiwari, S. Nagoji, and S. Tripathi. 2016. "Evidence of Anomalously Low  $\delta^{13}$ C of Marine Organic Matter in an Arctic Fjord." *Scientific Reports* 6: 1–9.
- Laurino, I. R. A., T. Z. Serafini, T. M. Costa, and R. A. Christofoletti. 2021. "The Role of Estuarine Macrofaunal Patterns for the Management of Marine Protected Areas in a Changing World." *Journal for Nature Conservation* 63: 126042.
- Lautenschlager, A. D., T. G. Matthews, and G. P. Quinn. 2014. "Utilization of Organic Matter by Invertebrates along an

Estuarine Gradient in an Intermittently Open Estuary." *Estuarine, Coastal and Shelf Science* 149: 232–43.

- Lee, S. Y. 1999. "The Effect of Mangrove Leaf Litter Enrichment on Macrobenthic Colonization of Defaunated Sandy Substrates." *Estuarine, Coastal and Shelf Science* 49: 703–12.
- Levin, L. A., W. Ekau, A. J. Gooday, F. Jorissen, J. J. Middelburg, S. W. A. Naqvi, C. Neira, N. N. Rabalais, and J. Zhang. 2009. "Effects of Natural and Human-Induced Hypoxia on Coastal Benthos." *Biogeosciences* 6: 2063–98.
- Li, Y., C. Fu, L. Zeng, Q. Zhou, H. Zhang, C. Tu, L. Li, and Y. Luo. 2021. "Black Carbon Contributes Substantially to Allochthonous Carbon Storage in Deltaic Vegetated Coastal Habitats." *Environmental Science & Technology* 55: 6495–504.
- Liao, Y., L. Shou, Y. Tang, A. Gao, Q. Chen, X. Yan, and J. Chen. 2018. "Influence of Two Nonindigenous Plants on Intertidal Macrobenthic Communities in Ximen Island Special Marine Protected Area, China." *Ecological Engineering* 112: 96–104.
- Liénart, C., A. R. Cirtwill, M. L. Hedgespeth, and C. Bradshaw. 2022. "A Sprinkling of Gold Dust: Pine Pollen as a Carbon Source in Baltic Sea Coastal Food Webs." *Limnology and Oceanography* 67: 53–65.
- Lillebø, A. I., M. R. Flindt, M. A. Pardal, P. G. Cardoso, S. M. Ferreira, and J. C. Marques. 2007. "The Faunal Role in the Degradation of the Common Intertidal Salt Marsh Plant *Scirpus maritimus.*" *Hydrobiologia* 579: 369–78.
- Lillebø, A. I., M. R. Flindt, M. Â. Pardal, and J. C. Marques. 1999. "The Effect of Macrofauna, Meiofauna and Microfauna on the Degradation of Spartina Maritima Detritus from a Salt Marsh Area." Acta Oecologica 20: 249–58.
- Lohrer, A. M., S. F. Thrush, C. J. Lundquist, K. Vopel, J. E. Hewitt, and P. E. Nicholls. 2006. "Deposition of Terrigenous Sediment on Subtidal Marine Macrobenthos: Response of Two Contrasting Community Types." *Marine Ecology Progress Series* 307: 115–25.
- Loreau, M., N. Mouquet, and R. D. Holt. 2003. "Meta-Ecosystems: A Theoretical Framework for a Spatial Ecosystem Ecology." *Ecology Letters* 6: 673–9.
- Louchouarn, P., M. Lucotte, and N. Farella. 1999. "Historical and Geographical Variations of Sources and Transport of Terrigenous Organic Matter within a Large-Scale Coastal Environment." Organic Geochemistry 30: 675–99.
- Mancinelli, G., and L. Rossi. 2002. "The Influence of Allochthonous Leaf Detritus on the Occurrence of Crustacean Detritivores in the Soft-Bottom Macrobenthos of the Po River Delta Area (Northwestern Adriatic Sea)." *Estuarine, Coastal and Shelf Science* 54: 849–61.
- Marcelina, Z., S. Adam, and R. Pierre. 2018. "Spatial and Temporal Variability of Organic Matter Sources and Food Web Structure across Benthic Habitats in a Low Diversity System (Southern Baltic Sea)." Journal of Sea Research 141: 47–60.
- Massicotte, P., E. Asmala, C. Stedmon, and S. Markager. 2017. "Global Distribution of Dissolved Organic Matter along the Aquatic Continuum: Across Rivers, Lakes and Oceans." *Science of the Total Environment* 609: 180–91.
- McClelland, J. W., R. M. Holmes, K. H. Dunton, and R. W. Macdonald. 2012. "The Arctic Ocean Estuary." *Estuaries and Coasts* 35: 353–68.
- McGovern, M., A. K. Pavlov, A. Deininger, M. A. Granskog, E. Leu, J. E. Søreide, and A. E. Poste. 2020. "Terrestrial Inputs Drive

