Characterizing the marine iodine cycle and its relationship to ocean deoxygenation in an Earth System model

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8 Abstract. Iodine abundance in marine carbonates (measured as an elemental ratio with calcium - I:Cal/Ca) is of 9 broad interest as a proxy for local/regional ocean redox. This connection arises because the speciation of iodine in 10 seawater <u>— in terms of</u> the balance between iodate (IO_3) and iodide (I) — is sensitive to the prevalence of oxic 11 vs. anoxic conditions. However, although I:CaI/Ca ratios are being increasingly commonly being measured in 12 ancient carbonate samples, a fully quantitative interpretation of this proxy is hindered by the scarcity of requires the 13 availability of a mechanistic and quantitative interpretative framework for the marine iodine cycle and that can 14 account forits sensitivity to the extent and intensity of ocean deoxygenation in the past. Here we present and 15 evaluate a representation of marine iodine cycling embedded in an Earth system model ('cGENIE') against both modern and paleo observations. In this framework, we account for IO_3^- uptake and reduction-release of I_2^- through the 16 17 biological pumpby primary producers, the occurrence of reduction of ambient IO_3 -reduction to I' in the water 18 column, plus the re-oxidation of I⁻ to IO₃⁻. We develop and test a variety of different plausible mechanistic 19 mechanisms for iodine reduction and oxidation transformation relationships between IO3- and I-and contrast model 20 projections against an updated compilation of observed dissolved IO₃⁻ and I⁻ concentrations in the present-day ocean. 21 In By optimizing the parameters controlling previously proposed mechanisms behind involved in marine iodine 22 cycling, we find that we can obtain broad matches to observed iodine speciation gradients in zonal surface 23 distribution, depth profiles, and oxygen deficient zones (ODZs). We-However, we also identify alternative, equally 24 well performing mechanisms which assume a more explicit mechanistic link between iodine transformation and 25 environment - an. This mechanistic ambiguity that highlights the need for more process-based studies on modern 26 marine iodine cycling. Finally, to help distinguish between competing representations of the marine iodine cycle 27 and, because our ultimate motivation is to further our ability to reconstruct ocean oxygenation in the geological past, 28 we conducted 'plausibility tests' of our various different model schemes against available I:CaI/Ca measurements 29 made on Cretaceous carbonates - a time of substantially depleted ocean oxygen availability compared to modern 30 and hence a strong test of our model. Overall, the simultaneous broad match we can achieve between modelled 31 iodine speciation and modern observations, and between forward-proxy modelled I:CaI/Ca and geological elemental 32 ratios, supports the application of our Earth system modelling in simulating the marine iodine cycle to help interpret 33 and constrain the redox evolution of past oceans. 34

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35 1. Introduction

36 Dissolved Iodine (I) in seawater is redox sensitive and as such, is a potential invaluable delineator of past ocean 37 deoxygenation. This arises directly from: (1) observations that the oxidized form of iodine (iodate, (IO3) is reduced 38 to iodide (I) under low oxygen conditions, and (2) that IO_3 in seawater is incorporated into carbonate lattice during 39 precipitation in proportion to its seawater abundance (whilst I' is not) (Lu et al., 2010; Podder et al., 2017; Kerisit et 40 al., 2018; Zhang et al., 2013; Hashim et al., 2022). Hence As a result, past ocean IO₃⁻ concentrations can be recorded 41 in coeval carbonates as I:Cal/Ca ratios, with the potential of for carbonate I:Cal/Ca to reflect the redox variation of 42 the ancient seawater (Lu et al., 2010). Indeed, the I:CaI/Ca ratio in marine carbonates is already widely being applied 43 widely as a paleoredox proxy, with studies employing it to explore the-variations in the dissolved oxygen ([O₂]) 44 concentration of seawater-variation throughout much of Earth history, from the Archean and through the Cenozoic 45 (Lu et al., 2010; Hardisty et al., 2014; Zhou et al., 2015; Lu et al., 2016; Edwards et al., 2018; Lu et al., 2018; Bowman 46 et al., 2020; Pohl et al., 2021; Wei et al., 2021; Ding et al., 2022; Shang et al., 2019; Liu et al., 2020; Fang et al., 2022; 47 Uahengo et al., 2020; Yu et al., 2022; Tang et al., 2023). The-However, realizing the full potential for H:Cal/Ca to 48 generate provide critical insights into how the oxygenation of the ocean has evolve through time, as well as the causes 49 and biological/ecological consequences of this, requires that we have an adequate understanding, not only of carbonate 50 IO₃⁻ incorporation, but of the <u>dynamics of the</u> marine iodine cycle in general.

51 Progress Considerable progress has been made over the past few decades towards the goal of understanding 52 the marine iodine cycle-in the past decades. Iodine has a relatively long residence time_in the ocean (~300 kyr; 53 Broecker and Peng, 1983), making leading to its concentration among being relatively constant throughout the global 54 ocean relatively constant(at_around 500_nM) (Elderfield and Truesdale, 1980; Truesdale et al., 2000; Chance et al., 55 2014). Although However, although the total concentration of dissolved iodine is relatively invariant, the two most 56 abundant species of dissolved iodine in the ocean, IO3⁻ and iodide I⁻, vary relative to each other depending on the 57 environment.- Today, IO3- is generally the dominant iodine species in oxygenated regions of the ocean, where it 58 representing represents total iodine nearly quantitatively below the euphotic zone. Within the euphotic zone, the 59 concentration of I⁻ occurs and generally increases in association with release during phytoplankton growth and 60 senescence (Hepach et al., 2020). Iodide is also more abundant in oxygen deficient zones (ODZ) often, but not 61 always quantitatively so (Truesdale et al., 2000; Rue et al., 1997; Cutter et al., 2018; Moriyasu et al., 2020; Farrenkopf 62 and Luther, 2002; Wong and Brewer, 1977; Truesdale et al., 2013; Rapp et al., 2020, 2019). Respectively, wWithin 63 the ODZ_S, IO₃⁻ is reduced to I⁻<u>-but not always quantitatively so</u>-and hence is present only at relativelyhas low 64 concentrations while I is abundant (Truesdale et al., 2000; Rue et al., 1997; Cutter et al., 2018; Moriyasu et al., 2020; 65 Farrenkopf and Luther, 2002; Wong and Brewer, 1977; Truesdale et al., 2013; Rapp et al., 2020, 2019)(Rue et al., 66 1997; Farrenkopf et al., 1997; Moriyasu et al., 2020; Rapp et al., 2019, 2020).

67 Although 10_3 is generally depleted in low- $[O_2]$ settings, the causal spatial relationship between seawater $[O_2]$ 68 and $[IO_3]$ is not simple and is <u>not</u> currently not-well understood. Recently published observations from <u>global oxygen</u> 69 deficient zones (ODZ_S) in the ocean reveals that the relationship between dissolved $[O_2]$ and $[IO_3]$ is not linear, but 70 instead it is possible that there is a certain $[O_2]$ or related redox threshold <u>associated with</u> triggering IO_3^- reduction 71 (Cutter et al., 2018; Moriyasu et al., 2020; Farrenkopf and Luther, 2002; Rue et al., 1997; Chapman, 1983). Formatted: Font: 12 pt

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Dissimilatory IO3⁻ reducing bacteria, as well as abiotic reduction with sulfide and dissolved Fe, have been identified 72 within ODZs (Farrenkopf et al., 1997; Councell et al., 1997; Jiang et al., 2023). In addition, slow oxidation-reduction 73 kinetics (Tsunogai, 1971; Hardisty et al., 2020; Schnur et al., 2024) imply the likelihood that in situ iodine signals 74 could be integrated across large-scale physical oceanography processes - including ocean currents and mixing 75 between water masses (Hardisty et al., 2021), and meaning that iodine species speciation reflects regional local regional 76 77 rather than strictly local-in-situ redox conditions (Lu et al., 2020b). Non-redox related processes, such as 78 phytoplankton-mediated IO3⁻ reduction and organic matter remineralization also exerts controls on iodine speciation 79 in the water column (Fig. 1; Elderfield and Truesdale, 1980; Wong et al., 1985; Luther and Campbell, 1991; Hepach 80 et al., 2020). Therefore, it is difficult to infer water column redox simply based on iodine speciation without 81 considering these interacting physical and biological effects.





Figure 1: The iodine cycle in marine oxygen deficient zones (ODZ) in cGENIE <u>including (1) IO₃' reduction to</u> <u>I', (2) re-oxidation of I' to IO₃', (3) photosynthetic IO₃' update by phytoplankton, and (4) I' releasement from</u> organic matter (OM) through remineralization. The <u>detailed</u> oxidation-reduction options ('threshold' and 'lifetime'/ 'reminO2lifetime'/ <u>''Fennel'</u>) are described in Section 2.2 and Table 1. Dashed arrows indicate variable processes during ensemble simulations. Note that the POI export is controlled by temperature (TDEP).

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92 Apart-Aside from the uncertainties associated with IO_3^- reduction, it is notable that the oxidants responsible 93 for IO_3^- formation during the <u>1</u> re-oxidation 1 are currently unknown, only that it is unlikely to be <u>free</u> O_2 , which is 94 not thermodynamically favorable to oxidize iodidne (Luther et al., 1995). A recent thermodynamic review indicates 95 that the reactive oxygen species (ROS) such as hydrogen peroxide and OH radicals can fully oxidize I^{\cdot} to IO₃^{\cdot}. Iodide 96 oxidation to IO3⁻ is a 6-electron transfer and other ROS, such as superoxide, are only thermodynamically favorable to 97 catalyze partial oxidation to intermediates (Luther, 2023). These ROS species have heterogenous distributions and 98 ambient ocean concentrations that are typically relatively low compared to iodine, supporting the likelihood of 99 temporally or spatially isolated high I⁻ oxidation rates despite of overall extremely slow rates (Schnur et al., 2024). 100 Additional support for spatially or temporally heterogenous I⁻ oxidation rates comes from recent experimental 101 observations of IO3⁻ production from I⁻ in nitrifying cultures (Hughes et al., 2021). Nitrification (oxidation of NH₄⁺ to 102 <u>NO₂ (and NO₃)</u> rates vary globally, with the highest values occurring in ODZs and the dissolved deep chlorophyl 103 maximum (summarized in Table 2 of Hughes et al., 2021). Regardless, nitrification or other specific mechanisms have 104 yet to be linked directly to I oxidation under normal marine conditions, leaving open the question of rates and locations 105 of I- oxidation.

106 Despite a growing understanding of I:Cal/Ca variations through geologic time, it remains challenging to 107 determine the mechanisms driving responsible for controlling the spatiotemporal patterns of marine [IO3-] and the 108 degree thathow these are linked to seawater oxygen and there have only been a few attempts to date to model the 109 marine iodine cycle. Given the prevailing uncertainty in the mechanisms governing the marine iodine cycle mentioned 110 above, and in conjunction with I:Cal/Ca being a relatively new proxy, it is perhaps not surprising that few attempts 111 have been made to model the marine iodine cycle. In a recent publication, a model was developed to simulate modern 112 ocean surface I distributions, with the aim of being able to improve tropospheric ozone models (Wadley et al., 2020). 113 This particular model was based around a relatively high horizontal ocean resolution (1° grid size) with a 3-layer 114 vertical upper water column. Iodine biogeochemical cycling was coupled with the nitrogen cycle, with the surface I 115 distribution sensitive to biological and hydrological factors including primary productivity, I:C ratio, oxidation, mixed 116 layer depth, advection, and freshwater flux. Because the Wadley et al., (2020) model was specifically focused on near-117 surface processes within the upper 500 m, it did not consider processes occurring within ODZs and hence is not 118 directly applicable to questions concerning the controls on I:Cal/Ca ratios. In contrast, a second model-based study 119 deliberately targeted paleoceanographic questions and incorporated an iodine cycle including redox-controlled 120 biogeochemical reactions into the 'cGENIE' Earth system model (Lu et al., 2018). The advantage for paleo studies 121 afforded by this particular approach is that the cGENIE model can take into account different continental 122 configurations, non-modern atmospheric composition (pO2, pCO2), and other boundary conditions that may have

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differed on ancient Earth relative to today (Ridgwell et al., 2007; Reinhard et al., 2016; Boscolo-Galazzo et al., 2021;
Remmelzwaal et al., 2019; Pohl et al., 2022; Reinhard and Planavsky, 2022).

125 Despite a growing understanding of I:Cal/Ca variations through geologic time, it remains challenging to 126 determine mechanisms driving spatiotemporal marine [IOx⁻] and the degree that these are linked to seawater oxygen. 127 Hence, the proxy is qualitative or semi-quantitative. Here, we calibrate the iodine cycle within the cGENIE Earth 128 System model to provide a mechanistic framework for interpreting ancient I:CaI/Ca variations. In this study, we build 129 on the work of Lu et al., (2018) and further develop and test a series of new potential parameterizations for water 130 column iodine oxidation and, reduction, cellular uptake, and release during remineralization (in addition to reduction 131 and transport associated with the biological pump). We also developed 3 criteria for assessing the model: (1) Statistical 132 133 location-dependent comparisons between the model and an iodine ocean observation data compilation. (2) Graphical 134 comparison of modeled and observed iodine across 3 illustrative iodine speciation gradients (depth profiles from 135 multiple ocean basins, latitudinal transects of surface waters, and across transects of the Eastern Tropical North Pacific 136 oxygen minimum zone (Moriyasu et al., 2020)). (3) Model applicability to ancient settings by comparing (also using 137 the M-score) projections of ocean surface I:CaI/Ca with published I:CaI/Ca data from the Cretaceous (Zhou et al., 138 2015).

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140 2. Model Description

141 2.1 The cGENIE Earth system modelling framework

142 cGENIE is a class of model known as an 'Earth system model with intermediate complexity' (EMIC)____a global 143 climate-carbon cycle model that simplifies one or more (typically physical elimaticclimate) components of the Earth 144 system. In the case of cGENIE, ocean circulation is solved for on a relatively low-resolution grid (here: an equal area 145 36×36 grid, which equates to 10° in longitude and latitude increments from 3° near the equator to 20° near the poles, 146 and with 16 non-equally spaced vertical levels). This is coupled to a 2D energy-moisture-balance-model (EMBM) and 147 a 2D dynamic-thermodynamic sea-ice model. The physics are described in (Marsh et al., 2011; Edwards and Marsh, 148 2005). We use a parameter calibration of seasonal pre-industrial climate following Cao et al., (2009).

149 The <u>Representation of the</u> primary factors controlling the oceanic iodine cycle _____specifically, biological 150 productivity, remineralization, and water column redox - follow Crichton et al., (2021)-Crichton et al., 2021. In this 151 configuration, _____are all represented in the model and described in Ridgwell et al., (2007). In that particular 152 configuration, the rate of organic matter export from the ocean surface is calculated based on just a single nutrient 153 (phosphate) control (together with modifiers reflecting ambient light-and, sea-ice cover, and temperature) and assumes 154 a Redfield-ratio stoichiometry (1:106) with carbon (Fig. 1). Organic matter is split partitioned into particulate (POM) 155 (33% of total export) and dissolved forms (DOM) (67%), with the former sinking down through the water column 156 where it is progressively remineralized according to a prescribed fixed 'decay' curveat a rate scaling with ambient 157 temperature, while the latter is physically transported by circulation and decays (is remineralized with a lifetime of 158 0.5 years). Here, we deviate from Ridgwell et al., (2007) (as well as the calibrated seasonal configuration of Cao et 159 al., 2009) by adopting a calibrated temperature dependence to both export production as well as the decay of POM in 160 the water column (described in Crichton et al., 2021 and Boscolo-Galazzo et al., 2021). When dissolved oxygen nears 161 depletion, sulphate (SO4²⁻) is assumed to be consumd as an electron acceptor supporting to support the remineralization 162 of organic matter (both POM and DOM)₃. The rate of POM remineralization in the water column is governed only by 163 ambient temperature which, in conjunction with a prescribed sinking rate, deterines the vertical distribution of solute 164 release and oxidant consumption. The relative availability of dissolved O_2 vs. $SO_4^{2^2}$ determines the proportion of 165 organic matter degraded by each electron acceptor. In this, the rate of which is governed by athe relative consumption 166 $of SO_4^{2-}$ is governed by a SO_4^{2-} half-saturation saturation limitation term as well as a dissolved oxygen (O₂) inhibition 167 term, while the rate that of oxic resperation of organic matter is restricted by an $[O_2]$ half-saturation limitation term (as 168 described in Reinhard et al., 2020). This deviates from the framework described in Ridgwell et al., (2007). The 169 difference is that, here, SO4²⁻-Sulphate can hence be consumed even before dissolved oxygen can become fully 170 depleted. Ambient temperature dictates the total fraction of POM that decays through both pathways per unit time and 171 within each ocean depth layer (Crichton et al., 2021), with local [O2] and [SO42-] determining the fractional split 172 between alternative pathways (Reinhard et al., 2020). For DOM, the a decay constant (here: 0.5 years) determines the 173 total fraction that decays is remineralized per unit time. It should be noted that currently, there is no published nitrogen 174 cycle in the cGENIE model framework and hence we do not consider nitrate reduction as part of the redox cascade 175 here.

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176 2.2 Marine iodine cycling in cGENIE

177 In the cGENIE model, iodine is present in three reservoirs: IO3- and I in the water column, and I incorporated in POM 178 (and DOM). We then consider four processes that transfer iodine between these reservoirs (summarized in Figure. 1): 179 (1) IO_3^- reduction in the water column, (2) I oxidation (also in the water column), (3) photosynthetic IO_3^- uptake (and 180 assumed intercellular reduction to I⁻), and (4) I⁻ release to seawater during the remineralization of POM (and DOM) 181 (Fig. 1). As dissolved tracersspecies, IO3⁻ and I⁻ are physically transported and mixed through ocean circulation (as is 182 Lincorporated into dissolved organic matter DOM), whereas iodine in POM settles vertically through the water column. 183 This is effectively the same overall framework as-used by Lu et al., (2018). Here-In this paper we re-assess this framework against an updated compilation of observed iodine speciation in the modern ocean and develop and test 184 185 alternative representations of IO3⁻ reduction (process (1), "threshold" threshold" threshold', 'inhibition' "inhibition'", 186 and <u>'reminSO4lifetime'</u>"reminSO4lifetime") and I⁻ re-oxidation (process (2), 'lifetime'"lifetime"'lifetime") 187 "Fennel""Fennel"; fennel", and 'reminO2lifetime": reminO2lifetime": reminO2lifetime"). Although we describe all 65 188 different parameterizations below for completeness and a number of different permutations of 3 IO3 reduction and 3 189 I re-oxidation processes (presented in S.I.), in this paper we will focus primarily on a single reduction parameterization 190 ('threshold') in combination with the 3 different re-oxidation schemes. 191 2.2.1, IO3⁻ reduction schemes, 192 'threshold'. In the numerical scheme of Lu et al., (2018),- when [O2] falls below a set concentration threshold, 193 IO_3^{-1} is immediately and quantitatively reduced to I⁻ (thereafter, we term this iodate reduction parameterization 194 "threshold" (threshold'). 195 **(inhibition'.** The "inhibition" inhibition' scheme links the IO₃ reduction rate with the ambient O₂ concentration. 196 Following the formulation for the rate of SO4²⁻ reduction in Reinhard et al., (2020), we apply an oxygen inhibition 197 term governed by a half-saturation constant. In devising this scheme, we note that while IO3⁻ reduction rates have

been determined experimentally, the quantitative relationship with $[O_2]$ (or other parameters) is unknown. The IO₃⁻ reduction under <u>"inhibition" inhibition</u> is mathematically described as:

$$d[IO_3^-]/dt = [IO_3^-] \times k_{red} \times \frac{k_{O_2}}{k_{O_2} + [O_2]}$$

(1)

in which k_{red} is the maximum first-order reduction rate of IO₃⁻, and k_{O2} is the half-saturation constant of O₂.
 <u>freminSO4lifetime</u>. Reduced sulfur (e.g. sulfides) is also suspected to play an important role in IO₃⁻ reduction

$$208 \qquad \qquad a[10_3]/at = [10_3] \times \frac{1}{\tau_{sul}} \times a[50_4]/at$$

209 in which τ_{sul} defines the rate constant parameter linking the IO₃⁻ and SO₄²⁻ reduction, while the $d[SO_4^{2-}SO_4]$ is 210 amount of SO₄²⁻ reduced during each model timestep.

211 2.2.2 I oxidation schemes

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213		"lifetime" 'lifetime'). In this scheme, I' oxidation follows the first-order reaction kinetics:	Formatted: Indent: Left: 0", Bulleted + Level: 1 +
214		$d[I^-]/dt = [I^-] \times \frac{1}{\tau} \tag{3}$	Aligned at: 0.25" + Indent at: 0.5"
215	•	where τ is the lifetime of I in seawater.	
216		۰ ــــ	Formatted: Indent: Left: 0.25"
217	•	<u>'Fennel'.</u> Given the overlapping redox potential between I and N (<u>e.g.</u> Rue et al., 1997; Cutter et al., 2018), we	Formatted: Font: Bold
218		explore the potential for a link between areas of I and nitrification. To simulate this, Given the potential link	Formatted: Indent: Left: 0", Don't add space between
219		between I ⁻ and nitrification, we devise an alternative <u>'Fennel'</u> "Fennel'' <u>fennel'</u> scheme, in which I' oxidation rates	paragraphs of the same style, Bulleted + Level: 1 +
220		vary as a function of ambient O_2 , increasing with ambient O_2 concentrations towards some hypothetical maximum	Formatted: Superscript
221		value following Michaelis-Menten kinetics (Fennel et al., 2005). In Fennel et al., (2005), this parameterization	romattea. Superscript
222		was originally devised for ammonia reoxidation. The form of this response is defined by the maximum reaction	
223		rate and O ₂ half-saturation constant (Fennel et al., 2005):	
224		$d[I^{-}]/dt = [I^{-}] \times k_{ox} \times \frac{[O_2]}{k_{fenn} + [O_2]} $ (4)	
225		in which k_{ox} defines the maximum rate constant of I ⁻ oxidation, while k_{fenn} is the O ₂ half-saturation constant.	
226	•	<u>'reminO2lifetime'.</u> Finally, in <u>'reminO2lifetime'''reminO2lifetime'''reminO2lifetime''</u> , we associate I' oxidation	Formatted: Font: Bold
227		with O ₂ consumption during remineralization. The logic behind this parameterization is the recent observation of	Formatted: Indent: Left: 0", Bulleted + Level: 1 +
228		I ^{\circ} oxidation to IO ₃ ^{$-$} catalyzed by bacteria, perhaps in association with ammonia oxidation (Hughes et al., 2021).	Aligned at: 0.25" + Indent at: 0.5"
229		Although the nitrogen cycle is not currently included in cGENIE, the NH_{4^+} oxidation can be scaled to OM	
230		remineralization (Martin et al., 2019) and hence to O2 consumption during remineralization. Under	
231		<u>'reminO2lifetime''reminO2lifetime'' reminO2lifetime'</u> , the lifetime of I' oxidation is inversely linked to O2	
232		consumption so that faster remineralizationwhich in the ocean leads to more intensive NH_4^+ oxidation	
233		enhances I oxidation. This I oxidation scheme follow this equation:	
234		$d[I^{-}]/dt = [I^{-}] \times \frac{1}{\tau_{02}} \times d[0_{2}]/dt $ (5)	
235	wh	here τ_{O2} is the rate constant parameter and d[O ₂] is the O ₂ consumption during remineralization during a single	
236	tin	nestep in the model.	
237	<u>2.2</u>	2.3 Biological reduction pump	
238	Th	e final <u>pair of coupled</u> processes in the marine iodine cycle framework concerns the processing of iodine directly	
239	thr	ough the biological pump. Phytoplankton-absorbed iodine is stored in the cell as IOa, I, or other forms, followed	Formatted: Subscript
240	by	release during senescence (Hepach et al., 2020). While there is some uncertainty as to whether IO3 ^{indate} reduction	Formatted: Superscript
241	is a	assimilatory or dissimilatory (Hepach et al., 2020), we adopt a comparable approach to nitrogen cycling (sequence:	Formatted: Superscript
242	NC	Da uptake, N incorporation into organic matter, remineralization and release as the reduced NHA+ form). We assume	Formatted: Subscript
243	<u>tha</u>	t it is necessary to clarify here that in order to we simplify the simulation, the modeled by assuming the IO t tracer	Formatted: Superscript
244	is	assimilated by phytoplankton and incorporated into POM during photosynthesis -In this, IO3- is taken up by	Formatted: Subscript
245	ph	ytoplankton and incorporated into OM during photosynthesis (Elderfield and Truesdale, 1980) before beingand	Formatted: Superscript
246	rel	eased as I ⁻ during remineralization and/or cell senescence (Wong et al., 2002; Hepach et al., 2020; Wong et al.,	Formatted: Subscript
247	19	85). cGENIE simulates these processes as a function of avia an assumed 'Redfield-ratio' of iodine to carbon (I:C	Formatted: Superscript

248 ratio) in OM. We note that while the value of I:C can be adjusted in the model is tunable, it is fixed currently assumed 249 fixed in value throughout the ocean. We discuss the merits of an optimized and uniform I:C compared to variable I:C 250 (e.g., Waddley et al., 2020) in more detail in the discussion.

251 2.3 Model-data comparisonevaluation

252 We used the model skill measure (M) (Watterson, 1996) to assess the performance of the marine iodine cycle in 253 cGENIE. A major advantage of the M-score is that it is calculated through location-dependent comparison (Fig. 2; Lu 254 et al., 2018). Another advantage of the M score is that it captures overall improvement of model performance relative 255 to model minus observation maps, since it is non-dimensional, and the higher M stands for better performance. For 256 comparison evaluating the marine iodine cycle in the cGENIE model with simulated distributions of iodine speciation, 257 we compiled oceanic iodine observation data from the literature (Fig. 2B; Table S4). This-Our dataset includes-builds 258 on the compiled dataset compilation of Chance et al., (2019) and Sherwen et al., (2019), which was used to calibrate 259 the Wadley et al., (2020) model, but includes more recent publications (referenced in Table S4) and is also expanded 260 to include the deep ocean and ODZ data. To avoid the influence of freshwater dilution and recycled iodine from the 261 sedimentary flux, we applied a filter which only keeps the measurements with total iodine (or IO₃ plus L) between 262 450 nM and 550 nM in the dataset. Note that the I⁻ measurements from the GP16 cruise in the ETSP are not included 263 for the comparison because of potential method considerations (see Cutter et al., 2018 and Moriyasu et al., 2023). 264 After these filtration methods filtering, the data were re-gridded by taking the average values according to the cGENIE 265 model grids. 266 We used the model skill measure (M-score) (Watterson, 1996) to assess the performance of the marine iodine 267 cycle in cGENIE compared to the gridded data. A major advantage of the M-score is that it is calculated through 268 location dependent comparison (Fig. 2; Lu et al., 2018). Another advantage of the M score is that it captures overall 269 improvement of model performance relative to model-minus-observation maps, since it is non-dimensional, and the

270 higher M stands for better performance. For each iodine speciation (hereby IO3- and I), a M-score M-score value is 271 calculated through comparing re-griddedgridded observations versus model results in each corresponding grid (Figure.

272 2). The synthesized M-score for iodine of each model experiment is calculated through averaging those for both I' and IO3⁻. The higher the M-score value the better the model-data performance.

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277 Figure 2: A), An example of location-dependent comparison between I- distributions in the cGENIE model 278 iodine data array and the regridded ocean observation. B). The sampling locations of iodine observation data 279 used for model-data comparison. Some coastal stations included in the figure are filtered out in the model-data comparison. The ETNP transect associated with Fig.6 is labeled as red box. The Ocean surface (uppermost ~81 280 281 m corresponding to the depth of the surface layer in the cGENIE model) sampling locations of IO₃ (A) and I 282 (B) field observations after data filtration for freshwater and sediment fluxes. The filtered observations were 283 then regridded according to the cGENIE 36×36×16 framework for model-data comparison of IO₃ (C) and I (D). The orange boxes in A and C surroundhighlight the grids of ETNP the oxygen deficient zone transect shown 284 285 <u>in Fig. 6Fig. 7.</u>

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287 2.4 Sensitivity analyses and model implementation

288 Because the relative roles of IO3⁻ reduction, I⁻ oxidation, and the shuttling of iodine through the biological pumpIO3⁻ 289 planktonic uptake in the water column are uncertain, we calibrate the parameters controlling these processes in 290 cGENIE by creating an ensemble of different parameter value combinations arranged controlling IO3 reduction and 291 I oxidation in a 2D regularly -spaced grid and then repeat the same 2D parameter ensemble for different assumptions 292 of regarding the biological pump (I:C) (Table 1, Fig. 3Fig. 4). The output of each ensemble member is then statistically 293 compared to our observational database. We assume the associated parameterization when the model reaches the best 294 M score of replicating modern ocean iodine distribution would also be applied to simulate iodine cycling in the past. 295 We focus on parameter ensembles testing the 3 different parameterizations for I oxidation (but only 'threshold' as the 296 IOs reduction parameterization) - 'lifetime-threshold', 'fennel-threshold', and 'reminO2lifetime-threshold'. (The 297 results of 2 additional parameterization-combinations - 'lifetime-reminSO4lifetime', and 'lifetime-inhibition' are 298 given in Table S1.) We discuss the reasons for selecting these specific parameterization-combinations in the 299 Discussion section. Finally, Toto explore whether the thecGENIE model-simulated dissolved oxygen distribution

