

# Leaf wax and Sr-Nd isotope evidence for high-latitude dust input to the central South China sea and its implication for fertilization

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1 **Leaf wax and Sr-Nd isotope evidence for high latitude dust input to the central South China Sea**  
2 **and its implication for fertilization**

3  
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18  
19 **Key Points:**

- 20 1. High concentrations of leaf-wax *n*-alkanes are measured in the South China Sea central basin
- 21 2. *n*-Alkane carbon preference index and hydrogen isotopic composition trace aeolian dust deposition from  
22 higher latitudes to the central basin
- 23 3. Aeolian dust may play a significant role in regulating deposition of terrestrial organic matter in the  
24 central South China Sea.

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27 **Abstract**

28       Recent time-series from sediment traps show abnormally high chlorophyll-*a* concentrations and  
29 primary productivity in the oligotrophic central South China Sea (SCS), especially during wintertime.  
30 Here we present new insights from compound-specific hydrogen isotopic analysis of leaf wax *n*-alkanes  
31 and Sr-Nd isotopes extracted from four basin-wide surface sediment transects. We find that the deepest  
32 surface sediments in the central basin contain the most depleted *n*-alkane hydrogen isotopes, which is  
33 suggestive of inputs from higher latitude soils in northern China. This is further supported by Sr-Nd  
34 isotopes of the same surface sediments. We propose that aeolian dust is transported by the winter  
35 monsoon and fertilizes the phytoplankton bloom in the central SCS. This process may have been  
36 enhanced during glacial periods due to a stronger winter monsoon, driving both vertical mixing and  
37 dust transport to the central basin.

38

39 *Keywords:* *n*-alkane, leaf wax, carbon and hydrogen isotopes, aeolian dust, Sr-Nd isotopes, South China  
40 Sea

41

42

43 **Plain Language Summary:**

44       Recent studies observe abnormally high winter primary productivity and nitrate concentrations  
45 in the surface waters of the central South China Sea. However, this is a nutrient limited region of the  
46 ocean, so the drivers of this primary productivity are unclear. Here we analyze leaf wax carbon and  
47 hydrogen isotopes, and Sr-Nd isotopes, at four shallow to deep water sediment transects to trace the  
48 sources of dust and organic matter in the sediments of the central basin. Our results suggest the central  
49 basin sediments receive significant terrestrial inputs of dust and nutrients from northern Asia via long-  
50 range aeolian transport (during the winter monsoon). These results give new insights to terrestrial-  
51 marine connections and the carbon cycle of the SCS. This process maybe a significant carbon sink in  
52 the present-day and during the past.

53

54 **1. Introduction**

55       The marine biological pump plays an important role in manipulating glacial-interglacial  
56 atmospheric CO<sub>2</sub>, with the majority of carbon uptake apparently occurring in high latitude oceans such  
57 as the Southern Ocean and subarctic Pacific (Brunelle et al., 2010; Martínez-García et al., 2014).  
58 Recently, Buchanan et al. (2019) used a global marine biogeochemical model to show that the low  
59 latitude ocean could be as important as high latitude locations for regulating atmospheric CO<sub>2</sub> during  
60 glacial periods, due to Fe-induced stimulation of dinitrogen (N<sub>2</sub>) fixation, strengthening the biological  
61 pump, and ultimately causing CO<sub>2</sub> drawdown during glacial periods. This modelling work is consistent  
62 with observations from ocean sediments cores that large quantities of sediments enriched in organic  
63 carbon are preserved in continental seas and basins along continental margins at low latitudes (Berner,  
64 1982; Hedges and Keil, 1995), suggesting an important role in the global carbon cycle (e.g., Dai et al.,  
65 2013; Liu et al., 2010).

66       The South China Sea (SCS) is the largest marginal sea of the Pacific Ocean. It receives more than  
67 700 million metric tons of fluvial sediments annually from surrounding rivers (Liu and Stattegger, 2014).  
68 However, recent evidence highlights anomalously high phytoplankton distributions in the central SCS  
69 where fluvial input is usually considered to be insignificant (Ma et al., 2013), indicating the central SCS  
70 could be a significant carbon sink (Hung et al., 2020). In particular, satellite observations show  
71 phytoplankton blooms, indicated by anomalously high concentrations of Chlorophyll-a, in the central

72 SCS during winter seasons (Ma et al., 2013; Ma et al., 2019). Export production from phytoplankton  
73 blooms contributes a major source of organic matter (OM) to sediments. Sinking particle fluxes  
74 including particulate organic matter (POM), calcium carbonate (CaCO<sub>3</sub>), opal, and lithogenic matter  
75 from long-term sediment traps show increased transportation and accumulation of biogenic materials  
76 during the winter, within the central basin, compared with northern shallower traps (Li et al., 2017; Ma  
77 et al., 2019; Priyadarshani et al., 2019; Zhang et al., 2019). Several 'bottom-up' driving mechanisms of  
78 this phenomenon have been proposed, including the intrusion of Kuroshio surface waters (Hung et al.,  
79 2007) and mesoscale eddies (Li et al., 2017) during the winter. However, the upwelling events induced  
80 by Kuroshio intrusions and meso-scale eddies only account for about 20% of total deposition observed  
81 in sediment traps between July 2012 and April 2013 (Zhang et al., 2019). Consequently, modern  
82 observations of enhanced winter primary productivity in the central SCS requires an additional  
83 mechanism for nutrient delivery.

