

# New insights into large-scale trends of apparent organic matter reactivity in marine sediments and patterns of benthic carbon transformation

Felipe S. Freitas<sup>1,3†</sup>, Philip A. Pika<sup>2,4††</sup>, Sabine Kasten<sup>5,6,7</sup>, Bo B. Jørgensen<sup>8</sup>, Jens Rassmann<sup>9</sup>, Christophe Rabouille<sup>9</sup>, Shaun Thomas<sup>10, †††</sup>, Henrik Sass<sup>10</sup>, Richard D. Pancost<sup>1,3</sup>, Sandra Arndt<sup>4</sup>

<sup>1</sup>Organic Geochemistry Unit, School of Earth Sciences & School of Chemistry, University of Bristol, Bristol, BS8 1TS, United Kingdom

<sup>2</sup>BRIDGE, School of Geographical Sciences, University of Bristol, Bristol, BS8 1RL, United Kingdom

<sup>3</sup>Cabot Institute for the Environment, University of Bristol, Bristol, BS8 1UH, United Kingdom

<sup>4</sup>Biogeochemistry and Earth System Modeling, Geosciences, Environment and Society Department, Université Libre de Bruxelles, Brussels, CP160/03 1050, Belgium

<sup>5</sup>Alfred Wegener Institute Helmholtz Centre for Polar and Marine Research, Bremerhaven, 27570, Germany

<sup>6</sup>Faculty of Geosciences, University of Bremen, 28359 Bremen, Germany

<sup>7</sup>MARUM – Center for Marine Environmental Sciences, University of Bremen, 28359 Bremen, Germany

<sup>8</sup>Section for Microbiology, Department of Biology, Aarhus University, 8000 Aarhus C, Denmark

<sup>9</sup>Laboratoire des Sciences du Climat et de l'Environnement, LSCE/IPSL, CEA-CNRS-UVSQ-Université Paris Saclay, 91198 Gif-sur-Yvette, France

<sup>10</sup>School of Earth and Ocean Sciences, Cardiff University, Cardiff, CF10 3AT, United Kingdom

† Current address: School of Earth Sciences, University of Bristol, Bristol, BS8 1RJ, United Kingdom

†† Current address: Department of Earth Sciences, VU University of Amsterdam, 1081 HV Amsterdam, The Netherlands

††† Current address: RSR Ltd, Parc Ty Glas, Llanishen, Cardiff, CF14 5DU, United Kingdom

25 *Correspondence to:* Felipe S. Freitas (felipe.salesdefreitas@bristol.ac.uk)

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## S1. Extended information on Study Sites

35 This section complements the information presented in Sect. 2 and Table 2 in the main manuscript. Table S1 displays further details on site descriptions regarding broad biological properties of each study site.

**Table S1. Oceanographic and biologic context of depositional environments.**

Site	Pelagic ecosystem structure	References
Severn estuary	Turbidity resulting from strong tidal dynamics restricts phytoplankton communities. In tidal flats microphytobenthos (mainly diatoms; but also, green algae, euglenophytes, and cyanobacteria) contributes to most of primary productivity.	Langston et al. (2010); Thomas (2014); Underwood (2010)
Rhone delta	Strong pulses of fresh water and sediments associated to flood events. Coccolithophorids are dominant year-round. During spring blooms, diatoms peak and dominates over cyanobacteria.	Antonelli et al. (2008); Cathalot et al. (2010); Zebracki et al. (2015); Uitz et al. (2012)
Aarhus Bay	Stable stratification separates upper oligotrophic wates from lower nutrient replete layers. Mixotrophic flagellates account for up to half of pigmented biomass in upper layers. Chl-a maxima develop at the pycnocline. Spring phytoplankton blooms dominated by diatoms.	Chen et al. (2017); Havskum and Riemann (1996); Jensen et al. (1990); Lomstein et al. (1990); Thingstad et al. (1996)
Arkona Basin	Long-term decrease in water transparency associated to eutrophication. Spring blooms are mainly dominated by diatoms and dinoflagellates. Cyanobacteria blooms occur in the end of summer.	Fleming and Kaitala (2006); Fleming-Lehtinen and Laamanen (2012); Zettler et al. (2007)
Helgoland Mud Area, North Sea	Spring blooms dominated by diatoms. Flagellates growth is restricted to summer months. Haptophytes represent a large contribution to phytoplankton community, although the timing and intensity of blooms vary in a less predictive way.	Hebbeln et al. (2003); Hickel et al. (1992); Riebesell (1993); Wiltshire and Manly (2004)
Skagerrak	Short-term and inter-annual variability in phytoplankton community is common. Chl-a displays a sub-surface maximum below pycnocline. Diatoms dominant during spring, whereas dinoflagellates are predominant during summer. Haptophytes account for half of nanophytoplankton. Cyanobacteria abundance is high in the central basin during summer.	Dahl and Johannessen (1998); Karlson et al. (1996); Richardson et al. (2003); Trimmer et al. (2013)
Arabian Sea	Monsoon regime drives hydrology and has a strong impact on the seasonality of primary productivity. Cyanobacteria dominates oligotrophic, warm upper waters. In subsurface waters eukaryotic phytoplankton are the dominant group and represents 50-80% of carbon biomass. Diatoms dominate coastal upwellings and are gradually replaced offshore by haptophytes.	Barlow et al. (1999); Cowie (2005); Latasa and Bidigare (1998); Rixen et al. (2019); Shalapyonok et al. (2001)
Bering Sea	Nutrient recycling and upwelling support autumn blooms. Ice melting triggers spring blooms. In the absence of sea ice, blooms are delayed until thermal stratification is established. Community composition is dominated by diatoms with small contribution of pico- and nanophytoplankton groups. Coccolithophores form large blooms during calm conditions.	Coyle et al. (2008); Gersonde (2009); Odate (1996); Stabeno and Hunt (2002); Stockwell et al. (2001)
Argentine Basin	Complex hydrology and water masses dynamics. Enhanced productivity at oceanographic fronts and Sub-Antarctic water upwellings. Haptophytes are the dominant group and dominant at shelf-break where strong mixing and nutrient renovation occur. Diatoms occur in low abundance. Other groups include picocyanobacteria, picoplanktonic coccoids, and flagellates.	Calliari et al. (2009); Carreto et al. (2003); Gayoso (1995); Peterson (1992)

