Sources and characteristics of terrestrial carbon in Holocene-scale sediments of the East Siberian Sea

Kirs Keskita1, Tommaso Tesi1,3,4, Lisa Bröder1,3, August Andersson1,3, Christof Pearce2,3, Martin Sköld5, Igor P. Semiletov6,7,8, Oleg V. Dudarev7,8 and Örjan Gustafsson1,3,*

1Department of Environmental Science and Analytical Chemistry, Stockholm University, Stockholm, SE 10691, Sweden
2Department of Geological Sciences, Stockholm University, Stockholm, SE 10691, Sweden
3Bolin Centre for Climate Research, Stockholm University, Stockholm, SE 10691, Sweden
4CNR-National Research Council of Italy, ISMAR-Marine Science Institute, Bologna, IT 40129, Italy
5Department of Mathematics, Stockholm University, Stockholm, SE 10691, Sweden
6International Arctic Research Center, University Alaska Fairbanks, Fairbanks, AK 99775, USA
7Pacific Oceanological Institute, Russian Academy of Sciences, Vladivostok, RU 690041, Russia
8Tomsk National Research Polytechnical University, Tomsk, RU 634050, Russia

*Correspondence to: Örjan Gustafsson (orjan.gustafsson@aces.su.se)

Abstract. Thawing of permafrost carbon (PF-C) due to climate warming can remobilise considerable amounts of terrestrial carbon from its long term storage to the marine environment. PF-C can be then buried in sediments or remineralised to CO2 with implications for the carbon-climate feedback. Studying historical sediment records during past natural climate changes can help to understand the response of permafrost to current climate warming. In this study two sediment cores collected from the East Siberian Sea were used to study terrestrial organic carbon sources, composition and degradation during the past ~9,500 cal yrs BP. The CuO-derived lignin and cutin products combined with δ13C suggest that there was a higher input of terrestrial organic carbon to the East Siberian Sea between ~9,500 and 8,200 cal yrs BP than in all later periods. This high input was likely caused by marine transgression and permafrost destabilisation in the early Holocene climatic optimum. Based on source apportionment modelling using dual-carbon isotope (Δ14C, δ13C) data, coastal erosion releasing old Pleistocene permafrost carbon was identified as a significant source of organic matter translocated to the East Siberian Sea during the Holocene.
1 Introduction

The amount of organic carbon (OC) stored in the northern circumpolar permafrost (PF) amounts to ~1300 Pg of which ~800 Pg OC is permafrost, seasonally thawing active-layer permafrost or taliks (Hugelius et al. 2014). Northern circumpolar soils thereby hold roughly half of the global soil OC pool (Tamocai et al., 2009). Modelled future climate scenarios predict increased amplified warming in the Arctic for the coming 100 years (IPCC, 2013). This will further destabilise permafrost, leading to increased delivery of terrestrial OC to the Arctic Ocean. The potential decomposition of this relict permafrost carbon (PF-C) and its subsequent release to the atmosphere as CO₂ or CH₄ constitutes a positive feedback to global warming (IPCC, 2013; Koven et al., 2011; Schuur et al., 2015; Shakhova et al., 2013, 2015, 2009; Vonk and Gustafsson, 2013). Considering the size of the Arctic PF-C pool it is important to better understand the dynamics and extent of its vulnerability to remobilisation in response to climate warming.

Many recent studies have focused on current carbon cycling in the Arctic land-ocean continuum (Anderson et al., 2009, 2011; Bröder et al., 2016a; Goñi et al., 2013; Goñi et al., 2000; Karlsson et al., 2015; Semiletov et al., 2016, 2011, 2012; Shakhova et al., 2010; Tesi et al., 2016a; Vonk et al., 2010; Winterfeld et al., 2015b) with possible linkages to climate change. Constraining how this system responded to earlier climate warming may help us to better predict the future response of PF-C and its climate couplings. The last glacial-interglacial transition constituted a major climate rearrangement on Earth. The increase in mean temperature coupled with sea level rise is thought to have profoundly destabilised PF-C and further released CO₂ to the atmosphere (Ciais et al. 2013; Crichton et al. 2016; Tesi et al. 2016a). Several studies have suggested that there was a warming-coupled translocation of terrestrial carbon during the climate warming that ended the last glacial period (e.g., Bauch et al. 2001; Ciais et al. 2013; Mueller-Lupp et al. 2000; Tesi et al. 2016a) similar to what is predicted to happen as a consequence of the anthropogenic climate change (Barnhart et al. 2014; Vonk and Gustafsson 2013).

Many of the previous Holocene timescale studies in the East Siberian Arctic Shelf (ESAS) have focused on the Laptev Sea (e.g., Bauch et al., 2001b; Mueller-Lupp et al., 2000; Tesi et al., 2016a). This study focuses on the East Siberian Sea (ESS) which has not yet been extensively studied in this aspect. The ESS receives terrestrial OC by coastal erosion, fluvial inflow and possibly sea bed erosion (Karlsson et al., 2016; Semiletov et al., 2005; Stein and Macdonald, 2004; Tesi et al., 2014, 2016b; Vonk et al., 2010). The coast of the ESS is dominated by carbon-rich Ice Complex Deposits (ICD) consisting of old Pleistocene material (Schirrmieister et al. 2011; Semiletov 1999a, 1999b; Vonk et al. 2012). These large ICD bluffs are vulnerable to coastal erosion (Semiletov et al., 2013; Stein and Macdonald 2004; Schirrmieister et al. 2011; Vonk et al. 2012). Coastal erosion can be further intensified with warming enhanced processes like loss of sea ice cover, increasing frequency of storms, degradation of ice-bonded coasts and sea level rise (Barnhart et al., 2014; Jones et al., 2009; Stein and Macdonald, 2004). The largest rivers directly emptying into the ESS are Indigirka and Kolyma with suspended matter discharge of 11.1 x 10¹² g yr⁻¹ and 123±19 x 10⁹ g yr⁻¹ (Gordeev, 2006; McClelland et al., 2016, respectively), with an input also from the Lena River. The Lena River drains into the Laptev Sea but its exported terrestrial OC is also transferred to the ESS via the Siberian Coastal Current (e.g., Alling et al., 2012; Sánchez-García et al., 2011). However, studies by Vonk et al. (2010, 2012) suggest that the contribution of ICD-PF erosion to the ESS sediment OC dominates over river discharge (ranging from 36 to 76 % in comparison to 5–35 %, respectively). Similar observation have been made in the Laptev Sea by Semiletov et al., (2005, 2011, 2012) and Vonk et al., (2012, 2014) concluding that the effect of the Lena River input is overall smaller than that from the coastal erosion.
In this study we investigate land-to-ocean transfer and fate of PF-C from the last post-glacial eustatic sea level rise until the present day. Our main objectives are to determine the sources and remobilisation fluxes of terrestrial OC as well as the composition and degradation status of the OC that was buried in ESS sediments during the Holocene. We characterise the OC composition by quantifying lignin phenols, cutin acids and other compounds yielded upon CuO oxidation to constrain the sources and degradation status of PF-C as well as the contribution of marine OC. Furthermore, we use a mixing model based on the isotopic composition ($\Delta^{14}C$, $\delta^{13}C$) of the deposited OC to quantify the contribution of three different sources: topsoil-PF from active-layer deepening, ICD-PF and marine plankton. Additionally, we study how OC deposition fluxes have changed over time in response to the sea level rise and Holocene warming.
2 Materials and methods

