TERRESTRIAL BROWNING FROM COLORED DISSOLVED ORGANIC MATTER (CDOM) CHANGES ALTERS THE SEASONAL PHENOLOGY PHYSICS AND BIOGEOCHEMISTRY OF THE COASTAL ARCTIC CARBON CYCLEMACKENZIE RIVER PLUME

Clement Bertin¹, Vincent Le Fouest², Dustin Carroll^{3,1}, Stephanie Dutkiewicz⁴, Dimitris Menemenlis¹, Atsushi Matsuoka⁵, Manfredi Manizza⁶, and Charles E. Miller¹

Correspondence: Clement Bertin (clement.maxime.bertin@jpl.nasa.gov)

Abstract. Arctic warming affects land-to-ocean fluxes of organic matter , through increased permafrost thaw, coastal erosion or river discharge, with significant impacts on coastal ecosystems and air-sea CO₂ fluxes. In this study, we modify a regional ECCO-Darwin ocean biogeochemistry simulation of the Mackenzie River region to include riverine export of colored dissolved organic matter (CDOM) and its effect on light attenuation, marine carbon cycling, and water-column heating from UV-A to visible light absorption. We find that CDOM light attenuation triggers both a two-week delay in the seasonal phytoplankton bloom and an increase in sea-surface temperature (SST) by 1.7°C. While the change in phytoplankton phenology has limited effect on air-sea CO₂ fluxes, the local increase in SST due to terrestrial browning-organic matter input switches the coastal zone from an annual sink of atmospheric CO₂ to a source (7.35 Gg C yr⁻¹). Our work suggests that the projected increase in terrestrial CDOM has strong implications for phytoplankton phenology and coastal air-sea carbon exchange in the Arctic.

10 1 Introduction

As anthropogenic emissions of carbon dioxide (CO_2) continue to increase (IPCC, 2023), it is critical to understand the time variability and future trajectory of the ocean carbon sink and its regional-scale response. The Arctic Ocean (AO) region constitutes an important sink of atmospheric CO_2 , estimated to be 116 ± 4 Tg C yr⁻¹ (Yasunaka et al., 2023), or roughly 7% of the global-ocean sink (Roobaert et al., 2019). The intense cooling of inflowing waters from adjacent seas and favorable conditions for phytoplankton growth result in elevated CO_2 uptake from increased CO_2 solubility and biological consumption, respectively. With Arctic air temperatures rising three to four time faster than the global mean due to the ice-albedo feedback

¹Jet Propulsion Laboratory, California Institute of Technology, Pasadena, CA, USA

²LIttoral ENvironnement et Sociétés (LIENSs) – UMR 7266, Bâtiment ILE, 2 rue Olympe de Gouges, 17000 La Rochelle, France

³Moss Landing Marine Laboratories, San José State University, Moss Landing, CA, USA

⁴Department of Earth Atmospheric and Planetary Sciences, Massachusetts Institute of Technology, Cambridge, MA, USA

⁵Institute for the Study of Earth, Oceans, and Space, University of New Hampshire, Durham, USA

⁶Geosciences Research Division, Scripps Institution of Oceanography, University of California San Diego, La Jolla, USA

(Rantanen et al., 2022), retreating sea-ice cover allows for a larger ocean surface area to be exposed to sunlight for longer periods of time (Bliss et al., 2019; Ardyna and Arrigo, 2020). As a result, AO Net Primary Production (NPP) increased by 90 Tg C (38%) from 1998–2012 (Ardyna and Arrigo, 2020; Lewis et al., 2020). Additionally, recent work by Terhaar et al. (2021) showed that a third of Arctic Ocean (AO) primary production is sustained by terrestrial fluxes from coastal erosion and rivers, resulting in-from large lateral fluxes of carbon and nutrients (Dittmar and Kattner, 2003; Le Fouest et al., 2013; Nielsen et al., 2022). However, the quantity and the composition of terrestrial matter exported to coastal regions is also impacted by climate change (Bertin et al., 2022; Mann et al., 2022; Tank et al., 2023), with potential to affect the bio-physical conditions of coastal AO waters.

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As Arctic river freshwater discharge increases (Feng et al., 2021), the quantity of terrestrial dissolved organic matter (DOM) exported to AO coastal peripheries is expected to increase. Due to complex molecular composition including aromatic cycles, DOM chemical composition depends on its origin and encompasses more than 20,000 molecular formulae (Dittmar et al., 2021). As it transitions from land to ocean, microbial activity and light alter DOM molecules, with their chemical composition being highly dependent on the transit through the terrestrial-aquatic environment (Cory et al., 2014; Cory and Kling, 2018). Once in coastal waters, the composition of riverine-derived DOM varies seasonally, likely being more labile (i.e., more easily degraded by microbes) during spring freshet (Spencer et al., 2009). A fraction of DOM, termed colored DOM (CDOM), possesses unique optical characteristics that enable it to efficiently absorb shortwave radiation — from ultraviolet (UV) to the visible light spectrum. In Arctic rivers, CDOM molecular weight and aromaticity increases with discharge (Mann et al., 2016), rendering it more resistant to degradation by marine bacteria (i.e., more refractory). Simultaneously, its interaction with light transforms CDOM either into 1) more-labile components of DOM (Osburn et al., 2009; Cory and Kling, 2018) or 2) directly into Dissolved Inorganic Carbon (DIC; Bélanger et al., 2006; Aarnos et al., 2018), which can promote CO₂ outgassing. By dampening light penetration into the water column, CDOM can drastically impact primary production (Bélanger et al., 2006; Li et al., 2024) (Li et al., 2024; Berezovski et al., 2025) and upper-ocean temperature (Hill, 2008; Kim et al., 2016; Soppa et al., 2019), which can also modulate air-sea CO₂ exchange. Consequently, air-sea CO₂ flux magnitude and direction in AO river plume regions remain highly uncertain, with both local-to-regional outgassing or uptake observed (Terhaar et al., 2019; Bertin et al., 2023; Roobaert et al., 2024). Additionally, as a result of global warming, accelerating permafrost thaw has the potential to change the composition of organic matter in coastal waters and therefore the coastal air-sea CO₂ fluxes via increased coastal erosion (Tanski et al., 2021; Nielsen et al., 2024) or river discharge (Mann et al., 2022). Thus, by a cascading effect, CDOM can locally amplify sea-ice melting due to increased sea-surface temperature (SST) from increased light attenuation (Pefanis et al., 2020). Therefore, understanding how terrestrial CDOM biophysical feedbacks influence coastal waters is critical to better characterize the consequences of climate change across Arctic coastal peripheries.

NPP in AO coastal regions also remains highly uncertain. The harsh polar conditions make it challenging to collect *in situ* in-situ observations and estimates from remote sensing are often contaminated by sea-ice, clouds, absence of light, and the high proportion of CDOM light absorption (Lewis and Arrigo, 2020; Li et al., 2024). Estimating NPP remotely

also requires several key assumptions regarding the vertical distribution of phytoplankton, since satellites only capture near-surface data (Arrigo et al., 2011; Silsbe et al., 2016). Current estimates suggest AO NPP ranges from 203–516 Tg C yr⁻¹ (Bélanger et al., 2013; Arrigo and Van Dijken, 2015), but these values are likely overestimated in coastal regions due to high CDOM concentrations. As a result, satellite estimates of air-sea CO₂ flux often fail to capture nearshore, river-plume regions (Bertin et al., 2023). To complement remote sensing, ocean biogeochemistry models (OBMs) permit full space-time coverage of AO coastal regions and can provide a mechanistic understanding of the processes that govern the air-sea CO₂ flux (Manizza et al., 2019; Mathis et al., 2022). Yet while most regional-scale OBMs now incorporate land-to-ocean nutrient transport (Terhaar et al., 2019; Lacroix et al., 2019; Lacroix et al., 2021; Savelli et al., 2025), their representation of the intricacies due to the CDOM feedbacks described above often remains partial or completely absent (though see e.g. Kim et al., 2018; Gnanadesikan et al., 2019; Pefanis et al., 2020).

In this study, we utilize a regional ocean-sea-ice-biogeochemistry model (ECCO-Darwin) to examine how riverine CDOM impacts the seasonal cycle of plankton biomass, phytoplankton biomass, primary productivity, and carbon cycling in the coastal AO. Our objectives are to 1) separate and explicitly quantify how CDOM's light attenuation properties affect both the physics and biogeochemistry in the river plume and 2) estimate how riverine CDOM modulates coastal air-sea CO₂ flux. Here we focus on the Southeastern Beaufort Sea (SBS), where the Mackenzie River discharges substantial freshwater and DOM into the AO (Bertin et al., 2022; Juhls et al., 2022). The remainder of this paper is structured as follows. First, we describe improvements made to the existing ECCO-Darwin regional configuration of the Southeastern Beaufort Sea (ED-SBS) regional set-up (Run_{strat} in Bertin et al., 2025) to incorporate CDOM processes and add riverine CDOM forcing. Second, we analyze the seasonal bio-physical conditions simulated by ED-SBS in the Mackenzie River plume. Third, we assess the impact of riverine CDOM on the physical characteristics of the plume region. Fourth, we analyze changes in phytoplankton phenology driven by riverine CDOM. Fifth, we estimate how CDOM impacts air-sea CO₂ flux within the plume region. Finally, we provide concluding remarks and suggestions for future work.

75 2 Methods

2.1 Explicit CDOM tracer parameterization

To simulate the coastal Arctic Ocean environment, we used the ED-SBS regional configuration, whose general numerical characteristics are fully detailed in Supporting Information Text S1 and in Bertin et al. (2023, 2025). ED-SBS simulated two marine dissolved organic carbon (DOC) pools with chemical properties representative of those found in the coastal AO: a semi-refractory pool (DOC_{sr}) characterizing the long-residence-time carbon loop with a lifetime of $\tau = 10$ years (Manizza et al., 2009), and a semi-labile pool (DOC_{sl}) characterizing the short-residence-time carbon loop with a lifetime of $\tau = 1$ month (including DOC molecules characterized by turnover rates ranging from weeks to months; Holmes et al., 2008; Spencer et al., 2015; Bertin et al., 2025). Land-to-sea forcing included daily discharge of freshwater and 6 biogeochemical tracers from the Mackenzie River, distributed over the three major Mackenzie Delta outlets: Shallow Bay (29.8%), Beluga Bay (37.6%), and

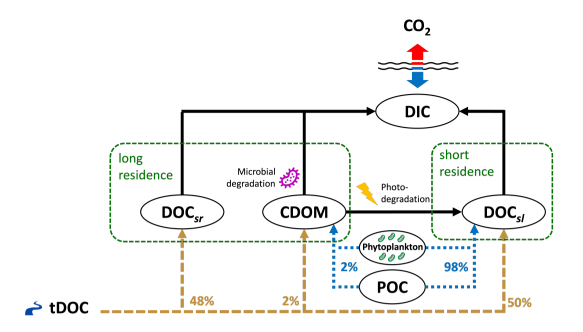


Figure 1. Conceptual diagram of dissolved carbon mass fluxes in the ED-SBS model. Mackenzie River terrestrial DOC (tDOC) mass flux (dashed brown lines) is distributed into marine DOC and CDOM pools according to the percentages shown in brown text. The result of phytoplankton grazing/mortality and particulate organic carbon (POC) dissolution is distributed over the DOC $_{sl}$ and CDOM pools (dotted blue lines).

Kugmallit Bay (32.6%) (Morley, 2012; Bertin et al., 2022). Freshwater discharge was driven by daily gauge measurements from the Arctic Great River Observatory (ArcticGRO; McClelland et al., 2023) and was linked to daily river temperature obtained from the Tokuda et al. (2019) dataset. Riverine concentrations of DOC, dissolved organic nitrogen (DON), dissolved organic phosphorus (DOP), dissolved silicate (DSi), dissolved inorganic carbon (DIC), and alkalinity (Alk) were forced as detailed in Bertin et al. (2025). As each export of dissolved organic constituents (DOC, DON & DOP) are estimated independently, the

In this study, we added an explicit "CDOM like" tracer to ED-SBS, expressed as a carbon mass concentration (mmol C m^{-3}), following the schematic shown in Figure 1. Terrestrial CDOM, which is observed to be non-labile (Blough and Del Vecchio, 2002; Aarnos et al., 2018), was added to the long-residence-time carbon loop of the model using the same microbial degradation rate turnover time as DOC_{sr} ($\tau = 10$ years). The CDOM tracer also interacted with the short-residence-time carbon loop by photochemical alteration of CDOM into more-labile carbon (Ward et al., 2017; Grunert et al., 2021; Clark et al., 2022). CDOM was photodegraded into DOC_{sl} with a maximum bleaching rate of 1/turnover time of 6 days (Dutkiewicz et al., 2015), which was modulated by ocean temperature and light intensity. Bleaching rate linearly increased from 0 when light intensity is 0 W m^{-2} to a maximum value (0.167 days $^{-1}$) when light is above 13 W m^{-2} (Dutkiewicz et al., 2015). CDOM

photodegradation rate corresponds to the bleaching rate modulated by a temperature function. When degraded into DOC_{sl} , CDOM products DON and DOP following the Redfield ratio of 120:16:1, allowing to represent the additional nutrient input generated by organic matter consumption. A fraction f_{CDOM} (= 2%) of mass fluxes received by DOC_{sl} through phytoplankton grazing/mortality and particulate organic carbon (POC) dissolution was also redistributed to CDOM.

In ED-SBS, Mackenzie River terrestrial DOC (tDOC) mass flux was equally distributed (50%) between semi-labile (DOC_{sl}) and semi-refractory (DOC_{sr}) DOC pools (based on recent estimates of the bioavailable tDOC fraction in the SBS, F. Joux, unpublished data from Nunataryuk field campaign; Lizotte et al., 2023). While 97% of DOC concentration variance is explained by CDOM absorption (Matsuoka et al., 2012), the mass concentration of riverine CDOM exported to SBS coastal waters remains unknown. As CDOM is part of the long-residence-time loop, we redistributed a percentage of tDOC mass flux from DOC_{sr} into the CDOM pool. After a sensitivity analysis (detailed in Appendix B), we set the ratio to 2% — re-partitioning Mackenzie River tDOC mass flux into 50%, 48%, and 2% DOC_{sl}, DOC_{sr}, and CDOM, respectively. Our ratio of total tDOC exported as CDOM falls in the lower range of estimates for the top 10 DOC exporting rivers (4–38% of tDOC; Aarnos et al., 2018). Finally, we generated CDOM initial and boundary conditions following the methods detailed in Supporting Information Text S2.

2.2 CDOM light attenuation relationship

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We first developed a new method for simulating CDOM light attenuation across the shortwave spectrum, from 320–735 nm. This allowed us to resolve the physical effect of CDOM light attenuation occurring in the UV-A (320–400 nm) and in the visible (400–735 nm) bands; the latter is often associated with Photosynthetically Active Radiation (PAR; spanning from 400–700 nm). An analysis of 31 CDOM spectral absorption measurements taken during the 2009 Malina campaign for different CDOM conditions across the SBS (see sampling locations in Supporting Information Figure S1; Matsuoka et al., 2012; Massicotte et al., 2021) revealed that $40\%\pm10$ (min:26–max:55) of light is absorbed by CDOM in the UV-A spectrum. These observations highlight the need to include full-band CDOM representation in OBMs, as most models only include light attenuation effects across PAR wavelengths. Note that in this study, we focus on light attenuation driven by CDOM absorption and disregard any backscattering effect from particulate matter. We acknowledge that the backscattering effect could play an important role in the SBS as the Mackenzie River is the Arctic's greatest exporter of particulate matter, but we aim here to build a foundation for determining the contribution of each component of terrestrial organic matter.

In our ED-SBS configuration, we approximated the relationship between CDOM light attenuation and its mass concentration (mmol C m⁻³) in high CDOM environments such as Arctic river-influenced waters. In this regard, we empirically estimated the CDOM diffuse attenuation coefficient (k_{CDOM} ; m⁻¹) from 31 in-situ measurements of the CDOM spectral absorption ($a_{CDOM}[\lambda]$; m⁻¹ nm⁻¹) across the SBS (Figure 2a.). The standard solar irradiance spectrum (ASTM G-173; U.S. D.O.E., 2005) was used as the reference shortwave solar spectrum at the surface ocean (Q_{sw_0} ; W m⁻² nm⁻¹) — Terms — terms are

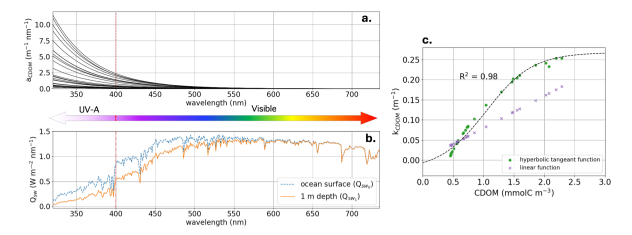


Figure 2. (a) In-situ CDOM spectral absorption measured over the Mackenzie Shelf during the 2009 Malina cruise for 31 water samples. (b) Shortwave solar spectrum (Q_{sw}) at the ocean surface (Q_{sw_0}) ; dashed blue line) and at 1-m depth after CDOM absorption (Q_{sw_1}) ; solid orange line). (c) CDOM attenuation (k_{CDOM}) relationship as it is described in Pefanis et al. (2020) (purple crosses) and in this study (green dots). The vertical red dashed line indicate the limit between UV-A and visible wavelength. Note that in Equation 1 we are computing the shortwave radiation absorbed from surface ocean to 1-m depth (Q_{sw_1}) , which results in the units being in W m⁻³ and hence k_{CDOM} having units of inverse meters.

listed in Table A1. We first calculated the shortwave spectrum attenuated at from the surface ocean to 1-m depth $(Q_{sw_1}; W m^{-2})^{-3}$ nm⁻¹) by multiplying Q_{sw_0} with $a_{CDOM}[\lambda]$ (Figure 2b.). Then, k_{CDOM} was retrieved by integrating Q_{sw_0} and Q_{sw_1} over the chosen wavelengths for each station using Equation 1.

$$k_{CDOM} = 1 - \frac{\int_{\lambda} Q_{sw_1} d\lambda}{\int_{\lambda} Q_{sw_0} d\lambda},\tag{1}$$

where lambda is the discrete wavelength (nm). Then, CDOM concentrations were estimated from a_{CDOM} [440nm] (m⁻¹) using the relationship from Neumann et al. (2020) (Equation 2).

$$CDOM = \frac{a_{CDOM}[440nm] + 0.2409}{Mc \times 0.0478},$$
(2)

where Mc is the carbon atomic mass ($Mc = 12.0107 \text{ g mol}^{-1}$). Finally, we fitted a hyperbolic tangent function (Equation 3) to obtain the relationship linking k_{CDOM} and CDOM concentrations across the range of conditions found in the SBS (Figure 2c).

$$k_{CDOM} = a \times \tanh(b \times \text{CDOM} + c) + d. \tag{3}$$

As shortwave radiation and PAR were simulated independently in the physical and biogeochemical components of the model, we calculated two different sets of parameters for the k_{CDOM} /CDOM concentration relationship for both components. Both relationships yielded an R² of \geq 0.98. Parameters fitted with the full shortwave spectrum (used in the physical component) were: a = -0.15, b = -1.31, c = 1.04, and d = 0.12. Parameters fitted with PAR (used in the biogeochemical component) were: a = -0.14, b = -1.18, c = 1.04, and d = 0.10.

