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## Evaluation of Under Sea-ice Phytoplankton Blooms in the Fully-Coupled, High-Resolution Regional Arctic System Model

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6 7 8 9	<sup>1</sup> Naval Postgraduate School <sup>2</sup> Institute of Oceanology Polish Academy of Sciences <sup>3</sup> Los Alamos National Laboratories <sup>4</sup> University of Alaska Fairbanks
10	Key Points:
11 12	• RASM reproduces the under sea-ice phytoplankton bloom observed in the Chukchi Sea in summer 2011
13 14	• Under sea-ice phytoplankton blooms are common in the Western Arctic and they occur under a specific combination of nutrient concentration and light
15 16	• High under sea-ice chlorophyll- <i>a</i> concentrations are also common in the Eastern Arctic, but they are at least in part a result of advection from open water blooms
17	upstream

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#### 18 Abstract

In July 2011, observations of a massive phytoplankton bloom in the ice-covered waters 19 of the western Chukchi Sea raised questions about the extent and frequency of under sea-20 ice blooms and their contribution to the carbon budget in the Arctic Ocean. To address 21 some of these questions, we use the fully-coupled, high-resolution Regional Arctic Sys-22 tem Model to simulate Arctic marine biogeochemistry over a thirty-year period. Our re-23 sults demonstrate the presence of massive under sea-ice blooms in the western Arctic not 24 only in summer of 2011 but annually throughout the simulation period. In addition, sim-25 ilar blooms, yet of lower magnitude occur annually in the eastern Arctic. We investigate 26 the constraints of nitrate concentration and photosynthetically available radiation (PAR) 27 on the initiation, evolution and cessation of under sea-ice blooms. Our results show that 28 increasing PAR reaching the ocean surface through the sea-ice in early summer, when 20 the majority of ice-covered Arctic waters have sufficient surface nitrate levels, is criti-30 cal to bloom initiation. However, the duration and cessation of under sea-ice blooms is 31 controlled by available nutrient concentrations as well as by the presence of sea-ice. Since 32 modeled critical PAR level are consistently exceeded in summer only in the western Arc-33 tic, we therefore conclude that the eastern Arctic blooms shown in our simulations did 34 not develop under sea ice, but were instead, at least in part, formed in open waters up-35 stream and subsequently advected by ocean currents beneath the sea ice. 36

#### <sup>37</sup> Plain Language Summary

In July 2011, scientists conducting research in the western Arctic Ocean observed 38 a large phytoplankton bloom under the sea-ice. Traditionally, such blooms were believed 39 to be rare. Using our state-of-the-art Arctic system model, were were able to demonstrate 40 that in fact, under sea-ice blooms have been occurring annually for the past several decades. 41 In the western Arctic, under sea-ice blooms begin when sufficient sunlight penetrates through 42 the sea-ice to the ocean, and end when nutrient concentrations become too low to sus-43 tain the phytoplankton. In the eastern Arctic, our model shows that under sea-ice blooms 44 still occur even when there is not enough sunlight penetrating the sea-ice. From this, we 45 conclude that phytoplankton blooms in the eastern Arctic begin in ice-free waters and 46 are advected beneath the sea-ice by ocean currents. 47

#### 48 1 Introduction

Marine phytoplankton have a strong effect on both the physical and the biologi-49 cal properties of the Arctic Ocean. In addition to its role in the regional carbon budget, 50 the presence of phytoplankton alters the optical properties of sea water, affecting wa-51 ter temperature, mixed layer depth, upper-ocean stratification, and sea-ice cover (Man-52 izza, Le Quere, Watson, & Buitenhuis, 2005). Phytoplankton also form the base of the 53 marine food web, supporting a wide variety of higher trophic organisms in pelagic com-54 munities (Grebmeier, Cooper, Feder, & Sirenko, 2006; Sigler et al., 2011), while the sink 55 of particulate organic matter produced by photosynthesis in the euphotic zone provides 56 the main food source in benthic communities (Grebmeier & Barry, 1991). 57