40 **S2. Extended information on Reaction-Transport Model description**

The Biogeochemical Reaction Network Simulator (BRNS) (Aguilera et al., 2005; Regnier et al., 2002) is an adaptive simulation environment that has been successfully employed to reproduce and quantify diagenetic processes in marine sediments across a wide range of depositional environments and timescales (Thullner et al., 2009; Wehrmann et al., 2013). It calculates concentration depth profiles of solid and dissolved species in marine sediments according to the vertically-resolved mass conservation equation of solid and dissolved species in porous media (Berner, 1980; Boudreau, 1997):

$$\frac{\partial \sigma C_i}{\partial t} = \frac{\partial}{\partial z} \left( D_{bio} \sigma \frac{\partial C_i}{\partial z} + D_i \sigma \frac{\partial C_i}{\partial z} \right) - \frac{\partial \sigma \omega C_i}{\partial z} + \alpha_i \sigma (C_i(0) - C_i) + \sum_n s_i^n r_n \quad (S1)$$

50 The first three terms on the right-hand side represent the transport process (bioturbation and molecular diffusion, advection, and bioirrigation; see Sect. S2.1), whereas the last term denotes the sum of all reactions (production and consumption; see Sect. S2.2) that affect species  $i$ . Table S2 provides a summary of all symbols employed here. In the following sections, we provide a detailed description of the model parameterization and solution.

**Table S2. Summary of model elements incorporated in the BRNS.**

Symbol	Description
	<i>Chemical species, i</i>
TOC	Total organic carbon
CH <sub>2</sub> O	Organic matter (simplified stoichiometry)
O <sub>2</sub>	Oxygen
NO <sub>3</sub> <sup>-</sup>	Nitrate
SO <sub>4</sub> <sup>2-</sup>	Sulfate
CH <sub>4</sub>	Methane
NH <sub>4</sub> <sup>+</sup>	Ammonium
PO <sub>4</sub> <sup>3-</sup>	Phosphate
HS <sup>-</sup>	Sulfides
	<i>Model parameters</i>
C <sub>i</sub>	Concentration of species $i$
$t$	Time
$z$	Sediment depth
$L$	Length of model domain
$T$	Temperature
$S$	Salinity
$h$	Water depth
	Porosity term
$\sigma$	Solid species, $\sigma = 1 - \varphi$ Dissolved species, $\sigma = \varphi$
$\varphi$	Sediment porosity
$\varphi_0$	Porosity at sediment-water interface
$\varphi_z$	Porosity at depth
$\varphi_\infty$	Porosity at greater depth
$\beta$	Porosity attenuation coefficient
$\omega$	Burial velocity – sedimentation rates
$\omega_0$	Burial velocity at sediment-water interface
$\omega_z$	Burial velocity at depth
$D_i$	Effective molecular diffusion coefficient of dissolved species $i$ at 0 °C
$D_i^*$	Corrected molecular diffusion coefficient of dissolved species $i$
$D_{bio}$	Bioturbation diffusion coefficient
$z_{bio}$	Depth of bioturbated zone – sediment mixed layer
$\alpha_i$	Bioirrigation rate
$\alpha_0$	Bioirrigation coefficient at sediment-water interface
$x_{irri}$	Bioirrigation attenuation depth length
$s_i^n$	Stoichiometric coefficient of specie $i$

$n$	Kinetically controlled reaction
$r_n$	Reaction rate
$K_i$	Half-saturation constant of species $i$
$f_i$	Primary redox reaction inhibition term of species $i$
$k_n$	First-order bimolecular rate constant of secondary redox reaction $r_n$
$x/y/z$	Stoichiometric constants
$age$	Age of organic matter
$\Gamma(v)$	gamma distribution
$a$	Reactive continuum model shaping parameter
$v$	Reactive continuum model scaling parameter
$om(k, t)$	Probability density function of OM distribution
$G_i$	Initial $TOC$ proportion in fraction $i$ – multi-G approximation of RCM
$k_i$	First-order degradation rate constant of fraction $i$
$g(k, 0)$	Initial fraction of $TOC$ characterized by a distinct reactivity
$G(k, 0)$	Initial fraction of $TOC$ within the reactivity range between 0 and $k$

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### S2.1. Transport parameters

The BRNS set-up used here accounts for sediment accumulation and compaction, molecular diffusion, bioturbation, and bioirrigation (see Eq. S1). Global transport parameter values are given in Table S3. Site-specific transport parameter values are given in the main manuscript (Table 5). Sediment porosity is assumed to decrease exponentially with depth due to sediment compaction:

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$$\varphi_z = \varphi_\infty + (\varphi_0 - \varphi_\infty) \exp(-\beta z) \quad (S2)$$

Consequently, the burial velocity is corrected for the effect of compaction assuming steady-state compaction (e.g., Berner, 1980):

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$$\omega_z = \omega_0(1 - \varphi_0)/(1 - \varphi_z) \quad (S3)$$

The diffusive fluxes are commonly quantified by means of Fick's first law of diffusion, which depends on the molecular diffusive coefficient  $D_i$  ( $i$  = dissolved species) (Boudreau, 1997; Burdige, 2006). The effective molecular diffusion coefficients are derived from Van Cappelen and Wang (1996). Here,  $D_i$  are corrected for temperature, salinity, and tortuosity. For solid species  $D_i = 0$ , whereas for dissolved species the corrected  $D_i^*$  is given by Boudreau (1997):

$$D_i^* = \frac{D_i(T,S)}{1 - \ln(\varphi^2)} \quad (S4)$$

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The model also accounts for the effect of sediment reworking by infaunal organisms in the bioturbated upper sediment layer ( $z < z_{bio}$ ). The process is generally described by a dispersive term with constant bioturbation diffusion coefficient  $D_{bio}$  (Boudreau, 1986). For bioturbated sites,  $D_{bio}$  was constrained based on an empirically derived relationship proposed by Middelburg et al. (1997):

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$$D_{bio} = 5.2 \cdot 10^{(0.7624 - 0.0003972 \cdot h)} \quad (S5)$$

For the Rhone delta,  $D_{bio}$  values were derived from Pastor et al. (2011).  $D_{bio}$  was kept constant within the bioturbated zone ( $z < z_{bio}$ ) then set to zero below  $z_{bio}$ . Bioturbation depth was fixed at 10 cm for most bioturbated sediments (Table 4), based on a compilation of mixed layer depths (Boudreau, 1994, 1998). At anoxic depositional environments (i.e., where  $O_2$

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concentrations are zero),  $z_{bio}$  was set to zero (Table 4). Similarly, the bioturbation diffusion coefficient  $D_{bio}$  was set to zero in anoxic environments. Bioirrigation describes the mixing by benthic macrofaunal organisms that build burrows or tubes in the sediment for feeding. It is parameterized as a nonlocal transport process with a nonlocal bioirrigation coefficient  $\alpha_i$ , which describes the exchange rates between the sediment-water interface and porewater at depth in bioirrigated zone of sediments (Aller, 1994; Aller and Aller, 1998; Burdige, 2006). The bioirrigation rate is given by (Thullner et al., 2009; Wehrmann et al., 2013):

$$\alpha_i = \alpha_0 \cdot \exp(-x/x_{irri}) \quad (S6)$$

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For solid species,  $\alpha_i$  is set to zero.

**Table S3. Global model transport parameter values implemented in the RTM. Effective molecular diffusion coefficients ( $D_i$ ) are given for  $T = 0$  °C and are corrected by site-specific temperature, salinity, and tortuosity.**

Parameter	Unit	Value	Reference
$\alpha_0$	yr <sup>-1</sup>	10	Thullner et al. (2009)
$x_{irri}$	cm	3.5	Thullner et al. (2009)
$D_{O_2}$	cm <sup>2</sup> yr <sup>-1</sup>	380.44	Van Cappellen and Wang (1996)
$D_{NO_3^-}$	cm <sup>2</sup> yr <sup>-1</sup>	394.58	Van Cappellen and Wang (1996)
$D_{SO_4^{2-}}$	cm <sup>2</sup> yr <sup>-1</sup>	173.92	Van Cappellen and Wang (1996)
$D_{NH_4^+}$	cm <sup>2</sup> yr <sup>-1</sup>	395.87	Van Cappellen and Wang (1996)
$D_{H_2S}$	cm <sup>2</sup> yr <sup>-1</sup>	331.61	Van Cappellen and Wang (1996)
$D_{CH_4(g)}$	cm <sup>2</sup> yr <sup>-1</sup>	263.93	Van Cappellen and Wang (1996)

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## S2.2. Reaction parameters

The reaction network implemented in the BRNS encompasses the most pertinent primary and secondary redox reactions found in the upper layers of marine sediments. Its formulation and parametrization builds on a number of previous studies that investigate diagenetic dynamics across several depositional environments and scales (Aguilera et al., 2005; Thullner et al., 2009; Van Cappellen and Wang, 1996; Wang and Van Cappellen, 1996; Wehrmann et al., 2013). It explicitly accounts for the heterotrophic degradation of OM coupled to the consumption of oxygen (aerobic OM degradation), nitrate (denitrification), sulfate (organoclastic sulfate reduction), as well as methanogenesis. Additionally, it accounts for nitrification, sulfide re-oxidation by  $O_2$ , anaerobic oxidation of methane (AOM) coupled to sulfate reduction and  $CH_4$  reoxidation by  $O_2$ . Due to the limited availability of data to constrain manganese oxide ( $MnO_2$ ) and iron hydroxide ( $Fe(OH)_3$ ) depositional fluxes, the model does not account for metal oxides reduction pathways. At certain depositional settings (e.g., Skagerrak), metal oxide pathways can be relatively important (e.g., Canfield et al., 1993; Rysgaard et al., 2001), particularly in continental margins that receive considerable inputs of iron (Beckler et al., 2016). Nevertheless, a previously published global assessment of the importance of metabolic pathways in marine sediments has found their contributions to the overall heterotrophic OM degradation to be negligible at a global scale (Thullner et al., 2009).

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OM heterotrophic degradation follows first-order degradation kinetics (see below; Sect. S2.3) and is coupled to the consumption of terminal electron acceptors (TEAs) (primary redox reactions,  $r_1 - r_4$ ; Table S4). The sequential utilization of TEAs is kinetically controlled by rate laws (Table S5) (e.g., Van Cappellen and Wang, 1996). Additionally, the reaction

120 network accounts for the reoxidation of reduced species produced during OM heterotrophic degradation (secondary redox reactions;  $r_5 - r_8$ ; Table S4). Following the classical approach, the rates of secondary redox reactions are described by bimolecular rate laws with the rate constant  $k_n$  (Van Cappellen and Wang, 1996). Global reaction parameter values are given in Table S6.

