



# Laminated sediments in the Bering Sea reveal atmospheric teleconnections to Greenland climate on millennial to decadal timescales during the last deglaciation

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**Abstract.** During the last glacial termination, the upper North Pacific Ocean underwent dramatic and rapid changes in oxygenation that lead to the transient intensification of oxygen minimum zones (OMZs), recorded by the widespread occurrence of laminated sediments on circum-Pacific continental margins. We present a new laminated sediment record from the mid-depth (1100 m) northern Bering Sea margin that provides insight into these deglacial OMZ maxima with exceptional, decadal-scale detail. Combined ultrahigh-resolution micro-X-ray-fluorescence (micro-XRF) data and sediment facies analysis of laminae reveal an alternation between predominantly terrigenous and diatom-dominated opal sedimentation. The diatomaceous laminae are interpreted to represent spring/summer productivity events related to the retreating sea ice margin. We identified five laminated sections in the deglacial part of our site. Lamina counts were carried out on these sections and correlated with the Bølling–Allerød and Preboreal phases in the North Greenland Ice Core (NGRIP) oxygen isotope record, indicating an annual deposition of individual lamina couplets (varves). The observed rapid decadal intensifications of anoxia, in particular within the Bølling–Allerød, are tightly coupled to short-term warm events through increases in regional export production. This dependence of laminae formation on warmer temperatures is underlined by a correlation with published Bering Sea sea surface temperature records and  $\delta^{18}\text{O}$  data of planktic foraminifera from the Gulf of Alaska. The rapidity of the observed changes strongly implies a close atmospheric teleconnection between North Pa-

cific and North Atlantic regions. We suggest that concomitant increases in export production and subsequent remineralization of organic matter in the Bering Sea, in combination with oxygen-poor waters entering the Bering Sea, drove down oxygen concentrations to values below  $0.1\text{ mL L}^{-1}$  and caused laminae preservation. Calculated benthic–planktic ventilation ages show no significant variations throughout the last deglaciation, indicating that changes in formation rates or differing sources of North Pacific mid-depth waters are not prime candidates for strengthening the OMZ at our site. The age models established by our correlation procedure allow for the determination of calendar age control points for the Bølling–Allerød and the Preboreal that are independent of the initial radiocarbon-based chronology. Resulting surface reservoir ages range within 730–990 yr during the Bølling–Allerød, 800–1100 yr in the Younger Dryas, and 765–775 yr for the Preboreal.

## 1 Introduction

The last glacial termination is characterized by the widespread decrease of oxygen concentrations at mid-depth in the world ocean (Jaccard and Galbraith, 2012). In the North Pacific, this prominent reduction led to the expansion of oxygen minimum zones (OMZs), culminating in the formation of anoxic, laminated sediment sequences at several continental margin locations (Fig. 1). Examples include the Santa Barbara Basin (e.g., Behl and Kennett, 1996), the Gulf

**Table 1.** Sediment cores from the Bering Sea containing laminated intervals during the last deglaciation. Depths are given in meter below sea level (m b.s.l.).

Sediment core	Latitude	Longitude	Depth (m b.s.l.)	Literature
SO202-10-5	52°44.57' N	179°50.87' E	1470	Gersonde (2012)
SO202-11-2	53°06.67' N	178°53.99' E	2703	Gersonde (2012)
SO202-12-1	54°03.04' N	179°05.24' E	2109	Gersonde (2012)
SO202-12-3	54°03.04' N	179°05.24' E	2109	Gersonde (2012)
SO202-13-5*	54°58.72' N	177°57.42' E	1382	Gersonde (2012)
SO202-14-3	56°47.19' N	178°49.45' E	3821	Gersonde (2012)
SO202-15-5	59°30.76' N	179°51.00' W	3129	Gersonde (2012)
SO202-17-1	60°07.39' N	179°27.95' W	1066	Gersonde (2012)
SO202-18-3	60°07.60' N	179°26.67' W	1111	Gersonde (2012); this study
SO202-18-6	60°07.60' N	179°26.61' W	1107	Gersonde (2012); this study
SO202-19-2	57°39.05' N	175°40.69' W	1752	Gersonde (2012)
SO202-20-1	55°08.68' N	171°04.95' W	2984	Gersonde (2012)
SO202-21-1	54°47.35' N	170°19.68' W	1911	Gersonde (2012)
SO202-22-2	54°34.45' N	168°48.66' W	1482	Gersonde (2012)
SO202-22-4	54°34.41' N	168°48.62' W	1476	Gersonde (2012)
SO201-2-114	59°13.87' N	166°59.32' E	1376	Max et al. (2012)
IODP U1339	54°40.02' N	169°58.902' W	1867.5	Takahashi et al. (2011)
IODP U1340	53°24.001' N	179°31.297' W	1294.6	Takahashi et al. (2011); Schlung et al. (2013)
IODP U1341	54°02.0025' N	179°0.49992' E	2139.5	Takahashi et al. (2011)
IODP U1342	54°49.699' N	176°55.003' E	818.6	Takahashi et al. (2011)
IODP U1343	57°33.399' N	175°48.966' W	1952.9	Takahashi et al. (2011)
IODP U1345	60°09.1917' N	179°28.204' W	1007.8	Takahashi et al. (2011)
HLY02-02-3JPC	60°07.674' N	179°26.508' W	1132	Cook et al. (2005)
HLY02-02-51JPC	54°33.192' N	168°40.014' W	1467	Cook et al. (2005); Caissie et al. (2010)
PC-23A	60°09.52' N	179°27.82' W	1002	Itaki et al. (2009); Kim et al. (2011)
PC-24A	60°15.70' N	179°25.34' W	852	Kim et al. (2011)
PC-25A	60°04.48' N	179°27.78' W	1152	Kim et al. (2011)
BOW-12A	53°23.47' N	179°33.47' W	1287	Okada et al. (2005); Tanaka and Takahashi et al. (2005)
UMK-3A	54°25.22' N	170°13.38' W	1892	Okada et al. (2005); Tanaka and Takahashi et al. (2005)

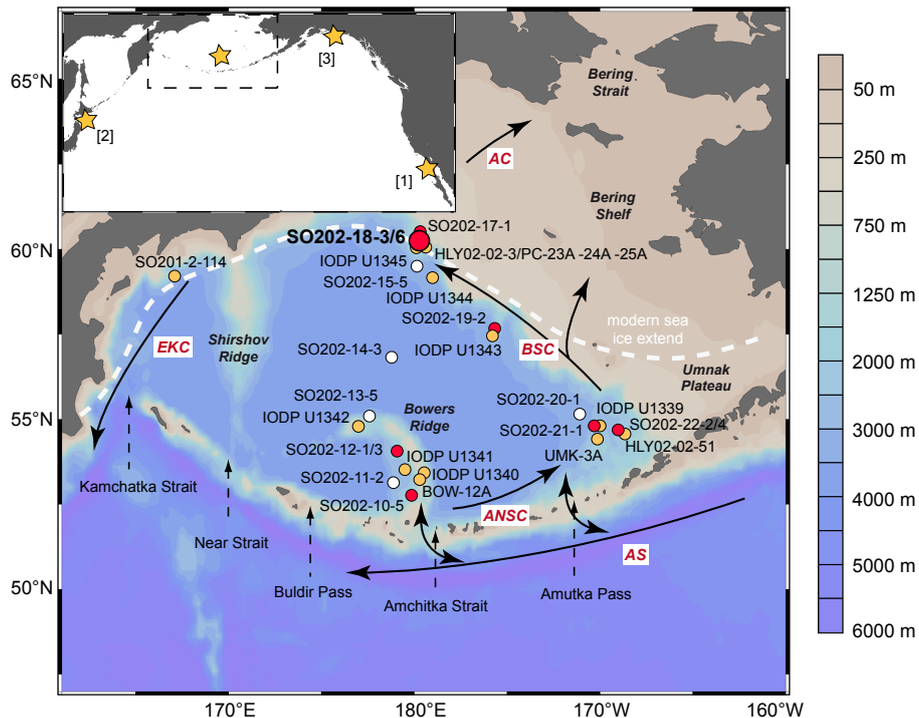
\* Magnetic susceptibility pattern and XRF data suggest that upper 15 ka are missing (Gersonde 2012).

of California (e.g., Kennett and Ingram, 1995; Zheng et al., 2000), the continental margins off Japan (Ikehara et al., 2006; Shibahara et al., 2007), the southeast Alaska Margin (Davies et al., 2011), and multiple sites located at intermediate depth in the Bering Sea (Fig. 1, Table 1 and cf., e.g., Cook et al., 2005; Itaki et al., 2009; Schlung et al., 2013).

The causes for the occurrence of laminations and hence significant decreases in oxygen concentration of bottom waters are discussed controversially. While some studies propose changes in the ventilation of intermediate waters and see no unambiguous evidence for higher export production (Kennett and Ingram, 1995; Zheng et al., 2000), others argue for increased export productivity that would contribute to anoxic conditions (Mix et al., 1999; Ortiz et al., 2004; Davies et al., 2011), or an interaction of both higher export production and ventilation changes (Cook et al., 2005; Shibahara et al., 2007). In records from lower North Pacific latitudes a strong coupling to North Atlantic climate was observed on millennial timescales in laminated sediments from the Santa Barbara Basin, which closely resemble Dansgaard–Oeschger–

type temperature changes of the Greenland Ice Sheet Project 2 (GISP2) ice core during the past 60 kyr (Kennett and Ingram, 1995; Hendy et al., 2002). Furthermore, a close atmospheric climate teleconnection between the North Atlantic and North Pacific has been proposed in several modeling studies (Mikolajewicz et al., 1997; Okumura et al., 2009). For the Bering Sea, it was shown that laminated sediments formed during warm phases of the Bølling–Allerød (14.64–12.85 ka) and the early Holocene (10.3–11.7 ka; e.g., Cook et al., 2005; Schlung et al., 2013). However, how closely laminated high-resolution sequences in the subarctic North Pacific are in fact linked to Greenland climate on shorter-than-millennial-scale changes remains unknown.

In principle, laminated sediments bear the potential to provide insights into climatic evolution and hemispheric coupling mechanisms with an extremely high temporal resolution (Hughen et al., 2000; Bronk Ramsey et al., 2012; Nakagawa et al., 2012). While it has been shown for laminated sediments from the Santa Barbara Basin and Saanich Inlet on Vancouver Island (Nederbragt and Thurow, 2001, 2005)



**Figure 1.** Bathymetric map of the Bering Sea with the position of laminated INOPEX cores (red circles) and INOPEX cores without laminations (white circles). Yellow circles mark the position of published laminated records in the Bering Sea (see Table 1 for details). The surface circulation is shown by black arrows. [AC]: Anadyr Current; [AS]: Alaska Stream; [ANSC]: Aleutian North Slope Current; [BSC]: Bering Slope Current; [EKC]: East Kamchatka Current. Dashed white line shows modern maximum sea ice extension (Reynolds et al., 2002). Dashed black arrows mark the gateways between the Bering Sea and North Pacific. Yellow stars on the small map mark location of laminated sediment records in [1] Gulf of California and Santa Barbara Basin (e.g., Behl and Kennett, 1996; Zheng et al., 2000), [2] near the island of Hokkaido (Ikehara et al., 2006; Shibahara et al., 2007), and [3] the southeast Alaska Margin (Davies et al., 2011). Map drawn with Ocean Data View (Schlitzer, 2002).