84 One such potential driving mechanism is the 'top-down' delivery of dust, loaded with nutrients,  
85 from higher latitudes by the East Asian Winter Monsoon (EAWM), which would stimulate nitrogen  
86 fixation and the biological pump. Tracers for aeolian dust in marine sediments include minerals (Blank  
87 et al., 1985; Liu et al., 2015), chemical components (Mcgee et al., 2016; Uematsu et al., 1983) and  
88 terrestrial organics (Bendle et al., 2007; Bendle et al., 2006; Boreddy et al., 2017). Clay minerals and  
89 grain sizes as well as chemical components have been discussed as possible input tracers for the SCS  
90 (Boulay et al., 2007; Liu et al., 2016), but the mixed signal of the fluvial input, deep water current  
91 transportation through Luzon strait, and/or the aeolian dust has made it difficult to confidently appoint  
92 the sources of sediments to the central SCS.

93 The molecular and isotopic composition of leaf wax *n*-alkanes in aerosols has been widely used to  
94 evaluate sources and pathways of airborne dust (Bendle et al., 2007; Ohkouchi et al., 1997; Schefuß et  
95 al., 2003). For example, the carbon isotopic composition of plant wax *n*-alkanes has been used to map  
96 the distribution of C<sub>3</sub>/C<sub>4</sub> plants in source regions, and to decipher variations in vegetation in the  
97 sediment record throughout the Quaternary (Jia et al., 2012; Li et al., 2015). Moreover, the δ<sup>2</sup>H values  
98 of modern leaf wax *n*-alkanes are well correlated with the δ<sup>2</sup>H of meteoric water at latitudinal scales  
99 (Rao et al., 2009) and are increasingly used in palaeohydrological reconstructions (Huang et al., 2018;  
100 Thomas et al., 2014). The δ<sup>2</sup>H values and carbon preference indices (CPIs; ratio of odd-to-even chained  
101 *n*-alkanes) of plant wax *n*-alkanes in East China surface soils exhibit a strong dependence on latitude

102 and the meteoric  $\delta^2\text{H}$  line (Rao et al., 2009). The SCS region is strongly influenced by the East Asian  
103 Monsoon, but whether the  $\delta^2\text{H}$  values of *n*-alkanes from surface sediments can be used to constrain  
104 sediment sources in the deep basin, which sits beyond the reach of most fluvial inputs, is still unknown.

105 We argue that the SCS represents a valuable research opportunity: local fluvial inputs are largely  
106 captured on the continental shelf, whilst the middle of central basin (>4000m) sits beyond the reach of  
107 most fluvial inputs. Thus, the deep SCS may capture a broad regional signal of aeolian dust inputs  
108 (relatively uncontaminated by fluvial or biogenic factors) from a position proximal to the Asian  
109 continent. We conduct a basin-wide survey of leaf wax molecular and isotopic distributions and  
110 radiogenic Sr-Nd isotopes in the surface sediments of the SCS, and contrast this data with observations  
111 and sedimentary of plankton groups in the SCS. This synthesis leads us to infer an increasing biological  
112 pump for the anomalous wintertime phytoplankton bloom in the SCS on glacial-interglacial timescales.

113

## 114 2. Samples and Methods

### 115 2.1 Sampling sites

116 A total of 62 surface sediment samples were collected from the SCS (Fig. 1), with water depths  
117 ranging from 30 to 4405m. The sampling transect therefore allows a comparison of the preservation of  
118 terrestrial organic matter between shallow and deep-water sediments. The samples were collected using  
119 a deep-sea sediment grab sampler or a box corer (0-5cm) and stored at -20°C prior to analysis.

### 120 2.2 TOC and Sr-Nd isotopic compositions analysis

121 For the TOC analysis, 1 gram sediment was decalcified with 2M HCl at room temperature for 24h,  
122 then rinsed with pure water until pH = 7. Samples were freeze dried and transferred into tin capsules,  
123 then analyzed by Elemental Analyzer. For the Sr-Nd isotopes analysis, 20 samples were digested in  
124 Teflon bombs with mixed agents of double distilled  $\text{HNO}_3$  and HF acid at 190 °C for 48h. Then, samples  
125 were detected using a Triton T1 thermal ionization mass spectrometer (TIMS) and a Neptune Plus multi-  
126 collector ICP-MS. The measured  $^{143}\text{N}/^{144}\text{Nd}$  and  $^{87}\text{Sr}/^{86}\text{Sr}$  ratios were normalized to  $^{146}\text{Nd}/^{144}\text{Nd} =$   
127  $0.7219$  and  $^{86}\text{Sr}/^{88}\text{Sr} = 0.1194$ , respectively. During the analysis, BCR-2 standard gave  $^{87}\text{Sr}/^{86}\text{Sr} =$   
128  $0.704989 \pm 8$  ( $2\sigma$ ) and  $^{143}\text{N}/^{144}\text{Nd} = 0.512644 \pm 2$  ( $2\sigma$ ). Nd results are calculated as  $\epsilon\text{Nd}(0) =$   
129  $[(^{143}\text{N}/^{144}\text{Nd})/0.512638] - 1 \times 10000$ , using the chondritic uniform reservoir value given by Jacobsen  
130 and Wasserburg (1980).

131 2.3 Lipid extraction and analysis.

132 Samples were freeze dried and homogenized with a pestle and mortar, then subjected to a  
133 methodology modified from Yang et al. (2014). Samples were ultrasonically extracted with an azeotrope  
134 of dichloromethane: MeOH (v/v 9:1) 5 times. All extracts were combined and collected after  
135 centrifugation. The combined extracts were concentrated to 1-2 mL using rotary evaporation and dried  
136 under a flow of N<sub>2</sub> gas. The total lipid extract was fractionated with *n*-hexane and MeOH into an apolar  
137 fraction (containing the *n*-alkanes) and a polar fraction.

