Pan-Arctic optical characteristics of colored dissolved organic matter: Tracing dissolved organic carbon in changing Arctic waters using satellite ocean color data

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ABSTRACT

Light absorption of the colored fraction of dissolved organic matter (CDOM) is a dominant optical component of the Arctic Ocean (AO). Here we show Pan-Arctic characteristics of CDOM light absorption for various Arctic regions covering both coastal and oceanic waters during the Tara Oceans Polar Circle expedition. The Siberian (or eastern) side of the AO is characterized by higher CDOM absorption values compared to the North American (or western) side. This is due to the difference in watersheds between the eastern and western sides of the AO and is consistent with an Arctic absorption database recently built by Matsuoka et al. (2014). A direct comparison between in situ and satellite data demonstrates that CDOM absorption is derived Arctic-wide from satellite ocean color data with an average uncertainty of 12% (root mean square error of 0.3 m⁻¹) using our previously published algorithm. For river-influenced coastal waters, we found a single and highly significant relationship between concentrations of dissolved organic carbon (DOC) and CDOM absorption (r² > 0.94) covering major Arctic river mouths. By applying this in situ relationship to satellite-derived CDOM absorption, DOC concentrations in the surface waters are estimated for river-influenced coastal waters with an average uncertainty of 28%. Implications for the monitoring of DOC concentrations in Arctic coastal waters are discussed.

1. Introduction

Colored dissolved organic matter (CDOM) refers to the fraction of the dissolved organic matter pool that absorbs light in water, and is quantified by its spectral absorption coefficient (m⁻¹). CDOM plays a variety of roles in ocean physical and biogeochemical processes (e.g., Moran and Zepp, 1997; Miller et al., 2002; Matsuoka et al., 2012, 2015) and provides the possibility to trace, using a simple optical measurement, the concentration of dissolved organic carbon (DOC) in seawater, the second largest reservoir of carbon in the ocean (Hansell, 2002). Strong correlations between DOC concentration and CDOM absorption have been reported for river-influenced coastal waters at all latitudes (e.g., Massicotte et al., in press and references therein), highlighting the potential usefulness of CDOM to study DOC distributions in these waters.

Despite the important role of CDOM in biogeochemical cycles and as a DOC proxy, our ability to document it in situ has been restricted by sea ice in the Arctic Ocean. Since the early 2000s, however, partly due to a significant reduction of summer extent of the Arctic ice pack, a number of in situ datasets have been acquired, and recently an Arctic seawater light absorption database was built to provide a synoptic view of the spatial and temporal variations of CDOM in the Arctic Ocean (Matsuoka...
et al., 2014). Still, a large part of the Siberian (or eastern) side of the Arctic Ocean (EAO), where high concentrations of organic carbon are delivered by river discharge (Raymond et al., 2007), was undersampled in the above-mentioned study, which impeded comparing optical properties between the EAO and the North American (or western) side of the Arctic Ocean (WAO). This is an important gap because a very large amount of organic carbon (400 Pg; Pg = 10^{15} g) is stored in the upper three meters of the northern Siberian soils (McGuire et al., 2009) and a significant fraction of this organic carbon may be delivered by river discharge into the Arctic Ocean, which might alter biogeochemical processes of the ocean (e.g., McGuire et al., 2009; IPCC, 2013). We lack knowledge about how the DOC budget of the Arctic Ocean will be modified as a consequence of ongoing global warming. Remote sensing of CDOM provides a powerful mean for tracing DOC in the Arctic Ocean and observing its dynamics and response to climate change from space.

The objectives of this study are therefore 1) to examine optical characteristics of CDOM in the different Arctic seas, 2) establish a relationship between DOC concentration and CDOM absorption coefficient that can be applied to wide range of river-influenced coastal waters, and 3) apply this relationship to satellite-derived CDOM absorption for estimating DOC concentrations from space. With known uncertainties, examples of monitoring of spatial-temporal variability in DOC concentrations for Arctic river-influenced coastal waters are shown at the end of the present study.

2. Materials and methods

The Tara Oceans Polar Circle expedition was conducted from 24 May to 5 November 2013 following a long transect in the Arctic Ocean (Fig. 1a). While the data presented in this study were mainly obtained from the Tara expedition, an Arctic absorption database built by Matsuoka et al. (2014) (hereafter referred to as the M2014 database), is also used here for comparison. Briefly, the M2014 database includes data from Western and Eastern Arctic Ocean covering Beaufort, Chukchi, Kara, and Laptev seas from May to October (Table 1).

To establish a DOC versus \(a_{\text{CDOM}}(443)\) relationship that is applicable to wide range of river-influenced coastal waters of the Arctic Ocean, we compiled and used publicly available data in addition to the Tara data (Section 2.1.4; Table 1). This provides a confidence of the general relationship in terms of statistics. Similarly, data obtained from MALINA, ICESCAPE2010, ICESCAPE2011 cruises included in the M2014 database (Matsuoka et al., 2014), in addition to Tara data, were also used to evaluate the performance of the CDOM algorithm developed by Matsuoka et al. (2013) (hereafter referred to as gsmA algorithm; see Sections 2.1.5 and 2.2.1).

In the present study, we refer waters having \(a_{\text{CDOM}}(443) < 0.1 \text{ m}^{-1}\) and \(> 0.1 \text{ m}^{-1}\) as oceanic and coastal waters, respectively. Of coastal waters, these waters are specifically referred to as river-influenced coastal waters when the \(a_{\text{CDOM}}(443)\) shows a significantly high correlation with salinity or DOC.

