2	Contrasting composition of terrigenous organic matter in
3	the dissolved, particulate and sedimentary organic carbon
4	pools on the outer East Siberian Arctic Shelf
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22 Abstract

23 Fluvial discharge and coastal erosion of the permafrost-dominated East Siberian Arctic delivers large quantities of terrigenous organic carbon (Terr-OC) to marine waters. 24 25 The composition and fate of the remobilized Terr-OC needs to be better constrained as it impacts the potential for a climate-carbon feedback. In the present study, the bulk isotope 26 $(\delta^{13}C \text{ and } \Delta^{14}C)$ and macromolecular (lignin-derived phenols) composition of the cross-shelf 27 exported organic carbon (OC) in different marine pools is evaluated. For this purpose, as part 28 of the SWERUS-C3 expedition (July-September 2014), sediment organic carbon (SOC) as 29 30 well as water column (from surface and near-bottom seawater) dissolved organic carbon 31 (DOC) and particulate organic carbon (POC) samples were collected along the outer shelves 32 of the Kara Sea, Laptev Sea and East Siberian Sea. The results show that the Lena River and the DOC may have a preferential role in the transport of Terr-OC to the outer shelf. DOC 33 concentrations (740-3600 µg/L) were one order of magnitude higher than POC (20-360 34 μ g/L), with higher concentrations towards to the Lena River plume. The δ^{13} C signatures in 35 the three carbon pools varied from -23.9±1.9‰ in the SOC, -26.1±1.2‰ in the DOC and -36 27.1±1.9‰ in the POC. The Δ^{14} C values ranged between -395±83‰ (SOC), -226±92‰ 37 (DOC) and -113±122‰ (POC). These stable and radiocarbon isotopes were also different 38 39 between the Laptev Sea and the East Siberian Sea. Both DOC and POC showed a depleted 40 and younger trend off the Lena River plume. Further, the Pacific inflow and the sea ice coverage, which works as a barrier preventing the input of "young" DOC and POC, seem to 41 have a strong influence in these carbon pools, presenting older and more enriched $\delta^{13}C$ 42 signatures under the sea ice extent. Lignin phenols exhibited higher OC-normalized 43 44 concentration in the SOC (0.10-2.34 mg/g OC) and DOC (0.08-2.40 mg/g OC) than in the POC (0.03-1.14 mg/g OC). The good relationship between lignin and Δ^{14} C signatures in the 45 DOC suggests that a significant fraction of the outer-shelf DOC comes from "young" Terr-46 OC. By contrast, the slightly negative correlation between lignin phenols and Δ^{14} C signatures 47 in POC, with higher lignin concentrations in older POC from near-bottom waters, may reflect 48 the off-shelf transport of OC from remobilized permafrost in the nepheloid layer. 49 Syringyl/vanillyl and cinnamyl/vannillyl phenols ratios presented distinct clustering between 50 DOC, POC and SOC, implying that those pools may be carrying different Terr-OC of 51 partially different origin. Moreover, 3,5-dihydroxybenzoic acid to vanillyl phenols ratios and 52 53 p-coumaric acid to ferulic acid ratios, used as a diagenetic indicators, enhanced in POC and SOC, suggesting more degradation within these pools. Overall, the key contrast between 54

- 55 enhanced lignin yields both in the youngest DOC and the oldest POC samples reflects a
- 56 significant decoupling of terrestrial OC sources and pathways.

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81 1. Introduction

82 Studies of terrestrial organic carbon (Terr-OC) in the Arctic Ocean are receiving increasing interest due to concerns about the consequences on the carbon cycle by amplified 83 84 climate change. The Eurasian Arctic Shelf is predicted to experience the highest increase in temperature on Earth, and its warming is even faster than predicted (Arndt et al., 2015; 85 86 Zwiers, 2002). The sources and the inputs of Terr-OC are likely to vary in the northern shelf margin due to the changing climate. It has been suggested that these changes may translocate 87 88 increasing amounts of Terr-OC to the coastal ocean (Vonk and Gustafsson, 2013). The Arctic tundra and taiga drainage basins represents roughly 50% of the global soil organic matter, 89 much within shallow permafrost (Gorham, 1991; Tarnocai et al., 2009), and 10-20% of the 90 global vegetation carbon with about 73% in Eurasia (McGuire et al., 2009; McGuire et al., 91 92 2010). Fluvial and erosional processes are expected to increase, as well as biomass cover, 93 resulting in higher input fluxes and changing composition of Terr-OC to the continental shelf 94 (Lantuit et al., 2013; Peterson et al., 2002; Sanchez-Garcia et al., 2014; Serreze et al., 2002). 95 In addition, those mechanisms would enhance the remobilization of permafrost carbon, potentially constituting a climate-carbon positive feedback, in terms of CO₂ outgassing from 96 97 degradation of thawing permafrost. Thus, it is essential to understand the sources, dynamical fate and composition of exported Terr-OC in order to assess its impact within the carbon 98 99 cycle.

100 The fate of Terr-OC in the dissolved organic carbon (DOC), particulate organic 101 carbon (POC) and sedimentary organic carbon (SOC) compartments of the marine system is still a matter of debate. Some studies have indicated a conservative behavior of DOC in the 102 103 Arctic Ocean with small influence on the ocean-atmosphere exchange of CO₂ (Amon and Meon, 2004; Dittmar and Kattner, 2003a; Köhler et al., 2003; McGuire et al., 2009), little or 104 105 no degradation in microbial incubations (Amon and Meon, 2004), and high concentrations of 106 lignin in the DOC pool (Amon and Benner, 2003; Amon et al., 2012; Lobbes et al., 2000). By 107 contrast, others suggest that DOC is highly degraded by photochemical oxidation or 108 microbial respiration in the water column or surface sediments (Alling et al., 2010; Benner 109 and Kaiser, 2011; Hernes and Benner, 2003; van Dongen et al., 2008b). Investigations of the particulate compartment indicate that POC degrades much faster than DOC, and just a small 110 111 fraction is transported off-shelf within the POC pool (Eglinton and Repeta, 2006; Sanchez-Garcia et al., 2011; van Dongen et al., 2008b). In addition, other processes on the wide and 112 113 shallow Arctic shelves such as hydrodynamic sorting, deposition, resuspension and uptake by

114 primary production may contribute to the dilution/dispersal of Terr-OC along the water and 115 sediment dispersal system (Stein and Macdonald, 2004; Tesi et al., 2016; Tesi et al., 2014). It seems that different pools of Terr-OC have different behavior and fate during remobilization 116 and transport. DOC and POC pools have much younger ¹⁴C ages than the deposited 117 sedimentary OC (Guo et al., 2007; Karlsson et al., 2016; Karlsson et al., 2011). Compound-118 specific radiocarbon analyses of lipid molecules and lignin phenols of surface sediments and 119 120 POC from major river mouths in the Arctic revealed marked age offsets between different Terr-OC pools (Feng et al., 2013; Vonk et al., 2010), but POC in the Eurasian rivers is not 121 122 well characterized. Further, we still have a very limited understanding of the composition and 123 cycling of Terr-OC in the Arctic Ocean.

The East Siberian Arctic Shelf (ESAS) is a particularly relevant region for 124 investigating the distribution and fate of Terr-OC in the DOC, POC and SOC pools. The 125 ESAS is the world's largest continental shelf and its adjacent basin is located in a region of 126 127 continuous and discontinuous permafrost. The extensive ESAS is quite shallow (~50 m 128 average depth) and receives massive amounts of Terr-OC (22±8 Tg OC/yr; Vonk et al., 2012). In the west, the Lena river and coastal erosion are the main inputs of OC (Laptev Sea 129 and western East Siberian Sea, W-ESS) (Charkin et al., 2011; Salvadó et al., 2015; Semiletov 130 et al., 2011; Tesi et al., 2014; Vonk et al., 2012). Alternatively, in the eastern East Siberian 131 Sea (E-ESS, from ~160°E to eastwards) marine phytoplankton represents an important source 132 133 of OC due to the influence of nutrient-rich Pacific inflow waters (Semiletov et al., 2005; Stein and Macdonald, 2004). Many investigations in the Arctic focused on characterizing the 134 composition and fate of riverine OC (Amon et al., 2012; Benner et al., 2005; Elmquist et al., 135 2008; Goni et al., 2000; Lobbes et al., 2000; van Dongen et al., 2008a; Winterfeld et al., 136 137 2015), sedimentary OC in the ESAS (Bröder et al., 2016; Karlsson et al., 2015; Salvadó et al., 2015; Tesi et al., 2014), and DOC and POC in the water column of the Eurasian Arctic Shelf 138 139 (Alling et al., 2010; Sanchez-Garcia et al., 2011). This is, however, the first study that characterizes collectively the DOC, POC and SOC pools along the outer shelf seeking to test 140 141 the hypothesis that carbon pools may carry different types of OC with different propensity toward off-shelf transport and degradation. The present study uses carbon isotopes and 142 macromolecular biomarkers to provide an extensive view of the composition and distribution 143 144 of Terr-OC along the outer ESAS, with the objective to evaluate the sources, degradation and 145 off-shelf transport of the DOC, POC and SOC pools.