In high-latitude environments such as the Arctic Ocean, phytoplankton growth is 58 strongly constrained by light availability. Because light penetration into the upper ocean 59 is attenuated by snow and sea-ice cover, it was generally believed until recently that phy-60 toplankton growth was limited to areas of open water, with negligible growth under the 61 sea-ice. However, under sea-ice phytoplankton blooms have been reported multiple times 62 over the past several decades (e.g. Fortier, Fortier, Michel, and Legendre (2002); Fukuchi 63 et al. (1989); Hill, Light, Steele, and Zimmerman (2018); Legendre, Ingram, and Poulin (1989)). In July 2011, an ICESCAPE (Impacts of Climate on EcoSystems and Chem-65 istry of the Arctic Pacific Environment) survey observed a massive phytoplankton bloom 66 beneath the sea ice in the northern Chukchi Sea (Arrigo et al., 2012). The phytoplank-67

ton biomass was observed to be four times higher beneath the sea-ice than in the sur-68 rounding open water. The bloom extended more than 100 km beneath the pack ice with 69 peak particulate organic carbon biomass located near the shelf break, underlying thick 70 sea ice. The species observed indicate that pelagic diatoms were dominate, with a much 71 smaller contribution from ice algae. The growth of this under sea-ice bloom was supported 72 by areas of thinner first-year ice and, particularly, by the presence of melt ponds that 73 allowed for greater penetration of light. Observational evidence suggests that this bloom 74 was not an isolated case, and that under sea-ice blooms maybe widespread on the Arc-75 tic continental shelves (Arrigo et al., 2014; Lowry, van Dijken, & Arrigo, 2014) 76

The ICESCAPE observations have sparked increased interest in under sea-ice phy-77 toplankton blooms. Several model studies have been performed to assess the physical 78 conditions that favor the development of such blooms. Palmer et al. (2014) used a 1-D 79 ecosystem model to demonstrate that sea-ice conditions, particularly melt pond prolif-80 eration, contributed to under sea-ice bloom formation due to the enhanced light trans-81 mission through melt pond-covered sea ice. A coupled ice-ocean model study by Zhang 82 et al. (2015) has demonstrated a link between simulated under sea-ice blooms and in-83 creased light availability due to decreased snow cover; however, the model used in the 84 study did not include melt ponds. Horvat et al. (2017) formulated a model based on the 85 Sverdrup critical depth hypothesis (Sverdrup, 1953), suggesting that under sea-ice blooms 86 can form when melt point fraction exceeds a critical value  $\Phi_c$ , but this study did not ad-87 dress nutrient availability, which was demonstrated to be important by Zhang et al. (2015). 88

The purpose of this study is to examine the temporal and spatial evolution of un-89 der sea-ice blooms in the fully-coupled, bio-physical, high-resolution Regional Arctic Sys-90 tem Model (RASM) from 1980 to 2011. The model has been expanded to include ma-91 rine biogeochemistry (mBGC) in its ocean and sea ice components, with the latter in-92 cluding multiple options for melt pond representation. As such, RASM is a powerful tool 93 to investigate air-sea and bio-physical coupling in presence of sea ice at seasonal to multi-94 decadal time scales. We first evaluate the model bio-physical skill by comparing the mod-95 eled chlorophyll-a (chl-a) results against observations of the phytoplankton bloom in the 96 Chukchi Sea during July 3-8 2011, as reported by Arrigo et al. (2012). Next, we exam-97 ine multi-decadal variability of the under sea-ice chl-a and nutrient distributions, as well 98 as photosynthetically available radiation (PAR) for three different decade-apart years 99 spanning the period from 1991-2011. Finally, we discuss the relative controls of light avail-100 ability and nutrient supply on the initiation and evolution of under sea-ice phytoplank-101 ton blooms in two selected regions of the western and eastern Arctic, and the contribu-102 tion of these blooms to total primary production (PP). 103

#### <sup>104</sup> 2 The Regional Arctic System Model

RASM is a high-resolution, fully coupled atmosphere-ice-ocean-land regional model 105 with a domain encompassing the entire marine cryosphere of the Northern Hemisphere. 106 including the major oceanic inflow and outflow pathways, with mid-latitude extensions 107 into the North Pacific and North Atlantic oceans to account for the passage of cyclones 108 into the Arctic Ocean (Figure 1). The components of RASM are the Weather Research 109 and Forecasting (WRF) model, the Variable Infiltration Capacity (VIC) land hydrology 110 model with the river routing scheme (RVIC), and the Los Alamos National Laboratory 111 (LANL) Parallel Ocean Program (POP) and Sea Ice (CICE) Models. The model reso-112 lution is 50 km for WRF and VIC, and  $1/12^{\circ}$  (approximately 9km) for POP and CICE. 113 RASM has been demonstrated to correspond well with observations in its representa-114 tion of the upper-ocean physical dynamics (DuVivier et al., 2016; Hamman et al., 2017; 115 Roberts et al., 2015), arctic climate (Cassano et al., 2017; Hamman et al., 2017) and pro-116 cesses across the coupled atmosphere–land–ocean–sea ice interface (Brunke et al., 2018). 117 Because this study focuses on marine biogeochemistry, only the details of POP and CICE 118 are further described below. 119