2.1 Background and study area

The East Siberian Sea (ESS) is located off the northeast Siberian coast between the Laptev Sea and the Chukchi Sea (Fig. 1). The ESS is one of the largest shelf seas (987,000 km²) in the Arctic Ocean as well as one of the shallowest (mean depth 52 m) (Jakobsson, 2002).

Thermokarst landscapes (i.e. thawing ice-rich permafrost) cover ~20 % (3.6 x 10⁶ km²) of the northern circumpolar permafrost region (Olefeldt et al., 2016). Ice Complex Deposit and thermokarst landscapes cover 2,400 km of the ESS coastline (Grigorieva 2003). The modern average rate of coastal retreat in the ESS and the adjacent Laptev Sea is 1–10 m yr⁻¹ (Grigoriev 2010), though locally, even higher retreat rates (up to 24 and 30 m yr⁻¹) have been reported in the most actively eroding parts (Kanevskiy et al., 2016; Romanovskii et al., 2004).

The coastal erosion rates have increased in the Arctic in recent decades (Barnhart et al., 2014; Günther et al., 2015; Jones et al., 2009). According to recent studies (e.g., Bröder et al., 2016a; Semiletov et al., 2013; Tesi et al. 2016b; Vonk et al. 2012) a large fraction of the remobilised PF-C is degraded during cross-shelf transport and released back to the contemporary carbon cycle. To better predict the consequences of the permafrost thaw, it is important to understand both the amount of remobilised organic carbon as well as its fate.

The shelf of the East Siberian Sea contains terrestrial permafrost forming during the sea level low of last glacial maximum (Jakobsson et al. 2014). During the Pleistocene-Holocene transition the ESAS was flooded when the sea level rose rapidly (Lambeck et al., 2014; Mueller-Lupp et al., 2000). This global marine transgression started ~20,000 cal yrs BP (Lambeck et al., 2014) and flooded the ESAS between ~11,000 to ~7,000 cal yrs BP (Bauch et al. 2001a; Mueller-Lupp 2000). The sampling site of the sediment core investigated in this study was flooded around 11,000 cal yrs BP (Lambeck et al., 2014). Post-glacial sea level rise with warming and wetting of the climate caused a major relocation of permafrost carbon from land to the Arctic Ocean (Bauch et al. 2001; Tesi et al. 2016a). Today the period with sea ice in the ESS is on average 3 months per year which is one of the reasons why the area remains fairly unstudied (Stein and Macdonald, 2004; Vetrov and Romankevich, 2004).

2.2 Sampling

A gravity core (called GC58) was collected in the East Siberian Sea at 54 m water depth as a part of the international SWERUS-C3 research expedition on i/b Oden in July–August 2014. The coring site (Leg 1, station 58, 74.4387° N, 166.0467° E) is located ~500 km from the modern shoreline (Fig. 1). An additional sediment core was collected at the same site (MUC58) using a sediment multicorer (Oktopus GmbH, Germany), which is specifically designed to preserve the sediment-water interface. The total length of GC58 was 78 cm while MUC58 was 32 cm long. The GC58 core was split in half during the expedition and kept refrigerated (~4°C). In the laboratory at Stockholm University, one half was subsampled at 1 cm intervals and kept frozen at -18°C. The multicore was sliced during the expedition at 1 cm intervals and then immediately frozen (-18°C). Prior to analyses, the samples were freeze-dried at the Department of Environmental Science and Analytical Chemistry, Stockholm University, Sweden.

2.3 ³²⁳⁰Pb dating

Radiogenic ³²³⁰Pb was analysed with a gamma-ray spectrometer (GRS) at the Department of Geology of the Swedish Museum of Natural History in Stockholm, Sweden. The GRS determines the decay energy of radioisotopes in counts per second by measuring gamma emission of the sample at a known energy level.
Prior to the GRS analysis, a subsample of approximately 10 g was homogenised and placed in a plastic container for at least three weeks to reach secular equilibrium between the radioisotopes of lead and radium ($^{210}$Pb and $^{226}$Ra, respectively). The samples were analysed for $^{210}$Pb (46.51 keV), $^{226}$Ra (186.05 keV) and $^{137}$Cs (661.66 keV) on an EG&G ORTEC® co-axial low energy photon spectrometer containing a High-Purity Germanium detector. The counting period for each sample lasted from 1–3 days depending on the amount of $^{210}$Pb in the sample. An externally calibrated U-series standard (pitchblende, Stackebo, Sweden) was used to determine the relative efficiency of the gamma detector system. For each sample a minimum of 350 counts was acquired. A blank (empty container) sample was measured to correct for the background activity. The original method is described in detail by Elmquist et al., (2007).

Two different models were used for the $^{210}$Pb dating: CRS (constant rate of supply) model which assumes a constant rate of supply of excess $^{210}$Pb fallout, and CIC (constant initial concentration) model which assumes constant initial concentration of excess $^{210}$Pb (Appleby and Oldfield, 1977).

2.4 Bayesian modelling of $^{14}$C ages for the chronology

For the age-depth model construction, molluscs retrieved from GC58 were analysed for their radiocarbon ($^{14}$C) content at the US-NSF National Ocean Sciences Accelerator Mass Spectrometry (NOSAMS) Facility at the Woods Hole Oceanographic Institution (WHOI), MA, USA. The analysis followed their standard procedures (Pearson et al., 1998) (Table 1).

