

1 **Assessing global-scale organic matter reactivity patterns in marine**
2 **sediments using a lognormal reactive continuum model**

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15

16 **Abstract**

17 Organic matter (OM) degradation in marine sediments is largely controlled by its
18 reactivity and profoundly affects the global carbon cycle. Yet, there is currently no general
19 framework that can constrain OM reactivity on a global scale. In this study, we propose a
20 reactive continuum model based on a lognormal distribution (*l*-RCM), where OM
21 reactivity is fully described by parameters μ (the mean reactivity of the initial OM bulk
22 mixture) and σ (the variance of OM components around the mean reactivity). We use the
23 *l*-RCM to inversely determine μ and σ at 123 sites across the global ocean. The results show

24 that the apparent OM reactivity ($\langle k \rangle = \mu \cdot \exp(\sigma^2/2)$) decreases with decreasing
25 sedimentation rate (ω) and show that OM reactivity is more than three orders of magnitude
26 higher in shelf than that in abyssal regions. Despite the general global trends, higher than
27 expected OM reactivity is observed in certain ocean regions characterized by great water
28 depth and/or pronounced oxygen minimum zones, such as the Eastern-Western Coastal
29 Equatorial Pacific and the Arabian Sea, emphasizing the complex control of the
30 depositional environment (e.g., OM flux, oxygen content in the water column) on benthic
31 OM reactivity. Notably, the *l*-RCM can also highlight the variability of OM reactivity in
32 these regions. Based on inverse modeling results in our dataset, we establish the significant
33 statistical relationships between $\langle k \rangle$ and ω , and further map the global OM reactivity
34 distribution. The novelty of this study lies in its unifying view, but also in contributing a
35 new framework that allows predicting OM reactivity in data-poor areas based on readily
36 available (or more easily obtainable) information. Such a framework is currently lacking
37 and limits our abilities to constrain OM reactivity in global biogeochemical and/or Earth
38 System Models.

39

40 **1 Introduction**

41 Marine sediments act as the ultimate sink for organic carbon. The size and reactivity of
42 the benthic organic matter (OM) reservoir is a critical component of the global carbon cycle
43 (Arndt et al., 2013). In particular, the reactivity of benthic OM imposes a substantial control
44 on the magnitude of benthic carbon export and burial over geological timescales due to the
45 recycling of organic carbon by dissimilatory microbial activity in the deep biosphere
46 (Boudreau, 1992; Zonneveld et al., 2010), the dissolution and precipitation of carbonates

47 (Meister et al., 2022; Nöthen and Kasten, 2011), and the production of methane (Dickens
48 et al., 2004; Whiticar, 1999). Decades of research have shown that OM reactivity is
49 controlled by both the nature of the OM (origin, composition and degradation state), as
50 well as its environmental and depositional conditions (e.g., redox conditions, sedimentation
51 rate, mineral protection, microbial community composition and biological mixing)
52 (Burdige, 2007; Egger et al., 2018; Hartnett et al., 1998; Hedges and Keil, 1995; Larowe
53 et al., 2020a; Zonneveld et al., 2010). However, due to the complex and dynamic nature of
54 the main controls on OM reactivity, the specific relative significance of these controlling
55 factors remains poorly quantified. Consequently, OM degradation models generally do not
56 explicitly describe the influence of environmental and depositional factors on OM
57 reactivity and its evolution but rather apply simplified parametrizations (Freitas et al., 2021;
58 Pika et al., 2021). Over the past decades, several models have been developed and
59 successfully used to quantify OM degradation in marine sediments. They can be broadly
60 divided into two groups: discrete models, such as the (multi) G model (Berner, 1964;
61 Jørgensen, 1978), and continuum models, such as the reactive continuum model (RCM)
62 (Boudreau and Ruddick, 1991) and the power model (Middelburg, 1989).

63 Discrete models divide the bulk OM pool into several discrete fractions, each with its
64 own constant reactivity (Fig.1A). The 1- G model is the earliest OM degradation model,
65 which is based on the assumption that OM degrades according to first order dynamics with
66 a single constant degradation rate constant (Berner, 1964). The multi- G model, on the other
67 hand, divides OM into several fractions, and each fraction is degraded according to a first-
68 order rate with a fraction-specific reactivity (Jørgensen, 1978). Although multi- G models
69 successfully fit observed OM degradation dynamics when comprehensive data sets are

70 available, their application on a global scale is complicated by the need to partition the OM
71 reactivity into a finite number of fractions and define their reactivities. A multi- G model
72 with n discrete OM fractions requires constraining $2n-1$ parameters and is, thus, over-
73 parametrized (Jørgensen, 1978). Nevertheless, because of its mathematical simplicity and
74 wide use, multi- G models have been used in a range of diagenetic models designed for the
75 global/regional scale (e.g., CANDI, MEDIA, MEDUSA, and OMEN SED) (Boudreau,
76 1996; Meysman et al., 2003; Munhoven, 2007; Pika et al., 2021). Constraining the $2n-1$
77 OM degradation model parameters for these global-scale applications is not
78 straightforward. Early strategies for constraining the reactivity of OM on a global scale
79 have focused on deriving empirical relationships between OM reactivity and single, easily
80 observable characteristics of the depositional environment (water depth, sedimentation rate,
81 or OM flux) (Arndt et al., 2013). However, poor statistically significant link between OM
82 reactivity and depositional environment could be established ($R^2 < 0.1$) after compiling
83 published multi- G model's parameters across a wide range of depositional environments,
84 model complexities as well sediment depths/ burial time scales (Arndt et al., 2013).

85 Reactive continuum models (RCMs) are an alternative to discrete models. They assume
86 that OM compounds are continuously distributed over a wide range of reactivities. The
87 degradation rate can be described as the sum of an infinite number of discrete fractions,
88 each degraded according to first-order kinetics (Boudreau and Ruddick, 1991), as

89
$$G(t) = \int_0^{\infty} G(0) \cdot g(k, 0) \cdot e^{-kt} dk \quad (1)$$

90 where $G(t)$ is OM content at time t , $G(0)$ is OM content at the sediment-water interface
91 (SWI), k is the first-order degradation rate constant, and $g(k,0)$ is the initial reactivity
92 distribution of OM at the SWI. The key to constructing an RCM is to select a continuum

93 distribution that describes the OM reactivity at the SWI (Fig. 1B). Considering the k value
 94 in Eq. 1 must be greater than zero ($k > 0$), some of the all-axial statistical distributions ($x \in$
 95 $(-\infty, +\infty)$) are not appropriate for constructing RCM (e.g., Normal distribution, Fig.1D₁).
 96 Boudreau and Ruddick. (1991), following Aris (1968) and Ho et al. (1987), proposed to
 97 use a Gamma distribution (γ -RCM, Fig.1D₂) due to its mathematical properties and its
 98 ability to capture the observed dynamics:

$$99 \quad g(k, 0) = \frac{a^\nu \cdot k^{\nu-1} \cdot e^{-ak}}{\Gamma(\nu)} \quad (2)$$

100 where a is the average age of the OM at the SWI, ν is the shape parameter, and $\Gamma(\nu)$ is the
 101 Gamma function. In addition, Middelburg. (1989) empirically derived a power law from a
 102 large data compilation of measured OM reactivity (Fig.1C), which is mathematically
 103 equivalent to the γ -RCM. The advantage of the continuum models over the discrete models
 104 is that they merely require constraining two free parameters to capture the widely observed
 105 continuous decrease in OM reactivity with degradation time/depth. Recently, γ -RCM has
 106 been used to inversely determine the free γ -RCM parameters, and thus benthic OM
 107 reactivity, from observed POC and sulfate depth profiles across a wide range of different
 108 depositional environments (Freitas et al., 2021). Although results revealed broad global
 109 patterns, no significant statistical relationship ($R^2 < 0.46$) between the parameters (a and ν)
 110 of the γ -RCM (Arndt et al., 2013) and characteristics of the depositional environment could
 111 be found, and constraining OM degradation model parameters on the global scale thus
 112 remains difficult.