#### 2.1 Physical ocean and sea-ice model

The POP and CICE configurations used in RASM are similar to the configuration 121 in the Community Earth System Model (CESM) version 1.1 (http://www.cesm.ucar 122 .edu) however, some adjustments have been made. In addition to changes needed to use 123 it as a regional model, POP has been modified to include a subgrid-scale brine rejection 124 parameterization of Jin, Deal, et al. (2012); Jin, Hutchins, Kawaguchi, and Kikuchi (2012), 125 which improves vertical ocean mixing under sea ice. Along the vertical axis, the model 126 is configured with 45 fixed-depth layers, including 7 layers in the top 42 m. The model 127 horizontal resolution of  $1/12^{\circ}$  is eddy-permitting across the entire RASM domain. The 128 combined effects of the fully coupled model, high spatio-temporal resolution and improved 129 parameterization of sub-grid scale bio-physical processes allowed reduction of biases in 130 physical and mBGC model outputs when compared to the coarse-resolution CESM model 131 (Jin et al., 2018).132

The CICE model (Roberts et al., 2018) in RASM is version 6, which includes, among 133 a number of changes, the latest column package modifications (Hunke, Lipscomb, Turner, 134 Jeffery, & Elliot, 2015, 2016). It has been configured to include mushy-layer thermody-135 namics (Turner & Hunke, 2015) and Elastic Anisotropic Plastic (EAP) sea ice rheology 136 (Wilchinsky & Feltham, 2004), as well as the explicit level ice pond parameterization 137 rather than the virtual melt ponds used in CESM. In addition, it uses new thermody-138 namic ocean coupling in which the basal freezing temperature is the same as the liquid 139 phase temperature within sea ice. RASM CICE uses five ice thickness categories, divided 140 at 0.65, 1.39, 2.47, 4.56 and 9.3 m. 141

The ice and ocean components are coupled using the coupler of Craig et al. (2012) with a coupling time step of 20 minutes to resolve sea ice-ocean inertial oscillations (Roberts et al., 2015)

The ocean and sea ice components were spun up for 78 model years, starting with 145 the initial conditions of no sea-ice and the ocean at rest. During this stage of initializa-146 tion, POP and CICE models were forced with CORE2 reanalysis (Large & Yeager, 2009). 147 Initial ocean temperature and salinity fields were from Polar science center Hydrographic 148 Climatology (PHC 3.0), (Steele, Morley, & Ermold, 2001). After the initial stage of spin 149 up the fully coupled version of RASM with bio-geochemical (BGC) components was run 150 for three years starting at the first of September 1979. This three-year period was re-151 peated three times in order to avoid the initial shock of any component, especially the 152 ocean and sea-ice BGC parts. The RASM production simulation, the results of which 153 are analyzed in this paper, was started in September 1979 and continued through the 154 end of 2018. The lateral boundary conditions at North Atlantic and Pacific sides utilized 155 temperature and salinity information (PHC 3.0) with 30-day restoring strength. The up-156 per and lateral atmospheric boundary conditions for the atmospheric model were based 157 on ERA-Interim reanalysis (Dee & Coauthors, 2011). 158

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#### 2.2 Marine biogeochemical model

The ocean BGC component in RASM is a medium-complexity Nutrients-Phytoplankton-160 Zooplankton-Detritus (NPZD) model (Moore, Doney, Kleypas, Glover, & Fung, 2002; 161 Moore, Doney, & Lindsay, 2004; Moore, Lindsay, Doney, Long, & Misumi, 2013). The 162 model has three phytoplankton categories: diatoms, small phytoplankton and diazotrophs, 163 with explicit carbon, iron (Fe) and chlorophyll-a (chl-a) pools for each category, as well 164 as an explicit Si pool for diatoms and an implicit CaCO<sub>3</sub> pool for small phytoplankton. 165 Other state variables are:  $NO_3 NH_4$ , Fe, Si, PO<sub>4</sub>, a herbivorous phytoplankton pool, 166 dissolved organic nitrogen, carbon, iron and phosphate (DON, DOC, DOFe, and DOP), 167 oxygen, dissolved inorgranic carbon (DIC) and alkalinity. 168