138 *n*-Alkanes were detected and identified by an Agilent 7890 gas chromatograph and 5975A mass  
139 spectrometer (GC-MS) equipped with a DB-5MS capillary column (60m × 0.25mm × 0.25μm). The  
140 alkane fraction was injected at a programmed temperature ramp of 3 °C/min<sup>-1</sup> from 70 to 300 °C and  
141 held at 300 °C for 30 min. Relative compound abundances were calculated by comparing corresponding  
142 MS (TIC) peak areas with internal standards of known concentration. Stable carbon and hydrogen  
143 isotope compositions of individual *n*-alkanes were determined followed the program of Huang et al.  
144 (2014) using a Finnigan Trace GC instrument attached to a Finnigan Delta Plus XP isotope ratio mass  
145 spectrometer. Duplicate analyses were used to confirm that the standard deviations of leaf wax carbon  
146 and hydrogen isotope determinations were better than ± 0.5‰ and 5‰ respectively. The δ<sup>13</sup>C and δ<sup>2</sup>H  
147 values are reported in the delta notation (‰) relative to Vienna Pee Dee Belemnite (VPDB) and Vienna  
148 Standard Mean Ocean Water (VSMOW), respectively.

149

150 **3. Results**

151 3.1 TOC and Sr-Nd isotope compositions

152 TOC values range from 0.12% to 0.82% (average 0.43%) in the SCS, and C/N values vary from  
153 1.8 to 8.7 (average 6.3). Both TOC and C/N decrease with distance offshore, with high values in the  
154 northern SCS, representing the higher organic matter deposits from fluvial inputs. However, in the  
155 central basin, the TOC and C/N show an increase, with values comparable to those of the northern SCS  
156 (Fig. 2A), suggesting a distinct source of terrestrial organic matter.

157 The <sup>87</sup>Sr/<sup>86</sup>Sr and <sup>143</sup>Nd/<sup>144</sup>Nd ratios range from 0.709161 to 0.712215 (average 0.712125) and from  
158 0.512058 to 0.512425 (average 0.512154), respectively. εNd(0) values range from -11.31 to -4.15  
159 (average -9.45). These values are consistent with previously published data of surface sediments from

160 the SCS (Liu et al., 2015, and references therein). Three samples from the northern and eastern SCS  
161 with the highest  $\epsilon\text{Nd}(0)$  (-6 to -4) derived from Luzon island (Liu et al., 2015).

### 162 3.2 Leaf wax molecular and isotopes distributions

163 The surface sediments contain *n*-alkanes characterized by a high odd-over-even carbon number  
164 ranging in carbon number from  $\text{C}_{16}$  to  $\text{C}_{35}$ , with  $\text{C}_{31}$  dominant (Fig. S1). This is a clear signature of  
165 terrestrial higher plant origin (Bush and McInerney, 2013; Eglinton and Hamilton, 1967). The  
166 concentrations of total long chain *n*-alkanes show large variations, with anomalously high  
167 concentrations in the deep central basin (Fig. 2B). CPI is typically around 2 in shallow surface sediments,  
168 but it increases abruptly to 8 in the deepest basin (Fig. 2C). Concentrations of long chain diols (LCDs)  
169 (algal biomarkers) range from 0.1 to 32.7 mg/g TOC dry sediments (Fig. 2D; Yang et al., 2020). It is  
170 notable that both *n*-alkanes and planktonic LCDs comprise an enhanced proportion of the total organic  
171 carbon (TOC) in the surface sediments deposited in the central basin, with TOC-normalized  
172 concentrations up to three orders of magnitude higher in deep sediments than in shallow sediments (Fig.  
173 3A-B).

174 Our *n*-alkane  $\delta^{13}\text{C}$  values shift from -29 to -31‰ from the northeastern coast to the central basin  
175 (Fig. 2E-F). The values in the deep central basin are similar to the northern SCS, and are highly variable  
176 (Fig. 3E). *n*-Alkane  $\delta^2\text{H}$  values vary from -140 to -160‰ in the northern SCS (Fig. 2G-H), but are  
177 depleted in the deep central basin, with minimum values around -200‰. The lowest values (-190 to -  
178 200 ‰) are distinct from the values observed in both Southern China catchment soils and the shallow  
179 sediments (Fig. 1, Pelejero et al., 2003; Rao et al., 2009).

180

## 181 4. Discussion

### 182 4.1 Sources of surface sediments in the central SCS

183 The Sr-Nd isotopes from SCS surface sediments and surrounding fluvial drainage systems have  
184 been well studied and were used to identify the sediment source provinces (Liu et al., 2016 and  
185 references therein). Three samples from the northern and eastern SCS have a more positive  $\epsilon\text{Nd}(0)$   
186 falling within the variation range of the Luzon island (Liu et al., 2016), indicating the possible influence  
187 of northern Luzon Arc material. The surrounding fluvial inputs have higher values of  $^{87}\text{Sr}/^{86}\text{Sr}$  (>7.2,  
188 Liu et al., 2016), which are distinguishable from our deep-water sediments, except for two samples  
189 from the Red River. However, the clay mineral assemblages showed limited influence of the Red River

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190 on the SCS central basin (Liu et al., 2016). Our data shows that the more positive  $\epsilon\text{Nd}(0)$  and lower  
191  $^{87}\text{Sr}/^{86}\text{Sr}$  isotopes from the central basin sediments are not consistent with, and thus may not originate  
192 from the surrounding fluvial drainage systems.

193 Instead, the  $^{87}\text{Sr}/^{86}\text{Sr}$  and  $\epsilon\text{Nd}(0)$  isotopes from the central basin are consistent with Asian dust  
194 regions, suggesting a source relationship (Fig. 4A). This result is in line with the geochemical analyses  
195 of sediments from Yongxing island in the SCS, which received significant high-latitude Asian aeolian  
196 dust inputs (Liu et al., 2014). Grain size of the sediments might exert an impact on the Sr-Nd isotopes  
197 as the Sr isotopic compositions are known to be strongly affected by the weathering, transportation and  
198 deposition (Chen et al., 2007). The grain size of the sediments ranges from 0-400  $\mu\text{m}$  (average medium  
199 diameter  $>5 \mu\text{m}$ ) in the northern and eastern SCS, but from 0-40  $\mu\text{m}$  (average medium diameter  $\sim 3\mu\text{m}$ )  
200 in the central SCS basin. Comparison of the grain size fractions suggests the Asian dust ( $<5 \mu\text{m}$ ) falling  
201 in the range of the SCS central basin is mainly contributed by the isotopic region B (i.e., the northern  
202 margin of Tibetan Plateau) and C (i.e., the Ordos Plateau) (Fig. 4B). A calculation based on a simple  
203 mass balance suggests 1:4 for the relative contribution of the isotopic regions B and C to the SCS central  
204 basin (Fig. 4B).

205 *n*-Alkane CPIs of  $\sim 2$  are consistent with those previously observed in shallow sediments from the  
206 northern SCS (about 1.4 to 2.9 from Xu et al., 2014), and somewhat lower than those from surrounding  
207 surface soils in Southern China (ca. 4, Rao et al., 2009; Luo et al., 2012), as well as northern SCS  
208 Holocene sediments (ca. 2.3, Pelejero, 2003). Lower CPIs in marine sediments than in source catchment  
209 soils suggest microbial degradation during riverine transportation (Ganeshram et al., 2011; Sun et al.,  
210 2005). However, in the deepest SCS sediments, CPIs  $>3$  are observed (Fig. 3C), with the highest CPI  
211 measured approaching 8. Such high values are not observed in any of the shallow water settings  
212 (Pelejero, 2003; Xu et al., 2014), suggesting that the terrestrial organic matter observed in this deep  
213 basin setting is unlikely to be of riverine origin. CPIs of surface soils in eastern and northern China  
214 show a strong relationship with latitude, with elevated values ( $> 5$ ) observed in higher latitudes (Rao et  
215 al., 2009). This high latitude CPI signature transported from the Asian dust area is also observed in the  
216 deep-sea surface sediments collected from the Central Pacific (Ohkouchi et al., 1997). Comparison of  
217 CPI values from SCS surface sediments with latitudinal soil profiles (Fig. S2), highlights that the high  
218 CPI values (ca. 8) in the deep basin match source locations at mid-high latitudes (about  $40^\circ\text{N}$ ) (Fig.  
219 4C), where the loess plateau (with average CPI at 12.3; Liu and Huang, 2005; Luo et al., 2012) and

220 Gobi desert are located. Although *n*-alkane CPI covaries with various different environmental factors  
221 (latitude, aridity and vegetation types) globally (Luo et al., 2012), latitude is still significant and  
222 correlates broadly with CPI values (Fig. S2), with higher CPI in high latitudes. We propose the  
223 distinctive CPI signature of the deepest SCS records are related to enhanced terrestrial organic matter  
224 contributions via aeolian dust transported directly from vegetation or soils (Chikaraishi and Naraoka,  
225 2003; Ning et al., 2005).

226 *n*-Alkane  $\delta^2\text{H}$  values in East China surface soils also exhibit a strong dependence on latitude, with  
227 lower values occurring at higher latitudes (Fig. 4D) (Rao et al., 2009). The  $\delta^2\text{H}$  of southern Chinese  
228 surface soils are around -160‰, consistent with northern SCS surface sediment values. However, the  
229  $\delta^2\text{H}$  values in the central basin are more depleted and closer to higher latitude soil values (according to  
230 the linear correlation between  $\delta^2\text{H}$  and latitude (Fig. 4D)), supporting the above suggestion that *n*-  
231 alkanes deposited in the central SCS are sourced from high-latitude dust inputs. The trend with water  
232 depth for *n*-alkane  $\delta^{13}\text{C}$  values in the SCS is more variable than for  $\delta^2\text{H}$ . *n*-Alkane  $\delta^{13}\text{C}$  is used to trace  
233 the relative contribution of  $\text{C}_3$  and  $\text{C}_4$  plant types. The *n*-alkane  $\delta^{13}\text{C}$  through the SCS thus likely records  
234 the  $\text{C}_3/\text{C}_4$  plant signal in terrestrial inputs to the SCS. The *n*-alkane  $\delta^{13}\text{C}$  in the SCS surface sediments  
235 is consistent with inputs from a diverse and mixed distribution of  $\text{C}_3$  and  $\text{C}_4$  plants extending from  
236 Southeast Asia throughout mainland China (Still et al., 2009). Thus in this context  $\delta^{13}\text{C}$  values are not  
237 as diagnostic as the corresponding  $\delta^2\text{H}$  values and CPIs for indicative of source regions.

238 Comparison of *n*-alkane  $\delta^2\text{H}$  and CPI data from the central basin with the available soil *n*-alkane  
239 data (this study and Rao et al., 2009, Fig. 4C-D), suggests significant soil inputs from  $>40^\circ\text{N}$ , ca.  
240 2000km to the north of the SCS region. High latitude arid and semi-arid regions in China and Asia,  
241 especially the Taklimakan desert (located at ca.  $35$  to  $45^\circ\text{N}$ ), are major sources of atmospheric dust in  
242 the Northern Hemisphere (add ref). During the winter monsoon, decreased winter precipitation allows  
243 more aeolian dust transport to the SCS, while only minimal inputs of entrained dust occurs during the  
244 summer monsoon, due to heavy summer precipitation and rainout of dust closer to source regions  
245 (Boulay et al., 2003; Tian et al., 2005). We conducted a seasonal back trajectory air masses model which  
246 simulated the seasonal organic matter transport pathway to the central SCS ( $13^\circ\text{N}$ ,  $115^\circ\text{E}$ ). The result  
247 shows a larger amount of dust transport from central Asia during the winter season compared to the  
248 summer season (Fig. S3), which is consistent with the results from Yongxin island (Xiao et al., 2017).  
249 Our *n*-alkane isotopic signatures (low  $\delta^2\text{H}$  values) are consistent with enhanced delivery of aeolian dust

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250 to the central SCS by the winter monsoon. The lipid profile results agree with the  $^{87}\text{Sr}/^{86}\text{Sr}$  and  $\epsilon\text{Nd}(0)$   
251 evidence, both confirm dust inputs from high latitude Asian source regions to the deep central basin of  
252 the SCS.

253

#### 254 4.2 Enhanced Aeolian dust input to the SCS central basin and its implications

255 Our results show both terrestrial leaf wax *n*-alkane and LCD concentrations increase in the central  
256 basin compared to northern SCS. Trace metal (Sr-Nd) isotopes from the same stations also constrain  
257 the original sources to predominantly mid-latitude Asian deserts (Fig. 4A-B). It is interesting to note  
258 that time-series sediment trap data from the SCS quantified the effects of the different processes on  
259 sinking particle fluxes, and highlighted that the northeast winter monsoon and associated aerosol  
260 deposition events played key roles in sediment deposition during the winter monsoon period (58.7%,  
261 Zhang et al., 2019, from October to April). Consequently, our investigations on surface sediments - and  
262 the reported time-series trap data - both indicate an important contribution of Asian dust to the SCS  
263 central basin.

264 Of significance is that Asian dust inputs are elevated during the winter season due to the occurrence  
265 of the enhanced Winter Asian Winter Monsoon. This is of importance for connections to the carbon-  
266 cycle, via processes such as fertilization of phytoplankton and the rapid deposition of organic matter.  
267 Both *n*-alkane and LCD concentrations are elevated in the surface sediments of the central basin  
268 compared with the northern SCS, and high *n*-alkane CPI values indicate relatively fresh organic matter  
269 deposition to the deepest water sites in the central SCS. These results are consistent with enhanced  
270 satellite Chl-*a* concentration and organic matter collected from time-series sediment traps in the central  
271 basin (Li et al., 2017; Ma et al., 2019; Priyadarshani et al., 2019; Zhang et al., 2019). Seasonal time-  
272 series data analysis shows higher OM deposition during the winter season when EAWM conditions  
273 prevail.

274 A “bottom-up” mechanism driving phytoplankton blooms and biomass in the marginal, stratified  
275 regions of the SCS has been well characterised (Chen, 2005; Tang et al., 1999; Li et al., 2017). During  
276 the winter season, the frequency of cold meso-scale eddies increases in the central SCS, which drives  
277 nutrient-rich subsurface waters to the surface and stimulates phytoplankton blooms in the oligotrophic  
278 SCS central basin (Chen, 2005; Tang et al., 1999). Coupled with the meso-scale eddies during winter  
279 season, vertical mixing in the upper water column is strongest in the central basin, and is about four

280 times deeper in winter than is seen in the rest of the year (Lu et al., 2020; Qu, 2001). Increased vertical  
281 mixing drives higher concentrations of nitrate to the surface layer and fertilizes the phytoplankton  
282 bloom in the central basin. This mechanism explains the higher nitrate concentration and primary  
283 production in the central basin compared to the northern SCS observed in satellite data. Logically,  
284 increased carbon export to the SCS during the winter monsoon will lead to the deposition of an outsized  
285 proportion of these sediments during that time interval. This bottom-up mechanism successfully  
286 explains the abnormal high chl-a and high OC fluxes at the central basin, but cannot reconcile the high  
287 terrestrial sourced *n*-alkane distribution in the central SCS.

288 High concentrations of high CPI *n*-alkanes which are depleted in  $\delta^2\text{H}$  demonstrate the importance  
289 of aeolian dust deposition in the transport of organic matter to the central SCS (Fig. 4C-D). We propose  
290 a “top-down” mechanism could be essential to explain the observed distributions of *n*-alkane and long  
291 chain diols in the surface sediments of the central SCS basin. Windborne dust particles containing both  
292 lithogenic material and land-derived lipids are significant in the rapid transfer of newly fixed organic  
293 carbon from the sea surface to the bottom (Ittekkot et al., 1992). The incorporation of minerals into  
294 biologically formed aggregates ensures the rapid deposition of fresh *n*-alkanes with higher CPI in the  
295 central basin. Meanwhile, modern observation studies suggest Asian dust events could enhance  
296 phytoplankton growth and primary production in Chinese marginal seas (Tan et al., 2011; Tan et al.,  
297 2012; Wang et al., 2012), which further supports evidence from TOC, carbonate, microfossil and lipid  
298 profiles studies (Thunell et al., 1992; Huang et al., 1997a; Huang et al., 1997b; Shiao et al., 2008; Ren  
299 et al., 2017). Long time *in situ* studies revealed aerosol deposition of dissolved inorganic nitrogen to  
300 the SCS, especially in the basin area, was approximately 20% on average (Kim et al., 2014; Shen et al.,  
301 2020; Gao et al., 2020). This atmospheric N deposition could support the primary production in the  
302 oligotrophic water of the SCS, which is characterized by limited nitrate.

303 Dust regulated iron supply might stimulate the nitrogen fixation, and is crucial in linking the  
304 biological cycling of iron to the assimilation of major nutrients and carbon fixation (Tagliabue et al.,  
305 2017). A recent model study shows Fe-induced stimulation of  $\text{N}_2$  fixation pathways could drive a  
306 considerable uptake of carbon dioxide in low latitude oceans during dusty glacial conditions (Buchanan  
307 et al., 2019). Gaye et al. (2009) proposed the  $\text{N}_2$  fixation contributed up to 20% to settling particle  
308 nitrogen in the deep SCS, about twice the estimated contribution in the northern SCS (Kao et al., 2012;  
309 Wong et al., 2007; Zhang et al., 2015). However, both foraminifera-bound nitrogen isotope records and

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310 ammonia oxidizing archaea records from the SCS (Ren et al., 2017; Dong et al., 2019) reconstruct lower  
311 N<sub>2</sub> fixation rates during glacial periods, despite the observed increase in dust deposition and productivity.  
312 Thus the role of Fe fertilization in regulating the SCS carbon and nitrogen cycles requires further  
313 investigation.

314 In summary our evidence from terrestrial leaf wax (*n*-alkane) and marine phytoplankton (LCDs)  
315 biomarkers is consistent with sediment trap time-series data (ref) and suggests a role for aerosol dust  
316 deposition in the winter phytoplankton blooms as observed by satellite in the central basin of the SCS  
317 (Ma et al., 2013). During the winter monsoon it appears that dust supplied from higher latitudes and  
318 vertical oceanic mixing supplies higher nutrients and triggers new production in the central SCS. Our  
319 results are also important for paleoclimate reconstructions, as this process would be expected to deliver  
320 enhanced supplies of terrestrial, nutrient bearing, dust during the glacial periods (Shiau et al., 2008; Ren  
321 et al., 2017). However, precise mechanisms and the relative importance of, for example, Fe fertilization  
322 in currently ambiguous. Thus more work, including *in-situ* monitoring in the central basin and model  
323 simulations are required to elucidate mechanisms, quantify fluxes and understand the importance of  
324 dust deposition at the air-water interface in low latitude marginal seas.

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## 326 5. Conclusions

327 Multiple lines of evidence based on high *n*-alkane CPI, depleted δ<sup>2</sup>H, and Sr-Nd isotope values  
328 in surface sediments highlight the transport and deposition of dust from high latitude regions to the  
329 central SCS basin. We conclude aeolian dust from northern China is transported by the East Asia Winter  
330 Monsoon and contributes (along with vertical mixing of nutrients) to the triggering of the winter  
331 phytoplankton in the central SCS. The intimate biotic-abiotic association triggered by dust supply could  
332 accelerate the organic deposition rate and thus has implications for the biological pump. Our results  
333 shows that nitrogen fixation in the marginal SCS central basin could be as important as vertical mixing,  
334 both in increasing primary production and for high sediment deposition rates. Thus the SCS central  
335 basin could be a significant, but hitherto overlooked, carbon sink during the present day and glacial  
336 periods, which merits further investigation.

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346 to the University of Birmingham

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348 **Data Availability Statement**

349 All of the original data has been uploaded as supplemental material and will be deposited publicly to  
350 the repository of Zenodo, once it is accepted for publishing. All the supporting data can be found in the cited  
351 references (Rao et al., 2009; Luo et al., 2012; Liu et al., 2007; Pettke et al., 2000; Defant et al., 1990; Jiang  
352 et al., 2013; Chen et al., 2007; Liu et al., 2014; Biscaye et al., 1997).