125 **Table S4. Reaction network governing heterotrophic organic matter degradation in marine sediments implemented in the Reaction-Transport Model (Adapted from Aguilera et al., 2005; Thullner et al., 2009; Wehrmann et al., 2013).**

Reaction Pathway	Stoichiometry	Reaction rate
<i>Primary redox reactions</i>		
$r_1$ Aerobic OM degradation	$(CH_2O)_x(NH_3)_y(H_3PO_4)_z + (x + 2y)O_2 + (y + 2z)HCO_3^-$ $\rightarrow (x + y + 2z)CO_2 + yNH_4^+ + zHPO_4^{2-}$ $+ (x + 2y + 2z)H_2O$	$r_1 = v \cdot (a + age)^{-1}$ $\cdot CH_2O$ $\cdot f_{O_2}$
$r_2$ Denitrification	$(CH_2O)_x(NH_3)_y(H_3PO_4)_z + \left(\frac{4x+3y}{5}\right)NO_3^-$ $\rightarrow \left(\frac{2x+4y}{5}\right)N_2 + \left(\frac{x-3y+10z}{5}\right)CO_2 + \left(\frac{4x+3y-10z}{5}\right)HCO_3^-$ $+ zPO_4^{2-} + \left(\frac{3x+6y+10z}{5}\right)H_2O$	$r_2 = v \cdot (a + age)^{-1}$ $\cdot CH_2O$ $\cdot f_{NO_3}$
$r_3$ Sulfate reduction	$(CH_2O)_x(NH_3)_y(H_3PO_4)_z + \frac{x}{2}SO_4^{2-} + (y - 2z)CO_2 + (y - 2z)H_2O$ $\rightarrow (x + y - 2z)HCO_3^- + yNH_4^+ + zHPO_4^{2-} + \frac{x}{2}H_2S$	$r_3 = v \cdot (a + age)^{-1}$ $\cdot CH_2O$ $\cdot f_{SO_4^{2-}}$
$r_4$ Methanogenesis	$(CH_2O)_x(NH_3)_y(H_3PO_4)_z + (y - 2z)H_2O$ $\rightarrow \left(\frac{x-2y+4z}{2}\right)CO_2 + (y - 2z)HCO_3^- + yNH_4^+ + zHPO_4^{2-}$ $+ \frac{x}{2}CH_4$	$r_4 = v \cdot (a + age)^{-1}$ $\cdot CH_2O$ $\cdot f_{CH_4}$
<i>Secondary redox reactions</i>		
$r_5$ Ammonium oxidation by oxygen	$NH_4^+ + 2O_2 + 2HCO_3^- \rightarrow NO_3^- + 2CO_2 + 3H_2O$	$r_5 = k_5 \cdot NH_4^+ \cdot O_2$
$r_6$ Sulfide oxidation by oxygen	$H_2S + 2O_2 + 2HCO_3^- \rightarrow SO_4^{2-} + 2CO_2 + 2H_2O$	$r_6 = k_6 \cdot (HS^- + H_2S) \cdot O_2$
$r_7$ Anaerobic oxidation of methane (AOM)	$CH_4 + CO_2 + SO_4^{2-} \rightarrow 2HCO_3^- + H_2S$	$r_7 = k_7 \cdot CH_4 \cdot SO_4^{2-}$
$r_8$ Methane oxidation by oxygen	$CH_4 + 2O_2 \rightarrow CO_2 + 2H_2O$	$r_8 = k_8 \cdot CH_4 \cdot O_2$

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**Table S5. Kinetic rate laws controlling the reaction network.**

Kinetic rate laws	
$r_1$	$\begin{cases} f_{O_2} = 1 \text{ for } O_2 > K_{O_2} \\ f_{O_2} = \frac{O_2}{K_{O_2}} \text{ for } O_2 \leq K_{O_2} \end{cases}$
$r_2$	$\begin{cases} f_{NO_3^-} = 0 \text{ for } f_{O_2} = 1 \\ f_{NO_3^-} = (1 - f_{O_2}) \text{ for } f_{O_2} < 1 \text{ and } NO_3^- > K_{NO_3^-} \\ f_{NO_3^-} = (1 - f_{O_2}) \frac{NO_3^-}{K_{NO_3^-}} \text{ for } f_{O_2} < 1 \text{ and } NO_3^- \leq K_{NO_3^-} \end{cases}$
$r_3$	$\begin{cases} f_{SO_4^{2-}} = 0 \text{ for } f_{O_2} + f_{NO_3^-} = 1 \\ f_{SO_4^{2-}} = (1 - f_{NO_3^-}) \text{ for } f_{NO_3^-} < 1 \text{ and } SO_4^{2-} > K_{SO_4^{2-}} \\ f_{SO_4^{2-}} = (1 - f_{NO_3^-}) \frac{SO_4^{2-}}{K_{SO_4^{2-}}} \text{ for } f_{NO_3^-} < 1 \text{ and } SO_4^{2-} \leq K_{SO_4^{2-}} \end{cases}$
$r_4$	$f_{CH_4} = \left(1 - (f_{O_2} + f_{NO_3^-} + f_{SO_4^{2-}})\right)$

140 **Table S6. Global reaction parameter values implemented in the RTM.**

Parameter	Unit	Value	Reference
$x/y/z$	–	106/16/1	Redfield (1934)
$v$	–	variable	Boudreau and Ruddick (1991)
$a$	yr	variable	Boudreau and Ruddick (1991)
$age$	yr	variable	Mogollón et al. (2012)
$K_{O_2}$	M	$8.0 \cdot 10^{-9}$	Van Cappellen and Wang (1996)
$K_{NO_3^-}$	M	$5.0 \cdot 10^{-9}$	Van Cappellen and Wang (1996)
$K_{SO_4^{2-}}$	M	$1.0 \cdot 10^{-7}$	Van Cappellen and Wang (1996)
$k_5$	$M^{-1} yr^{-1}$	$1.0 \cdot 10^7$	Dale et al. (2011)
$k_6$	$M^{-1} yr^{-1}$	$1.0 \cdot 10^9$	Dale et al. (2011)
$k_7$	$M^{-1} yr^{-1}$	$5.0 \cdot 10^6$	Dale et al. (2011)
$k_8$	$M^{-1} yr^{-1}$	$1.0 \cdot 10^{13}$	Dale et al. (2011)