The column package version of CICE includes two BGC parameterizations. One 169 is a skeletal layer (SKL) parameterization in which all biological processes are assumed 170 to be confined to a 3-cm layer at the bottom of the ice. The other is a vertical (ZBGC) 171 parameterization in which biological activity is distributed throughout the ice column. 172 In this study, we use the ZBGC parameterization for all our simulations. The model in-173 cludes three algal categories (diatoms, small phytoplankton and *Phaeocystis sp*), two dis-174 solved organic carbon tracers (polysaccharids and lipids), a dissolved organic nitrogen 175 tracer, NO<sub>3</sub>, NH<sub>4</sub>, SiO<sub>3</sub>, dissolved Fe (FeD), dimethylsulfide (DMS), and dissolved and 176 particulate dimethylsulfoniopropionoate (DMSPd and DMSPp). Additional details on 177 the sea ice BGC component can be found in Jeffery et al. (2020). 178

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#### <sup>179</sup> 3 Conditions required for under-ice blooms

#### 3.1 Nutrient requirements

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Here we focus on nitrate as the limiting nutrient for under-ice blooms. A bloom occurs when phytoplankton growth rate exceeds the loss rate. If we assume that phytoplankton cell carbon to nitrogen ratios are fixed, then algal growth rate  $(G_{NO_3})$  and loss rate  $(L_{NO_3})$  are determined by nitrate uptake, and bloom permitting conditions occur when

$$L_{NO_3} = G_{NO_3} = \mu_{max} \left( \frac{NO_3}{NO_3 + \kappa_{NO_3}} \right), \tag{1}$$

where  $\mu_{max}$  is the maximum algal growth rate, and  $\kappa_{NO_3} = 2.5 \text{ mmol/m}^3$  is the half-saturation constant for nitrate uptake. The critical value of nitrate concentration necessary to produce a bloom can then be calculated as

$$NO_3 = \frac{L_{NO_3} \times \kappa_{NO_3}}{\mu_{max} \left(1 - \frac{L_{NO_3}}{\mu_{max}}\right)}.$$
(2)

 $L_{NO_3} \text{ and } \mu_{max} \text{ are both temperature-dependent quantities, with the temperature }$   $dependence being defined in RASM as T_{dep} = 2.0^{((T+273.16)-(30.0+273.16))/10.0}. Because$  our analysis focuses on chl-a in the surface layer, T is assumed to be -1.8° C as a rep-  $resentative surface water temperature under the ice. Thus, \mu_{max} can be defined as PC_{ref}T_{dep},$   $where PC_{ref}=4.8 \text{ days}^{-1} \text{ is the maximum diatom growth rate at } T_{ref}=30°C \text{ and } L_{NO_3}$   $= mort \times T_{dep}, \text{ where mort}=0.15 \text{ day}^{-1} \text{ is the diatom mortality rate. Given these assumptions, the critical nitrate concentration required to permit a bloom is NO_3=0.08 \text{ mmol/m}^3.$ 

#### 3.2 Light requirements

When sufficient nutrients are available, the PAR becomes the limiting factor for 197 under-ice blooms. In their model study based on the Sverdrup critical depth hypothe-198 sis (Sverdrup, 1953), Horvat et al. (2017) related PAR beneath the ice to melt ponds con-199 centration, computing a critical melt pond fraction that would be necessary for an under-200 ice bloom to occur. Here we adopt a number of Horvat et al.'s assumptions, as well as 201 their use of Sverdrup's hypothesis, to estimate the critical PAR levels necessary for an 202 under-ice bloom. While RASM does not include PAR among its variables, it does include 203 shortwave radiation through the ice to the ocean surface. We therefore use the 0.43 PAR 204 to shortwave ratio estimated by Olofsson, Van Laake, and Eklundh (2007) to determine 205  $I_0$ , or PAR at the ocean surface under the ice. PAR at depth z can then be computed 206 as 207

$$I(z) = I_0 e^{\kappa_w z},\tag{3}$$

where  $\kappa_w = 1.2 \text{ m}^{-1}$  is the bulk irradiance coefficient of PAR in clear water (Pegau, 2002).