147 **2. Materials and methods**

148 **2.1 Study area**

149 The ESAS is the widest, shallowest and, by area, largest continental shelf in the World Ocean. It comprises 40% of the Arctic shelf and 20% of the Acrtic Ocean (Stein and 150 Macdonald, 2004). This study focuses on the outer shelf of ESAS seas (Laptev Sea and East 151 Siberian Sea) and the Kara Sea (Figure 1). The Kara Sea has an area of $880 \cdot 10^3$ km² and a 152 mean depth of 110 metres. It receives a large amount of fresh water mainly from the Ob river. 153 The Laptev Sea, between ~110°E and 140°E, covers almost $500 \cdot 10^3$ km² and has an average 154 water depth of 50 m. This sea receives large amounts of freshwater (\sim 745 km³·yr⁻¹) mainly 155 transported by the Lena river (566 km³·yr⁻¹) (Cooper et al., 2008; Semiletov et al., 2000), but 156 most of the TerrOC that enters the Laptev Sea is coming from coastal erosion of late 157 Pleistocene ice complex deposits (53±5%) (Semiletov et al., 2011; Vonk et al., 2012). The 158 East Siberian Sea has an average water depth of 58 m, and is the largest and most ice-bound 159 shelf sea of the Arctic Ocean (Stein and Macdonald, 2004). It extends from 140°E to 180°E 160 covering an area of $987 \cdot 10^3$ km², and receives freshwater inputs from the Indigirka and 161 Kolyma rivers. This sea exhibits two physical and biogeochemical regimes. The eastern East 162 Siberian Sea (E-ESS, from ~160°E to ~180°E), which is influenced by the Pacific inflow 163 waters, and primary production represents an important source of OC (Semiletov et al., 2005; 164 Stein and Macdonald, 2004). And the western East Siberian Sea (W-ESS), between ~140°E 165 166 and ~160°E, where river runoff and coastal erosion of thawing permafrost supply the major 167 part of OC, but there is also a relatively high marine productivity, particularly in certain polynya regions. 168

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170 **2.2 Sampling**

171 A comprehensive set of samples was obtained during July-August 2014 as part of the international Swedish-Russian-US investigation of the Carbon-Climate-Cryospshere 172 Interactions in the East Siberian Arctic Ocean (SWERUS-C3) expedition onboard I/B Oden. 173 174 The first of two SWERUS-C3 2014 legs was an extensive 45-day campaign of complex geophysical and hydrogeochemical sampling including at-sea analysis. The sample collection 175 176 for this study consisted of four types of samples along the outer ESAS (Figure 1): i) POC from high volume filtration on 293 mm glass fibre filter (GF/F; Whatman Inc.) with a 177 178 nominal 0.7 µm cut-off, ii) POC from 47 mm Teflon filters (1 µm cut-off), iii) DOC isolated

with solid phase extraction (SPE) cartridges, iv) surface sediment samples collected with amulticorer.

181 Surface and near-bottom waters (5m above bottom) were sampled and filtered through high volume 293 mm GF/F filters (pre-combusted for 5h at 500°C). Samples were filtered 182 either directly from the seawater intake (SWI) or by pumping water from 1000L tanks filled 183 184 from the SWI or from a submersible pump. The systems were connected to an electronic flow meter, in the flow path below the filter, and a pressure meter situated directly above the GF/F 185 filter holder. We maintained the flow to about 8.5 $L \cdot min^{-1}$, and stopped filtering before the 186 backpressure reached 1 bar to avoid cell lysing. After sampling of the particulate fraction, the 187 188 GF/F filters were folded, put in a pre-combusted aluminium foil and stored at -20°C. Since GF/F filters are not compatible with the alkaline hydrolysis of the CuO oxidation protocol to 189 analyse lignin-derived phenols, POC samples were also obtained on 47 mm Teflon filters in 190 191 order to analyse lignin-derived phenols in POC. We placed the Teflon filters in the filtration 192 unit and applied a positive pressure flow with a peristaltic pump at a flow rate of 25 ml·min⁻¹. 193 POC samples in Teflon filters were folded in two, placed in petri dishes and stored frozen (-194 20°C) until laboratory analysis.

195 The dissolved fraction of organic matter was isolated by high-volume SPE cartridges 196 containing 10 g of sorption material composed of octadecyl carbon moieties (C_{18}) chemically 197 bonded to a silica support (C₁₈-SPE Mega-Bond Elut; Agilent) (Louchouarn et al., 2000). 198 Cartridges were preconditioned with 5 resin volumes of methanol followed by 5 resin 199 volumes of acidified (pH 2) Milli-Q Plus UV water. The water samples, previously filtered with GF/F, were acidified to pH 2 using reagent-grade concentrated HCl and pumped through 200 201 the SPE cartridge with a peristaltic pump and silicone tubing. By this method, the water (~30L) was delivered directly into the headspace of the SPE cartridge and forced by pressure 202 through the sorbent at a flow rate of 100 mL min⁻¹. Thereafter, we rinsed each SPE cartridge 203 with 1L of acidified (pH2) Milli-Q Plus UV water to remove residual salts. Sample cartridges 204 205 were packed in aluminum foil and stored at 4°C until further processing.

Sediment cores were collected at water depths of 40 to 3120 m with an 8-tube multicorer (Oktopus GmbH, Germany), which was developed to collect samples of the seabed with an undisturbed sediment-water interface. The liners were made of polycarbonate and were 60 cm long with a 10 cm diameter. The multicorer was deployed with full weight (head weight about 500 kg) at a speed of 0.5 m/s near the seabed. To increase recoveries, the multicorer was left for 1 minute on the seafloor. The cores were sectioned on low resolution 212 (1cm intervals; shelf stations <200m water depth) or on high resolution (0.5cm intervals; 213 slope and rise stations >200m water depth), and sediment samples were transferred into 214 plastic bags and stored in the freezer (-20°C). This study focuses on surface sediments (0-1 215 cm).

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217 **2.3 Bulk elemental and isotope analysis**

The analyses of organic carbon content, $\delta^{13}C$ and $\Delta^{14}C$ in the DOC, POC and SOC 218 pools have been described earlier (Karlsson et al., 2011; Louchouarn et al., 2000). Briefly, 219 220 DOC was determined onboard after GF/F filtration by high-temperature catalytic oxidation 221 (Shimadzu TOC-L_{CPH}). In the laboratory, SPE cartridges were eluted with 50 mL of methanol. Then, we subsampled 0.5-1mL of the eluent, depending on DOC concentrations, 222 and placed it in smooth wall tin capsules for liquids (6 x 12 mm, Elemental Microanalysis, 223 Devon, UK). For organic carbon content and δ^{13} C composition of POC and SOC, GF/F filters 224 and surface sediment samples were subsampled and liquid acidified with HCl (1.5M) to 225 remove carbonates. The analyses were performed in triplicates using a Carlo Erba NC2500 226 elemental analyzer connected via a split interface to a Finnigan MAT Delta Plus mass 227 spectrometer at the Stable Isotope Laboratory of the Department of Geological Sciences at 228 Stockholm University. Some subsamples, after similar preparation steps, were analyzed for 229 its radiocarbon content (Δ^{14} C) at the US-NSF National Ocean Sciences Accelerator Mass 230 Spectrometry (NOSAMS) Facility at Woods Hole Oceanographic Institution. Uncertainties of 231 Δ^{14} C, δ^{13} C, and OC analyses were ±0.002 (fraction modern error), ±0.1‰, and ±2% of the 232 measured OC content, respectively. 233

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235 **2.4 Lignin phenols analysis**

The quantification of lignin-derived phenols in DOC (SPE eluents), POC (Teflon 236 filters), and SOC (surface sediment samples) was performed as described in detail by 237 Louchouarn et al 2000 and Tesi et al., 2014. Briefly, for the analysis of dissolved lignin, 5-15 238 239 mL of elution samples (1-2 mg OC equivalent) were reduced to dryness under a stream of nitrogen in Teflon tubes. The dried samples were then oxidized under alkaline oxygen free 240 241 conditions (degassed NaOH solution, 8%) (Goni and Montgomery, 2000) with an addition of 10 mg of glucose to prevent superoxidation of the lignin polymer and spiked with recovery 242 standards (trans-cinnamic acid and ethyl vanillin). Samples were then acidified, extracted 243

twice with ethyl acetate and concentrated under vacuum at 60°C. The same oxidation and
extraction procedure was also used for POC (Teflon filters) and SOC samples, but without
the addition of glucose in sediment samples.