**S2.3 Organic matter degradation model**

OM is composed of a complex and dynamic mixture of compounds that are distributed over a wide, continuous spectrum of reactivities. Thus, OM degradation is described by the reactive continuum model (RCM) (Boudreau and Ruddick, 1991), which assumes a continuous distribution of OM compounds over the entire reactivity spectrum. The RCM assumes that the initial distribution of OM compounds over the reactivity spectrum follows a gamma distribution that is completely determined by two free parameters:  $a$  is the average lifetime of the more reactive components of bulk OM, and  $v$  represents the dimensionless scaling parameter of the distribution near  $k = 0$ . As such, the RCM approach requires the definition of two parameters that will define the shape of the OM distribution over reactivity  $k$ :

$$k(z) = \frac{v}{a + age(z)} \quad (S7)$$

$age(z)$  denotes the age of the sediment layer at depth  $z$ . For non-bioturbated sediments ( $z > z_{bio}$ ) the burial  $age(z)$  can be calculated as a function of the burial velocity:

$$age(z) = age_0 + \frac{(1-\varphi)z + \beta^{-1}(\varphi_0 - \varphi_\infty)(\exp(-\beta z) - 1)}{\omega(1-\varphi_0)} \quad (S8)$$

160 However, within the bioturbated upper sediment layers, the age distribution of reactive species is controlled by both  
 165 sedimentation, bioturbation, and the reactivity  $k$  of reactive species (Meile and Van Cappellen, 2005). As such, the RCM  
 approach cannot easily be applied to the upper mixed sediment layers. Therefore, within the bioturbated layer ( $z < z_{bio}$ ), the  
 RCM is approximated by a discrete multi-G model (200 fractions) to circumvent the difficulty of quantifying OM ages within  
 bioturbated sediments (Dale et al., 2015; Meile and Van Cappellen, 2005). The multi-G model approach divides the bulk OM  
 into 200 discrete compound classes  $i$  each degrading according to first-order kinetics with a degradation rate constant  $k_i$   
 (Jørgensen, 1978). The degradation rate of the bulk OM,  $R_{TOC}$ , and the initial concentration of OM in compound class  $i$ ,  
 $TOC_i(0)$  are thus given by:

$$C_{ox}(z) = \sum_{i=1}^{200} k_i \cdot TOC_i(z) \quad (S9)$$

$$170 \quad TOC_i(0) = G_i \cdot TOC \quad (S10)$$

The initial proportion of  $TOC$  in fraction  $i$ ,  $G_i$ , as well as the compound class specific reactivity rate-constant  $k_i$  can be  
 determined from the initial probability density function  $om(k, t)$  that provides the concentration of  $TOC$  having a degradability  
 $k$  and  $k + dk$  at time 0. The initial fraction of  $TOC$  characterized by a distinct reactivity  $k$  is given by  $g(k, 0)$ :

$$175 \quad g(k, 0) = \frac{om(k, 0)}{TOC_0} = \frac{a^v \cdot k^{v-1} \cdot e^{-a \cdot k}}{\Gamma(v)} \quad (S11)$$

The initial fraction of  $TOC$  within the reactivity range between 0 and  $k$  is given by integrating Eq. S12 (if  $a, v, k > 0$ ):

$$180 \quad G(k, 0) = \int_0^k g(0, k) dk = \int_0^k \frac{a^v \cdot k^{v-1} \cdot e^{-a \cdot k}}{\Gamma(v)} dk = \left( \frac{1 - \Gamma(v, a \cdot k)}{\Gamma(v)} \right) \quad (S12)$$

Where  $\Gamma(v, a \cdot k)$  denotes the inverse gamma distribution.

185 Within the bioturbated sediment layer, the RCM was approximated by dividing the reactivity range  $k = [10^{-15}, 10^{(-\log(a)+2)}]$   
 into 200 equal reactivity bins,  $k_j$ . The initial fraction  $G_i$  of  $TOC$  within reactivity bin  $k_{j-1}$  and  $k_j$  (and thus with reactivity  $k_i =$   
 $k_{j-1} + \frac{k_j - k_{j-1}}{2}$ ) in the 200G-model is then calculated as:

$$G_i = G(k_j, 0) - G(k_{j-1}, 0) \quad (S13)$$

190 The least and the most reactive fractions  $G_1$  and  $G_{200}$  with reactivity  $k_1 = 10^{-15} \text{ yr}^{-1}$  and  $k_{200} = 10^{-\log(a)+2} \text{ yr}^{-1}$ , respectively,  
 are calculated based on the incomplete gamma distribution:

$$G_1 = \int_{k_1}^{\infty} g(k_1, 0) dk = \frac{\Gamma(v, a \cdot k_1)}{\Gamma(v)} \quad (S14)$$

$$195 \quad G_{200} = \int_{k_{200}}^{\infty} g(k_{200}, 0) dk = \frac{\Gamma(v, a \cdot k_{200})}{\Gamma(v)} \quad (S15)$$

Once  $G_i$  and  $k_i$  are determined, the steady-state analytical solution of the diffusion-advection-reaction equation (Boudreau, 1997) for OM in the bioturbated zone can then be calculated by:

$$TOC(z) = \sum_{i=1}^{200} A_i \cdot \exp(a_i \cdot z) + B_i \cdot \exp(b_i \cdot z) \quad (S16)$$

with:

$$a_i = \frac{\omega - \sqrt{\omega^2 + 4 \cdot D_{bio} \cdot k_i}}{2 \cdot D_{bio}} \quad (S17)$$

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and

$$b_i = \frac{\omega + \sqrt{\omega^2 + 4 \cdot D_{bio} \cdot k_i}}{2 \cdot D_{bio}} \quad (S18)$$

210  $A_i$  and  $B_i$  denote integration constants that can be determined by defining appropriate boundary conditions (Boudreau, 1997) for OM at the upper and lower boundaries. Below the bioturbated zone, the depth evolution of  $TOC$  is determined by the RCM formulation.

