INTRODUCTION

Biogeochemical cycles in marine sediments are driven by aerobic and anaerobic microbial processes (Canfield et al. 2005). Rates of these processes are controlled by a number of key physical, chemical and biological factors, including reactivity and burial of organic matter and solid electron acceptors by accretion and biogenic reworking, as well as supply of dissolved electron acceptors (e.g. $O_2$, $NO_3^-$, and $SO_4^{2-}$) and removal of inhibitory metabolites (e.g. $NH_4^+$ and $H_2S$) by solute diffusion, advection, and bioirrigation (Kristensen et al. 2012, Aller 2014). Some of these factors may appear independent at first glance (e.g. particle burial versus solute transport), but they are in fact all coupled through transport-reaction processes within the sediment matrix.

The burial rate of organic matter into oceanic sediments undisturbed by benthic fauna is typically $<1$ cm yr$^{-1}$ (Van Weering et al. 1987, Schimmelmann et al. 2016). Faunal reworking enhances this process considerably, particularly within the upper 6–10 cm of sediments.

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seds (DeMaster et al. 1985, Boudreau 1998, Teal et al. 2008). The fate of sedimentary organic matter is therefore largely controlled by microbial processes in a subsurface chemical environment, where reactions strongly depend on the delivery of electron acceptors (Meile et al. 2005, van de Velde & Meysman 2016). The ultimate electron acceptor in sediments underlying anoxic water column is O2, but much organic carbon oxidation in reality occurs away from the oxic zone by anaerobic processes using other oxidized compounds (NO3−, Mn(IV), Fe(III), and SO42−) as electron acceptors (Canfield et al. 2005). Although the aptitude to accept electrons and thus the energy output from oxidation processes varies among electron acceptors, their quantitative role for the total carbon oxidation is primarily controlled by their availability.

Most burrow-dwelling macrofauna ventilate their burrows and thereby cause marked bioirrigation, which strongly affects transport conditions within sediments (Kristensen et al. 2012). Thus, bioirrigation may enhance solute transport by up to an order of magnitude relative to molecular diffusion. The actual extent of the enhancement depends primarily on functional traits of the infauna and sediment type (Kristensen 2001, Meysman et al. 2006, Shull et al. 2009). Bioirrigation usually decreases in an exponential fashion with depth in the sediment (Martin & Banta 1992, Meile et al. 2001, Forster et al. 2003) in accordance with the declining depth distribution of benthic fauna (Dauwe et al. 1998, Weissberger et al. 2008).

While dissolved electron acceptors, e.g. NO3− and SO42−, are delivered through diffusive and bioirrigation-driven advective solute transport in the sediment porewater, oxidized metals utilized as electron acceptors are solid in form. Hence, they cannot be transported downward by diffusion, and must be buried as solids through sediment accretion or biogenic reworking (Aller 1990). Alternatively, the reduced Mn2+ and Fe2+ forms can be oxidized within the subsurface sediment by O2 translocated into macrofauna burrows (Kostka et al. 2002). Thus, Mn(IV) and Fe(III) are unimportant electron acceptors in laminated sediments, but they are typically dominant in bioturbated deposition areas with high availability and recycling efficiency (Thamdrup 2000, Vandieken et al. 2006).

Studies of sediment bioirrigation usually estimate pore-water transport from tracer profiles. Conservative tracers such as bromide or tracers with known reaction kinetics such as 222Rn are often used for this purpose (Martin & Banta 1992, Forster et al. 2003, Shull et al. 2009). Also, profiles of reactive dissolved compounds such as CO2 and NH4+ have been used to assess bioirrigation given that the depth distribution of reaction rates is known (Meile et al. 2005). Several transport-reaction models can be applied for this purpose (Boudreau 1997). Bioirrigation can be modeled as enhanced diffusion, but non-local exchange describes the process better when burrow ventilation moves solutes rapidly over long distances within the sediment and/or induces advective flow or radial diffusion across burrow walls (Kristensen & Hansen 1999, Meysman et al. 2006, Shull et al. 2009).

The objectives of this study were to assess microbial carbon oxidation and macrofaunal bioirrigation in the upper 20 cm of the sediment along a Skagerrak−Kattegat−Belt Sea depth transect. Emphasis was on (1) the role of bioirrigation by macrofauna communities for the distribution of reactions and solutes in the sediment and (2) the partitioning of electron acceptors (O2, NO3−, Mn(IV), Fe(III), and SO42−) for carbon oxidation. The examined area is well-studied with respect to sediment biogeochemistry (e.g. Canfield et al. 1993a,b, Thamdrup et al. 1994, Rysgaard et al. 2001) and benthic fauna distribution (e.g. Rosenberg 1995, Rosenberg et al. 1996, Dauwe et al. 1998) through a number of cruises over the years. However, no studies have yet determined in detail the magnitude and biogeochemical impact of bioirrigation caused by macrobenthic burrow ventilation.

MATERIALS AND METHODS

Study area and fauna sampling

Samplings were done in late August and early September 2014 on board the R/V Aurora at 4 stations in the Skagerrak–Kattegat–Belt Sea area (Fig. 1). Stations AU1 (58° 6.21′ N, 9° 49.35′ E) and AU2 (58° 2.96′ N, 9° 7.46′ E) were located at 586 and 318 m water depth, respectively, in areas of Skagerrak where the sediment is known to be rich in iron and manganese (Canfield et al. 1993a,b). AU3 (57° 48.39′ N, 11° 3.19′ E) was located at 45 m water depth in a deposition area of the northern Kattegat where the sediment is affected by intensive bioturbation, mostly due to brittle stars (Rosenberg 1995, Jensen et al. 2003); whereas AU4 (55° 0.26′ N, 10° 6.49′ E) was located at 38 m water depth in an area of the southern Little Belt where benthic fauna is suppressed by frequent and recent O2 deficiency in the bottom water (HELCOM 2003).

All sediment from 3 box cores (40 × 40 × 60 cm) sampled at each station was sieved through a 1 mm mesh on deck for fauna quantification. The material
retained from each box core was preserved in 4% formaldehyde on board, and later in the laboratory sorted carefully to separate preserved benthic animals from organic and inorganic debris. Recovered animals were stored in 70% ethanol for later identification to lowest possible taxon and for counting. The fauna was also classified into functional groups according to mobility and feeding behavior. Unfortunately, no biomass was determined due to loss of samples during a major laboratory renovation.

Flux measurements

Four box cores were sampled at each station for on-deck flux incubations and vertical sediment profiling. One sediment sub-core was taken with an acrylic glass core liner (33 cm long and 8 cm diameter) to about 20 cm depth from each box core; leaving space for ~10 cm of overlying water. The cores were inspected for traces of bioirrigation (i.e. excessive depth distribution of oxidized sediment), and placed with open tops in a 50 l tank filled with bottom water from the respective sampling sites. Temperature, salinity and O2 were maintained at levels similar to those recorded near the bottom by the ship’s CTD (Table 1). Use of bottom water assured the correct salinity, while in situ temperature was kept (±1°C) by thermostatted cooling. Since O2 in the bottom water at AU1–3 was close to air saturation, no special efforts were necessary to adjust the O2 level. The tank was in these cases simply stirred by aeration with compressed air. Aeration for AU4 was performed with a N2/air mixture to obtain an O2 level close to 33% of air saturation, as measured in the bottom water of this site. The cores were left in darkness to rest for 4 h before initiating flux incubations. Water circulation inside the core liners was assured by an external rotating magnet (60 rpm) driving internal magnetic stirring bars placed in the overlying water phase of each core.