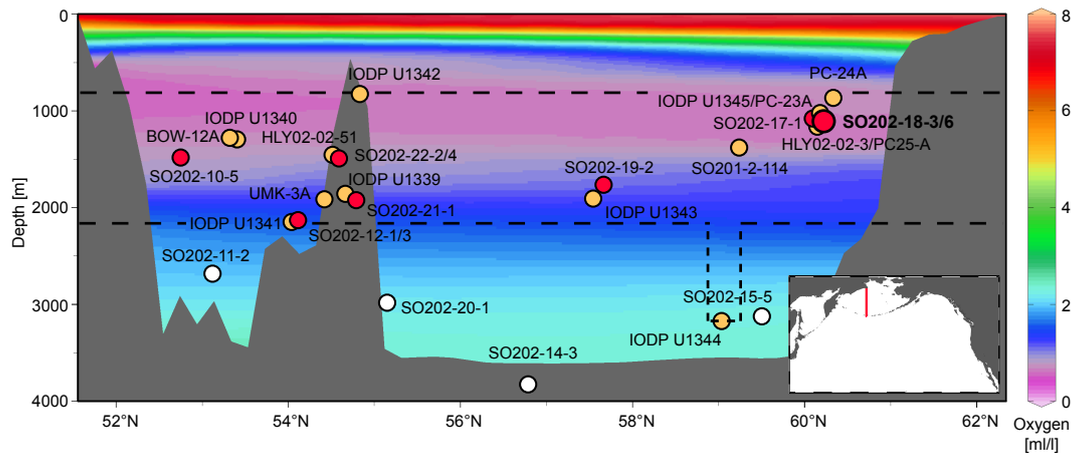
that each pair of laminae represents 1 year of deposition, the nature of laminations in the Bering Sea is largely unconstrained, with cases being made in previous studies for them either representing annual depositional events (Cook et al., 2005) or with no evidence found for such annual resolution (Caissie et al., 2010). Unfortunately, no laminae counts from Bering Sea sequences exist so far that shed light on this question.

In this paper, we investigate two episodically laminated sediment records from almost identical locations on the northern Bering Slope with high temporal resolution. These sites were selected from a collection of sediment cores retrieved during expedition SO202 INOPEX (Innovative North Pacific Experiment; Gersonde, 2012) to decipher deglacial changes in OMZ dynamics. With an accelerator mass spectrometry (AMS)  $^{14}\text{C}$ -anchored core chronology and detailed laminae counts, we give information on whether laminae were likely deposited annually and to what extent our records correspond to North Greenland Ice Core (NGRIP)  $\delta^{18}\text{O}$  time series, especially during the warm phases of the Bølling–Allerød (GI-1; Andersen et al., 2006; Rasmussen et al., 2006) and the Preboreal (11.65–10.70 ka;

Mangerud et al., 1974; van der Plicht et al., 2004). High-resolution X-ray fluorescence (micro-XRF) data and sediment X-ray images provide information about formation processes of discrete laminae. We compare our results to existing seasonal sea surface temperature (SST) records from the Bering Sea and Greenland ice core data to deduce connections between hemispheric temperature changes and anoxia occurrence. Calculated benthic–planktic (B–P) ventilation ages will show whether water mass changes contribute to laminae formation.

## 2 Study area and oceanographic setting

The Bering Sea is the third-largest marginal sea in the world and comprises a broad continental shelf (Fig. 1). This shelf is seasonally covered with sea ice, with maximum extension occurring between March and April, including our core location, and minimum sea ice extension between July and September (Niebauer, 1980; Danielson et al., 2011). These strong seasonal contrasts are reflected in the SSTs, which range between 8 °C (summer) and 0.6 °C (winter) at our site



**Figure 2.** North–south profile through the Bering Sea and annual mean oxygen levels (Garcia et al., 2009). Red circles mark cores from this study containing laminated sediments; white circles mark cores from the INOPEX expedition without laminations. Core SO202-13-5 is not shown as the upper 15 kyr seems to be missing (Gersonde, 2012). Yellow circles mark published sediment records containing laminated sediments (see Table 1 for details). Dashed lines represent maximum and minimum water depth of the OMZ in the Bering Sea for the last deglaciation based on the occurrence of laminated sediment cores. Note that the deepening of the OMZ down to 3173 m water depth might be a local phenomenon (see text for details). Transect drawn with Ocean Data View (Schlitzer, 2002).

(Levitus and Boyer, 1994), and are induced by the interplay between the Siberian High and the Aleutian Low (Niebauer, 1980). Surface water masses are influenced by the Alaska Stream (AS), which enters the Bering Sea through several straits between the Aleutian Islands. The surface circulation is characterized by a counterclockwise gyre, formed by the East Kamchatka Current (EKC), the Bering Slope Current (BSC), and the Aleutian Northern Slope Current (ANSC), the lattermost being the link between the inflowing water from the North Pacific and the BSC. The BSC contributes to the northward-flowing Anadyr Current (AC) and feeds the EKC. Surface waters leave the Bering Sea with the southward-flowing EKC to the North Pacific. Deepwater exchange occurs via Kamchatka Strait and Near Strait, but only Kamchatka Strait is deeper than 2000 m and allows inflow of deep Pacific waters. Modern mid-depth to deep waters (ca. 200–2500 m water depth) within the Bering Sea itself are not well investigated but thought to be rather homogenous in their physical and chemical characteristics and mainly represent a mixture of North Pacific Deep and Intermediate Water (Tsunogai et al., 1979; Luchin et al., 1999). Primary production is high, especially along the edge of the eastern continental shelf and is caused by tidal mixing and transverse circulation, which gave this area the name “Green Belt” (Springer et al., 1996). Biological productivity is dominated by siliceous plankton, with diatoms as major constituents of biogenic opal (Takahashi et al., 2002). Although no persistent mid-depth anoxia exist in the Bering Sea today, an OMZ develops between 900 and 1000 m water depth (Fig. 2, Garcia et al., 2010). In contrast to other permanent OMZs – e.g., in the eastern North and South Pacific – the one in the Bering Sea exhibits a significant seasonality, with lowest oxygen con-

centrations occurring during winter season (Paulmier and Ruiz-Pino, 2009). Throughout the year, oxygen concentrations in the core depth of the OMZ range between 0.4 and 0.7 mL L<sup>-1</sup> (Garcia et al., 2010).

### 3 Material and methods

#### 3.1 Sediment records

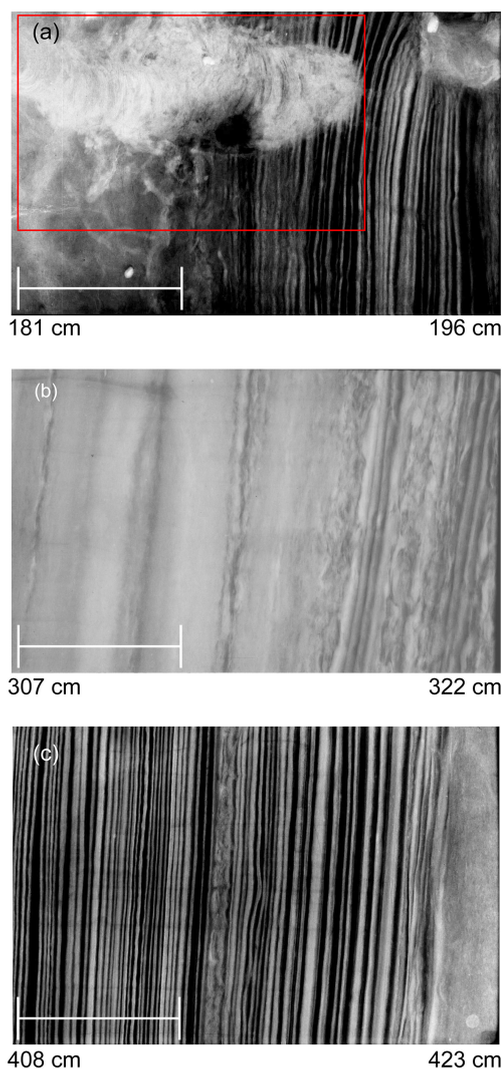
During R/V *Sonne* cruise SO202-INOPEX, 15 sediment cores were recovered from the Bering Sea in 2009 (Fig. 1) on a depth transect from 1066 to 3821 m water depth (Fig. 2). Of these cores, eight yielded laminated sequences. In this study, we use two cores from an almost identical position: piston core SO202-18-3 (60°07.60' N, 179°26.67' W; water depth: 1111 m; 10.5 cm core diameter) and neighboring kasten core SO202-18-6 (60°07.60' N, 179°26.61' W; water depth: 1107 m, 30 cm long rectangular sides). Both were retrieved off the large shelf in the northern Bering Sea, and high-resolution echo sound sub-bottom profiling data show that both coring devices penetrated to the same sediment intervals. The sediments consist of diatom-bearing silty to sandy clay. A prominent feature in both cores is the occurrence of several, meter-thick laminated sequences consisting of alternating laminae of lighter pure diatom ooze and darker diatomaceous mud. Also, several ash layers appear in both cores. Detailed core descriptions and sediment echo sounding data from the site are provided in Gersonde (2012).

### 3.2 Geochemical and X-ray image analyses

In order to obtain high-resolution geochemical data sets of major element composition, non-destructive XRF measurements were performed at the Alfred Wegener Institute with an Avaatech XRF core scanner at 1 cm resolution for both sediment cores. Each core segment was scanned three times with tube voltages of 10, 30, and 50 kV and counting times of 10, 15, and 30 s, respectively. In a second step, laminated sequences were scanned with up to 200  $\mu\text{m}$  resolution at the Leibniz Institute for Baltic Sea Research using an ITRAX core scanner from Cox Analytical Systems and 45 kV Cr-tube voltage and 15 s counting time. Radiography images on the kasten core were obtained by cutting out  $25 \times 10 \times 0.5$  cm sediment slices that were wrapped into plastic foil and sealed vacuum-tight. For the radiographies, we used the cabinet X-ray system Faxitron Series from Hewlett Packard with 40 kV, 3 A and 3 min exposure time. For the piston core, X-ray images were taken with the ITRAX core scanner on split sediment surfaces before the XRF scan, with tube voltages between 50 and 60 kV, a tube current of 30 mA, and exposure times between 1000 and 1200 ms depending on the thickness and density of the sediment. The radiographic images from the ITRAX scanner are radiographic positives, in which core segments of lower density appear as light intervals and vice versa (Croudace et al., 2006). Biogenic opal was measured by the sequential leaching method after Müller and Schneider (1993), total carbon content was measured with an Elementar Vario III CNS analyzer, and total organic carbon content with an Eltra CS-2000 Carbon Sulfur element analyzer after removal of carbonate. Sediment samples were also corrected for the pore water salt content (Kuhn, 2013).  $\text{CaCO}_3$  was calculated as the difference between the former two multiplied by 8.333.

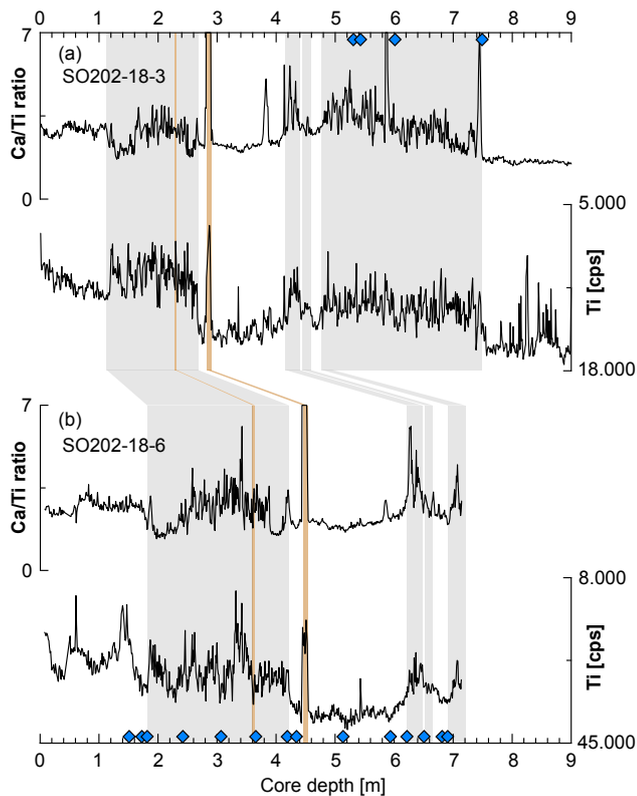
### 3.3 Radiocarbon dating

For AMS  $^{14}\text{C}$  analyses, freeze-dried sediment slices of 2–2.5 cm thickness were washed over a 63  $\mu\text{m}$  mesh-size sieve and dried after washing at 30 °C. At least 1.7 mg of monospecific samples of the planktic foraminifera *Neogloboquadrina pachyderma* sinistral were picked from the 125–250  $\mu\text{m}$  fraction. This foraminifera is a subsurface-dwelling species that lives in the North Pacific between 50 and 200 m water depth (Kuroyanagi et al., 2002; Bauch et al., 2002). Radiocarbon dating on benthic foraminifera were carried out on mixed benthic, mostly shallow infaunal species (like *Uvigerina* spp., *Elphidium* spp, *Nonionella* spp.; cf. Buben-shchikova et al., 2008) from the fraction  $> 250 \mu\text{m}$ . All radiocarbon measurements were performed by the National Ocean Sciences Accelerator Mass Spectrometry (NOSAMS) facility at the Woods Hole Oceanographic Institution. Ages are reported following established conventions (Stuiver and Polach, 1977). All radiocarbon dates were converted into calendar ages with the calibration software Calib 7.0 (Stu-