247 Prior to the analyses, extracts were re-dissolved in pyridine and derivatized. Target compounds were quantified on a gas-chromatograph mass spectrometer (GC-EI-MS, Agilent) 248 using a DB1-MS capillary column (30m x 250µm, 0.25µm stationary phase thickness, 249 Agilent J&W) for separation. Quantification of lignin phenols, benzoic acids, and p-250 hydroxybenzenes was achieved using the response factors of external standards. All reported 251 concentrations of CuO oxidation products were reported in mg of biomarker per g OC. Some 252 sediment samples (SWE-1, SWE-4, SWE-6, SWE-14, SWE-23, SWE-24) were also analysed 253 by Bröder et al. (2016). The small differences in lignin phenols results may stem from the 254 different injections and calibrations used. 255

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257 **3. Results and discussion**

258 **3.1 Elemental Composition and Distribution of DOC, POC and SOC**

DOC in the water column of the outer ESAS during SWERUS-2014 expedition was 259 one order of magnitude higher than POC. The DOC concentrations ranged from 740 to 3600 260 $\mu g \cdot L^{-1}$ (mean of 1400±790 $\mu g \cdot L^{-1}$) and POC varied between 20 and 360 $\mu g \cdot L^{-1}$ (mean of 261 $110\pm80\mu$ g·L⁻¹) (Table 1, Figure 2A, Figure 3). Those values are in the same range as previous 262 studies in the Siberian Arctic Seas (Alling et al., 2010; Benner et al., 2005; Sanchez-Garcia et 263 al., 2011). Whereas DOC showed the highest values in surface waters of the Laptev Sea 264 $(2000\pm1100 \ \mu g \cdot L^{-1})$, particularly off the Lena river mouth, POC concentrations were slightly 265 higher in the Kara Sea $(290\pm86\mu g \cdot L^{-1})$ and the E-ESS $(150\pm92\mu g \cdot L^{-1})$ with no significant 266 differences between surface and near-bottom waters. SOC values in surface sediments from 267 the same stations presented higher concentrations in the E-ESS (1.32±0.42%), but also 268 exhibited an increase in the Laptev Sea $(1.21\pm0.26\%)$ (Figure 3). This is in the lower range of 269 what was previously reported in the inner-shelf of the ESAS (Charkin et al., 2011; Karlsson 270 et al., 2015; Karlsson et al., 2011; Tesi et al., 2014; Vonk et al., 2012), suggesting either 271 degradation of Terr-OC or sediment sorting during the across-shelf transport as discussed in 272 273 Tesi et al. 2014 and Bröder et al. 2016. Alternatively, higher POC and SOC values in the E-ESS may be related to the higher marine productivity in that region due to the Pacific water 274 275 influence (Semiletov et al., 2005; Stein and Macdonald, 2004).

276 The resulting bulk ratios in the DOC_{SPE} fraction indicate terrestrially dominated organic matter sources. The OC/TN (TN = organic nitrogen + inorganic nitrogen) of DOC_{SPE} 277 ranged between 14 and 43 (mean of 28 ± 8.4) without significant differences between surface 278 279 and near-bottom waters (Table 1). Those ratios showed decreasing trends off the Lena river 280 plume with higher ratios in the Laptev Sea and W-ESS. The same pattern and similar ratios were observed in the inner-shelf of the ESAS (Karlsson et al., 2016). Moreover, these values 281 282 are in the same range as OC/TN ratios of DOC in Eurasian Arctic rivers, which varied between 23 and 69 (Lobbes et al., 2000), and the high OC/TN ratios (>40) of DOC collected 283 284 from the Kara Sea (Köhler et al., 2003; Opsahl et al., 1999). Marine organic matter has OC/TN values around 6-8 and terrestrial derived organic matter OC/TN ratios higher than 15 285 (Baldock et al., 1992; Hedges et al., 1986; Hedges and Oades, 1997). The OC/TN ratios in 286 the particulate and sedimentary compartments were much lower than in the DOC_{SPE}. Those 287 ratios ranged between 5 and 12 (mean of 7 ± 1.7) in the POC and from 6 to 8 (mean of 7 ± 0.5) 288 in the SOC. Similar OC/TN values were observed in the inner-shelf of the ESAS and Arctic 289 rivers in the particulate fraction (McClelland et al., 2016; Sanchez-Garcia et al., 2011). 290 However, these lower OC/TN ratios are at odds with e.g. δ^{13} C-OC and may be influenced by 291 selective degradation of labile carbonaceous forms (Hugelius and Kuhry, 2009), and/or 292 adsorption of inorganic nitrogen (e.g. ammonium) derived from decomposition of organic 293 294 matter (Sanchez-Garcia et al., 2011; Schubert and Calvert, 2001).

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296 **3.2 Stable Carbon and Radiocarbon Isotopes**

The east-to-west extension of the depleted $\delta^{13}C$ signatures reflects a strong influence 297 of the Lena River (Figure 3), both in the Laptev Sea and the ESS. The δ^{13} C signatures in the 298 three carbon pools ranged from -23.9±1.9‰ in the SOC, -26.1±1.2‰ in the DOC_{SPE} and -299 27.1±1.9‰ in the POC, with no significant differences between surface and bottom waters 300 (Tables 1, 2 and 3). The more depleted δ^{13} C-POC is consistent with marine productivity 301 using excess dissolved inorganic carbon (DIC) from the Lena river, which is more depleted 302 than marine DIC (Alling et al., 2012; Semiletov et al., 2016). This mechanism also explains 303 similarly depleted δ^{13} C-POC in the Lena plume far offshore in the Laptev Sea and ESS that 304 matched with depletion of other nutrients (Alling et al., 2010; Sanchez-Garcia et al., 2011). 305 The distribution of δ^{13} C-SOC was more homogeneous reflecting average over time in the 306 307 surface sediment regime. Only the concentration of DOC presented a good correlation with

308 δ^{13} C-DOC_{SPE} signatures, which indicates that higher concentrations of DOC come from 309 terrigenous sources (Figure 3).

The radiocarbon ages of DOC_{SPE} and POC showed a depleted and younger trend off 310 the Lena River plume. The Δ^{14} C signals ranged between -395±83‰ (SOC), -226±92‰ 311 (DOC_{SPE}) and -113±122‰ (POC) presenting contrasting offsets between the Laptev Sea and 312 the East Siberian Sea, particularly in the E-ESS (Tables 1, 2 and 3; Figure 4). The older and 313 enriched $\delta^{13}C$ signatures in the outer-shelf of the ESS may reflect the influence of sea ice 314 coverage and the Pacific inflow from the East. We suggest that the sea ice would work as a 315 barrier preventing the direct terrigenous input from inland and reinforcing the influence of 316 Pacific waters. The more enriched Δ^{14} C signatures in POC than in DOC are in accordance 317 with previous studies in the Arctic Ocean (Griffith et al., 2012) and the Southern Ocean 318 319 (Druffel and Bauer, 2000), which reflect likely a dominant marine source in the particulate carbon pool. The radiocarbon signatures in the DOC pool of the outer-shelf of the ESAS are 320 321 older than those observed in the Lena River (>39‰) (Raymond et al., 2007), but younger or 322 similar (in the outer and eastern stations) than those reported in surface waters of the Canada 323 Basin (<-216‰) (Arctic Ocean) (Griffith et al, 2012), reflecting the inputs of Pacific waters. 324 The considerable change in age within the DOC pool during the cross-shelf transport is likely due to mixing with older marine DOC. The SOC pool does not present marked west-east 325 distribution of Δ^{14} C as observed in DOC_{SPE} and POC. The SOC also depicts older signatures 326 near the New Siberian Islands. A recent study from the same area at the land-ocean interface 327 presented older signatures in the POC than in the DOC (Karlsson et al., 2016), suggesting 328 that thawing permafrost was transported preferentially within the POC pool. Therefore, our 329 330 results support the hypothesis that remobilized permafrost preferentially settles out close to 331 land, and then it is transported off-shelf through sediment resuspension-redeposition events. The older signals in the dissolved fraction of the ice-covered regions are consistent with a 332 more recalcitrant OC in the dissolved pool of the Arctic Ocean (Follett et al., 2014; Griffith et 333 334 al., 2012). It seems that the ice extent boundary works as a barrier that prevents the input of young DOC coming from the buoyant freshwater plume of the Lena river (Figure 4). It is 335 important to point out that near-bottom waters presented more depleted and similar $\Delta^{14}C$ 336 signatures in both DOC_{SPE} and POC ($-258\pm94\%$ and $-250\pm83\%$, respectively) than in surface 337 waters (-213±93‰ and -57±86‰, respectively) (Figure 5; Tables 1 and 2), suggesting the 338 same older and terrigenous source of OC in both pools. Those contrasting age offsets 339

between surface and near-bottom waters, particularly for the POC fraction, may reflect theoff-shelf transport of OC translocated over long distances from thawing permafrost.

DOC was the only carbon pool that presented good correlations with Δ^{14} C and δ^{13} C 342 343 data (Figure 6). Those relationships are consistent with previous observations in the Arctic Ocean (Amon et al., 2012; Benner et al., 2004; Schreiner et al., 2013). The correlation 344 between δ^{13} C and DOC (r²=0.68), with more depleted values for samples with a high DOC 345 concentration, and vice versa, reflects both mixing and the transport of the terrigenous DOC 346 347 along the shelf. Processes such as hydrodynamic sorting, deposition, resuspension and uptake by primary production may contribute to the dispersal and processing of the OC in the ESAS. 348 On the other hand, the relationship between Δ^{14} C and DOC (r²=0.87) represents both mixing 349 and the source of the terrigenous DOC, where samples with higher DOC concentrations are 350 composed by young Terr-OC and lower DOC concentrations by old and refractory Terr-OC. 351 Overall, these findings are direct evidence that a large proportion of DOC exported to the 352 353 outer shelf comes from young and fresh vascular plant material.

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355 **3.3 Lignin-Derived Phenols**

Lignin-derived phenols are exclusively synthesized by vascular plants and account for 356 one third of the organic matrix of wood, grasses, needles, and herbage, therefore, they have 357 been extensively used to characterize the pathway of terrestrial matter in the marine 358 environment (Louchouarn et al., 1999; Pasqual et al., 2013; Tesi et al., 2014). The carbon-359 normalized lignin content (mg/g OC) refers to the sum of vanillyl, syringyl and cinnamyl 360 phenols. DOC_{SPE} samples presented lignin concentration on the same order as the 361 corresponding underlying sediments (0.10-2.34 and 0.08-2.40 mg/g OC, respectively). By 362 363 contrast, the particulate carbon pool had slightly lower OC-normalized lignin concentrations between 0.03 and 1.14 mg/g OC (Figure 4; Tables 1 and 2). Some surface POC samples, the 364 365 ones in the Kara Sea, showed more lignin than in the DOC, but those concentrations were relatively low compared to the higher lignin yields observed in the dissolved and sedimentary 366 367 OC pools from the ESAS. Lignin levels are relativity small when compared to the actual river/coastal erosion input (Amon et al., 2012; Lobbes et al., 2000), but in the same range to 368 369 lignin values in SOC and DOC pools from previous studies in ESAS sediments (Karlsson et 370 al., 2015; Tesi et al., 2014) and the polar surface water of the Arctic Ocean (Benner et al., 371 2005). These are the first POC-lignin data in the Arctic Ocean. Lignin concentrations 372 exhibited contrasting offsets between surface and near-bottom waters, particularly in the POC

pool. DOC_{SPE} presented similar lignin concentrations in surface and near-bottom waters, 373 except for the more concentrated samples closer to the Lena river mouth. Conversely, POC 374 375 showed enhanced levels in all near-bottom water samples (from 0.17 to 1.14 mg/g OC), with 376 even higher concentrations than in the dissolved pool (from 0.16 to 0.91 mg/g OC) (Figure 377 2B; Tables 1 and 2). Those vertical lignin dissimilarities in the water column were not observed in the total OC of the dissolved and particulate fractions. However, Δ^{14} C-OC also 378 showed offsets in the particulate pool. While DOC_{SPE} depicted similar Δ^{14} C signatures in 379 both surface and near-bottom waters (Figure 5B), POC was much older in near-bottom waters 380 381 (Figure 5A). Hence, these findings suggest that particulate old OC with high concentrations of lignin, probably coming from thawing permafrost, is mainly transported off-shelf in near-382 bottom waters by resuspension and remobilization of the SOC pool. 383

384 Lignin phenols exhibited decreasing OC-normalized concentrations with increasing distance from the Lena river plume in the DOC and SOC pools and in both surface and near-385 386 bottom waters (Figure 4). Previous studies in ESAS for other biomarkers have also reported 387 decreasing across-shelf trends of terrestrial organic matter with increasing distance from the coast (Selver et al., 2015; Tesi et al., 2014). Several studies reported minimal degradation of 388 389 DOM across the broad Eurasian shelves (Dittmar and Kattner, 2003b; Kattner et al., 1999; Köhler et al., 2003). With such a scenario, our off-shelf decreasing lignin concentrations in 390 391 DOC, POC and SOC pools may be interpreted to result from dilution with marine organic matter during transport and/or hydrodynamic sorting along the water and sediment dispersal 392 393 system. However, other studies found that terrestrial DOC in this ESAS shelf sea system was degraded, with a first-order removal rate constant of 0.3 yr^{-1} (Alling et al., 2010). Recent 394 395 studies also suggested high reactivity of lignin in rivers (Benner and Kaiser, 2011; Fichot and Benner, 2014; Ward et al., 2013) and in offshoreward direction across ESAS (Bröder et al., 396 397 2016; Tesi et al., 2014). If this instead is the dominating process, the decreasing trend in the 398 current study may also be due to degradation.

Our results depicted a strong positive relationship between lignin phenols and total dissolved organic content within the 35 DOC_{SPE} samples analysed along the outer ESAS (r = 0.89) (Figure 7). There were also significant correlations between OC-normalized concentrations of lignin phenols and δ^{13} C (r = 0.66) and Δ^{14} C (r = 0.78) in the DOC_{SPE} pool. These data is consistent with the modern radiocarbon ages of DOC observed in Arctic rivers (Benner et al., 2004; Benner et al., 2005; Karlsson et al., 2016), which also demonstrated a general agreement between lignin phenols and Δ^{14} C signatures as traces of terrrigenous DOC. Lignin phenols were found in old OC from permafrost (Tesi et al., 2014). Compound-specific radiocarbon analyses of lignin phenols from sediments off major river mouths in ESAS indicated that those macromolecules were younger than sedimentary bulk OC (Feng et al., 2013). This is consistent with lignin compounds derived from both sources, and the higher lignin content from younger DOC_{SPE} likely coming from either recently produced vascular plant material or from contemporary topsoil.

By contrast, the slightly negative correlation between lignin phenols and $\Delta^{14}C$ 412 signatures in POC (r = 0.53) (Figure 7), with higher lignin concentrations in older POC, 413 suggests that those macromolecules are coming from remobilized older permafrost carbon. 414 415 Those results are consistent with previous findings indicating that OC from thawed permafrost is transported preferentially within the particulate carbon pool (Karlsson et al., 416 417 2016). There was no relationship between lignin content and bulk POC and SOC, which suggests that both pools are composed by a mixture of marine and terrestrial organic carbon. 418 419 Taken together, whereas "young" Terr-OC is transported mainly within the surface dissolved 420 fraction, near-bottom POC and SOC carries off-shelf preferentially old OC from remobilized permafrost. 421

422

423 **3.4 Biomarker indications of sources of DOC, POC and SOC**

The ratios of individual or classes of lignin phenols are frequently used to infer the 424 types of plants yielding the phenols and to what extent the organic matter has been oxidized. 425 Vanillyl phenols (vanillin, acetovanillone and vanillic acid) are ubiquitous in lignin, while 426 427 syringyl phenols (syringaldehyde, acetosyringone and syringic acid) derive only from angiosperms (Hedges and Mann, 1979; Spencer et al., 2008). Ratios of syringyl to vanillyl 428 429 (S/V) phenols indicate contribution of angiosperm and gymnosperm vegetation to Terr-OC. Our low S/V ratios (from 0.14 to 1.05) indicate gymnosperm vegetation as the most 430 431 important source of lignin (Figure 8A; Tables 1, 2 and 3). However, the high S/V ratios in the easternmost samples, particularly within the DOC and SOC, reflect a higher source apportion 432 of tundra plants (Lobbes et al., 2000). Elevated values of S/V were also reported in sediments 433 434 and dissolved organic carbon from the inner-shelf of the same study area (Karlsson et al., 435 2016; Tesi et al., 2014). The fact that the Indigirka and Kolyma watersheds are north of the Arctic Cirle with a general shift to flowering tundra plants could explain the elevated S/V 436 437 ratios in the E-ESS.