Fluxes of O2 (total oxygen uptake, TOU), dissolved inorganic carbon (DIC: CO2 + HCO3− + CO32−), NH4+, and NO3− across the sediment–water interface were determined from 18 h (AU1), 13 h (AU2), 9 h (AU3) and 2 h (AU4) dark incubations. The cores were sealed with acrylic glass lids during flux incubations while maintaining water stirring. O2 was analyzed at the start and end by a fiber-optic O2 dipping probe connected to a Microx 4 (PreSens) oxygen meter through an otherwise sealed port in the lids. Initial and final water samples were taken through the port in the lids as well. Subsamples for DIC were preserved with 50 μl of saturated HgCl2 in 5 ml glass Exetainers (Labco) and stored at 5°C until analysis by the flow injection/diffusion cell technique (Hall & Aller 1992). Subsamples for NH4+ and NO3− were GF/F filtered and stored frozen (−20°C) in 20 ml plastic vials until analysis on a Quickchem 8500 Flow Injection Analyzer (Lachat Instruments) according to the protocols of Bower & Holm-Hansen (1980) for NH4+ and Armstrong et al. (1967) for NO3−. Fluxes were calculated from the difference between initial and final water concentrations.

Porewater and solid phase profiles

When flux measurements were terminated, Br− (NaBr) was added to the tank water at a concentration of ~10 mM for determination of bioirrigation in...
flux cores. Cores were incubated in the Br⁻ enriched water with open tops for 1 to 2 d before porewater extraction and solid phase analyses. Cores were sliced into 1 cm intervals to 4 cm depth followed by 2 cm intervals to 18 cm depth (slices 6–8, 10–12 and 14–16 cm were discarded). Porewater was extracted by centrifuging half of each slice in 50 ml centrifuge tubes at 1200 × g for 10 min. The supernatant was GF/F filtered and separated into subsamples. Of these, 1 ml was stored in Eppendorf tubes and preserved with 10 μl saturated HgCl₂ for later DIC analysis as described above, 1 ml was stored at 5°C for later Br⁻, Cl⁻ and SO₄²⁻ analysis by ion chromatography (ICS-2000, Dionex) and the rest (>3 ml) was stored frozen for later NH₄⁺ analysis as described above.

Supplementary cores were sampled using a 100 cm long Rumohr corer (8 cm diameter) to test if 2 d on-board handling of flux cores before porewater extraction impacted the results. Porewater from Rumohr cores was extracted immediately after sampling by inserting Rhizons (Rhizosphere Research) into pre-drilled holes at 10 cm intervals in the upper 30 cm of the core-liner. The extracted porewater was analyzed for DIC, SO₄²⁻ and NH₄⁺ as described above.

Solid phase parameters were determined on sediment subsamples from core sectioning. Wet density of sediment from each slice was measured as the wet weight (ww) of a known sediment volume. Subsequently, subsamples were analyzed for water content by drying at 100°C for 24 h. The dried sediments were acidified with HCl fumes to remove inorganic carbon and analyzed for organic C and N content with an elemental analyzer (Flash EA 2000 Series, Thermo Analytical). Reactive solid phase Fe and Mn were obtained by extracting 100–300 mg fresh sediment subsamples in 5 ml of 0.5 M HCl for 30 min under frequent shaking followed by centrifugation at 1200 × g for 10 min (Lovley & Phillips 1987). The supernatant was stored at 5°C until analysis for Fe(II) and total Fe by transferring 50 μl subsamples to 2 ml of 0.02 % Ferrozine solution without and with 1% of the reducing agent hydroxylamine (10 g l⁻¹), respectively. All solutions were analyzed spectrophotometrically at 562 nm, and reactive amorphous Fe(III) was operationally defined as the difference between total extractable Fe and Fe(II). Mn²⁺ in the extracts was analyzed by flame atomic absorption spectrometry (AAnalyst 100, Perkin Elmer).

NH₄⁺ adsorption coefficients (K_{NH4}⁰) were determined according to Holmboe & Kristensen (2002). Briefly, a series of sediment slurries (~6 g sediment mixed with 30 ml seawater), with NH₄⁺ adjusted to about 0, 1, 2 and 4 mM for each sediment type, were incubated in the dark for 2 d before centrifugation at 1200 × g for 10 min. The supernatant was GF/F filtered and stored frozen (~20°C) until NH₄⁺ analysis as previously described. The centrifuged sediment was homogenized, and exchangeable NH₄⁺ was extracted by transferring 2 g sediment subsamples to 5 ml of 2 M KCl for 30 min at 5°C followed by centrifugation at 1200 × g for 10 min. The supernatant was GF/F filtered and stored frozen (~20°C) until NH₄⁺ analysis as described above. The dimensionless linear adsorption coefficient, K_{NH4}⁰ was calculated according to Krom & Berner (1980), where K_{NH4} is the slope from a plot of extracted NH₄⁺ vs. porewater NH₄⁺.

**O₂ microprofiles**

Vertical microprofiles of O₂ were measured on subcores taken from box cores using polycarbonate core liners (25 cm long and 5 cm diameter). Cores were stored on deck at in situ O₂ and temperature for 1–5 h before measurements. Since preliminary profiles from AU4 showed O₂ depletion at the sediment–water interface, no further O₂ microprofiles were made at this station. Microprofiles at the other stations were measured using custom-built Clarke-type microelectrodes with internal reference and guard (Revsbech 1989). Two to 4 profiles were measured on each of 2 to 3 cores from each station.

**Anoxic incubations**

Vertical profiles of anaerobic carbon oxidation (Cₐ), sulfate reduction (SR), manganese reduction (MnR), iron reduction (FeR), and ammonification (Nₐ) were quantified by closed anoxic incubations. Sediment remaining from the depths 0–2, 2–4, 8–10 and 16–18 cm after slicing the 4 flux cores at each station was immediately used for the anoxic incubations. The sediment slices from each depth interval were pooled (a total of ~200 ml), homogenized and transferred in air into eight 20 ml glass scintillation vials (jars). Rapid handling (few minutes) assured limited oxidation of the sediment. The jars were then sealed gas-tight without headspace to prevent O₂ intrusion, and incubated in darkness at in situ temperatures (see Table 1).

Two jars from each depth were sacrificed at 7 d (AU4) to 20 d (AU1) intervals for determination of changes in porewater DIC, SO₄²⁻, and NH₄⁺ concentrations, as well as solid phase Mn, Fe(II), and Fe(III). Porewater was extracted by centrifuging the jars at
1200 \times g$ for 10 min. The GF/F filtered porewater subsamples were handled and analyzed for DIC, $\text{SO}_4^{2-}$ and $\text{NH}_4^+$ as described above. The sediment remaining after porewater extraction was analyzed for reactive Mn, Fe(II) and Fe(III) as described above. The majority of extracted Mn was assumed to be reactive Mn(IV) available for microbial reduction (Kristiansen et al. 2002). Reaction rates ($\text{Cox}$, SRR, and $\text{Nmin}$) in nmol cm$^{-3}$ d$^{-1}$ were calculated from a linear fit of concentration changes in the time series of samples. $\text{Nmin}$ was corrected for adsorption, using the appropriate $K_{\text{eqal}}$ values as described above. MnR and FeR were determined as solid-phase reactive Mn(IV) and Fe(III) depletion over time.

**Data analysis**

Pielou’s evenness, based on the Shannon diversity index, was used to describe how numerically equal the species in the benthic fauna communities were at the examined sampling stations (Heip et al. 1998):

\[
\text{Shannon diversity: } H' = -\sum_{i=1}^{R} p_i \ln p_i \quad (1)
\]

\[
\text{Pielou’s evenness: } J' = \frac{H'}{H_{\text{max}}} = \frac{H'}{\ln R} \quad (2)
\]

where $p_i$ is the proportional abundance of species $i$, and $R$ is the species richness.

Bioirrigation in flux cores on board the ship was determined from vertical $\text{Br}^-$ profiles by a 1D diffusion model using transient transport equations (Martin & Banta 1992, Forster et al. 2003) with exponentially decreasing non-local transport in the bioturbated zone (zone 1: $0 \leq x \leq L_1$):

\[
\frac{\partial C_1}{\partial t} = D_1 \frac{\partial^2 C_1}{\partial x^2} - \alpha_0 e^{-bx} (C_1 - C_0) \quad (3)
\]

and diffusive transport below this depth (zone 2: $L_1 \leq x \leq L_2$):

\[
\frac{\partial C_2}{\partial t} = D_1 \frac{\partial^2 C_2}{\partial x^2} \quad (4)
\]

where $C_0$ is the concentration of $\text{Br}^-$ in the overlying water (mM); $C_1$ and $C_2$ are the $\text{Br}^-$ concentrations in the porewater (mM) at depth $x$ of zone 1 and zone 2; $L_1$ is the depth of the bioturbated zone (cm) and $L_2$ is the depth of total sediment column (20 cm); $D_1$ is the sediment diffusion coefficient (cm$^2$ d$^{-1}$) after correction of molecular diffusion for porosity (Li & Gregory 1974); $\alpha_0$ is the non-local transport coefficient (d$^{-1}$) at $x = 0$; and $b$ is the attenuation coefficient (cm$^{-1}$) of non-local exchange within the bioturbated zone. The initial and boundary conditions are:

\[
t = 0, \ 0 < x < L_2, \ C_1 = C_2 = C_{\text{initial}}
\]

\[
t > 0, \ x = 0, \ C_1 = C_0
\]

\[
t > 0, \ x = L_1, \ \frac{\partial C_1}{\partial x} = \frac{\partial C_2}{\partial x} \text{ and } C_1 = C_2
\]

\[
t > 0, \ x = L_2, \ \frac{\partial C_2}{\partial x} = 0
\]

where $C_{\text{initial}}$ is the background porewater $\text{Br}^-$ concentration before any transport occurs. The equations were solved in MATLAB using a finite element code by first applying a Galerkin approximation in space (Skeel & Berzins 1990) and then time integration with a variable-step, variable-order solver based on numerical differentiation formulas of orders 1 to 5 (Shampine & Reichelt 1997, Shampine et al. 1999).

Finally, the constants $\alpha_0$, $b$ and $L_1$ were determined by fitting, using a trust-region-reflective least-squares algorithm (Coleman & Li 1994, 1996) to minimize the sum of square errors for the measured pairs.

A double-exponential decay model was used to describe the depth-dependent reaction pattern determined by the anoxic jar incubations. The total reactive organic pool ($G_T$) in the sediment was assumed to be divided into 2 decomposable fractions. Accordingly, a 2-$G$ depth-dependent approach ($\frac{dG_T}{dx}$, Westrich & Berner 1984) should be capable of describing the diagenesis. Each fraction diminishes with depth in the sediment at a rate defined by the product of its concentration and first-order depth attenuation constant (Boudreau 1997). The total organic matter degradation is then equal to the sum of the 2-$G$ depth-dependent rates according to:

\[
\frac{dG_T}{dx} = -a_1 G_1 - a_2 G_2 \quad (nmol \ cm^{-3} \ cm^{-1}) \quad (5)
\]

where $a_1$ and $a_2$ are the down-core attenuation constants (cm$^{-1}$), and $x$ is depth in the sediment (cm). Pool sizes ($G$) can be substituted with reaction rates ($R = \frac{dG}{dt}$) by transforming Eq. (5) into:

\[
R(x) = R_1 e^{-a_1 x} + R_2 e^{-a_2 x} \quad (nmol \ cm^{-3} \ d^{-1}) \quad (6)
\]

where $R(x)$ is the total reaction rate at depth $x$, $R_1$ is the rate of the fast pool at depth $x = 0$, and $R_2$ is the rate of the slow pool at depth $x = 0$. The down-core attenuation constant is in reality the first-order decay constant (d$^{-1}$) divided by the burial velocity, or sedimentation rate ($\omega$; cm d$^{-1}$). Diagenetic reactions involving DIC, $\text{SO}_4^{2-}$ and $\text{NH}_4^+$ derived from the anoxic
incubations were fitted to Eq. (6) to obtain the depth attenuation of $C_{\text{ox}}$, $SR$, $N_{\text{min}}$ in sediments at the selected stations along the Skagerrak–Kattegat–Belt Sea depth transect.

Solute concentrations or generated in sediments by the 2-G depth-dependent diagenetic reactions (e.g. DIC, $\text{SO}_4^{2-}$, and $\text{NH}_4^+$) are subject to transport by molecular diffusion, advection, and mixing processes caused by physical or biogenic activity. By assuming steady state and ignoring sedimentation, the concentration of any solute, $C(x,t)$, in the porewater (mM) can be described by the classic 1D diffusion-reaction model of Berner (1980). However, in the presence of bioturbating benthos, the model must be modified to account for the enhanced, but with depth exponentially decreasing, bioturbation transport induced by fauna ventilation. A transport-reaction model with a non-local exchange module for use in bioirrigated sediments was applied here according to the 2-layer formalism of Aller (1982).

For $0 < x < L_1$:

$$\frac{\partial C_1}{\partial t} = \frac{D_s}{(1+K)} \frac{\partial^2 C_1}{\partial x^2} - \alpha_0 e^{-bx} (C_1 - C_0) + R_1 e^{-a_1 x} + R_2 e^{-a_2 x} = 0$$

For $L_1 < x < \infty$:

$$\frac{\partial C_2}{\partial t} = \frac{D_s}{(1+K)} \frac{\partial^2 C_2}{\partial x^2} + R_1 e^{-a_1 x} + R_2 e^{-a_2 x} = 0$$

The boundary conditions are:

$$x = 0, \quad C = C_0$$

$$x = L_1, \quad \frac{\partial C_1}{\partial x} = \frac{\partial C_2}{\partial x} \quad \text{and} \quad C_1 = C_2$$

$$x = \infty, \quad \frac{\partial C_2}{\partial x} = 0$$

The constants $D_s$ (sediment diffusion coefficient), $K$ (dimensionless adsorption coefficient), $C_0$ (overlying water concentration), $R_1$, $R_2$, $a_1$, $a_2$ (reaction rates and attenuation coefficients), and $L_1$ (visually detected depth of bioirrigation) were derived from the above analyses and modelling, while the unknown constants $\alpha_0$ (non-local transport coefficient at the sediment-water interface) and $b$ (depth attenuation constant for non-local exchange) were determined from best fit of the model to measured porewater profiles. The differential equations were solved in MATLAB with a finite difference code that implemented the 3-stage Lobatto IIIa formula (Kierzenka & Shampine 2001). The constants $\alpha_0$ and $b$ were then determined by fitting, using a trust-region-reflective least-squares algorithm (Coleman & Li 1994, 1996) to minimize the sum of square errors for the measured pairs.

### RESULTS

#### Fauna composition

The deep Skagerrak stations (AU1–2) had less than half the abundance of benthic fauna than the roughly 2100 ind. m$^{-2}$ found at the shallower Kattegat station (AU3) (Table 2). The Belt station (AU4), on the other hand, was azoic due to recent episodes of bottom-water $O_2$ deficiency. The species richness ranged from 17 at AU1 to 28 at AU2. Pielou’s evenness was higher at AU2 than AU1 and AU3 due to distinct dominance of 1 ($\text{Spiochaetopterus typicus}$, $>50\%$) or 2 ($\text{Amphiura filiformis}$, $>50\%$, and $\text{Kurtiella bidens}$, $>30\%$) species, respectively, at the latter stations. The partitioning of taxa varied considerably, with dominance of polychaetes ($>60\%$) at AU1, molluscs ($<80\%$) at AU2, and echinoderms ($>50\%$) at AU3. The predominant species also governed the partitioning of feeding types, as for example the subsurface deposit-feeding $\text{Thysanura equalis}$ and $\text{Yoldiella lucida}$ almost exclusively accounted for the $>60\%$ occurrence of this feeding type at AU2. Similarly, the dominant species at AU1 and AU3 were responsible for the high level of suspension-feeding ($>60$ and $>90\%$, respectively) at these stations. Conversely, neither the taxa nor the feeding types seem to influence the bioturbation types. Gallery- or tube-forming species were most common ($>60\%$) at AU1, whereas surficial burrowers dominated at AU2 ($>80\%$) and AU3 ($<100\%$). It should be noted, however, that the deep-burrowing and gallery-forming ghost shrimp $\text{Callianassa tyrreni}$ was present at AU3 with a minimum density ($\pm SD)$ of 16 ± 4 ind. m$^{-2}$.

The unfortunate lack of biomass determination in the present study was compensated using literature values from the same area. Rosenberg (1995) and Rosenberg et al. (1996) quantified the benthic fauna at 23 stations in Skagerrak from 69 to 682 m water depth. Based on their data on abundance and biomass, the average individual biomass decreases with water depth according to the following equation:

$$W = 89.12 \times e^{-3.75 \times d}, \quad r^2 = 0.573$$

where $W$ is weight (mg ww) and $d$ is water depth (m). Accordingly, the individual biomass at AU1–3 was estimated to 10, 27 and 75 mg ww, respectively, resulting in a total biomass increase with decreasing depth from 8 to 159 g ww m$^{-2}$ at the 3 sampling stations (Table 2).
Table 2. Abundance (± SD), biomass and species richness (± SD) as well as Pielou's evenness (J') for benthic fauna at the 4 sampling stations along the Skagerrak–Kattegat–Belt Sea depth transect. The abundance is partitioned into: (1) Major taxonomic groups (Polychaeta, Mollusca, Crustacea and Echinodermata) with indications of the 2 most dominant species. (2) Feeding types (SF: suspension feeder; SDF: surface deposit-feeder; SSDF: subsurface deposit-feeder; P: predator). (3) Bioturbation types (SB: surficial burrower [<5 cm]; GB: gallery/tube burrower [>5 cm]). WW: wet face deposit-feeder; SSDF: subsurface deposit-feeder; P: predator).

Table 3. Sediment characteristics at the 4 stations along the examined Skagerrak–Kattegat–Belt Sea depth transect in late August and early September 2014. Values for 3 depth intervals are given. POC: particulate organic carbon; dw: dry weight; –: data not available.

Solid phase sediment profiles

The sediments at AU1 and AU2 had similar appearance and consisted of primarily silt with a wet density of ~1.3 g cm\(^{-3}\) (Table 3). Both sediments had a distinct brownish oxidized zone in the upper 5–10 cm that gradually faded 10–15 cm downcore and extended deepest at AU1. Sediment from AU1 appeared slightly muddier than that from AU2 as indicated by the higher porosity and organic content. AU3 sediment consisted mostly of fine sand with a density of 1.6–1.7 g cm\(^{-3}\). An oxidized zone extended to 5–6 cm depth at this station with occasional oxidized traces, i.e. burrows of *C. tyrhena*, down to at least 20 cm depth in the sediment. Porosity and organic content of the sediment decreased with depth at all stations, primarily in the upper 6 cm. Lowest particulate organic carbon (POC) content was observed at AU3, approaching about half the level measured at AU1 and AU2. The sediment at AU4 was soft and organic-rich mud with a density of 1.1 g cm\(^{-3}\). It appeared completely anoxic and sulfidic, with no oxidized sediment except for a thin (<1 mm) brown film at the surface. The organic C:N ratio was slightly lower at AU2 (11–12) than at AU1 and AU3 (13–14) and higher than at AU4 (9–10). NH\(_4^+\) adsorption coefficients (\(K_{NH4}\)) were close to 1 at all stations and sediment depths, except for markedly lower values in the upper 2 cm.

Solid-phase Fe was generally dominated by high levels of Fe(II), increasing with depth in the sediment at all stations, reaching ~110, ~60, ~80 and ~90 μmol cm\(^{-3}\) at AU1–4, respectively (Fig. 2). Fe(III) only surpassed Fe(II) in the upper strongly oxidized 5 cm at AU1 (up to 80 μmol cm\(^{-3}\)), where Fe(II) was close to 0. Below this depth, Fe(III) decreased to around...
and below 10 μmol cm−3. The near-surface level of Fe(II) and Fe(III) was similar at AU2 and AU3 (15–20 and 25–30 μmol cm−3, respectively). Fe(III) decreased rapidly within the upper 5 cm at these stations to trace levels at AU2, and 5–10 μmol cm−3 at AU3. The anoxic AU4 only showed traces of Fe(III). Reactive solid Mn was high at AU1, showing an increase with depth from ~10 to ~32 μmol cm−3 in the upper 2–3 cm followed by a decrease to ~20 μmol cm−3 from 6 to 10 cm depth (Fig. 3). The other stations had only traces of solid Mn (2–4 μmol cm−3) with no noticeable depth pattern.