To account for natural differences in the amount of $^{14}$C in the atmosphere as well as differences between the marine environment and the atmosphere (e.g., Stuiver and Braziunas 1993), all $^{14}$C data were calibrated with the Marine13 calibration curve. The offset in the local reservoir age was taken into account by using a $\Delta R$ of 50±100 years. Since there are no $\Delta R$ values for the ESS in the literature, this $\Delta R$ value was taken from a study in the Laptev Sea (Bauch et al. 2001a). The radiocarbon dates are reported in conventional radiocarbon ages (cal yrs BP) (Stuiver and Polach 1977).

The age model of the core was built with the OxCal v4.2 program based on the radiocarbon dated molluscs and a depositional model ($P_{\text{sequence}}, k = 0.5$) (Bronk Ramsey 2008; Bronk Ramsey and Lee 2013). Also, the base of the adjacent multicore dated with $^{210}$Pb was used in the model. The $^{210}$Pb date used was an average age (50 yrs BP) from the two $^{210}$Pb dating models (CRS, CIC) for the bottom layer (12.5 cm) of the multicore (Supplementary Table S3). The age model of GC58 was constructed with a Bayesian statistics approach using the reservoir age ($\Delta R$) and the depth as a prior model and measured radiocarbon dates as likelihoods. The posterior probability densities were acquired with a Markov Chain Monte Carlo procedure which calculates possible distributions in order to date each sediment layer using the given prior model and likelihoods (Bronk Ramsey 2008).

Sampling with a heavy gravity corer often disturbs the sediment-water interface and thereby causes losses of the surface sediments. The organic carbon (OC) content of GC58 was therefore compared to the OC content of the adjacent MUC58 to identify for the possible loss. According to the comparison, the top 3 cm were likely lost in GC58 (Supplementary Fig. S1), and thus corrected for.
2.5 Alkaline CuO oxidation

Microwave assisted alkaline CuO oxidation was carried out using the method by Goñi and Montgomery (2000). Each homogenised subsample of around 300 mg was mixed with 300 mg of cupric oxide (CuO) and 50 mg of ammonium iron (II) sulphate hexahydrate ((NH₄)₂Fe(SO₄)₂·6H₂O). After thorough mixing, nitrogen-purged 2M NaOH was added to each sample. Alkaline oxidation was performed with an UltraWave Milestone 215 microwave digestion system at 150°C for 90 min.

A known amount of internal recovery standards (ethyl-vanillin, cinnamic acid) was added to the CuO reaction products and then acidified to pH 1 with concentrated HCl (35 %). The CuO reaction products were repeatedly extracted using ethyl acetate (EtOAc). Anhydrous sodium sulphate (NaSO₄) was added to remove the remaining water. The extracts were dried in a CentriVap (Christ RVC 2-25) at 60°C, re-dissolved in pyridine and stored in a freezer (-18°C) until further analysis.

Finally, the samples were analysed with a gas chromatograph mass spectrometer (GC-MS, Agilent 7820A) using a DB5-MS capillary column (60 m x 250 μm, 0.25 μm stationary phase thickness, Agilent J&W) at an initial temperature of 60°C, followed by a ramp of 5°C/min until reaching 300°C. Prior to the GC-MS analysis, the extracts were derivatised with bis-trimethylsilyl trifluoroacetamide (BSTFA) + 1 % trimethylchlorosilane (TMCS) to silylate exchangeable hydrogens. The quantification of the samples was based on the comparison of the key ions to commercially available standards. Concentrations of CuO oxidation products were normalised to the organic carbon content of the sample and are reported as mg g⁻¹ OC.

2.6 Bulk organic carbon and stable carbon isotope analyses

For the total organic carbon content (TOC), the total nitrogen content (TN) and the stable carbon isotope analysis (δ¹³C) of TOC, subsamples of 10–15 mg were homogenised and placed in silver capsules, acidified with 1.5M HCl to remove carbonates and then dried at 60°C. The TOC, TN and δ¹³C-TOC were quantified with an elemental analyser Carlo Erba NC2500 connected via a split interface to a Finnigan MAT Delta V mass spectrometer at the Stable Isotope Laboratory of the Department of Geological Sciences at Stockholm University.

For radiocarbon (¹⁴C) analysis of the bulk organic carbon, subsamples of sediment were acidified with 1.5M HCl and sent to NOSAMS. To account for the time between the deposition and the measurement, the ¹⁴C dates were calibrated with the Eq. (1) using the age data derived from the age model. The bulk radiocarbon data are reported as Δ¹⁴C (Stuiver and Polach 1977).

\[
\Delta^{14}C = (Fm \times e^{\lambda(1950-Ye)} - 1) \times 1000
\]

where Fm is the Fraction Modern, \(\lambda\) is 1/mean life of radiocarbon= 1/8267 and Ye is the year of collection derived from the age model (Stuiver and Polach, 1977).

2.7 Source apportionment

The carbon isotope fingerprint of OC (Δ¹³C, δ¹³C) can be used to quantitatively diagnose the relative contribution of topsoil-PF, ICD-PF and marine OC assuming isotopic mass balance (e.g., Vonk et al., 2012). In other words, the carbon isotopic signatures may help to understand whether the OC comes from coastal erosion as a result of the post-glacial warming and sea level rise, active-layer deepening of permafrost carbon in the watershed (as a response to the post-glacial warming) or sedimentation of marine phytoplankton. These different
sources have a natural variability in their isotopic composition (end-members). This variability needs to be taken into account to correctly estimate the relative source contributions and the associated uncertainties (e.g., Andersson, 2011). In previous studies a Bayesian Markov Chain Monte Carlo (MCMC) driven approach has been used to effectively estimate the relative source contributions for individual data points (Andersson et al., 2015; Tesi et al., 2016a). Here, we expand this approach to include the time-dependence of the down-core isotopic signatures, taking an advantage of the relatively small variability of the $7\delta^{13}\text{C}$ data points, whilst also using the 10 $\Delta^{13}\text{C}$ points. The time-dependence of different proportions was taken into account by following the approach of Parnell et al. (2012). The method is described in detail in the Supplementary Methods.