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300	imparteds any particular bias to the tuned iodine cycle, we repeated the model ensembles for each of the 3
301	parameterization-combinations but, continually restoring the 3D pattern of $[O_2]$ in the model to that of the World
302	Ocean Atlas 18 (WOA18) climatology (Garcia et al., 2018). The model ensembles are summarized in Table 1.
303	Each ensemble member was run for a total of 2,000 years and each starts from the same initial state, which
304	was an experiment run for 10,000 years to equilibrium using a random set of iodine parameters within the ranges in
305	Table 1. Running the modelseach ensemble member for 2000 years minimizes the CPU time but was also found to be
306	more than sufficient to allow iodine inventories to equilibrate to new steady states. To explore whether the model
307	simulated dissolved oxygen distribution imparted any particular bias to the tuned iodine cycle, we repeated the model
308	ensembles, continually restoring the 3D pattern of [O ₂] in the model to that of the World Ocean Atlas 18 (WOA18)
309	elimatology (Garcia et al., 2018). The output of each ensemble member is then statistically compared to our
310	observational database.
311	
312	2.5 Evaluation against geological observations
313	Parameter tuning, and the ability to reproduce modern observations, does not by itself offer any guarantee that spatial
314	patterns are being simulated for the 'correct' mechanistic reason-(i.e., specific set and relative importance of
315	mechanisms). This is even more pertinent in the context of the application of a modern-tuned model to paleo-redox
316	reconstruction. To test whether ourquantify to what degree the calibrated parameterization-combinations for the
317	modern marine-new iodine cycle hadhave predictive power in the geological past, we carried out a deep-time
318	plausibility test.
319	For the paleo plausibility test, we adopted the Cretaceous, pre-OAE2 (ca. 93 Ma) configuration (continental
320	arrangement and ocean bathymetry, wind stress and velocity, and zonal average planetary albedo boundary conditions)
321	from of Monteiro et al., (2012). We choose this particular geological interval because the controls on ocean redox have
322	been previously evaluated using the cGENIE model (Monteiro et al., 2012; Hülse et al., 2019), the oceanic conditions
323	are much more extensively dysoxic and anoxic than present-day and hence represent a relatively severe test of the
324	model iodine cycle, and- a number of I:CaI/Ca proxy measurements are available (Zhou et al., 2015). In order to
325	evaluate the same configuration of the iodine cycle as optimized in this study, we also substituted the temperature-
326	independent representation of biological export production and fixed remineralization profile of POM in the water
327	column (i.e., Ridgwell et al., (2007)) for the temperature-dependent scheme of Crichton et al., (2021) used in our
328	modern calibration. However, in substituting the biological pump scheme in the model we changealter the ocean redox
329	landscape compared with e.g., Monteiro et al., (2012). We therefore testexplore a range of different assumptions
330	regarding the ocean PO ₄ inventory at the time as a means of generating a range of different plausible states of ocean
331	oxygenation. In this, we test: 0.2, 0.4, 0.6, 0.8, 1.0, and 1.5 times the mean modern concentration (2.15 µM). We run
332	the model with each of the best-fit (highest M-score) sets of parameter values associated with the main 35 different
333	parameterization-combinations (but focus on the results of the same 32 additional parameterizationcombinations we
334	did for comparison against modern were run and are presented in SI), and for each of the varying PO4 inventory
335	assumptions, and for 10,000 years to steady-state. Major cation concentrations were adjusted to a value more

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357	dependency	with temperature	(Fig	. S3 ir	1 Zhou	et al.,	2014). For	our	Cretaceous	model	calibration,	we apply	the K	D
		*													_

358 based on local temperature (at each grid point associated with a sampling section) simulated by cGENIE. Beyond

359 <u>temperature, we acknowledge that IO_3^- incorporation into carbonate lattice through substitution $IO_3^- + Na^+ \leftrightarrow CO_3^{2-}$ </u>

 $\frac{+ Ca^{2+} \text{ is controlled by [Na^+], [CO_3^{2-}], and [Ca^{2+}] (Podder et al., 2017). However, either quantifying these ions during}{2}$

361 the Cretaceous seawater or quantitative calculation of ion substitution dynamics requires further constraints. Although

362 <u>uncertainties are inevitable, we assume our temperature-controlled [IO₃]-to-I/Ca conversion based on current</u>

363 quantitative knowledge meets the requirement for Cretaceous model-data comparison. We extracted simulated I/Ca

364 values from the model grid points corresponding to the sections reported by Zhou et al., (2015) and calculated the M-

365 <u>score.</u>

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combinations here (Table 1) "lifetime threshold": "Fennel threshold": "Fennel threshold": and

"reminO2lifetime threshold": (A detailed justification and discussion for selecting these

parameterization combinations is included in the Discussion section.)

Table 1. The cGENIE iodine redox options and the associated range of parameters of these options. The detailed introduction of each parameter is described in section 2.2.2 and the plausibility of these parameter ranges is discussed in 4.1.1. Note that the oxidation rate constant k in 'Fennel' is in unit of year'l in the model configuration, which is the reciprocal of the 'lifetime'. A detailed table containing all considered parameterization ranges can be found in Table S1.

-	Parameter	ization-		lodine oxidation	parameters	lodine reduction parameters		
_	combination descrip	<u>ination</u> Parameter description 'life (y		'reminO2lifeti me' (× 10⁻⁵ mol/kg)	'Fennel' (Inhibition constant/ μM O ₂)	'threshold' (μM O₂)	I:C ratio (× 10 ⁻⁴ mol/mol)	
	Simulation	cGENIE O ₂	10-170	١	١	1-110	0.5-3.5	
	threshold'1	WOA	10-170	١	١	1-110	0.5-3.5	
-	<u>'fennel-</u>	cGENIE O ₂ .	10-170 (1/k)	0.01- 1	20	1-110	0.5-3.5	
	tion 2	WOA.	10-170 (1/k)	0.01- 1	20	1-110	0.5-3.5	
-	<u>'reminO2lifetime</u>	cGENIE O ₂	١	0.01- 1	١	1-100	0.5-3.5	
	threshold'Simula tion 3	WOA	١	0.01- 1	١	1-100	0.5-3.5	

Results 385 3.

387

419

386 In this section, we start by summarizing the overall statistical outcome of the tuning, then present a series of spatial

analysis comparisons for each of the highest M-score ensemble members. The spatial analyses progressively reduce 388 in scale, moving from global surface distributions (Section 3.2), to global and basin-specific water column profiles

389 (Section 3.3), and finally to spatial comparisons for a specific ODZ region (Section 3.4). Our lastfinal comparisonset

- 390 of results (Section 3.5) is based on are of the modeled OAE-2Cretaceous scenarios using the best parameterizations
- 391 from the modern, which then are compared to carbonate I:Cal/Ca datavalues measured in the rock record.

392 3.1. Model skill score

393 The M-score values achieved across the complete ensemble for each of the 3 main parameterization-combinations are 394 illustrated shown in Fig. 3Fig. 4. The sensitivity test shows the and illustrate how the statistical fit is M scoreM-scores 395 are sensitive to all three of the main parameters for the ensembles in Fig. 3. Higher model skill scores are usually 396 reached when "threshold" threshold" is tuned to $10 \ \mu M$ [O₂] for all the ensembles, including both model-simulated 397 [O₂] and WOA-forced [O₂]. For the ensembles, "lifetime-threshold" lifetime-threshold and "Fennel-398 threshold": fennel-threshold', the highest M-scoreM-scores are similar - 0.305 and 0.308, respectively (Table 2). 399 Both these ensembles have the highest performance when "threshold" threshold", "lifetime" lifetime", and I:C ratio are tuned to 10 μ M [O₂], 50 years, and 1.5×10^{-4} mol/mol, respectively, which is generally consistent with observations 400 401 (Lu et al., 2016, 2020b; Tsunogai, 1971; Elderfield and Truesdale, 1980) (discussed in more detail later). The model 402 performance of "reminO2lifetime-threshold" reminO2lifetime-threshold' is less good than the other two 403 combinations. with the best M score M-score of 0.266 when "threshold" 'threshold'. 404 <u>"reminO2lifetime"</u>; reminO2lifetime", and I:C ratio are tuned to 10 μ M O₂, 1 × 10⁻⁶ mol/kg, and 3.5 × 10⁻⁴ mol/mol, 405 respectively (Table 2, Fig. 3Fig. 4). We note that for each parameterization-combination, the highest possible M 406 score<u>M-score</u> achievable by tuning improves when [O₂] is forced to that of the World Ocean Atlas 18 (WOA18) 407 climatology (Garcia et al., 2018). 'reminO2lifetime''reminO2lifetime''reminO2lifetime threshold'

408 3.2. Meridional surface I⁻ distribution

409 Figure 4-5 shows a comparison between the observed latitudinal distribution of [I⁻] at the surface and as simulated by 410 the model for each parameterization-combination (for the respective best M-score ensemble member). Note that the 411 observations (Section 2.3) are binned to the corresponding model grid cells and as such, reflect averages over the 412 upper-most 80 m of the water column. This represents a reduction from 1338 to 141 surface ocean data points. We 413 find that the surface ocean [I⁻] in the model shows a trend of increasing values towards low latitudes, broadly consistent 414 with observations (Chance et al., 2014) (Fig. 4Fig. 5). The "lifetime threshold" 'lifetime-threshold' and "Fennel-415 threshold"'fennel-threshold' show similar latitudinal trends, but both overestimate the surface I' in the mid-low 416 latitudes in the southern hemisphere. The 'reminO2lifetime-threshold'22 ensemble produces better estimation of 417 meridional surface [I-] trend, although overestimates [I-] in the tropical surface ocean compared to the other two 418 ensembles (Fig. 4Fig. 5).

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420

421Figure 45: Modeled latitudinal surface iodide distribution compared with observation with the cGENIE422simulated [O2] and the [O2] restoring forcing. The elevated [I⁻] observed and modeled in low latitudes is the423result of phytoplankton reduction in the surface ocean. Note that the I⁻ distribution simulated by <u>``</u>lifetime-424threshold<u>``</u> and <u>``</u>fennel-threshold<u>``</u> are close but not identical.

425

426 3.3. Global and basin-specific iodine depth distributions

427 Comparisons between the observed distributions of I^{\cdot} and IO₃⁻ seawater concentrations among the global ocean and 428 the Atlantic and Pacific Oceans are presented in Fig. 5Fig. 6. Again, we re-gridded the iodine observations (see: 429 Section 2.4) and selected sub-sets of the data that lay in either Atlantic or Pacific basins, contrasting with the

430 corresponding model values at those locations. We find only relatively minor differences between the best M-score

431 ensemble member of each of all three parameterization-combinations, and all show increased $[IO_3^-]$ and decreased $[I^-]$ 432 with increased depth below the photic zone in the Atlantic and Pacific basins, as well as globally (Fig. 5Fig. 6). The 433 modeled depth profile broadly matches with observation in the Atlantic and deep Pacific Ocean, except the 434 underestimated subsurface peak of [I⁻] observed in the Pacific and overestimated [IOs⁻] in the deep Pacific (Fig. 5Fig. 435 $\underline{6}$). This mismatch of subsurface I⁻ peak is probably the result of sampling bias, with most of the Pacific iodine 436 observations from ODZs in the Eastern Tropical North Pacific (ETNP) and the Eastern Tropical South Pacific (ETSP). 437 For example, in model depth profiles masked to only include grid points with corresponding observations, the modelled Pacific depth profiles show a clear mid-depth ODZ feature (Fig. <u>\$859</u>). 438





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441 442 443 444

Figure 56: Modeled averaged iodine (including iodate and iodide) depth profile among global ocean, the Pacific, and, the Atlantic compared with observation. The surface I⁻ enrichment among the ocean basins is caused by phytoplankton reduction. The subsurface (~500m) I⁻ enrichment is the result of sampling bias <u>that since</u> most of the observations are from the ETNP and ETSP ODZs (see main text for details).

446 **3.4.** Iodine distribution within ODZs

447 To assess the model ability to simulate iodine cycling in marine low oxygen environments, we compared distributions 448 of oxygen and iodine species in the ETNP (Fig. 6Fig. 7). The O2 transects amongst all model simulations are the same 449 because we only changed the parameterizations of the iodine cycle between ensembles and ensemble members (i.e., they all simulate the same biological pump in the ocean). All the three chosen best-performance-experiments show 450 451 similar iodine anomalies (IO3- depletion) in the ETNP, fitting the general feature of the observation. Other 452 parameterizations did not replicate the ODZ (Fig. S4). However, even under the "best-fitting" parameters, compared 453 to the observations, the ODZ feature in the model is underestimated both in intensity and in areal extent compared to 454 the observations (Fig. 7). Notably, compared to [O₂] measured in the ETNP transect, the model underestimates the 455 extent of the ODZ. Severe deoxygenation below 50μ M [O₂] was observed in relatively shallow depths between 100-456 200m in the ETNP, and this ODZ extends for more than 3000 km towards off-shore from Mexican coast (Fig. 6Fig. 457 7). Although cGENIE simulates the O2-deficient pattern in the ETNP, the extent of the ODZ is underestimated. The 458 simulated oxycline is ~200 m deeper than the observation and the [O2] variation is gradual. The ODZ below 20 µM 459 [O2] in the model is limited to a small spatial extent within 1000 km offshore, which is much smaller than that in the

- 460 observation (Fig. 6Fig. 7).
- 461



467

Figure 67: Modeled (contour) and observed (colored dots) west-to-east transect (location shown in Fig. 1) of O2 (top row) and IO₃ and O₂(bottom rows) in the ETNP. Note that the WOA-forced [O₂] models simulate a larger extent of IO3 anomaly, which better matches the observation. The left-hand panel contours are model results 466 based on cGENIE-simulated [O₂] while contours on the right are model results from WOA-forced [O₂].

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468 All the three chosen best performance experiments show similar iodine anomalies (IO3⁻ depletion) in the 469 ETNP, fitting the general feature of the observation. Other parameterizations did not replicate the ODZ (Fig. S4). 470 However, even under the "best fitting" parameters, compared to the observations, the ODZ feature in the model is 471 underestimated both in intensity and in areal extent compared to the observations (Fig. 6). The observed IO₂-depletion 472 zone ([IO₃⁻] <100_nM) occurs in shallower depths between 100-500_m, matching the shallow ODZ and spans ~2000 473 km offshore; however, the modeled IO37 depletion zones in the ETNP are located in 400 700_m, and only extends 474 within 1000 km from the shore.

475 We also ran model ensembles forcing cGENIE to restore the modern ocean [O2] annual average climatology 476 to that of the WOA18 (Garcia et al., 2018) (Fig. 6Fig. 7). Now the Under these conditions, the subsurface IO3⁻ depletion 477 zone in the ETNP ODZ in all three ensembles extends ~2000 km offshore and spans across 100-1000 m in depth (Fig. 478 67). The shallow and extended ODZ iodine distribution in the ETNP better matches the observation compared to non-479 O2 restoration simulations.