2.1. In situ data

2.1.1. CDOM absorption

2.1.1.1. UltraPath measurements. Light absorption coefficients of CDOM using an UltraPath (World Precision Instruments, Inc.) was determined by following the protocols proposed by Bricaud et al. (2010) and Matsuoka et al. (2012). To avoid repeating the protocols, only relevant points to the present study are recalled here. A sample was collected daily from a surface CTD/Niskin bottle or the in-line system flow-through (when no CTD deployment took place) into glass bottles pre-rinsed with MilliQ water. The sample bottles were covered with aluminum foil to avoid a potential effect of light degradation on CDOM in the water. These samples were filtered within a few hours after the sampling using 0.2 \(\mu\)m GHP filters (Acrodisc Inc.) pre-rinsed with 200 ml of Milli-Q water. Absorbance spectra of filtrates were measured from 200 to 727 nm with 1 nm increments relative to a salt solution that was used as a reference. The reference was prepared to have a similar salinity as samples (± 2 salinity units) using Milli-Q water and granular NaCl precombusted in an oven (at 450 °C for 4 h). Abnormally high absorbance values in the near infrared spectral domain were sometimes observed due to the presence of air bubbles in the cell of the sample. These suspicious spectra were removed prior to analysis. While significant effort was made to minimize the difference in temperature and salinity between a sample and reference water during our cruise, this was challenging especially for areas in which waters showed a large salinity gradient with different water temperature. The temperature difference was minimized by placing both the reference and the filtrates at 4°C in the dark for up to 1 h. The salinity difference was minimized by subtracting the mean value of \(OD_{\text{DOM}}(\lambda)\) between 683 and 687 nm (\(OD_{\text{DOM}}\)) from the whole spectrum following Babin et al. (2003) and CDOM absorption coefficients \(a_{\text{CDOM}}(\lambda, \text{m}^{-1})\) were calculated as follows:

\[
a_{\text{CDOM}}(\lambda) = 2.303 \frac{OD_{\text{DOM}}(\lambda) - OD_{\text{DOM}}(\text{broad})}{l}
\]

where 2.303 is a factor for converting base e to base 10 logarithms, and \(l\) is the optical pathlength (m). A 2 m optical pathlength was used for the measurement, except for water having high CDOM content (> 2.0 \(\text{ m}^{-1}\); Matsuoka, unpublished data) where 0.1 m pathlength was used. Replicates of CDOM absorption spectra were averaged for each measurement.

We acknowledge that scattering due to colloids might have influenced absorption measurement using a long pathlength (> 0.5 m; Flöge et al., 2009), which in turn might have influenced CDOM spectra when applying null-correction. However, for Mackenzie river mouth where the highest particle concentrations of the Arctic Ocean are observed (Holmes et al., 2002), Matsuoka et al. (2012) showed that CDOM absorption measurements using an UltraPath with 2-m pathlength were within the instrument resolution from those measured using a Perkin-Elmer Lambda-19 spectrophotometer with 10-cm cell. These results suggest that the impact of colloidal scattering on our absorption measurement can be considered as negligible in the present study.

2.1.1.2. ac-s measurements. To obtain CDOM measurements between those done with the UltraPath, we used a hyper spectral ac-s instrument installed in an automated flow-through system (red crosses in Fig. 1a). Hereafter CDOM absorption coefficients provided by the ac-s are noted as \(a_{\text{ac-s}}(\lambda)\) (in \(\text{m}^{-1}\)).

Filtered water was pumped through the ac-s (WET Labs) for 10 min per hour (see Boss et al., 2013 for the ac-s setup). The last minute of every 10 min filtered cycle was median-binned and saved. No clean water spectra for calibration were collected with the ac-s during the expedition, and instrument was cleaned weekly. Therefore, the spectra potentially contained errors due to instrument drift (it is acknowledged that the ac family of instrument drifts in time as presented by Boss et al., 2013). Here, null-correction on the entire spectral range was applied.

The ac-s was then interpolated with 1-nm increments. To match the corresponding UltraPath measurements in this range. The calibration for ac-s spectra were made by computing and offset between UltraPath and dissolved ac-s measurements done within 25 min:

\[
\text{Offset} = a_{\text{ac-s}}(\lambda) - \text{median}(a_{\text{ac-s}}(\lambda))
\]

where \(a_{\text{ac-s}}(\lambda)\) and \(a_{\text{ac-s}}(\lambda)\) represent CDOM absorption using UltraPath (UP) and ac-s, respectively. There were 59 such matchups within 12 h and 20 km from an UP measurement of dissolved matter, the nearest calibration spectra (in time) was used to correct the spectra:

\[
a_{\text{ac-s}}(\lambda) = a_{\text{ac-s}}(\lambda) + \text{Offset(near)}
\]

A spectral slope \(\text{SCDOM, nm}^{-1}\) was calculated by fitting an
exponential equation to the data in the spectral domain (i.e., 400–406 to 500 nm) as follows:

\[ a_{\text{CDOM}}(\lambda) = a_{\text{CDOM}}^{(443)} e^{-a_{\text{CDOM}}(443)} \, e^{-a_{\text{CDOM}}(\lambda)} \]  

The spectral range was chosen to be as similar as possible relatively to the one for the M2014 database (350 to 500 nm; Matsuoka et al., 2014) while avoiding the range from 500 to 550 nm for ac-s measurements where the signal was, on occasion, within the uncertainty of the measurement (not shown). Using the MALINA data, which exhibited a large variability in \( a_{\text{CDOM}}(\lambda) \) (see Matsuoka et al., 2012 for details), it was possible to establish a DOC versus \( a_{\text{CDOM}}(443) \) relationship (see Matsuoka et al., 2012 for details).