Seasonality in Organic Matter and Nutrient Biogeochemistry in a High Arctic Fjord System (Isfjorden, Svalbard)." *Frontiers in Marine Science* 7: 542563.

- McGovern, M., A. E. Poste, E. Oug, P. E. Renaud, and H. C. Trannum. 2020. "Riverine Impacts on Benthic Biodiversity and Functional Traits: A Comparison of Two Sub-Arctic Fjords." *Estuarine, Coastal and Shelf Science* 240: 106774.
- McInerney, F. A., and S. L. Wing. 2011. "The Paleocene-Eocene Thermal Maximum: A Perturbation of Carbon Cycle, Climate, and Biosphere with Implications for the Future." *Annual Review of Earth and Planetary Sciences* 39: 489–516.
- McLeod, R. J., and S. R. Wing. 2009. "Strong Pathways for Incorporation of Terrestrially Derived Organic Matter into Benthic Communities." *Estuarine, Coastal and Shelf Science* 82: 645–53.
- McTigue, N. D., and K. H. Dunton. 2014. "Trophodynamics and Organic Matter Assimilation Pathways in the Northeast Chukchi Sea, Alaska." *Deep Sea Research Part II: Topical Studies in Oceanography* 102: 84–96.
- Meybeck, M. 1982. "Carbon, Nitrogen, and Phosphorus Transport by World Rivers." *American Journal of Science* 282: 401–50.
- Middelburg, J. J., and P. M. J. Herman. 2007. "Organic Matter Processing in Tidal Estuaries." *Marine Chemistry* 106: 127–47.
- Misic, C., L. Gaozza, M. Petrillo, and A. Covazzi Harriague. 2016. "The Allochthonous Material Input in the Trophodynamic System of the Shelf Sediments of the Gulf of Tigullio (Ligurian Sea, NW Mediterranean)." *Marine Environmental Research* 120: 9–19.
- Mudge, S. M., and C. E. Duce. 2005. "Identifying the Source, Transport Path and Sinks of Sewage Derived Organic Matter." *Environmental Pollution* 136: 209–20.
- Munguia, P., and A. F. Ojanguren. 2015. "Bridging the Gap in Marine and Terrestrial Studies." *Ecosphere* 6: 1–4.
- Nakagawa, S., G. Samarasinghe, N. R. Haddaway, M. J. Westgate, R. E. O'Dea, D. W. A. Noble, and M. Lagisz. 2019. "Research Weaving: Visualizing the Future of Research Synthesis." *Trends in Ecology and Evolution* 34: 224–38.
- Nasi, F., R. Auriemma, F. Relitti, M. Bazzaro, D. Cassin, and T. Cibic. 2020. "Structural and Functional Response of Coastal Macrofaunal Community to Terrigenous Input from the Po River (Northern Adriatic Sea)." *Estuarine, Coastal and Shelf Science* 235: 106548.
- Nasi, F., L. Ferrante, F. Alvisi, E. Bonsdorff, R. Auriemma, and T. Cibic. 2020. "Macrofaunal Bioturbation Attributes in Relation to Riverine Influence: What Can We Learn from the Po River Lagoonal System (Adriatic Sea)? Estuarine." Coastal and Shelf Science 232: 106405.
- Neira, C., L. A. Levin, E. D. Grosholz, and G. Mendoza. 2007. "Influence of Invasive Spartina Growth Stages on Associated Macrofaunal Communities." *Biological Invasions* 9: 975–93.
- Nelson, N. B., and D. A. Siegel. 2013. "The Global Distribution and Dynamics of Chromophoric Dissolved Organic Matter." *Annual Review of Marine Science* 5: 447–76.
- Netto, S. A., and F. Gallucci. 2003. "Meiofauna and Macrofauna Communities in a Mangrove from the Island of Santa Catarina, South Brazil." *Hydrobiologia* 505: 159–70.
- Netto, S. A., and P. C. Lana. 1997. "Intertidal Zonation of Benthic Macrofauna in a Subtropical Salt Marsh and Nearby Unvegetated Flat (SE, Brazil)." *Hydrobiologia* 353: 171–80.