The end-member values for the three source classes were taken from the literature (Bröder et al., 2016b; Tesi et al., 2016a) topsoil-PF ($\Delta^{13}\text{C}=-232±147 \text{‰}$, $\delta^{13}\text{C}=-26.95±1.17 \text{‰}$, mean ± standard deviation), representing thaw of the active-layer of permafrost; marine OC ($\Delta^{13}\text{C}=-50±12 \text{‰}$, $\delta^{13}\text{C}=-20.97±2.56 \text{‰}$), resulting from primary production of phytoplankton; and ICD-PF ($\Delta^{13}\text{C}=-940±31 \text{‰}$, $\delta^{13}\text{C}=-26.3±0.63 \text{‰}$), resembling the old Pleistocene material from coastal erosion. The end-member value for ICD-PF was corrected with Eq. (1) to account for the age of the deposition.

2.8 Grain size analysis

Prior to the grain size analysis subsamples of sieved (500 µm) sediments from GC58 were homogenised. The grain size analysis was done with a Malvern Mastersizer 3000 laser diffraction particle size analyser, which can measure particles between 10 nm and 3.5 mm. Sodium hexametaphosphate (10 %) was used to disaggregate the particles suspended in deionised water. To further aid the disaggregation, all samples were exposed to ultrasound for 60 s and allowed to disperse in continuous flow for 3 min in total (including 60 s of ultrasonication) prior to the measurements. To control the concentration of the sample in the flow during the measurements, the obscurity was kept between 5–15 %. High sample obscurity (i.e. high concentration) would cause multiple light scatterings, thus distorting the results. Each sample was analysed in five replicates. The measurements were carried out at the Department of Geological Sciences at Stockholm University, Sweden.
3 Results and Discussion

3.1 Age chronology of the core

The deepest part of the sediment core GC58 dates back ~9,500 cal yrs BP i.e. to the early Holocene. The age-depth model shows an evident hiatus in the middle of the core between 39.5 cm and 40.5 cm resulting in an age gap of ~6,500 years (~8,200–1,700 cal yrs BP) (Fig. 2). In addition, there is a shorter gap in the chronology between ~9,300 and ~8,500 cal yrs BP. In studies from the adjacent Laptev Sea such age discrepancies have not been observed (Bauch et al. 2001a; Bauch et al. 2001b; Tesi et al. 2016a). It therefore seems likely that there has been a local event causing the removal of sediment layers. There might not have been accumulation during those periods, or the age gap could be a condensed unit of sediment. Although any actual sediment transport processes giving rise to such a putative total halt in sedimentation is rather elusive and unlikely. Since the whole East Siberian Arctic Shelf (ESAS) is a very shallow shelf where sea ice is formed (Conlan et al., 1998; Jakobsson, 2002), a likely explanation for an age gap is ice scouring as observed in the Laptev Sea (Ananyev et al., 2016). An ice scouring event could have formed a gouge at the sea bottom that later was refilled with sediment (Barnes et al. 1984).

The accumulation rates of GC58 obtained from the $^{14}$C measurements vary between 0.2 and 1.4 mm yr$^{-1}$ (17.0–138.9 cm kyr$^{-1}$) and mass accumulation rates (MAR) spanned 0.02–0.1 g cm$^{-2}$ yr$^{-1}$. Bauch et al. (2001a) have reported similar sedimentation rates (0.1–2.6 mm yr$^{-1}$) from the outer shelf of the Laptev Sea around the same time period. The linear sedimentation rate for the adjacent sediment core MUC58 derived from $^{210}$Pb dating is 1.3 mm yr$^{-1}$ and an average MAR 0.03 g cm$^{-2}$ yr$^{-1}$. Similar accumulation rates with $^{210}$Pb dated sediment cores have been reported in other studies from the East Siberian Sea: 1.1–1.6 mm yr$^{-1}$ (Vonk et al. 2012) and 1.4–1.5 mm yr$^{-1}$ (Bröder et al., 2016b). The slight difference in accumulation rates using $^{210}$Pb chronology compared to $^{14}$C may be due to active biological mixing giving higher accumulation rates for the shorter time scale of more surficial sediments (Baskaran et al. 2016; Boudreau 1994).

3.2 Sediment grain size, stable carbon isotopes and biomarker composition of organic matter

Grain size can be used to describe the depositional environment. The sediment core GC58 consists mostly of clay and silt, with a fraction of sand (Supplementary Fig. S2). The higher sand content that is observed at ~8,500 cal yrs BP may reflect a higher-energy depositional regime likely due to proceeding marine transgression and energetic coastal dynamics. Bauch et al. (2001a) have reported a shift from sandy silt to clayey silt around 7,400 cal yrs BP from a sediment core collected in the eastern Laptev Sea. They attribute this change to the end of the sea level rise and the establishing of more stable conditions. The GC58 sediment core has a hiatus at that time period but has a similar clayey silt composition at the top part of the core (~1,700 cal yrs BP until today). This may indicate comparably similar stable conditions in the East Siberian Sea in the last 1,700 cal yrs BP.

The total organic carbon (TOC) concentrations in GC58 vary from 0.5 to 1.1% (Supplementary Table S1) with the highest TOC content in the surface sediments. These data agree with average TOC contents reported for the East Siberian Sea (Semiletov et al., 2005; Stein and Macdonald, 2004; Vetrov and Romankevich, 2004; Vonk et al., 2012). The OC fluxes for GC58 calculated with the $^{14}$C age-model (covering ~9,500 cal yrs BP) range between 1.2 and 10.9 g m$^{-2}$ yr$^{-1}$ (Fig. 3a). The OC fluxes for MUC58 calculated with the $^{210}$Pb chronology (covering the most recent ~100 yrs) are similar and vary from 0.4 to 6.1 g m$^{-2}$ yr$^{-1}$ (Supplementary Table S2). The OC fluxes show an increasing trend from the bottom of the core toward the top in both cores. A similar trend has been reported by (Bröder et al., 2016b) from the East Siberian Sea using two $^{210}$Pb-dated sediment cores. For GC58, the high OC flux at the very top of the core is likely related to the
merging of the two dating systems ($^{14}$C and $^{210}$Pb), which causes a higher sediment accumulation rate at the top of the core and thus higher fluxes.