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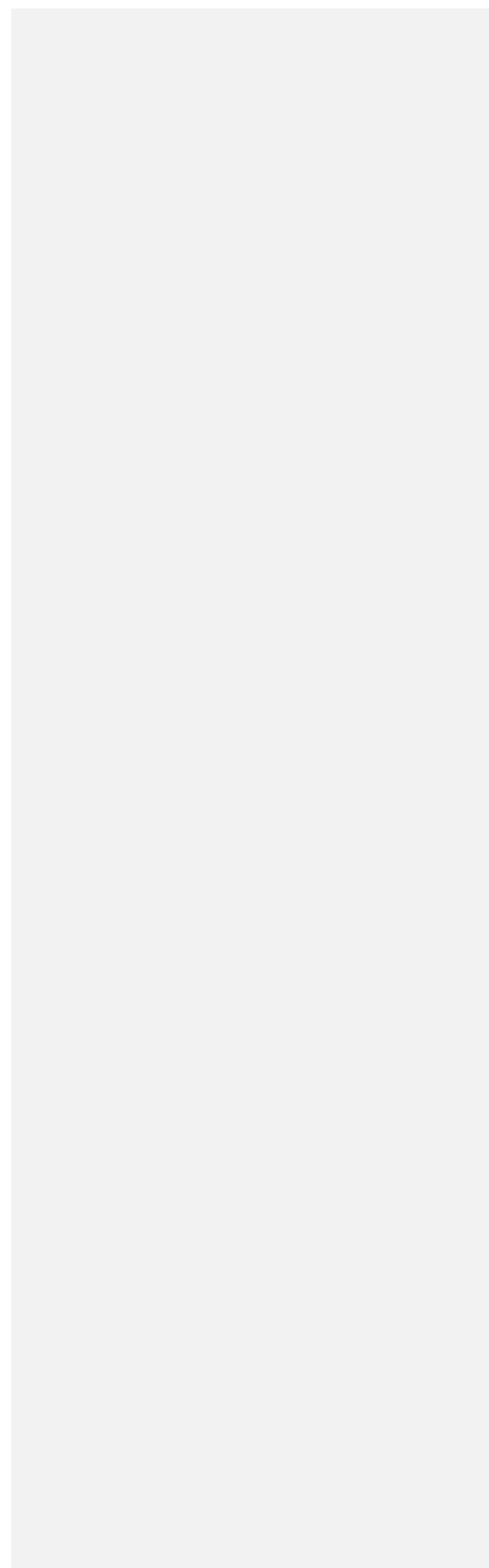
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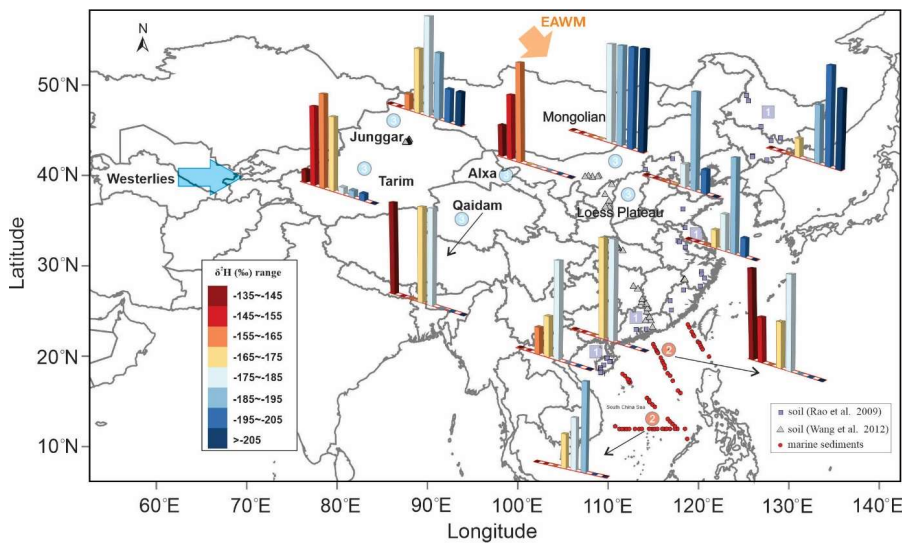


549 **Figure Captions**

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551 Fig. 1 Distribution of samples used in this study: a) new surface sediment analyses from the South China  
552 Sea (red dots); b) surface soil samples previously reported by Rao et al. (2009) and Luo et al. (2012) in  
553 the Chinese mainland. The frequency histograms show the geographical distribution of the hydrogen  
554 isotopic composition of the C<sub>31</sub> n-alkane in : 1) northeastern China (purple squares, Rao et al., 2009);  
555 2) the shallow and central SCS (red circles, this study) and; 3) the Loess Plateau and Gobi deserts (blue  
556 circles, this study). The sequential color gradient (from red to blue) represents heavy to light hydrogen  
557 isotope values.

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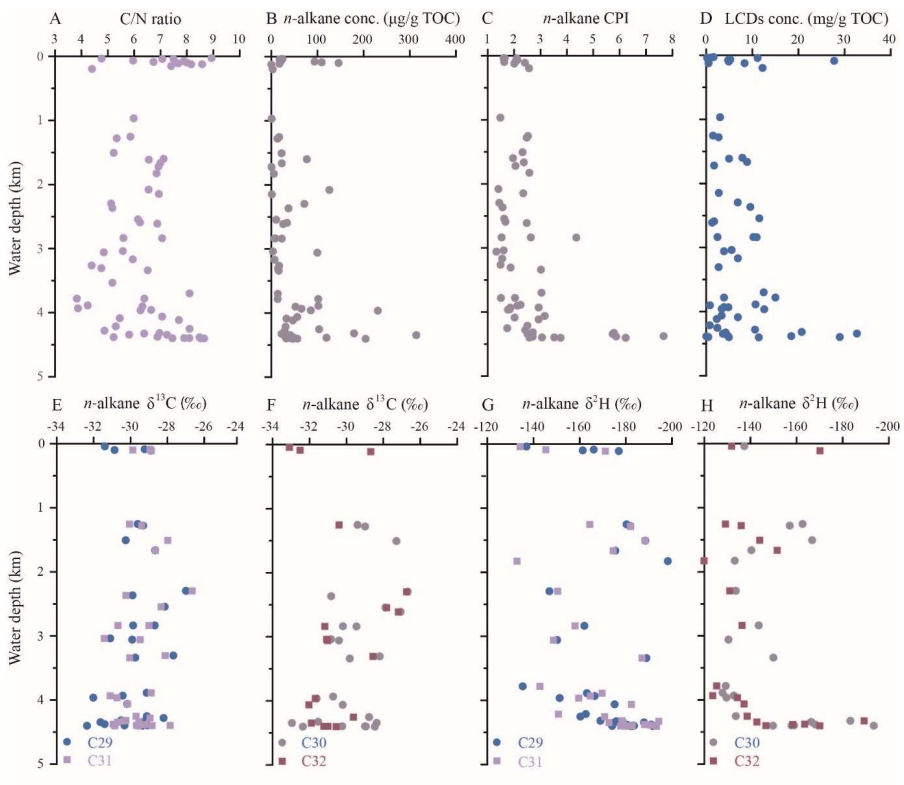
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576 Fig. 2. Water depth distribution of molecular and elemental parameters measured in SCS surface  
 577 sediments. Marked changes are notable at water depths below 4000 m. a) C/N ratio, b-c) Concentrations  
 578 of *n*-alkanes and CPI values, d) Concentrations of long chain diols, e-h) Odd ( $C_{29}$  and  $C_{31}$ ) and even  
 579 ( $C_{30}$  and  $C_{32}$ ) chains hydrogen and carbon alkane isotopes.  
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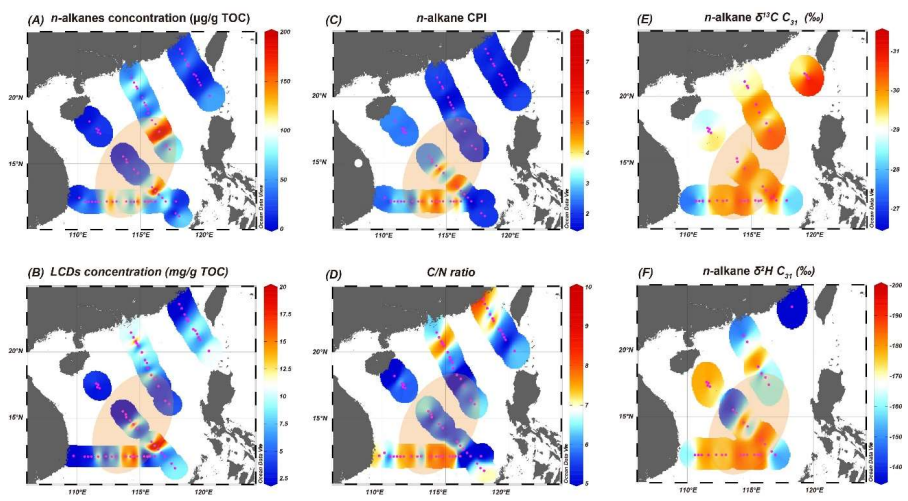
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Fig. 3 Spatial distribution of molecular and elemental parameters from the northern to the central basin of the SCS. Distinct values (e.g. high *n*-alkane concentrations, high CPI, isotopically light) are highlighted in the central basin. a-b) Concentrations of *n*-alkane and long chain diols, c) *n*-alkane CPI value, d) C/N ratio, e-f) carbon and hydrogen isotopes of C<sub>31</sub> *n*-alkane. The shaded circle roughly show the deep basin with water depth deeper than 2400m.