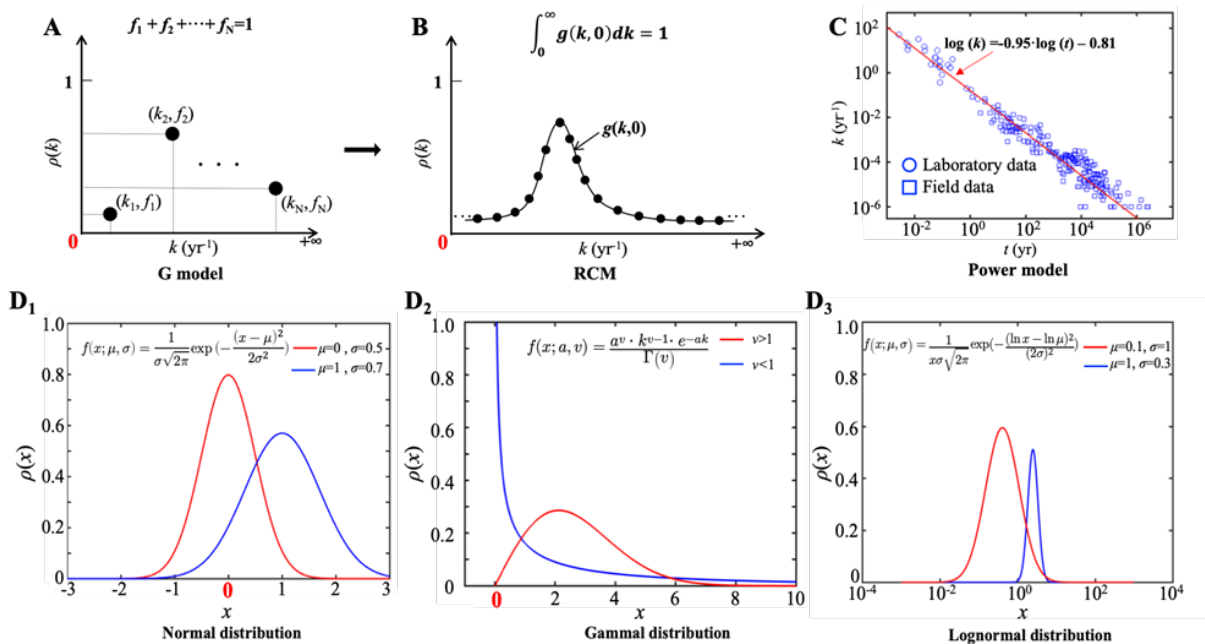
113 Here, we present an RCM based on a lognormal distribution (Forney and Rothman,
 114 2012b):

$$115 \quad g(k, 0) = \frac{1}{k \cdot \sigma \cdot \sqrt{2\pi}} \cdot e^{-(\ln k - \ln \mu)^2 / (2\sigma^2)} \quad (3)$$

116 where $\ln \mu$ is the mean of $\ln k$, and σ^2 is the variance of $\ln k$ (Fig.1D₃). Parameter μ
 117 determines the mean reactivity of the initial OM bulk mixture and parameter σ reflects the
 118 spread of OM components around the mean reactivity.

119 The lognormal distribution is formed by the multiplicative effects of random variables,
 120 which is commonly observed in nature (e.g., the radioactivity of elements in the crust, the
 121 incubation period of infectious diseases, and ecological species abundance) (Limpert et al.,
 122 2001). In the ocean system, the rates of ocean primary production and biological carbon
 123 export also fit the lognormal distribution (Cael et al., 2018). The degradation of OM in
 124 natural ecosystems is controlled by a network of biologically, physically, and chemically
 125 driven processes (Forney and Rothman, 2014), so the variables raised from such
 126 multiplicative processes are often followed by a lognormal distribution. Forney and
 127 Rothman (2012b) showed that litter bag OM incubation data is indeed best described by a
 128 lognormal distribution of rates.

129



130 **Figure 1. Schematic diagram of different OM degradation models.** A: G model, B:
131 RCM, C: Power model and D: Common continuum distribution functions. The x coordinate
132 denotes the variation range of values, and the y coordinate denotes the probability density
133 distribution (ρ) (D₁: the Normal distribution, a typical all-axis distribution, D₂: the Gamma
134 distribution, a typical semi-axis ($x>0$) distribution, and D₃: the Lognormal distribution, a
135 typical semi-axis ($x>0$) distribution).

136

137 In this study, we first compared the l -RCM with other OM degradation models and
138 analyzed the advantages of the l -RCM in describing the OM reactivity distribution. Then
139 we simulated OM degradation in marine sediment at 123 global sites using the l -RCM.
140 Based on inverse modeling results in our dataset, we established the empirical formulas of
141 OM reactivity vs sedimentation rate and further mapped the global OM reactivity
142 distribution. This study provides a new framework for assessing OM reactivity on
143 regional/global scales and predicting OM reactivity in data-poor areas based on easily
144 obtainable environmental parameters (e.g., sedimentation).

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146 **2 Materials and methods**

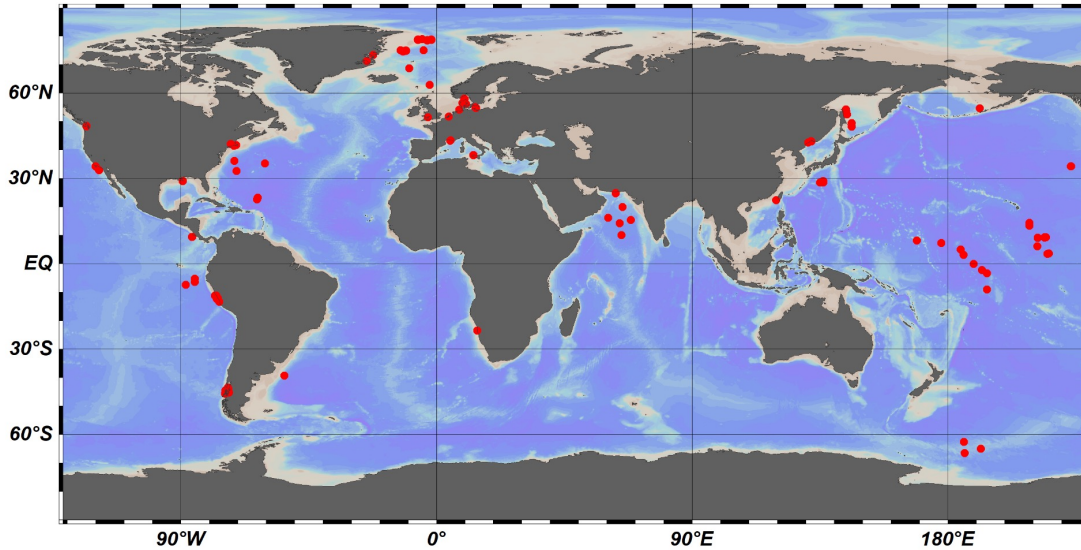
147 **2.1 OM degradation model approach**