2.3 CDOM biophysical feedback

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We included the effect of CDOM on light attenuation in the biogeochemical component of the model (which already included light attenuation by water and Chlorophyll-a (Chl-a)). PAR intensity (I(z), W m⁻²), at depth z is calculated according to the following equation:

$$I(z) = (1 - f_{ice}) \times 0.4 \times I_{sw} \times e^{-[k_w + k_{chl} \times chl(z) + k_{CDOM}(z)]dz}, \tag{4}$$

- where I_{sw} (W m⁻²) is the shortwave downwelling irradiance (input from the physical component of the model), for which 40% is considered as PAR, f_{ice} is the ice-cover fraction, k_w is the diffuse attenuation coefficient for pure seawater ($k_w = 0.04 \text{ m}^{-1}$), k_{chl} is the Chl-a diffuse attenuation coefficient ($k_{chl} = 0.04 \text{ m}^2 \text{ mg Chl-a}^{-1}$), chl(z) (mg Chl-a m⁻³) is the total concentration in Chl-a at depth z, and k_{CDOM} is the diffuse attenuation coefficient for CDOM at depth z.
- We included the biophysical feedback of CDOM light attenuation ocean warming by including k_{CDOM} , integrated over the entire shortwave spectra in the physical component of the model (see section 2.2). The physical component of the model already included the thermal effect of light attenuation by seawater, calculating a downwelling light decay profile (dk_{sw} ; 1-D) based on Jerlov water types (Paulson and Simpson, 1977) and decreasing from a value of 1 at the ocean surface to 0 near the seafloor. We included the thermal effect of CDOM light attenuation by calculating a CDOM light decay profile (dk_{CDOM}) based on k_{CDOM} (equation 5), also decreasing from 1 to 0. As CDOM concentrations are variable in space, the resulting light decay profile produces a 3-D field.

$$\begin{cases}
dk_{CDOM}(0) = 1 \\
dk_{CDOM}(z) = dk_{CDOM}(z-1) \times e^{-k_{CDOM}d(z-1)},
\end{cases}$$
(5)

where $dk_{CDOM}(0)$, the decay at the surface ocean (0-m depth) is set to 1, since simulated light has not yet been affected CDOM and z-1 is the depth of the vertical grid cell above z. The dk_{CDOM} calculation is then propagated from the ocean surface to the seafloor, as its value at depth z depends on all the values above. We then multiplied both decay profile to yield the total decay profile (dk_{tot} ; 3-D) as follows:

$$dk_{tot}(z) = dk_{sw}(z) \times dk_{CDOM}(z). \tag{6}$$

The setup described above represents a significant advancement over the previous model development by Pefanis et al. (2020). We included an updated took the advantage of an extensive in-situ carbon dataset collected in 2009 to update the parameterization of CDOM mass fluxes as they transition between short and long-residence-time carbon loops, where it was previously represented using a single DOC pool (Dutkiewicz et al., 2015). We also revisited the k_{CDOM}/CDOM relationship, transitioning from a linear to a hyperbolic tangent relationship (see Figure 2c.) This is particularly relevant for river plume regions where CDOM concentration reaches high values. Finally, our developments included the heating contribution of CDOM UV-A absorption, which contributes to roughly 40% of CDOM light absorption in the Mackenzie shelf region. The ED-SBS setup presented here is thus able to better represent the terrestrial browning effect on Arctic coastal regions.

3 Results

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The simulations presented herein include all model improvements detailed above (Run_{full}), *i.e.*, a CDOM tracer communicating with two DOC pools; CDOM light attenuation as a hyperbolic tangent function, including UV-A attenuation heating effect; and riverine input (see Table 1). For the remainder of the study, we focus our analysis on the 2012 – different from parameterization year (2009) – for two reasons: 1) sea-ice area showed a major reduction during this year (Parkinson and Comiso, 2013) and 2) previous results by Pefanis et al. (2020) focus on this specific year. However, all simulations were performed with the same forcings over 5 years (2008–2012) to mitigate spin-up effects in processes directly affected by the inclusion of CDOM, such as dissolved carbon (DOC and DIC) concentration, or indirectly affected such as nutrient stock through changes in primary productivity. We also limit our analysis to the Mackenzie River plume region, which we define by the time-mean sea-surface salinity (SSS) isohaline of 27 (Supporting Information Figure S1).

We also compute metrics that describe sea-ice phenology, as defined in Bliss et al. (2019); these metrics are then spatially averaged over the plume region. The day of opening (DofO) and the day of closing (DofC) are respectively the first and last

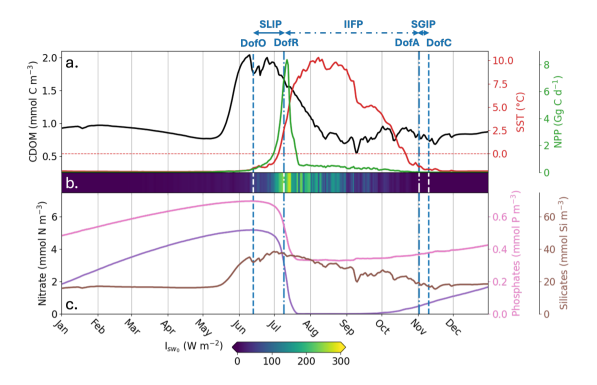


Figure 3. Spatially-averaged surface-ocean parameters simulated by $\operatorname{Run}_{full}$ in the Mackenzie River plume during 2012. Parameters shown are: (a) CDOM concentration (mmol C m^{-3} ; black line), SST (°C; red line), NPP (Gg C d^{-1} ; green line), (b) shortwave downwelling irradiance at the ocean surface ($\operatorname{I}_s w0$; W m^{-2}), (c) nitrate concentration (mmol N m^{-3} ; purple line), phosphate concentration (mmol P m^{-3} ; pink line) and, silicate concentration (mmol Si m^{-3} ; brown line). The vertical dotted blue lines show the spatial-mean day of opening (DofO) and day of closing (DofC) and the vertical dashed-dotted blue lines show the spatial-mean day of retreat (DofR) and day of advance (DofA). Sea ice melting periods are shown consecutively, the seasonal loss of ice period (SLIP), the inner ice-free period (IIFP), and the seasonal gain of ice period (SGIP).

days when sea-ice concentration is below 80%. The day of retreat (DofR) and the day of advance (DofA) are respectively the first and last days when sea-ice concentration is below 15%. The period between these two days is the inner ice-free period (IIFP) or open-water period. The period between DofO and DofR is defined as the seasonal loss of ice period (SLIP) and the period between DofA and DofC is the seasonal gain of ice period (SGIP). The above metrics are summarized in a schematic (see Appendix C) and are also indicated on the top of the following figures.

3.1 Mackenzie River plume seasonal phenology

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We first describe the seasonal phenology of several important physical and biogeochemical variables in the simulated Mackenzie River plume. In the river plume, $\operatorname{Run}_{full}$ simulates an average surface CDOM concentration of 0.85 ± 0.08 mmol C m⁻³

Table 1. Characteristics of the simulations tested in this study.

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Experiment name	\mathbf{k}_{CDOM}	CDOM heating	CDOM river input
Run_{full}	hyperbolic tangent	UV-A & visible	yes
$\operatorname{Run}_{noriv}$	hyperbolic tangent	UV-A & visible	no
Run_{lin}	linear	visible (PAR)	yes
$\operatorname{Run}_{ctrl}$	off	off	yes
Run_{light}	hyperbolic tangent	off	yes

Changes to Run full are highlighted in bold.

from August to May, with a peak of 2.04 mmol C m⁻³ during the spring freshet, followed by declining concentrations in July (Figure 3, black line). With regard to the sea-ice phenology in the river plume, the model simulates an open-water period of ~4 months (115 days), with SLIP and SGIP lasting 1 month (June 13 to July 9) and 1 week (November 2 to 10), respectively. From January to June, the SST is on average near the seawater freezing temperature (-1.93°C) and slowly starts heating up in June with increasing shortwave downwelling irradiance at the ocean surface (I_{sw0}; Figure 3b.) and accelerating freshwater discharge. In July, ocean-surface shortwave downwelling irradiance reaches a maximum, rapidly heating SST until it reaches a peak value of 10.3°C on August 8. Then, temperatures slowly cool until the end of SGIP. Phytoplankton rapidly bloom during the SLIP period, with a peak in surface NPP of 8.35 Gg C d⁻¹ occurring two days after DofR. The production period — defined as the duration when NPP exceeds half of its maximum — lasts 7 days and coincides with the period when subsurface light is the most intense. Nitrate and phosphate are quickly consumed during the phytoplankton bloom until the nitrate stock is depleted. Nutrient stocks are replenished through vertical mixing, advective transport, and remineralization from October to June. The simulated silicate tracer is directly connected to DSi riverine mass flux and therefore increases with elevated runoff.

Within the Mackenzie River plume region, Run full full captures the mean SST amplitude and variability during the openwater period depicted by observations (Figure D1). The model underestimates SST by 17% from mid-July to mid-September. This is due to a later simulated SLIP, which delays surface-ocean heating and causes simulated SST to increase later in the season. Run full full also reasonably reproduces the amplitude of the phytoplankton bloom observed by remote sensing, as the simulated surface-ocean Chl-a peaks at approximately the same concentration as reported by Lewis et al. (2020). However, the model underestimates the bloom's duration, simulating a bloom that lasts only half as long as observed by satellite. This discrepancy arises from the model's later simulated SLIP (similar to its SST behavior) and the rapid depletion of nitrates during the late open-water period. A more detailed and comprehensive model-data evaluation for 2012 is provided in Appendix D.