- Netto, S. A., and P. C. Lana. 1999. "The Role of Above- and Below-Ground Components of *Spartina alterniflora* (Loisel) and Detritus Biomass in Structuring Macrobenthic Associations of Paranagua Bay (SE, Brazil)." *Hydrobiologia* 400: 167–77.
- Norkko, A., S. F. Thrush, J. E. Hewitt, V. J. Cummings, J. Norkko, J. I. Ellis, G. A. Funnell, D. Schultz, and I. MacDonald. 2002.
  "Smothering of Estuarine Sandflats by Terrigenous Clay: The Role of Wind-Wave Disturbance and Bioturbation in Site-Dependent Macrofaunal Recovery." *Marine Ecology Progress Series* 234: 23–41.
- Nuñez, M. A., M. C. Chiuffo, A. Pauchard, and R. D. Zenni. 2021. "Making Ecology Really Global." *Trends in Ecology & Evolution* 36: 766–9.
- Nuwer, J. M., and R. G. Keil. 2005. "Sedimentary Organic Matter Geochemistry of Clayoquot Sound, Vancouver Island, British Columbia." *Limnology and Oceanography* 50: 1119–28.
- Olin, J. A., N. E. Hussey, S. A. Rush, G. R. Poulakis, C. A. Simpfendorfer, M. R. Heupel, and A. T. Fisk. 2013. "Seasonal Variability in Stable Isotopes of Estuarine Consumers under Different Freshwater Flow Regimes." *Marine Ecology Progress Series* 487: 55–69.
- Painter, S. C., D. J. Lapworth, E. M. S. Woodward, S. Kroeger, C. D. Evans, D. J. Mayor, and R. J. Sanders. 2018. "Terrestrial Dissolved Organic Matter Distribution in the North Sea." *Science of the Total Environment* 630: 630–47.
- Panagos, P., P. Borrelli, K. Meusburger, B. Yu, A. Klik, K. J. Lim, J. E. Yang, et al. 2017. "Global Rainfall Erosivity Assessment Based on High-Temporal Resolution Rainfall Records." *Scientific Reports* 7: 1–12.
- Pardo, J. 2023. "pardojcf/TerOMacrofauna\_Review\_Pardo-et-al.: TerOMacrofauna\_Review\_Pardo-et-al.-Bibliometric data (v1.0.0)." Zenodo. Dataset. https://doi.org/10.5281/zenodo. 7702058.
- Parker, R., L. Benson, C. Graves, S. Kröger, and R. Vieira. 2021. "Carbon Stocks and Accumulation Analysis for Secretary of State (SoS) Region: (2020)." Cefas Project Report for Defra. 42 pp.
- Pearson, T., and R. Rosenberg. 1978. "Macrobenthic Succession in Relation to Organic Enrichment and Pollution of the Marine Environment." Oceanography and Marine Biology: An Annual Review 16: 229–311.
- Polis, G. A., W. B. Anderson, and R. D. Holt. 1997. "Toward an Integration of Landscape and Food Web Ecology: The Dynamics of Spatially Subsidized Food Webs." *Annual Review* of Ecology and Systematics 28: 289–316.
- Qiao, J., H. Bao, D. Huang, D.-W. Li, T.-Y. Lee, J.-C. Huang, and S.-J. Kao. 2019. "Runoff-Driven Export of Terrigenous Particulate Organic Matter from a Small Mountainous River: Sources, Fluxes and Comparisons among Different Rivers." *Biogeochemistry* 147: 71–86.
- Quiroga, E., P. Ortiz, R. González-Saldías, B. Reid, F. J. Tapia, I. Pérez-Santos, L. Rebolledo, et al. 2016. "Seasonal Benthic Patterns in a Glacial Patagonian Fford: the Role of Suspended Sediment and Terrestrial Organic Matter." *Marine Ecology Progress Series* 561: 31–50.
- Renz, J. R., M. Powilleit, M. Gogina, M. L. Zettler, C. Morys, and S. Forster. 2018. "Community Bioirrigation Potential (BIPc), an Index to Quantify the Potential for Solute Exchange at the

Sediment-Water Interface." *Marine Environmental Research* 141: 214–24.

- Rezek, R. J., B. Lebreton, B. Sterba-Boatwright, and J. B. Pollack. 2017. "Ecological Structure and Function in a Restored Versus Natural Salt Marsh." *PLoS One* 12: e0189871.
- Rezende, C. E., W. C. Pfeiffer, L. A. Martinelli, E. Tsamakis, J. I. Hedges, and R. G. Keil. 2010. "Lignin Phenols Used to Infer Organic Matter Sources to Sepetiba Bay—RJ, Brasil." *Estuarine, Coastal and Shelf Science* 87: 479–86.
- Richoux, N. B., and P. W. Froneman. 2007. "Assessment of Spatial Variation in Carbon Utilization by Benthic and Pelagic Invertebrates in a Temperate South African Estuary Using Stable Isotope Signatures." *Estuarine, Coastal and Shelf Science* 71: 545–58.
- Riera, P., and P. Richard. 1997. "Temporal Variation of  $\delta^{13}$ C in Particulate Organic Matter and Oyster *Crassostrea gigas* in Marennes-Oleron Bay (France): Effect of Freshwater Inflow." *Marine Ecology Progress Series* 147: 105–15.
- Riisgård, H. U., and P. S. Larsen. 2010. "Particle Capture Mechanisms in Suspension-Feeding Invertebrates." *Marine Ecology Progress Series* 418: 255–93.
- Rodil, I. F., P. Lucena-Moya, T. Tamelander, J. Norkko, and A. Norkko. 2020. "Seasonal Variability in Benthic–Pelagic Coupling: Quantifying Organic Matter Inputs to the Seafloor and Benthic Macrofauna Using a Multi-Marker Approach." *Frontiers in Marine Science* 7: 1–18.
- Rühl, S., C. Thompson, A. M. Queirós, and S. Widdicombe. 2020. "Missing Links in the Study of Solute and Particle Exchange between the Sea Floor and Water Column." *ICES Journal of Marine Science* 77: 1602–16.
- Saito, K., J. E. Walsh, A. Bring, R. Brown, A. Shiklomanov, D. Yang, K. Saito, et al. 2021. "Future Trajectory of Arctic System Evolution." In Arctic Hydrology, Permafrost and Ecosystems, edited by D. Yang and D. L. Kane, 893–914. Cham: Springer.
- Salen-Picard, C., D. Arlhac, and E. Alliot. 2003. "Responses of a Mediterranean Soft Bottom Community to Short-Term (1993–1996) Hydrological Changes in the Rhone River." *Marine Environmental Research* 55: 409–27.
- Salen-Picard, C., A. M. Darnaude, D. Arlhac, and M. L. Harmelin-Vivien. 2002. "Fluctuations of Macrobenthic Populations: A Link between Climate-Driven River Run-Off and Sole Fishery Yields in the Gulf of Lions." *Oecologia* 133: 380–8.
- Sampaio, L., A. M. Rodrigues, and V. Quintino. 2010. "Carbon and Nitrogen Stable Isotopes in Coastal Benthic Populations under Multiple Organic Enrichment Sources." *Marine Pollution Bulletin* 60: 1790–802.
- Sarà, G., M. De Pirro, C. Romano, P. Rumolo, M. Sprovieri, and A. Mazzola. 2007. "Sources of Organic Matter for Intertidal Consumers on Ascophyllum-Shores (SW Iceland): A Multi-Stable Isotope Approach." *Helgoland Marine Research* 61: 297–302.
- Sarker, S., M. Masud-Ul-Alam, M. S. Hossain, S. Rahman Chowdhury, and S. M. Sharifuzzaman. 2021. "A Review of Bioturbation and Sediment Organic Geochemistry in Mangroves." *Geological Journal* 56: 2439–50.
- Sarmiento, J. L., and N. Gruber. 2013. Ocean Biogeochemical Dynamics. Princeton, NJ: Princeton University Press.
- Schell, D. M. 1983. "Carbon-13 and Carbon-14 Abundances in Alaskan Aquatic Organisms: Delayed Production from Peat in Arctic Food Webs." *Science* 219: 1068–71.

- Shilla, D., and J. Routh. 2017. "Using Biochemical and Isotopic Tracers to Characterise Organic Matter Sources and Their Incorporation into Estuarine Food Webs (Rufiji Delta, Tanzania)." Chemistry and Ecology 33: 893–917.
- Simoneit, B. R. T. 2006. "Atmospheric Transport of Terrestrial Organic Matter to the Sea." In *Handbook of Environmental Chemistry*, Volume 2: Reactions and Processes, Part N, edited by O. Hutzinger, 165–208. Berlin: Springer-Verlag.
- Smeaton, C., X. Cui, T. S. Bianchi, A. G. Cage, J. A. Howe, and W. E. N. Austin. 2021. "The Evolution of a Coastal Carbon Store over the Last Millennium." *Quaternary Science Reviews* 266: 107081.
- Smith, R. W., T. S. Bianchi, M. Allison, C. Savage, and V. Galy. 2015. "High Rates of Organic Carbon Burial in Fjord Sediments Globally." *Nature Geoscience* 8: 450–3.
- Snelgrove, P. V. R., K. Soetaert, M. Solan, S. Thrush, C. L. Wei, R. Danovaro, R. W. Fulweiler, et al. 2018. "Global Carbon Cycling on a Heterogeneous Seafloor." *Trends in Ecology and Evolution* 33: 96–105.
- Sobrinho, R. d. L., M. C. Bernardes, C. E. de Rezende, J. H. Kim, S. Schouten, and J. S. S. Damsté. 2021. "A Multiproxy Approach to Characterize the Sedimentation of Organic Carbon in the Amazon Continental Shelf." *Marine Chemistry* 232: 103961.
- Sokołowski, A., A. Szczepańska, P. Richard, M. Kędra, M. Wołowicz, and J. M. Węsławski. 2014. "Trophic Structure of the Macrobenthic Community of Hornsund, Spitsbergen, Based on the Determination of Stable Carbon and Nitrogen Isotopic Signatures." *Polar Biology* 37: 1247–60.
- Solan, M., E. M. Bennett, P. J. Mumby, J. Leyland, and J. A. Godbold. 2020. "Benthic-Based Contributions to Climate Change Mitigation and Adaptation." *Philosophical Transactions of the Royal Society B: Biological Sciences* 375: 20190107.
- Solan, M., E. R. Ward, E. L. White, E. E. Hibberd, C. Cassidy, J. M. Schuster, R. Hale, and J. A. Godbold. 2019. "Worldwide Measurements of Bioturbation Intensity, Ventilation Rate, and the Mixing Depth of Marine Sediments." *Scientific Data* 6: 1–6.
- Stasko, A., B. Bluhm, C. Michel, P. Archambault, A. Majewski, J. Reist, H. Swanson, and M. Power. 2018. "Benthic-Pelagic Trophic Coupling in an Arctic Marine Food Web along Vertical Water Mass and Organic Matter Gradients." *Marine Ecology Progress Series* 594: 1–19.
- Stoler, A. B., and R. A. Relyea. 2020. "Reviewing the Role of Plant Litter Inputs to Forested Wetland Ecosystems: Leafing through the Literature." *Ecological Monographs* 90: e01400.
- Sun, S., E. Schefuß, S. Mulitza, C. M. Chiessi, A. O. Sawakuchi, M. Zabel, P. A. Baker, J. Hefter, and G. Mollenhauer. 2017. "Origin and Processing of Terrestrial Organic Carbon in the Amazon System: Lignin Phenols in River, Shelf, and Fan Sediments." *Biogeosciences* 14: 2495–512.
- Szczepanek, M., M. J. Silberberger, K. Koziorowska-Makuch, E. Nobili, and M. Kędra. 2021. "The Response of Coastal Macrobenthic Food-Web Structure to Seasonal and Regional Variability in Organic Matter Properties." *Ecological Indicators* 132: 108326.
- Tagesson, T., G. Schurgers, S. Horion, P. Ciais, F. Tian, M. Brandt, A. Ahlström, et al. 2020. "Recent Divergence in the Contributions of Tropical and Boreal Forests to the Terrestrial Carbon Sink." *Nature Ecology & Evolution* 4: 202–9.