148 We constructed an RCM with lognormal distribution (l -RCM) to simulate the OM
149 degradation in marine sediments. The $g(k,0)$ we used in Eq. 1 is the lognormal distribution
150 (Eq.3). Because of the tail of $g(k,0)$, the mean rate constant for bulk OM degradation or the
151 apparent degradation rate of the bulk OM ($\langle k \rangle$) is greater than the median μ , as follows:

$$152 \quad \langle k \rangle = \int_0^{\infty} k \cdot g(k,0) dk = \mu \cdot e^{\sigma^2/2} \quad (4)$$

153 **2.2 Inverse model approach**

154 Here, we used 123 published datasets of OM depth profiles across a wide range of
 155 different depositional environments that have been sourced from published literature
 156 (Middelburg, 1989; Arndt et al., 2013; Middelburg et al., 1997) and the IODP database
 157 (Fig.2, Supplementary Table S1) to inversely determine the μ and σ parameters. We also
 158 analyzed a small number ($n=12$) of laboratory experiment data on OM degradation
 159 (Middelburg, 1989), as well as OM degradation data ($n=16$) from terrestrial soils (Katsev
 160 and Crowe, 2015). We followed the inverse modeling approach by Forney et al.(2012a) to
 161 identify the best-fitting parameters μ and σ based on the Newton method.



162 **Figure 2. Global distribution of investigated sites.**

163
 164
 165 Notably, the burial time was correlated with the porosity. A simple exponential function
 166 was used to describe porosity in sediments:

$$167 \quad \varphi(x) = \varphi_0 \cdot e^{-\lambda x} \quad (5)$$

168 where φ_0 is the values of porosity at the SWI, λ is the attenuation coefficient, and x is
 169 depth. Considering the compaction impacts on OM degradation, the burial time
 170 corresponding to each depth in the OM profile can be calculated as:

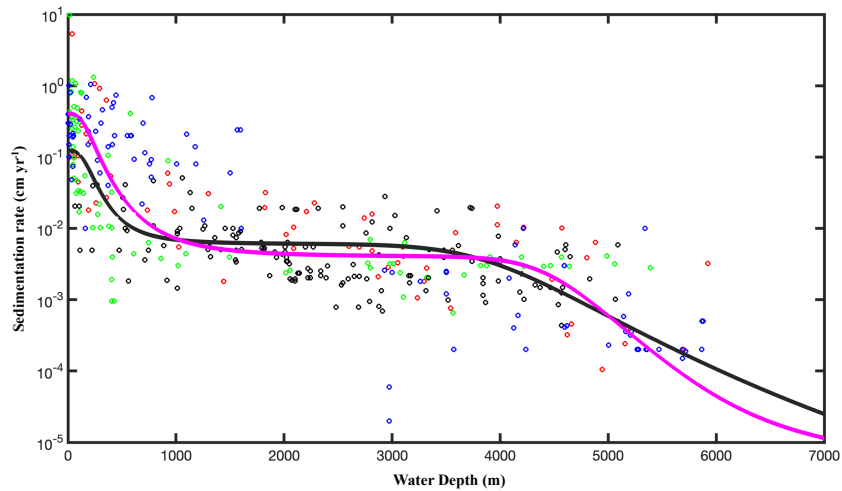
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$$t(x) = \int_0^x \omega^{-1} dx = \frac{x}{\omega_f} + \frac{(\varphi_0 - \varphi_f)}{(1 - \varphi_f) \cdot \lambda \cdot \omega_f} \cdot (e^{-\lambda \cdot x} - 1) \quad (6)$$

172 where φ_f is the values of porosity at larger depths, calculated from Eq. 5 and the pre-set
 173 simulation depth. If the porosity data were not available, the global set as: shelf regions (φ_0 :
 174 0.45, λ : 0.5×10^{-3}), slope regions (φ_0 : 0.74, λ : 1.7×10^{-4}), and abyssal regions (φ_0 : 0.7, λ :
 175 0.85×10^{-3}) (LaRowe et al., 2020b).

176 **2.3 Global upscaling of sedimentation rate**

177 The inversely determined μ , σ couples of all investigated sites were then used in a linear
 178 regression method to derive the empirical relationships between OM parameters μ , σ , $\langle k \rangle$
 179 and the local sedimentation rates (ω). A correction factor was applied to account for the
 180 skewness bias inherent in the back conversion from a log-log transformed linear regression
 181 model to arithmetic units. The newly derived empirical relationships between $\langle k \rangle$ and ω
 182 were then used to calculate global maps of OM reactivity at the SWI on a $1^\circ \times 1^\circ$ grid cell
 183 of the world ocean. At each grid point, ω was estimated based on the empirical relationship
 184 between ω (ω in cm yr^{-1}) and the water depth (z in m) (Eq.7, Fig.3), derived from 260
 185 observations on the global continental shelves (Burwicz et al., 2011), complemented here
 186 by an extra 360 sites including abyss regions (data from Arndt et al. (2013), Egger et al.
 187 (2018)).

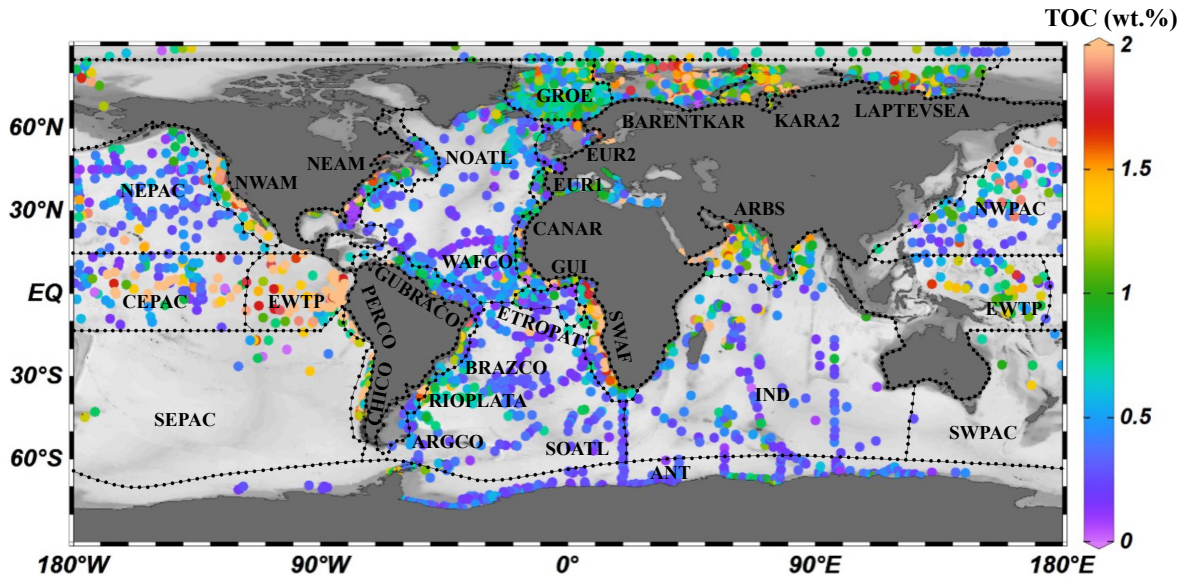
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$$\omega(z) = \frac{0.4}{1 + \left(\frac{z}{200}\right)^{3.5}} + \frac{0.004}{1 + \left(\frac{z}{4500}\right)^{17}} \quad (7)$$



189
 190 **Figure 3. Relationship between Sedimentation rate (w) and water depth (z in m).** The
 191 data are taken from Arndt et al. (2013) (black circles), Egger et al.(2018) (pink circles),
 192 Betts and Holland (1991) (red circles), Colman and Holland (2000) (green circles), and
 193 Seiter et al. (2004) (blue circles). The pink line is the fitting result according to Eq. 7
 194 ($R^2=0.57$), and the black line is the fit obtained from the data of Burwicz et al. (2001)
 195 ($R^2=0.43$).

196

197 Considering the geographic differences in depositional environments and to describe the
 198 global distribution of sedimentary OM reactivity in more detail, we divided the global
 199 ocean into 30 different regions (Table 2, Fig.4) using 5600 single measured data of OM
 200 content in global surface sediment (<5 cm sediment depth) and the previously used
 201 combined qualitative and quantitative geostatistical methods (Seiter et al., 2004).



202

203 **Figure 4. The 30 different regions of the global ocean were divided using 5600 single**
 204 **measured data of OM content (wt.%) of surface sediments**

205

206 **3 Results and discussion**

207 **3.1 OM reactivity distribution described by the γ -RCM and the l -RCM**

208 To compare OM reactivity distribution described by the l -RCM and the γ -RCM, we
 209 determined the best fit to the eight OM datasets reported by Boudreau and Ruddick. (1991).
 210 The results show that both RCMs fit the data equally well, as illustrated by the high
 211 coefficient of determination for each fit ($R^2 > 0.9$, Table 1 and Fig.5). However, the l -RCM
 212 and the γ -RCM differ in their ability to find a unique solution and in their respective
 213 probability density functions of OM reactivity ($\rho(k)$). For example, Fig.6A and 6B show
 214 the best-fit OM profiles for two contrasting sites: BX-6 on the shelf and DSDP 58 in the
 215 abyssal region. The inversely determined parameters at the two sites are $\mu = 2.23 \times 10^{-3} \text{ yr}^{-1}$,
 216 $\sigma = 2.03$ at BX-6, and $\mu = 6.11 \times 10^{-5} \text{ yr}^{-1}$, $\sigma = 1.66$ at DSDP 58 by the l -RCM. At BX-6, the

217 best-fitting parameters by the γ -RCM are $\nu = 0.278$ and $a = 22.5$, and at DSDP 58, $\nu = 1.08$
218 and $a = 20224$. According to the parameter sensitivity analysis, the R^2 of the fitted results
219 remains greater than 0.9 when a and ν change substantially simultaneously (Fig.6D,
220 Supplementary Table S2, Fig.S1, S2, and S3). As a result, different combinations of a and
221 ν can fit the data equally well. For example, simultaneously increasing ν and a ($\nu = 0.5$ and
222 $a = 53$) at site BX-6 or decreasing ν and a ($\nu = 0.5$ and $a = 4024$) at site DSDP 58 lead to a
223 slight change in R^2 . Adding additional measured data, such as depth profiles of porewater
224 sulfate and methane concentrations, can help find a unique solution (Freitas et al., 2021).
225 In contrast, the best-fit parameters μ and σ are unique in the l -RCM, and even small changes
226 in either parameter can lead to abysmal fitting results (Fig.6D). The second difference
227 between the two models concerns the shape of the probability distribution $\rho(k)$. Statistically,
228 the features of the Gamma distribution vary with the value of ν . If $\nu < 1$, $\rho(k)$ tends to positive
229 infinity when k approaches zero. In contrast, if $\nu > 1$, $\rho(k)$ tends to zero when k approaches
230 zero. Hence, the characteristics of the Gamma distribution under different ν values are
231 difficult to visually compare the OM reactivity distributions at site BX-6 ($\nu < 1$) and DSDP
232 58 ($\nu > 1$) (Fig.6C). Compared with γ -RCM, the l -RCM can better distinguish OM reactivity
233 distribution at different sites.

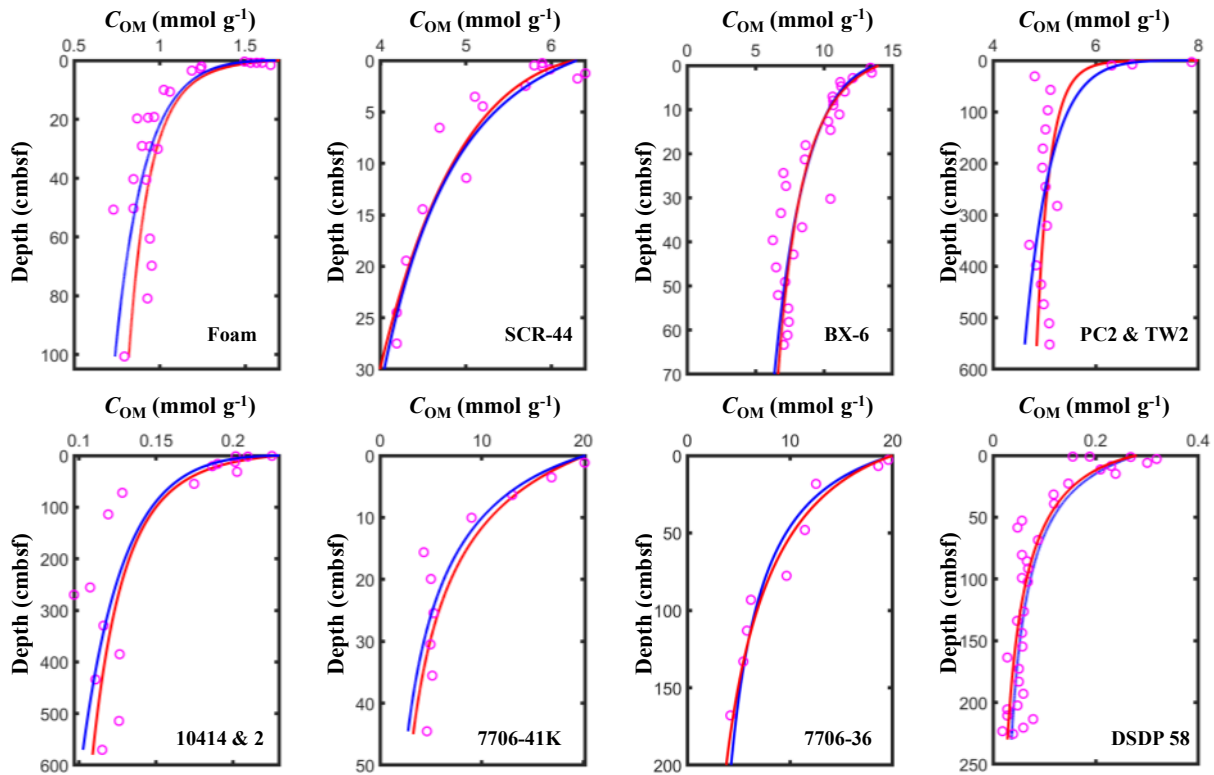
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235 **Table 1. List of model parameters and coefficients of determination (R^2) for the**
236 **fitting result of γ -RCM and l -RCM.**