### Table 1

Summary of CDOM absorption coefficients (m\(^{-1}\)), DOC concentrations (\(\mu\)M), and \( R_{rs}(\lambda) \) (sr\(^{-1}\)) data used in the present study. ID #1 is the main dataset used in the present study. ID #2 is a large optical database established by Matsuoka et al. (2014) (M2014 database) and is used for comparison. IDs #3–7 are only used to establish a DOC versus \( a_{\text{CDOM}}(443) \) relationship.

<table>
<thead>
<tr>
<th>ID #</th>
<th>Dataset</th>
<th>Location</th>
<th>Period</th>
<th>Variable</th>
<th>( N )</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Tara</td>
<td>Pan-Arctic Ocean</td>
<td>2013-05-24–2013-11-05</td>
<td>CDOM (%) ( R_{rs}(\lambda) )</td>
<td>919 (20)</td>
<td>This study</td>
</tr>
<tr>
<td>2</td>
<td>M2014 database</td>
<td>Beaufort, Chukchi, Kara, and Laptev Seas</td>
<td>2002-05-05–2011-10-13</td>
<td>CDOM (%), CDOM ( R_{rs}(\lambda) )</td>
<td>797 (77)</td>
<td>Matsuoka et al. (2014)</td>
</tr>
</tbody>
</table>

\* 919 of data were obtained from ac-s measurements that were calibrated against UltraPath ones as described in Section 2.1.1.
was demonstrated that the difference in the two spectral slopes using the two different spectral ranges was low (root mean square error, RMSE = 0.0012 nm⁻¹, N = 381). To confirm that the RMSE does not influence our results, we used a bootstrap method which randomly adds uncertainties (both positive and negative of variable magnitude independent at each wavelength) based on the RMSE (i.e., 0.0012 nm⁻¹) as follows. For each aCDOM(λ) spectrum, a maximum error (ɛmax) was calculated using aCDOM(443) and SCDOM ± 0.0012 at 400 nm. The positive and negative ɛmax was then randomly added to aCDOM(λ) from 400 to 500 nm. A SCDOM was calculated to the ɛmax-included data as described above. This analysis was repeated a thousand times. The statistics associated with the new SCDOM were obtained and compared to those in the Table 2. Results were not significantly different from the values presented in the Table 2 (p = 0.10, t-test).

Prior to the analysis, we removed all the spectra not within a 1 day temporal window around an UltraPath measured spectra (in total, 919 ac-s spectra were retained).

2.1.2. CDOM fluorescence

An Aquatic Laser Fluorescence Analyzer (ALFA, WETLabs; Chekalyuk et al., 2014; Chekalyuk and Hafez, 2013; Chekalyuk et al., 2016) was installed as part of the Tara flow-through system. The ALFA instrument provides dual-wavelength excitation (405 and 514 nm) of laser-stimulated emission (LSE) for spectral and temporal fluorescence analysis. Spectral deconvolution (SDC) was used to examine fluorescence associated with a specific constituent and Raman scattering in water (R). The Raman scattering intensity was used for fluorescence normalization to provide adjustment for potential variability of the excitation source intensity and water optical properties (Chekalyuk and Hafez, 2008). We analyzed the ALFA SDC assessment of the CDOM fluorescence/Raman ratios measured using laser excitation at 405 nm and 514 nm. The ALFA sensor was cleaned weekly following the manufacturer recommended protocol.

2.1.3. DOC concentration

Surface water samples were collected directly from either the Rosette Niskin bottle or from the flow-through system (at the same time as the UP measurements; see Fig. 1a) into acid-clean 60 ml HDPE Nalgene bottles through pre-combusted GF/F filters (450 °C for 4 h; e.g., Carlson et al., 2010; Shen et al., 2012). Samples were then immediately stored at −20 °C. At the end of the cruise the samples were shipped to the University of South Carolina for analysis. Concentrations of dissolved organic carbon (DOC) were determined using the high-temperature combustion method and a Shimazu total organic carbon (TOC-VCSN) analyzer (Shen et al., 2016). In total, 20 samples were collected for DOC analyses.

2.1.4. Additional CDOM absorption and DOC concentrations data

While the dataset obtained from the Tara cruise provide a large number of CDOM absorption spectra, the number of the corresponding DOC concentrations data is limited (i.e., N = 20). To augment our dataset for our analysis, we added publicly available datasets for both CDOM absorption and DOC concentrations data from Retamal et al. (2007), Osburn et al. (2009), Matsuoka et al. (2012), Goncalves-Araujo et al. (2015), and Osburn et al. (2016) (hereafter referred to as R2007, O2009, M2012, G2015, and O2016, respectively; see Table 1). These data cover major Arctic river mouths including both Siberian and North American sides.

CDOM absorption was determined using a traditional spectrophotometer with a 5 or 10 cm quartz cell, except for M12 where an UltraPath was used as described above (section UltraPath measurements in Section 2.1.1). DOC concentrations were determined based on high temperature oxidation using a TOC analyzer for the mentioned data.