481 482 483

484

Table 2. The performance of the cGENIE iodine simulations and associated parameterization when the model reaches the best global <u>M scoreM-score</u>. Note that the oxidation rate constant k in 'Fennel' is in unit of year⁻¹ in the model configuration, which is also the reciprocal of the 'lifetime'. The full model performance is summarized in Table S2. Note that the lifetime in 'Fennel' is parameterized as k (year⁻¹) = 1/lifetime.

lodine lodine oxidation parameters reduction I:C ratio (× 10⁻⁴ Parameterization-combination parameters Model skill Parameter description 'Fennel' score 'lifetime' 'reminO2lifetime' 'threshold mol/mol) (Inhibition constant/ (years) (× 10⁻⁵ mol/kg) (µM O₂) μM O₂) 'lifetimecGENIE O₂ 50 ١ ١ 10 1.5 0.305 threshold'Simulati on 1 WOA 50 ١ ١ 10 1.5 0.385 50 20 10 1.5 0.308 cGENIE O₂ ١ 'fennel-(1/k) threshold'Simulati 10 on 2 ١ 20 10 3.5 WOA 0.385 (1/k) cGENIE O₂ ١ 0.1 ١ 10 3.5 0.266 'reminO2lifetimethreshold'Simulati on 3 WOA ١ 0.1 ١ 10 3.5 0.365

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517	4. Discussion
518	In summary: we ran-presented the results of cGENIE Earth system model ensembles (both with internally calculated
519	and WOA-imposed [O2] distribution) ensembles for five-for 3 parameterization-combinations of iodine cycling that
20	showed the best performance parameters (but summarized a total of 5 different parameterization-combinations in
21	Table S2) in the cGENIE Earth system model (both with internally calculated and WOA imposed [O2] distribution)
22	and presented the results for 3 of them that showed the best performance. We analyzed the performance of the
23	ensembles via: (1) <u>M score</u> for the model-data match of both $[I^{-}]$ and $[IO_{3}^{-}]$ across the entire ocean, (2)
24	qualitative model-observation comparison of latitudinal surface ocean distributions of [I ⁻], (3) inspection of averaged
25	depth profiles in global and individual ocean basins for both $[I^{-}]$ and $[IO_{3}^{-}]$, (4) inspection of iodine transects across
26	the across the Eastern Tropical North Pacific (ETNP) ODZ, and (5) M score M-score for model and carbonate I:CaI/Ca
27	observations (Zhou et al., 2015)_for pre-OAE2the Cretaceous.
28	4.1. Overall model skill score comparison
29	Two broad observations emerge from the M score M-score comparison. First, the 1st-order kinetic iodine oxidation
30	associated ensembles ('lifetime-threshold' and 'Fennel-threshold') have the highest M-score (Table 2),
31	under both cGENIE-simulated [O2] and WOA-forced [O2]. This is consistent with previous observations of 1st-order
32	kinetics for I ⁻ oxidation (Tsunogai, 1971). Second, the simulations with WOA-forced [O ₂] produce significantly
33	higher $\frac{M \text{ score}M \text{ -score}s}{M \text{ -score}}$ than that of the cGENIE-simulated [O ₂] field (at least ~0.08 of improvement; Table 2).
34	Despite a 1st-order non-O2 dependent oxidation mechanism providing the highest M-scoreM-scores, the WOA vs
35	internally model-generated dissolved oxygen distributions comparison highlights the O2 and related redox
36	dependency of the iodine cycle from the perspective of IO3 ⁻ reduction. Each of these factors are discussed in the
37	following Section 4.1.1.
38	Notably, our M score M-score results also demonstrate a potentially minor role, if any, for iodine cycling
39	with DOC. As described in 2.3.1, the "reminO2lifetime"'reminO2lifetime' and the "reminSO4lifetime" seales I
40	oxidation and IO3 ⁻ reduction with O2 consumption and SO4 reduction during the remineralization of POC (or POC
1	and DOC), respectively. An alternative (with DOC remineralization) scenario was tested to combine the iodine
12	redox reactions with DOC remineralization in cGENIE (Table S1). Compared to the default settings, adding the
43	DOC remineralization control to the I-oxidation ("reminO2lifetime" reminO2lifetime") or IO3-reduction
4	("reminSO4lifetime") does not increase the M scoreM-score of the model. More than the overall M scoreM-score,
45	the simulation of latitudinal I ⁻ trend in the defaulted no DOC controlled iodine cycle models performs better in
16	replicating the depth profiles, as well as the IO3 ⁻ depletion feature in the ODZ (discussed in later sections),
+0	

A credible representation of the marine iodine cycle requires not only that observations can be replicated, but that
 replication occurs whenin achieving a fit to observations, tuned parameter values fall within real-world ranges. In this

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section, we discuss the validity-plausibility of our best-fit (maximized M-score) parameter values. For the iodine cycle,
these parameters include are: the O₂ threshold, <u>I:C ratio, and the parameter (depending on the specific</u>
parameterization-combination) controlling the I⁻ oxidation rate, and the cellular I:C ratio.

555 Our model M scoreM-score is highest with an [O2] reduction threshold of 10_uM (Fig. 3Fig. 4 and Table 2). 556 Although it is generally well accepted that IO3⁻ is reduced in low oxygen settings (Luther, 1991; Rue et al., 1997; 557 Wong et al., 1985; Wong and Brewer, 1977; Farrenkopf and Luther, 2002), the degree of O₂ depletion that triggers 558 IO_3^- reduction is still unclear. A relatively high threshold for triggering IO_3^- reduction is shared based on 559 comparison between planktic and benthic foraminiferal I-Cal/Ca and ambient [O2] (20-70 µM O2; Lu et al., 2016, 560 2020a). These [O₂] thresholds are similar to that determined in a previous cGENIE-based iodine cycle study (30 µM) 561 (Lu et al., 2018), but it is difficult to directly compare this to our results because of differences in the model 562 representation of the ocean biological pump, the iodine observational data-set, and model-data comparison methods 563 utilized.

564 Many of the studies suggesting a relatively high $[O_2]$ threshold is are based on evaluations the relationship 565 of $[IO_3^-]$ and $-[O_2]$ within the upper chemocline of ODZs; however, evaluation of $[O_2^-]$ and $[IO_3^-]$ from ODZs throughout the entire water column suggest the potential for IO₃⁻ persistence within the low oxygen cores of ODZs. 566 567 Specifically, IO3⁻ accumulation is observed within the ETNP and ETSP at depths where [O2] is close to or below the 568 detection limit of the sensors which is reported near 1 µM (Hardisty et al., 2021). In addition, it is worth noting that 569 the kinetics of IO_3^- reduction is heterogeneous both within and between ODZs. For example, a transect evaluating IO_3^- 570 reduction rates in the ETNP observed rapid rates in the upper oxycline, where $[O_2]$ was near ~12 μ M, but the potential 571 for sluggish rates in the ODZ cores, where $[O_2]$ was below detection. In an early study of the Arabian Sea, IO_3^- was 572 reduced rapidly within the ODZ core. Together, these suggest IO3 reduction may be controlled by factors beyond O2 573 (Hardisty et al., 2021; Farrenkopf and Luther, 2002). For example, IO3 is likely formed in high [O2], non-ODZ water 574 masses but can be sustained upon transport or mixing within oligotrophic, offshore ODZ regions where organic 575 supplies are more limited (Hardisty et al., 2021). A comparison to the N cycle would also indicate a low [O2] 576 threshold—__for example, denitrification has a sub-µM [O2] threshold and has a similar redox potential with IO3-577 reduction (Dalsgaard et al., 2014; Thamdrup et al., 2012). A sub-µM [O2] threshold for IO3⁻ reduction could explain 578 the $[IO_3]$ variations observed in ODZ cores with $[O_2]$ below the μ M detection limits of sensors; however, iodine 579 speciation has yet to be analyzed alongside $[O_2]$ measurements via sensors with sub- μ M detection limits, such as 580 STOX sensors. Regardless, our 10 μ M [O₂] threshold based on maximizing the <u>M-scoreM-score</u> reflects a global 581 average value and clearly falls within the ranges of oceanographic observations.

For both our study and that of Lu et al., (2018), an I⁻ lifetime of 50 years maximizes model performance. However, Lu et al., (2018) chose to implement a slightly lower value of 40 years for their paleo-application because it reflected the slowest rate observed in the literature at that time (Tsunogai, 1971). Notably, though IO₃⁻ is the most abundant marine iodine species, its production from I⁻ has never been <u>unambiguously</u> observed under normal marine conditions. This has acted as a major hinderance on providing direct constraints. That said, our <u>eGENIE-model-based</u> estimate is consistent with a multitude of other constraints that indicate that I⁻ oxidation to IO₃⁻ undergoes extremely slow kinetics. The I⁻ oxidation rates calculated through indirect methods including mass balance and seasonal iodine 589 speciation changes (Tsunogai, 1971; Campos et al., 1996; Truesdale et al., 2001; He et al., 2013; Edwards and 590 Truesdale, 1997; Žic et al., 2013; Moriyasu et al., 2023) or through radiogenic tracer spiked incubations (Hardisty et 591 al., 2020; Schnur et al., 2024; Ştreangă et al., 2024) have a wide range of variation from 1.5_nM/yr to 670_nM/yr. The 592 lifetime in cGENIE is 50 years, which can be approximately converted to the zeroth order rate of <9 nM/yr, falling in 593 the lower end of the previous studies.

594 Our best-fit I:C ratio is 1.5×10⁻⁴ mol/mol, and this value which is in agreement with plankton measurements 595 and mass-balance calculations (Chance et al., 2010; Elderfield and Truesdale, 1980). In the photic zone, IO3⁻ is taken up by phytoplankton and incorporated into their cellular structures followed by subsequent conversion into I- (Hepach 596 597 et al., 2020). Due to this, it is assumed that IO3⁻ removal in the surface layer of the ocean is a function of organic 598 carbon fixation during primary productivity according to Redfield-like ratios (Campos et al., 1996; Chance et al., 599 2010). Of the parameters incorporated into the model, in theory, I:C should probably be the best constrained. However, 600 published I:C ratios based on field and laboratory measurements vary over several orders of magnitude (10⁻⁵ to 10⁻³) 601 (Elderfield and Truesdale, 1980; Campos et al., 1996; Hepach et al., 2020; Chance et al., 2010). To limit the number 602 of model simulations and size of the ensembles, we only varied I:C between 5×10^{-5} and 3.5×10^{-4} mol/mol (in 603 increments of, increasing by 1.0×10^{-4}), which covers the range indicated by previous studies (Elderfield and Truesdale, 604 1980).

605 It is unlikely that the I:C value is constant across the global ocean due to differences in phytoplankton 606 compositions and other factors, such as temperature (Wadley et al., 2020). In cGENIE, most of the elevated (> 200 607 nM) surface [I⁻] values over 200 nM is present occur in the ETSP and the northern Indian Ocean, representing the 608 effect of high primary productivity and transformation of IO₃⁻ to I⁻ via the rapid recycling DOM 'loss term' in the 609 representation of biological export in the model (Fig. S6). The mismatch underestimation overestimation of tropical I 610 with tbetween the model and the observations probably hints that the I:C ratio is not constant in the ocean, as which 611 is also hypothesized by Wadley et al., (2020), although in the absence of an explicit representation of primary 612 productivity in the model and lack of a spatially variable f-ratios (Laws et al., 2000)_(implicitly, the f-ratio is can be 613 considered to be 0.33 everywhere in cGENIE). In testing a fixed, spatially uniform I:C, Wadley et al. (2020) 614 underestimated surface [I⁻] in low latitudes and overestimated concentrations in mid-latitudes. Based on their model-615 observation comparison, they hypothesized that the I:C ratio decreases systematically with sea surface temperature 616 (SST) (Wadley et al., 2020). Until more constraints are developed on spatial variability and associated driving factors 617 for I:C, a generalized approach of a globally uniform I:C remains the most parsimonious assumption, especially considering cGENIE's intended extrapolation to ancient settings. 618

619 4.1.2. Comparing alternative iodide oxidation parameterizations in cGENIE

Due to similarity in redox potentials, the iodine cycling in the ocean has been hypothesized to be linked to the cycling of nitrogen (Rue et al., 1997). Nitrification promoting I^{\circ} oxidation to IO₃^{\circ} has been hypothesized inferred from in field studies (Truesdale et al., 2001; Žic et al., 2013), and more recently has been linked via observation of I^{\circ} oxidation to IO₃^{\circ} by ammonia oxidizing bacteria in laboratory environments (Hughes et al., 2021). We further note that Wadley et al., (2020) explicitly link I^{\circ} oxidation to NH₄⁺ oxidation in their surface ocean iodine cycle model. **Formatted:** Indent: First line: 0.5", Don't add space between paragraphs of the same style

625 As an alternative to the first-order lifetime oxidation parameterization used here and in Lu et al., (2018) and 626 in the current absence of a full nitrogen cycle (and hence explicit NH_4^+ oxidation) in cGENIE, we also applied a 627 strategy ("reminO2lifetime") which links I oxidation to organic carbon remineralization and the consumption rate 628 of dissolved oxygen. The reasoning behind this is that the O₂ consumption rate in the model implicitly reflects bacterial 629 oxidizing activity in the water column and hence the potential for I⁻ to be oxidized to IO₃".

We find that the overall model performance involved with the <u>'reminO2lifetime'</u> is lower than other experiments where I⁻ oxidation is ubiquitously oxidized according to a parametrized lifetime, or <u>'lifetime-threshold'</u> (maximum <u>M-score 0.266 vs. 0.305/0.308 under cGENIE simulated [O₂]) (Fig. 3Fig. 4</u> and Table 2). However, despite slightly lower <u>M-score M-scores</u>, the <u>'reminO2lifetime''</u> <u>scheme</u> generally replicates the latitudinal surface [I⁻] trend, the depth profiles in the ocean basins, and the ODZ transect (Figs. 4-6).

635 Under the default-'lifetime'- scheme, I' will oxidize in the whole ocean regardless of the concentration (or 636 even presence/absence) of ambient O2. This scenario might hence not perform well in replicating the ocean iodine 637 cycling at intervals during the Phanerozoic when ODZs were highly expanded, especially for O2 lower than the IO3-638 reduction threshold which should inhibit I oxidation or particularly during the Precambrian when the majority of the 639 ocean was ferruginous or euxinic and highly depleted in O2 (for example, summarized by Lyons et al., 2014). Ideally, 640 for application to in paleoceanographic studies, I oxidation should be linked to ambient an [O2] related I oxidation 641 alternate scheme is required in some way. Although thermodynamics theory suggests O_2 does not directly drive I 642 oxidation (Luther et al., 1995), field studies in ODZs indeed observed low [O2] inhibits this process (Farrenkopf and 643 Luther, 2002; Moriyasu et al., 2020). We hence developed and tested variable I⁻ oxidation kinetics, with the ambient 644 dissolved O₂ providing an inhibition of the rate of oxidation based on the enzymatic nitrification scheme of Fennel et 645 al., (2005).

646 Since most of the ocean is fully oxygenated today, the there is little difference of in M scoreM-scores between 647 "lifetime"______ and "Fennel" oxidation models are minor parameterizations (0.305 vs. 0.308, Table 2). The parameter valuess associated with the highest M score<u>M-scores in each between two oxidation options scheme are also very</u> 648 649 close to each other, except with the exception that 'Fennel'"Fennel"'fennel' oxidation together with the modelunder 650 WOA forcing, has a higher optimal I:C ratio $(3.5 \times 10^{-4} \text{ mol/mol})$ and faster saturated I⁻oxidation kinetics (0.1 yr⁻¹ vs. 651 0.02 yr+ in other ensembles). The parameter differences between the 'Fennel'"Fennel" WOA ensemble and 652 other models ensembles make sense because the faster oxidation rate compensates the increased I⁻production through 653 the higher biotic uptake rate. Despite the very close M-scores, 'Fennel' oxidation has aunder WOA-forcing has a 654 higher optimal I:C ratio (3.5), \times 10⁴ mol/mol) and faster saturated I oxidation kinetics (0.1 yr⁻¹ vs. 0.02 yr⁻¹ in other 655 ensembles), Such parameter differences between 'Fennel'-WOA and other ensembles reflect compensation between 656 faster oxidation rate (2) in Fig. 1) and increased I production through the higher biotic uptake rate (3) in Fig. 1). The 657 pre-OAE2 simulations are particularly illustrative of this tradeoff and are discussed in more detail in section 4.3. 658 In summary, all the three parameterization-combinations produce high and comparable M scoreM-scores 659 and similar parameters (oxidation, reduction, and I:C) associated with these M-scores (Table 2). Although 660 direct field-based evidence to evaluate the controlling parameters of 'reminO2lifetime'-is absent, the parameters 661 controlling the other model scheme are consistent with previous studies.

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662 4.2. Beyond the M-score: model-data comparison across iodine gradients

663 As applied here, the M-score provides a quantitative measure that describes the overall model global 664 performance and allows us to directly compare the implications of parameter value and parameterization choices. 665 However, the a M score M score based on a global dataset overlooks can obscure regional gradients that may be important for both paleo- and modern oceanographic research. Indeed, amongst all the various ensembles we ran as 666 667 part of this study (Table S1), only 'lifetime-threshold', "'lifetime-Fennel'", and 'reminO2lifetime-threshold'" 668 performed sufficiently well in replicating the modern oceanic iodine gradients (Figs. S2-S4) and informed our decision 669 tothat we decided to focus only on these 3 parameterization-combinations-here. We now discuss this in more detail 670 below.