438 Cinnamyl phenols (p-coumaric acid, ferulic acid) are predominantly found in herbaceous tissues, and the ratio cinnamyl over vanillyl (C/V) has been used to distinguish 439 440 woody lignin from other sources (Goni and Hedges, 1992; Hedges and Mann, 1979). C/V 441 ratios did not show a specific trend along the east to west data set. Similar results were 442 observed previously in inner-shelf sediments and in the colloidal DOC fraction from the ESAS (Karlsson et al., 2016). As we also analysed lignin phenols in the particulate fraction, 443 444 we could see that C/V ratios were slightly higher in POC (0.64±0.42) than in SOC (0.37±0.18) (Tables 1, 2 and 3), possibly reflecting more herbaceous plants or sphagnum 445 446 moss source in the particulate pool and more woody lignin in the sedimentary carbon. Those 447 ratios should always be carefully interpreted as photooxidation and microbial degradation can alter the original compositions (Hedges and Prahl, 1993; Opsahl and Benner, 1995). 448 However, degradation lowers both S/V and C/V ratios, but mostly C/V. If degradation was 449 indeed the main process we would expect a correlation between S/V and C/V. Therefore, 450 while some degradation cannot be excluded, the observed differences in lignin phenols 451 between carbon pools likely reflect a different source. Regarding the classical source plot of 452 S/V versus C/V, our data set distributes along a line between angiosperm leaves and grasses 453 454 and gymnosperm wood, suggesting that little amounts of non-woody angiosperm tissues are 455 mixing with large amounts of gymnosperm woods in these samples (Figure 8A). Overall, this plot underlines distinct clustering between OC pools, suggesting that angiosperms are mainly 456 457 transported by SOC and gymnosperms by POC.

458 p-hydroxybenzoic acids (P) can originate from different sources, while phydroxyacetophenone (Pn) has only been detected in terrigenous organic matter, particularly 459 460 in peat and sphagnum (Williams et al., 1998), while p-hydroxybenzaldehyde (Pl) and phydroxybenzoic acid (Pd) can also derive from marine sources (Goni and Hedges, 1995). The 461 Pn/P ratios observed in DOC (0.08-0.37), SOC (0.06-0.17) and POC (0.02-0.14) suggest that 462 the OC in the dissolved pool contains a larger terrestrial component than in the particulate 463 and sediment pools (Tables 1, 2 and 3). Those ratios present a slight east-to-west trend with 464 465 higher values off the Lena river plume. Similar trends and results were observed by Karlsson et al., 2016 in the colloidal OC along the ESAS coast (0.15-30). Amon et al., 2012 466 characterized the chemical composition of DOC in Arctic rivers and reported Pn/P ratios in 467 the same range, for instance, those ratios in the Lena, Indigirka and Kolyma rivers varied 468 between 0.30 and 0.39. Further, P/V ratios presented an opposite trend to Pn/P ratios with 469 470 much higher values in the POC (11.9 \pm 9) than in the DOC (1.4 \pm 0.6) and SOC pools (4 \pm 3.6)

471 (Tables, 1, 2, 3). These ratios are in agreement with the relationships of DOC and Δ^{14} C and 472 δ^{13} C presented above, which indicate that the DOC exported off-shelf is mainly "young" and 473 terrestrial. Nevertheless, it is important to note that these proxies should be interpreted 474 carefully as P products account for less than 0.1-0.2% of the bulk OC, and in some samples 475 the particulate and sedimentary pools have more lignin yield than the dissolved OC.

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477 **3.5 Indicators of Terr-OC degradation across the OC continuum**

The relative abundances of some lignin phenols provide information about the 478 479 diagenetic alteration of Terr-OC. The acid/aldehyde ratios of syringyl (Sd/Sl) and vanillyl 480 (Vd/Vl) have been utilized as indicators of the relative degradation of the plant matter contribution, as aldehydes degrade faster than corresponding acids (Goni and Hedges, 1992; 481 482 Hedges et al., 1986). However, some caution should be applied in the interpretation as source signals are more varied than originally thought, and fractionation occurs during 483 484 leaching/adsorption processes (Benner et al., 1990; Hernes et al., 2007). Our data showed Sd/Sl and Vd/Vl ratios higher in DOC_{SPE} (1.9±0.6 and 2.0±0.7, respectively), than in SOC 485 486 $(0.9\pm0.2 \text{ and } 1.1\pm0.3)$ and POC $(0.4\pm0.1 \text{ and } 0.5\pm0.2)$ indicating the presence of highly 487 oxidized lignin in the dissolved pool (Tables 1, 2 and 3; Figure 8B). It is important to notice 488 that the ranges of Sd/Sl and Vd/Vl ratios in this study were relatively broad and with clear 489 clusters between carbon pools. POC ratios were lower than the underlying sediments and the 490 dissolved carbon pool presented very high ratios (Figure 8B). Those ratios are in accordance with global studies on POC (Hernes and Benner, 2002; Lobbes et al., 2000; Winterfeld et al., 491 2015), sediments (Goni and Montgomery, 2000; Goni et al., 2005; Tesi et al., 2014) and 492 493 DOC (Amon et al., 2012; Hernes and Benner, 2002; Lobbes et al., 2000), which also found higher ratios in the dissolved than in the particulate phase. The elevated Sd/Sl and Vd/Vl in 494 495 the dissolved fraction, as well as the enhanced ratios in the SOC pool, may reflect leaching/adsorption processes (Hernes et al., 2007; Houel et al., 2006). 496

Another proxy commonly used to determine the degradation of Terr-OC is the ratio between 3,5-dihydroxybenzoic acid and vanillyl phenols (3,5-Bd/V) (Farella et al., 2001; Houel et al., 2006; Otto and Simpson, 2006; Prahl et al., 1994). Since 3,5-Bd is highly resistant to degradation (Dickens et al., 2007) while vanillyl phenols are very susceptible to degradation, higher values of 3,5-Bd/V are indicative of more degraded Terr-OC. Our results presented opposite patterns than the ones observed by Sd/Sl and Vd/Vl with higher 3,5-Bd/V ratios in POC (1.7 ± 0.7) and SOC (1.0 ± 0.6) and lower ratios in DOC_{SPE} (0.7 ± 0.3) (Tables 1, 2 504 and 3; Figure 9B). These values are in accordance with those in DOC from ESAS rivers (0.4-505 0.7) (Amon et al., 2012) and the colloidal fraction from the ESAS land-ocean interface (0.4-506 0.8) (Karlsson et al., 2016). In addition, SOC ratios are consistent with those observed in 507 surficial sediments from the same area (0.2-1.3) (Tesi et al., 2014). However, we could not 508 find previous studies to compare our 3,5-Bd/V ratios in POC. The higher ratios in POC suggest that Terr-OC is more degraded in the particulate fraction than in the other carbon 509 510 pools of the outer ESAS. We should also consider that those ratios could support the role of a source change as observed with S/V and C/V proxies, and macroalgal sources of 3,5-Bd 511 512 might be significant in selected marine systems comprising minimal fractions of terrigenous 513 organic matter (Goni and Hedges, 1995). The 3,5-Bd/V ratios in DOC_{SPE}, POC and SOC 514 depicted a slightly increasing tendency in the eastern samples (Figure 9B). Previous studies in sediments and the colloidal fraction from the ESAS also reported the same trend (Karlsson et 515 516 al., 2016; Tesi et al., 2014) reflecting the Pacific inflow from the east of more marine and/or degraded OC. We consider in our study that this degradation proxy is more reliable than 517 Sd/S1 and Vd/V1 ratios as it is not affected by the leaching/adsorption processes between 518 carbon pools. Therefore, the Terr-OC in the ESAS is more degraded in the POC and SOC 519 520 pools.