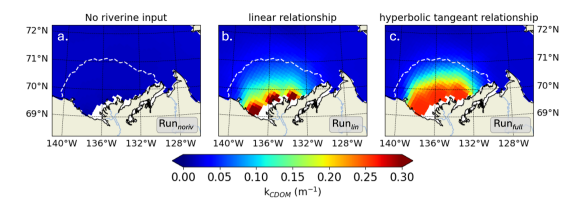


Figure 4. For July 2012, time-mean CDOM diffuse attenuation coefficient (k_{CDOM} , m⁻¹) for (a) Run_{noriv}, (b) Run_{lin}, and (c) Run_{full}. The white dashed line marks the time-mean spatial extent of the Mackenzie River plume.

3.2 Adding riverine CDOM to ED-SBS

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We next explore how the inclusion of riverine CDOM impacts light attenuation characteristics on the Mackenzie River shelf by comparing $\operatorname{Run}_{full}$ (presented above) to two similar set-ups: 1) excluding CDOM riverine forcing ($\operatorname{Run}_{noriv}$; autochthonous CDOM only) and 2) using a linear CDOM light attenuation only in visible light (similar to Pefanis et al. (2020); Run_{lin}). We analyze the differences for the month of July, when shortwave downwelling irradiance (I_{sw}) is maximum and terrestrial CDOM is more likely to affect the biophysical characteristics of the plume region. The simulation excluding river mass flux exhibits a space-time mean k_{CDOM} of $0.02~\mathrm{m}^{-1}$ ($\operatorname{Run}_{noriv}$) in the plume region (Figure 4a). Including riverine CDOM increases k_{CDOM} to $0.13~\mathrm{m}^{-1}$ and $0.16\mathrm{m}^{-1}$ when using a linear (Run_{lin}) and hyperbolic tangent ($\operatorname{Run}_{full}$) relationship with CDOM, respectively. In the vicinity of the river mouth, k_{CDOM} reaches values 6.5 to 8 times higher than simulations without riverine CDOM forcing, highlighting the importance including the riverine CDOM effect on light in the nearshore region. When using a linear relationship, k_{CDOM} increases as CDOM concentration increases, triggering high values (>0.3 m^{-1} with a maximum at $0.59~\mathrm{m}^{-1}$) in the direct vicinity of the river mouth, with a sharp transition to lower values further offshore (<0.2 m^{-1}) (Figure 4b.). When using the hyperbolic tangent relationship, k_{CDOM} is capped to $0.26~\mathrm{m}^{-1}$, given the k_{CDOM} /CDOM relationship fitted with in-situ observations (see Figure 2c.). As a result, CDOM attenuation is more evenly spread along the nearshore region (Figure 4c.).

3.3 Riverine CDOM biophysical feedback

We now examine how riverine CDOM influenced the physical conditions of the SBS during 2012, introducing a control simulation (Run_{ctrl}) that differs from Run_{full} by turning off both CDOM light attenuation (Section 2.2) and its effect on seawater

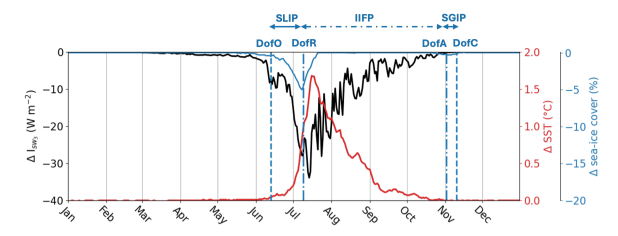


Figure 5. Difference in subsurface shortwave downwelling irradiance at 3m depth (I_{sw_3} in W m^{-2} ; black line), SST (°C; red line) and seaice concentration (%; blue line) between Run_{full} and Run_{ctrl} . The vertical dotted blue lines show the spatial-mean Day of Opening (DofO) and Day of Closing (DofC) and the vertical dashed-dotted blue lines show the spatial-mean Day of Retreat (DofR) and Day of Advance (DofA) simulated by Run_{full} .

heating (section 2.3) (see Table 1).

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In the river plume, $\operatorname{Run}_{full}$ simulates a peak of surface CDOM concentration during the spring freshet, which coincides with the SLIP and the increase in surface-ocean shortwave downwelling irradiance (Figure 3). As a result, the subsurface shortwave irradiance (I_{sw_3}) — defined as the shortwave irradiance ($\operatorname{W} m^{-2}$) below the model surface layer (3-m depth) — decreases by 13.4 W m^{-2} (40%) on average during the SLIP (Figure 5) compared to the simulation without CDOM effects ($\operatorname{Run}_{ctrl}$). CDOM light attenuation in the plume region then triggers an additional SST increase ($\operatorname{\Delta SST}$ up to 1°C), driving a decrease in sea-ice cover by up to 5% (Figure 5). We note a delay of 1 day in the DofR in $\operatorname{Run}_{full}$ compared to $\operatorname{Run}_{ctrl}$ (not shown), demonstrating the limited influence of riverine CDOM on sea-ice phenology. Terrestrial CDOM has a maximum impact on the physical condition of the plume one week after the DofR, with a 45% decrease in subsurface shortwave downwelling irradiance and an increase of by up to 1.68°C (Figure 5). Finally, the impact of riverine CDOM gradually diminishes as the tracer becomes diluted in the open ocean during the IIFP.

Following the approach in subsection 3.2, we analyze the influence of the kCDOM parameterization on the river plume's temperature by comparing the changes in SST simulated by Run_{noriv} , Run_{lin} , and Run_{full} , relative to Run_{ctrl} . We focus on the month of July, when CDOM has the greatest impact on SST in the Mackenzie River plume (Figure 6). In Run_{noriv} , the change in CDOM heating relative to Run_{ctrl} is solely attributed to marine CDOM produced by phytoplankton grazing and mortality. The spatially-averaged change in SST due to phytoplankton-generated CDOM, based on the improved CDOM-carbon loop connection (see section 2.1), is 0.45 ± 0.09 °C. The specific contribution of riverine CDOM leads to increases of

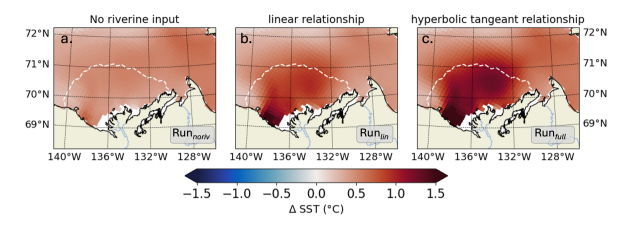


Figure 6. Time-mean SST differences (°C) for July 2012 between (a) Run_{noriv} , (b) Run_{lin} , and (c) Run_{full} , relative to the baseline simulation which excludes the CDOM effect (Run_{ctrl}). The white line in each panel indicates the mean extent of the Mackenzie River plume.

84% $(0.83 \pm 0.24^{\circ}\text{C}, \text{Run}_{lin})$ and 144% $(1.10 \pm 0.28^{\circ}\text{C}, \text{Run}_{full})$ using the linear and hyperbolic tangent k_{CDOM}/CDOM relationships, respectively. We note that the k_{CDOM} relationship in Run_{lin} only considers a classic linear CDOM warming effect resulting from PAR attenuation, emphasizing the dominant role of UV-A in SST warming.

3.4 CDOM effect on marine primary productivity

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In the remainder of the study, we explore the specific effects of CDOM light attenuation and ocean heating on the coastal ecosystems primary producers and the carbon cycle, focusing on the biological and solubility pump. From here, we only focus on three simulations: Run_{full}, Run_{light}, and Run_{ctrl}. The later two simulations deviate from Run_{full} by turning off aspects of the CDOM light absorption (see Table 1): In Run_{ctrl}, we turn off both CDOM light attenuation (Section 2.2) and its effect on seawater heating (section 2.3). In Run_{light}, we turn off only the CDOM heating effect (section 2.3) but include its effect on light attenuation. We then disentangle the individual impacts of light attenuation and their influence on ocean temperature over seasonal timescales.

Annual surface-ocean NPP integrated in the river plume region remains similar across simulations, whether including the influence of CDOM on light and temperature ($\operatorname{Run}_{full}$) or not ($\operatorname{Run}_{ctrl}$), yielding 0.10 and 0.13 Tg C yr⁻¹, respectively. However, a mean delay of 15 ± 3 (min: 9-max: 23) days occurs in the seasonal phytoplankton bloom, defined here as the day when Chl-a reaches its peak value. The surface-ocean NPP maximum, initially occurring in the middle of SLIP, is delayed to DofR by the end of the sea ice melt season due to CDOM (Figure 7a). Introducing both CDOM light and biophysical parameterizations ($\operatorname{Run}_{full}$) results in a 85% increase in peak NPP, with 78% attributed to the change in CDOM/light interactions and 7% to increasing SST. However, the production period — defined as the duration when NPP exceeds half of its maximum —

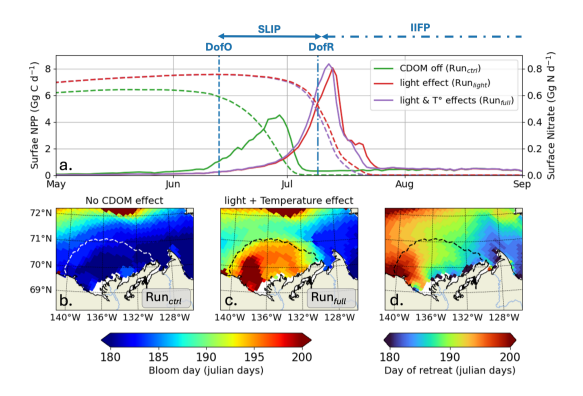


Figure 7. (a) Surface-ocean NPP (Gg G d⁻¹; thick lines) and nitrate stock (Gg N d⁻¹; dashed lines) simulated by ED-SBS without the CDOM light attenuation effect (green line; Run_{ctrl}), including 1) CDOM PAR light attenuation (red line; Run_{light}) and 2) CDOM light attenuation and ocean warming effect (purple line; Run_{full}). Maps of bloom day (julian days) (b) without the CDOM light attenuation effect, (c) including CDOM PAR light attenuation, and (d) map of DofR (julian days) simulated by EDS-SBS. Dotted line on the maps show the time-mean SSS isohaline of 27; vertical dotted and dashed-dotted blue lines show the DofO and DofA, respectively.

decreases from 12 to 5 days, thereby explaining the similar annual NPP.