671 4.2.1. Meridional surface [I⁻] gradient

672 All the 3 main parameterization-combinations summarized in Table 2, as well as the observations, show enrichment 673 of I' in the surface ocean at low latitudes (Fig. 4Fig. 5). The pathway of transforming IO_A into I' in these oxidized 674 waters is through primary productivity in the photic zone, which resulteds in I accumulation within the mixed layer 675 (Chance et al., 2014). In the low latitudinal surface ocean, weaker vertical mixing resulting from warmer surface 676 temperatures allows L accumulation in the shallow mixed layer (Chance et al., 2014; Moriyaus et al., 2023). - such 677 vertical mixing feature is also simulated by cGENIE (Fig. S7). The pathway of transforming IO3- into I- in these 678 oxidized waters is through primary productivity in the photic zone, which is temperature dependent (Chance et al., 679 2014). A recent north south transect showed the highest surface I enrichment in the oligotrophic, permanently 680 stratified tropical stations (Moriyasu et al., 2023). Therefore, the IO3- flux from deep waters through seasonal mixing 681 may be an important balance to in-situ IO3- reduction rate by primary producers in the high latitudes (Chance et al., 682 2014; Moriyasu et al., 2023). Importantly, this temperature stratification and related vertical mixing trend is also 683 simulated by cGENIE (Fig. S7).-The cGENIE model generates the general pattern of latitudinal surface I⁻ distribution 684 pattern as well; however, overestimation especially in low latitudes may exist, especially in the tropical surface where 685 [I⁻] are is close to 500 nM among all the cGENIE-O2 models (Fig. 4Fig. 5).

686 The cGENIE (internally generated oxygen distributions) vs WOA (imposed distributions) O2 comparison 687 provides evidence that I generated in low [O2] settings may broadly enhance [I] in oxygenated photic waters, with lower and closer-to-observations [I⁻] values in the WOA tunings (Fig. 4Fig. 5). This includes 'lifetime-threshold', 688 689 where O2 only impacts the reductive portion of the iodine cycle, but also the "Fennel" and "reminO2lifetime" 690 where rates of I⁻ oxidation is are also [O₂] dependent. More specifically, most of the elevated (over > 200 nM) surface 691 [I] in cGENIE, occurs in the ETSP and the northern Indian Ocean and corresponds to locations of high primary 692 productivity (Fig. S6). Since the surface ocean [O2] in the model is never below 200 µM, O2-dependent IO3⁻ reduction 693 at the ocean surface is unlikely. Instead, ex situ transport from proximal regions of subsurface anoxia is the most 694 probable source of elevated I. Indeed, the most prominent regions of I enrichment in the model occur near the Peruvian coast and in the Arabian Sea, where ODZs lie below the surface (Fig. S6). More detailed data-model 695 696 comparison among these two areas is limited because the observation data are few (e.g., Farrenkopf and Luther, 2002 697 and Rapp et al., 2020). In contrast, the meridional trend of I⁻ in the surface Atlantic Ocean, where ODZs are less

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developed, exhibits better agreement with both the observation and the Wadley et al., (2020) model (Fig. S6). The overestimation of tropical ocean surface [I⁻] by cGENIE is hence likely to be associated with deficiencies in the simulation of ODZ oxygenation.

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 ImportantlyThat said, modeled overestimations in of surface ocean [I⁻] may be difficult to confirm verify

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 given current observational data densities. Specifically, aIn comparing comparison of observational data to model

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 latitudinal trends which have been masked to only include model grid points with corresponding observations,

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 although-show__the same broad trend of increasing [I⁻] in the low latitudes exists, there-but with are fewer so

 705
 calledincidents of apparent model over-estimation "overestimations" (Fig. \$758). More observations in surface ocean

 706
 [I⁻] from low latitudes is required are needed to better assess the validity of elevated modelled surface ocean [I⁻] in

 707
 some regions.

708 4.2.2. Global and basin-specific iodine depth profiles

All the modelsiodine cycle schemes - (both cGENIE-[O₂] and WOA-forced-) generate a decrease in [I⁻] and increase in [IO₃⁻] from the euphotic zone down to the deep abyssal zone across ocean basins, matching the primary-productiondriven pattern (Fig. 5Fig. 6). As discussed in the previous section, this surface maxima of [I⁻] in the oxygenated water column is the result of biologically mediated reduction or release during cell senescence of phytoplankton. Below the photic zone, [I⁻] is close to zero and IO₃⁻ becomes the dominant species. The deep ocean is mostly oxygenated and has longer water residence times (several millennia, Matsumoto, (2007)) compared to the I⁻ lifetime (<40 years, Tsunogai, (1971)), thus facilitating I⁻ oxidation in the absence of IO₃⁻ reduction in ODZs.

716 We note that there are multiple general discrepancies between observations and data-model output as well as 717 differences between the results of WOA and vs. cGENIE-[O2] parameter tuning. In general, all models reproduce the 718 global average better, relative to the basin-specific profiles. We suggest that the global averaged profiles are a better 719 test of the cGENIE simulations because of sampling biases associated with individual basins. For example, the 720 discrepancy between the model and the observations is are prominent in the Pacific (Fig. 5Fig. 6). The observed Pacific 721 subsurface $[I^-]$ peak mirrors the $[IO_3^-]$ minima that occurs at depths where ODZs are present. This ODZ feature in the 722 averaged Pacific observation profile is likely the result of sampling bias since most of the observations from the Pacific 723 were are from the ETNP (Rue et al., 1997; Moriyasu et al., 2020), and thus do not reflecting the overall iodine 724 distribution in the Pacific Ocean (Fig. S859). As for the meridional trends, sampling bias is again demonstrated in 725 #This conclusion is supported by-depth profiles masked to only include modeled grid points with corresponding 726 observation data. For example, there is a , with a clear mid-depth ODZ feature in the modeled-masked model Pacific 727 depth profiless due to extensive ODZ studies in this region (Fig. 5859). A similar example is from the Indian Oce 728 which we do not show, since most iodine subsurface data come from the ODZ, not normal marine, localities. All this 729 said, while the general features of iodine speciation with depth are generally similar, our data compilation indicates the potential for some basin-specific variations which require further research to validate and mechanistically 730 understand. 731

732 4.2.3. Iodine distribution within ODZs

733 One of the major goals of calibrating the iodine cycle in cGENIE is to be able to simulate the iodine behavior 734 associated with ancient low oxygen settings. To assess this potential, we analyzed model performance for the ETNP 735 (Rue et al., 1997; Moriyasu et al., 2020) where observational data are abundant and the areal extent of the ODZ is 736 sufficiently large to be reflected in the model grid (Fig. 6). Importantly, it should still be noted that cGENIE is best 737 applied to understanding broad scale processes and thus the scope of the ETNP ODZ transect comparison is likely too 738 fine resolution to expect a good match. That said, the The simulated reduction in IO3- to I generally overlaps with the 739 extent of the ODZ (Fig. 6), which provides support for the use of cGENIE to understand the broad distribution of 740 ancient [IO3] and [O2]. Other nNon-threshold model parameterization-combinations (not discussed here but shown in 741 Fig. S4) did not replicate the ODZ feature in iodine speciation.

742 Across-In all model configurations assuming cGENIE-[O2], the most prominent discrepancy is an 743 underestimation of the spatial extent and intensity of the IO3- depletion zone in the ETNP (Fig. 6) because - a 744 <u>consequence</u> of simulated subsurface O_2 deficient area is being notably narrower than that compared to WOA 745 climatology (Fig. 6, Fig. S5). There This is principally a consequence of relatively weak continental margin upwelling 746 and Equatorial divergence, itself a consequence of the low resolution (both horizontally and vertically) model grid 747 that the 3D ocean circulation component is implemented on together with its simplified physics (Marsh et al., 2011) 748 Marsh et al., 2011. Another consequence of the low resolution nature of the cGENIE model grid is that the extent of 749 the ETNP is of the order of the size of an individual grid box and are multiple factors that might affect the performance 750 of simulating the O2 cycle in cGENIE. The model might underestimate the intensity of gyres in the North and Tropical 751 Pacific, resulting in the narrowed ODZs in these areas. Also, the pattern of upwelled nutrients into the surficial 752 Tropical Pacific needs to be tuned to better replicate the productivity and the O2 consumption during remineralization. 753 Another source of uncertainty is that the short term processes, such as seasonal or El Niño driven ODZ variations at 754 the time of sample collection are not replicated in the cGENIE simulations. In addition, due to the low special 755 resolution of cGENIE, the wholeentire ETNP only covers the longitude range of three grids in the model framework 756 (Figure. 1). Meanwhile, the depth resolution is elose toca. 100-200 m per layer -in the surfaceupper ocean, which also 757 limits the finer simulation of ODZ features. Awareness of these uncertainties limitations highlights the importance of 758 the usefocusing use of the model-to understand on regional and global oxygenation features - as supported by proxy 759 constraints and their mechanistic framework during paleoceanographic reconstructions, as opposed to 760 overinterpreting unconstrained high-resolution or local features. We are indeed aware of these uncertainties associated 761 with O2 simulation in cGENIE. However, as a model targeted to assist paleoredox reconstructions, broadly 762 reconstructing ODZ features in a coarse spatial resolution is acceptable.

763 Other data-model misfits might have arisen as a result ofmay be due to-additional IO₃⁻ reduction 764 dependencies not explicitly accounted for in the model. As discussed above, shipboard incubation and radiogenic-765 tracer-spiked rate calculation suggest that IO_3^- reduction is slow in the offshore ETNP ODZ (Hardisty et al., 2021). 766 This could explain why measurable IO_3^- is present in the core of the ETNP ODZ (Fig. 6). This is further exacerbated 767 in the Eastern Tropical South Pacific ODZ, where $[IO_3^-]$ remains above 250 nM in some studies (Cutter et al., 2018) 768 but near detection limits in in others (Rapp et al., 2020), suggesting extreme spatiotemporal variability related to 769 currently unconstrained factors. Further, while the capability of microbes using IO_3^- as an electron acceptor for Commented [AR17]: 10.5194/gmd-4-957-2011 Commented [MOU18R17]: Keyi: add to citation list. 770oxidizing organic matter has been confirmed in laboratory culture experiments (Councell et al., 1997; Reyes-Umana771et al., 2021; Yamazaki et al., 2020; Amachi et al., 2007; Farrenkopf et al., 1997), no study to date has demonstrated772non-O2 dependent controls driving variable IO_3^- reduction rates.

773 An important factor contributing to elevated [I⁻] in ODZs is benthic fluxes and reduction occurring within 774 the uppermost layers of marine sediments (akin to denitrification). To help account for this in our M-scoreM-score 775 and model calibration (see: methods section), 'excess' iodine (>500 nM total iodine) was filtered from our 776 observational dataset. The excess iodine originated from the sediment flux has been observed in ODZ water columns 777 contacting anoxic sediments (Chapman, 1983; Farrenkopf and Luther, 2002; Cutter et al., 2018; Moriyasu et al., 2020; 778 Scholz et al., 2024). We note that excess iodine is occurs explicitly as I⁻, reflecting the limited or lack of oxygen within 779 the ODZ, and is a local-regional phenomenon not yet observed_to persist beyond ODZ settings. Future work can focus 780 on understanding the degree, if any, that excess I' is oxidized to IO3, and hence impacts the I/Ca paleoredox proxy. 781 As such, since our goal is a paleoceanography focused model, cGENIE does not incorporate the benthic flux of I-782 because only IO3⁻ is tracked via the I:CaL/Ca paleoredox proxy.

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784 **4.3.** Applicability of the cGENIE marine iodine cycle to paleo-redox reconstruction

785 Parameter tuning, and the ability to reproduce modern observations, does not by itself offer any guarantee that spatial 786 patterns are being simulated for the 'correct' reason (i.e., specific set and relative importance of mechanisms). This is 787 even more pertinent in the context of the application of a modern-tuned model to paleo redox reconstruction. To test 788 whether our new iodine cycle had predictive power in the geological past, we carried out a deep-time plausibility test. 789 For the paleo plausibility test, we adopted the Cretaceous, pre-OAE2 (ca. 93 Ma) configuration (continental 790 arrangement and ocean bathymetry, wind stress and velocity, and zonal average planetary albedo boundary conditions) 791 from Monteiro et al., (2012). We choose this particular geological interval because the controls on ocean redox have 792 been previously evaluated using the GENIE model (Monteiro et al., 2012; Hülse et al., 2019), the oceanic conditions 793 are much more extensively dysoxic and anoxic than present-day and hence represent a relatively severe test of the 794 model iodine cycle, and a number of I:Cal/Ca proxy measurements are available (Zhou et al., 2015). In order to 795 evaluate the same configuration of the iodine cycle as optimized in this study, we also substituted the temperature 796 independent representation of biological export production and fixed remineralization profile of POM in the water Ridgwell et al., (2007)) for the temperature dependent 797 scheme of Crichton et al. (2021) Ho 798 substituting the biological pump scheme in the model we change the ocean redox landscape compared with e.g., 799 Monteiro et al., (2012). We therefore test a range of different assumptions regarding the ocean PO4 inventory at the 800 time as a means of generating a range of different plausible states of ocean oxygenation. In this, we test: 0.2, 0.4, 0.6, 801 0.8, 1.0, and 1.5 times the mean modern concentration (2.15 mM). We run the model with each of the best-fit (highest 802 M-score) sets of parameter values associated with the 5 different parameterization combinations (but focus on the 803 results of the same 3 combinations we did for comparison against modern), and for each of the varying PO4 inventory 804 assumptions, for 10,000 years to steady-state.

805





844 Most of the parameterization combinations tested reach their highest M scoreM-scores under the assumption 845 of 0.6 0.8 × modern [PO4] (Fig. S9S10). Previous analysis using the same climatological configuration of the GENIE 846 model indicated a PO+ inventory of 1.0 x modern was most consistent with geological redox-related observations prior to OAE2 (Monteiro et al., 2012). However, our assumption here of temperature dependent POM export and 847 848 remineralization leads to higher export and shallower more intense ODZs compared to temperature invariant biological scheme (see: Crichton et al., (2021)). Hence, for a similar degree of ocean anoxia, we would expect the 849 850 need for a slightly lower nutrient inventory, as we indeed find. In terms of the I:CaI/Ca M-score, we find the parameterization-combinations "reminO2lifetime-851 852 hold"<u>'reminO2lifetime_threshold'</u> and "fennel_threshold"<u>'fennel_threshold'</u> better replicate the geological 853 observations compared to the "lifetime threshold" lifetime threshold. In general, WOA derived parameter sets

perform better than those derived from cGENIE [O₂], again hinting at the importance of reducing the uncertainties in
 simulating the modern O₂ cycle in cGENIE. These observations are also largely independent of the ocean PO₄
 inventory assumption. Although we tested additional parameters in this study (Table S1), only those combinations

857 listed in Table 1 ("lifetime - threshold", "lifetime reminO2lifetime", "lifetime - Fennel") perform well in replicating 858 iodine gradients within the modern ocean (Figs. S2-4). Although combinations of parameterizations such as 859 "reminO2lifetime threshold"<u>terminO2lifetime threshold</u> with DOC remineralization, "lifetime inhibition", and 860 "lifetime reminSO4lifetime", also produce elevated M scoreM-scores, these combinations do not perform well in 861 replicating iodine gradients within the modern ocean (Figs. S2-4). Thus, until there is a better mechanistic 862 understanding of IO3- reduction in the modern ocean, the safest choice is arguably to apply the parameterization that 863 best reproduces modern observations and hence retain use of the "threshold" IO3- reduction 864 parameterization for paleo applications.