- Tang, M., and E. Kristensen. 2010. "Associations between Macrobenthos and Invasive Cordgrass, Spartina anglica, in the Danish Wadden Sea." *Helgoland Marine Research* 64: 321–9.
- Tanimura, A., W. Liu, K. Yamada, T. Kishida, and H. Toyohara. 2013. "Animal Cellulases with a Focus on Aquatic Invertebrates." *Fisheries Science* 79: 1–13.
- Topçu, N. E., E. Turgay, R. E. Yardımcı, B. Topaloğlu, A. Yüksek, T. M. Steinum, S. Karataş, et al. 2019. "Impact of Excessive Sedimentation Caused by Anthropogenic Activities on Benthic Suspension Feeders in the Sea of Marmara." Journal of the Marine Biological Association of the United Kingdom 99: 1075–86.
- Trannum, H. C., H. Gundersen, E. Oug, B. Rygg, and K. M. Norderhaug. 2018. "Soft Bottom Benthos and Responses to Climate Variation and Eutrophication in Skagerrak." *Journal* of Sea Research 141: 83–98.
- Turetsky, M. R., B. W. Abbott, M. C. Jones, K. Walter Anthony, D. Olefeldt, E. A. G. Schuur, C. Koven, et al. 2019. "Permafrost Collapse Is Accelerating Carbon Release." *Nature* 569 (7754): 32–4.
- Turnewitsch, R., B. Domeyer, and G. Graf. 2007. "Experimental Evidence for an Effect of Early-Diagenetic Interaction between Labile and Refractory Marine Sedimentary Organic Matter on Nitrogen Dynamics." *Journal of Sea Research* 57: 270–80.
- van de Velde, S. J., S. Hidalgo-Martinez, I. Callebaut, G. Antler, R. K. James, M. Leermakers, and F. J. R. Meysman. 2020.
  "Burrowing Fauna Mediate Alternative Stable States in the Redox Cycling of Salt Marsh Sediments." *Geochimica et Cosmochimica Acta* 276: 31–49.
- Van Hoey, G., K. Guilini, M. Rabaut, M. Vincx, and S. Degraer. 2008. "Ecological Implications of the Presence of the Tube-Building Polychaete Lanice conchilega on Soft-Bottom Benthic Ecosystems." Marine Biology 154: 1009–19.
- van Nugteren, P., L. Moodley, G. J. Brummer, C. H. R. Heip, P. M. J. Herman, and J. J. Middelburg. 2009. "Seafloor Ecosystem Functioning: The Importance of Organic Matter Priming." *Marine Biology* 156: 2277–87.
- Volkenborn, N., C. Meile, L. Polerecky, C. A. Pilditch, A. Norkko, J. Norkko, J. E. Hewitt, S. F. Thrush, D. S. Wethey, and S. A. Woodin. 2012. "Intermittent Bioirrigation and Oxygen Dynamics in Permeable Sediments: An Experimental and Modeling Study of Three Tellinid Bivalves." *Journal of Marine Research* 70: 794–823.
- Volkenborn, N., L. Polerecky, S. I. C. Hedtkamp, J. E. E. Van Beusekom, and D. De Beer. 2007. "Bioturbation and Bioirrigation Extend the Open Exchange Regions in Permeable Sediments." *Limnology and Oceanography* 52: 1898–909.
- Wakeham, S. G., and E. A. Canuel. 2006. "Degradation and Preservation of Organic Matter in Marine Sediments." In *Handbook of Environmental Chemistry*, Volume 2: Reactions and Processes, Part N 295–321. Berlin: Springer.
- Ward, J. E., and S. E. Shumway. 2004. "Separating the Grain from the Chaff: Particle Selection in Suspension- and Deposit-Feeding Bivalves." *Journal of Experimental Marine Biology and Ecology* 300: 83–130.
- Wassmann, P., and K. Olli. 2005. Drainage Basin Nutrient Inputs and Eutrophication: Anintegrated Approach. Norway: University of Tromsø.