Two cinnamyl phenols, *p*-coumaric acid (pCd) and ferulic acid (Fd), are additional 521 522 CuO oxidation products of lignin that are particularly abundant in grasses and many herbaceous tissues. The two phenols differ by a presence of a methoxyl group, and this may 523 524 explain the preferential degradation of ferrulic acid (Opsahl and Benner, 1998). Therefore, 525 pCd/Fd ratio has been used as a diagenetic indicator (Amon et al., 2012; Houel et al., 2006). 526 In this data set pCd/Fd ratios follow the same pattern as the ones observed in 3,5-Bd/V ratios 527 with higher values in POC and SOC and a slightly increasing tendency in the eastern SOC samples (Tables 1, 2 and 3; Figure 9C). This strengthens the hypothesis that POC and SOC 528 529 are more degraded than DOC.

The strong relationship between lignin concentrations and the ¹⁴C-age of DOC_{SPE} also reflects the role of diagenetic processes. The younger the marine DOC is, the higher is the concentration of lignin (Figure 7). Those relationships are consistent with previous observations in the Arctic Ocean where the age of DOC decreased with increasing concentration of lignin (Benner et al., 2004). These results suggest that a large proportion of DOC exported to the outer shelf of the ESAS, off the Lena river, comes from recently produced vascular plant material with little exposure to microbial degradation. Whereas most 537 of terrigenous POC settles out close to land and is transported through repeated cycles of 538 deposition and resuspension across the shelf, DOC is dispersed further out onto the EAS with 539 variable extends of conservative mixing.

540

541 **4. Conclusions**

542 This extensive study provides improved understanding on the sources and composition of Terr-OC in the DOC, POC and SOC pools in the extensive outer ESAS. The 543 distribution of a wide variety of bulk (δ^{13} C and Δ^{14} C) and macromolecular proxies (lignin-544 545 derived phenols) reflects a strong influence of the Lena river on the outer shelf, both in the 546 Laptev Sea and the western ESS. These findings demonstrate that a large proportion of the surface DOC exported off-shelf comes from "young" and fresh vascular plant material. The 547 older and more enriched $\delta^{13}C$ signatures in the E-ESS and its higher POC and SOC 548 concentrations suggest a greater influence of sea ice coverage and the Pacific inflow. Near-549 bottom waters present more depleted Δ^{14} C signatures and higher concentrations of lignin, 550 particularly within the POC pool. This is a key evidence of decoupling of the POC and DOC 551 552 pools and reflects the off-shelf transport of permafrost-derived OC in the nepheloid layer, through repeated cycles of deposition and resuspension across the shelf. The ratios of S/V 553 554 indicate gymnosperm vegetation as the most important source of lignin, and increasing S/V 555 ratios in the easternmost samples reflect a relatively higher source contribution of tundra plants. Moreover, the opposite trends in the Pn/P and P/V ratios also indicate that DOC 556 primarily contain a terrigenous OC. Taking together S/V and C/V ratios we observe distinct 557 clustering between DOC, POC and SOC, suggesting that those pools are carrying Terr-OC of 558 partially different origin. Regarding the degradation state of Terr-OC, lignin-phenols 559 fingerprints are presenting contrasting results. While acid/aldehyde ratios are higher for 560 DOC, possibly due to fractionation during leaching, 3,5-Bd/V and pCd/Fd ratios were 561 enhanced in POC and SOC, suggesting more degradation. If this hypothesis is true, the 562 563 remobilized OC from permafrost, which is mainly transported within POC and SOC pools, 564 could experience less burial and more mineralization than the DOC pool. The high abundance 565 of Terr-OC in the outer ESAS, particularly in the dissolved and sedimentary carbon pools, is a clear indicator of the magnitude of shelf to basin transport. Overall, the results are a key 566 evidence for decoupling of the DOC, POC and SOC pools in the ESAS and elucidate the off-567 shelf transport of permafrost-derived OC in the particulate pool of near-bottom waters. 568

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ID	Region	Lat	Long	DOC^1	C/N	$\delta^{13}\!C$	$\Delta^{14} C$	Lignin ²	S/V	C/V	Sd/S1	Vd/Vl	3.5Bd/V	Pn/P	P/V
DOC	C-swi ³														
T-1	KS	79.8	67.9	882	14	-24.7		0.1	0.4	0.5	2.6	3.1	1.2	0.1	2.6
T-2	KS	81.7	75.3	906	15	-24.6		0.1	0.4	0.6	1.8	2.1	1.3	0.2	2.7
T-3	LS	81.3	109.4	809	18	-25.1		0.3	0.5	0.3	1.6	1.9	0.8	0.3	1.5
T-4	LS	81.0	112.9	1171	18	-25.4		0.2	0.4	0.3	2.0	2.4	0.8	0.2	1.5
1	LS	78.9	125.2	737	18	-24.9	-279	0.2	0.5	0.4	1.7	1.9	1.0	0.3	1.9
4	LS	77.8	126.7	1183	19	-25.9	-214	0.4	0.4	0.2	2.7	2.7	0.6	0.2	1.0
6	LS	77.1	127.4	1127	18	-26.0									
13	LS	76.8	125.9	1326	22	-26.3									
14	LS	76.9	127.8	1418	25	-26.8		0.5	0.4	0.2	1.7	2.0	0.4	0.3	0.9
23	LS	76.2	129.3	3428	43	-27.8		2.4	0.3	0.2	1.5	1.5	0.2	0.2	0.4
24	LS	75.6	129.6	3347	42	-27.9		2.1	0.3	0.2	1.6	1.5	0.3	0.2	0.4
25	LS	76.0	130.7	3598	43	-27.9	-19	2.0	0.3	0.2	1.9	1.7	0.3	0.1	0.4
26	LS	76.5	132.0	3428	40	-27.6									
27	LS	76.9	132.2	2159	36	-27.4									
28	LS	77.3	134.8	2650	39	-27.6	-90	1.2	0.3	0.2	1.6	1.7	0.3	0.2	0.5
29	LS	77.8	136.7	1695	33	-27.0									
39	W-ESS	77.7	141.4	2038	36	-27.4		0.8	0.3	0.2	1.7	1.4	0.5	0.2	0.6
40	W-ESS	77.6	145.8	2191	36	-27.2		0.7	0.4	0.3	2.0	1.8	0.3	0.1	0.6
41	W-ESS	77.0	148.3	1373	35	-29.0									
44	W-ESS	76.3	146.0	1289	26	-26.4	-160	0.4	0.4	0.3	1.9	1.7	0.5	0.3	1.1
45	W-ESS	76.4	148.1	1078	30	-26.9		0.3	0.5	0.4	2.8	3.0	0.7	0.2	1.4
46	W-ESS	76.4	149.9	935	25	-26.3									
48	W-ESS	76.5	150.8	995	22	-26.2		0.2	0.5	0.5	2.9	3.1	1.0	0.2	1.7
49	W-ESS	76.5	156.9	1264	23	-25.4		0.1	0.6	0.5	2.8	3.6	0.9	0.1	1.9
50	W-ESS	75.8	158.5	1086	21	-25.3	-262	0.2	0.5	0.5	3.1	3.1	1.0	0.1	1.8
52	E-ESS	74.1	160.6	881	24	-25.3	-288	0.1	0.4	0.1	0.6	1.0	0.6	0.3	2.6
56	E-ESS	74.6	161.9	922	21	-24.9									
57	E-ESS	74.4	163.7	847	23	-24.5		0.2	0.7	0.9	1.3	1.8	0.9	0.2	1.9
58	E-ESS	74.4	166.2	854	23	-24.7		0.1	0.5	0.5	3.5	3.8	1.3	0.1	2.2
59	E-ESS	74.4	168.5	748	25	-25.0									
60	E-ESS	73.4	169.5	853	21	-24.6	-268	0.2	0.7	0.7	1.6	1.9	0.6	0.2	1.7
61	E-ESS	74.1	170.9	890	26	-24.4	-278	0.2	0.6	0.8	2.3	2.8	1.5	0.2	2.1
63	E-ESS	74.7	172.4	823	27	-25.3		0.2	0.6	0.8	1.9	2.4	0.7	0.3	1.7
66	E-ESS	75.9	174.3	862	26	-26.0	-270	0.2	0.6	0.9	1.7	2.3	0.9	0.3	1.5
DOC	C-sub ⁴														
13	LS	76.8	125.9	1208	25	-26.7									
14	LS	76.9	127.8	1216	26	-26.5		0.2	0.5	0.5	2.5	2.5	0.7	0.3	1.4
23	LS	76.2	129.3	1181	30	-27.0									
25	LS	76.0	130.7	1072	35	-27.2		0.9	0.4	0.4	1.4	1.2	0.4	0.3	0.7
27	LS	76.9	132.2	1072	30	-26.3									
28	LS	77.3	134.8	1711	36	-27.4	-171	0.9	0.4	0.3	1.2	0.9	0.3	0.4	0.6
29	LS	77.8	136.7	916	33	-26.6									
39	W-ESS	77.7	141.4	2170	46	-28.6									
40	W-ESS	77.6	145.8	2047	36	-27.6		0.6	0.4	0.4	1.7	1.4	0.4	0.4	0.7
41	W-ESS	77.0	148.3	1526	42	-28.0									
44	W-ESS	76.3	146.0	1259	32	-27.4	-188	0.3	0.6	0.7	1.2	1.3	0.5	0.5	1.1
46	W-ESS	76.4	149.9	1076	19	-25.7									
48	W-ESS	76.5	150.8	991	26	-26.4		0.2	0.6	0.7	2.1	2.1	0.6	0.6	1.4
50	W-ESS	75.8	158.5	973	20	-26.9		0.2	0.6	0.7	1.4	1.7	0.6	0.6	1.5
52	E-ESS	74.1	160.6	938	20	-25.3	-307	0.2	0.7	0.7	1.2	1.4	0.7	0.7	1.7
56	E-ESS	74.6	161.9	911	21	-25.4									
59	E-ESS	74.4	168.5	724	20	-24.9									
60	E-ESS	73.4	169.5	869	20	-25.4	-366	0.3	0.7	0.7	1.2	1.4	0.6	0.6	1.6
63	E-ESS	74.7	172.4	816	18	-24.6		0.2	0.7	0.7	1.1	1.4	0.6	0.6	1.8

Table 1. Composition of surface and near-bottom DOC samples collected in the outer Eurasian 854 855 Arctic Shelf.

¹DOC concentrations ($\mu g \cdot L^{-1}$) ²Lignin OC-normalized concentrations ($mg \cdot g^{-1}$ OC) ³swi, seawater intake samples (surface water samples at 8 m depth)

859 ⁴sub, samples obtained by submersible pump (near-bottom water samples, 5 m above bottom)

ID	Region	Lat	Long	POC ¹	C/N	$\delta^{13}C$	$\Delta^{14}C$	Lignin ²	S/V	C/V	Sd/S1	Vd/Vl	3.5Bd/V	Pn/P	P/V
POC-s	swi ³														
T-1	KS	79.8	67.9	232	5.1	-26.9		0.2	0.4	0.7	0.3	0.4	1.9	0.1	5.7
T-2	KS	81.7	75.3	353	6.0	-24.7		0.2	0.4	1.6	0.3	0.6	2.7	0.1	10.6
T-3	LS	81.3	109.4	85	6.5	-28.2		0.7	0.1	0.2	0.3	0.3	0.9	0.1	2.8
T-4	LS	81.0	112.9	89	5.9	-28.0		0.6	0.3	0.8	0.3	0.5	2.3	0.1	3.5
1	LS	78.9	125.2	70	4.9	-28.6	14	0.1	0.3	0.5	0.5	0.4	1.4	0.1	14.6
4	LS	77.8	126.7	138	6.3	-28.8	11	0.1	0.3	0.7	0.5	0.6	3.0	0.1	13.1
6	LS	77.1	127.4	100	5.4	-26.8									
13	LS	76.8	125.9	51	5.4	-27.5									
14	LS	76.9	127.8	54	4.9	-27.9		0.4	0.3	0.4	0.3	0.3	1.6	0.1	3.0
23	LS	76.2	129.3	40	5.1	-29.0		0.7	0.5	0.4	0.8	1.1	0.7	0.1	3.0
24	LS	75.6	129.6	67	5.4	-28.4		0.6	0.4	0.2	0.6	1.0	0.4	0.1	1.6
25	LS	76.0	130.7	117	5.4	-30.2	5	0.2	0.4	0.3	0.6	0.9	0.7	0.1	2.0
26	LS	76.5	132.0	78	5.0	-29.5									
27	LS	76.9	132.2	96	5.1	-29.6									
28	LS	77.3	134.8	45	5.2	-29.3	-75	0.2	0.4	0.4	0.4	0.6	0.8	0.1	5.2
29	LS	77.8	136.7	58	5.1	-28.7									
39	W-ESS	77.7	141.4	69	8.1	-28.0		0.2	0.5	0.8	0.2	0.4	1.8	0.1	5.6
40	W-ESS	77.6	145.8	85	8.6	-28.5		0.1	0.3	0.4	0.4	0.4	1.2	0.1	4.3
41	W-ESS	77.0	148.3	66	7.0	-27.8									
44	W-ESS	76.3	146.0	63	7.9	-28.8	64	0.1	0.5	0.3	0.2	0.3	1.2	0.0	4.8
45	W-ESS	76.4	148.1	61	8.4	-27.5									
46	W-ESS	76.4	149.9	27	6.7	-27.5									
48	W-ESS	76.5	150.8	52	7.0	-27.3		0.1	0.3	0.6	0.4	0.6	1.6	0.0	13.0
49	W-ESS	76.5	156.9	138	9.1	-25.9		0.05	0.3	0.7	0.4	0.6	1.8	0.0	21.7
50	W-ESS	75.8	158.5	96	9.3	-25.6	-102	0.05	0.3	0.6	0.4	0.5	1.3	0.0	14.1
52	E-ESS	74.1	160.6	120	8.1	-24.4	-94	0.1	0.6	1.0	0.3	0.5	2.4	0.1	22.2
56	E-ESS	74.6	161.9	151	9.2	-24.0									
57	E-ESS	74.4	163.7	160	10.9	-23.0		0.1	0.3	0.7	0.3	0.5	2.1	0.1	14.7
58	E-ESS	74.4	166.2	60	7.3	-23.7		0.1	0.4	0.8	0.4	0.7	2.5	0.04	19.3
59	E-ESS	74.4	168.5	286	10.2	-27.4									
60	E-ESS	73.4	169.5	109	5.9	-25.4	-90	0.05	0.4	0.9	0.5	0.4	1.6	0.04	31.5
61	E-ESS	74.1	170.9	231	8.0	-24.9	-69	0.03	0.7	2.4	0.3	0.4	3.2	0.1	38.9
63	E-ESS	74.7	172.4	67	7.4	-24.4		0.1	0.4	0.4	0.5	0.4	1.0	0.0	9.8
66	E-ESS	75.9	174.3	20	6.7	-27.4	-240	0.2	0.6	0.5	0.3	0.4	1.5	0.0	15.3
POC-s	sub^4														
13	LS	76.8	125.9	38	6.7	-29.0									
14	LS	76.9	127.8	47	6.0	-30.9		0.4	0.6	0.9	0.3	0.6	2.3	0.1	10.41
23	LS	76.2	129.3	99	11.7	-27.2									
25	LS	76.0	130.7	131	10.1	-26.8		1.1	0.8	0.2	0.3	0.5	0.8	0.1	2.12
27	LS	76.9	132.2	54	7.9	-26.3									
28	LS	77.3	134.8	48	6.6	-28.1	-365	0.9	0.5	0.9	0.4	0.7	3.0	0.1	4.03
29	LS	77.8	136.7	60	6.4	-25.9									
39	W-ESS	77.7	141.4	93	6.9	-26.7									
40	W-ESS	77.6	145.8	52	7.2	-26.4		0.7	0.3	0.3	0.3	0.5	1.9	0.1	4.60
41	W-ESS	77.0	148.3	173	11.3	-27.5									
44	W-ESS	76.3	146.0	95	6.5	-28.1	-193	0.4	0.4	0.5	0.4	0.4	2.3	0.1	10.52
46	W-ESS	76.4	149.9	171	5.8	-25.8									
48	W-ESS	76.5	150.8	78	6.1	-26.2		0.3	0.4	0.5	0.2	0.4	1.6	0.0	23.50
50	W-ESS	75.8	158.5	166	6.3	-28.4		0.3	0.3	0.3	0.3	0.4	1.1	0.0	13.10
52	E-ESS	74.1	160.6	72	6.4	-27.1	-258	0.3	0.2	0.5	0.4	0.5	2.0	0.0	16.58
56	E-ESS	74.6	161.9	116	8.9	-25.9									
59	E-ESS	74.4	168.5	124	7.6	-26.4									
60	E-ESS	73.4	169.5	356	7.4	-27.5	-185	0.2	0.7	0.9	0.3	0.4	1.8	0.1	18.7
63	E-ESS	74.7	172.4	168	7.0	-25.0		0.2	0.6	0.6	0.2	0.3	1.9	0.0	21.9

Table 2. Composition of surface and near-bottom POC samples collected in the outer Eurasian 860 861 Arctic Shelf.

¹POC concentrations ($\mu g \cdot L^{-1}$) ²Lignin OC-normalized concentrations ($mg \cdot g^{-1}$ OC) ³swi, seawater intake samples (surface water samples at 8 m depth)

864 865 ⁴sub, samples obtained by submersible pump (near-bottom water samples, 5 m above bottom)

ID	Region	Lat	Long	Depth ¹	SOC^2	C/N	$\delta^{13}C$	$\Delta^{14} C$	Lignin ³	S/V	C/V	Sd/S1	Vd/Vl	3.5Bd/V	Pn/P	P/V
SOC																
1^*	LS	78.9	125.2	-3120	1.0	7.1	-22.3	-418	0.6	0.7	0.3	0.6	0.9	0.6	0.1	1.6
4^*	LS	77.8	126.7	-2186	1.3	6.8	-22.5	-428	0.4	0.6	0.4	1.0	1.3	1.3	0.1	2.2
6	LS	77.1	127.4	-92	0.8	6.7	-23.2									
13	LS	76.8	125.9	-74	1.3	7.4	-24.1									
14	LS	76.9	127.8	-64	0.9	6.4	-24.3	-314	0.8	0.6	0.2	0.9	1.2	0.7	0.1	1.6
23	LS	76.2	129.3	-56	1.6	7.6	-25.0	-333	0.6	0.5	0.2	1.0	1.3	0.7	0.1	1.5
24	LS	75.6	129.6	-46	1.1	6.9	-24.8	-284	1.8	0.6	0.3	1.0	1.3	0.5	0.2	1.1
25	LS	76.0	130.7	-53	1.6	8.4	-25.5		2.4	0.6	0.3	0.8	1.0	0.3	0.2	0.9
26	LS	76.5	132.0	-52	1.2	7.9	-24.4	-441								
27	LS	76.9	132.2	-44	1.4	7.5	-24.2									
28	LS	77.3	134.8	-49	1.4	7.1	-23.8	-421	1.0	0.5	0.3	1.1	1.4	0.6	0.1	1.4
29	LS	77.8	136.7	-57	1.1	6.9	-23.4	-427								
39	W-ESS	77.7	141.4	-45	0.5	7.9	-24.0		0.8	0.5	0.2	0.9	1.2	0.6	0.1	1.4
40	W-ESS	77.6	145.8	-47	0.4	7.1	-23.7	-457	0.7	0.6	0.2	1.0	1.3	0.7	0.1	1.7
41	W-ESS	77.0	148.3	-40	0.3	7.7										
44	W-ESS	76.3	146.0	-43	1.2	7.9	-24.8	-484	1.4	0.6	0.4	0.7	1.0	0.5	0.2	1.4
45	W-ESS	76.4	148.1	-40	1.0	7.7	-24.4									
46	W-ESS	76.4	149.9	-40	1.1	7.2	-24.7	-463								
48	W-ESS	76.5	150.8	-40	1.4	7.4	-25.8	-345	0.3	0.5	0.3	1.2	1.2	1.3	0.1	3.8
49	W-ESS	76.5	156.9	-47	1.3	6.6	-23.6	-375	0.2	0.6	0.3	1.3	2.0	1.9	0.1	6.3
50	W-ESS	75.8	158.5	-44	1.2	6.7	-24.6	-523	0.5	0.5	0.2	0.9	1.2	0.7	0.1	2.0
52	E-ESS	74.1	160.6	-46	0.8	7.1	-23.9	-550	0.3	0.4	0.2	0.8	0.9	0.9	0.1	3.3
56	E-ESS	74.6	161.9	-48	1.1	7.3	-23.7									
57	E-ESS	74.4	163.7	-52	1.6	7.0	-24.2	-326	0.1	0.6	0.3	0.8	1.1	1.9	0.1	7.6
58	E-ESS	74.4	166.2	-54	1.7	7.5	-23.8	-296	0.1	0.7	0.5	0.8	1.4	2.4	0.1	13.3
59	E-ESS	74.4	168.5	-54	1.7	6.7	-23.5	-307								
60	E-ESS	73.4	169.5	-43	0.9	7.6	-24.0	-472	0.7	1.0	0.6	0.7	0.8	0.5	0.1	2.9
61	E-ESS	74.1	170.9	-51	1.8	7.0	-24.2	-318	0.3	0.9	0.8	0.9	1.0	0.9	0.1	5.3
63	E-ESS	74.7	172.4	-64	1.7	7.3	-22.7	-251	0.1	0.8	0.8	0.7	1.1	1.9	0.1	11.3
66*	E-ESS	75.9	174.3	-239	0.8	5.6	-21.0	-448	0.1	0.8	0.4	0.3	0.5	1.4	0.1	8.4

Table 3. Composition of surface sediment samples collected in the outer Eurasian Arctic Shelf. 866

867 868 869 ¹Water depth (m) ²Percentage of sedimentary organic carbon ³Lignin OC-normalized concentrations (mg·g⁻¹ OC)

870 *Cores sectioned on high resolution (0.5cm intervals)

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880 Figures

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Figure 1. Map of the Eurasian Arctic Shelf including the Kara, Laptev and East Siberian seas
(E-ESS, eastern East Siberian Sea; W-ESS, western East Siberian Sea). SWI samples,
seawater intake samples (surface water samples at 8 m depth); SUB samples, samples
obtained by submersible pump (near-bottom water samples, 5 m above bottom).



Figure 2. Longitudinal distribution of organic matter content in water and sediment samples. A) Organic carbon concentrations (μ g/L) in DOC (green circles) and POC (blue squares). The left y-axis is for DOC and the right y-axis for POC. B) Lignin concentrations (mg/g OC) in DOC_{SPE} (green circles), POC (blue squares) and SOC (red diamonds); swi, seawater intake samples (surface water samples at 8 m depth); sub, samples obtained by submersible pump (near-bottom water samples, 5 m above bottom). Dash line indicate the latitude of the Lena River mouth.

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Figure 3. Spatial distribution of organic carbon concentrations and δ^{13} C signatures in the DOC, POC (surface water samples at 8 m depth) and SOC pools of the ESAS.





Figure 4. Spatial distribution of lignin (mg/g OC) and Δ^{14} C signatures in the DOC_{SPE}, POC 912 (surface water samples at 8 m depth) and SOC pools of the ESAS. Sea ice (%) during the first 913 sampling day in the ESAS (15/7/2015).



Figure 5. Longitudinal distribution of Δ^{14} C signatures in DOC_{SPE} (green circles), POC (blue squares) and SOC (red diamonds); swi, sea water intake samples (surface water samples at 8 m depth); sub, samples obtained by submersible pump (near-bottom water samples, 5 m above bottom). Dash line indicate the latitude of the Lena River mouth.



Figure 6. Relationships between organic carbon and Δ^{14} C and δ^{13} C signatures in the DOC (green circles), POC (blue squares) and SOC (red diamonds).



Figure 7. Correlations between lignin concentrations (mg/g OC) and organic carbon, δ^{13} C and Δ^{14} C signatures in DOC (green circles), POC (blue squares) and SOC (red diamonds).



998Figure 8. Lignin-phenols ratios in DOC_{SPE} (green circles), POC (blue squares) and SOC (red999diamonds). A) Classical source plot of syringyl/vanillyl (S/V) vs. cinnamyl/vannillyl (C/V).1000Typical ranges for woody and non-woody tissues of both angiosperm and gymnosperm1001vegetation are indicated as boxes in the graph (Goñi et al., 2000). B) The acid/aldehyde ratios1002of syringyl (Sd/Sl) vs. vanillyl (Vd/Vl).



Figure 9. Lignin proxies of Terr-OC and relative degradation state of DOC_{SPE} (green circles), 1016 POC (blue squares) and SOC (red diamonds). A) Ratios between 3,5-dihydroxybenzoic acid 1017 and vanillyl phenols (3,5-Bd/V). B) Ratios between *p*-coumaric acid and ferulic acid 1018 (pCd/Fd).