865 The pre-OAE2 comparison is revealing because it-the observations encapsulates a strong gradient between 866 high and very low I:CaI/Ca (Fig. 7Fig. 3), reflecting respectively, high and low surface ocean concentrations of IO3-867 in the model. All three of the parameterization-combinations we focus on here (just focusing on with WOA-derived 868 parameter values) correctly lead to veryare capable of reproducing low I:CaI/Ca values in the lower latitudinal sections 869 (Demerara Rise, Tarfaya, and Raia del Pedale; Fig. 9), although with a tendency to slightly overestimate seawater 870 IO_3^- depletion (cross-plots in Fig. 9). Low ocean surface $[IO_3^-]$ is duoccurs in the model as a result of e to the existence 871 of a circum-Equatorial band of intense sub-surface anoxia and short transport time to the surface (and hence limited 872 oxidation). In the higher latitudinal sections, including Newfoundland, Eastbourne, and South Ferriby, I:Cal/Ca values 873 tend to be underestimated to varying degrees (Fig. 9). It is possible that the total iodine inventory has varied through 874 Earth history relative to the modern day value (~500 nM), which was adopted for our Cretaceous model. Indeed, the 875 overall underestimated I/Ca by cGENIE might be the result of overall higher Cretaceous total iodine inventory (Zhou 876 et al., 2015; Lu et al., 2018). However, such difference is easily masked by local regional redox variation and is thus 877 difficult to track (Zhou et al., 2015). Due to the lack of evidence otherwise, we assume the average total iodine during 878 the Cretaceous is close to the modern, and the consistent I/Ca underestimation is caused by uncertainty in model 879 simulation. Compared However, compared to the 'lifetime' parameterization, both 'reminO2lifetime' and 'Ffennel' 880 oxidation simulate the I:Cal/Ca values in these high latitudinal sections rather better, with the regression closer to the 881 1:1 line (dotted in Fig. 9). We find this relationship instructional for understanding controls in the modern iodine cycle, 882 which we discuss in more detail below.



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 Figure 9. The pre-OAE2 I:Cal/Ca field derived from cGENIE [IO3-] simulations, and the comparison between

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 modeled and observed I:Cal/Ca from sections.

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We first note that both 'lifetime' and 'fennel' iodine oxidation parameterizations in conjunction with a threshold of iodate reduction and internally generated GENIE-[O₂], give rise to identical parameter values (Table 2). We infer that this is because the modern ocean is predominately well-oxygenated and hence there is little inhibition of I⁻ oxidation in practice. In the Cretaceous environment, although I⁻ oxidation inhibition should be widespread, the M-scores are similar (Fig. 8). The rate of I⁻ oxidation in well oxygenated seawater is likely then critical in explaining elevated <u>I:Cal/Ca</u> values at higher Cretaceous latitudes. However, simply decreasing the lifetime in the modern ocean

893 would result in an under estimation of surface ocean [I']. The 'fennel-threshold' combination under WOA-[O2] reveals 894 interior directly through the biological pump and elevated cellular I:C $(3.5 \times 10^{-4} \text{ mol/mol} \text{ vs. } 1.5 \times 10^{-4} \text{ mol/mol})$. In the 895 896 Cretaceous ocean this combination allows for both sharper latitudinal gradients in [IO3] (and hence I:CaI/Ca) to 897 develop, as well as steeper vertical gradients which allow for non-zero I:CaI/Ca values at low latitudes to be captured 898 (cross-plot in Fig. 9). This slight enhancement of the upper ocean [IO3'] gradient is also apparent in the present-day 899 analysis (Fig. 5Fig. 6). The combination of 'reminO2lifetime' with a reduction threshold works similarly - - a shorter 900 lifetime for I' under oxic conditions offset in the modern ocean by elevated cellular I:C (Table 2). However, in this 901 case, our gridded parameter search identifies the trade-off as producing the highest M-score for both modelled and 902 WOA oxygen distributions.

903 What we learn from this is that the cGENIE iodine cycle tuned to modern observations has predictive power 904 under a very different state of ocean oxygenation (and circulation and operation of biological pump). However, this 905 is not true for every choice of parameterization, and the simple 'lifetime-threshold' combination, which when 906 calibrated was statistically almost the best representation of the iodine cycle, was unable to reproduce the latitudinal 907 H:CaI/Ca gradients in the Cretaceous ocean. Shortening the lifetime (and adding an inhibition term) together with 908 increasing the assumed cellular I:C, could maintains fidelity in simulating the modern ocean fidelity whilst much 909 better capturing Cretaceous I:Cal/Ca. That even better representations of Cretaceous I:Cal/Ca were possible but at the 910 expense of reproducing modern observations adequately hints that improvements in our mechanistic understanding 911 are needed, although all of the above assumes that the simulation of the Cretaceous redox landscape is plausible.

912 One caveat to our paleo comparison is that becasuse the residence time of iodine in the ocean is only around 913 300 kyr, the dissolved iodine inventory of the Cretaceous ocean could have deviated substantially from the modern 914 seawater value that we assume here (500 nM). It is possible that the total iodine inventory has varied through Earth 915 history relative to the modern day value (~500 nM),. Indeed, the One possible explanation for the overall 916 underestimation of ed I:CaI/Ca by cGENIE might then be the result of overall higher that the Cretaceous total iodine 917 inventory was higher than modern (Zhou et al., 2015; Lu et al., 2018). Even a moderate increase (ca. 20-40%) in the 918 ocean iodine inventory (which we did not test here) would presumably act to increase the slope of the regression lines 919 for the parameterization-combinations 'fennel-threshold' and 'reminO2lifetime-threshold' and bring them close to 920 the 1:1 line (Figure. 9). However, such difference is easily masked by local regional redox variation and is unlikely a 921 primary control on IO2-availability in reducing oceans (Zhou et al., 2015). Due to the lack of evidence otherwise, we 922 assume the average total iodine during the Cretaceous is close to the modern, and the consistent I:Cal/Ca 923 underestimation is caused by uncertainty in model simulation. Another uncertainty concerns the potential impact on 924 carbonate I/Ca ratios of different-from-modern dissolved calcium concentrations in the Cretaceous ocean. The calcite 925 crystal growth experiments of Zhou et al. [2014] on which we base our model-simulated I/Ca values were carried out 926 in synthetic solutions and not seawater. The partition coefficient K_D was calculated as [I/Ca]/[IO₃] and as a result our 927 model formulation is independent of the ambient Ca²⁺ concentration. New I/Ca carbonate precipitation experiments 928 that explicitly account for the geological range in seawater composition would help increase confidence in the model-929 simulated I/Ca values and hence improve the model simulation comparison possible with geological observations.

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Choice of marine iodine cycle representation in cGENIE 4.4.

932 The risk in tuning model parameters to fit some data target is always that the processes being tuned are distorted to 933 accommodate an underlying structural error with the model. In this paper we illustrated how sub-regional-scale ocean 934 oxygenation features such as the Eastern Tropical North Pacific ODZ (Figure. 67) are poorly represented at the 935 resolution of the cGENIE Earth system model, and how this can lead to deficiencies in the simulation of the marine 936 iodine cycle in these regions. Indeed, we found an improved statistical fit to observed iodine speciation data when we 937 imposed a dissolved oxygen climatology to the model the grid rather than use the internally -general model $[O_2]$ fields 938 (Table 2). In the case of the 'Fennel' parameterization-combination, we end up with two different sets of parameter 939 values – one associated with the best fit to iodine speciation data under cGENIE projected $[O_2]$, and one under WOA 940 imposed [O₂]. Selecting this particular scheme for paleo applications would appear to create a conundrum – whether 941 (a) to choose the cGENIE $[O_2]$ derived parameter value set₇ and accept that model deficiencies were being implicitly 942 corrected for in simulating marine iodine cycling and hence imposing a potential bias on paleo I/Ca, or (b) accept 943 WOA [O2] derived parameter values. The choice would arguably be to accept the less potentially biased parameter 944 value set (b), particularly as the relatively small-scale ODZ features of the modern ocean may be expanded to the 945 regional-to-global scale in deeper time oceans and hence the redox features driving iodate reduction would become 946 much more tractable for a model of the resolution of cGENIE. 947 However, we note that this choice does not exist for parameterization-combinations 'lifetime-threshold' and 948 'reminO2lifetime-threshold'. In both cases the sets of optimal parameter values are the same, whether derived from 949 <u>cGENIE</u> projected $[O_2]$ or WOA imposed $[O_2]$ (Table 2). This gives us confidence that both these schemes are 950 relatively agnostic to the details of how the modern oxygen cycle is simulated, and that errors in the simulation of sub-951 regional-scale ocean oxygenation features are not critical. (We note that iodine speciation data is still fitted better 952 under WOA imposed [O2].) The choice of how to best represent the marine iodine cycle in cGENIE is then a choice 953 between 'lifetime-threshold', which has been used in a similar form previously (Lu et al., 2018) and is capable of the 954 best fit to modern observations, or 'reminO2lifetime-threshold'. While 'lifetime-threshold' might be preferable for 955 more recent geological time and ocean ODZs similar to the present-day, it fails to account for I/Ca contrasts under 956 more extreme redox gradients (Figure. 9). Although more difficult to interpret mechanistically, in suppressing the rate 957 of I oxidation under low ambient [O2], 'reminO2lifetime-threshold' appears to be the more appropriate paleo choice. 958 We note that to achieve improved simulation of sub-regional-scale features of the ocean redox landscape, 959 future paleo cGENIE model I/Ca applications could make use of the coupled GENIE-PLASIM climate configuration 960 (Holden et al., 2016). (Holden et al., 2016). In this, the ocean circulation model is implemented on a higher resolution 961 grid $(64 \times 32 \text{ with } 32 \text{ vertical levels})$ and includes a coupled atmospheric GCM component (hence enabling interannual 962 variability). It is also capable of being configured with different continental configurations for paleo climate questions 963 (e.g. Keery et al., 2018). 964

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965 5. Conclusions

966 Using the cGENIE Earth system model, we performed a series of ensemble experiments to determine --parameter 967 searches for suitable parameterizations to represent the marine iodine cycle and identified the best performing 968 combinations of parameter values combinations in each case. We found that the optimized parameter values 969 associated with IO3⁻ planktonic uptake, water column IO3⁻ reduction and I⁻ oxidation, plus IO3⁻ planktonic uptake (and 970 subsequence I release during remineralization), are all fell within the range of field and experiment observations and 971 hence could be considered plausible. Three iodine cycling parameterization _combinations, 'lifetime-threshold', 972 'reminO2lifetime-threshold', and 'fennel-threshold' emerged were considered as viable candidates following our tests 973 of the global ocean model M-score, and model-data comparisons made across specific iodine gradients 974 (euphotic latitudinal distribution, depth distribution, and ODZ distribution). We further evaluated the plausibility of 975 our parameterizations and their paleo and ocean deoxygenation applicability by contrasting forward-proxy model generated I:CaI/Ca values against observations, taking the (pre-OAE2) Cretaceous redox landscape as a case study. 976 977 While some model-data discrepancies emerge for both modern and paleo, these highlight that future observational 978 and/or experimental work is necessary to better constrain modern iodine cycling mechanisms and related 979 spatiotemporal heterogeneities. While we further identified highlighted the importance of improving the simulation 980 of dissolved oxygen distributions in models, equally, we also found that our conclusions regarding preferred 981 parameterizations and even specific parameter values, was not overly dependent on the specific details of the simulated modern OMZs. Overall, our analysis gives us a degree of confidence that carbonate I:CaI/Ca ratios can be interpreted 982 983 in terms of past ocean oxygenation using models such as 'cGENIE'. While while some model-data discrepancies 984 emerge for both modern and paleo, these highlight that future observational and/or experimental work is necessary to 985 better constrain modern iodine cycling mechanisms and related spatiotemporal heterogeneities but that representations 986 of the marine iodine cycle in an Earth system model based on modern observations and mechanistic knowledge appear 987 transferrable to interpreting the geological record of the I/Ca oxygenation proxy. 988