- Welsh, D. T. 2003. "It's a Dirty Job but Someone Has to Do It: The Role of Marine Benthic Macrofauna in Organic Matter Turnover and Nutrient Recycling to the Water Column." *Chemistry and Ecology* 19: 321–42.
- Werry, J., and S. Y. Lee. 2005. "Grapsid Crabs Mediate Link between Mangrove Litter Production and Estuarine Planktonic Food Chains." *Marine Ecology Progress Series* 293: 165–76.
- Witt, V., C. Wild, and S. Uthickea. 2012. "Terrestrial Runoff Controls the Bacterial Community Composition of Biofilms along a Water Quality Gradient in the Great Barrier Reef." *Applied and Environmental Microbiology* 78: 7786–91.
- Yakushev, E. V., E. A. Protsenko, J. Bruggeman, P. Wallhead, S. V. Pakhomova, S. K. Yakubov, R. G. J. Bellerby, and R. M. Couture. 2017. "Bottom RedOx Model (BROM v.1.1): A Coupled Benthic-Pelagic Model for Simulation of Water and Sediment Biogeochemistry." *Geoscientific Model Development* 10: 453–82.
- Yool, A., A. P. Martin, T. R. Anderson, B. J. Bett, D. O. B. Jones, and H. A. Ruhl. 2017. "Big in the Benthos: Future Change of Seafloor Community Biomass in a Global, Body Size-Resolved Model." *Global Change Biology* 23: 3554–66.
- Zapata-Hernández, G., J. Sellanes, C. Mayr, and P. Muñoz. 2014. "Benthic Food Web Structure in the Comau Fjord, Chile (~42°S):

Preliminary Assessment Including a Site with Chemosynthetic Activity." *Progress in Oceanography* 129: 149–58.

Zapata-Hernández, G., J. Sellanes, M. Thiel, C. Henríquez, S. Hernández, J. C. C. Fernández, and E. Hajdu. 2016.
"Community Structure and Trophic Ecology of Megabenthic Fauna from the Deep Basins in the Interior Sea of Chiloé, Chile (41–43°S)." Continental Shelf Research 130: 47–67.

#### SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

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