**Anaerobic reaction rates and sediment-water fluxes**

Reaction rates (Cox, SR, and Nmin) obtained from the anoxic jar incubations varied among stations and generally decreased strongly with depth in the sediment (Fig. 4). Cox and SR near the sediment surface at the shallow AU3 and AU4 were an order of magnitude higher than at AU1 and ~2 times higher than at AU2. The declining anaerobic rates with depth in the upper 18 cm of the sediment fitted well at all stations to a depth-dependent double exponential decay pattern according to Eq. (6) (Table 4). Down-core attenuation of Cox, SR and Nmin was generally higher at the more reactive shallow than the deep stations with rates approaching almost the same level below 15 cm depth at all stations. MnR and FeR behaved differently, with highest MnR at AU1 and highest FeR at AU3 (Fig. 5), followed by a strong decrease with sediment depth from about 25 and 140 nmol cm−3 d−1, respectively, near the surface to <5 and <30 nmol cm−3 d−1 at 18 cm depth. MnR at AU2 was irregular and always below 10 nmol cm−3 d−1, while MnR was almost 0 at AU3 and AU4. FeR showed similar depth attenuation at AU1 and AU2 with a level about 50 nmol cm−3 d−1 lower than AU3 at all depths. FeR was almost 0 throughout the sediment column at the anoxic AU4.

Area-specific reaction rates in the depth zone of interest were estimated by integrating Eq. (6) to 20 cm depth (Table 4). Cox decreased almost 10-fold with increasing water depth from AU4 to AU1, while SR decreased almost 30-fold within the same depth range. The Cox:SR ratio around 2 at AU4 indicates that carbon oxidation at that station was dominated by SR. The high Cox:SR ratios at the other stations, particularly that of 6.9 for AU1, indicate that processes other than SR contributed to DIC production. Accordingly, the range of FeR was 4–15 mmol m−2 d−1 from AU1 to AU3, while MnR was almost 2 and 1 mmol m−2 d−1 at AU1 and AU2, respectively. Almost no FeR and MnR were detected at AU4. Nmin was lowest at AU1, with area-specific rates reaching ~20% of those at AU2 and AU4, and only 10% of that at AU3 (Table 4). The Cox:Nmin ratios were close to the Redfield ratio with values ranging from 5 to 13.5.

Solute fluxes also reflected the water depth, showing an order of magnitude lower TOU and DIC flux...
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values in deep than in shallow water (Table 5). The respiratory quotient, RQ (ratio between DIC efflux and TOU), was highest at AU3 and lowest at AU2 and AU4 with a range of 1.4 to 2. The DIC fluxes at AU1, AU3 and AU4 (Table 5) were about 3 times higher than the depth integrated jar rates (Table 4), whereas the flux at AU2 was 20% higher than the corresponding jar rate. This discrepancy indicates that DIC efflux at AU2 for some reason may be underestimated. A slight NH₄⁺ uptake was evident at AU1 and AU2, while AU3 and AU4 showed a 20 to 30-fold higher release. NO₃⁻ fluxes were low and erratic at AU1–3, while AU4 had a high uptake. NH₄⁺ + NO₃⁻ effluxes (negative for AU2) were lower than the depth-integrated NH₄⁺ production (N_{min}) in jars at AU1–3, while AU4 exhibited almost 3 times higher efflux than depth integrated rates.

### Table 4. Parameters obtained by fitting down-core reactivity of carbon oxidation (Cox), sulfate reduction (SR), and ammonification (N_{min}) to Eq. (6). The 0–20 cm depth integrated reactions (mmol m⁻² d⁻¹) are indicated. The Cox:N_{min} and Cox:SR ratios are shown for comparison. R₁ and R₂ are given as nmol cm⁻³ d⁻¹; a₁ and a₂ are given as cm⁻¹.

<table>
<thead>
<tr>
<th></th>
<th>AU1</th>
<th>AU2</th>
<th>AU3</th>
<th>AU4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cox (r²)</td>
<td>1.00</td>
<td>0.997</td>
<td>0.999</td>
<td>0.965</td>
</tr>
<tr>
<td>R₁</td>
<td>31.16</td>
<td>76.90</td>
<td>629.15</td>
<td>272.18</td>
</tr>
<tr>
<td>a₁</td>
<td>1.816</td>
<td>0.336</td>
<td>1.339</td>
<td>0.552</td>
</tr>
<tr>
<td>R₂</td>
<td>18.85</td>
<td>27.44</td>
<td>53.07</td>
<td>102.74</td>
</tr>
<tr>
<td>a₂</td>
<td>0.078</td>
<td>0.046</td>
<td>0.069</td>
<td>0.087</td>
</tr>
<tr>
<td>0–20 cm</td>
<td>1.65</td>
<td>5.21</td>
<td>10.12</td>
<td>14.67</td>
</tr>
</tbody>
</table>

| SR (r²) | 0.969 | 1.000 | 0.999 | 0.983 |
| R₁     | 3.18  | 45.36 | 58.44 | 50.69 |
| a₁     | 0.355 | 0.478 | 0.165 | 0.202 |
| R₂     | 1.25  | 7.20  | 8.18  | 37.10 |
| a₂     | 0.020 | 0.029 | 0.054 | 0.062 |
| 0–20 cm | 0.240 | 1.722 | 4.378 | 6.717 |

| Cox:SR | 6.88 | 3.02 | 2.31 | 2.18 |
| FeR     |      |      |      |      |
| 0–20 cm | 4.24 | 6.84 | 14.65| 0.31 |

| MnR     |      |      |      |      |
| 0–20 cm | 1.73 | 1.12 | 0    | 0    |

| N_{min} (r²) | 0.987 | 0.994 | 0.998 | 0.996 |
| R₁     | 3.56  | 19.16 | 29.74 | 35.63 |
| a₁     | 1.096 | 0.368 | 0.514 | 0.910 |
| R₂     | 2.04  | 3.68  | 14.76 | 6.69  |
| a₂     | 0.094 | 0.061 | 0.082 | 0.074 |
| 0–20 cm | 0.216 | 0.945 | 2.029 | 1.090 |

| Cox:N_{min} | 7.64 | 5.51 | 4.99 | 13.46 |

Fig. 4. Depth-dependence of (a) anaerobic Cox (dissolved organic carbon [DIC] production), (b) SR (sulfate reduction), and (c) N_{min} (NH₄⁺ production) in the sediment at the 4 stations (AU1–4). Symbols indicate the calculated rates (slope ± SD, n = 4) from anaerobic incubations and curves represent the best fit to the double-exponential model shown in Eq. (4).