Lignin phenols and cutin acids are useful proxies for tracing carbon of terrestrial origin because both compounds are solely biosynthesised in terrestrial plants. Lignin is an essential component in cell walls of vascular plants (Higuchi, 1971), while cutin is a lipid polyester, which forms a protective wax layer on epidermal cells of leaves and needles with other lipids (e.g., Kunst and Samuels 2003). These compounds have been widely used in recent studies of terrestrial OC in the Arctic (e.g., Amon et al., 2012; Bröder et al., 2016b; Gotli et al., 2013; Tesi et al., 2014). Both lignin and cutin fluxes show a similar trend with the highest fluxes at the bottom of the core (~9,500 cal yrs BP) indicating a high proportion of terrestrial organic matter (Fig. 3b). The large variability in the fluxes between ~9,500 and ~8,200 cal yrs BP compared to the latest ~1,700 cal yrs BP suggests that the system was more dynamic at that time. The rapid decrease in both lignin and cutin fluxes proposes a change from terrestrially dominated to marine dominated input at ~8,400 cal yrs BP in this part of the East Siberian Sea. Bauch et al. (2001b) suggested a similar regime shift from terrestrial to marine in the Laptev Sea between ~8,900 and ~8,400 cal yrs BP based on the occurrence of bivalves and benthic foraminiferal species. The same process affecting OC fluxes is likely causing also higher lignin and cutin fluxes at the top of GC58. The overall decrease in lignin and cutin fluxes as well as concentrations (Supplementary Table S3) in time is likely due to increasing hydrodynamic sorting and degradation during transport as transport times from the coast became longer because of the marine transgression (Fig. 3a). Bröder et al., (2016a) have observed a similar strong decrease in the amount of terrestrial organic carbon depositions with increasing distance from the coast in the Laptev Sea. A recent study by Tesi et al. (2016b) shows that the largest particles, rich in lignin (i.e. plant debris), tend to be preferentially buried close to the shore and with cross-shelf transport of lignin occurring overwhelmingly bound to fine particles (with low settling velocities) (i.e. of the total lignin deposited to the marine environment only a fraction, ~4–5 %, travels across the shelf).

Other useful indicators of the marine input in organic matter are CuO oxidation derived low-molecular weight fatty acids (LMW-FA). They are mainly found in phytoplankton but also in other organisms such as bacteria and algae (Gotli and Hedges, 1995). Especially C16FA:1 together with C14FA and C16FA serve as proxies for marine OC as they are highly abundant in marine sediments and very low in concentrations in ICD-PF and topsoil-PF (Gotli and Hedges, 1995; Tesi et al., 2014). The highest fluxes of LMW-FA are observed for the very top of the core (Fig. 3c), indicating a larger proportion of marine OC. The values decrease rapidly down-core as marine FA are readily degraded (e.g., Bröder et al., 2016b; Canuel and Martens, 1996). This trend may also be influenced by the change in input from terrestrial to marine dominated sources.

The stable isotopic composition of bulk OC ($\delta^{13}$C) may be used to distinguish between marine and terrestrial organic matter (Fry and Sherr, 1984). The $\delta^{13}$C values for C3-photosynthesised terrestrial carbon are between -23 to -30 ‰, whereas marine carbon has a less depleted $\delta^{13}$C signature between -18 ‰ and 24 ‰ (e.g., Fry and Sherr, 1984). However, these end-member values may differ depending on the region, especially in the Arctic where open water and sea ice phytoplankton exhibits different isotopic fingerprints (Kohlbach et al., 2016). The $\delta^{13}$C values for GC58 range from -23 to -25 ‰ (Fig. 3d) with the most depleted values (i.e. most terrestrial) between ~9,500 and ~8,200 cal yrs BP, and the least depleted values (i.e. most marine) from ~1,700 cal yrs BP until the modern time. Mueller-Lupp et al. (2000 and references within) have argued that $\delta^{13}$C values in sediments of the Arctic Ocean can have a terrestrial overprint in $\delta^{13}$C composition caused by the rapid degradation of planktonic organic matter i.e. the amount of marine organic matter of the total organic matter
pool in the Arctic is relatively low. Yet, the gradual change in δ\(^{13}\)C indicates that the contribution of marine organic matter is greater at the top of the core where the δ\(^{13}\)C values are less depleted.

It is notable that the values for all the different parameters shown in Fig. 3 on both sides of the age gap (between ~8,200 and ~1,700 cal yrs BP) are near-continuous in spite of the ~6,500 year hiatus (except for the bulk Δ\(^{14}\)C OC values). This could be explained by bioturbation, mixing the older part of the core with the newer deposits, thus resulting in an apparent continuity in property values across the hiatus. The Δ\(^{14}\)C values suggest that there was more δ\(^{13}\)C depleted material deposited ~1,600 cal yrs BP ago, causing a drop in the Δ\(^{13}\)C values. Though more likely, as the Δ\(^{14}\)C values are dependent on time, any uncertainty in the age model would have an effect on the Δ\(^{13}\)C values.

3.3 Degradation status of terrestrial organic matter

Lignin phenols provide insight to the degradation status of the deposited terrestrial organic matter. The acid-to-aldehyde ratios of lignin phenols, syringic acid to syringaldehyde (Sd/Sl) and vanillic acid to vanillin (Vd/Vl), have been used to study degradation of lignin (e.g., Benner and Opsahl, 1995; Hedges et al., 1988). As acids are more abundant in relation to aldehydes in degraded lignin, higher ratios mean more degraded lignin (Goñi et al., 1993). Both Sd/Sl and Vd/Vl ratios show great variability throughout the core (Fig. 4a), especially for the top part of the core. The variability at the core top may reflect the analytical uncertainty caused by very low lignin concentrations. In addition, Goñi et al. (2000) and Tesi et al. (2014) have argued that the acid to aldehyde ratios of lignin phenols might not serve as good degradation proxy for Arctic Ocean sediments as the material entering the marine environment might have experienced degradation prior to entering the marine system.