Core	γ -RCM			l -RCM		
	ν (-)	a (yr)	R^2	μ (yr ⁻¹)	σ (-)	R^2
Foam	0.152	4.2	0.930	2.2×10^{-3}	3.725	0.923
SCR-44	0.202	70.4	0.929	4.4×10^{-4}	2.706	0.922
BX-6	0.278	22.5	0.929	2.24×10^{-3}	2.031	0.936
PC2&TW2	0.052	0.16	0.937	5.5×10^{-5}	6.688	0.947
10141&2	0.193	10184	0.935	1.9×10^{-6}	3.289	0.936
7706-41K	0.910	141.3	0.974	9.5×10^{-3}	0.899	0.972

7706-36	0.804	231.7	0.978	4.79×10^{-4}	1.089	0.980
DSDP58	1.080	20224	0.917	6.11×10^{-5}	1.663	0.921

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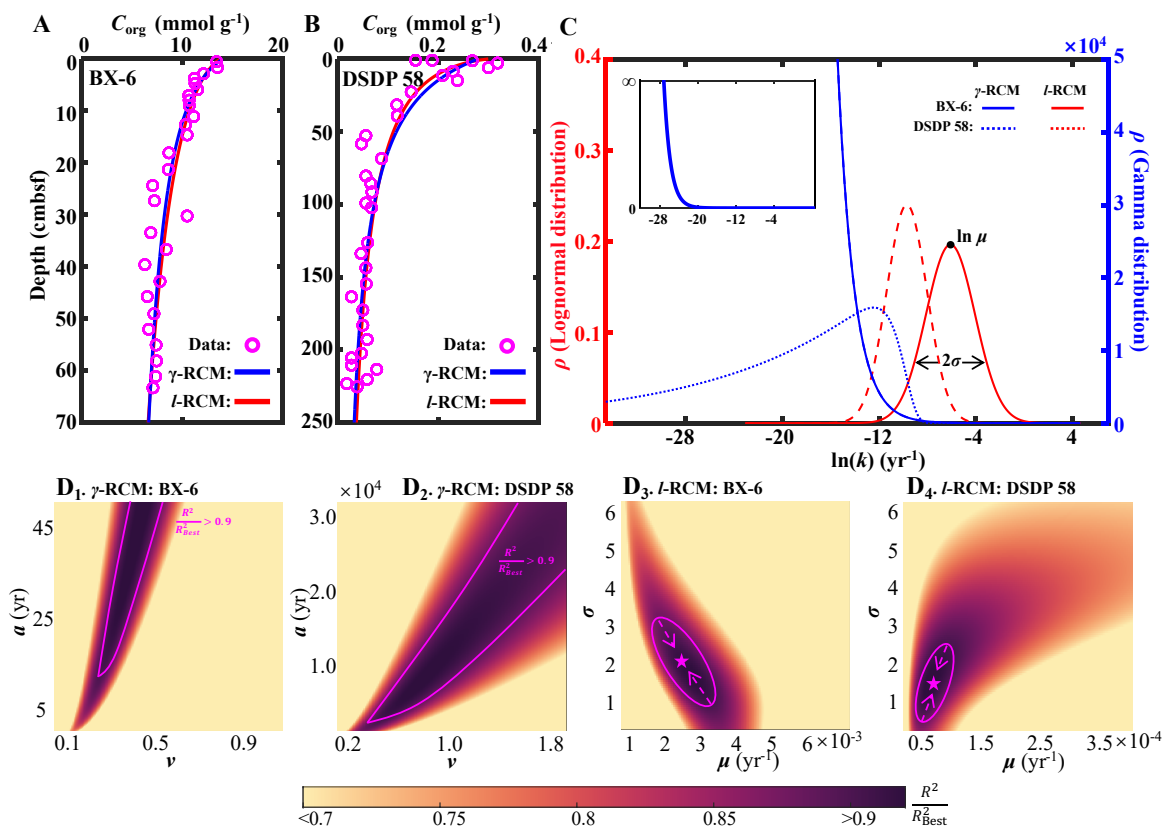
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Figure 5. Fitting results of the l -RCM and the γ -RCM. The pink dots are the measured

240

OM data, the red lines are l -RCM fitting results, and the blue lines are γ -RCM fitting results.

241



242

243 **Figure 6. Comparison of *l*-RCM and γ -RCM.** **A, B:** the fitting results of the *l*-RCM and
 244 the γ -RCM for site BX-6 and DSDP 58. **C:** OM reactivity distribution from *l*-RCM and γ -
 245 RCM. Top inset, Gamma distribution at site BX-6 at a larger y-axis. **D:** Distribution of
 246 R^2/R_{Best}^2 for parameter sensitivity analysis of the γ -RCM and the *l*-RCM at sites BX-6 and
 247 DSDP 58. The pink lines in the D₁ and D₂ denote the range that $R^2/R_{\text{Best}}^2 > 0.9$ in the γ -RCM.
 248 The R^2/R_{Best}^2 in the *l*-RCM converges as the pink arrows in the D₃ and D₄, ultimately
 249 reaching the best fitting results as the pink pentagrams.

250

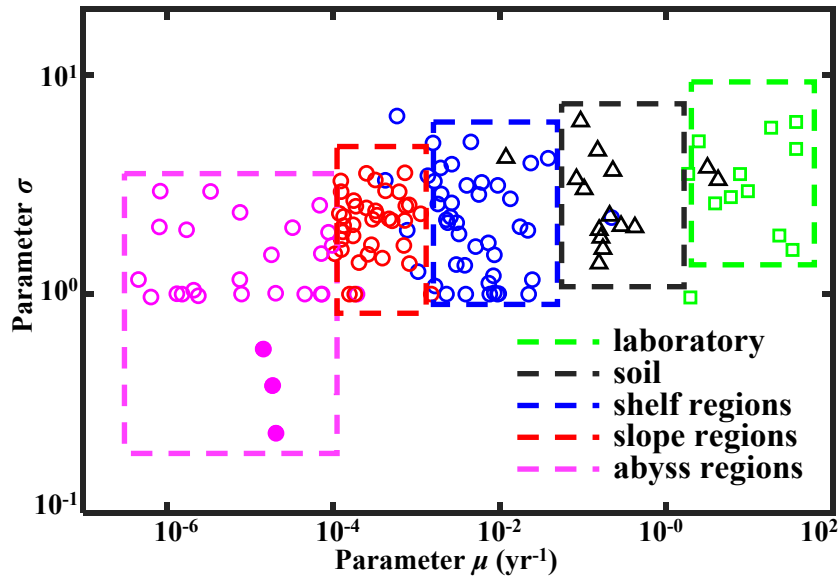
251 3.2 Regional distribution of OM reactivity

252 In the *l*-RCM, parameter μ represents the mean reactivity of the OM fractions, which
 253 dominates the rate of OM degradation (Supplementary Fig.S2), and parameter σ describes
 254 the homogeneity of OM fractions, with larger σ value indicating more heterogeneous
 255 mixture of OM (Forney et al., 2012b). The inverse determination of the *l*-RCM parameters