Fig. 5. Depth-dependence of (a) MnR (manganese reduction), and (b) FeR (iron reduction) in the sediment at the 4 stations (AU1–4). Symbols indicate the calculated rates (slope ± SD, n = 4) from anaerobic incubations.
Vertical solute profiles and non-local exchange

O₂ profiles at AU1–3 decreased steeply in a parabolic fashion with depth in the sediment (Fig. 6). The depth penetrations of 15–18 mm at AU1, ~8 mm at AU2 and ~4 mm at AU3 were inversely related to sediment reactivity. The profiles were smoother and more regular at AU1 and AU2 than at AU3 due to the near-surface activity of *Amphiura filiformis* at the latter station. The diffusive O₂ uptake (DOU) at AU1−3 corresponded to 62, 47 and 59%, respectively, of the TOU at these stations.

Porewater profiles of DIC, SO₄²⁻ and NH₄⁺ (Fig. 7) reflected the balance between reactions and transport. The porewater data from flux cores and Rumohr cores agreed well and fitted excellently to Eqs. (7) & (8), indicating that the on-board handling of flux cores had only limited impact on porewater solutes. The lack of any fauna and bioirrigation at AU4 was clearly evident from the perfect fit to all solutes using molecular diffusion as the only transport process. Curve fitting of the data from AU1−3, however, required the consideration of bioirrigation in the form of non-local exchange. AU2 and AU3 had almost identical surface intercepts of non-local exchange, with α₀ of 0.24 to 0.34 d⁻¹ and 0.22 to 0.38 d⁻¹, respectively (Fig. 7); the lowest values for both stations were obtained for DIC and highest for NH₄⁺. The depth attenuation (b) of bioirrigation was twice as fast at AU2 than at AU3, resulting in negligible non-local exchange below ~15 cm at the former and below ~30 cm depth at the latter station. The surface intercept of non-local exchange at AU1 attained a level that was only ~15% of those at the other 2 bioturbated stations for all solutes and with attenuation similar to that at AU3. Accordingly, non-local exchange at AU1 was negligible below ~10 cm depth.

Bioirrigation estimates obtained from the on-deck Br⁻ incubations differed strongly from those described above by diagenetic modeling (Figs. 7 & 8). Although the bioirrigation depth limits by the 2 approaches were similar for AU1 and AU2, the surface non-local exchange coefficients (α₀) obtained by the Br⁻ approach were only ¼ of those obtained from the diagenetic model. The opposite was evident for AU3 with 3−4 times higher α from the Br⁻ approach in the upper 5 cm and virtually 0 below 5 cm, while the diagenetic approach revealed significant bioirrigation down to almost 30 cm depth. Nevertheless, depth attenuation (b) of bioirrigation was consistently 20−30 times slower in the bioturbated zone when using Br⁻ compared with diagenetic modeling.

## DISCUSSION

### Controls of fluxes and reaction rates

The 4 Skagerrak–Kattegat–Belt Sea stations examined in this study evidently represent a wide range of environmental conditions and are as such ideal for the comparison of factors controlling solute fluxes and reaction rates. The reactivity of organic matter is clearly impacted by water depth and distance from land, with up to 1 order of magnitude higher fluxes and reaction rates at shallow (AU3 and AU4) than at deep stations (AU1 and AU2). This pattern is obviously related to differences in the amount, origin (Hernes & Benner 2006) and age (Kaiser & Benner 2009) of deposited organic matter. Thus, more reactive fractions of the organic debris disappear in transit when deposited at >200 m compared with...
40–50 m water depth. According to McDonnell et al. (2015), the reactivity of organic particles decreases by 75% due to decay while sinking 350 m in the water column, which is in the same order of magnitude as the presently observed depth attenuation of benthic metabolism (Table 5). The unexpected excess organic matter in the sediment at the deep AU1 and AU2 compared with the shallow AU3 may originate from horizontal transport of poorly reactive organic matter (Dauwe et al. 1998, Ståhl et al. 2004). Accordingly, it has been proposed that the deep Skagerrak is a deposition area for fine particles transported from coastal areas to the south via the Jutland Current (Van Weering et al. 1987, Ståhl et al. 2004, Gyllencreutz & Kissel 2006).

The stable and almost air-saturated oxic bottom water at the non-stratified Skagerrak (AU1 and AU2) and Kattegat (AU3) locations allow an innate balance between aerobic and anaerobic processes, partly driven by an abundant fauna, that maintains sediment biogeochemistry at near steady-state. AU4 in the Little Belt, on the other hand, is obviously unstable with transient state fluxes as it suffers from frequent bottom water O₂ deficiency (HELCOM 2003, Conley et al. 2009), due to high benthic O₂ demand combined with more or less permanent water column stratification (Diaz & Rosenberg 2008, Lehmann et al. 2014).

The double-exponential decay profiles of C₀ₓ, SR and Nₜₙ, recorded for all stations in the present study (Fig. 4) are comparable with those obtained previously in the Kattegat–Skagerrak area (Canfield et al. 1993a, Rysgaard et al. 2001, Jensen et al. 2003), but the 0–20 cm depth-integrated rates of, particularly, C₀ₓ (Table 4) are generally lower than the corresponding DIC fluxes (Table 5). The discrepancy is partly a consequence of excluding the O₂-containing surface layer and benthic fauna by isolation of sediment during anoxic incubations (Hansen & Kristensen 1998, Kristensen et al. 1999). This shortcoming is particularly important at AU1–3, since carbon oxidation may
slow down by up to an order of magnitude when oxic sediment turns anoxic and benthic fauna is eliminated (Kristensen et al. 1995, Kristensen & Holmer 2001). Furthermore, depth integration of Cox, SR and N_{min} to only 20 cm depth may underestimate total sediment metabolism by ignoring reactions in the deep biosphere below, although the majority of carbon oxidation takes place in the upper sediment (Flury et al. 2016). On the other hand, the lower release of NH_4^+ observed from flux measurements at AU1−3 (Table 5) as compared to depth-integrated N_{min} (Table 4) is probably due to coupled nitrification−denitrification (Rysgaard et al. 2001) and anammox (Trimmer et al. 2013) that was absent in the anoxic incubations.

The low DOU estimated from O_2 microprofiles indicates that active benthic fauna in flux cores from AU1−3 is responsible for about half of TOU, which is in accordance with Glud (2008). The direct contribution of benthic fauna respiration to TOU at AU1−3 can be estimated from the fauna biomass using a universal biomass-specific rate of 1.5 to 2.0 μmol O_2 (g ww)^{-1} h^{-1} for benthic fauna at 15°C (Kristensen 1989, Vopel et al. 2003, Schröer et al. 2009). By applying a typical Q_{10} of 2 for benthic fauna (Andresen & Kristensen 2002), the direct fauna respiration at the sampled stations is estimated to account for 4 to 6% at AU1, 7 to 9% at AU2 and 41 to 55% at AU3 of TOU. The fauna contribution is low at AU1 and AU2 compared to published values of 20 to 50% (Christensen et al. 2000, Papaspyrou et al. 2010), while that at AU3 agrees well with the literature. Most of the excess TOU relative to DOU is therefore of fauna-stimulated microbial origin at AU1 and AU2, while there is limited room for a fauna-induced microbial component at AU3 as also reported for this location by Vopel et al. (2003).

Benthic fauna and bioirrigation

Fauna abundance and species richness observed in this study at the deep Skagerrak stations, AU1 and AU2 (Table 2), were similar to, but in the low range of, those found by Josefson (1985) and Rosenberg et al. (1996) for trench and slope locations in the same area. Feeding types among studies were similar with dominance of suspension-feeders near the trench (AU1) and subsurface deposit-feeders at the slope (AU2). Fauna composition at the shallow Kattegat station, AU3, was comparable to the shallow slope examined by Rosenberg (1995). Both studies found dominance of Amphiura sp. and Kurtiella bidentata, but the species richness and total abundance in the present study was in the low range of those found by Rosenberg (1995). Also in this case, the feeding types matched well between the 2 studies with complete dominance of suspension feeders. The absence of benthic fauna at AU4 was clearly caused by bottom water hypoxia (Hansen et al. 2014).

The relatively low macrofaunal biomass and the small mean individual weight in the organic-rich sediments of the deep Skagerrak are probably due to low nutritional value of the depositing organic matter (Dauwe et al. 1998). Rosenberg et al. (1996) observed a vertical discontinuity in species composition and trophic groups at about 400 m water depth (slope and deep basin areas), and attributed this to variations in sedimentation rate and organic matter supply. Subsurface-deposit feeders should therefore dominate below 400 m depth as they are generally adapted to a poor food source (Lopez & Levinton 1987) by high sediment processing rates, selective ingestion of digestible particles and effective absorption of nutrients (Dauwe et al. 1998). Conversely, suspension feeders dominate in shallow areas where frequent
resuspension and deposition events provide ample suspended food (Rosenberg et al. 1996). However, the dominance of the suspension feeder *Spiochaetopterus typicus* at the deep AU1 seems to contradict this general trend. Rosenberg et al. (1996) also noted this apparent conflict and suggested that *S. typicus* under conditions of low particle concentration may switch to surface deposit-feeding, as also reported by Fauchald & Jumars (1979). Dauwe et al. (1998) similarly found that the vertical distribution and trophic structure of macrofauna in the North Sea–Skagerrak area reflected differences in quality of the organic matter. They found that most organisms at shallow water depth feed on freshly deposited or suspended material as interface or suspension feeders, while deeper Skagerrak sediments mainly support small-sized deeply penetrating deposit feeders.

The diagenetic transport-reaction modelling approach (Eqs. 7 & 8) revealed that the 3 faunated stations (AU1–3) all were subject to substantial bioirrigation (Fig. 7). Non-local exchange of the same magnitude and depth attenuation as estimated here have been observed previously in other off-shore sediments (Martin & Banta 1992, Forster et al. 2003, Grigg et al. 2005). The similar profile shape of non-local exchange coefficients (α<sub>c</sub>, α<sub>n</sub>, and α<sub>SR</sub>; see Fig. 7) when estimated from fits of COX, SR and N<sub>min</sub> (Fig. 4) to porewater solute distributions (Fig. 7) confirms the validity of the diagenetic approach despite the above-mentioned critical assumptions regarding the depth-dependent reaction patterns. The slight solute-specific variations in the magnitude of α, with consistently higher values for NH<sub>4</sub><sup>-</sup> than for DIC and SO<sub>4</sub><sup>2-</sup>, are in accordance with the diffusion-dependent predictions of Meile et al. (2005). The differences in non-local exchange observed among stations, on the other hand, were due to the functionality of the benthic fauna. The high near-surface α<sub>n</sub> at AU3 was obviously caused by the high abundance and biomass of *Amphiura filiformis* (Rosenberg 1995), while the deep burrowing *Callianassa tyrrhena* was responsible for the relatively slow depth attenuation of α. AU2 exhibited almost the same α<sub>n</sub> as AU3 due to the near-surface dominance of bivalves with high capacity of ventilation. However, the lack of deeper burrowing species at AU2 caused the rapid attenuation of α with depth in the sediment. The overall low non-local exchange at AU1 was a consequence of low faunal activity due partly to food limitation and partly to low temperature, and dominance of small deep-burrowing and tube-forming polychaetes. These cause limited near-surface bioirrigation, but maintain continued activity deep into the sediment as indicated by the slow depth attenuation.

Although the use of the inert tracer Br<sup>-</sup> is an elegant and easy approach to quantify bioirrigation and non-local exchange in sediments (Martin & Banta 1992, Forster et al. 2003), it was invalid for on-deck cores in the present study. The non-local exchange coefficients obtained from transient transport modelling of Br<sup>-</sup> profiles (Eqs. 3 & 4) after on-deck incubations were either negligible (AU1 and AU2) or very high and confined to the upper 5 cm (AU3) (Fig. 7). In contrast to the diagenetic approach (Eqs. 7 & 8), that is based on the true *in situ* conditions in the sediment before sampling, the Br<sup>-</sup> method relies on bioirrigation performed by the fauna during on-deck incubations using relatively small cores. The true bioirrigation of deep tube- or burrow-dwelling species seems to be underestimated by this method due to incomplete sampling, compression and chopping of burrows and/or animals during the coring procedure (Blomqvist 1991, Eleftheriou & Moore 2005). These damaged conditions cannot be repaired by the benthos within the short time available (hours) before start of on-deck incubations. For example, bioirrigation by *Callianassa tyrrhena* at AU3 was not included as this ghost shrimp inhabits burrows deeper than the coring depth. Furthermore, the benthic fauna may be disturbed by vibrations from the ship during incubations. For example, Solan et al. (2016) found that even underwater broadband sound fields can alter the activity of benthic fauna and affect their fluid and particle transport. Since the exact stress response is species-specific, ship vibrations may have slowed down the activity of the species from AU1 and AU2, while *Amphiura filiformis* from AU3 maintained or even increased its bioirrigation activity only in the top 5 cm. It is therefore not recommended to use the Br<sup>-</sup> tracer approach on newly extracted sediment cores, particularly not onboard a ship. The Br<sup>-</sup> method should be restricted to laboratory incubations on pre-adapted sediment cores or homogenized sediment with a known quantity of benthic fauna added (Quintana et al. 2007). In both cases, several days must pass from core retrieval or establishment to initiation of incubation.