**Figure 3.** Examples of three different sediment facies in SO202-18-3/6 as shown in radiographs. (a) Bioturbated Facies (left side) and Bioturbated Lamination Facies (middle section) at the end of Preboreal laminations, (b) Layered Facies, and (c) Laminated Facies during the Preboreal. All radiographs are shown with increasing core depth to the right. The core depths of each radiograph sections are indicated below the images. The red box in (a) marks a bioturbational feature that led to an age reversal in a  $^{14}\text{C}$  age in core SO202-18-6, 180–182.5 cm. The scale bar on each radiograph marks a 5 cm interval.

iver and Reimer, 1993) and the INTCAL13 calibration curve (Reimer et al., 2013). For the initial age conversion, a planktic reservoir age of  $R = 700$  yr was chosen, in line with previous works (Max et al., 2012; Rella et al., 2012; Itaki et al., 2009; Cook et al., 2005). Changes in deep-water ventilation ages were derived by calculating benthic–planktic ventilation ages, which are the difference between coeval benthic and planktic foraminiferal  $^{14}\text{C}$  ages from the same respective core depth. All AMS  $^{14}\text{C}$  measurements and



**Figure 4.** Inter-core correlation between piston core SO202-18-3 (a) and kasten core SO202-18-6 (b) based on XRF-scanning data of the Ca/Ti ratio and Ti concentrations (given in counts per second). Gray bars mark the occurrence of Layered and Laminated Facies; brown bars mark ash layers. Note inverse scale of Ti concentrations. Blue diamonds mark planktic AMS  $^{14}\text{C}$  ages.

calibration results are provided in Table 2. One  $^{14}\text{C}$  result of the depth interval 180–182.5 cm in SO202-18-6 was discarded from further analysis, as it shows an age reversal that is likely attributed to a localized, bioturbation feature, which is clearly visible in the X-ray image from this depth interval (cf. Fig. 3a). For our other laminated cores from the Bering Sea an initial age model to determine Termination I was constructed based on correlation of XRF results (Supplement Fig. S1).

### 3.4 Sediment classification and laminae counts

Laminae counts were performed manually on X-ray images, as the single laminae, due to their varying densities, are clearly visible in these images, compared to digital photos of the core surface. In addition, we combined the X-ray images with chlorine (Cl) counts from micro-XRF scans as a proxy for water contents (Tjallingii et al., 2007). These counts responded sensitively to density changes within the different laminae, as light and dark laminae contain different amounts of pore water. The combination of X-ray images and high-resolution ITRAX data has been previously used and shown

to be a suitable method for laminae counting (e.g., Staff et al., 2012). We always counted couplets of one light and one dark lamina. Laminae counting proceeded until no single laminae were visible anymore in the X-ray images, e.g., through the effect of bioturbation. The deeper laminated sequences were counted on piston core SO202-18-3, as they were not fully recovered in core SO202-18-6. For the upper laminated sections, we concentrated on core SO202-18-6, which has a better core quality during this interval. Based on repetitive laminae counts from different core section and by different investigators, the counting error is estimated to be  $\pm 5$  laminae couplets for respective, ca. 1 m long, laminated sequences. This error estimation is used as counting uncertainties for the prominent laminated units that we identified in our cores (cf. Sect. 4.3). As we do not have a continuous record of laminated sediments, we cannot estimate a cumulative maximum counting error, which increases with depth, as done for the Greenland ice core record (Rasmussen et al., 2006).

For this study, the sediment was classified into four different types of sediment facies: (1) “Laminated Facies”, (2) “Layered Facies”, (3) “Bioturbated Lamination Facies”, and a (4) “Bioturbated Facies”. A comparable approach was applied previously in other Pacific locations on laminated sediments from the Santa Barbara Basin (Behl and Kennett, 1996) and the Japan Sea (Watanabe et al., 2007) in order to assign different types of sediment to different oxygen concentrations. The differentiation into these categories was done by visual inspection of the X-ray images (Fig. 3): the Laminated Facies consists of distinct and undisturbed sub-millimeter-scale up to 6 mm thick laminations with clear boundaries that can be counted and do not show signs of burrowing or other disturbances. In the Layered Facies, boundaries between biogenic and siliciclastic parts of the laminations are not clearly discernible or disappearing and centimeter-thick, undisturbed layers of mostly siliciclastic material are formed instead of clearly alternating sedimentary patterns as in the Laminated Facies. The thickness of these facies in our sediment core ranges from several centimeters down to 1 cm. In the Bioturbated Lamination Facies, originally laminated structure is partially disturbed through burrowing organisms. Lastly, the Bioturbated Facies does not show any traces of laminations or layers, since it has been completely homogenized by benthic organisms.

## 4 Results and discussion

### 4.1 Initial $^{14}\text{C}$ -supported age model and regional stratigraphic context

In order to correlate cores SO202-18-3 and SO202-18-6, the occurrence of two distinct ash layers, Ca/Ti ratios, and Ti counts from XRF scanning, as well as the occurrence of the laminated sections, were used (Fig. 4). In the XRF scans the

**Table 2.** AMS  $^{14}\text{C}$  ages, calibrated calendar ages with  $1\sigma$  ranges, reservoir ages (res. age), and calculated benthic–planktic ventilation ages (ventil. age). Marked bold/italic res. ages are deduced by NGRIP correlation/layer counting. For comparison calendar ages with a constant reservoir age of  $R = 700$  yr are also shown. Nps: *Neogloboquadrina pachyderma* sinistral; Uvi: *Uvigerina* spp.; mb: mixed benthic foraminifera.

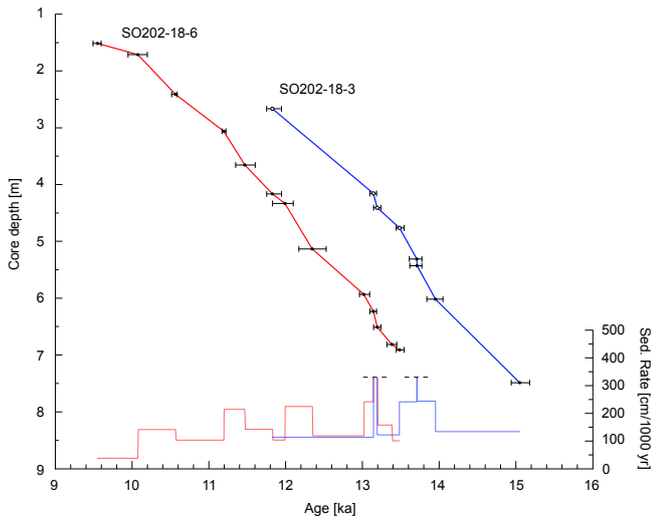
Core	Sample ID	Depth (cm)	Species	$^{14}\text{C}$ age (yr)	Res. age (yr)	Ventil. ages (yr)	Error ventil. ages $\pm$ (yr)	Calendar age (yr)	Calendar age (yr) $1\sigma$ range	Calendar age (yr) with 700 yr res.age
SO202-18-3	OS-93967	530–532	Nps	12 600 $\pm$ 50	<b>870</b>	650	100	13 539	13 472–13 579	13 710
	OS-93973	530–532	mb	13 250 $\pm$ 50						
	OS-93968	542–544	Nps	12 600 $\pm$ 45	<b>820</b>	850	100	13 612	13 551–13 711	13 712
	OS-93881	542–544	mb	13 450 $\pm$ 55						
	OS-93978	601–603	Nps	12 800 $\pm$ 65	<b>730</b>	700	115	13 918	13 809–14 005	13 954
	OS-93969	601–603	mb	13 500 $\pm$ 50						
	OS-92056	747.5–750	Nps	13 350 $\pm$ 55	<b>880</b>	600	110	14 640	14 419–14 856	15 049
	OS-92023	747.5–750	Uvi	13 950 $\pm$ 55						
SO202-18-6	OS-96110	150–152.5	Nps	9280 $\pm$ 65	770	680	115	9505	9476–9539	9550
	OS-96032	150–152.5	mb	9960 $\pm$ 50						
	OS-93972	170–172.5	Nps	9650 $\pm$ 35	770	800	85	10 029	9917–10 150	10 075
	OS-93974	170–172.5	mb	10 450 $\pm$ 50						
	OS-94161 <sup>a</sup>	180–182.5	Nps	9520 $\pm$ 80	770	1030	130	9766	9604–9890	9884
	OS-96033	180–182.5	mb	10 550 $\pm$ 50						
	OS-85754	240–242.5	Nps	10 050 $\pm$ 25	770	450	80	10 476	10 422–10 518	10 566
	OS-90768	240–242.5	mb	10 500 $\pm$ 55						
	OS-90732	305–307.5	Nps	10 450 $\pm$ 40	770	600	90	11 126	10 907–11 194	11 195
	OS-92014	305–307.5	mb	11 050 $\pm$ 50						
	OS-85755	364–366.5	Nps	10 700 $\pm$ 35	<b>765</b>	700	80	11 324	11 263–11 388	11 470
	OS-90699	364–366.5	mb	11 400 $\pm$ 45						
	OS-85756 <sup>b</sup>	415–417.5	Nps	10 850 $\pm$ 25	<b>775</b>	450	75	11 650	11 506–11 750	11 828
	OS-90698 <sup>b</sup>	415–417.5	mb	11 300 $\pm$ 50						
	OS-96111 <sup>b</sup>	432–434.5	Nps	10 950 $\pm$ 55	830	600	115	11 745	11 614–11 946	11 992
	OS-96112 <sup>b</sup>	432–434.5	mb	11 550 $\pm$ 60						
	OS-94120 <sup>b</sup>	512–514.5	Nps	11 150 $\pm$ 65	800	650	125	12 210	12 083–12 382	12 348
	OS-96034 <sup>b</sup>	512–514.5	mb	11 800 $\pm$ 60						
	OS-96095 <sup>b</sup>	592–594.5	Nps	11 850 $\pm$ 60	1100	450	140	12 685	12 654–12 724	13 020
	OS-96035 <sup>b</sup>	592–594.5	mb	12 300 $\pm$ 80						
	OS-85757	622–624.5	Nps	12 000 $\pm$ 35	<b>990</b>	650	100	12 869	12 795–12 935	13 144
	OS-90733	622–624.5	mb	12 650 $\pm$ 65						
	OS-85758	650–652.5	Nps	12 050 $\pm$ 30	<b>970</b>	700	75	12 955	12 897–13 028	13 194
	OS-92049	650–652.5	mb	12 750 $\pm$ 45						
	OS-96107	680–682.5	Nps	12 250 $\pm$ 70	860	600	140	13 230	13 150–13 293	13 384
	OS-96106	680–682.5	mb	12 850 $\pm$ 70						
OS-85759	690–692.5	Nps	12 350 $\pm$ 30	<b>880</b>	650	90	13 318	13 274–13 353	13 483	
OS-92057	690–692.5	mb	13 000 $\pm$ 60							

<sup>a</sup> Age reversal. <sup>b</sup> Derived from Max et al. (2014).

laminated and layered sequences in both cores are characterized by higher Ca/Ti ratios and lower Ti counts (Fig. 4), with similar peak structure especially in the Ca/Ti ratios. Due to the shorter length of the kasten core, the lowermost laminated section was only partly recovered in core SO202-18-6. In total, piston core SO202-18-3 contains 461 cm of low-oxygen sediment sequences (i.e., Laminated and Layered Facies), and kasten core SO202-18-6 about 300 cm. We correlated the onset of laminated and non-laminated sections as basic lithostratigraphic units, by visual inspection and XRF-scanning records between cores SO202-18-3 and SO202-18-6. As an independent correlation marker we used two tephra layers identified in both cores (Table 3 and Gersonde, 2012). This enabled us to transfer  $^{14}\text{C}$  ages of core SO202-18-6 to corresponding depths of core SO202-18-3, where the upper part was not sampled. The correlated core depths of core SO202-18-3 are given in Supplement Ta-

ble S1. An overview of the different sedimentary facies with according core depths is given in Table 3; the correlation pattern is shown in Fig. 4. Other laminated INOPEX cores from the Bering Sea reveal similar trends in the XRF data (Supplement Fig. S1).