The ratio of 3,5-dihydrobenzoic acid to vanillyl phenols (3,5-Bd/V) is another proxy used to constrain the degradation status of terrestrial organic matter in sediments (e.g., Hedges et al. 1988; Tesi et al. 2014; Tesi et al. 2016a). Specifically, this proxy is used to distinguish diagenetically-altered mineral soil OC from relatively fresh vascular plant debris (Farella et al. 2001; Louchouarn et al. 1999; Prahl et al. 1994). The only source of 3,5-Bd in the marine environment is from brown algae which are not common in the study area (Goñi and Hedges, 1995; Tesi et al., 2014). The low 3,5-Bd/V ratio at the bottom of the core (~9,500–8,200 cal yrs BP) implies that the organic matter that was deposited in that period was relatively undegraded (Fig. 4b). The extent of degradation gradually increases toward the top of the core. However, hydrodynamic sorting may affect the degradation values as the largest particles of fresh vascular plant debris are likely buried close to the coast (Tesi et al., 2016b). The input of organic matter was higher before ~8,200 cal yrs BP, presumably due to coastal erosion caused by the marine transgression. When sediments are quickly buried they can serve as a more effective sink for terrestrial organic matter (Hilton et al., 2015). As the material is less degraded and the sedimentation rates are high in GC58 between ~9,500 and ~8,200 cal yrs BP, the input of organic matter was likely high causing it to be quickly buried. Similar high input of terrestrial material has been observed in the Laptev Sea ~11,000 cal yrs BP (Tesi et al. 2016a).

The location of the study site is currently ~500 km offshore so transport time and thereby the oxygen exposure time of the organic matter in the benthic compartment is now longer than in the earlier phase of the Holocene. The longer distance from the coast allows more time for organic matter to degrade before burial (Bröder et al., 2016a). Hartnett et al. (1998) have also shown that the burial efficiency of organic carbon decreases as a function of oxygen exposure time. The same trend can be seen in the fraction remaining lignin (\(\text{F}_{\text{lignin/terrOC}}\)) i.e. the amount of lignin as a ratio of the observed and expected (assuming conservative mixing i.e. no degradation) concentrations of lignin and terrestrial OC (terrOC) (see Supplementary Methods for details). In
GC58 the flignin/terrOC decreases down-core likely as a result of the proceeding marine transgression (Supplementary Fig. S3). This trend suggests that with longer transport time the lignin degradation is more extensive due to the protracted oxygen exposure time and hydrodynamic sorting (Keil et al., 2004; Tesi et al., 2016a). We estimated this lateral transport time to be ~1.4 kyr longer at modern times than at the beginning of the Holocene for the station GC58 (Supplementary Fig. S4). To model the lateral transport times, we used the flignin/terrOC with individual degradation rates for terrOC and lignin (Bröder et al. 2017, submitted) (see Supplementary Methods).

### 3.4 Dual-isotope based source apportionment of OC

The source apportionment results show that most of the organic matter originates from coastal erosion since ICD-PF material is the largest fraction (41–91 %) throughout the core (Fig. 5). Earlier studies demonstrated that the decay of fresh marine organic matter is more rapid compared to degradation of terrestrial organic matter (Karlsson et al., 2011, 2015; Salvadó et al., 2016; Vonk et al., 2010). This may lead to selective preservation of terrestrial organic matter in the sediments of the East Siberian Arctic Shelf (Karlsson et al. 2011, 2015; Vonk et al. 2010). The proportion of old terrestrial organic matter might also be greater in Arctic sediments due to generally low primary production in the area (Stein and Macdonald, 2004). The contribution of topsoil-PF is fairly low throughout the core (3–23 %). This may be due to the location of GC58 between the two major rivers (Kolyma and Indigirka) resulting in relatively low amounts of fluvial inflow depositing topsoil permafrost. To further interpret our results within a larger context of PF-C destabilisation during post-glacial warming, we compared our results with another transgressive deposit collected in the Laptev Sea (PC23, Fig. 1, Tesi et al. 2016a). For the Laptev Sea (PC23), there was a predominant influence of watershed-sourced material via river discharge during the onset of the Holocene, followed by a similar contributions of marine OC and ICD-PF fractions (both sources varying between 31 and 56 %) from ~8,300 cal yrs BP to present. For the East Siberian Sea (GC58), the contribution of ICD-PF is more prominent for the same time period, indicating a higher significance of coastal erosion for the East Siberian Sea compared to the Laptev Sea (Fig. 6), especially when compared to the early Holocene signature. Topsoil-PF fractions in PC23 are slightly higher (8–25 %) than in GC58 (3–23 %) from ~8,300 cal yrs BP to current day. The difference is likely caused by a strong influence of the Lena River at the sampling location of PC23 and less fluvial inflow to GC58 due to location farther away from the mouths of the Lena, Kolyma and Indigirka rivers.

When the shoreline was farther seaward during the early Holocene, the core PC23 from the Laptev Sea experienced a large influence of Lena River derived material (80–90 %) (Tesi et al. 2016a). This material was supplied to the Laptev Sea in response to the deglaciation and associated active-layer deepening in the watershed (Tesi et al. 2016a). Although the record of GC58 does not go back in time to the glacial-interglacial transition at the very onset of the Holocene, our results suggest that coastal erosion was likely the dominant process affecting the permafrost carbon supply and deposition also at that time. This seems likely, especially when considering the location of the core GC58 in between the rivers, and as has been observed in modern day shallower sediments in the East Siberian Sea (Bröder et al., 2016b; Vonk et al., 2012).
3.5 Biomarker indications of sources of terrestrial organic matter

The lignin fingerprint of organic matter sources in GC58 is consistent with the dual-carbon isotope modelling. Here we focus on the cinnamyl to vanillyl phenols and syringyl to vanillyl phenols ratios (C/V and S/V, respectively). The C/V ratio can be used to differentiate between woody (i.e. shrubs and trees) and non-woody (i.e. leaves, needles, grasses) plant tissues as origin of the terrestrial OC since cinnamyl phenols are produced only in non-woody vascular plant tissues (Hedges et al., 1988). Moreover, the S/V ratio differentiates between gymnosperms (conifers) and angiosperms (flowering plants) as syringyl phenols are produced solely in angiosperms (Hedges et al., 1988). Thereby higher S/V ratios mean more contribution from angiosperm plants.

The S/V and C/V ratios in GC58 show that the terrestrial material transported to the ESS originates mainly from soft tissue material (i.e. grasses and leaves) both from angiosperm and gymnosperm plants (Fig. 7). The lignin fingerprint of old Pleistocene material (ICD-PF) is characterised by high ratios of both C/V and S/V i.e. a high abundance of soft plant tissues from the tundra steppe vegetation (e.g. grass-like material) (Tesi et al. 2014; Winterfeld et al. 2015). Observations from the Laptev Sea (sediment core PC23, Fig. 1) reveal a much stronger influence from woody material indicating a watershed source, likely from the Lena River, rather than from coastal erosion (Fig. 7). It should be noted that the lignin phenols are susceptible to degradation. Cinnamyl phenols in particular are known to degrade fairly fast, which may lower the C/V ratios (Benner and Opsahl, 1995). However, even considering degradation effects, the relatively high C/V and S/V values that characterise GC58, indicate grass-type material typical of tundra/steppe biome and ICD-PF deposits (Tesi et al., 2014; Winterfeld et al., 2015a).
4 Conclusions

This down-core study provides new insights into terrestrial carbon dynamics in the East Siberian Sea (ESS) from the early Holocene warming period until the present. Our results suggest a high input of terrestrial organic carbon to the ESS during the last glacial-interglacial period caused by permafrost destabilisation. This material was mainly characterised as relict Pleistocene permafrost deposited via coastal erosion as a result of the sea level regression.

The flux rates of both lignin and cutin compounds show a declining trend from the early Holocene until today, suggesting a change from mainly terrestrial to marine dominated input. The same change can be seen in the stable carbon isotope ($\delta^{13}$C) data, which imply a regime shift from terrestrial to more marine dominated sediment input at ~8,400 cal yrs BP.

The source apportionment data highlights the importance of coastal erosion as a terrestrial carbon source to this region of the ESS throughout the Holocene. This is supported by the lignin composition, which suggests that the terrestrial carbon in the sediment core GC58 consists mainly of soft tissues of plants (i.e. grasses), typical for tundra/steppe vegetation during the Pleistocene. Both the biomarker and grain size data imply that the conditions have been more stable in the ESS in the past ~1,700 cal yrs BP compared to the early Holocene.

The comparison of the source apportionment results ($\delta^{13}$C, $\Delta^{14}$C) and the lignin fingerprint (C/V and S/V ratios) for the sediment cores GC58 and PC23 show a difference in the carbon sources between the East Siberian Sea and the adjacent Laptev Sea. The relict Pleistocene permafrost, mostly originating from coastal erosion, may be more dominant in the ESS than in the Laptev Sea. Data for the sediment core PC23 show that the Laptev Sea instead had a relatively high input of terrestrial carbon from the watershed, which is likely due to the influence of the Lena River.

The accelerating coastal erosion rates along the Siberian coast and amplified warming in the Arctic predicted by many climate models are likely to cause permafrost destabilisation and remobilisation of terrestrial carbon to the marine environment, as observed in the beginning of the Holocene. To better understand the consequences of the permafrost thawing processes, the extent of degradation of terrestrial carbon in the marine environment should be better constrained. Also, to improve the understanding of the processes in the ESS and in the whole Arctic region more historical down-core studies would be needed.
Author contributions

T. Tesi and Ö. Gustafsson conceived and designed the research project. T. Tesi, L. Bröder, I. Semiletov, O. Dudarev and Ö. Gustafsson collected the samples with the help from the IB/RV Oden crew. C. Pearce and K. Keskitalo developed the age-depth model of GC58. K. Keskitalo carried out all chemical and geological analyses on GC58 and MUC58. M. Sköld and A. Andersson ran the MCMC simulation for the OC source apportionment. A. Andersson estimated the lateral transport times. K. Keskitalo wrote the paper and produced the figures with input from all the co-authors.

Competing interests

The authors declare that they have no conflict of interest.
Acknowledgments

We thank the crew and personnel of IB/RV Oden. We thank Rienk Smittenberg for the use of the microwave extraction facilities. We also thank Carina Jakobsson, Heike Siegmund and Karin Wallner for their help with the laboratory analyses. This study was supported by the Knut and Alice Wallenberg Foundation (KAW contract 2011.0027), the Swedish Research Council (VR contract 621-2004-4039 and 621-2007-4631), the Nordic Council of Ministers Cryosphere-Climate-Carbon Initiative (project Defrost, contract 23001) and the European Research Council (ERC-AdG project CC-TOP #695331). Additionally, I. Semiletov thanks the Russian Government for financial support (mega-grant under contract #14.Z50.31.0012). O. Dudarev thanks the Russian Science Foundation for financial support (No. 15-17-20032). T. Tesi acknowledges EU financial support as a Marie Curie fellow (contract no. PIEF-GA-2011-300259). Contribution no. 1916 of ISMAR-CNR Sede di Bologna. L. Bröder acknowledges financial support from the Climate Research School of the Bolin Centre for Climate Research. C. Pearce received funding from the Danish Council for Independent Research / Natural Science (project DFF-4002-00098/FNU). M. Sköld acknowledges financial support from the Swedish Research Council (Grant 2013:05204).
References


Table 1. Radiocarbon (\(^{14}\)C) ages of the molluscs retrieved from the sediment core GC58. The \(^{14}\)C ages are shown in years BP with an age error (yrs) and as calibrated \(^{14}\)C ages (cal yrs BP) with one standard deviation (±1σ) of the individual \(^{14}\)C dates. Also shown the \(\delta^{13}\)C (‰) values of the molluscs.