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By early June, surface-ocean nutrient stocks are replenished through vertical mixing, advective transport, and remineralization that primarily occurred during winter – Note that the differences in May surface nitrate concentrations observed in Figure 7a are related to changes in stock replenishment over the spin up period due to CDOM inclusion. High sea-ice concentrations during most of the year result in light availability being primary limiting factor for phytoplankton growth, with temperature as a background limitation (See Appendix E). As the season progresses into SLIP the sea-ice concentration decreases, leading to higher light penetration into upper-ocean waters. In Run_{ctrl} , this allows phytoplankton to utilize nutrients and initiates a bloom (Supporting Information Figure S2a.) that persists until the nitrate stock is entirely consumed and thus limits further phytoplankton growth. However, by early June, riverine CDOM (Run_{light}) drives additional light attenuation, counterbalancing the increased light penetration resulting from sea-ice loss (see Figure 5), hence slowing down the bloom initiation and delaying it by roughly 2 weeks (see Figure 7 and Supporting Information Figure S2b). Consequently, phytoplankton bloom latter in

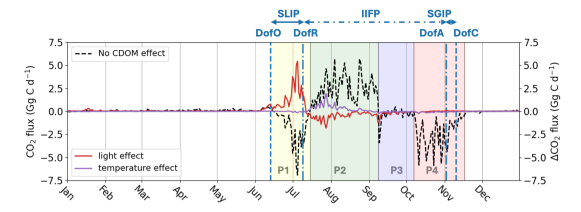


Figure 8. Air-sea CO_2 flux $(Gg \ C \ d^{-1})$ simulated by Run_{ctrl} in the plume region without CDOM biophysical feedback effects (black dotted line) and the change in air-sea CO_2 flux $(Gg \ C \ d^{-1})$ induced by the PAR light attenuation effect (red thick line) and warming effect (purple thick line). The vertical dotted blue lines show the average DofO and DofC and the vertical dashed-dotted blue lines show the average DofR and DofA. Phases with a switch in air-sea CO_2 flux simulated by Run_{ctrl} are indicated by four colors (P1: yellow, P2: green, P3: blue, and P4: red)

the season until the nitrate stock is exhausted and again limits further growth. We find an east-west gradient in the maximum bloom day (Figure 7c), correlated with the DofR (Figure 7d). This supports our hypothesis that light attenuation from riverine CDOM export complements light attenuation from sea-ice during the melting period and delays the seasonal phytoplankton bloom until the open-water period.

3.5 CDOM effect on coastal air-sea CO₂ fluxes

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In the absence of CDOM effects (light and heating effect), Run_{ctrl} results in a net annual CO_2 sink of -11.40 Gg C yr⁻¹ within the plume region. Over seasonal timescales, air-sea CO_2 exchange occurs from DofO to DofC, with four distinct phases (Figure 8). The following figures show the net air-sea CO_2 flux, integrated within the river plume region over the time period considered. The initial phase, starting from DofO and extending to one week after DofR, exhibits a substantial net CO_2 sink of -53.2 Gg C, which is attributed to phytoplankton growth (see Figure 7). Following this, the second phase, which spans two months at the onset of the inner ice-free period (IIFP), is marked by a significant net CO_2 outgassing of 137.9 Gg C — this results from the decline in phytoplankton abundance and heightened local concentrations of DIC/DOC from river discharge (Bertin et al., 2023). Subsequently, a less-variable, one-month long phase follows, characterized by a delicate balance (air-sea CO_2 flux near 0 Gg C d^{-1}) that results in a moderate net uptake of -10.3 Gg C. The third phase, stating in early October and extending to one week after DofC, exhibits a strong net CO_2 sink of -99.2 Gg C. During this last phase, phytoplankton decline due to depleted nitrate levels and DIC/DOC concentrations return to background levels as river discharge diminishes.

Over seasonal timescales, substantial changes in the timing and patterns of air-sea CO₂ flux occur during the two initial phases due to the inclusion of CDOM effects. As a result of CDOM light attenuation, we observe a delay in phytoplankton activity from the first phase (prior to DofR) to the subsequent phase (Figure 7), leading to a 79% reduction in simulated CO₂ uptake during phase 1 (+42.0 Gg C; Figure 8). Furthermore, the increase in SST due to CDOM is minimal during this period (Figure 5), resulting in an negligible impact on net air-sea CO₂ flux (0.4 Gg C).

As the phytoplankton bloom simulated by Run_{full} peaks at the onset of the second phase, CDOM light attenuation reduces net CO₂ outgassing by 47.0 Gg C. However, the warming effect of SST counteracts the reduced CO₂ outgassing (caused by phytoplankton growth), driving a CO₂ outgassing of 19.8 Gg C during this period. Consequently, the net CO₂ outgassing for this period is reduced by 27.2 Gg C. Comparing the loss in CO₂ uptake on the first period (42.0 Gg C) and the gain in CO₂ uptake (-27.2 Gg C), the reduction in the CO₂ sink during the first period is 14.8 Gg C higher than the gain in the second period.

Thus, changes in CO₂ fluxes during these two periods represent 80% of the annual net loss in CO₂ sink. As a consequence, when including the CDOM bio-physical feedback (Run_{full}), the plume switches to a net annual CO₂ outgassing of 7.35 Gg C yr⁻¹. We show here that, despite the greater effect of light attenuation on the magnitude and sign of air-sea CO₂ flux, the temperature effect is the dominant contributor in the transition of the plume from a sink to a source of CO₂, as it dampens the increased CO₂ uptake due to phytoplankton growth in early summer.

340 4 Discussion

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Assessing air-sea CO₂ fluxes in Arctic coastal environments remains challenging, as the carbon cycle and ecosystems are affected by a wide range of physical and biogeochemical processes that span the land-ocean continuum. As 11% of the global river discharge is fluxed into the Arctic Ocean (McClelland et al., 2012), coastal waters are highly influenced by terrestrial browning (Lewis et al., 2020; Li et al., 2024), motivating the need to include this effect in ocean biogeochemistry models. In this study, we develop a new regional-scale ECCO-Darwin model that simulates 1) the impact of marine CDOM on the physical properties of the water column (Kim et al., 2018; Gnanadesikan et al., 2019; Pefanis et al., 2020) and 2) the interaction of terrestrial CDOM with the marine carbon cycle (Neumann et al., 2021; Clark et al., 2022).

Our model includes CDOM light attenuation (k_{CDOM}) as a hyperbolic tangent function of CDOM concentration, estimated from in-situ observations of CDOM spectral absorption from 280–750 nm on the Mackenzie Shelf. Using this relationship, simulated CDOM in the plume region compares reasonably well with both in-situ and satellite measurements (Matsuoka et al., 2012, 2017; Massicotte et al., 2021, see Appendix B). Furthermore, we show that using a hyperbolic tangent for k_{CDOM} limits the effect of CDOM light attenuation in high CDOM concentration regions, allowing for the light attenuation from CDOM to be distributed more evenly along the nearshore region (Figure 4). Based on these results, we suggest that similar relationships be used in future models that aim to realistically represent coastal regions where CDOM concentrations reach high values. Additionally, this relationship was calculated from CDOM absorption integrated over the entire shortwave spectra, which in-

cludes the UV light absorption component, which is estimated to contribute up to 40% of CDOM absorption on the Mackenzie Shelf. Therefore, our study considers the complete effect of CDOM attenuation on ocean heating, inducing a 36% increase in the seasonal cycle of SST compared to previous methods (CDOM heating from PAR and k_{CDOM} as a linear relationship; see Run_{lin} and Gnanadesikan et al., 2019; Pefanis et al., 2020; Neumann et al., 2021).

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Many ocean biogeochemistry models now incorporate land-to-ocean nutrient fluxes (Terhaar et al., 2019; Lacroix et al., 2021; ?) (Terhaar et al., 2019; Lacroix et al., 2021; Savelli et al., 2025), however, ocean circulation and physics often shape-drive the biogeochemical state without possible biogeochemical feedbacks the potential feedback of biogeochemistry on physics. In 365 Arctic coastal regions, CDOM absorption has been reported to be a significant factor in the ocean heat budget (Hill, 2008; Soppa et al., 2019), but models still fail to include this feature. We find that including the CDOM heating effect in ED-SBS improved the model's ability to simulate the space-averaged SST observed during the early open-water season (Good et al., 2020, See Appendix D). We further show that riverine CDOM absorption contributes to a 1.7°C increase in SST in the Mackenzie River plume, which is consistent with the increased seasonal amplitude previously reported for the AO (Gnanadesikan et al., 370 2019). The maximum increase occurs at the onset of the open-water season (0.2°C/day), which is the same order of magnitude as observed in the Laptev Sea (Soppa et al., 2019). Although our model includes a component of CDOM generated by phytoplankton mortality and its associated light attenuation, we lack light attenuation by Chl-a (Dutkiewicz et al., 2019), which has been shown to increase the SST signal by $\sim 0.5^{\circ}$ C along the Arctic continental shelves (Lengaigne et al., 2009). We note that the simulated increase in SST has a limited impact on sea ice, as we only observe only a 5% decrease in sea-ice cover and a 375 change in DOR by a single day.