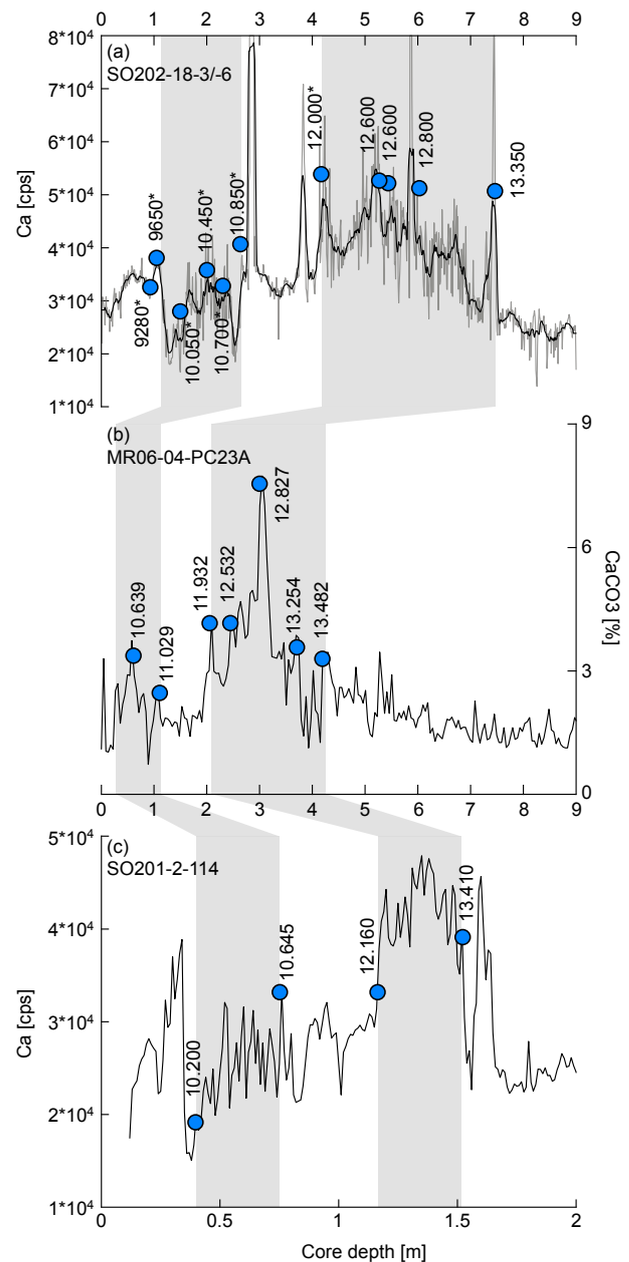
To establish a chronostratigraphic framework for the combined core SO202-18-3/6, we converted the individual planktic AMS  $^{14}\text{C}$  ages to calendar ages by using the INTCAL13 calibration curve (Reimer et al., 2013). In this first age control step, we used a constant reservoir age of  $R = 700$  yr (Table 2), in line with most previous studies (Gorbarenko et al., 2005; Itaki et al., 2009; Max et al., 2012). However, marine reservoir ages likely have varied over the glacial termination in the western North Pacific (Sarnthein et al., 2007). We address the potential changes in local reservoir ages in the discussion of Sect. 4.3. An age–depth relationship (Fig. 5) was established by linear interpolation between



**Figure 5.** Age–depth relationship of cores SO202-18-3 (blue curve) and SO202-18-6 (red curve) based on use of constant reservoir ages of 700 yr. Error bars show 1 $\sigma$  ranges of <sup>14</sup>C results. White circles in blue curve show ages derived by core-to-core correlation. Light red and blue curves show sedimentation rates. Note the cut-off in the sedimentation rate at 325 cm 1000 yr<sup>−1</sup> marked by a dashed line.

individual planktic AMS <sup>14</sup>C age control points through their reported median probability ages (Telford et al., 2004). According to this age model, linear sedimentation rates mostly range between 150 and 250 cm ka<sup>−1</sup>.

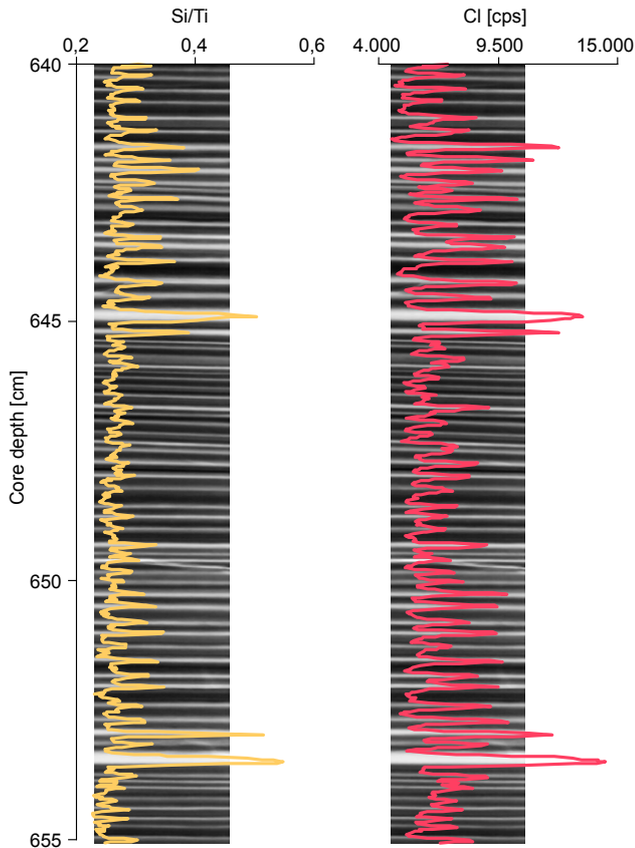
To incorporate the radiocarbon-based age model of our sediment record into an established regional stratigraphic framework for the last deglaciation, we used calcium carbonate concentrations, represented by Ca XRF counts in our records and compared those with earlier published sediment records. Deglacial sediment sequences in the North Pacific region are characterized by the occurrence of two prominent calcium carbonate peaks that have been radiocarbon-dated and assigned to glacial terminations Ia and Ib in the open North Pacific (Keigwin et al., 1992; Galbraith et al., 2007), the Bering Sea (Gorbarenko et al., 2005), and the Okhotsk Sea (Keigwin, 1998; Max et al., 2012). Our Ca concentration records closely correspond to previously published carbonate data (Itaki et al., 2009; Max et al., 2012; Rella et al., 2012). In those studies, following other published records (Cook et al., 2005; Kim et al., 2011; Schlung et al., 2013), the lower laminated sequence was assigned to the Bølling–Allerød interstadial, whereas the upper laminated interval was identified to represent the early Holocene. In particular, the 11-point smoothed Ca counts curve of core SO202-18-3 (Fig. 6a) closely resembles the lower-resolved CaCO<sub>3</sub> record of nearby core PC23A (Fig. 6b, Itaki et al., 2009; Rella et al., 2012) in the laminated intervals, with similar peak structures. The corresponding uncorrected AMS <sup>14</sup>C ages of planktic foraminifera measured on the individual carbonate peaks in SO202-18-3 and SO202-18-6 are nearly



**Figure 6.** Regional stratigraphic framework based on established basin-wide carbonate peak patterns (Max et al., 2012; Keigwin et al., 1992). Comparison of core SO202-18-3 (this study) to published Bering Sea records with radiocarbon age control and laminated sediment sections. (a) Ca XRF counts of SO202-18-3; thick black line shows the 11-point moving average of the Ca data. <sup>14</sup>C ages in this graph marked with asterisks are from SO202-18-6 and were transferred to the corresponding depths of SO202-18-3 through core-to-core correlation. One <sup>14</sup>C age is not plotted, as it shows an age reversal (see text for details). (b) Core MR06-04-PC23A (Itaki et al., 2009; Rella et al., 2012). (c) Core SO201-2-114 (Max et al., 2012). Blue circles denote radiocarbon dates age with raw, uncalibrated <sup>14</sup>C ages. Gray shading indicates anoxic (laminated/layered) core sections.

**Table 3.** Core depths of the different facies types occurring in cores SO202-18-3 and SO202-18-6: lam: Laminated Facies; lay: Layered Facies; biolam: Bioturbated Lamination Facies; bio: Bioturbated Facies; ash: ash layer. Depth intervals of prominent laminated units (TI-BLU1–5) are marked bold/italic.

Core	Depth (cm)	Facies	Core	Depth (cm)	Facies
SO202-18-3	0–111.82	bio	SO202-18-6	0–183.91	bio
	111.82–120.07	biolam		183.91–191.84	biolam
	120.07–125.52	lam		191.84–193.5	lam
	125.52–126.6	lay		193.5–208.57	biolam
	126.6–128.31	lam		208.57–214	lam
	128.31–132.57	lay		214–256.42	lay
	132.57–137.1	lam		256.42–258.49	lam
	137.7–157.45	lay		258.49–261.04	lay
	157.45–159.62	lam		261.04–263.09	lam
	159.62–163.47	lay		263.09–269.33	lay
	163.47–169.37	lam		269.33–270.18	lam
	169.37–173.4	lay		270.18–280.83	lay
	173.4–174.28	lam		280.83–300.33	lam
	174.28–180.66	lay		300.33–317.67	lay
	180.66–196.91	lam		317.67–322	lam
	196.91–200.62	lay		322–324.84	lay
	200.62–201.99	lam		<b>324.84–360.25</b>	<b>lam</b>
	201.99–203.92	lay		<b>360.25–362.49</b>	<b>ash</b>
	<b>203.92–226.7</b>	<b>lam</b>		<b>362.49–421.05</b>	<b>lam</b>
	<b>226.7–231.39</b>	<b>ash</b>		421.05–448	bio
	<b>231.39–266.27</b>	<b>lam</b>		448–453.57	ash
	266.27–282.2	bio		453.57–620.02	bio
	282.2–287.5	ash		620.02–629.85	biolam
	287.5–415.75	bio		<b>629.85–647.39</b>	<b>lam</b>
	415.75–421.18	biolam		647.39–651.74	bio
	<b>421.18–423.43</b>	<b>lam</b>		651.74–665.68	biolam
	<b>423.43–424.89</b>	<b>lay</b>		665.68–690.75	bio
	<b>424.89–440.78</b>	<b>lam</b>		690.75–698.8	biolam
	440.78–444.08	bio		698.8–725	lam
	444.08–458.83	biolam			
	458.83–476.15	bio			
	476.15–483.59	biolam			
	<b>483.59–503.4</b>	<b>lam</b>			
	<b>503.4–504.52</b>	<b>lay</b>			
	<b>504.52–509.7</b>	<b>lam</b>			
	<b>509.7–510.36</b>	<b>lay</b>			
	<b>510.36–527.42</b>	<b>lam</b>			
	527.42–528.85	lay			
	528.85–532.46	lam			
	532.46–533.48	lay			
	533.48–535.33	lam			
	535.33–536	lay			
	536–538.5	lam			
	538.5–540.73	lay			
	540.73–542.72	lam			
	542.72–543.84	lay			
	<b>543.84–561.75</b>	<b>lam</b>			
	<b>561.75–563.2</b>	<b>lay</b>			
	<b>563.2–580.76</b>	<b>lam</b>			
	<b>580.76–581.77</b>	<b>lay</b>			
	<b>581.77–602.81</b>	<b>lam</b>			
	602.81–608.87	lay			
	608.87–615.45	lam			
	615.45–618.16	lay			
	<b>618.16–747.08</b>	<b>lam</b>			
	747.08–1849	bio			



**Figure 7.** Example micro-XRF Si/Ti and Cl counts and X-ray images of SO202-18-3, 640–655 cm core depth. Note that less dense laminae are shown in white. The white laminae with high amounts of well-preserved diatoms represent the spring/summer bloom, while the dark, denser laminae contain high amounts of terrigenous material.

identical to those dated in core PC23A, i.e., mostly differing by less than  $100^{14}\text{C yr}$ . Though it is less densely dated, core SO201-2-114 with laminated sediment sections from the western Bering Sea margin (Max et al., 2012) also closely matches our carbonate peak pattern and uncorrected  $^{14}\text{C}$  ages (Fig. 6c).