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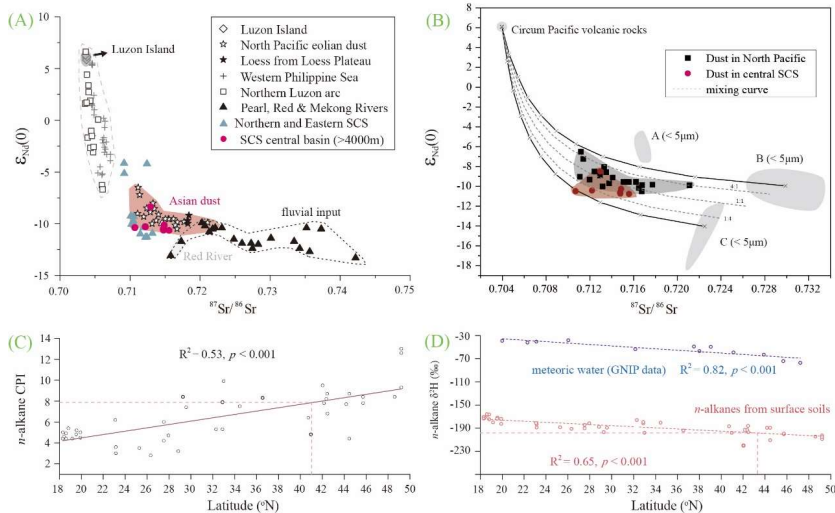
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625 Fig. 4 Latitudinal gradients for Sr-Nd isotopes, CPI values and hydrogen isotopes of *n*-alkanes. These  
626 parameters are used to infer the source of dust which has been long-range transported to the deep, central  
627 basin of the SCS. A) Correlation between  $\epsilon_{Nd}$  and  $^{87}Sr/^{86}Sr$  of SCS surface sediments, modified from  
628 Liu et al. (2014). Shaded areas show potential sources for material in the central basin. Data of the Pearl,  
629 Red and Mekong rivers from Liu et al. (2007); northern Pacific ocean dust samples are from Pettke et  
630 al. (2000). Luzon Arc and Luzon Island samples are from Defant et al. (1990), Philippine Sea samples  
631 are from Jiang et al. (2013), Loess samples are from Chen et al. (2007). B) Sr-Nd isotopic compositions  
632 of ancient dust falls in North Pacific, and SCS central basin, together with the mixing curve between  
633 the  $<5 \mu m$  silicate fractions of the Chinese deserts (Chen et al., 2007) and the volcanic end-member  
634 used in Biscaye et al. (1997). The grey dashed lines means the mixing fraction based on end member B  
635 and C C), the variation of the CPI values of the *n*-alkanes of the surface soils from different latitude  
636 East China reported by Rao et al. (2009). The dashed line in red shows the maximum of CPI values  
637 identified in marine sediments of the deepest water in SCS, with the interception with the regression  
638 line (red line) showing the potential dust source; D), the variation of the deuterium content in meteoric  
639 water (the original data are available at the website of Global Network of Isotopes in Precipitation  
640 (GNIP), <http://isohis.iaea.org>) and in *n*-alkanes of surface soils in East China reported by Rao et al.  
641 (2009). The lowermost perpendicular red dashed lines indicate the most depleted  $\delta^2H$  values of the *n*-  
642 alkanes identified in marine sediments of the deepest waters in the SCS. The interception with the  
643 regression line (dashed line in blue) shows the general latitude of the potential dust source.

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