2.1.5. Remote sensing reflectance

We followed the NASA Ocean Optics Protocols (Mueller and Austin, 1995) and Hooker et al., 2013 to measure in-water upwelled radiance (Lw, µW cm⁻² nm⁻¹ sr⁻¹) and downward irradiance (Ed, µW cm⁻² nm⁻¹). These data were obtained at 19 wavelengths ranging from 320 to 780 nm (10 nm full-width at half-maximum, FWHM). A compact-optical profiling system (C-OPS, Biospherical Instruments Inc.) (Morrow et al., 2010) was deployed at 29, 36, and 19 stations for Tara Arctic, MALINA, and ICESCAPE2010 cruises, respectively. For the ICESCAPE2011 cruise, a profiling reflectance radiometer series 800 (PRR-800, Biospherical Instruments Inc.; 10 nm FWHM) was deployed at 24 stations. The above-water downward irradiance, so-called the global solar irradiance (Es: sum of direct and diffuse components) were used to correct Ed and Lu data for change in the incident light field during water column profiling. The effect of ship shadow was minimized by deploying the in-water spectroradiometer away from the main ship body. Only data having tilt angles < 5° were used for analysis in the present study, as recommended by Hooker et al. (2013). Subsurface Lu values at null depth (i.e., Lu(0−, λ)) were calculated using the slope and intercept by fitting the least-squares linear equation to the log-transformed upwelled radiance versus z for MALINA, ICESCAPE2010, and ICESCAPE2011 data. The depth interval was chosen following the method described by Antoine et al. (2013). Briefly, a centered depth z0 and +/− Δz was first set within homogeneous surface layer. The Δz was then increased to have the best statistical fit but was chosen to be as minimal as possible within the homogeneous surface layer. Data within this depth interval were used to extrapolate Lu(z, λ) to Lu(0−, λ). For Tara data, loss extrapolation was performed. The remote sensing reflectance, Rs(λ) is defined to be equal to 0.54 Lu(0−, λ)/Es(λ), where λ indicates wavelength. In the present study, Rs(λ) was at six wavelengths (i.e, 412, 443, 490, 532, 555, and 670 nm corresponding approximately to the ocean color bands of the Moderate-Resolution Imaging Spectroradiometer (MODIS) onboard the Aqua satellite, Aqua/MODIS) were used to derive aCDOM(λ) at 443 nm using gsmA algorithm. Prior to analysis, we removed the water Raman contribution to Rs(λ) using the method of Lee et al. (2013). The impact of Raman scattering on Rs(λ) data is examined in Appendix A.1.

2.2. Satellite ocean color data

2.2.1. Match-up analysis

Aqua/MODIS Rs(λ) data (λ = 412, 443, 488, 531, 555, and 667 nm; 15 nm and 10 nm FWHM for 412–443 and 488–667 nm, respectively) were obtained from the NASA Ocean Biology Processing Group (OBPG) satellite to in situ validation system (Bailey and Werdell, 2006; http://seabass.gsfc.nasa.gov/search). In this study, data from two different reprocessing versions (R2013.1 and R2014.0, respectively) were used to evaluate aCDOM(443) retrievals using gsmA algorithm.
Knowing that Aqua/MODIS has issues with radiometric degradation since 2011 [https://oceancolor.gsfc.nasa.gov/reprocessing/r2014/aqua/], we chose to include two Aqua/MODIS reprocessing data sets so as to examine how this degradation – and any residual calibration errors and differences that might persist between and within the two processings - might influence our results. No such a radiometric issue has been currently widely revealed for NPP/VIIRS, so the latest data were used in the present study.

\[ a_{\text{CDOM}}(443) \] was retrieved by optimizing the difference between measured \( R_L(\lambda) \) and \( R_L(\lambda) \) calculated using absorption (including unknown \( a_{\text{CDOM}}(443) \)) and backscattering coefficients, and coefficients associated with geometry of the sun. The Aqua/MODIS-derived \( a_{\text{CDOM}}(443) \) estimates were then directly compared to in situ \( a_{\text{CDOM}}(443) \) measurements and the performance of the algorithm was evaluated individually for each processing version. We used data within \(+/- 3\) h from satellite overpass within solar and sensor zenith angles of 74 and 56°, respectively for this match-up analysis (Bailey and Werdell, 2006; IOCCG, 2015). To be consistent with in situ C-OPS data, the water Raman contribution to satellite-derived \( R_L(\lambda) \) was removed prior to the analysis using the method presented by Lee et al. (2013).

2.2.2. Estimating DOC concentration from space

To illustrate spatio-temporal variability in satellite-derived DOC concentrations in Arctic river mouths, Aqua/MODIS Level 2 (L2) \( R_L(\lambda) \) swath images at 1-km nadir spatial resolution were obtained from the NASA ocean color website (http://oceandata.sci.gsfc.nasa.gov/MODIS-Aqua/L2). Cloud-free swath images were selected and were used to retrieve \( a_{\text{CDOM}}(443) \) using gsmA algorithm. Because of the polar orbit of earth observing satellites, several swath images are available per day at polar regions. Swath images within 2 h around solar noon were used to make a daily composite \( a_{\text{CDOM}}(443) \) image at 1 km spatial resolution. By applying an empirical relationship between DOC concentrations and \( a_{\text{CDOM}}(443) \) to satellite-derived CDOM absorption (described in Section 3.2), we estimated DOC concentrations for river-influenced coastal waters. The uncertainty in our estimates of DOC concentrations was determined according to sensitivity analysis as described in Appendix A.2.

2.3. Evaluation functions

We used evaluation functions proposed by Bailey and Werdell (2006) to evaluate the performance of the retrievals of \( a_{\text{CDOM}}(443) \) using gsmA algorithm applied to in situ or satellite \( R_L(\lambda) \) data. They include median of satellite to in situ ratio (\( R_t \)), the semi-interquantile range (SIQR), the median absolute percent error (MPE), and root mean square error (RMSE) defined as follows:

\[
R_t = \text{median} \left( \frac{X_{\text{mod}}}{X_{\text{obs}}} \right)
\]

\[
\text{SIQR} = \frac{Q_3 - Q_1}{2}
\]

\[
\text{MPE} = \text{median} \left( 100 \times \frac{X_{\text{mod}} - X_{\text{obs}}}{X_{\text{mod}} + X_{\text{obs}}} \right)
\]

\[
\text{RMSE} = \sqrt{\frac{\sum_{n=1}^{N} (X_{\text{mod}} - X_{\text{obs}})^2}{N}}
\]

where \( X_{\text{mod}} \) and \( X_{\text{obs}} \) represent vectors of estimated and measured \( a_{\text{CDOM}}(443) \), respectively. \( Q_1 \) and \( Q_3 \) represent the 25th and 75th percentile, respectively. In addition, model II linear regression based on ranged major axis (RMA) method was performed following Legendre and Legendre (1998). In this study, log-transformed data were used for these evaluations except for \( R_L(\lambda) \) data (see Appendix A.1).

3. Results and discussion

3.1. CDOM absorption characteristics at the Pan-Arctic Ocean

CDOM absorption values fall in the range of the M2014 database (Fig. 1b). The frequency of high \( a_{\text{CDOM}}(443) \) values (> 1.0 \( \text{m}^{-1} \)) was somewhat higher compared to M2014, due mainly to higher CDOM contents observed in the Yenisei and Ob River mouths (Aas et al., 2002; Hessen et al., 2010; Heim et al., 2014; Matsuoka et al., 2014). Overall, lower median \( a_{\text{CDOM}}(443) \) values were observed in the present study compared to the M2014 (0.037 and 0.058 \( \text{m}^{-1} \) for the Tara and the M2014, respectively; Table 2). \( \text{SCDOM} \) Values in this study were similarly distributed to those in the M2014 database (Fig. 1c; medians of \( \text{SCDOM} \)
values for this study and the M2014 database were 0.0187 and 0.0183 nm$^{-1}$, respectively; Table 2). These results confirm that CDOM measurements obtained during the Tara Arctic expedition are representative of the range observed in the Arctic Ocean.

Spatial distribution patterns of CDOM absorption properties were examined in a wide spatial coverage including both oceanic and coastal waters (Fig. 2). In the Norwegian and Greenland Seas, and west coast of Greenland, $a_{\text{DOM}}(443)$ values were remarkably low (< 0.1 m$^{-1}$). This result is consistent with that reported by Kirk (1994). In contrast, high $a_{\text{DOM}}(443)$ values were observed in river-influenced coastal areas of the Siberian side (up to 2.1 m$^{-1}$) and on the Canadian side (up to 0.6 m$^{-1}$). The high $a_{\text{DOM}}(443)$ values corresponded with low salinity in these waters (Fig. 2c). Indeed, $a_{\text{DOM}}(443)$ values were tightly correlated with salinity for all river mouths (Fig. 3a), indicating that high CDOM contents were delivered by rivers. This type of conservative behavior of the $a_{\text{DOM}}(443)$ versus salinity relationship in river-influenced coastal waters has been reported elsewhere (e.g., Jerlov, 1976; Monahan and Pybus, 1978; Nieke et al., 1997; Bélanger et al., 2006; Retamal et al., 2007; Mannino et al., 2008; Matsuoka et al., 2012). For WAO, the $a_{\text{DOM}}(443)$ versus salinity relationship established in September in the present study was similar to the one obtained in the southern Beaufort Sea in August reported by Matsuoka et al. (2012) (blue dashed line in Fig. 3a; $p > 0.5$). Our $a_{\text{DOM}}(443)$-salinity relationship for the Laptev Sea obtained in August was also similar to that obtained from August to September and reported by Heim et al. (2014). Our $a_{\text{DOM}}(443)$ versus salinity relationship obtained in July for the Kara Sea was close to the relationship observed in September by Aas et al. (2002), although the slope was lower in the present study. These results confirm previous studies; while the conservative physical mixing plays a major role in the behavior of $a_{\text{DOM}}(443)$-salinity relationship at low salinity (< 25), this relationship can vary seasonally and inter-annually (particularly for the case of Kara Sea). Data points that deviated from the mixing line likely come from waters experiencing degradation of organic matter due to either physical (Alling et al., 2010) and/or biogeochemical processes (Ortega-Retuerta et al., 2012; Asmala et al., 2014; Matsuoka et al., 2015).

The spatial distribution pattern of $S_{\text{DOM}}$ was more variable than $a_{\text{DOM}}(443)$ (Fig. 3b). Given that $S_{\text{DOM}}$ is influenced by complex physico-biogeochemical effects (i.e., mixing, photo-bleaching, microbial activity; Nelson and Siegel, 2002), it is difficult to explain the spatial variability using $S_{\text{DOM}}$ and salinity alone. Instead, $S_{\text{DOM}}$ versus $a_{\text{DOM}}(443)$ relationship could provide a rough idea of characteristics of dissolved organic matter (Stedmon et al., 2011) when compared to that in the literature. For example, Matsuoka et al. (2011) (M11) using data obtained from oceanic waters suggested photo-bleaching is a major effect controlling $S_{\text{DOM}}$ versus $a_{\text{DOM}}(443)$ relationship. Our data points showing $a_{\text{DOM}}(443)$ lower than 0.1 m$^{-1}$ are generally distributed within the 95% confidence interval (gray lines in Fig. 3b). At $a_{\text{DOM}}(443) > 0.1$ m$^{-1}$ (obtained from coastal environments), most of our data fall within the range obtained in coastal waters by Matsuoka et al. (2012) (M12; black dashed line in Fig. 3b). These results suggest that a majority of our data can be explained by river-influence and/or photo-bleaching. Data points outside M11 and M12 might be attributed to either sea ice melt (Amon, 2004; Matsuoka et al., 2012), microbial effect (Matsuoka et al., 2015), physical mixing (Retamal et al., 2008), or their combined effect. Further work using chemical identification technique is required to ascertain these effects.

The Raman-corrected CDOM fluorescence at 405 nm of the ALFA was highly correlated with the CDOM absorption measurement at 405 nm ($r^2 = 0.99$; Fig. 4). This result highlights the informative potential of CDOM fluorescence, the technology of choice on autonomous platforms. Note that the relationship between CDOM absorption at 514 nm and the Raman-corrected CDOM fluorescence stimulated at 514 nm were less robust ($r^2 = 0.88$; plot not shown), possibly affected by the spectral overlap between CDOM and phycobiliprotein (PBP) pigment fluorescence bands (Chekaluk and Hafez, 2008). The ratios of $\alpha_{\text{DOM}}(\lambda_2)$ to $F_{\text{DOM}}(\lambda_2)$ ($\lambda_1$ or $\lambda_2$ represents either 405 nm or 514 nm) versus $S_{\text{DOM}}$ did not show clear pattern in this study (not shown). While corresponding absorption and fluorescence data were obtained only at two wavelengths in this study, hyper-spectral fluorescence measurements combined with coincident absorption measurements may provide potential for characterizing both sources and compositions of CDOM (e.g., Stedmon and Markager, 2005; Guéguen et al., 2012; Para et al., 2012; Goncalves-Araujo et al., 2015).

3.2. Monitoring DOC concentrations from satellite remote sensing

There is growing concern regarding the modifications in DOC fluxes
originating from permafrost thaw and the consequences in biogeochemical processes in Arctic coastal waters, particularly its microbial consumption and the potential release of carbon dioxide (CO₂) into the atmosphere (IPCC, 2007, 2013; McGuire et al., 2009). DOC concentrations are highly correlated with CDOM absorption for river-influenced coastal waters (e.g., Matsuoka et al., 2012, 2014; Heim et al., 2014). In addition to mechanisms controlling CDOM absorption properties (see Section 3.1), tracing these variations temporally and geographically is important to better understand the carbon cycle.

To trace DOC dynamics, we recently developed a semi-analytical algorithm for Arctic waters (i.e., gsmA algorithm) that can be applied to satellite ocean color Rrs(λ) data such as those provided by Aqua/MODIS and the Suomi National Polar-orbiting Partnership (NPP) Visible Infrared Radiometer Suite (VIIRS) (Matsuoka et al., 2013, 2014). This algorithm includes two steps: 1) first retrieving aCDOM(443) value and then 2) estimating DOC concentrations. The present re-evaluation of this algorithm using in situ Rrs(λ) (i.e., C-OPS) data obtained from both EAO and WAO shows that aCDOM(443) value can be derived with 7.1, 0.176, and 0.060 for MPE, RMSE, and SIQR, respectively (Fig. 5; Table 3). Rt and slope were close to 1 and coefficient of determination was high (r² = 0.86). In addition, a direct comparison of in situ measurements with satellite-derived aCDOM(443) (i.e., match-up analysis) demonstrates that the MPEs of aCDOM(443) estimate for the two Aqua/MODIS processing versions (R2013.1 and R2014.0) are 9.0 and 11.6%, respectively (Fig. 6, Table 3). Other statistical measures also show the reasonable aCDOM(443) estimate (Rt close to 1, reasonable SIQR and RMSE). It is interesting to note that slope for R2014.0 reduced and was reasonable aCDOM(443) estimate (Rt close to 1, reasonable SIQR and RMSE), indicating that residual errors in satellite instrument calibrations do not strongly influence our results. A MPE of 8.5% was obtained for NPP/VIIRS. All these results confirm the reasonable performance of gsmA algorithm for Arctic waters. Note that coefficients of determination for all satellite sensors are relatively low and the slope for R2013.1 is higher than 1, due mainly to the small dynamic range of aCDOM(443) used for the analysis (Table 3).

Table 3

<table>
<thead>
<tr>
<th>Retrievals</th>
<th>Rt</th>
<th>SIQR</th>
<th>%MPE</th>
<th>RMSE</th>
<th>Slope</th>
<th>Intercept</th>
<th>r²</th>
<th>Range of in situ aCDOM(443)</th>
<th>N</th>
</tr>
</thead>
<tbody>
<tr>
<td>aCDOM(443)_C-OPS</td>
<td>1.030</td>
<td>0.060</td>
<td>7.1</td>
<td>0.176</td>
<td>1.01</td>
<td>−0.01</td>
<td>0.86</td>
<td>0.012–1.082</td>
<td>103</td>
</tr>
<tr>
<td>aCDOM(443)_MODIS (R2013.1)</td>
<td>1.038</td>
<td>0.114</td>
<td>9.0</td>
<td>0.275</td>
<td>1.26</td>
<td>0.36</td>
<td>0.53</td>
<td>0.013–0.100</td>
<td>32</td>
</tr>
<tr>
<td>aCDOM(443)_MODIS (R2014.0)</td>
<td>1.039</td>
<td>0.090</td>
<td>11.6</td>
<td>0.292</td>
<td>1.13</td>
<td>0.14</td>
<td>0.51</td>
<td>0.013–0.475</td>
<td>31</td>
</tr>
<tr>
<td>aCDOM(443)_VIIRS</td>
<td>0.997</td>
<td>0.063</td>
<td>8.5</td>
<td>0.303</td>
<td>0.17</td>
<td>−1.36</td>
<td>0.01</td>
<td>0.013–0.101</td>
<td>12</td>
</tr>
</tbody>
</table>

*** p < 0.0001.
274 μM) and were back to similar values as observed on 19 July 2011. In terms of absolute DOC concentrations estimated using satellite ocean color data, our estimates fall within the ones reported in the literature for four and major Arctic river mouths (i.e., Ob’, Yenisey, Lena, and Mackenzie river mouths; Amon, 2004; Hessen et al., 2010; Matsuoka et al., 2012; Orek et al., 2013). These results suggest that our approach to obtain DOC concentrations from ocean color data is reasonable, highlighting that the continuous monitoring of DOC concentrations is now possible with a known uncertainty.

Note that for some images, pixels near river mouths are masked out (for example 10 August 2011 for Lena river mouth on Fig. 8), which points out atmospheric correction problems for river mouths. This problem should be solved in the future using approaches such as those applied by Doxaran et al. (2015) and Matsuoka et al. (2016).

4. Conclusions

In the present study, we show that CDOM absorption properties vary significantly in various environments of the Pan-Arctic Ocean using continuous sampling instruments installed in a flow-through mode such as ac-s and ALFA. Our results suggest that establishing the relationship between CDOM absorption and its spectral slope could be useful for examining effects of terrestrial input, photobleaching, and microbial activity as outlined by Stedmon et al. (2011). Combined with spectral CDOM fluorometry, it may help to better understand variability in quality of dissolved organic matter. To facilitate discriminating the biogeochemical effects on CDOM, the optical methods need to be ascertained with a robust technique such as chemical identification.

Match-ups with satellite data demonstrate that CDOM absorption at 443 nm is derived using satellite ocean color data with an average uncertainty of 12% (RMSE of 0.3 m$^{-1}$), and DOC concentration is derived with an average uncertainty of 28%. Long-term in situ river discharge data are now publicly available thanks to several organizations (e.g., US Geological Survey, Environment Canada) and international projects such as the Pan-Arctic River Transport of Nutrients, Organic matter, and Suspended sediments (PARTNERS, e.g., Holmes et al., 2012). Satellite ocean color data have accumulated nearly 20 years. When combining these data together, DOC fluxes for river can further be estimated from space. This approach could be used to examine the trend of DOC fluxes in recent decades for Arctic rivers as a consequence of permafrost thaw and increase in river discharge (McGuire et al., 2009; IPCC, 2013).

Satellite estimate of DOC might contribute to investigate recent
Fig. 8. Estimates of concentrations of DOC in the surface layer for Laptev Sea using Aqua/MODIS ocean color data. Three relatively cloud-free images were selected for 2010 (left), 2011 (middle), and 2012 (right). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

Fig. 9. Same for Fig. 8 but for Kara Sea. Three relatively cloud-free images were selected for 2010 (left), 2011 (middle), and 2012 (right).
modifications in Arctic DOC cycling and its impact on the global CO2 flux when combined with in situ observations and a numerical model. Regarding in situ observations, recent findings showed that old permafrost-origin DOC is much more labile than what was previously thought and can be rapidly utilized by heterotrophic bacteria (Mann et al., 2015; Spencer et al., 2015). This apparently contradictory fact is likely explained by high levels of aliphatics included in the permafrost that is rapidly utilized by microbes (Spencer et al., 2015). This result implies a possible increase in CO2 release to the atmosphere that is rapidly utilized by microbes (Spencer et al., 2015). This fact is likely explained by high levels of aliphatics included in the permafrost, which has led to improve the performance of a numerical model that includes the microbial loop (Le Fouest et al., 2013). With the associated uncertainties, future modification in carbon cycle of the Arctic Ocean can be better assessed. This important task remains to be made in further studies.

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Appendix A.1. Effect of Raman scattering on \( R_{\lambda}(\lambda) \) data and \( a_{\text{CDOM}}(443) \) estimate

We examined the effect of Raman scattering on \( R_{\lambda}(\lambda) \) data and \( a_{\text{CDOM}}(443) \) estimate. The evaluation functions described in Section 2.3 were used for in situ \( R_{\lambda}(\lambda) \) data with \( R_{\lambda}^{\text{mod}}(\lambda), \text{sr}^{-1} \) and without Raman correction \( R_{\lambda}^{\text{obs}}(\lambda), \text{sr}^{-1} \); \( a_{\text{CDOM}}(\lambda) \) and \( a_{\text{CDOM}}^{\text{obs}}(\lambda) \) can be regarded as \( X^{\text{mod}} \) and \( X^{\text{obs}} \), respectively in Eq. 5–8 (but in linear scale). \( a_{\text{CDOM}}(443) \) was further estimated using either \( a_{\text{CDOM}}(\lambda) \) or \( a_{\text{CDOM}}^{\text{obs}}(\lambda) \) independently and the difference in the \( a_{\text{CDOM}}(443) \) estimates was evaluated as described in Section 2.3.

Results showed that the correction removed the effect of Raman scattering on \( R_{\lambda}(\lambda) \) as shown in Rt and regression slopes < 1:1 line (Table A1). The corresponding MPE of \( R_{\lambda}(\lambda) \) ranged from about 2% at 412 nm to about 5% at 670 nm. The SIQR varied from 0.002 sr\(^{-1} \) at 412 nm to 0.010 sr\(^{-1} \) at 670 nm. Given high quality of \( R_{\lambda}(\lambda) \) (the error < 5% is targeted) is required to derive meaningful geophysical values, this result suggests non-negligible contribution of Raman scattering on \( R_{\lambda}(\lambda) \) data.