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989 990	Competing interests. The contact author has declared that none of the authors has any competing interests.		
991	Author contributions. KC, AR, and DH conceptualized the research presented in this paper. DH and AR acquired		
992	funding to support the study. AR developed iodine tracer and associated biogeochemical mechanisms in cGENIE. KC		
993	and DH designed model performance under the modern ocean configurations. KC compiled the modern ocean iodine		
994	database and the Cretaceous I:Cal/Ca data. KC ran the modern-ocean cGENIE analysis and performed model-data		
995	comparison. AR performed model-data evaluation for Cretaceous configurations. KC prepared the manuscript with		
996	contributions from all co-authors.		
997			
998	Acknowledgements. Funding support for DH and KC comes from NSF OCE 1829406. AR acknowledges support from		
999	National Science Foundation grants EAR-2121165 and OCE- 2244897, and the NASA Interdisciplinary Consortia		
1000	for Astrobiology Research (ICAR) Program (80NSSC21K0594).		
1001			
1002	Code availability. The code for the version of the 'muffin' release of the cGENIE Earth system model used in 👘 🔸	<	F
1003	this paper, is tagged as v0.9.54, and is assigned a DOI: 10.5281/zenodo.13376310.		F
1004	Configuration files for the specific experiments presented in the paper can be found in the		
1005	directory: genie-userconfigs/PUBS/published/Cheng_et_al.Biogeosciences.2024. Details of		
1006	the experiments, plus the command line needed to run each one, are given in the readme.txt		
1007	file in that directory. All other configuration files and boundary conditions are provided as part		
1008	of the code release.		
1009	A manual detailing code installation, basic model configuration, tutorials covering various		
1010	aspects of model configuration, experimental design, and output, plus the processing of		
1011	results, is assigned a DOI: 10.5281/zenodo.1407657. The code for the version of the 'muffin' release of		
1012	the cGENIE Earth system model used in this paper, is tagged as v0.9.13, and is assigned a DOI:		
1013	10.5281/zenodo.3999080.		
1014	Configuration files for the specific experiments presented in the paper can be found in the directory:		
1015	genie-userconfigs/PUBS/chengetal.BG.2020. Details of the experiments, plus the command line		
1016	needed to run each one, are given in the readme.txt file in that directory. All other configuration		
1017	files and boundary conditions are provided as part of the code release.		
1018	A manual detailing code installation, basic model configuration, tutorials covering various		
1019	aspects of model configuration, experimental design, and output, plus the processing of results, is		
1020	assigned a DOI: 10.5281/zenodo.4305997.		C
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- 1022 Amachi, S., Kawaguchi, N., Muramatsu, Y., Tsuchiya, S., Watanabe, Y., Shinoyama, H., and Fujii, T.:
- 1023 Dissimilatory iodate reduction by marine Pseudomonas sp. strain SCT, Appl. Environ. Microbiol., 73, 5725–5730,
- 1024 https://doi.org/10.1128/AEM.00241-07, 2007.
- 1025 Boscolo-Galazzo, F., Crichton, K. A., Ridgwell, A., Mawbey, E. M., Wade, B. S., and Pearson, P. N.: Temperature
- 1026 controls carbon cycling and biological evolution in the ocean twilight zone, Science (80-.)., 371, 1148–1152,
 1027 https://doi.org/10.1126/science.abb6643, 2021.
- 1028 Bowman, C. N., Lindskog, A., Kozik, N. P., Richbourg, C. G., Owens, J. D., and Young, S. A.: Integrated
- 1029 sedimentary, biotic, and paleoredox dynamics from multiple localities in southern Laurentia during the late Silurian
- 1030 (Ludfordian) extinction event, Palaeogeogr. Palaeoclimatol. Palaeoecol., 553, 109799,
- 1031 https://doi.org/10.1016/j.palaeo.2020.109799, 2020.
- 1032 Broecker, W. S. and Peng, T. H.: Tracers in the Sea., https://doi.org/10.1016/0016-7037(83)90075-3, 1983.
- 1033 Campos, M. L. A. M., Farrenkopf, A. M., Jickells, T. D., and Luther, G. W.: A comparison of dissolved iodine
- 1034 cycling at the Bermuda Atlantic Time-series station and Hawaii Ocean Time-series station, Deep. Res. Part II Top.
- 1035 Stud. Oceanogr., 43, 455–466, https://doi.org/10.1016/0967-0645(95)00100-x, 1996.
- 1036 Chance, R., Weston, K., Baker, A. R., Hughes, C., Malin, G., Carpenter, L., Meredith, M. P., Clarke, A., Jickells, T.
- 1037 D., Mann, P., and Rossetti, H.: Seasonal and interannual variation of dissolved iodine speciation at a coastal
- 1038 Antarctic site, Mar. Chem., 118, 171–181, https://doi.org/10.1016/j.marchem.2009.11.009, 2010.
- 1039 Chance, R., Baker, A. R., Carpenter, L., and Jickells, T. D.: The distribution of iodide at the sea surface, Environ.
- 1040 Sci. Process. Impacts, 16, 1841–1859, https://doi.org/10.1039/c4em00139g, 2014.
- 1041 Chance, R., Tinel, L., Sherwen, T., Baker, A., Bell, T., Brindle, J., Campos, M., Croot, P., Ducklow, H., He, P.,
- 1042 Hoogakker, B., Hopkins, F., Hughes, C., Jickells, T., Loades, D., Reyes Macaya, D., Mahajan, A., Malin, G.,
- 1043 Phillips, D., Sinha, A., Sarkar, A., Roberts, I., Roy, R., Song, X., Winklebauer, H., Wuttig, K., Yang, M., Zhou, P.,
- and Carpenter, L.: Global sea-surface iodide observations, 1967-2018., https://doi.org/doi:10.5285/7e77d6b9-83fb41e0-e053-6c86abc069d0, 2019.
- 1046 Chapman, P.: Changes in iodine speciation in the Benguela current upwelling system, Deep Sea Res. Part A,
- 1047 Oceanogr. Res. Pap., 30, 1247–1259, https://doi.org/10.1016/0198-0149(83)90083-3, 1983.
- Councell, T. B., Landa, E. R., and Lovley, D. R.: Microbial reduction of iodate, Water. Air. Soil Pollut., 100, 99–
 106, 1997.
- 1050 Crichton, K. A., Wilson, J. D., Ridgwell, A., and Pearson, P. N. P. N.: Calibration of temperature-dependent ocean
- 1051 microbial processes in the cGENIE.muffin (v0.9.13) Earth system model, Geosci. Model Dev., 14, 125–149,
- 1052 https://doi.org/10.5194/gmd-14-125-2021, 2021.
- 1053 Cutter, G. A., Moffett, J. G., Nielsdóttir, M. C., and Sanial, V.: Multiple oxidation state trace elements in suboxic
- 1054 waters off Peru: In situ redox processes and advective/diffusive horizontal transport, Mar. Chem., 201, 77–89,
- 1055 https://doi.org/10.1016/j.marchem.2018.01.003, 2018.
- 1056 Dalsgaard, T., Stewart, F. J., Thamdrup, B., De Brabandere, L., Revsbech, N. P., Ulloa, O., Canfield, D. E., and
- 1057 Delong, E. F.: Oxygen at nanomolar levels reversibly suppresses process rates and gene expression in anammox and
- 1058 denitrification in the oxygen minimum zone off Northern Chile, MBio, 5, 1–14,

- 1059 https://doi.org/10.1128/mBio.01966-14, 2014.
- 1060 Ding, Y., Sun, W., Liu, S., Xie, J., Tang, D., Zhou, X., Zhou, L., Li, Z., Song, J., Li, Z., Xu, H., Tang, P., Liu, K.,
- 1061 Li, W., and Chen, D.: Low oxygen levels with high redox heterogeneity in the late Ediacaran shallow ocean:
- 1062 Constraints from I/(Ca + Mg) and Ce/Ce* of the Dengying Formation, South China, Geobiology, 20, 790–809,
- 1063 https://doi.org/10.1111/gbi.12520, 2022.
- 1064 Edwards, A. and Truesdale, V. W.: Regeneration of inorganic iodine species in loch etive, a natural leaky incubator,
- 1065 Estuar. Coast. Shelf Sci., 45, 357–366, https://doi.org/10.1006/ecss.1996.0185, 1997.
- 1066 Edwards, C. T., Fike, D. A., Saltzman, M. R., Lu, W., and Lu, Z.: Evidence for local and global redox conditions at
- 1067 an Early Ordovician (Tremadocian) mass extinction, Earth Planet. Sci. Lett., 481, 125–135,
- 1068 https://doi.org/10.1016/j.epsl.2017.10.002, 2018.
- Edwards, N. R. and Marsh, R.: Uncertainties due to transport-parameter sensitivity in an efficient 3-D ocean-climate
 model, Clim. Dyn., 415–433, https://doi.org/10.1007/s00382-004-0508-8, 2005.
- 10/0 model, Chin. Dyn., 415–455, https://doi.org/10.1007/s00502-004-0500-0, 2005.
- 1071 Elderfield, H. and Truesdale, V. W.: On the biophilic nature of iodine in seawater, Earth Planet. Sci. Lett., 50, 105–
- 1072 114, https://doi.org/10.1016/0012-821X(80)90122-3, 1980.
- 1073 Fang, H., Tang, D., Shi, X., Zhou, L., Zhou, X., Wu, M., Song, H., and Riding, R.: Early Mesoproterozoic Ca-
- 1074 carbonate precipitates record fluctuations in shallow marine oxygenation, Precambrian Res., 373, 106630,
- 1075 https://doi.org/10.1016/j.precamres.2022.106630, 2022.
- 1076 Farrenkopf, A. M. and Luther, G. W.: Iodine chemistry reflects productivity and denitrification in the Arabian Sea:
- Evidence for flux of dissolved species from sediments of western India into the OMZ, Deep. Res. Part II Top. Stud.
 Oceanogr., 49, 2303–2318, https://doi.org/10.1016/S0967-0645(02)00038-3, 2002.
- 1079 Farrenkopf, A. M., Dollhopf, M. E., Chadhain, S. N., Luther, G. W., and Nealson, K. H.: Reduction of iodate in
- 1080 seawater during Arabian Sea shipboard incubations and in laboratory cultures of the marine bacterium Shewanella
- 1081 putrefaciens strain MR-4, Mar. Chem., 57, 347–354, https://doi.org/10.1016/S0304-4203(97)00039-X, 1997.
- Fennel, K., Follows, M., and Falkowski, P. G.: The co-evolution of the nitrogen, carbon and oxygen cycles in the
 Proterozoic ocean, Am. J. Sci., 305, 526–545, 2005.
- 1084 Garcia, H. E., Weathers, K., Paver, C. R., Smolyar, I., Boyer, T. P., Locarnini, R. A., Zweng, M. M., Mishonov, A.
- 1085 V., Baranova, O. K., Seidov, D., and Reagan, J. R.: Volume 3: Dissolved Oxygen, Apparent Oxygen Utilization, and
- 1086 Oxygen Saturation. A. Mishonov Technical Ed.; NOAA Atlas NESDIS 83, World Ocean Atlas 2018, 38 pp., 2018.
- 1087 Hardisty, D. S., Lu, Z., Planavsky, N. J., Bekker, A., Philippot, P., Zhou, X., and Lyons, T. W.: An iodine record of
- Paleoproterozoic surface ocean oxygenation, Geology, 42, 619-622, https://doi.org/10.1130/G35439.1, 2014.
- 1089 Hardisty, D. S., Lu, Z., Bekker, A., Diamond, C. W., Gill, B. C., Jiang, G., Kah, L. C., Knoll, A. H., Loyd, S. J.,
- 1090 Osburn, M. R., Planavsky, N. J., Wang, C., Zhou, X., and Lyons, T. W.: Perspectives on Proterozoic surface ocean
- redox from iodine contents in ancient and recent carbonate, Earth Planet. Sci. Lett., 463, 159–170,
- 1092 https://doi.org/10.1016/j.epsl.2017.01.032, 2017.
- 1093 Hardisty, D. S., Horner, T. J., Wankel, S. D., Blusztajn, J., and Nielsen, S. G.: Experimental observations of marine
- 1094 iodide oxidation using a novel sparge-interface MC-ICP-MS technique, Chem. Geol., 532, 119360,
- 1095 https://doi.org/10.1016/j.chemgeo.2019.119360, 2020.

- 1096 Hardisty, D. S., Horner, T. J., Evans, N., Moriyasu, R., Babbin, A. R., Wankel, S. D., Moffett, J. W., and Nielsen, S.
- 1097 G.: Limited iodate reduction in shipboard seawater incubations from the Eastern Tropical North Pacific oxygen
- 1098 deficient zone, Earth Planet. Sci. Lett., 554, 116676, https://doi.org/10.1016/j.epsl.2020.116676, 2021.
- 1099 Hashim, M. S., Burke, J. E., Hardisty, D. S., and Kaczmarek, S. E.: Iodine incorporation into dolomite:
- 1100 Experimental constraints and implications for the iodine redox proxy and Proterozoic Ocean, Geochim. Cosmochim.
- 1101 Acta, 338, 365–381, https://doi.org/10.1016/j.gca.2022.10.027, 2022.
- 1102 He, P., Hou, X., and Aldahan, A.: Iodine isotopes species fingerprinting environmental conditions in surface water
- along the northeastern Atlantic Ocean, Sci. Rep., 3, 1–9, https://doi.org/10.1038/srep02685, 2013.
- 1104 Hepach, H., Hughes, C., Hogg, K., Collings, S., and Chance, R.: Senescence as the main driver of iodide release
- from a diverse range of marine phytoplankton, Biogeosciences, 17, 2453–2471, https://doi.org/10.5194/bg-17-24532020, 2020.
- 1107 Holden, P. B., Edwards, N. R., Fraedrich, K., Kirk, E., Lunkeit, F., and Zhu, X.: PLASIM-GENIE v1.0: A new
- intermediate complexity AOGCM, Geosci. Model Dev., 9, 3347–3361, https://doi.org/10.5194/gmd-9-3347-2016,
 2016.
- 1110 Hughes, C., Barton, E., Hepach, H., Chance, R., Wadley, M. R., Pickering, M. D., Hogg, K., Pommerening-r, A.,
- 1111 Stevens, D. P., and Jickells, T. D.: Oxidation of iodide to iodate by cultures of marine ammonia-oxidising bacteria,
- 1112 Mar. Chem., 234, https://doi.org/10.1016/j.marchem.2021.104000, 2021.
- 1113 Hülse, D., Arndt, S., and Ridgwell, A.: Mitigation of Extreme Ocean Anoxic Event Conditions by Organic Matter
- 1114 Sulfurization, Paleoceanogr. Paleoclimatology, 34, 476–489, https://doi.org/10.1029/2018PA003470, 2019.
- 1115 Jia-zhong, Z. and Whitfield, M.: KINETICS OF INORGANIC REDOX REACTIONS IN SEAWATER I . The
- 1116 reduction of iodate by bisulphide Micro-organisms play a dominant role in the diagenesis of organic-rich sediments .
- The oxidative breakdown of the organic matter, with the accom- panying reductio, Science (80-.)., 19, 121–137,
 1118 1986.
- 1119 Jiang, Z., Cui, M., Qian, L., Jiang, Y., Shi, L., Dong, Y., Li, J., and Wang, Y.: Abiotic and Biotic Reduction of
- 1120 Iodate Driven by Shewanella oneidensis MR-1, Environ. Sci. Technol., https://doi.org/10.1021/acs.est.3c06490,
 1121 2023.
- 1122 Keery, J. S., Holden, P. B., and Edwards, N. R.: Sensitivity of the Eocene climate to CO2 and orbital variability,
- 1123 Clim. Past, 14, 215–238, https://doi.org/10.5194/cp-14-215-2018, 2018.
- 1124 Kerisit, S. N., Smith, F. N., Saslow, S. A., Hoover, M. E., Lawter, A. R., and Qafoku, N. P.: Incorporation Modes of
- 1125 Iodate in Calcite, Environ. Sci. Technol., 52, 5902–5910, https://doi.org/10.1021/acs.est.8b00339, 2018.
- 1126 Laws, E. A., Falkowski, P. G., Smith, W. O., Ducklow, H., and James J McCarthy: Temperature effects on export
- 1127 production in the open ocean, Global Biogeochem. Cycles, 14, 1231–1246, 2000.
- 1128 Liu, A., Tang, D., Shi, X., Zhou, X., Zhou, L., Shang, M., Li, Y., and Fang, H.: Mesoproterozoic oxygenated deep
- 1129 seawater recorded by early diagenetic carbonate concretions from the Member IV of the Xiamaling Formation,
- 1130 North China, Precambrian Res., 341, 105667, https://doi.org/10.1016/j.precamres.2020.105667, 2020.
- 1131 Lu, W., Ridgwell, A., Thomas, E., Hardisty, D. S., Luo, G., Algeo, T. J., Saltzman, M. R., Gill, B. C., Shen, Y.,
- 1132 Ling, H. F., Edwards, C. T., Whalen, M. T., Zhou, X., Gutchess, K. M., Jin, L., Rickaby, R. E. M., Jenkyns, H. C.,