<table>
<thead>
<tr>
<th>Corrected depth* (cm)</th>
<th>NOSAMS Accession nr.</th>
<th>Type</th>
<th>Age (^{14})C (yrs BP)</th>
<th>Age error (yrs)</th>
<th>(\delta^{13})C (‰)</th>
<th>Age (^{14})C (Cal yrs BP)</th>
<th>1σ</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.5</td>
<td>OS-119395</td>
<td>Mollusc, fragments</td>
<td>895</td>
<td>25</td>
<td>0.55</td>
<td>455</td>
<td>92</td>
</tr>
<tr>
<td>8.5</td>
<td>OS-120688</td>
<td>Mollusc, fragments</td>
<td>&gt;Modern</td>
<td>-</td>
<td>1.70</td>
<td>45</td>
<td>32</td>
</tr>
<tr>
<td>34.5</td>
<td>OS-120689</td>
<td>Mollusc, fragments</td>
<td>2,260</td>
<td>20</td>
<td>1.55</td>
<td>1,806</td>
<td>125</td>
</tr>
<tr>
<td>39.5</td>
<td>OS-120690</td>
<td>Mollusc, fragments</td>
<td>2,210</td>
<td>15</td>
<td>1.55</td>
<td>1,746</td>
<td>122</td>
</tr>
<tr>
<td>47.5</td>
<td>OS-123161</td>
<td>Mollusc, fragments</td>
<td>7,960</td>
<td>35</td>
<td>0.90</td>
<td>8,372</td>
<td>110</td>
</tr>
<tr>
<td>51.5</td>
<td>OS-119396</td>
<td>Mollusc, fragments</td>
<td>8,010</td>
<td>25</td>
<td>1.06</td>
<td>8,429</td>
<td>112</td>
</tr>
<tr>
<td>54.5</td>
<td>OS-120691</td>
<td>Mollusc, fragments</td>
<td>8,020</td>
<td>20</td>
<td>0.49</td>
<td>8,441</td>
<td>113</td>
</tr>
<tr>
<td>65.5</td>
<td>OS-119397</td>
<td>Mollusc, <em>Macoma calcarea</em></td>
<td>8,780</td>
<td>25</td>
<td>-2.46</td>
<td>9,372</td>
<td>117</td>
</tr>
<tr>
<td>72.5</td>
<td>OS-120692</td>
<td>Mollusc, fragments</td>
<td>8,880</td>
<td>20</td>
<td>-0.91</td>
<td>9,499</td>
<td>122</td>
</tr>
<tr>
<td>78.5</td>
<td>OS-120693</td>
<td>Mollusc, fragments</td>
<td>8,950</td>
<td>25</td>
<td>-0.79</td>
<td>9,595</td>
<td>132</td>
</tr>
</tbody>
</table>

*Corrected depth is the original depth + 3 cm to account for core top loss during sampling (Sect 2.4).
Figure 1. Map of the Eastern Siberian Arctic Shelf showing the location of the sampling site (Station SWERUS C3-1-58) (Schlitzer, R., Ocean Data View, http://odv.awi.de, 2015). Also shown in the map is the location of the sediment core PC23 (Station SWERUS C3-1-23, Tesi et al., 2016a). The red line marks the isobath (34 m water depth) which is approximately where the coast line was in the beginning of the sediment archive (GC58) ~9,500 cal yrs BP (Lambeck et al., 2014).
Figure 2. An age-depth model of the sediment core GC58 based on radiocarbon ($^{14}$C) dated molluscs (see Table 1) and $^{210}$Pb (base of a multicore collected at the same location). All the modelled dates were calibrated with Marine13 calibration curve. A $\Delta R$ value of 50±100 yrs was used to account for the differences in the local reservoir age. The core GC58 dates back ~9,500 cal yrs BP.
Figure 3. Organic matter composition of the sediment core GC58. The x-axis has breaks due to gaps in the sediment chronology. (a) Organic carbon fluxes (g m$^{-2}$ yr$^{-1}$) were high at the bottom of the core. The high fluxes at the top of the core are likely related to the merging of two dating systems ($^{210}$Pb and $^{14}$C, see Sect. 3.2). The sea level rose rapidly in the early Holocene (Lambeck et al., 2014). (b) Both lignin and cutin fluxes (mg m$^{-2}$ yr$^{-1}$) decrease toward the core top. High fluxes at the top of the core are influenced by the OC fluxes and likely do not show an actual increase in the fluxes of lignin and cutin (see Sect. 3.2). (c) Low molecular weight fatty acids (LMW-FA) show an influence of marine organic matter at the top of the core. (d) The δ$^{13}$C (‰) values illustrate a gradual shift from terrestrial dominated to more marine dominated input of organic matter towards the core top. The Δ$^{14}$C (‰) values (corrected for the time between the deposition and the measurement) show that the bulk organic carbon is older at the bottom of the core than at the core top. The drop in the Δ$^{14}$C values ~1,700 cal yrs BP is likely an artefact caused by the age model used to correct for the Δ$^{14}$C values.
Figure 4. Degradation proxies for terrestrial organic carbon in the sediment core GC58. The x-axis has breaks due to gaps in the sediment chronology. (a) Syringyl acid to syringaldehyde (Sd/Sl) and vanillic acid to vanillin (Vd/Vl) ratios are a lignin-phenol based degradation proxy. (b) Also the ratio of 3,5-dihydrobenzoic acid to vanillyl phenols (3,5-Bd/V) provides information on degradation of terrestrial organic carbon. Higher values imply more degraded material for all the ratios as illustrated with the turquoise arrow. The 3,5-Bd/V values suggest a gradual increase in degradation from the bottom of the core to the top.
Figure 5. Dual-carbon isotope ($\delta^{13}$C, $\Delta^{14}$C) based source apportionment of organic carbon (OC) illustrates fractions (%; mean ± SD) of old Pleistocene permafrost (ICD-PF) in brown, thaw of active-layer permafrost (topsoil-PF) in green and primary production (marine OC) in blue of the sediment core GC58. The ICD-PF is the dominant fraction throughout the core.
Figure 6. Lignin composition of the sediment core GC58 (black circles). The ratio between cinnamyl and vanillyl phenols (C/V) is used as a proxy to distinguish between soft and woody plant tissues. The ratio of syringyl to vanillyl phenols (S/V) indicates the difference between gymnosperm and angiosperm plants. The boxes indicate typical values for S/V and C/V ratios characterising different plant material (ranges from Goñi and Montgomery, 2000). Measured S/V and C/V ratios for Ice Complex Deposit permafrost (ICD-PF) are shown with green triangles (Winterfeld et al., 2015a) and with an orange square (±standard deviation) (Tesi et al., 2014). Measured S/V and C/V ratios for topsoil-PF (Lena River POC) are illustrated with orange diamonds (Winterfeld et al., 2015a). Also shown the lignin composition of the sediment core PC23 (blue diamonds) from the Laptev Sea.
Figure 7. Dual-carbon isotope ($\delta^{13}C$, $\Delta^{14}C$) composition of the sediment cores GC58 and PC23. Topsoil-PF refers to organic matter from the active-layer of permafrost, ICD-PF to relict Pleistocene Ice Complex Deposit permafrost (Yedoma) and marine OC to organic matter from primary production. The end-member values for different sources are taken from the literature (Bröder et al., 2016b; Tesi et al., 2016a). The green arrow points to the direction from the bottom to the top of the core (GC58).