Independent to the above-mentioned analysis, we performed a sensitivity analysis to examine the Raman effect on \( R_{\lambda}(\lambda) \) using a radiative transfer simulation (Hydrolight version 5.3, hereafter referred to as HES5.3). Simulations were done for a typical Arctic waters (chl a concentration = 0.1 mg m\(^{-3} \), 0.01 m\(^{-1} \) < \( a_{\text{CDOM}}(443) \) < 0.04 m\(^{-1} \), and 0.1 g m\(^{-3} \) < suspended particle matter concentration < 1.0 g m\(^{-3} \)) based on Arctic optical parameters documented by Matsuoka et al. (2011, the present study), which produced HES5.3-simulated \( R_{\lambda}(\lambda) \) including and without Raman effect. Result showed that Raman scattering has an impact on \( R_{\lambda}(\lambda) \) more than 3% particularly in the green-red spectral domain. This result is consistent with our analysis.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Rt</th>
<th>SIQR</th>
<th>%MPE</th>
<th>Slope</th>
<th>Intercept</th>
<th>( r^2 )</th>
<th>RMSE</th>
<th>N</th>
</tr>
</thead>
<tbody>
<tr>
<td>( R_{\lambda}(412) )</td>
<td>0.981</td>
<td>0.002</td>
<td>1.95</td>
<td>0.976</td>
<td>0.000</td>
<td>1.00E-04</td>
<td>103</td>
<td></td>
</tr>
<tr>
<td>( R_{\lambda}(443) )</td>
<td>0.978</td>
<td>0.002</td>
<td>2.20</td>
<td>0.976</td>
<td>0.000</td>
<td>1.00E-04</td>
<td>103</td>
<td></td>
</tr>
<tr>
<td>( R_{\lambda}(490) )</td>
<td>0.973</td>
<td>0.006</td>
<td>2.73</td>
<td>0.986</td>
<td>0.000</td>
<td>1.00E-04</td>
<td>103</td>
<td></td>
</tr>
<tr>
<td>( R_{\lambda}(555) )</td>
<td>0.963</td>
<td>0.010</td>
<td>3.71</td>
<td>0.986</td>
<td>0.000</td>
<td>1.00E-04</td>
<td>103</td>
<td></td>
</tr>
<tr>
<td>( R_{\lambda}(670) )</td>
<td>0.962</td>
<td>0.010</td>
<td>3.85</td>
<td>0.981</td>
<td>0.000</td>
<td>1.00E-04</td>
<td>103</td>
<td></td>
</tr>
<tr>
<td>( a_{\text{CDOM}}(443) )</td>
<td>0.999</td>
<td>0.013</td>
<td>1.32</td>
<td>0.987</td>
<td>-0.012</td>
<td>0.93E-03</td>
<td>0.237</td>
<td>103</td>
</tr>
</tbody>
</table>

* * * p < 0.0001.
The significant contribution of Raman scattering on DOC concentrations is required to provide a sense of confidence in the estimates. A similar statistical analysis performed by Matsuoka et al. (2016) was adopted in the present study. In essence, mean and the standard deviation of the intercept (int) and the slope (slo) for the log-transformed DOC versus aCDOM(443) relationship established in the present study were used to generate their normality of distribution by using 105 of iteration for each of the variables (the int or the slo each is a vector having 105 of data, hereafter referred to as vector in bold). The number of data (i.e., 105) was determined so as to obtain a standard error of the DOC estimate < 0.1% (in black; right axis of Fig. A1). Similarly, normality of distribution of aCDOM(443) was also generated using the mean of the measured value (obtained from Fig. 7) and the standard deviation was set to be 12% of the mean (the uncertainty of satellite retrievals of aCDOM(443); see Section 3.2 and Table 3; the percent difference was high (r² = 0.93, N = 103). This result suggests the minor impact of Raman scattering on aCDOM(443) estimate when using gsmA algorithm. It should be noted however that the few percent difference in R₀(λ) could have a significant impact on a geophysical value when using other algorithms. This needs to be determined in the further work.

Appendix A.2. Uncertainty analysis

The uncertainty in estimates of DOC concentrations is required to provide a sense of confidence in the estimates. A similar statistical analysis performed by Matsuoka et al. (2016) was adopted in the present study. In essence, mean and the standard deviation of the intercept (int) and the slope (slo) for the log-transformed DOC versus aCDOM(443) relationship established in the present study were used to generate their normality of distribution by using 105 of iteration for each of the variables (the int or the slo each is a vector having 105 of data, hereafter referred to as int and slo; vector in bold). The number of data (i.e., 105) was determined so as to obtain a standard error of the DOC estimate < 0.1% (in black; right axis of Fig. A1). Similarly, normality of distribution of aCDOM(443) was also generated using the mean of the measured value (obtained from Fig. 7) and the standard deviation was set to be 12% of the mean (the uncertainty of satellite retrievals of aCDOM(443); see Section 3.2 and Table 3; the aCDOM(443) is also a vector having 105 of data). The int, slo, and aCDOM(443) values were then introduced into our DOC versus aCDOM(443) relationship to estimate DOC concentrations (105 of DOC estimates). The coefficient of variation (CV) for the DOC estimates was calculated to be 28% (in red; left axis of Fig. A1) and determined as the uncertainty in the present study.

References


Fig. A1. Statistical analysis for determining uncertainty of satellite estimates of DOC concentrations. Left (red) and right (black) axes represent coefficient of variation (CV, in percent) and standard error (SE) relative to the absolute DOC estimate (in percent) as a function of the number of data points, respectively. Red and black dashed lines represent 98% in CV and 0.1% in SE relative to DOC estimates, respectively. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)