## 215 **S2.5 Model solution**

Transport and reaction equations were solved sequentially. Firstly, the diffusion term was discretized at each time-step of the numerical integration using the semi-implicit Crank-Nicholson scheme. This was followed by the calculation of the advective transport, using a 3<sup>rd</sup> order accurate total variation diminishing algorithm with flux limiters (Regnier et al., 1998). The reaction network was subsequently solved. The mass-conservation equation (Eq. S1) was discretized on an uneven grid (Boudreau, 1997):

220

$$z(n) = \frac{L((\xi_n^2 + \xi_c^2)^{0.5} - \xi_c)}{(L^2 + \xi_c^2)^{0.5} - \xi_c} \quad (S19)$$

where  $z(n)$  is the depth of the  $n^{th}$  grid point,  $L$  denotes the length of the model domain,  $\xi_n$  is a point in a hypothetical grid, and  $\xi_c$  is depth relative to which  $z(n)$  is quadratically distributed for  $\xi_n \gg \xi_c$  and linearly distributed for  $\xi_n \ll \xi_c$ .  $L$  and  $\xi_n$  were chosen so that the grid size,  $\Delta z$ , increases downcore from SWI to a maximum of  $L$ . The size of model domain  $L$  was fixed at 1,000 cm for all sites, except for the Bering Sea, in which the model domain is extended to 1,500 cm, due to the low sedimentation rate assumed for this site (Table 2). This choice is based on initial tests and ensures that the model domain covers the diagenetically most active zone, thus reducing the influence of biogeochemical dynamics in underlying sediments on biogeochemical dynamics within the model domain. BRNS was run until steady state ( $\Delta t < 0.01$ ) was reached:

230

$$run\ time = 2 \cdot \frac{L}{\omega} \quad (S20)$$

### S3. Establishing a minimum dataset of observations for the determination of apparent organic matter reactivity

We use an inverse model approach to extract the optimal OM reactivity parameter set  $a$  and  $v$  (i.e., the apparent OM reactivity) by assuming that the rank of a parameter set depends on the similarity between simulated and measured data. However, all inverse model approaches and, thus the quality of the inversely determined parameters are subject to limitations, such as core top loss, uniqueness of fit, unresolved processes, transient dynamics. These limitations can be alleviated by using comprehensive, multi-component observational data sets for the inverse modelling approach. However, especially on the global scale, data availability is often limited. Consequently, it is important to identify a minimum set of observational data that are widely available and comparably easy to measure. To identify such a minimum observational data set, we here tested performance of different artificial porewater data sets in determining apparent OM reactivity. Using an artificial dataset (Table S7; Fig. S1 and S2) that was generated by using a defined OM reactivity, we ran the inverse model approach (see main manuscript, Sect. 3.2.4) using 1) *TOC* profiles only and 2) *TOC* and  $SO_4^{2-}$  profiles.

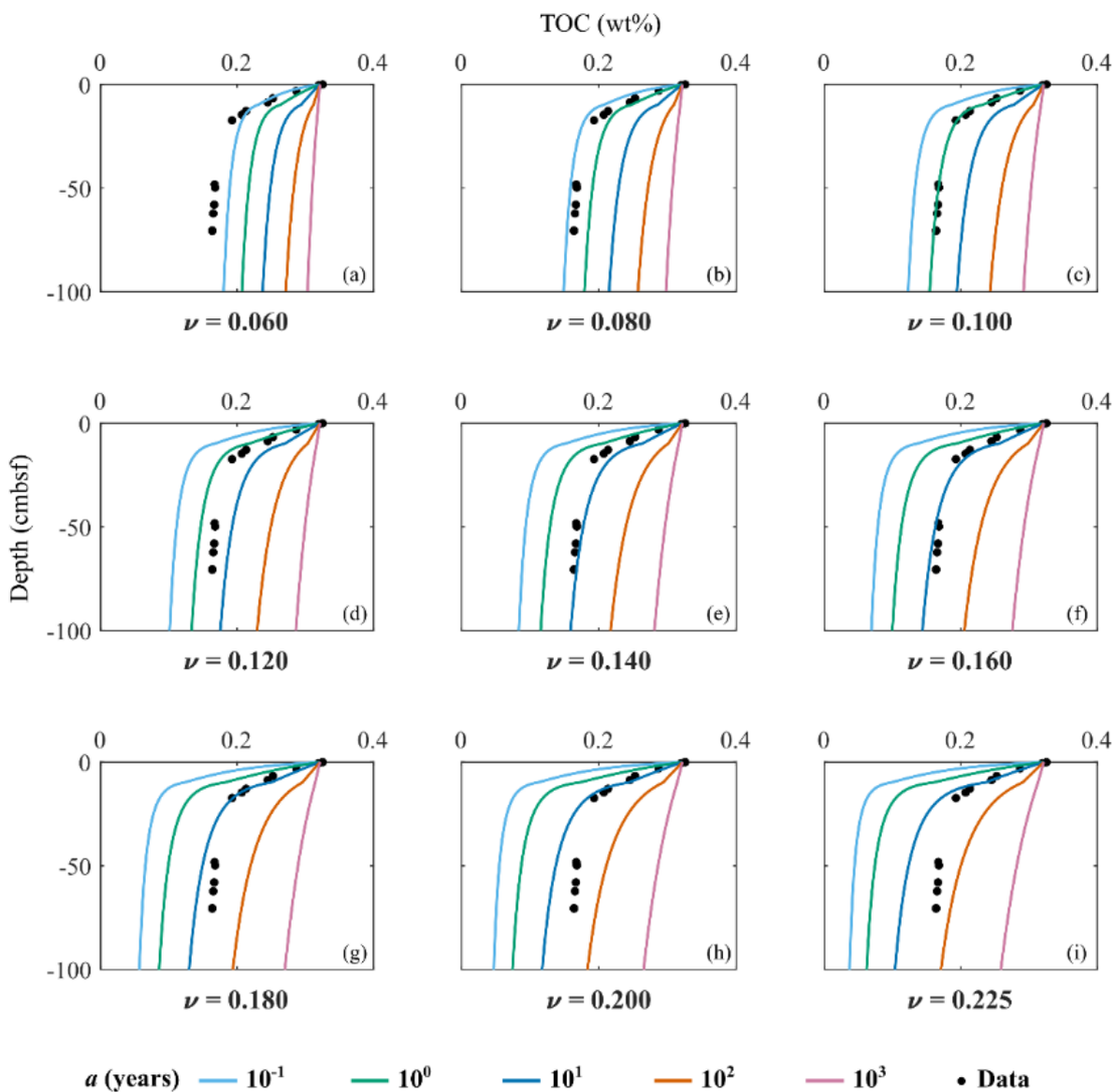
**Table S7. Model parameters and boundary conditions adopted in the sensitivity analyses used to determine the minimum dataset necessary to constrain apparent organic matter reactivity  $k$  based on the Reactive Continuum Model parameters  $a$  and  $v$ . Model parameters omitted here follow those in Tables S2 and S6.**

<i>Parameter</i>	<i>Unit</i>	<i>Value</i>	<i>Parameter</i>	<i>Unit</i>	<i>Value</i>	<i>Parameter</i>	<i>Unit</i>	<i>Value</i>
<i>Transport</i>			<i>Reactions</i>			<i>Boundary conditions</i>		
$L$	cm	150	$x/y/z$	–	106/16/1	<i>TOC</i>	wt%	0.32
$\alpha_0$	yr <sup>-1</sup>	10	$a$	yr	[10 <sup>-3</sup> – 10 <sup>7</sup> ]	$O_2$	μM	325
$x_{irr}$	cm	3.5	$v$	–	[10 <sup>-1</sup> – 10 <sup>0</sup> ]	$NO_3^-$	μM	9
$z_{bio}$	cm	10				$SO_4^{2-}$	mM	28
$D_{bio}$	cm <sup>2</sup> yr <sup>-1</sup>	23.98				$NH_4^+$	μM	0
$\phi$	–	0.47				$HS^-$	μM	0
$\omega$	cm yr <sup>-1</sup>	0.06				$CH_4$	μM	0
$T$	°C	0.65						
$S$	–	35						
$h$	m	250						

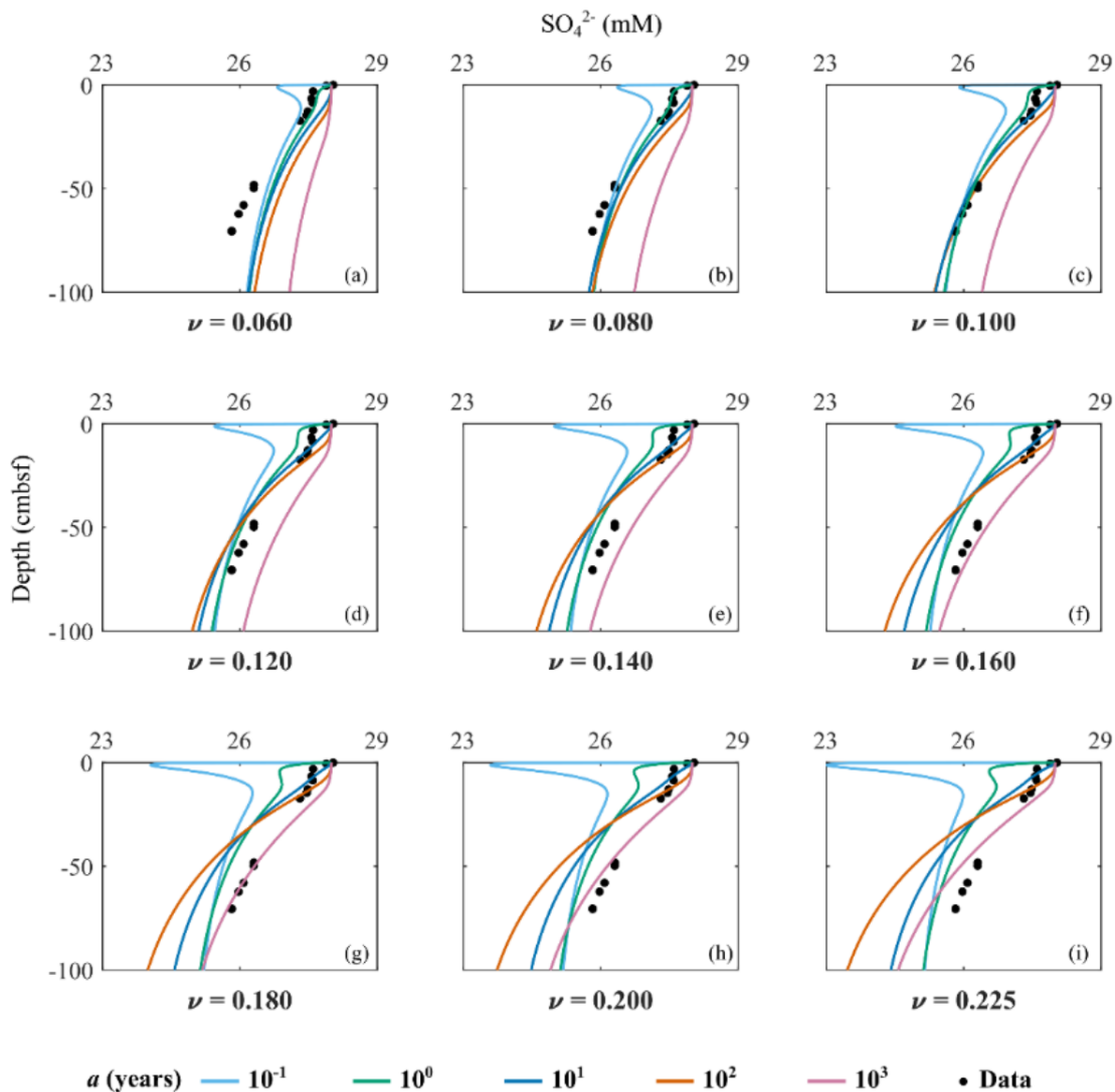
Our sensitivity analysis confirms that when *TOC* is considered as a single constraint, multiple pairs of  $a$  and  $v$  produce very similar *TOC* depth-profiles that fit the artificial data equally well (Fig. S1). Model results show that including  $SO_4^{2-}$  depth profiles add an additional constraint (Fig. S2) and facilitates the identification of a best fit  $a$  and  $v$  parameter set. This is illustrated by the Taylor diagrams (Taylor, 2001), which summarizes the statistical fitting ( $r$ , RMSD, and SD) (Fig. S3). In addition, because changes in  $a$ – $v$  exert different effects on *TOC* and  $SO_4^{2-}$  depth profiles (Fig. S1 and S2), including these two species reduces the impact of the  $a$ – $v$  correlation on the uniqueness of fit (Fig. S3).