**Electron acceptor availability and the role of bioturbation**

The relative importance of aerobic microbial carbon oxidation, estimated as the difference between DIC flux and the sum of all anaerobic jar reactions,
varies among stations in the Skagerrak–Kattegat area, ranging from 8% at AU2 to 55–60% at AU1 and AU3 of total sediment DIC flux. Although the low aerobic contribution at AU2 may substantiate an underestimate of DIC fluxes at this station, similar variability has been reported for the same area with an average of typically ~50% (Rysgaard et al. 2001). Utilization of electron acceptors driving anaerobic carbon oxidation gradually switches to more oxidized forms with increasing water depth and distance from land in accordance with the decreasing rates of $C_{ox}$ (Table 6). A quite consistent pattern actually emerges when all available literature data on the partitioning of anaerobic electron acceptors in sediments are compiled as a function of water depth in the Skagerrak–Kattegat–Belt Sea area (Fig. 9). Thus, SR dominates (>50%) in shallow water (below 40 m depth), decreases to <50% when moving to deeper water, particularly below 400 m depth, and reaches 0 around 700 m depth. Conversely, FeR is virtually 0 below 40 m water depth and increases to ~50% below this depth and down to about 600 m, but returns to 0 again around 700 m depth. MnR, on the other hand, is close to 0 down to at least 500 m water depth and rapidly increases to complete dominance at 700 m. The contribution of denitrification is generally below 10% at all depths. These depth patterns and proportions are within the range previously reported for continental margin sediments with high organic matter flux (Thamdrup 2000, Canfield et al. 2005). Particularly the dominance of MnR and FeR is a striking feature of deep sediments rich in Mn(IV) and Fe(III) (Aller 1990, Canfield et al. 1993a, Hyun et al. 2017). While the availability of the electron acceptors $\text{NO}_3^-$ and metal oxides (Mn(IV) and Fe(III)) is dependent on the presence of $\text{O}_2$ in the overlying water, SR can be supported via diffusion of $\text{SO}_4^{2-}$ into the sediment from high concentrations in the overlying seawater. Accordingly, the intense carbon oxidation in the very reactive sediments at AU4 underlying a hypoxic water column is completely dominated by SR.

### Table 6. Partitioning of anaerobic electron acceptors at the 4 stations along the examined Skagerrak–Kattegat–Belt Sea depth transect in late August and early September 2014. Values are given as % of total anaerobic carbon oxidation ($C_{ox}$) according to Eq. (6) (Table 4). Denitrification (Denitr) was derived from Rysgaard et al. (2001) for AU1–3 and as the influx of $\text{NO}_3^-$ at AU4, and converted to C-units by multiplication with 4/5. $\text{SO}_4$ reduction (SR) was obtained by integrating Eq. (6) below the oxic zone and to 20 cm depth (Table 4), and converted to C-units by multiplying by 2. Fe reduction (FeR) and Mn reduction (MnR) was obtained by depth-integrating the results in Fig. 6 below the oxic zone and to 20 cm, and converted to C-units by multiplying by 1/4 and 1/2, respectively.

<table>
<thead>
<tr>
<th>Station</th>
<th>MnR</th>
<th>Denitr</th>
<th>FeR</th>
<th>SR</th>
</tr>
</thead>
<tbody>
<tr>
<td>AU1</td>
<td>33.1</td>
<td>8.4</td>
<td>40.3</td>
<td>18.3</td>
</tr>
<tr>
<td>AU2</td>
<td>8.6</td>
<td>12.6</td>
<td>26.2</td>
<td>52.7</td>
</tr>
<tr>
<td>AU3</td>
<td>0</td>
<td>3.0</td>
<td>28.6</td>
<td>68.4</td>
</tr>
<tr>
<td>AU4</td>
<td>0</td>
<td>3.1</td>
<td>5.3</td>
<td>91.5</td>
</tr>
</tbody>
</table>

Fig. 9. Partitioning of anaerobic electron acceptors for carbon oxidation in sediments as a function of depth along the Kattegat–Skagerrak transition zone. Data stem from Canfield et al. (1993a), Rysgaard et al. (2001) and from the present study. Den: denitrification; SR: sulfate reduction; FeR: iron reduction; MnR: manganese reduction.
al. 2013). It is not known to what extent reworking, as suggested by Canfield et al. (1993a), or bioirrigation (Kristensen et al. 2012) contributes to the inventory of these important electron acceptors in the Kattegat and Skagerrak area. It has been argued that the delivery of metal oxides within the upper 10 cm of marine sediment is caused solely by biogenic particle mixing (Aller 1990, Canfield et al. 1993a). Hence, Canfield et al. (1993a) and Hyun et al. (2017) estimated that the measured MnR in Skagerrak and the East Sea (east of Korea) could be supported at a biodeffusion (reworking) coefficient ($D_R$) of 19 and 10 cm$^2$ yr$^{-1}$, respectively. A similar calculation based on the present data reveals a need for $D_R$ at AU1 of ~23 cm$^2$ yr$^{-1}$ that must reach down to about 10 cm as judged from the depth distribution of Mn(IV). However, $D_R$ in sediments at comparable water depths as in the Skagerrak rarely exceeds 5 cm$^2$ y$^{-1}$ and is commonly below 1 cm$^2$ y$^{-1}$ (Middelburg et al. 1997, Lecroart et al. 2010). In fact, $D_R$ at AU1 has recently been measured to <1 cm$^2$ y$^{-1}$ (L. Deng, C. C. Su & M. A. Lever pers. comm.). The discrepancy becomes much worse when FeR is considered at AU2 and AU3 where the required $D_R$ approaches 100 cm$^2$ y$^{-1}$ as also estimated by Canfield et al. (1993a). This greatly exceeds the measured values of 4–17 cm$^2$ y$^{-1}$ (L. Deng, C. C. Su & M. A. Lever pers. comm.). Accordingly, other reoxidation processes such as ventilation-driven downward translocation of O$_2$ are required. Ventilation is several orders of magnitude faster than particle reworking (Aller 1982, Kristensen 2001), and the oxygen brought down into burrows through bioirrigation will readily oxidize Fe$^{2+}$ and Mn$^{2+}$ along the burrow wall and turn the sediment into a mosaic of oxidized burrow walls. At high abundances, these may even merge to form a continuous oxidized layer to 10 cm depth in the sediment (Norkko et al. 2011). Beam et al. (2018) similarly concluded that bioirrigation is the primary driver for rapid oxidation of Fe$^{2+}$ deep in marine sediments. The metal oxides subsequently become available for microbial reduction, for example when the dynamic inhabitants abandon burrows or die. The exact partitioning between MnR and FeR then solely depends on the availability and oxidation of these metals in the sediment.

Concluding remarks

The reactivity of organic matter along the Skagerrak–Kattegat–Belt Sea transect is clearly impacted by water depth and distance from land with up to 1 order of magnitude higher fluxes and reaction rates at shallow than deep areas. The vertical distribution of reactions and partitioning of electron acceptors for carbon oxidation in sediments along this transect is dependent on the balance between the reducing capacity of the sediment due to reactions and downward transport of O$_2$. Thus, the relatively high rates of MnR and FeR in deep water were controlled by rapid reoxidation of metals by enhanced downward translocation of O$_2$ into the low reactive anoxic sediment through ventilation-driven bioirrigation by the infauna.

The present study also concludes that quantification of bioirrigation of newly retrieved sediment cores onboard a ship should preferably be done by diagenetic transport-reaction modelling rather than the widely used Br$^-$ incubation technique. However, it is suggested that future studies look closer into the comparison of methods to estimate bioirrigation to confirm this supposition. Furthermore, there is a strong need for research on the role of individual fauna species and functional groups for bioirrigation and reoxidation of metals in deep sediments. If successful, the obtained results can be valuable in future evaluations and mapping of key biogeochemical mechanisms and conditions in oceanic sediments.

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