- 1133 Lyons, T. W., Lenton, T. M., Kump, L. R., and Lu, Z.: Late inception of a resiliently oxygenated upper ocean,
- 1134 Science (80-.)., 361, 174–177, https://doi.org/10.1126/science.aar5372, 2018.
- 1135 Lu, W., Rickaby, R. E. M., Hoogakker, B. A. A., Rathburn, A. E., Burkett, A. M., Dickson, A. J., Martínez-Méndez,
- 1136 G., Hillenbrand, C. D., Zhou, X., Thomas, E., and Lu, Z.: I/Ca in epifaunal benthic foraminifera: A semi-
- 1137 quantitative proxy for bottom water oxygen in a multi-proxy compilation for glacial ocean deoxygenation, Earth
- 1138 Planet. Sci. Lett., 533, https://doi.org/10.1016/j.epsl.2019.116055, 2020a.
- 1139 Lu, W., Dickson, A. J., Thomas, E., Rickaby, R. E. M., Chapman, P., and Lu, Z.: Refining the planktic foraminiferal
- 1140 I/Ca proxy: Results from the Southeast Atlantic Ocean, Geochim. Cosmochim. Acta, 287, 318-327,
- 1141 https://doi.org/10.1016/j.gca.2019.10.025, 2020b.
- 1142 Lu, Z., Jenkyns, H. C., and Rickaby, R. E. M.: Iodine to calcium ratios in marine carbonate as a paleo-redox proxy
- 1143 during oceanic anoxic events, Geology, 38, 1107–1110, https://doi.org/10.1130/G31145.1, 2010.
- 1144 Lu, Z., Hoogakker, B. A. A., Hillenbrand, C.-D., Zhou, X., Thomas, E., Gutchess, K. M., Lu, W., Jones, L., and
- 1145 Rickaby, R. E. M.: Oxygen depletion recorded in upper waters of the glacial Southern Ocean, Nat. Commun., 7, 1–
- 1146 8, https://doi.org/10.1038/ncomms11146, 2016.
- Luther, G. W.: Review on the physical chemistry of iodine transformations in the oceans, Front. Mar. Sci., 10, 1–16,
 https://doi.org/10.3389/fmars.2023.1085618, 2023.
- 1149 Luther, G. W. and Campbell, T.: Iodine speciation in the water column of the Black Sea, Deep Sea Res. Part A.
- 1150 Oceanogr. Res. Pap., 38, S875–S882, https://doi.org/10.1016/s0198-0149(10)80014-7, 1991.
- 1151 Luther, G. W., Wu, J., and Cullen, J. B.: Redox Chemistry of Iodine in Seawater, Adv. Chem., 135–155, 1995.
- 1152 Luther, W.: Iodine Chemistry in the Water the Chesapeake Bay : Evidence Iodine Forms Column of for Organic,
- 1153 Esruarine, Coast. Shelf Sci., 32, 267–279, 1991.
- 1154 Marsh, R., Müller, S. A., Yool, A., and Edwards, N. R.: Incorporation of the C-GOLDSTEIN efficient climate
- 1155 model into the GENIE framework: "eb_go_gs" configurations of GENIE, Geosci. Model Dev., 4, 957-
- 1156 992, https://doi.org/10.5194/gmd-4-957-2011, 2011.
- 1157 Martin, T. S., Primeau, F., and Casciotti, K. L.: Modeling oceanic nitrate and nitrite concentrations and isotopes
- 1158 using a 3-D inverse N cycle model, Biogeosciences, 347–367, 2019.
- 1159 Matsumoto, K.: Radiocarbon-based circulation age of the world oceans, J. Geophys. Res. Ocean., 112, 1–7,
- 1160 https://doi.org/10.1029/2007JC004095, 2007.
- 1161 Monteiro, F. M., Pancost, R. D., Ridgwell, A., and Donnadieu, Y.: Nutrients as the dominant control on the spread
- 1162 of anoxia and euxinia across the Cenomanian-Turonian oceanic anoxic event (OAE2): Model-data comparison,
- 1163 Paleoceanography, 27, https://doi.org/10.1029/2012PA002351, 2012.
- 1164 Moriyasu, R., Evans, N., Bolster, K. M., Hardisty, D. S., and Moffett, J. W.: The Distribution and Redox Speciation
- 1165 of Iodine in the Eastern Tropical North Pacific Ocean, Global Biogeochem. Cycles, 34, 1–23,
- 1166 https://doi.org/10.1029/2019GB006302, 2020.
- 1167 Moriyasu, R., Bolster, K. M., Hardisty, D. S., Kadko, D. C., Stephens, M. P., and Moffett, J. W.: Meridional survey
- 1168 of the Central Pacific reveals iodide accumulation in equatorial surface waters and benthic sources in the abyssal
- 1169 plain, Global Biogeochem. Cycles, https://doi.org/10.1029/2021GB007300, 2023.

- 1170 Panchuk, K., Ridgwell, A., and Kump, L. R.: Sedimentary response to Paleocene-Eocene thermal maximum carbon
- 1171 release: A model-data comparison, Geology, 36, 315–318, https://doi.org/10.1130/G24474A.1, 2008.
- 1172 Podder, J., Lin, J., Sun, W., Botis, S. M., Tse, J., Chen, N., Hu, Y., Li, D., Seaman, J., and Pan, Y.: Iodate in calcite
- 1173 and vaterite: Insights from synchrotron X-ray absorption spectroscopy and first-principles calculations, Geochim.
- 1174 Cosmochim. Acta, 198, 218–228, https://doi.org/10.1016/j.gca.2016.11.032, 2017.
- 1175 Pohl, A., Lu, Z., Lu, W., Stockey, R. G., Elrick, M., Li, M., Desrochers, A., Shen, Y., He, R., Finnegan, S., and
- 1176 Ridgwell, A.: Vertical decoupling in Late Ordovician anoxia due to reorganization of ocean circulation, Nat.
- 1177 Geosci., 14, 868–873, https://doi.org/10.1038/s41561-021-00843-9, 2021.
- 1178 Pohl, A., Ridgwell, A., Stockey, R. G., Thomazo, C., Keane, A., Vennin, E., and Scotese, C. R.: Continental
- 1179 configuration controls ocean oxygenation during the Phanerozoic, Nature, 608, 523–527,
- 1180 https://doi.org/10.1038/s41586-022-05018-z, 2022.
- Rapp, I., Schlosser, C., Barraqueta, J. M., Wenzel, B., Lüdke, J., and Scholten, J.: Controls on redox-sensitive trace
 metals in the Mauritanian oxygen minimum zone, 4157–4182, 2019.
- 1183 Rapp, I., Schlosser, C., Browning, T. J., Wolf, F., Le Moigne, F. A. C., Gledhill, M., and Achterberg, E. P.: El Niño-
- 1184 Driven Oxygenation Impacts Peruvian Shelf Iron Supply to the South Pacific Ocean, Geophys. Res. Lett., 47,
- 1185 https://doi.org/10.1029/2019GL086631, 2020.
- 1186 Reinhard, C. T. and Planavsky, N. J.: The History of Ocean Oxygenation, Ann. Rev. Mar. Sci., 14, 331–353,
- 1187 https://doi.org/10.1146/annurev-marine-031721-104005, 2022.
- 1188 Reinhard, C. T., Planavsky, N. J., Olson, S. L., Lyons, T. W., and Erwin, D. H.: Earth's oxygen cycle and the
- evolution of animal life, Proc. Natl. Acad. Sci. U. S. A., 113, 8933–8938, https://doi.org/10.1073/pnas.1521544113,
 2016.
- 1191 Reinhard, C. T., Olson, S. L., Turner, S. K., Pälike, C., Kanzaki, Y., and Ridgwell, A.: Oceanic and atmospheric
- methane cycling in the cGENIE Earth system model, Geosci. Model Dev., 1–45, https://doi.org/10.5194/gmd-202032, 2020.
- 1194 Remmelzwaal, S. R. C., Dixon, S., Parkinson, I. J., Schmidt, D. N., Monteiro, F. M., Sexton, P., Fehr, M. A.,
- 1195 Peacock, C., Donnadieu, Y., and James, R. H.: Investigating Ocean Deoxygenation During the PETM Through the
- 1196 Cr Isotopic Signature of Foraminifera, Paleoceanogr. Paleoclimatology, 34, 917–929,
- 1197 https://doi.org/10.1029/2018PA003372, 2019.
- 1198 Reyes-Umana, V., Henning, Z., Lee, K., Barnum, T. P., and Coates, J. D.: Genetic and phylogenetic analysis of
- 1199 dissimilatory iodate-reducing bacteria identifies potential niches across the world's oceans, ISME J.,
- 1200 https://doi.org/10.1038/s41396-021-01034-5, 2021.
- 1201 Ridgwell, A., Hargreaves, J. C., Edwards, N. R., Annan, J. D., Lenton, T. M., Marsh, R., Yool, A., and Watson, A.:
- 1202 Marine geochemical data assimilation in an efficient Earth system model of global biogeochemical cycling,
- 1203 Biogeosciences, 4, 87–104, https://doi.org/10.5194/bg-4-87-2007, 2007.
- 1204 Rue, E. L., Smith, G. J., Cutter, G. A., and Bruland, K. W.: The response of trace element redox couples to suboxic
- conditions in the water column, Deep. Res. Part I Oceanogr. Res. Pap., 44, 113–134, https://doi.org/10.1016/S0967 0637(96)00088-X, 1997.

- 1207 Schnur, A. A., Sutherland, K. M., Hansel, C. M., and Hardisty, D. S.: Rates and pathways of iodine speciation
- 1208 transformations at the Bermuda Atlantic Time Series, Front. Mar. Sci., 1–14,
- 1209 https://doi.org/10.3389/fmars.2023.1272870, 2024.
- 1210 Scholz, F., Hardisty, D. S., and Dale, A. W.: Early Diagenetic Controls on Sedimentary Iodine Release and Iodine-
- 1211 To-Organic Carbon Ratios in the Paleo-Record, Global Biogeochem. Cycles, 38,
- 1212 https://doi.org/10.1029/2023GB007919, 2024.
- 1213 Shang, M., Tang, D., Shi, X., Zhou, L., Zhou, X., Song, H., and Jiang, G.: A pulse of oxygen increase in the early
- 1214 Mesoproterozoic ocean at ca. 1.57–1.56 Ga, Earth Planet. Sci. Lett., 527, 115797,
- 1215 https://doi.org/10.1016/j.epsl.2019.115797, 2019.
- 1216 Ştreangă, I.-M., Repeta, D. J., Blusztajn, J. S., and Horner, T. J.: Speciation and cycling of iodine in the subtropical
- 1217 North Paci fi c Ocean, Front. Mar. Sci., 1–15, https://doi.org/10.3389/fmars.2023.1272968, 2024.
- 1218 Tang, D., Fang, H., Shi, X., Liang, L., Zhou, L., Xie, B., Huang, K., Zhou, X., Wu, M., and Riding, R.:
- 1219 Mesoproterozoic Molar Tooth Structure Related to Increased Marine Oxygenation, J. Geophys. Res.
- 1220 Biogeosciences, 128, 1–18, https://doi.org/10.1029/2022jg007077, 2023.
- 1221 Thamdrup, B., Dalsgaard, T., and Revsbech, N. P.: Widespread functional anoxia in the oxygen minimum zone of
- 1222 the Eastern South Pacific, Deep. Res. Part I Oceanogr. Res. Pap., 65, 36–45,
- 1223 https://doi.org/10.1016/j.dsr.2012.03.001, 2012.
- 1224 Truesdale, V. W., Bale, A. J., and Woodward, E. M. S.: The meridional distribution of dissolved iodine in near-
- 1225 surface waters of the Atlantic Ocean, Prog. Oceanogr., 45, 387–400, https://doi.org/10.1016/S0079-6611(00)000091226 4, 2000.
- 1227 Truesdale, V. W., Watts, S. F., and Rendell, A. R.: On the possibility of iodide oxidation in the near-surface of the
- 1228 Black Sea and its implications to iodine in the general ocean, Deep. Res. Part I, 48, 2397–2412, 2001.
- 1229 Truesdale, V. W., Nausch, G., and Waite, T. J.: The effects of the 2001 Barotropic intrusion of bottom-water upon
- 1230 the vertical distribution of inorganic iodine in the Gotland Deep, Cont. Shelf Res., 55, 155–167,
- 1231 https://doi.org/10.1016/j.csr.2013.01.005, 2013.
- 1232 Tsunogai, S.: Iodine in the deep water of the ocean, Deep. Res., 18, 913–919, 1971.
- 1233 Uahengo, C. I., Shi, X., Jiang, G., and Vatuva, A.: Transient shallow-ocean oxidation associated with the late
- 1234 Ediacaran Nama skeletal fauna: Evidence from iodine contents of the Lower Nama Group, southern Namibia,
- 1235 Precambrian Res., 343, 105732, https://doi.org/10.1016/j.precamres.2020.105732, 2020.
- 1236 Wadley, M. R., Stevens, D. P., Jickells, T. D., Hughes, C., Chance, R., Hepach, H., Tinel, L., and Carpenter, L. J.: A
- 1237 Global Model for Iodine Speciation in the Upper Ocean, Global Biogeochem. Cycles, 34, 0–3,
- 1238 https://doi.org/10.1029/2019GB006467, 2020.
- 1239 Watterson, I. G.: Non-dimensional measures of climate model performance, Int. J. Climatol., 16, 379–391,
- 1240 https://doi.org/10.1002/(SICI)1097-0088(199604)16:4<379::AID-JOC18>3.0.CO;2-U, 1996.
- 1241 Wei, B., Tang, D., Shi, X., Lechte, M., Zhou, L., Zhou, X., and Song, H.: A Pulsed Oxygenation in Terminal
- 1242 Paleoproterozoic Ocean: Evidence From the Transition Between the Chuanlinggou and Tuanshanzi Formations,
- 1243 North China, Geochemistry, Geophys. Geosystems, 22, 1–23, https://doi.org/10.1029/2020GC009612, 2021.

- 1244 Wong, G. T. F. and Brewer, P. G.: The marine chemistry of iodine in anoxic basins, Geochim. Cosmochim. Acta,
- 1245 41, 151–159, https://doi.org/10.1016/0016-7037(77)90195-8, 1977.
- 1246 Wong, G. T. F., Takayanagi, K., and Todd, J. F.: Dissolved iodine in waters overlying and in the Orca Basin, Gulf of
- 1247 Mexico, Mar. Chem., 17, 177–183, https://doi.org/10.1016/0304-4203(85)90072-6, 1985.
- 1248 Wong, G. T. F., Piumsomboon, A. U., Dunstan, W. M., Wong, G. T. F., Piumsomboon, A. U., and Dunstan, W. M.:
- 1249 The transformation of iodate to iodide in marine phytoplankton cultures, Mar. Geol. Prog. Ser., 237, 27–39, 2002.
- 1250 Yamazaki, C., Kashiwa, S., Horiuchi, A., and Kasahara, Y.: A novel dimethylsulfoxide reductase family of
- 1251 molybdenum enzyme , Idr , is involved in iodate respiration by Pseudomonas sp . SCT, 22, 2196-2212,
- 1252 https://doi.org/10.1111/1462-2920.14988, 2020.
- 1253 Yu, Y., Chen, Y., Li, D., and Su, J.: A transient oxygen increase in the Mesoproterozoic ocean at ~1.44 Ga:
- 1254 Geochemical evidence from the Tieling Formation, North China Platform, Precambrian Res., 369, 106527,
- 1255 https://doi.org/10.1016/j.precamres.2021.106527, 2022.
- 1256 Zhang, S., Xu, C., Creeley, D., Ho, Y. F., Li, H. P., Grandbois, R., Schwehr, K. A., Kaplan, D. I., Yeager, C. M.,
- 1257 Wellman, D., and Santschi, P. H.: Response to comment on "iodine-129 and iodine-127 speciation in groundwater at
- 1258 hanford site, U.S.: Iodate incorporation into calcite," Environ. Sci. Technol., 47, 13205-13206,
- 1259 https://doi.org/10.1021/es4046132, 2013.
- 1260 Zhou, X., Thomas, E., Rickaby, R. E. M., Winguth, A. M. E., and Lu, Z.: I/Ca evidence for upper ocean
- 1261 deoxygenation during the PETM, Paleoceanography, 29, 964–975, https://doi.org/10.1002/2014PA002702, 2014.
- 1262 Zhou, X., Jenkyns, H. C., Owens, J. D., Junium, C. K., Zheng, X. Y., Sageman, B. B., Hardisty, D. S., Lyons, T. W.,
- 1263 Ridgwell, A., and Lu, Z.: Upper ocean oxygenation dynamics from I/Ca ratios during the Cenomanian-Turonian
- 1264 OAE 2, Paleoceanography, 30, 510–526, https://doi.org/10.1002/2014PA002741, 2015.
- 1265 Žic, V., Carić, M., and Ciglenečki, I.: The impact of natural water column mixing on iodine and nutrient speciation
- 1266 in a eutrophic anchialine pond (Rogoznica Lake, Croatia), Estuar. Coast. Shelf Sci., 133, 260–272,
- 1267 https://doi.org/10.1016/j.ecss.2013.09.008, 2013.
- 1268