The observed close similarities between the cores of this study and previously published records give us confidence to assign our Laminated and Layered Facies sections to the deglacial phases of the Bølling–Allerød and Preboreal, and the bioturbated intermediary sediment sections to the Younger Dryas (GS-1) interval. Based on this assumption we progressed to subsequent laminae counts and analyses, in order to assess the nature of deglacial and Preboreal anoxia occurrences on the Bering Sea margin and their temporal relationship with regional ocean proxy time series and the NGRIP ice core reference record (NGRIP-Members et al., 2004).

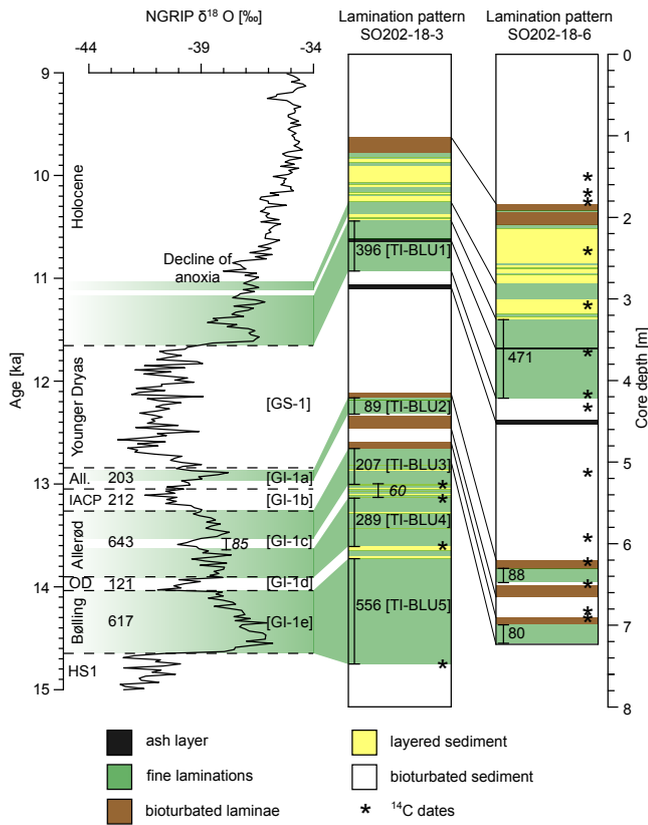
The comparison to other laminated sediment cores from the Bering Sea also reveals that the OMZ expanded to greater

water depths, as sediment cores that lie below the lower boundary of the modern moderate OMZ of 900 m water depth show laminations during the last deglaciation (Fig. 2). The maximum vertical expansion of the OMZ according to the lamination occurrence in sediment cores extended from 818 m (laminated core U1342; Takahashi et al., 2011) to a maximum of 3173 m (laminated core U1344; Takahashi et al., 2011). However, such deep expansion of the deglacial OMZ might be a localized phenomenon of the core U1344 site, as a comparable depth was not recorded as laminated in cores retrieved during the INOPEX cruise. There, the deepest laminated cores are from 2109 m (SO202-12), during the last deglaciation (Figs. 1, 2; Gersonde, 2012).

#### 4.2 Core site lamination pattern based on XRF and X-ray results

The micro-XRF data and radiographs reveal that the laminae of pure diatom ooze have a lower density due to the high concentration of well-preserved diatoms. This also leads to higher Si/Ti ratios and Cl counts (Fig. 7), the latter being the effect of the high porosity of the material and thus higher pore water content. Through the effect of dilution, the diatom-rich laminated sequences also lead to generally lower Ti counts within these intervals (Fig. 4). Microscopic analyses of selected smear slides from both cores indicate that Si-rich laminae comprise well-preserved diatom species such as *Fragilariopsis oceanica*, *F. cylindrus*, *Bacterosira bathyomphala* and *Porosira glacialis*, which are sea-ice- or cold-water-related species in the modern Bering Sea (Fig. S2; Ren et al., 2014). Preliminary diatom countings revealed concentrations of up to 76 % for the sea-ice-related diatoms *F. oceanica* and *F. cylindrus* in the diatom species assemblages of these laminae. Detailed counting results of selected smear slides are given in Supplement Table S2. Phytoplankton blooms at ice edges during spring are common in the Bering Sea today (Niebauer and Alexander, 1985). Accordingly, we interpret the Si-rich layers as diatom-productivity-dominated time intervals related to the spring/summer sea ice break-up phase, whereas their denser counterparts are lower in biogenic Si and contain relatively large amounts of terrigenous detrital material, rather reflecting the autumn/winter sedimentation.

The X-ray images further reveal that the transition from bioturbated sediment to laminations occurs abruptly, especially at the Younger Dryas–Holocene transition in both cores and at the onset of the Bølling in core SO202-18-3. However, traces of bioturbation can reach several centimeters downcore at the transition from laminated to bioturbated sediment (Fig. 3a). According to our sediment analysis there are repeated, thicker intervals of laminated sediment sequences in our cores. We call these “Termination I Bering Sea Lamination Units” with consecutive numbering from the younger to older sequences (TI-BLU1–5). In core SO202-18-3, TI-BLU3–5 are separated by thinner sediment



**Figure 8.** Correlation between lamination pattern in cores SO202-18-3/6 to 20 yr average NGRIP ice core data (Rasmussen et al., 2006). Dashed lines and captions on the right side of NGRIP data mark the events according to the GICC05 chronology, and their duration is given with the numerical numbers between the dashed lines. The left side of the stable isotope graph shows the commonly used terminology of the Blytt–Sernander sequence (Mangerud et al., 1974) for comparison. The italic number inside GI-1c gives the duration of a short cold period inside the Allerød that we also identified in our core SO202-18-3. Green colors in the NGRIP data mark warm phases that correspond to the laminated sediment intervals of our sediment cores. On the right side the lamination patterns of SO202-18-3 and SO202-18-6 are shown. Bioturbated Facies are shown in white, Bioturbated Lamination Facies in brown, Layered Facies in yellow, Laminated Facies in green, and ash layers in black. Black lines between both cores mark correlated sediment facies. Laminated sections that were correlated with warm periods in the ice core record are labeled TI-BLU1–5. Asterisk mark <sup>14</sup>C dates, and numbers and vertical black lines inside the lamination patterns give results of laminae counts for the TI-BLU intervals. Additionally, laminae countings of a short cold period between TI-BLU4 and TI-BLU3 that was identified during our correlation are given in italic letters (see text for details). Note that thick laminated sections do not occur after 11 ka. (All.): Allerød; (IACP): Intra-Allerød Cold Period; (HS1): Heinrich Stadial I; (OD): Older Dryas.

sequences containing additional Layered Facies sediments, while the Bioturbated Laminae Facies as well as the Bioturbated Facies intermittently occur in both cores between

TI-BLU1 and TI-BLU3 (Fig. 8). Small intervals of laminated sediments occur also above TI-BLU1; however we did not include those parts as separately numbered units in the deglacial terminology presented here. At the current stage, we decided to concentrate our efforts on the deglacial section of our record.

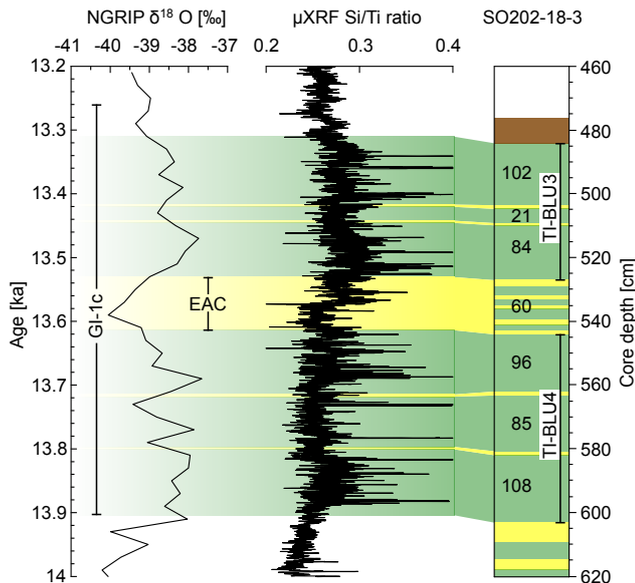
### 4.3 A floating laminae-counted chronology anchored to the NGRIP δ<sup>18</sup>O time series

Based on our initially established radiocarbon-anchored and regionally correlated stratigraphic framework for site SO202-18-3/6, we performed a detailed counting of the lamination pattern in the deglacial section of site SO202-18-3/6, assuming one laminae couplet represents 1 year of deposition. Following this assumption we refer to a laminae couplet as a “varve” hereafter. We compared the occurrences of the identified five TI-BLU sections, and the number of varves counted therein, to the NGRIP oxygen stable isotope record (20 yr resolution) as a Northern Hemisphere climatic reference record. For this exercise, we used two anchor dates for the marine-ice-core correlation: (1) the onset of laminations (TI-BLU5) at the start of the Bølling (or GI-1e) and (2) the re-establishment of laminations at the Younger Dryas–Holocene boundary (onset TI-BLU1). In addition, we follow the established assumption that laminated sediments correspond to the warmest interstadial phases (e.g., Behl and Kennett, 1996; Rella et al., 2012). In contrast, layered and bioturbated sediments correspond to colder periods.

The results of the varve counts in the sections TI-BLU1–5 are illustrated in Fig. 8. Our observed alternation of laminated TI-BLU intervals and Layered or Bioturbated Facies sections closely resemble submillennial-scale warm–cold oscillation originally identified in the Greenland ice core δ<sup>18</sup>O record and described in the deglacial chronology by the INTIMATE working group (Rasmussen et al., 2006; Blockley et al., 2011). The number of counted varves seemingly corresponds to time intervals that lie above a certain minimum in the δ<sup>18</sup>O NGRIP data of around 39 ‰ (Fig. 8). In the following, we compared our deglacial time series of Bering Sea events with the respective Greenland sequence of events (GI-1e to GS-1/Holocene). From the older towards the younger section of our cores we counted the following intervals.

Section TI-BLU5 consists of 556 ± 6 varves. Based on our defined first chronostratigraphic anchor point, these TI-BLU5 varves correspond to the Bølling (GI-1e, Fig. 8), which comprises 617 yr on the GICC05 timescale. This ca. 90 % temporal coverage argues for the occurrence of annually deposited, laminated sediments at our site. Based on the assessment that cold intervals are not laminated, the following mainly Layered Facies section between TI-BLU5 and TI-BLU4 would then be assigned to the small cooling of the Older Dryas (GI-1d) in the GICC05 chronology.

Section TI-BLU4 consists of 289 ± 3 varves and correlates to a first warm phase in the NGRIP Allerød (GI-



**Figure 9.** Detailed lamination pattern of core SO202-18-3 covering the Allerød (GI-1c) interval with a correlation with NGRIP ice core data. Left panel: NGRIP 20 yr average  $\delta^{18}\text{O}$  record. EAC marks the “Early Allerød Cooling”, a small cooling event not named in the GICC05 chronology. Middle panel: micro-XRF Si/Ti ratios as a proxy for productivity, with higher ratios indicative of higher productivity. Right panel: Bering Sea lamination pattern with varve counts. Bioturbated Facies is shown in white, Bioturbated Lamination Facies in brown, Layered Facies in yellow, and Laminated Facies in green as in Fig. 8.

1c), ranging from 13.90 to 13.62 ka, thus indicating a nearly complete overlap with our laminae counts. While no formal subdivision has been established within the Allerød (GI-1c) so far (e.g., by the INTIMATE working group; Blockley et al., 2011), we partition this time interval as we can identify short cold periods within this warm phase, which are imprinted as Layered Facies intersections in otherwise Laminated Facies parts in our core and can be correlated with short cold spells in the 20 yr averaged NGRIP  $\delta^{18}\text{O}$  time series. The following sequence between TI-BLU4 and TI-BLU3 contains several Layered Facies intervals and is assigned to a short, intra-GI-1c cooling period in the NGRIP  $\delta^{18}\text{O}$  record that lasts about 85 yr (13.62–13.53 ka) and which we tentatively call Early Allerød Cooling (EAC, Fig. 9). Despite the intermittent accumulation of Layered Facies sediments within the EAC, we counted  $60 \pm 1$  varves and thus cover 80 % of the potential reference time. The detailed inspection of TI-BLU4 before the EAC and TI-BLU3 after the EAC reveals that these two intervals are both interrupted by multiple, thin (i.e., less than 2 cm thick) Layered Facies sediment sections at ca. 13.80, 13.72, 13.44, and 13.42 ka (Fig. 9). Based on our varve counts these short intervals correspond to rapid, inter-decadal cooling intervals in GI-1c with durations of less than 60 yr each. This lamination

pattern is surprisingly similar to the distant NGRIP record, in particular with regard to the bi-decadal resolution of the ice core record’s  $\delta^{18}\text{O}$  time series. This might also be a reason for the slight differences between the temperatures decreases in NGRIP and the position of these thin Layered Facies sections.

Within the uppermost part of interval TI-BLU3 (Fig. 8) the effect of downward bioturbation slightly decreases the precision of laminae counts. In total,  $207 \pm 2$  varves were counted in this interval. However, when correlated with the upper part of the Allerød (GI-1c), ranging from 13.53 to 13.26 ka, our counts still cover more than 77 % of the NGRIP reference time interval under an annual deposition scenario. We acknowledge that varves might have been lost in the counting process due to the upper completely bioturbated boundary. The cold period on top of TI-BLU3 matches the entire Intra-Allerød Cold Period, or IACP (GI-1b). This colder interval in its youngest part shows the first Bioturbated Facies sediments and a short interval of Bioturbated Laminae Facies.

Thereafter, the Laminated Facies section begins, identified as TI-BLU2 (Fig. 8). We counted  $89 (\pm 1)$  varves in this section and assigned these to NGRIP Allerød (GI-1a). Compared to the GICC05 chronology that provides a duration of  $203 \pm 71$  yr for this time interval, our TI-BLU2 section is at face value lacking a considerable part of preserved laminae, or years. To resolve this discrepancy, we consider the following explanation: the TI-BLU2 interval is bound both on the upper and lower boundary by intervals with the Bioturbated Laminae Facies, which we were not able to count reliably. Thus, we presume that our reported varve counts in TI-BLU2 are skewed towards lower numbers than have been deposited originally. In fact, for a “true” count of all originally deposited varves, we would have to add the neighboring upper and lower core intervals of the Bioturbated Laminae Facies (5.43 and 18.75 cm, respectively; cf. Table 3) to our counts in the Laminated Facies section (18.13 cm, Table 3, all values taken from core SO202-18-3). If we roughly extrapolated the potential number of “uncounted varves” from these Bioturbated Laminae Facies based on core depths and average sedimentation rate/thickness of laminae within the Laminated Facies, we would have to add ca.  $119 \pm 12$  varves to our counted 89 varves (in depth interval 421.18–440.78 cm, Table 3, core SO202-18-3), thus bringing the total of originally deposited varves to  $208 \pm 13$ . This value would then be in excellent agreement with the GICC05-based duration of 203 yr for GI-1a. As a result, we are rather confident that the proposed annual sedimentation regime for the laminae persisted throughout the entire warm phases of the Bølling–Allerød (GI-1). Moreover, a potentially slightly diachronous GS-1 inception as shown by differing proxy time series within the NGRIP ice core might hamper a direct annual one-to-one correlation in this particular time interval (e.g., Ruth et al., 2007; Steffensen et al., 2008). The subsequent interval between TI-BLU2 and TI-BLU1 corresponds to the Younger Dryas (GS-1), and the occurrence of the Bio-

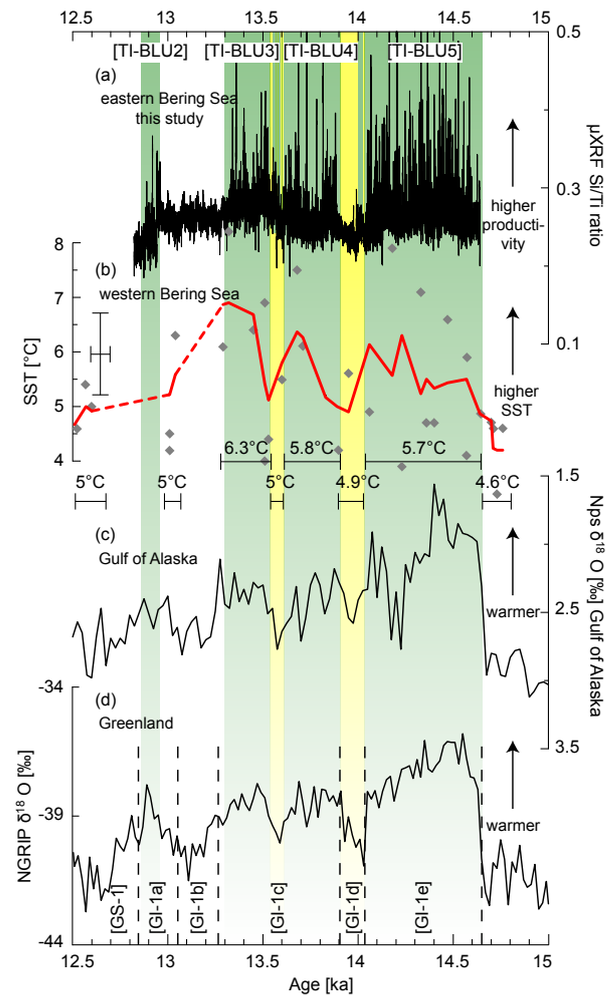
turbated Facies during this cold interval prevented laminae counts.

For the onset of TI-BLU1 we assume an age of 11.65 ka, based on the GICC05 chronology (Fig. 8), in line with our defined anchor point. For TI-BLU1 (11.65–11.8 ka) we, unfortunately, cannot establish a clear upper chronostratigraphic boundary, because cold and warm oscillations identified in the NGRIP ice core during the Preboreal are at present difficult to reconcile with our Bering Sea time series (Fig. 8). We note, however, that neither are all early Holocene climatic oscillations unambiguously recorded in all Greenland ice cores (Rasmussen et al., 2007). Thus, we defined the upper end of TI-BLU1 through laminae counts with a total number of  $471 \pm 4$  varves (Fig. 8).

Taken together, our laminae counts show strong support for the hypothesis that laminae are deposited annually. As a result, our achieved correlation allows for the defining of sections which are directly tied to GICC05-derived calendar ages. These independent calendar ages deduced by correlation with the NGRIP record can in turn be used to estimate regional surface reservoir ages (i.e.,  $R$ ) for radiocarbon dating (Table 2). Our calculated reservoir ages range within 730–990 yr in the Bølling–Allerød (mean reservoir age: 875 yr), 800–1100 yr in the Younger Dryas (mean reservoir age: 910 yr) and 765–775 yr in the Preboreal (mean reservoir age: 770 yr) (cf. Table 2, directly calculated  $R$  values in bold/italic letters). These are qualitatively consistent with earlier studies that suggested varying reservoir ages in the North Pacific during the last deglaciation (Gebhardt et al., 2008; Sarnthein et al., 2007). Our calculated values for local reservoir ages are in line with studies that inferred surface reservoir ages close to 730 yr during the last deglaciation, with variations of about  $\pm 200$  yr in the eastern North Pacific (Lund et al., 2011), and with reservoir age estimates of 700–800 yr for the British Columbia coast (Southon and Fedje, 2003). However, we note that regional reservoir ages reconstructed for the eastern North Pacific and western North Pacific during the Bølling–Allerød (GI-1) derived by the “plateau tuning method” (Sarnthein et al., 2007) are younger than our results – with 440 and 720 yr, respectively – pointing to the possibility of distinct regional or even localized oceanographic reorganizations in the upper and mid-depth water column during the last deglaciation (Sarnthein et al., 2013).

#### 4.4 Teleconnections and forcing mechanisms for rapid climate signal transfer

As a result of the close correspondence between the counted lamination patterns and the Greenland ice core data (Figs. 8, 9), we assume that the preservation of laminations is tied to warm surface temperature intervals concomitant with higher export production, superimposed on the already-warm background temperature within GI-1. This co-occurrence of higher export production with warmer temperatures is also



**Figure 10.** Comparison of laminated core SO202-18-3 with Bering Sea SST records, planktic  $\delta^{18}\text{O}$  data from the Gulf of Alaska, and Greenland temperature changes across the Bølling–Allerød time interval. (a) Micro-XRF Si/Ti ratios of SO202-18-3. (b) In red the stacked SST record (three-point running mean) of cores SO201-2-77, SO201-2-2-85, and SO201-2-101 from the Shirshov Ridge, western Bering Sea. The gray squares depict the individual SST measurements (Max et al., 2012). Dashed parts in the SST record indicate intervals with insufficient temperature data. The age model used in this plot is the published model of Max et al. (2012). The numerical temperatures below graph (b) are the calculated mean SSTs from the marked warm and cold intervals, and the horizontal bars indicate the temporal range of the average SST. The error bars on the left side of (b) show the uncertainties of SST measurements and radiocarbon datings. (c)  $\delta^{18}\text{O}$  data of planktic foraminifera *Neogloboquadrina pachyderma sinistral* from the Gulf of Alaska as a proxy of long-term changes in annual average SST (Praetorius and Mix, 2014). Note that we used the age model after Praetorius and Mix (2014). (d) NGRIP stable isotope data as reference record. The occurrence of mainly Laminated Facies, Layered Facies, and Bioturbated Facies in SO202-18-3 is shown in green, yellow, and white, respectively. Laminated sections in SO202-18-3 are labeled TI-BLU as in the text.

evident on decadal timescales by a slight decline of the Si/Ti ratios in our proposed EAC (Fig. 9) and on centennial scales during the Older Dryas (GI-1d) and IACP (GI-1b; Fig. 10a). The Si/Ti ratios are robust indicators of bio-siliceous export production, as they are not susceptible to post-depositional alteration by sedimentary redox conditions.