Following Horvat et al. (2017), we assume a constant, depth-independent phytoplankton decay rate  $\Gamma$  (m<sup>-1</sup>s<sup>-1</sup>) throughout the mixed layer. We then determine the phytoplankton growth rate as

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$$G(D) = \frac{M}{D} \int_{-D}^{0} I(z) dz \tag{4}$$

where D is the mixed layer depth, and M is a coefficient relating phytoplankton growth rate to PAR availability, so that  $\Gamma/M$  is the compensation irradiance. For our calculations, we used the Eastern Arctic compensation irradiance estimates of Regaudie-de-Gioux and Duarte (2010) and set  $\Gamma/M = 1.3$  mol quanta m<sup>-2</sup> day<sup>-1</sup>, which was multiplied by a conversion factor for PAR from sunlight of 2.5 W mol<sup>-1</sup> day<sup>-1</sup> to give us  $\Gamma/M = 3.25$ W m<sup>-2</sup>.

As discussed in the section above, bloom conditions require that the phytoplankton growth rate exceeds the loss rate. We can therefore determine the critical value for  $I_0$  by setting  $G(D) = \Gamma$  and solving for  $I_0$  to get

$$I_0 = D\kappa_w \frac{\Gamma}{M} (1 - e^{-\kappa_{wD}})^{-1}.$$
(5)

#### 4 Model results

Under sea-ice blooms in the Arctic are composed primarily of diatoms, both in ob-222 servations (Arrigo et al., 2014) and in RASM. We therefore focus on diatoms in our eval-223 uation of model results. All references to chl-a concentration and primary production 224 in this paper refer specifically to diatom values. Figure 2 (a) shows modeled surface chl-225 a distribution in the northern Chukchi Sea during July 3-8 2011, corresponding to the 226 dates when an under-ice bloom was detected in the region during the 2011 ICESCAPE 227 survey (Arrigo et al. 2012). While a bloom is present in the model in the north-west cor-228 ner of the region, it is located further north than the observed bloom, and the modeled 229 chl-a concentrations are lower than the observed concentrations. Modeled ice concen-230 trations for this time period (shown as red contours in Figure 2) indicate that the mod-231 eled sea ice has retreated further north compared to the satellite-observed sea ice con-232 ditions, and surface nitrate concentrations (not shown) are near zero throughout the re-233 gion, suggesting that the model bloom has peaked earlier in the season and has consumed 234 most available nutrients. The chl-a distribution for Jun 20-24 2011 (Figure 2 (b)), when 235 modeled sea ice extent was similar to observed extent during the ICESCAPE cruise, shows 236 improved correspondence to the observed bloom, with the location and spatial extent 237 of the modeled bloom being similar to observations. However, while the maximum mod-238 eled chl-a value for the region is  $21.45 \text{ mg/m}^3$ , the observed values reach as high as 64.7239  $mg/m^3$ . Point-to-point comparison of modeled versus observed chl-a values is shown in 240 Figure 3. The discrepancy between maximum modeled and observed values might be due 241 to several reasons. One is the fact that that model surface atmospheric conditions, such 242 as winds, clouds, radiative fluxes which force sea ice, are not prescribed from a reanal-243 ysis but 'predicted' from an active atmospheric model in the fully coupled configuration 244 of RASM. Another possible reason could be the fact that the ocean model's horizontal 245 resolution, while eddy-permitting, is not eddy-resolving and likely doesn't fully capture 246 the mesoscale ocean dynamics, hence also small-scale local chl-a gradients that are seen 247 in the observations. Additional discrepancies in the modeled sea ice cover might be re-248 lated to inadequate coupling of horizontal momentum transfer across the atmosphere -249 sea ice - ocean interface. 250

When considering model results over a larger region of the western Arctic (Figure 4), it is clear that that the full extent of the 2011 under sea-ice bloom in the western Arctic was significantly larger than the area covered by the ICESCAPE observations. This <sup>254</sup> bloom appears annually throughout the simulation period, indicating that under-ice blooms <sup>255</sup> are not a recent phenomenon. At the same time model results reveal relatively signif-<sup>256</sup> icant variability in the under sea-ice bloom distribution, magnitude and timing. To demon-<sup>257</sup> strate this, Figures 4, 6 and 8 are shown with the modeled surface chl-*a* distributions <sup>258</sup> for the western Arctic bloom (WB) region, designated as the region between latitudes <sup>259</sup> 70° N and 78° N and longitudes 150° E to150° W, for the years 2011, 2001 and 1991.