By adding CDOM light attenuation to ED-SBS, we are also able to improve simulated also observe a change in the phytoplankton bloom phenology in of the Mackenzie River plumeeompared to Lewis and Arrigo (2020) satellite observations (See Appendix D). During the freshet season (early June), in Run full full, riverine CDOM triggers a small difference in light limitation (see Appendix E), which delays the phytoplankton bloom by two weeks to the end of melting season. As a result of increased light penetrating the water column, the simulated phytoplankton bloom amplitude is 85% higher and 1 week shorter due to rapid nitrate consumption. In the plume region, we further observe a westward gradient in the phytoplankton bloom peak day, which is correlated with the day of sea-ice retreat (Figure 7 c. and d.). These results highlight that the coupling between CDOM and sea-ice have an important play a dominant role in shaping phytoplankton phenology, while the CDOM heating effect has a second order effect.

Although the model successfully simulates the timing of the bloom peak compared to Lewis et al. (2020), we Further comparing simulated Chl and primary productivity with satellite observation (Lewis et al., 2020) in the Mackenzie River plume (Note that this time-series are calculated where observations are available; more details in Appendix D), both Run_{ctrl} and Run_{full} overestimate the average maximum in surface Chl by 55% and 62%, respectively (Figure 9a). However, Run_{full} better simulates the spatial distribution of surface Chl especially in the vicinity of the coast (see Figure D1). Run_{full} also successfully

simulate the maximum in NPP observed by satellite (28 Gg C d⁻¹) while Run_{ctrl} underestimate it by 13%. However, with respect to the initiation of the bloom Run_{ctrl} better matches observations (surface Chl and NPP), where Run_{full} bloom initiate with 2 to 3 weeks delay. Looking into more details on the sea ice melting behavior (SLIP equivalent with or without CDOM), we find that ED-SBS exhibits a shorter 2012 SLIP, with a 24-days delay in the DofO and a 16-day delay in the DofR. We therefore acknowledge that the combination of sea-ice and CDOM light attenuation (Run_{full}) triggers the correct phenology in phytoplankton bloom initiation with respect to sea ice melting, but the incorrect timing as the bloom initiates 3 weeks later due to delayed DofO. This behavior in relation to the sea ice is confirmed in the comparison of averaged SST in the Mackenzie River plume (more details in Appendix D). As observed melting season starts mid-May, SST rises in early June when DofR approaches, while simulated melting season only kicks in mid-June – 1 month after the observations – allowing a rise in SST by the beginning of July. Finally, the observed production remains high latter in the open-water season (August to September); ED-SBS is not able to sustain a high rate of primary productivity during this period as nitrate is entirely consumed, shutting down the bloom. The low simulated levels of Chla after the bloom could be attributed to a match-mismatch with zooplankton being a general closure term in the model.

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We argue that including CDOM does not necessarily improve the phytoplankton phenology in the Mackenzie River plume compared to observations but does enhance its behavior regarding to sea ice melting. Furthermore, biophysical feedback of CDOM on water heating plays a non-negligible role in simulating SST in the region. We note that further improvements to the sea-ice model and its interaction with phytoplankton are would be required to accurately simulate the initiation of the bloom. 410 Importantly, ED-SBS does not consistently represent under-ice blooms, which have been shown to be an important processes in Arctic phytoplankton phenology (Ardyna and Arrigo, 2020; Ardyna et al., 2020). The next version of ED-SBS, which will have high horizontal (~ 1 km) and vertical resolution (~ 1 m at the surface), will permit improved representation of fine-scale sea-ice dynamics, such as cracks, leads, and specific features of the Mackenzie Delta such as the Stamukhi (Carmack and Macdonald, 2002; Matsuoka et al., 2016)—this work is presently underway. Including melt ponds in future version of the model will also be necessary to improve the initiation of the phytoplankton bloom, as there their impact on light penetration through sea-ice has been reported to be important for the development of under-ice blooms (Clement Kinney et al., 2023). (Ardyna and Arrigo, 2020; Clement Kinney et al., 2023). This might have an important effect as early snow melt and sea ice breakup as shown to enhance algal export in the Beaufort Sea (Nadaï et al., 2021). Finally, as a consequence of climate change and delayed sea-ice freeze-up, Arctic phytoplankton phenology has been reported to transition to double bloom characteristics (Manizza et al., 2023); with a spring bloom initiated by under-ice blooms and low-light-adapted diatoms followed by an au-420 tumn bloom characterized by low nitrogen ecosystems low-nitrogen adapted phytoplankton (Ardyna and Arrigo, 2020). The inclusion of the later ecosystem latter ecosystem components in ED-SBS could improve the phytoplankton representation in the latter open-water period, as our ecosystem is rapidly limited by nitrate concentrations. Furthermore, this hypothesis aligns with previous work by Choi et al. (2024), who demonstrated showed that the inclusion of a nitrogen fixer (not dependent on 425 nitrate) better represents could explain the secondary fall bloom.

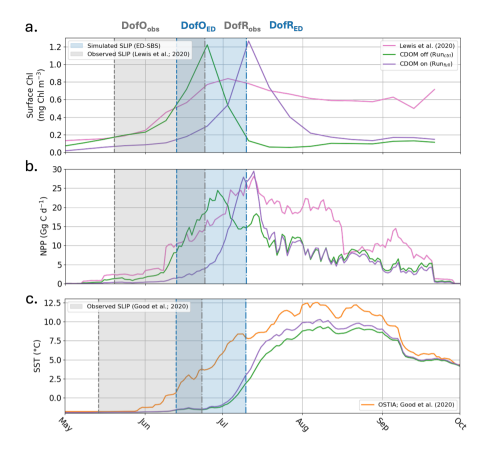


Figure 9. 2012 (a) averaged Surface Chl-a (mg Chl-a m^{-3}), (b) integrated NPP (Gg C d^{-1}), and (c) averaged SST (°C) over the Mackenzie River plume region for Run_{ctrl} (green line), Run_{full} (purple line), satellite observations (pink line Lewis et al., 2020), and in-situ/satellite observations (orange line Good et al., 2020). The blue (grey) area indicates the simulated (observed) seasonal loss of ice period.

While the CDOM heating effect has a limited impact on phytoplankton phenology, its role in modulating air-sea CO₂ fluxes is crucial, especially for the annual budget. With the inclusion of CDOM light attenuation, and as a consequence of two week weeks delay of the phytoplankton bloom, the strong CO₂ uptake that occurs during the melting period (without CDOM effect) disappears and shifts into a dampening of the early open-water period CO₂ outgassing (Figure 8). Over annual timescales, this results in a decrease in the net CO₂ sink of 4.6 Gg C yr⁻¹, with the Mackenzie River plume region remaining a CO₂ sink. However, the inclusion of the CDOM heating effect and the 1.7°C increase in SST at the onset of the open-water season promotes an increase in CO₂ outgassing due to reduced pCO₂ solubility, which balances the decrease in CO₂ outgassing driven by the phytoplankton bloom. Annually, CDOM heating promotes a 14.1 Gg C yr⁻¹ decrease in CO₂ uptake and switches the Mackenzie River plume region to a net CO₂ outgassing of 7.35 Gg C yr⁻¹. Although the contribution of Mackenzie River to the Arctic CO₂ budget is small (Yasunaka et al., 2023), we demonstrate that CDOM is an important factor contributing to CO₂ fluxes in coastal regions. In the future, the projected increase in terrestrial organic matter fluxes may drive elevated CDOM

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levels in Arctic coastal regions, thus affecting the solubility pump and local marine ecosystems (Nguyen et al., 2022). This effect is likely to be even more important in the Eurasian Basin, where terrestrial CDOM export is more pronounced (Stedmon et al., 2011).

Our study focuses on the Mackenzie River plume region, which is the main contributor of particulate organic carbon (POC) at the pan-Arctic scale (McClelland et al., 2016). Similarly to CDOM, terrestrial POC fluxes are likely to increase in the future with increased runoff, permafrost thaw, and coastal erosion (Doxaran et al., 2015; Nielsen et al., 2024; Sauerland et al., 2025). The additional amount of carbon exported to the coastal water by erosion alone could decrease the Arctic Ocean CO₂ uptake 445 by 7–14% by 2100 (Nielsen et al., 2024). Particulate organic matter also decreases the light available for primary production through absorption and backscattering (Stramski et al., 2004; Wozniak and Dera, 2007), potentially having an even stronger effect on the phenology. The effect of particle light backscattering may however overtake the effect of absorption, likely driving a decrease in coastal CO₂ uptake mainly by lower phytoplankton production rather than increased heat as shown with CDOM in this study. We acknowledge that ED-SBS does not account for terrestrial POC mass flux and the effect of suspended particulate 450 matter on the attenuation of light (backscattering effect), which might be significant in this region (Lizotte et al., 2023). However, we aim here to focus on the effect of the dissolved fraction and do not explore further assumptions regarding the particulate fraction effect, since our model does not account for solid sedimentation parameterization and bottom-sediment/seawater interactions. Future work will focus on the addition of a sediment model to fill this gap (Sulpis et al., 2022). Combined with the 455 new Plankton, Aerosol, Cloud, ocean Ecosystem (PACE) satellite mission, which includes a hyperspectral imaging radiometer, the next generation of ED-SBS will be able to disentangle signature of Chl, CDOM, and particulate matter to better estimate coastal Arctic CO₂ fluxes. Finally, the Estimating the Circulation and Climate of the Ocean (ECCO) framework has paved the way for the use of adjoint modeling at the global-ocean scale (Brix et al., 2015; Carroll et al., 2022). Future simulations may involve the use of adjoint modeling to optimize ED-SBS based on available physical and biogeochemical observations in the 460 SBS.

5 Conclusions

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We have developed a new regional model (ED-SBS) which includes terrestrial CDOM export from the Mackenzie River. The CDOM component interacts with the marine earbon cycle (DOC and DIC) and dissolved carbon pool and its feedback on the physical properties of the water column, such as light intensity and temperature. In particular, our model simulates UV-light absorption, which has been thus far ignored in model studies and is estimated to contribute to 40% of the light absorption in the SBS. We also suggest a new CDOM attenuation relationship as a hyperbolic tangent of CDOM concentration, which is able to better simulate light absorption in high CDOM concentration environments, such as river plumes.