256 μ and σ across the wide range of different depositional environments allows quantitative
257 insights into OM reactivity and provides essential information on the main environmental
258 controls on OM reactivity. Fig.7 illustrates the inversely determined μ - σ for all 123 depth
259 profiles of marine sediment POC investigated in this study and compares them with
260 inversely determined parameters from published soil and laboratory incubation data. It
261 highlights the large inter- and intraregional variability of best-fit μ (10^{-6} – 10^2 yr⁻¹) and σ
262 (0.2–6). However, despite the large variability, it also reveals broad global patterns in μ
263 and σ . Notably, best-fit μ - σ couples form environmental clusters along a μ gradient, with
264 the highest μ being determined for laboratory degradation experiments of fresh
265 phytoplankton (Garber, 1984; Westrich and Berner, 1984) ($\mu=10^0$ – 10^2 yr⁻¹), followed by
266 soil incubation under natural (Katsev and Crowe, 2015), yet still idealized conditions
267 ($\mu=10^0$ – 10^1 yr⁻¹), while OM degraded in marine sediments generally reveals lower
268 inversely determined $\mu < 10^0$ yr⁻¹. The higher μ values determined for soil OM seemingly
269 contradict the widely accepted notion that soil OM is generally less reactive than marine
270 OM (Larowe et al., 2020a; Zonneveld et al., 2010). However, this apparent contradiction
271 can be explained by the idealized conditions of the incubation experiments (e.g., only one
272 type of material, some of which had nitrogen added), as well as the degradation state of the
273 investigated OM. Although soil OM is structurally less reactive (Hedges and Keil, 1995;
274 Zonneveld et al., 2010), the soil incubation experiments were conducted with initially
275 undegraded material. In contrast, OM deposited in marine sediments consists of a complex
276 mixture of OM from autochthonous and allochthonous sources that is altered to various
277 degrees during transit from its source to the sediment (Hewson et al., 2012).

278 In addition to the difference between incubation data and field observations, Fig.7 also
279 reveals a three order of magnitude decrease in inversely determined μ for OM from the
280 shelf (10^{-3} – 10^{-1} yr⁻¹) to the slope (10^{-4} – 10^{-3} yr⁻¹), and ultimately abyssal regions ($<10^{-4}$ yr⁻¹).
281 In addition, shelf and slope regions also generally reveal a larger σ (1–3), while abyssal
282 regions display a narrower σ range (0.5–1). This observed progressive decrease in μ and σ
283 from the shelf to the abyssal ocean confirms previously observed patterns (Arndt et al.,
284 2013; Freitas et al., 2021; Zonneveld et al., 2010) and reflects the interaction between OM
285 structure (or its source) and the degree of alteration/pre-processing as OM transits from its
286 original source to the ultimate sedimentary sink. In the dynamic shelf regions, highly
287 variable OM loads from different sources, including *in-situ* produced marine OM, laterally
288 transported, pre-processed terrestrial or marine OM, are often physically protected from
289 further erosion/deposition cycles due to high suspended sediment loads (Arndt et al., 2013;
290 Larowe et al., 2020a). As a result, benthic OM is composed of a complex mixture of fresh
291 and pre-aged compounds of highly variable (hence larger σ of the initial distribution), yet
292 generally higher reactivity. On the upper and mid-continental slopes, intensive lateral
293 and/or vertical transport processes or the abrupt relocation of sediment result in similar
294 complex mixtures of OM (hence similar σ of the initial distribution) (Larowe et al., 2020a).
295 However, transport timescales are often longer due to the greater water depths and distance
296 to land. The deposited OM is generally more degraded and thus less reactive than in shelf
297 environments. In contrast, benthic OM in abyssal regions is mainly derived from marine
298 production (Rowe and Staresinic, 1979; Larowe et al., 2020a). During its slow settling
299 through the water column, highly reactive OM compounds are rapidly degraded, and only
300 the less reactive compounds persist and settle onto the sediment (Dunne et al., 2007). The

301 values of μ and σ in the abyssal regions are thus significantly smaller than in the shelf and
 302 slope regions. The decrease of μ and σ from the shelf to abyssal regions reveals a decline
 303 in reactivity during lateral transport of OM, where μ mainly controls the overall reactivity
 304 and σ indicates the coverage of the main component of OM.



305
 306 **Figure 7. Regional distribution of OM reactivity.** Distribution of parameters σ and μ in
 307 different regions. Pink solid circles denote fitting results of sites in the NEPAC with
 308 extremely low OM reactivity.

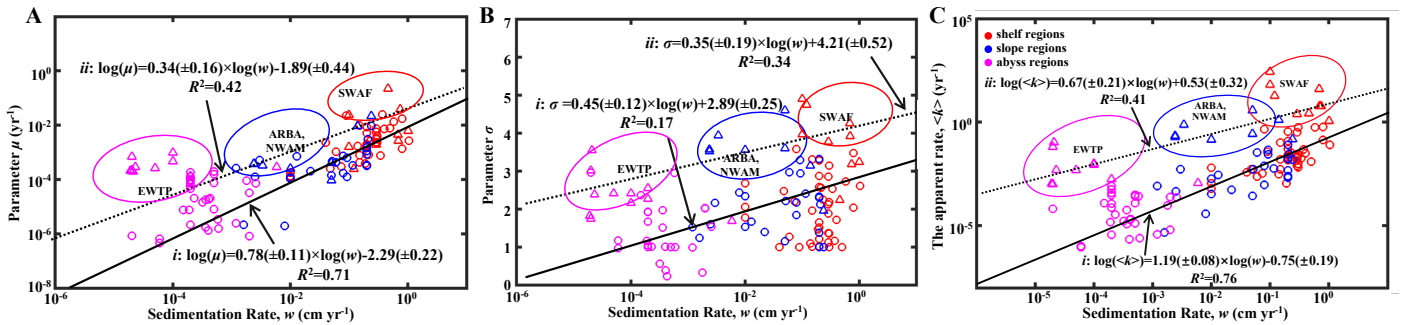
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310 3.3 Global distribution patterns of OM reactivity

311 Parameters μ and σ together control the degradation process of OM, which can be further
 312 described by the apparent degradation rate of the bulk OM ($\langle k \rangle$). Sedimentation rate (ω)
 313 is a widely observed and comparably easy to measure proxy for local depositional
 314 conditions with sizable global data sets or empirical formulas available (Burwicz et al.,
 315 2011). Fig. 8A, 8B and 8C show the global decreasing trend of μ , σ and $\langle k \rangle$ with ω for the
 316 general sea regions (shelf (<200m), slope (200–2000m), and abyss (>2000m)). The active