To further test the relationship between lamination occurrence and SST, we used lower-resolved alkenone-based SST reconstructions of three cores from the Shirshov Ridge in the western Bering Sea (Max et al., 2012) to infer SST information at least on a basin scale. We used these SSTs as no data are available directly from Laminated and Layered Facies sediment intervals of our sites. We are aware that the alkenone data may represent a seasonal signal, most likely a summer temperature signal (Seki et al., 2007). We merged the individual site reconstructions presented in Max et al. (2012) into one stacked, three-point running mean SST record (Fig. 10b) to consolidate the regional signal. Hereby, we used the published age model of Max et al. (2012). This age model is mainly based on high-resolution scans of color  $b^*$  values, a proven proxy for biogenic productivity variations (Nürnberg and Tiedemann, 2004). The close correlation between high color  $b^*$  values and lamination occurrence during the last deglaciation was also shown for nearby core PC23A (Kim et al., 2011). The standard error of the originally used SST calibration is  $\pm 1.5^\circ\text{C}$  (Max et al., 2012). Despite the lower resolution, these SST data indicate corresponding temperature variations throughout the Bølling–Allerød (GI-1), and SST maxima broadly correlate with the occurrence of laminated intervals and warmer Greenland air temperatures. The stacked SST record from the Bølling–Allerød (GI-1) reveals temperature differences between the cold phases of the Older Dryas (GI-1d), the small EAC as well as the IACP (GI-1b), and the warm laminated periods. Mean SSTs during the colder Bølling–Allerød intervals in the Bering Sea were around  $5^\circ\text{C}$ , and increased to  $6^\circ\text{C}$  in the warmer phases (Fig. 10b). Further support comes from a stacked  $\delta^{18}\text{O}$  record of the planktic foraminifera *Neogloboquadrina pachyderma* sinistral of the Gulf of Alaska (GoA), which was interpreted as a proxy for changes of annual SSTs (Fig. 10c; Praetorius and Mix, 2014). This record with decadal-scale resolution clearly shows the warm–cold oscillations during the Bølling–Allerød (GI-1) that are visible in our lamination sequence and in the NGRIP record (Fig. 10d). We observe a close correspondence between the GoA and our Bering Sea record even for short-term events like the observed EAC event, reflected in higher  $\delta^{18}\text{O}$  GoA values (i.e., lower SSTs). These temperature trends from several locations in the North Pacific in principle support our assumption that anoxia occurrence is connected to warmer SSTs; however, additional annually resolved SST proxy time series from our sites would be needed to prove a causal relationship between higher SSTs and the occurrence of anoxia.

The clear imprint of even such short time periods like the low-amplitude EAC between 13.62 and 13.53 ka and the

Older Dryas in the Bering Sea lamination pattern strongly argues for a close atmospheric coupling between the North Atlantic and the North Pacific without significant time lags. Several previous works have provided evidence for millennial-scale variations in Bering Sea biogenic productivity and upper-ocean characteristics during the last glacial termination that occur in phase with both Greenland and east Asian paleoclimatic reference records (Rella et al., 2012; Itaki et al., 2009; Max et al., 2012; Caissie et al., 2010; Riethdorf et al., 2013b). This in-phase behavior of laminated sediment deposition in warm phases of the deglacial Bølling–Allerød (GI-1) interstadial is also known from several locations around the North Pacific Rim (Jaccard and Galbraith, 2012), albeit not on shorter decadal timescales as implied by our study. The oceanographic changes recorded in our sediment core occur with rapidity, comparable to the fast climate changes reported from ice cores (Taylor et al., 1997; Steffensen et al., 2008) or varved lake sediments (Kossler et al., 2011; Brauer et al., 2008).

In the past, studies using general circulation models investigated the teleconnection between the North Atlantic and the North Pacific and possible linkages to changes in the Atlantic Meridional Overturning Circulation (e.g., Mikolajewicz et al., 1997; Vellinga and Wood, 2002; Okumura et al., 2009). In these numerical model experiments cold periods like Heinrich Stadial 1 and the Younger Dryas are simulated by freshwater forcing in the North Atlantic, which leads not only to a weakening of the overturning cell but also to significantly colder temperatures over both the North Atlantic and North Pacific regions. However, while the response over the North Pacific is common in many models, the transfer mechanisms are less well understood, especially on decadal timescales. One important climate determinant for variations in the Bering Sea ocean–atmosphere system is the Aleutian Low as a main action center in the circulation of the Northern Hemisphere (Rodionov et al., 2007). Changes in the strength and location of this system have a direct effect on winter air and sea surface temperatures and thus sea ice occurrence, which in turn influence the strength of the phytoplankton bloom season. Several studies suggest a strong Aleutian Low is associated with warm winters in the Bering Sea (Niebauer, 1983), and that winter temperatures in the Bering Sea are sensitive to the position of the Aleutian Low (Rodionov et al., 2007). It has further been shown that the Aleutian Low varies on decadal timescales and is also associated with variability of the Arctic Oscillation (Overland et al., 1999). We suggest that decadal changes in position and strength of the Aleutian Low lead to colder and warmer winters on decadal timescales. The colder (warmer) winters are connected with increased (decreased) sea ice cover, which is highly sensitive to temperature changes and fosters decreased (increased) diatom blooming events.

#### 4.5 Characteristics and causes of millennial- to decadal-scale OMZ development

The occurrence of laminations serve as a direct indicator for oxygenation decreases below the threshold for most marine benthic life and the development of severe and persistent anoxia in Pacific intermediate water masses. We follow previous works (Behl and Kennett, 1996) in assigning different oxygen concentrations to the different facies in our record. These are as follows:  $< 0.1 \text{ mL L}^{-1} \text{ O}_2$  excluding any burrowing macrofauna for Laminated Facies;  $0.1 \text{ mL L}^{-1} \text{ O}_2$  for Layered Facies, which allows only meiofaunal bioturbation that just diffuses the laminations on millimeter scales but does not lead to deeper bioturbation by larger organisms;  $0.1\text{--}0.2 \text{ mL L}^{-1} \text{ O}_2$  for Bioturbated Laminae permitting partial homogenization; and  $> 0.3 \text{ mL L}^{-1} \text{ O}_2$  for the Bioturbated Facies allowing bioturbating macrofauna (Behl and Kennett, 1996).

##### 4.5.1 Deglacial strengthening and Holocene decline of anoxia in the Bering Sea

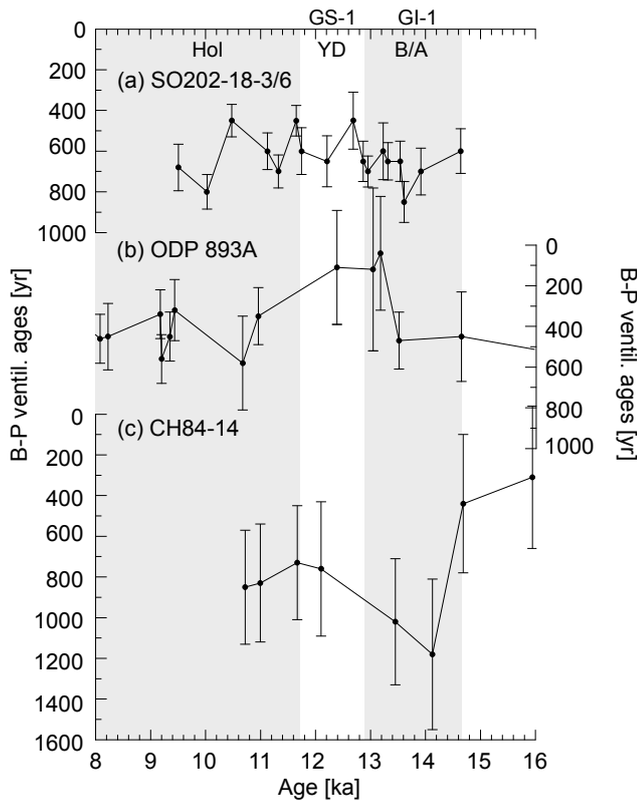
Our facies analysis reveals millennial-scale changes between stadial, well-oxygenated (bioturbated) phases and interstadial, mainly anoxic (laminated/layered) phases during the last glacial termination, consistent with recent assessments (Jaccard and Galbraith, 2012). In our records, Bioturbated Facies occur on millennial scales during the cold stadial Younger Dryas (GS-1) phase and the relatively cold IACP (GI-1b), indicating improved oxygen concentrations in the bottom water at our site. These periods are contrasted with the two interstadial dysoxic–anoxic sediment facies during the entire Bølling (GI-1e)–Allerød (GI-1c) and Preboreal (Fig. 8).

Notably, superimposed on these interstadial warm phases we recorded multiple, rapid facies changes on inter-decadal timescales between Layered and Laminated Facies (Fig. 9). These changes exhibit a shorter-term, subtler process that modulates the mid-depth deglacial Bering Sea oxygen concentration. Though Layered Facies do not show a clear expression of alternating seasonal laminae like in laminated sections, they are devoid of major bioturbation, thus evidencing persistent anoxic to dysoxic conditions. These patterns lead us to distinguish between millennial-scale larger oxygenation changes (between oxic and dysoxic–anoxic) and superimposed decadal-scale smaller fluctuations (between dysoxic and anoxic) as distinct patterns in OMZ strength.

The Laminated and Layered Facies disappear after the Preboreal in the Holocene, indicating both a vertically contracting and abating OMZ in the Bering Sea compared to deglacial conditions (Fig. 8). As the increasing NGRIP  $\delta^{18}\text{O}$  data would suggest ongoing deposition of laminations, changes in the environmental background conditions must have been distinct. Multiple causes are likely. Firstly, alkenone-based SST data show a temperature decrease of

more than  $3^\circ\text{C}$  in the Bering Sea throughout the early Holocene, which might lead to reduced phytoplankton productivity due to shorter and colder summers (Caissie et al., 2010; Riethdorf et al., 2013a). However, SSTs are still higher compared to the Bølling–Allerød, pointing to additional factors besides SSTs that might influence the Holocene OMZ development. Secondly, another potential cause could be a change in the characteristics of mid-depth water masses at our sites in the Bering Sea, either in age (and thus nutrient inventories) or in oxygenation compared to the mid-late Holocene. Published ventilation ages reveal no clear Holocene changes in the North Pacific (Duplessy et al., 1989; Ahagon et al., 2003), and the occurrence of oxygen-poor, Layered Facies together with rather unchanged B–P ventilation ages suggests that oxygen depletion of mid-depth waters is not caused by the presence of upwelled older, oxygen-poor North Pacific deep-water masses (cf. 4.5.2). In support of this notion, evidence from Gulf of Alaska sites that lie in the upstream part of waters entering the Bering Sea show oxygenation increases from anoxic–suboxic towards suboxic–oxic conditions around 10.9–9.2 ka (Addison et al., 2012; Davies et al., 2011), broadly in line with ceasing anoxia at our site SO202-18-3/6.

An additional factor for the establishment of higher oxygenation potentially is the opening of the Bering Strait at around 11 ka (Elias et al., 1996; Hu et al., 2010). Today, the northward annual through-flow of relatively warm, nutrient-rich North Pacific water into the Arctic Ocean is around 0.8 sverdrup (Woodgate et al., 2010). Roughly, during the time of the Bering Strait opening at 11 ka the occurrence of Layered Facies increases until laminations finally disappear at 10 ka. As Layered Facies represent slightly higher oxygen concentrations compared to laminated sequences, this change might represent a trend towards decreasing productivity, beginning with the opening of the Bering Strait. Likely, the effect of this gateway opening was not instantaneous but rather steadily increasing and amplified with ongoing sea level rise. Today the gateway influences the lower-productivity ecosystems of the Chukchi and Beaufort seas with the advection of high nutrient Pacific waters (Grebmeier et al., 2006). As it has further been suggested that higher primary production in the Chukchi Sea also during the Holocene could result from the greater nutrient supply from the Bering Sea (e.g., Keigwin et al., 2006), we suggest the following mechanism: before 11 ka Bering Sea surface ocean circulation was substantially different from modern patterns, with an absent or weakened Anadyr Current and differing expression of the Bering Slope Current, also leading to different interactions with the Alaskan Stream and the open North Pacific. These changes potentially caused a more intense gyre circulation within the Bering Sea, enhanced trapping and recirculation of nutrients and thus higher productivity. We further suggest that the opening of this shallow gateway also led to a decline in the surface stratification as warm, nutrient-rich surface waters were transported to the Arctic Sea. Such



**Figure 11.** Compilation of  $^{14}\text{C}$  paleo-ventilation ages (a) for core SO202-18-3/6, derived from benthic–planktic foraminifera age differences compared to results from (b) the Santa Barbara Basin (Ingram and Kennett, 1995; Roark et al., 2003) and (c) off Japan (Duplessy et al., 1989). Error bars show radiocarbon dating errors. For this comparison, planktic  $^{14}\text{C}$  ages of Japan and the Santa Barbara Basin were recalculated using the new INTCAL13 calibration curve and reservoir ages of Sarnthein et al. (2007) for the Santa Barbara Basin. Hol: Holocene; YD: Younger Dryas; B/A: Bølling–Allerød

a trend for declining stratification after the Preboreal in the Bering Sea can be tentatively assumed based on differences between alkenone and Mg/Ca-based SST starting at about 9 ka (Riethdorf et al., 2013a). As a second factor, the coastline retreated significantly further away from the core position during the flooding of the shallow northern shelf areas in connection with ongoing sea level rise. Thus, the influence of the fluvial input, delivering nutrients and enhancing surface stratification, diminished.

#### 4.5.2 Millennial-scale changes in intermediate water ventilation

To identify the potential impact of North Pacific intermediate water circulation and ventilation changes on longer-term, millennial-scale oxygen variations on our sites, we calculated benthic–planktic ventilation ages (Fig. 11). In principle, older B–P ages could indicate the transport of higher-nutrient, lower-oxygen water masses, such as North Pacific

Deep Water (NPDW) to the Bering Sea and their entrainment into the upper, mid-depth waters, thereby supporting OMZ strengthening. However, our B–P ages show only moderate oscillations throughout the deglacial sequence, with values between 470 and 700  $^{14}\text{C}$  yr, all significantly below values reported for Pacific Deep Water of  $1530 \pm 230$   $^{14}\text{C}$  yr (Lund et al., 2011). Additionally, B–P age oscillations at our sites do not reflect stadial–interstadial changes in ventilation and are with variations of 100–200 yr substantially smaller in amplitude than those reported from distant, more southern mid-depth locations, e.g., off Japan and the Santa Barbara Basin for the same time interval (Fig. 11; cf. Duplessy et al., 1989; Ahagon et al., 2003; Roark et al., 2003; Kennett and Ingram, 1995). Note that for this comparison planktic  $^{14}\text{C}$  ages of Japan and the Santa Barbara Basin were recalculated using the new INTCAL13 calibration curve and reservoir ages of Sarnthein et al. (2007) for the Santa Barbara Basin. One potential reason for the small oscillations could be that the Bering Sea is remote from the formation area of the most prominent mid-depth water mass, North Pacific Intermediate Water (NPIW), which has a strong influence on the ventilation off Japan. Our B–P ventilation age data do not support a collapse of subarctic North Pacific mid-depth water circulation during GI-1 and the Preboreal. As a result, we do not consider that physical forcing leading to NPIW formation rate or source region changes play a decisive role in the development and vertical expansion of interstadial anoxia at mid-depth in the Bering Sea.

However, for the Gulf of California and the Santa Barbara Basin it was suggested that phases of oxic and dysoxic conditions were controlled by changes in oxygen concentration of upper intermediate water (Kennett and Ingram, 1995; Hendy et al., 2002). Based on analyses of laminated sediments from the North American continental margin, Zheng et al. (2000) suggested suppressed ventilation at higher latitudes of the North Pacific during the Bølling–Allerød. In line with these assumptions, epibenthic  $\delta^{13}\text{C}$  data from the Alaska Margin (Davies et al., 2011) and from the modern formation regions of NPIW (Max et al., 2014) show major decreases in mid-depth oxygenation during the Bølling–Allerød. We thus assume that changes in NPIW oxygen concentrations are an important factor for conditioning the mid-depth waters in the Bering Sea on millennial timescales. Oxygen decreases can be caused by high export production and subsequent higher mid-depth remineralization of exported organic matter along the pathway of NPIW in the Pacific subarctic gyre without significant changes in the formation rate of NPIW (Crusius et al., 2004). This scenario would be in line with our radiocarbon B–P age ventilation data and is supported by a number of studies indicating widespread export productivity peaks during the Bølling–Allerød and the Preboreal throughout the North Pacific (Brunelle et al., 2010; Kohfeld and Chase, 2011), combined with reductions in oxygen concentrations along the pathway of NPIW throughout the North Pacific (Hendy and Pedersen, 2006; McKay et al., 2005).

#### 4.5.3 Modulation of OMZ strength during the Bølling–Allerød (GI-1) and Preboreal

Increased export production through higher biogenic productivity leads to an intensification of the OMZ through organic matter degradation. Generally, higher primary productivity is indicated by various productivity proxies. Increased Ca/Ti ratios (Fig. 4) and carbonate mass accumulation rates (Fig. S3) might be caused by higher carbonate production through coccolithophorids (Okazaki et al., 2005; Khim et al., 2010). Further, the low Ti concentrations during the laminated intervals, higher biogenic opal mass accumulation rate (Fig. S3), and high Si/Ti ratios in the biogenic laminae (Fig. 7) point to increased seasonal export production. Regionally increased bio-siliceous productivity during the Preboreal and Bølling–Allerød was also shown for the southern Bering Sea (Gorbarenko et al., 2005) and in nearby, partly laminated, core PC23A (Fig. 1; Khim et al., 2010; Katsuki et al., 2014). Several environmental factors can influence the biological productivity and OMZ development.

1. Higher bottom water temperatures at our site during the warmer Bølling–Allerød phases would increase the remineralization of organic matter at the seafloor, as metabolic rates of organisms increase with higher temperatures (Matsumoto et al., 2007). Such a process could work independently of surface ocean processes as a positive feedback, contributing to elevated oxygen consumption in intermediate waters. Unfortunately, regional bottom water temperature reconstructions from North Pacific sites are to our knowledge not available to provide information on the importance of such a process. However, a qualitative assessment of potential temperature changes based on published benthic oxygen isotope records from comparable water depths shows no unambiguous trends. On Shirshov Ridge, values show a glacial–interglacial amplitude change of 1.95 ‰ (Riethdorf et al., 2013), and in the eastern Bering Sea of ca. 1.4 ‰ (Rella et al., 2012).
2. During peak glaciation, the biological productivity was restricted by a shorter and colder summer that led to the spatial extension of sea ice (Takahashi, 2005; Nakatsuka et al., 1995). Higher temperatures during the Bølling–Allerød and since the onset of the Holocene (Max et al., 2012; Riethdorf et al., 2013a) would have supported an earlier retreat of sea ice and a change from longer and more pronounced, sometimes multi-year sea ice cover during cold phases like Heinrich Stadial 1 and Younger Dryas to shorter, seasonal sea ice seasons. This is in line with evidence based on diatom assemblages from the Umnak Plateau (Caissie et al., 2010), which indicates during the Bølling–Allerød a shift from perennial to seasonal sea ice and a pattern similar to the ice-related bloom in the marginal ice zone on the Bering Shelf today.

3. Regionally elevated rates of primary productivity ultimately require high-nutrient surface water concentrations. On the NE Bering Sea margin, these can be delivered through several sources. High meltwater input by fluvial runoff from retreating North American hinterland glaciers would provide additional nutrients through lateral transport of terrigenous material, most likely delivered during warmer episodes (Sancetta et al., 1984; Itaki et al., 2009). In addition, nutrient increase was likely enhanced to a significant extent by the deglacial sea level rise, leading to the flooding of the previously exposed, wide, shallow proximal northern Bering Sea shelf. This process would have provided additional nutrients compared to the modern situation, as suggested for similar settings with laminated sediments from the southeast Alaska Margin (Davies et al., 2011). Another process influencing the amount of biological productivity is light, depending on upper ocean stratification, which can constitute the limiting factor in biogenic productivity especially in high latitudes (Kohfeld et al., 2005). Before the Bølling–Allerød, enhanced upper-ocean mixing potentially increased light limitation by a deeper mixed layer. With the onset of the Bølling, input of meltwater from the North American ice sheets stratified the upper water column and fostered biological productivity (Lam et al., 2013; Katsuki et al., 2014) as it would enable phytoplankton to prolong their growth season and better use the more abundant bio-available nutrients in the upper photic zone (Niebauer et al., 1995; Niebauer and Alexander, 1985). Such changes in freshwater-induced stratification might also work fast enough to explain the observed decadal-scale correspondence of fine laminations to higher temperatures (Fig. 10).

In summary, we presume that major forcing of OMZ was likely a combination of sea ice changes, with less sea ice during the warm phases and variations in ocean stratification with a more stratified upper water due to freshwater input.

However, the anoxic–dysoxic bottom waters during the Older Dryas (GI-1d) and the EAC did not ameliorate to a point that would permit strong bioturbation, but just enabled the observed meiofauna-induced dispersal of annual layer structures on millimeter scales. Had instead the former been the case, the layered structure of the sediment would have been destroyed by larger-scale macrofauna bioturbation. A further increase in oxygenation, despite the lower productivity in these cold phases, was likely prevented by the still comparably low oxygen concentrations of mid-depth water entering the Bering Sea.

Taken together, we see two major processes that contribute to the strengthening of the OMZ in the Bering Sea and thus the formation of laminated sediments. Based on rather constant ventilation ages, we observe millennial-scale changes in the NPIW oxygen concentrations

without significant variations in the formation rate of NPIW. This millennial-scale pattern, however, apparently did not drive oxygen concentrations below the critical threshold for anoxic conditions and lamina formation (i.e.,  $< 0.1 \text{ mL L}^{-1}$ ). Oxygen concentrations decreased to that extent only in combination with higher regional export production and subsequent remineralization of organic matter during the warm phases of the Bølling–Allerød. Our results corroborate earlier hypotheses that argued for oxygen drawdown in the North Pacific by a combination of higher organic matter export and lower  $\text{O}_2$  concentrations at intermediate waters (Zheng et al., 2000; Crusius et al., 2004). Other possible forcing mechanisms connected to indices like the Pacific Decadal Oscillation or to solar forcing are currently under discussion (e.g., Katsuki et al., 2014) and at present cannot be corroborated based on the data presented in this story.

## 5 Summary and conclusions

Two laminated cores from the northern Bering Slope located in intermediate water depths allow for a tight coupling to Greenland ice core data on decadal timescales, especially for the deglacial Bølling–Allerød phase (GI-1, 14.64–12.85 ka). By correlating a suite of laminated sediment intervals to NGRIP oxygen isotope data, we established an age model partly independent from radiocarbon dating, which in turn was used to calculate marine reservoir ages. Mean reservoir ages are 875 yr for the Bølling–Allerød (GI-1), 910 yr for the Younger Dryas (GS-1), and 770 yr for the Preboreal. The correlation and laminae counts implied the presence of annually layered sediment sections (varves). The combined information from sediment facies analysis and micro-scale high-resolution XRF scanning showed that the laminations represent an annual alternation of high bio-siliceous spring/summer blooming events driven by the sea ice retreat and a rather siliciclastic background sedimentation.

Our results show for the first time that the mid-depth water column in the Bering Sea reacted quickly on inter-decadal timescales to changes in upper-ocean characteristics, particularly in regard to sea ice dynamics, surface water stratification, and sea surface temperatures. Potential implications for the future arise from these observations of rapid changes. It is unclear whether oxygen concentrations could drop again to anoxic levels if temperature increases and sea ice decreases continue along present trends (e.g., Brown and Arigo, 2012; Grebmeier, 2006), leading to the expansion of subarctic Pacific “death zones”. This would create significant consequences and large-scale ecosystem readjustments.

On millennial, or stadial to interstadial, timescales relatively constant benthic–planktic ventilation ages reveal that the OMZ development is more likely caused by basin-wide mid-depth remineralization of organic matter in intermediate waters, rather than changes in formation rate or sources of intermediate water masses. On decadal timescales, the for-

mation of laminated sediments during the Bølling–Allerød seems to be effectively coupled to warm temperatures, as expressed in NGRIP ice core  $\delta^{18}\text{O}$  and Bering Sea SST data. This tight correlation with warm phases during the Bølling–Allerød in the ice core data implies a close atmospheric teleconnection to North Atlantic and Greenland climate. During these warm intervals, rapid sea ice and temperature-induced decadal–interannual-scale export productivity changes lead to fine modulations of OMZ strength between anoxic and dysoxic conditions. Comparison with other laminated cores from the Bering Sea shows that during the last deglaciation the OMZ expanded to greater water depths, possibly of more than 3000 m at some locations. The disappearance of the laminations during the Holocene might be coupled to the opening of the Bering Strait.

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