In the plume region, we find that not including the coupled effects of light attenuation from sea-ice cover and riverine CDOM export leads to an earlier simulated seasonal phytoplankton bloom (2 weeks). By including riverine CDOM influence, the bloom occurs after the melting season, where light conditions are optimal, with a simulated phytoplankton bloom 85% higher than simulations without effect of CDOM, but also 1 week shorter due to quicker consumption of nitrate. We further find that including the riverine CDOM biophysical feedback switches the net CO₂ sink in the plume region from -11.40 Gg C yr⁻¹ (without CDOM effects) to a net outgassing of 7.35 Gg C yr⁻¹. Although the change in phytoplankton phenology has limited impact on the air-sea CO₂ fluxes, we find that the simulated outgassing is driven by reduction in pCO₂ solubility resulting from a 1.7°C increase in SST. Our modeling study demonstrates the importance of CDOM biophysical feedback in Arctic river plume regions, and the strong implications of CDOM radiative heating on pCO₂ solubility and air-sea CO₂ fluxes. In the context of climate change, we suggest that future increases in terrestrial organic matter exports could substantially affect ecosystems and air-sea CO₂ fluxes in shallow Arctic coastal regions where CDOM export is high.

Appendix A: List of terms

Table A1. List of terms used in this study

Terms	Abbreviation	Unit	Definition
Shortwave solar spectrum	Q_{sw}	$\mathrm{W}~\mathrm{m}^{-2}~\mathrm{nm}^{-1}$	Solar irradiance spectrum – at the surface of the ocean
			it corresponds to ASTM G-173 standard spectrum
Shortwave downwelling irradiance	I_{sw}	${\rm W}~{\rm m}^{-2}$	Integrated solar irradiance used in the physical
			component of the model
CDOM diffuse attenuation coefficient	k_{CDOM}	m^{-1}	loss of light intensity through CDOM.
CDOM absorption	a_{CDOM}	$\mathrm{m}^{-1}~\mathrm{nm}^{-1}$	loss of light absorbed by CDOM for each wavelength

Appendix B: Terrestrial CDOM ratio validation

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We set the percentage of DOC_{sr} redistributed into the CDOM pool by performing a sensitivity experiment. Three different parameterizations of the riverine CDOM input were tested: Marine CDOM tracer is forced at the Mackenzie River mouth by a.

1%, b. 2%, and c. 4% of the total riverine tDOC mass flux. This percentage is subtracted from DOC_{sr} to CDOM as detailed in section 2.1 (Figure 1). Then, we compared the simulated light CDOM absorption (a_{CDOM}[λ]) derived from simulated CDOM concentration (see equation 1) (see equation B1; Dutkiewicz et al., 2015) in the Mackenzie river plume, with in-situ observations of a_{CDOM}[440nm] measured during the Malina Campaign (see location of station in Figure S1 Matsuoka et al., 2012; Massicotte et al., 2021) and remotely-sensed a_{CDOM}[443nm] (Matsuoka et al., 2017). We then calculated the CDOM light absorption at a specific wave length from the simulated CDOM concentration using the following equation (Dutkiewicz et al., 2015):

$$a_{CDOM}[\lambda] = C_{CDOM} \times e^{-S_{CDOM}(\lambda - \lambda_0)} \times CDOM, \tag{B1}$$

where λ_0 is the reference waveband ($\lambda_0 = 450$ nm), C_{CDOM} is the CDOM absorption at λ_0 ($C_{CDOM} = 0.18$ m² mmolC⁻¹), and S_{CDOM} is the CDOM absorption spectral slope ($S_{CDOM} = 0.018$ m⁻¹).

We used 4 comparison metrics to compare retrieved CDOM absorption ($a_{CDOM}[\lambda]$) from simulated CDOM against observations: the median (\pm standard deviation), the correlation coefficient (r), the median percent error (MPE) and the unbiased root-mean-square error (MPE). Additional information and equations for the comparison metrics are detailed in Supporting Information Text S3. We find that changing the percentage of tDOC forcing CDOM as no impact on the correlation coefficient (Table B1). Size of discrepancies between the simulated and observed values (URMSE) are equivalent when riverine CDOM takes 1% or 2% of tDOC input but increases by 55 to 117% when forcing is set to 4%. The MPE increases by 40% to 89% when doubling the tDOC exported to CDOM from 1% to 2% and increases from 84% to 109% when doubling the tDOC exported to CDOM from 2% to 4%. The median of $a_{CDOM}[\lambda]$ is 0.08 ± 0.26 and 0.03 ± 0.25 m⁻¹ for in-situ and satellite observations, respectively. With 4% and 2% of tDOC forcing the CDOM pool, the simulated median of $a_{CDOM}[\lambda]$ is respectively fourfold (and doubled compared to observations). The simulated median is closer to observations when forcing with 1% of riverine tDOC. Comparing the time-mean 2009 CDOM absorption in the Mackenzie River plume region (Figure B1), the model forced with 2% of tDOC best fits the satellite data within the river plume area, while the model forced with 1% of tDOC results in a consistent underestimate. According to metrics and the comparison of time-mean $a_{CDOM}[\lambda]$ fields, parameterization b. was selected as the method best able to reproduce observed CDOM in the Mackenzie River plume region.

Table B1. Comparison metrics between simulated and observed $a_{CDOM}[\lambda]$ (m⁻¹

CDOM forcing parameterization	Observations	n	r	MPE	URMSE	$Median_{obs} \pm std$	$Median_{mod} \pm std$
1% of tDOC	Malina	18	0.78	45.10	0.18	0.08 ± 0.26	0.10 ± 0.14
	AMODIS	15,250	0.65	101.73	0.20	0.03 ± 0.25	0.06 ± 0.10
2% of tDOC	Malina	18	0.78	76.00	0.18	0.08 ± 0.26	0.16 ± 0.28
	AMODIS	15,250	0.65	192.06	0.19	0.03 ± 0.25	0.08 ± 0.20
4% of tDOC	Malina	18	0.79	159.20	0.39	0.08 ± 0.26	0.30 ± 0.56
	AMODIS	15,250	0.65	353.57	0.31	0.03 ± 0.25	0.14 ± 0.41

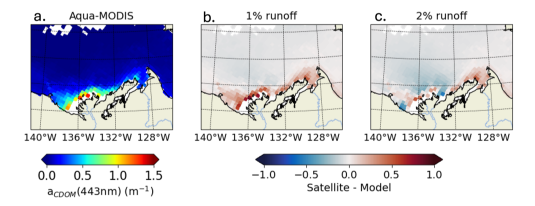


Figure B1. 2009 annual-mean CDOM absorption at 443 nm from (a) remotely-sensed observations and differences from simulated CDOM fields with (b) 1% and (c) 2% of tDOC redistributed into CDOM tracer. Simulated CDOM absorption is compared with satellite observations that are space-time colocated with the simulations.

Appendix C: Sea-ice phenology parameters

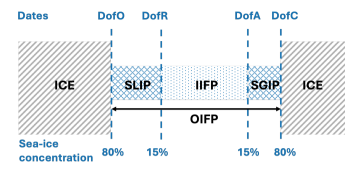


Figure C1. Conceptual diagram of sea ice seasonal evolution from spring/summer retreat (left) through fall/winter advance (right). Adapted from Bliss et al. (2019)

Appendix D: Comparison with observations

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We compared the 2012 weekly surface-ocean Chl-a (mg Chl-a m⁻³) and daily primary production (mg C m⁻² d⁻¹) simulated by ED-SBS (Run*etrl* and Run*full* and Run*full*) with satellite observations (Lewis et al., 2020) from Lewis et al. (2020) data estimated by the AOReg emp algorithm. We also compared simulated daily-mean SST (°C) for 2012 in both models with *in-situ* satellite observations (OSTIA Good et al., 2020). As both observational products have a finer horizontal resolution than grid-spacing compared to ED-SBS, we bin averaged the observations within each model grid cell. We then calculated the spatially-averaged value within the Mackenzie River plume region where satellite data were available to assess the model's ability to represent these observations (Figure 9 and D1). Note that the number of observations (n) available within the river plume area (n_{max}=281 for the entire area) varies in time. As both observational products provide sea-ice concentration data, we also calculated observed sea-ice phenology metrics to compare with our model simulations.

Within the Mackenzie River plumeregion, both simulations (Runctrl and Runfull) generally reproduce the amplitude of the phytoplankton bloom, since simulated surface-ocean Chl-a peaks roughly at the same concentration as observations; simulated NPP is underestimated by 25% (Figure D1a. and b.). By adding CDOM effects in Note that satellite observations from Lewis et al. (2020) AOReg, emp algorithm are the most suitable for model-observation comparison in the Mackenzie river plume; as they improve CDOM pollution removal in Chl estimates, using different fits of the Chl/R_{RS} (remote sensing reflectance) relationship for offshore region and shelf seas (where isobath < 1000 m). However, the model (Runfull), the phytoplankton bloom peaks during same week as observations (it peaks 2 weeks earlier without CDOM). This demonstrates that including CDOM effects improves ED-SBS's ability to represent phytoplankton bloom phenology. However, ED-SBS still does not capture the complete duration of the phytoplankton bloom as it simulates a bloom that lasts half as long compared to satellite observations. This difference is due to the sea-ice model's inability to realistically simulate the timing of seasonal loss of ice period (SLIP = DofR-DoO), Based on observations, the phytoplankton bloom initiates during the SLIP and peaks after the DofR (Figure D1b., pink line and grey area). In Run full, the model is able to reproduce this phenology in Beaufort Shelf being shorter than other Arctic Seas, the Mackenzie River plume spreads out further away from the Shelf. We then observed abrupt increase in Lewis et al. (2020) Chl estimates along the shelf (1000 m isobath) due to the start of the phytoplankton bloom (Figure D1b., purple line and blue area), but as sea-ice melt and break-up occurs later the simulation, the model is not able to match the start time of the observed phytoplankton bloom. In contrast, Runctrl (Figure D1b., green line) is able to match the initiation of the bloom only because the bloom peaks during the sea-ice melting period. Finally, the observed production remains high latter in the open-water season (August to September); ED-SBS is not able to sustain a high rate of productivity during this period as nitrate is entirely consumed, shunting down the bloomchange in regional Chl/R_{RS} fit (see Figure S3) and decided to remove offshore data from the comparison. This only affecting a small portion of the Northwestern plume region.