In addition to the western Arctic bloom region described above, a second bloom of similar duration but of lower chl-a is simulated by the model in the eastern Arctic, and it also varies in distribution, magnitude and timing. Figures 5, 7 and 9 show the surface chl-a distributions within the eastern Arctic bloom (EB), designated as the region between latitudes 75° N and 85° N and longitudes 0° to 90° E, for the same years 2011, 2001 and 1991 as in the case of and for comparison with the WB results.

#### <sup>266</sup> 5 Discussion

To summarize the above results, the left side of Figure 10 shows the time series 267 of spatially-integrated surface chl-a for both the EB and WB regions during May, June 268 and July of 1991, 2001 and 2011. Previous satellite observations suggest that Arctic spring 269 phytoplankton blooms are beginning to occur earlier in the year due to earlier sea-ice 270 breakup and decreasing sea-ice concentrations in early summer (Kahru, Brotas, Manzano-271 Sarabia, & Mitchell, 2011). Our results show a similar pattern for the eastern bloom, 272 which reached peak chl-a levels during June 19-23 in 1991, June 12-16 in 2001, and June 273 7-11 in 2011. However, the western bloom does not show the same pattern, with the 2001 274 peak (June 27-July 1) occurring later that the 1991 peak (June 22-26). 275

For both regions, the post-peak decline of total under sea-ice chl-a is affected by 276 decreasing sea-ice coverage as well as by phytoplankton mortality. The EB region shows 277 similar coverage for all three years of our analysis, with approximately 50% of the re-278 gion still being covered by sea-ice by the end of July. In particular, 2001 and 2011 both 279 show EB chl-a total decreasing from mid-June through July at a faster rate that the sea-280 ice coverage, indicating that the decrease is due primarily to phytoplankton loss. This 281 loss cannot be explained by nutrient depletion, as is discussed below. By contrast, the 282 rate of total chl-a decrease in the WB region corresponds more closely to the rate of sea-283 ice decrease, particularly in 2011, when under sea-ice chl-a declines to near zero at the 284 same rate that the region becomes ice-free, suggesting that little to no actual phytoplank-285 ton loss is taking place. This is further supported by the dashed lines in the top panel 286 of Figure 11, which show the PP for the entire WB remaining relatively constant after 287 the under sea-ice bloom peaks. 288

Rows 2-4 in Figures 4 to 9 show the PAR (row 2) nutrient (row 3) and combined 289 nutrient and PAR (row 4) conditions for the western and eastern under-ice blooms. For 290 both regions, the period of May 15-19 in all years has been designated as "pre-bloom." 291 while the period of July 21-25 was designated as "post-bloom." Red areas in the figures 292 indicate the regions where nutrient and light conditions meet the critical requirements 293 for bloom formation, as discussed in sections 3.1 and 3.2. During the pre-bloom period, 294 nitrate concentrations are above the critical threshold throughout the ice-covered Arc-295 tic, reflecting the build-up of nutrients during the winter. As the blooms progress, nu-296 trient concentrations become reduced, with the greater decrease occurring in the west-297 ern bloom region; the eastern and central Arctic remain nutrient-replete even into the 298 post-bloom period. Therefore, the beginning of the under sea-ice blooms is triggered by 200 PAR availability, after which the blooms persist until the available nutrients are depleted 300 or until the region becomes ice-free (at which point the bloom is no longer considered 301 an under sea-ice bloom). The critical PAR criteria of Equation 5 can thus allow us to 302 distinguish between true under-ice blooms and blooms that originally formed in open 303 water and were subsequently advected beneath the ice. The majority of EB area in RASM 304

does not meet the critical PAR criteria in 2011 or 1991. However, the entire EB region still has a bloom in all three years of our study, suggesting an advective origin. This conclusion is consistent with the study of Johnsen et al. (2018), in which an under-ice bloom was observed in May 2010 northwest of Svalbard in our EB region and attributed to advection, with northward flowing water masses and southward flowing sea ice.