$CH_4$  profiles potentially offer an additional qualitative constraint when the data are available. The dynamics of  $SO_4^{2-}$  and  $CH_4$  are solely controlled by OM degradation and AOM (e.g., Bowles et al., 2014; Egger et al., 2018; Regnier et al., 2011), although their distributions can be also affected by bioirrigation in particularly shallow sulfate-methane transition zones (SMTZs) (e.g., Dale et al., 2019). In anoxic settings and deeply buried sediments,  $SO_4^{2-}$  is the dominant TEA and  $CH_4$  is the most common reduced species (Jørgensen et al., 2019a, b). Thus, a combination of *TOC*,  $SO_4^{2-}$ , and  $CH_4$  (if available to verify the depths of

265 the SMTZ) depth profiles incorporate the information contained in the observed benthic sulfur and carbon dynamics and is sufficient to extract robust estimates of apparent OM reactivity and its evolution from the sediment-water interface down to the SMTZ.



270 **Figure S1.** Total organic carbon depth profiles yielded by the sensitivity analysis for an ensemble of scaling parameter  $\nu$  and shaping parameter  $\alpha$  (see Table S3). Based on *TOC* alone multiple pairs of  $\alpha$  and  $\nu$  would be extracted from the same depth-profile.



275

**Figure S2.** Sulfate depth profiles yielded by the sensitivity analysis for an ensemble of scaling parameter  $\nu$  and shaping parameter  $a$  (see Table S3). Considering  $SO_4^{2-}$  alongside  $TOC$  improves the determination of  $a$  and  $\nu$  by excluding those pairs that only fit  $TOC$ .

280

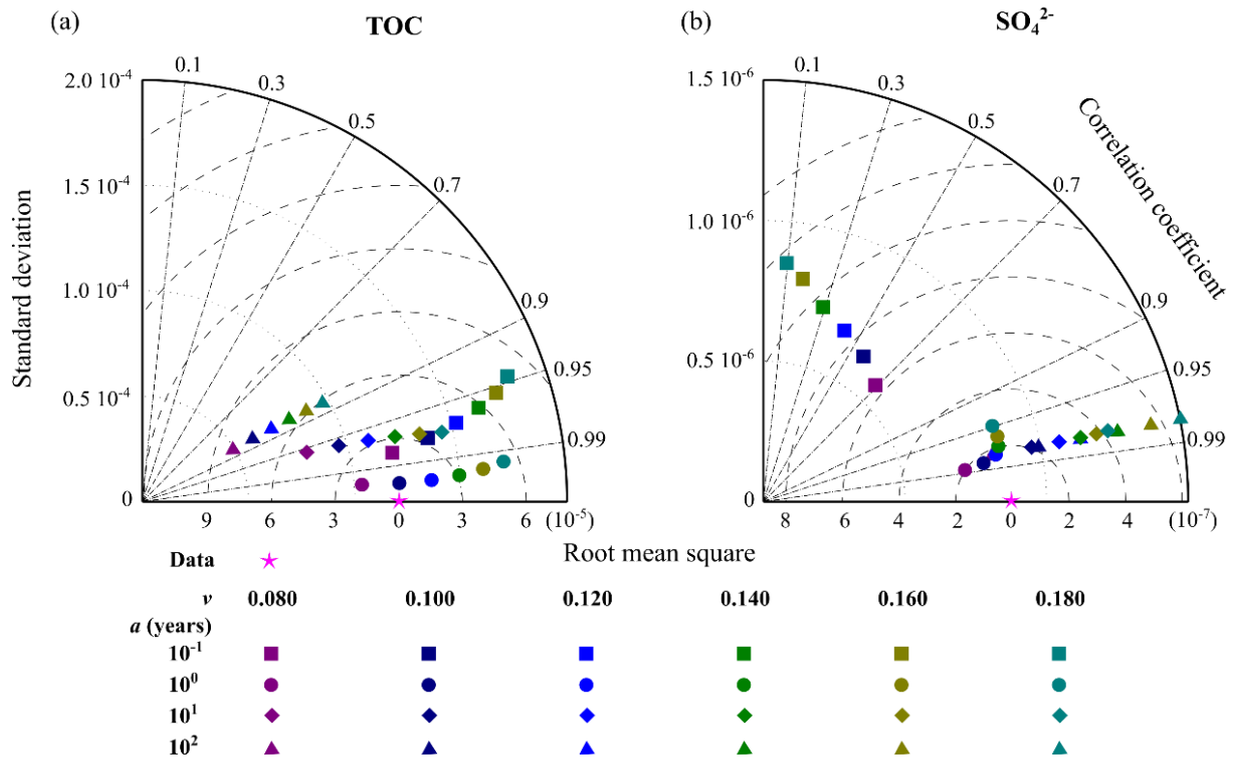


Figure S3. Sensitivity analysis best-fit based on Reactive Continuum model parameters  $a$  and  $v$  for (a) total organic carbon and (b) sulfate. The adoption of two species (i.e.,  $\text{TOC}$  and  $\text{SO}_4^{2-}$ ) relieves the uncertainties in constraining organic matter reactivity parameters.

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