ED-SBS generally represents the spatially-averaged SST amplitude and variability in the Mackenzie River plume region during the open-water period compared to observations (Figure D1b9c.). The model underestimates SST by 25% from mid-July

to mid-September when CDOM is not included (Runetrl_{stel}). Adding CDOM effects improves simulated SST by decreasing this underestimate to 17% over the same period. However, similar to phytoplankton, we observe a delay in the surface-ocean heating, which is linked to later simulated SLIP. In the observations, SST starts increasing halfway through the melting season (Figure D19c., orange line and grey area). In both simulations, SST also increases halfway through the melting season (Figure D19c., purple and grey lines and blue area), but the SLIP occurs later in the season and thus surface-ocean warming also occurs later. We observe that this difference in June SST warming manifests mainly in the northwestern section of the plume where observed sea-ice melting occurs first (not shown). Simulated sea-ice first melts in the eastern section of the plume and propagates westward triggering a substantial difference in northwestern plume SST throughout the season compared to observations (Figure D1a,b,c). However, the simulated eastern plume SST is comparable to observation improving by 13% the correlation coefficient in the Mackenzie river plume. This confirms that sea-ice plays an important role in the ability of ED-SBS and expect to improve the sea-ice phenology by including fine-scale processes that drive sea-ice melting, such as explicit representation of leads and cracks.

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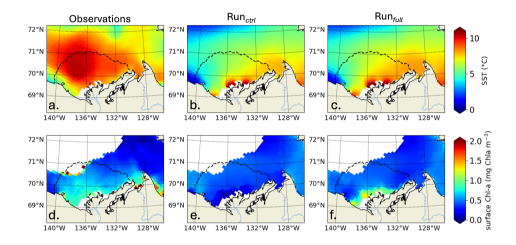


Figure D1. SST (a°C) averaged from July to September 2012 (top row) and Surface Chl-a (mg Chl-a m^{-3})—averaged from June to July 2012 (blower row) surface NPP-for observations (mg C m^{-3} d^{-1} a and d), and Run_{ctrl} (c and e)SST (C) spatially averaged over the Mackenzie River plume region for Runctrl (green line), Runfull full (purple line), satellite observations (pink line Lewis et al., 2020), c and in-situ/satellite observations (orange line Good et al., 2020). The blue (greyf)area indicates—the simulated (observed) seasonal loss dotted line shows the plume region; time-mean SSS isobaline of ice period; 27.

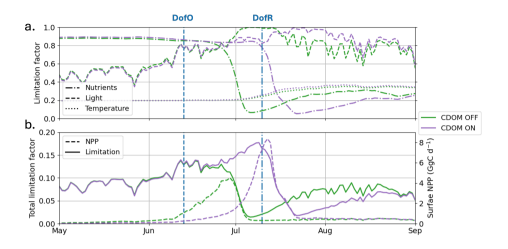


Figure E1. Simulated limitation factors for nutrients (solid lines), light (dashed lines), and temperature (dash-dotted lines), averaged between the two phytoplankton functional types and in the plume region for Run_{ctrl} (green lines) and Run_{full} (purple lines).

In ED-SBS, phytoplankton growth for each species (j) is limited by light, temperature, and nutrient availability. The three limitation factors (equations E1, E2, and E3), which yield values between 0 and 1, are combined (multiplied) to provide the total phytoplankton limitation factor. In this study, we average both phytoplankton type limitation factors in the plume region (Figure E1) to analyze the parameters affecting phytoplankton growth.

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The factor with the lowest value is generally considered as the factor limiting phytoplankton growth. In the Arctic Ocean, as ocean temperatures are typically low, temperature is a consistent limitation factor year-round, shaping the background state of phytoplankton growth. In the SBS, the temperature limitation factor (γ^{temp} ; dashed dotted lines in Figure E1) in the the plume region ranges from 0.2 in winter to 0.4 during the open-water season. This seasonal change is mainly due to increased light penetration as a result of melting of sea ice and mixing of Mackenzie River-derived freshwater into the coastal ocean. The inclusion of CDOM heating has a limited influence on phytoplankton limitation. Therefore, in this study temperature limitation drives a consistent dampening effect in phytoplankton growth but does not influence phytoplankton phenology.

In winter, elevated sea-ice cover causes high light limitation, with the spatially-averaged factor ranging between 0.4–0.6 in the plume region (Figure E1). Additionally, nutrients concentrations are high (see Figure 7), with a nutrient limitation factor over 0.8. Therefore, phytoplankton growth is limited by physics (both light and temperature). As sea ice begins to melt and break up (DofO), the light limitation factor increases up to 0.8 — triggering the start of phytoplankton growth in both simulations. By early June, as riverine CDOM spreads in the plume region, a difference of 0.02 in the light limitation occurs between the simulations with (Run_{full}) and without (Run_{ctrl}) CDOM effects. This small difference is sufficient to trigger a

slight difference in phytoplankton growth at this time, which allows the bloom to initiate in Run_{ctrl} (green dotted line; Figure E1b.) and thus delays the phytoplankton bloom by two weeks. In both simulation, elevated phytoplankton growth (mid-June or early July for Run_{ctrl} and Run_{full} , respectively) consumes nutrients, rapidly decreasing the nutrient limitation factor to \sim 0 and arresting the phytoplankton bloom. Therefore, as the nutrient limitation factor exceeds the temperature limitation factor (on July 3rd and July 15th without and with CDOM, respectively), the phytoplankton growth becomes primarily limited by nutrients. Analyzing each nutrient's (nitrate, phosphate, and silicate) limitation factor (not shown), we find that nitrate is the primarily limiting nutrient in the plume region.

In ED-SBS, the limitation factors for light (Equation E1), temperature (Equation E2), and nutrients (Equation E3) are computed using the following equations:

$$\begin{cases} \gamma_j^{light} = 1 - \exp\left(\frac{\langle \alpha I \rangle_j Chl - a: C_j}{P_j^{Cm}}\right) & \text{if } I_{tot} > I_{min} \\ \gamma_j^{light} = 0 & \text{else}, \end{cases}$$
 (E1)

where I_{tot} and I_{min} are the total and minimum light intensity for phytoplankton growth (W m⁻²), respectively, α is the Chl-a specific initial slope of the photosynthesis-light curve for each species, $Chl - a : C_j$ is the maximum Chl-a carbon ratio for each species, and P_i^{Cm} is the maximum growth rate.

$$\gamma_j^{temp} = c^{Arr} \left[e^{A_e^{Ar} ((T+273.15) - T_{ref}^{Ar^{-1}}) e^{-e_{2j|T-T_j^{opt}|^{p_j}}}} \right]_{>10^{-10}}, \tag{E2}$$

where T is the ocean temperature; c^{Arr} , A_e^{Ar} and T_{ref}^{Ar} the pseudo-Arrenhius equation coefficients set to 0.5882, -4000K, and 293.15K, respectively, and T_i^{opt} is the optimal phytoplankton temperature for each species.

The nutrient limitation factor takes the value of the most limiting factor between Phosphorus, Nitrogen, Silicate and Iron:

$$\gamma_{j}^{nut} = min(\gamma_{j}^{P}, \gamma_{j}^{N}, \gamma_{j}^{Si}, \gamma_{j}^{Fe}) \text{ and } \gamma_{j}^{N} = [\gamma_{j}^{NO_{3}} + \gamma_{j}^{NO_{2}} + \gamma_{j}^{NH_{4}}]_{0}^{1}$$
 (E3)

As nitrate is the primary limiting nutrient in this study, we describe below the nitrate limitation factor:

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$$\gamma_j^{N0_3} = \frac{N0_3}{N0_3 + k_j^{N0_3}} e^{-\sigma_j^{amm} NH_4},$$
 (E4)

where $k_j^{N0_3}$ is the half-saturation concentration for nitrate limitation and $\sigma_j^{amm}NH_4$ is the coefficient for NH_4 inhibition of $N0_3$ uptake.

Code and data availability. Model code and platform-independent instructions for running ED-SBS simulations are available at https://github.com/MITgcm-contrib/ecco_darwin/tree/master/regions/mac_delta/llc270/biogeochem_setup/carroll_2020_ecosystem/CDOM_setup.

Model output from all simulations described in this study (Run_{ctrl}, Run_{full}, Run_{light}, Run_{noriv}, and Run_{lin}) are available on the ECCO Drive: https://ecco.jpl.nasa.gov/drive/files/ECCO2/LLC270/Mac_Delta/CDOMsetup. ED-SBS forcing files are also available on the ECCO Drive at https://ecco.jpl.nasa.gov/drive/files/ECCO2/LLC270/Mac_Delta. Note that to access the ECCO Drive files users must register for a free Earthdata account at https://urs.earthdata.nasa.gov/users/new.

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 Software: CB. Supervision: VLF, DM and CM. Funding acquisition: VLF, DM and CM. Writing – original draft preparation: CB. Writing – review and editing: CB, VLF, DC, SD, DM, AM, MM and CM.

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