317 OM fractions (e.g., sugars and proteins) are preferentially exhausted during the lateral
318 transport of OM from the shelf to the abyssal regions, leading to a decrease in the mean
319 OM reactivity (μ , Fig. 8A), and thus OM is mainly composed of refractory components (σ ,
320 Fig. 8B). Due to the multiple sources of OM in the shelf regions, including fresh and older
321 OM imported laterally by inland rivers, and OM settled from the euphotic layer (LaRowe
322 et al., 2020a), the values of the values of μ , σ and $\langle k \rangle$ fluctuates significantly. However,
323 the general trend is superimposed by a large variability and apparent reactivity $\langle k \rangle$ in
324 specific environments, notably deviating from this generally observed trend. More
325 specifically, higher μ and σ values and, thus, higher OM reactivities occur in the Eastern-
326 Western Coastal Equatorial Pacific (EWEP), Southwestern-Africa continental margin
327 (SWAF), Northwestern-America continental margin (NWAM), and the Arabian Sea
328 (ARBS) regions. These results are completely consistent with prior observations and model
329 results (Arndt et al., 2013) and can be directly linked to the prevailing water-column redox
330 and depositional conditions. High benthic OM reactivities have previously been reported
331 for depositional environments that are characterized by a dominance of marine algal OM
332 (Hammond et al., 1996) and strong lateral transport processes (e.g., SWAF, NWAM)
333 (Arndt et al., 2013). Consequently, the larger values of all μ and σ , and $\langle k \rangle$ occur in the
334 inverse modelling results for these depositional environments (Fig. 8). Furthermore, the
335 reactivity of sedimentary OM is considerably influenced by oxygen content or more
336 precisely, by oxygen exposure time in the water column and at the seafloor (Aller, 1994;
337 Hartnett et al., 1998; Hedges and Keil, 1995; Mollenhauer et al., 2003; Zonneveld et al.,
338 2010). Lower oxygen concentrations, as present in these regions in the form of pronounced
339 oxygen minimum zones (OMZs), will slow down the degradation of OM both in the water

340 column and at the sediment surface (Jørgensen et al., 2022). This enables the burial of more
 341 reactive OM into the sediments and thus results in the occurrence of high sedimentary OM
 342 reactivity in these regions despite great water depth (e.g., ARBS, EWTP) (Arndt et al.,
 343 2013; Bogus et al., 2012; Ingole et al., 2010; Luff et al., 2000; Volz et al., 2018). The *l*-
 344 RCM not only captures the broad patterns of OM reactivity across the global seafloor even
 345 better than previous models, but also provides statistically more significant relationships
 346 between OM reactivity ($\langle k \rangle$) and sedimentation rate (ω) than inversely determined
 347 parameters of γ -RCM ($R^2 < 0.46$) and discrete models ($R^2 < 0.1$) (Arndt et al., 2013).
 348 Considering that no robust quantitative framework exists at this stage to predict OM
 349 reactivity as a function of easily observable environmental parameters, the *l*-RCM provides
 350 an excellent first-order predictor and a step forward in assessing the global distribution
 351 patterns of OM reactivity, despite the poor relationship between $\langle k \rangle$ and ω for these special
 352 regions (e.g., EWTP, SWAF, NWAM, and ARBS).



354 **Figure 8. Global distribution patterns of OM reactivity.** A. Log-log plot of ω and μ . B.
 355 Log-log plot of ω and σ . C. Log-log plot of ω and $\langle k \rangle$. The solid black line (i) denotes
 356 linear regression for shelf, slope, and abyssal regions. The black dotted line (ii) denotes
 357 linear regression for high OM reactivity regions, including the EWTP, ARBS, NWAM,
 358 and SWAF regions.

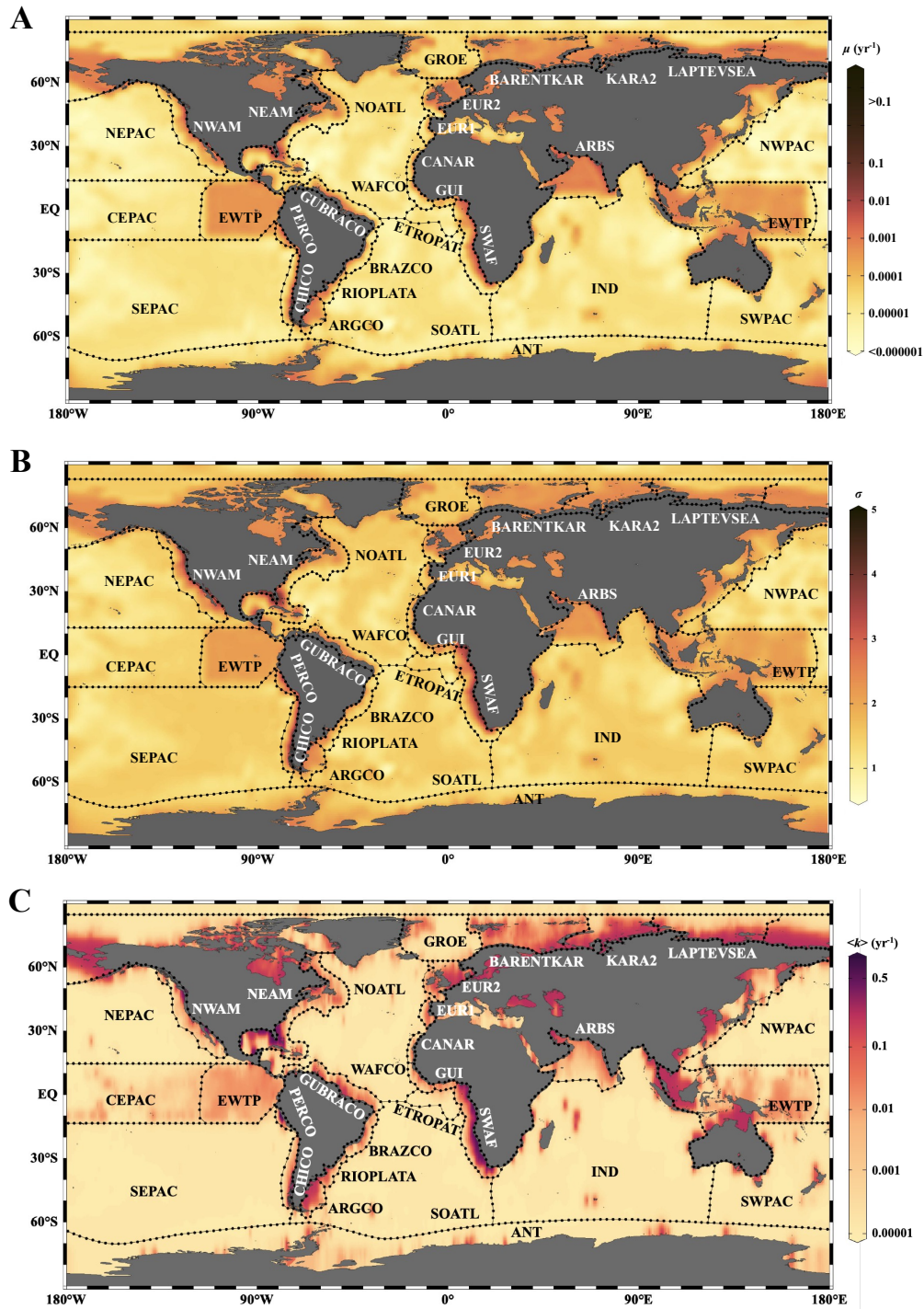
359

360 Based on the empirical relationships in Fig.8 (*i* for the general water depth-related
361 regions, *ii* for the specific regions (EWTP, ARBS, NWAM, and SWAF)), and the water
362 depth– ω relationship (Eq.5), we finally derived, to our knowledge, the world’s first map
363 of the global distribution of parameter μ , σ , and $\langle k \rangle$ (Fig.9). Using the relationship between
364 water depth, ω , and $\langle k \rangle$ (Fig.3 and Fig.8C), we further estimated the mean apparent OM
365 reactivity ($\langle K_{\text{region}} \rangle$) in the 30 regions of global ocean (Table 2). Furthermore, the
366 heterogeneity of the OM reactivity distribution in global marine sediments is well
367 illustrated in Fig. 9. Specifically, higher μ (Fig.9A), σ (Fig.9B), and OM reactivity (Fig.9C)
368 is reflected in shelf regions, particularly in northern Atlantic provinces with high latitudes
369 (e.g., Barents Sea ($\langle K_{\text{region}} \rangle \approx 0.02 \text{ yr}^{-1}$), Laptev Sea ($\langle K_{\text{region}} \rangle \approx 0.03 \text{ yr}^{-1}$), and Kara Sea
370 ($\langle K_{\text{region}} \rangle \approx 0.01 \text{ yr}^{-1}$)), due to shallower water depths and high OM fluxes from inland
371 (Burwicz et al., 2011; Seiter et al., 2004). Besides that, the global map also highlights the
372 extremely low OM reactivity, especially in some regions, as indicated by the absence of
373 sulfate-methane transition (SMT) (e.g., the NE-Pacific, NEPAC) (Eggert et al., 2018) and
374 central ocean gyre regions (e.g., South Pacific Gyre) (LaRowe et al., 2020b). Deeper water
375 depth (>5000m), relatively low OM content (~0.2wt.%), and the old OM age (>10⁴ years)
376 result in comparably lower μ and σ values (Fig.9A and 9B) and, thus, extremely low benthic
377 OM reactivity ($\langle K_{\text{region}} \rangle \approx 10^{-4} \text{ yr}^{-1}$) (Kallmeyer et al., 2012, Müller and Suess, 1979).
378 Normally, greater water depth enhances oxygen exposure time for OM degradation, and
379 thereby reduce the reactivity of OM arriving at the seafloor, as reflected in the smaller μ
380 values (Fig. 9A). In ocean areas characterized by pronounced OMZs, however, due to
381 strong coastal upwelling or a high export rate of plankton-derived OM, the inhibition of
382 OM degradation processes in the water column results in the preservation of

383 heterogeneously mixed OM components (both active and refractory), as reflected in the
384 larger σ values (Fig. 9B), leading to higher than expected OM reactivity in specific regions
385 despite greater water depths (e.g., ARBS and EWTP ($\langle K_{\text{region}} \rangle \approx 0.01 \text{ yr}^{-1}$) (Fig. 9C). Thus,
386 the *l*-RCM provides a new framework not only for identifying the differences in OM
387 reactivity between regions, but also for assessing regional/global OM reactivity patterns
388 using easily obtainable information (e.g., sedimentation).

389 OM reactivity exerts an important control on the relative significance of OM degradation
390 pathways in marine sediments. In oxic environments, OM will be mainly respired
391 aerobically and through denitrification, whereas deeper within the sediment, it will mainly
392 be decomposed through anaerobic pathways such as sulfate reduction and methanogenesis
393 (Regnier et al., 2011). Therefore, further work should be conducted to simulate the
394 associated biogeochemical processes using the *l*-RCM to better quantify OM degradation
395 and burial in marine sediments on regional or global scales.

396



397

398 **Figure 9. Distribution of μ (A), σ (B), and $\langle k \rangle$ (C) in the global ocean with $1^\circ \times 1^\circ$**
 399 **resolution.**

400

401 **Table 2. Abbreviations of regions in this paper (Seiter et al., 2004), and their area,**
 402 **mean water depth, mean OM content in surface sediment (<5 cm), and apparent OM**
 403 **degradation rata ($\langle K_{\text{region}} \rangle$).**

Abbreviation	Region	water depth ^a (m)	Mean OM (wt.%)	$\langle K_{\text{region}} \rangle$ (yr ⁻¹)
SWAF	SW-Africa continental margin	334	2.5	0.48542
NWAM	NW-America continental margin	731	1.7	0.12695
ARBS	Arabian Sea	1600	1.4	0.08182
EWTP	East-West Coastal Equatorial Pacific	3662	1.2	0.01587
ANT	South Polar Sea	1300	0.3	0.00029
ARGCO	Argentina continental margin	1859	0.3	0.00026
BARENTKAR	Barents Sea and Kara Sea	224	1.1	0.02081
BRAZCO	Brazil continental margin	1051	0.5	0.00034
CANAR	Canaries	1190	0.6	0.00031
CEPAC	Central Equatorial Pacific	5022	0.3	0.00002
CHICO	Chile continental margin	1444	1.5	0.00028
ETROPAT	Eastern tropical Atlantic	2253	0.7	0.00026
EUR1	N-European continental margin	1290	0.8	0.00029
EUR2	S-European continental margin	974	0.3	0.00037
GROE	Northern Nordic Sea	1563	0.7	0.00027
GUBRACO	SE-America continental margin	1844	0.4	0.00026
GUI	Gulf of Guinea	1586	1.1	0.00027
INA	Indian Ocean deep sea	4042	0.4	0.00021
KARA2	Kara Sea	281	1.2	0.01111
LAPTEVSEA	Laptev Sea	190	0.9	0.02964
NEAM	NE-America continental margin	1045	0.9	0.00034
NEPAC	NE-Pacific	4463	0.4	0.00012
NOATL	Northern Atlantic	2161	0.4	0.00026
NWPAC	NW-Pacific	4898	0.6	0.00004
PERCO	Peru continental margin	1020	4.8	0.00035
RIOPLATA	Rio de la Plata mouth	1784	0.8	0.00026
SEPAC	SE-Pacific	3952	0.5	0.00022
SOATL	Southern Atlantic	3592	0.4	0.00024
SWPAC	SW-Pacific	3153	0.8	0.00025
WAFCO	W-Africa continental margin	1982	0.6	0.00026

404 ^awater depth and mean OM content are based on the average depth and OM content of the
 405 sites in each region of Fig.4.

406

407 **4 Conclusions**

408 Compared with previous OM degradation models, the *l*-RCM presented here not only
409 well fits OM depth-content profiles, but also better represents the distribution of OM
410 reactivity by the parameters μ and σ . We use the *l*-RCM to inversely determine μ and σ at
411 123 sites across the global ocean, including shelf, slope, and abyssal regions. Our results
412 show that the apparent OM reactivity ($\langle k \rangle = \mu \cdot \exp(\sigma^2/2)$) decreases with decreasing
413 sedimentation rate (ω), and that OM reactivity is more than three orders of magnitude
414 higher in shelf than in abyssal regions. Due to the complex depositional environments (e.g.,
415 oxygen minimum zones), OM reactivity is higher than predicted in some specific regions
416 (e.g., the NWAM, SWAF, ARBS, and EWTP), which was also captured by the *l*-RCM in
417 these regions. Based on two empirical relationships between the OM reactivity ($\langle k \rangle$) and
418 sedimentation rate (ω), we obtained the global OM reactivity distribution patterns and
419 finally mapped the global OM reactivity distribution. The reactivity of OM serving as fuel
420 for microbial activity in marine sediments firmly controls the degradation pathways and
421 metabolism rates. Thus, the *l*-RCM has direct implications on the constraints for OM
422 degradation and burial in marine sediments on regional or global scales.

423

424 **References**

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614

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623 **Author contributions**

624 S.X. and B.L. designed the study and performed the research with S.A., S.K., and Z.W.;

625 All authors discussed the results and commented on the manuscript.

626 **Competing interests**

627 The authors declare that they have no competing interests.