The advective origin of the EB is further supported by the differences in the chl-310 a and PP time series for the two blooms. The April chl-a totals integrated over the top 311 150 m for the WB (Figure 10 (a)) are near zero for all years shown. In 2011, the chl-a312 totals drop to zero again by August, while in 2001 and 1991 the totals drop below  $5 \times 10^7$ 313 kg but remain non-zero. This is consistent with the PP time series for the region (Fig-314 ure 11 (a)), which begins near zero in April for all years and decreases to zero again in 315 August for 2011 but not for the other two years. In contrast, the EB chl-a totals for all 316 three years begin at approximately  $1.5 \times 10^7$  kg in April and do not drop below  $2 \times 10^7$ 317 kg throughout the time series. Likewise, the PP for the EB remains non-zero for the en-318 tire time span shown in Figure 11. In fact, the full-year PP time series for the EB (not 319 shown) indicates positive PP totals in the top 150 m starting in mid-February. The EB 320 region does not have sufficient light to support photosynthesis that early in the year, in-321 dicating that the chl-a and PP totals must be the result of a bloom advected from an 322 ice-free location farther south. While the maximum PP totals for the EB region are ap-323 proximately half of the WB totals, the early start and long duration of the EB still make 324 it a significant factor in the total PP for the Arctic region. 325

RASM simulations indicate that favorable PAR conditions for under-ice blooms have 326 existed in the western Arctic at least as far back as 1991, allowing massive blooms to oc-327 cur annually on the shelf and along the shelf break in that region. Satellite-derived es-328 timates of sea-ice thickness have indicated that Arctic sea ice has been growing thinner 329 since at least 1982, as multi-year ice is replaced by first-year ice (Maslanik et al., 2007). In addition, the presence and extent of melt ponds on the surface of the sea ice in the 331 Western Arctic has been increasing over the last few decades (Hutchings & Faber, 2018). 332 The model results suggest that if these trends continue, PAR penetration through sea-333 ice to the ocean surface will increase, leading to larger and earlier-occurring under sea-334 ice blooms, with a corresponding increase in Arctic primary production and nutrient con-335 sumption. In situ sampling in the EB and WB regions in the next few years would serve 336 to confirm these results and provide a clearer picture of the effects of sea-ice reduction 337 on Arctic Ocean biogeochemistry. 338

#### <sup>339</sup> Figure Captions

Figure 1: The atmosphere/land and ice/ocean domains of the Regional Arctic System Model

Figure 2: Modeled surface chl-a distribution in the northern Chukchi Sea during July

- 343 3-8 2011 (a) and Jun 20-24 2011 (b). Circles represent the locations and observed sur-
- $_{344}$  face chl-*a* concentrations for hydrographic stations sampled during the July 2011 ICESCAPE

cruise (Arrigo et al., 2012). Red lines indicate modeled ice concentration; green lines in

both panels indicate observed ice concentration from satellite during the ICESCAPE cruise.

Figure 3: Surface chl-a distributions for the hydrographic stations shown in Figure 2 and for the corresponding model grid cells.

Figure 4: Top row shows the modeled ocean surface chl-*a* distributions before, at peak chl-*a*, and after the Western Arctic Bloom during 2011 in the region where ice fraction is greater than 50%. Red areas in the second row indicate the regions where PAR through the ice to the ocean surface exceeds the critical value as determined in Section 3.2. Red areas in the third row indicate the regions where surface nitrate concentration exceeds the critical value as determined in Section 3.1. Red areas in the bottom row indicate the

- regions where both PAR and nitrate exceed the their critical values. Pink contour in-
- dicates the region of the 2011 ICESCAPE cruise (Arrigo et al., 2012)
- <sup>357</sup> Figure 5: Same as in Figure 5, but for the Eastern Arctic Bloom.
- Figure 6: Same as in Figure 5, but for the Western Arctic Bloom in 2001.
- Figure 7: Same as in Figure 5, but for the Eastern Arctic Bloom in 2001.
- Figure 8: Same as in Figure 5, but for the Western Arctic Bloom in 1991.
- Figure 9: Same as in Figure 5, but for the Eastern Arctic Bloom in 1991.

Figure 10: Time series of spatially integrated surface chl-a (green lines) and percentage

sea-ice area (black lines) for the eastern and western under sea-ice bloom areas for the

years 1991, 2001 and 2001. Vertical green bars delineate the 5-day period surrounding

- the date of the chl-a maximum for each time series. Vertical gray bars delineate July 3-
- <sup>366</sup> 8 2011. Dashed lines indicate the pre-bloom and post-bloom periods as shown in the first and last columns of Figures 4.0
- and last columns of Figures 4-9

Figure 11: Spatially-integrated model primary production for the WB and EB regions for May, June and July of 1991, 2001 and 2001. Dashed lines represent primary production for the entire region. