

Characteristics of colored dissolved organic matter (CDOM) in the Western Arctic Ocean: relationships with microbial activities

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62 Abstract

Colored dissolved organic matter (CDOM), a significant fraction of dissolved organic carbon 63 (DOC), plays various roles in physical and biogeochemical processes in natural waters. In the Arctic 64 65 Ocean, CDOM is abundant because of major input by large rivers. To better understand the processes that drive variations in CDOM, light absorption coefficients of CDOM $[a_{CDOM}(\lambda), m^{-1}]$ 66 67 were extensively documented together with temperature, salinity, chlorophyll a, nitrate concentrations, and bacterial production (BP) and abundance (BA) in the Western Arctic Ocean 68 69 (WAO) from early to late summer as part of the MALINA and the ICESCAPE expeditions. The 70 data set covered contrasting situations, from bloom to post-bloom conditions and from river-71 influenced to oceanic water masses. While CDOM photobleaching occurred in the surface layer (< 72 20 m), we observed significantly lower spectral slopes for CDOM absorption spectra (S_{CDOM}) in 73 addition to higher $a_{CDOM}(440)$ in the layer below (intermediate layer: 30.7 < salinity < 33.9). In 74 particular, the low S_{CDOM} values were found in the Chukchi Sea and the western part of the Beaufort 75 Sea, which coincided with high BP and BA values. Considering the high primary production 76 observed in these areas during our cruises (Arrigo et al., 2012), we hypothesize that S_{CDOM} 77 variations reflect the degradation of phytoplankton that is associated with heterotrophic bacterial 78 activity. In our datasets, a simple regression analysis showed that S_{CDOM} was significantly correlated

with BP and BA. A principal component analysis further supported this conclusion. From our field observations, it was shown that variations in $a_{CDOM}(440)$ and S_{CDOM} result to a large extent from bacterial activity, at least in the WAO.

82 1. Introduction

83 Examining the budget of dissolved organic carbon (DOC) in the Arctic Ocean is crucial to 84 improving our understanding of modifications in the carbon cycle resulting from ongoing global 85 warming. While this warming likely induces thawing of permafrost containing a huge amount of 86 DOC, which is subsequently delivered by river discharge into the Arctic Ocean (Peterson et al., 87 2002; McClelland et al., 2006; Raymond et al., 2007), the long-term trend in the DOC budget of the Pan-Arctic Ocean has yet to be established. In recent studies, Matsuoka et al. (2013, 2014) 88 developed a semi-analytical algorithm to quantitatively estimate DOC concentrations for Arctic 89 90 coastal waters using satellite remote sensing data, which allows the continuous monitoring of variability in DOC concentrations. In contrast, knowledge about the production and the removal 91 92 processes gained from field observations using traditional methods is limited temporally and geographically (Bussmann, 1999; Garneau et al., 2008; Kirchman et al., 2009; Ortega-Retuerta et 93 94 al., 2012). This prevents us from understanding the balance between these processes.

95 Light absorption by the colored fraction of dissolved organic matter (CDOM) provides useful information about biogeochemical processes (Carder et al., 1989; Nelson et al., 1998; Miller et al., 96 2002; Nelson et al., 2004, 2007; Matsuoka et al., 2012). While microbial activity is highly variable 97 98 in natural environments (e.g., Azam et al., 1983), the link between heterotrophic bacterial 99 production and CDOM absorption was reported in the Sargasso Sea (Nelson et al., 1998). More 100 recently, Matsuoka et al. (2012) suggested that lower spectral slopes of CDOM absorption spectra 101 observed in some Arctic Ocean water masses resulted from heterotrophic microbial activity. 102 However, a direct relationship between CDOM and bacteria has not been well documented.

103 The objective of this study is to examine the relationships between CDOM absorption properties 104 and heterotrophic bacterial production (BP) and abundance (BA) in the Western Arctic Ocean

105 (WAO) as well as their link with environmental variables (i.e., temperature, salinity, and nitrate and106 chlorophyll *a* concentrations).

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108 2. Datasets and methods

109 Data were collected during three cruises in the WAO (Figure 1): the joint France-Canada-USA 110 Arctic campaign, MALINA (30 July to 27 August 2009), and the National Aeronautics and Space 111 Administration (NASA) ICESCAPE cruises in 2010 and 2011 (referred to as ICESCAPE 2010: 15 112 June to 22 July 2010 and ICESCAPE 2011: 25 June to 29 July 2011, respectively). While waters in the Mackenzie shelf-basin area during the MALINA cruise were oligotrophic and typical of post-113 114 bloom conditions (Ortega-Retuerta et al., 2012), waters in the Chukchi-Beaufort Seas during the 115 ICESCAPE cruises were highly productive (Arrigo et al., 2012). To examine the general 116 characteristics in the WAO, these three datasets were combined and used in this study.

117 Temperature and salinity profiles were obtained using a SBE-911 plus (Seabird) conductivity-118 temperature-depth (CTD) probe. Chlorophyll *a* (chl *a*) and phaeopigment concentrations (mg m⁻³) 119 were determined fluorometrically (Holm-Hansen et al., 1965). Nitrate concentrations (NO₃, µmol 120 kg⁻¹) were measured following Grasshoff et al. (1999) for MALINA and Armstrong et al. (1967) for 121 ICESCAPE cruises. Oxygen concentrations (Oxy, µmol kg⁻¹) were measured following Carpenter 122 (1965) with modifications by Culberson et al. (1991).

123 Apparent oxygen utilization (AOU, μ mol kg⁻¹) was calculated by referring solubility of oxygen 124 for well-mixed winter waters having temperature and salinity of -1.8 °C and 31, respectively. These 125 reference data were proven to be valid for western Arctic waters (Matsuoka et al., 2012).

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127 2.1. CDOM absorption

Details of the method used for measurement of CDOM absorption are documented in Matsuoka et al. (2012). Briefly, water samples were filtered using 0.2 μ m pore-size filters immediately after sampling. Absorption coefficients of CDOM ($a_{CDOM}(\lambda)$, m⁻¹) were determined from 200 to 735 nm

131 in 1-nm increments using a liquid waveguide system (UltraPath, World Precision Instruments, Inc.).

132 The spectral slope of $a_{CDOM}(\lambda)$ (S_{CDOM}, nm⁻¹) was calculated by nonlinear regression of the data 133 from 350 to 500 nm (Babin et al., 2003; Matsuoka et al., 2011, 2012).

134 A previous study (Helms et al., 2008) showed that the spectral slopes ($S_{275-295}$ and $S_{350-400}$, nm⁻¹ 135 for both) corresponding to two distinct wavelength ranges (i.e., 275-295 nm and 350-400 nm, 136 respectively) and their ratio (S_R , dimensionless) provide insights into sources (e.g., marine or 137 terrestrial sources; Carder et al., 1989; Nelson et al., 2007) and/or local processes affecting the 138 CDOM distribution (e.g., lateral transport, vertical mixing, photo-bleaching, heterotrophic bacterial 139 alteration; Nelson et al., 1998, 2004; Matsuoka et al., 2012; Yamashita et al., 2013). These slope 140 parameters were calculated for our Arctic datasets by fitting a linear model to CDOM absorption 141 coefficients in the two distinct spectral ranges (Helms et al., 2008).

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143 2.2. Bacterial abundance

144 Heterotrophic prokaryotes, including bacteria and archaea, are abbreviated throughout the 145 manuscript as "bacteria". Samples were fixed with glutaraldehyde (0.25% final concentration) and 146 stored at -80°C until processing. During MALINA, bacterial cells (BA) were counted aboard the 147 ship by flow cytometry using a FACS ARIA (Becton, Dickinson and company) equipped with 488 148 nm and 633 nm lasers and a standard filter setup (Ortega-Retuerta et al., 2012). During ICESCAPE 149 2010, bacterial cells were counted by flow cytometry at the home laboratory using a BD FACS 150 Calibur Flow Cytometer (Becton, Dickinson and company). During ICESCAPE 2011, bacterial 151 cells were counted aboard the ship using an Accuri C6 (Becton, Dickinson and company) equipped 152 with a 488 nm laser. In all cases, samples were thawed and SYBR Green-I was added at a final 153 dilution of 1:10,000. Samples were incubated in the dark for 15 min before analysis. Bacterial cells 154 were identified on a plot of green fluorescence (515-545 nm) versus right angle light scatter (SSC), 155 using the green fluorescence as a threshold parameter. High nucleic acid (HNA) and low nucleic 156 acid (LNA) bacteria were discriminated according to their green fluorescence and counted

- 157 separately (Marie et al., 1997). HNA cells have often been considered as active bacteria (Gasol et al.,
- 158 1999).
- 159
- 160 2.3. Bacterial production

161 Bacterial production (BP) was measured following Ortega-Retuerta et al. (2012). Briefly BP was measured by ³H-leucine incorporation (Smith and Azam, 1992). Samples (1.5 mL in triplicate plus 162 one killed control) were added to sterile microcentrifuge tubes, containing 20-30 nM [4,5-³H]-163 164 leucine. This concentration was sufficient to saturate bacterial leucine uptake (data not shown). 165 Incorporation rates were measured after 2-h incubations at *in situ* temperature, and incubations were 166 stopped by the addition of trichloroacetic acid (5% final concentration). Leucine incorporation rates 167 were converted to carbon production using the conversion factor of 1.5 kg C produced per mole of 168 leucine incorporated (Kirchman, 1993), considering no isotope dilution.

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170 2.4. Statistical analyses

171 2.4.1. Regression analysis

To examine a direct relationship between two variables, several regression analyses were performed. Because the two variables were random (or not controlled), Model II regression was applied in this study (Legendre and Legendre, 1998).

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176 2.4.2. Principal component analysis

For a dataset containing several variables, a multiple regression analysis is not always the best method for examining the relationships among the variables. Principal component analysis (PCA) is preferable because it summarizes, in a few dimensions, most of the variability present in a dispersion matrix of a large number of variables (Legendre and Legendre, 1998), and has been applied to a large number of oceanographic studies (e.g., Legendre and Legendre, 1998; Uitz et al., 2008; Suzuki et al., 2012). We thus applied this method to our datasets.

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184 2.5. Sea ice concentration

185 Daily sea ice concentrations data acquired by the Defense Meteorological Satellite Program 186 (DMSP) SSM/I passive microwave sensor (25-km spatial resolution) were downloaded from the 187 National Snow and Ice Data Center (NSIDC) at 188 ftp://sidads.colorado.edu/pub/DATASETS/nsidc0051_gsfc_nasateam_seaice/. Daily images (29 for 189 MALINA, 38 for ICESCAPE2010, and 35 for ICESCAPE2011) were averaged to generate an 190 image of mean sea ice concentrations for each cruise (Figure 2). crile

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3. Results and discussion 192

193 3.1. Vertical distribution of CDOM

194 To examine the vertical distribution of CDOM absorption properties in the WAO, a_{CDOM}(440) and S_{CDOM} were plotted against depth (Figure 3). At depth < 150 m, a_{CDOM} (440) values were highly 195 196 variable for early to middle summer (ICESCAPE2010 and 2011 data: blue crosses and red 197 diamonds in Figure 3a and c). Values for late summer (MALINA data: black circles in Figure 3a), however, were less variable and low near the surface (down to 0.0197 m⁻¹), except for river waters 198 that showed significantly higher values (up to 1.08 m^{-1} ; see arrow in Figure 3a). At depth > 150 m, 199 all data tended to decrease with depth, approaching $0.0277 \pm 0.0025 \text{ m}^{-1}$ (dotted rectangle in Figure 200 3c). This type of profile was similar to that reported by Guéguen et al. (2012), who suggested that 201 202 the maximal values around 150 m are associated with microbial activity (see sections 3.3 and 3.4). 203 While determination of the origin of CDOM (e.g., production by phytoplankton, heterotrophic 204 bacteria, etc) is still challenging using our dataset alone, we acknowledge that phytoplankton 205 especially at the deep chlorophyll maximum is one of the CDOM sources (Matsuoka et al, 2012).

Similarly to $a_{CDOM}(440)$, S_{CDOM} values for early-middle summer varied widely at depths < 150 206 207 m (Figure 3b). In contrast, those values showed less variability in late summer and were highest near the surface (up to 0.022 nm⁻¹; Figure 3b and d). At depths > 150 m, all S_{CDOM} values 208

approached $0.0167 \pm 0.0005 \text{ nm}^{-1}$ with increasing depth (dotted rectangle in Figure 3d).

210 For surface waters, our results thus showed that $a_{CDOM}(440)$ values decreased in association with 211 increases in S_{CDOM} from early to late summer (for up to 2.5 months). It is well known that as a 212 result of solar irradiation, high-weight molecules are transported into low-weight molecules that 213 absorb light in the shorter spectral wavelengths, a phenomenon called photo-bleaching. The spectral 214 slope therefore increases after the photo-bleaching (Twardowski and Donaghay, 2002). The 215 timescale is likely one to three months (Granskog et al., 2009). Satellite-derived sea ice 216 concentration images further showed that surface waters at most of our sampling stations were ice-217 free and therefore exposed to solar irradiation (Figure 2). All these results demonstrate photo-218 bleaching occurred during our observations.

For deep waters, CDOM absorption properties were stable (Figure 3a and b). Because CDOM in these waters is considered to be biologically unavailable or refractory, those absorption values can be considered as an end-member for refractory CDOM. If so, the proportion of CDOM lability could be optically quantified when the other end-member from labile CDOM is obtained. Further work is necessary to examine this issue.

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225 **3.2. Relationship between CDOM and hydrography**

226 Because salinity can be a good proxy of Arctic Ocean hydrography (e.g., Carmack et al., 1989; 227 Macdonald et al., 1989; Matsuoka et al., 2012), relationships between CDOM absorption properties 228 (i.e., $a_{CDOM}(440)$ and S_{CDOM}) and salinity were examined (Figure 4). At salinity < 28, two types of 229 waters were distinguished: 1) river waters with a strong negative correlation between $a_{CDOM}(440)$ 230 and salinity and 2) ice-melt waters showing both low $a_{CDOM}(440)$ and salinity (Figure 4a; see also 231 Matsuoka et al., 2012). These waters were observed in the Mackenzie shelf-basin area in the late 232 summer (black circles: MALINA cruise); river waters samples were collected in the Mackenzie 233 River mouth, while ice melt waters samples were taken near the sea ice, far away from the river 234 mouth (> 300 km). Data points at salinities < 28 located in the Chukchi-Beaufort Seas in mid-

summer (red diamonds: ICESCAPE 2011 cruise) were not far away from the negative a_{CDOM}(440)

236 versus salinity relationship for MALINA cruise (Figures 4a and b). In this salinity range, S_{CDOM}

exhibited low variability (0.0196 \pm 0.0011; Figure 4b), which is consistent with the value reported

238 by Matsuoka et al. (2012) (0.0192 \pm 0.0011).

239 At salinities > 28, $a_{CDOM}(440)$ values for the MALINA cruise tended to be lower than those for 240 the ICESCAPE 2010&2011 cruises (early-middle summer) conducted in the Chukchi-Beaufort Seas 241 (Figure 4c). This was especially true in the surface layer (i.e., 28 < salinity < 30.7), where the lower 242 $a_{CDOM}(440)$ values for the MALINA cruise corresponded to slightly but significantly higher S_{CDOM} values (0.0189 \pm 0.0006) compared to those for the ICESCAPE 2010&2011 cruises (0.0182 \pm 243 244 0.0004 and 0.0183 \pm 0.0012, respectively) (T-test, p < 0.0001; Figure 4c and d), which suggests 245 that photo-bleaching occurred from early-middle to late summer (Nelson et al., 1998; Twardowski 246 and Donaghay, 2002, Matsuoka et al., 2011; see also Figure 3). In other words, CDOM in the 247 Chukchi-Beaufort Seas observed during the early-middle summer cruises might have been 248 relatively new because of in situ production and/or input from the Bering Sea (Matsuoka et al., 249 2011; Shen et al., 2012).

The negative relationship between $a_{CDOM}(440)$ and salinity in the Lower Halocline Water (LHW: 33.9 < salinity < 34.7) and Atlantic Layer (AL: salinity > 34.7; our water samples were always collected at depths shallower than 850 m) was very similar among cruises (Figure 4c). This relationship was likely stable across ice-free seasons and areas of the WAO. So the $a_{CDOM}(440)$ versus salinity relationship could be specific to these waters.

In the intermediate layer between the surface and the LHW+AL layers (i.e., 30.7 < salinity <
33.9), a_{CDOM}(440) values for the ICESCAPE 2010&2011 cruise in the Chukchi-Beaufort Seas were
significantly higher than those for MALINA in the Mackenzie shelf-basin area (T-test, *p* < 0.0001).
Correspondingly, S_{CDOM} values in the intermediate layer were much lower for the ICESCAPE
2010&2011 cruises than for the MALINA cruise (T-test, *p* < 0.0001; Figure 4d). Based on field
observations, Nelson et al. (1998) suggested that bacteria produce CDOM when taking up dissolved 10

organic carbon (DOC), which is also consistent with results from laboratory experiments. Moran et al. (2000), Helms et al. (2008), and Ortega-Retuerta et al. (2009) observed that heterotrophic bacterial activity is associated with a decrease in the spectral slope of CDOM absorption over time. Note, however, that this result is contrary to the one obtained by Nelson et al., 2004. This contrast might originate from differences in substrate of organic matter utilized by bacteria and/or in distinct bacterial assemblages. Considering these findings, we hypothesized that the significantly low S_{CDOM} values observed in this study are associated with heterotrophic microbial activity.

268 To test this hypothesis, a vertical section of CDOM absorption properties, as well as salinity and 269 phaeopigments in the intermediate layer (i.e., 30.7 < salinity < 33.9), along the transect from the 270 Kotzebue Sound (KS) to the Chukchi Hotspot (CH) was further analyzed (Figure 5). The transect 271 covered both river-influenced and biologically productive areas sampled during the ICESCAPE 272 2010 cruise (see Figure 1 for location). Waters showing high $a_{CDOM}(440)$ and low salinity values 273 were observed in the surface layer of the KS, indicating river-influenced waters (Figure 5a and b). 274 Further offshore, this trend was no longer observed in the surface layer of the CH. Relatively high 275 a_{CDOM}(440) values, corresponding to significantly low S_{CDOM} values, were observed near the bottom 276 of the CH. Interestingly, the low S_{CDOM} values were associated with high concentrations of phaeopigments ($r^2 = 0.70$, p < 0.0001; Figure 5c and d); BP measurements are not shown here 277 278 because of the limited number of data points along this transect. Because phaeopigments reflect 279 degradation products of phytoplankton intermediated by bacteria, this result partly supports our 280 hypothesis that both the high a_{CDOM}(440) and low S_{CDOM} values observed in this study resulted from 281 heterotrophic microbial activity.

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3.3. Relationship between the spectral slope of CDOM absorption and bacteria

To examine the direct relationship between the spectral slope of CDOM absorption and bacteria, S_{CDOM} was regressed against BP and BA (Figure 6; chl *a* concentrations, temperature and nitrate concentrations were also regressed against S_{CDOM}). There was a weak but significant negative

correlation between BP and S_{CDOM} (Figure 6a; $r^2 = 0.20$, p < 0.0001). The coefficient of determination for this relationship was the second highest following the BP versus chl *a* relationship $(r^2 = 0.24, p < 0.0001;$ Table 1). Temperature showed the third highest correlation with slightly lower coefficient of determination ($r^2 = 0.18, p < 0.0001$). Significant correlations were not found for BP versus salinity or nitrate concentrations.

S_{CDOM} was negatively correlated with BA (Figure 6b; $r^2 = 0.26$, p < 0.0001). The coefficient of determination for this relationship was the highest, followed by BA versus chl *a* ($r^2 = 0.22$, p < 0.0001; Table 1). Similarly to BP, temperature showed the third highest correlation with BA ($r^2 = 0.17$, p < 0.0001). No significant correlations were found between BA and either salinity or NO₃ concentration. These results suggest that low S_{CDOM} values were generally associated with high BP and BA in our environments.

A similar regression analysis was performed for the spectral parameters $S_{275-295}$, $S_{350-400}$, and S_R , as done by Helms et al. (2008). Although results using $S_{350-400}$ were similar to those using S_{CDOM} (Table 2), none of them revealed higher correlations with BP and BA than S_{CDOM} ; S_{CDOM} is a better variable to reflect bacterial activity.

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303 **3.4. Multiple relationships among variables**

304 To examine relationships among several variables, a PCA analysis was performed. The extracted 305 axes of the PCA (Figure 7) can be interpreted as follows. Principal Component 1 (PC1) made the 306 largest contribution, explaining 30.9 % of total variance. The positive component was strongly 307 related to chl a, a_{CDOM}(440), BA, and BP and had a weak relationship with temperature and salinity. 308 Thus, the positive PC1 was considered to reflect production of organic matter. More interestingly, 309 S_{CDOM} alone showed the opposite trend compared to the above-mentioned variables (i.e., production 310 of organic matter), suggesting that negative PC1 might reflect the decomposition of organic matter. 311 Principal Component 2 (PC2) explained 21.4 % of total variance. The negative component was 312 strongly related to NO₃ concentration and apparent oxygen utilization (AOU) and was weakly

related to salinity. NO₃ and AOU were tightly correlated, consistent with recent findings by Matsuoka et al. (2012) for these two variables in the salinity range of 30.7-33.9 (or approximately 50-200 m depth; see their Figures 3c and d). Thus, negative PC2 was considered to reflect the aphotic zone. In contrast, the positive PC2 might reflect the euphotic zone, which is further supported by the fact that this component was weak but related to the production of organic matter.

Note that result of salinity in the PCA analysis is not surprising and can be explained as follows. First, salinity exhibited an opposite trend compared to that shown by S_{CDOM} . This general trend is shown in Figure 4. Second, this variable had a negative direction in PC2. This result is consistent with our discussion that negative PC2 represents the aphotic zone, showing higher salinity. Furthermore, Figure 7 provides important information: Because only S_{CDOM} showed opposite direction compared to parameters related to production of organic matter, this variable might be considered to reflect the degradation products of organic matter.

As expected, S_{CDOM} was negatively correlated with BP and BA (Figure 6), which is consistent with the result of simple regression analysis (Figure 6 and Table 1). Similarly, S_{CDOM} was negatively correlated with chl *a* and phaeopigments (p < 0.0001 for both). By taking into account the findings in Figures 5-7, our results suggest that variations in S_{CDOM} reflect the degradation of phytoplankton that is associated with heterotrophic bacterial activity.

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331 4. Conclusions

This study demonstrated that variations in the spectral slope pf CDOM absorption (S_{CDOM}) are partly explained by bacterial production (BP) and bacterial abundance (BA) variations. A simple regression analysis showed that S_{CDOM} was related to both BA and BP, which was further supported by PCA analysis. Bacterial abundance and production is likely dependent on bioavailability of DOM (e.g., Moran et al., 2000; Helms et al., 2008; Ortega-Retuerta et al., 2009). The spectral slope of CDOM reflects to some extent the level of DOC availability (Moran et al., 2000; Helms et al., 2008; Ortega-Retuerta et al., 2009). Therefore, it is consistent that S_{CDOM} is significantly correlated

339 with bacterial activity. Further work is necessary to better understand changes in CDOM cycling in

the Arctic Ocean.

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489 Figure captions

- 490 Figure 1. Locations of sampling stations for ICESCAPE 2010 (blue crosses), ICESCAPE 2011 (red
- diamonds), and MALINA (black circles) cruises in the Arctic Ocean. A transect from
 Kotzebue Sound (KS) to the Chukchi Hotspot (CH) is shown as a black line. Vertical
 sections of CDOM absorption properties as well as salinity and phaeopigment
 concentrations along this transect in the intermediate layer (i.e., 30.7 < salinity < 33.9) are
 shown in Figure 5.

496 Figure 2. Mean sea ice concentration images provided by the using satellite microwave sensor,

497 DMSP SSM/I, during (a) MALINA, (b) ICESCAPE2010, and (c) ICESCAPE2011 cruises.

These images were generated by averaging daily images available during each cruise.
Sampling stations are also displayed with white circles, yellow crosses, and red diamonds,
respectively.

Figure 3. Vertical profiles of (a) CDOM absorption coefficients at 440 nm ($a_{CDOM}(440)$, m⁻¹) and (b) their spectral slope (S_{CDOM} , nm⁻¹). X-axis for (a) is log-transformed to show variability in both low and high $a_{CDOM}(440)$ values. Mean profiles of (c) $a_{CDOM}(440)$ and (d) S_{CDOM} with 10-m intervals. Standard deviations are shown as horizontal bars.

505 Figure 4. Upper panels: CDOM absorption coefficients at 440 nm as a function of salinity (S) for

506(a) the whole salinity range and (c) for $S \ge 28$. A linear fit provided by Matsuoka et al.507(2012) is shown in grey. Data points along this fit correspond to river-influenced waters.508Data points for ice melt waters are shown in the circle in (a). Lower panels: spectral slope of509CDOM absorption coefficients, S_{CDOM} as a function of salinity for (b) the whole salinity510range and (d) for $S \ge 28$.

- Figure 5. Vertical sections of (a) salinity, CDOM absorption properties (b) a_{CDOM}(440) and (c)
 S_{CDOM}, and (d) phaeopigment concentrations along the transect from Kotzebue Sound (KS)
 to the Chukchi Hotspot (CH).
- 514 Figure 6. Relationship between S_{CDOM} and (a) bacterial production (BP, $\mu g \ C \ m^{-3} \ d^{-1}$) and (b)

515 bacterial abundance (BA, cells ml^{-1}).

516 Figure 7. Biplot of principal component analysis (PCA). The important features of this plot are as 517 follows: 1) direction of each arrow represents contribution to principal component 1 (PC1: 518 x-axis) and 2 (PC2: y-axis) for a given variable and 2) magnitude of an arrow represents the 519 strength of the variable to the components. For example, BP and chl a concentrations 520 showed a similar direction, suggesting they are related to each other and to positive PC1. 521 This result is consistent with the direct regression analysis (Figure 6 and Table 1). The 522 positive PC1 reflects production of organic matter (section 3.4). Thus, BP and chl a 523 concentrations can be considered as contributors to the production of organic matter. See 524 section 3.4 for details.

525

Table 1. Summary of the Model II regression analysis for bacterial production (BP, μ gC m⁻³ d⁻¹) and abundance (BA, cells ml⁻¹) as a function of chlorophyll *a* (chl *a*, mg m⁻³) concentrations, spectral slope of CDOM (S_{CDOM}, nm⁻¹), temperature (T, degrees C), salinity, and nitrate (NO₃, μ mol kg⁻¹) concentrations. A total of 133 samples were used for each regression analysis.

Parameter	Statistics	Log ₁₀ (BP)	$Log_{10}(BA)$
Log ₁₀ (chl <i>a</i>)	r^2	0.24	0.22
~O×	Slope	0.40	0.15
C.U	p-value	< 0.0001	< 0.0001
S _{CDOM}	r^2	0.20	0.26
	Slope	-169.9	-73.65
	p-value	< 0.0001	< 0.0001
$Log_{10}(T+2)^{*}$	r^2	0.18	0.17
	Slope	0.70	0.26
	p-value	< 0.0001	< 0.0001
Salinity	r^2	0.04	0.10

	Slope	0.06	0.04
	p-value	< 0.05	< 0.01
NO ₃	r^2	0.04	0.05
	Slope	-0.03	-0.01
	p-value	< 0.05	< 0.01

^{*}For conversion to a base 10 logarithm, 2 was added to T.

533 Table 2. Determination coefficients between bacterial variables (BP or BA) and spectral slope

534 parameters ($S_{275-295}$, $S_{350-400}$, and its ratio S_R) proposed by Helms et al. (2008).

	r^2	Log ₁₀ (BP)	Log ₁₀ (BA)
	S ₂₇₅₋₂₉₅	0.11	0.12
	S ₃₅₀₋₄₀₀	0.19	0.23
	S_R	0.03	0.04
Rcce	Q'LE		













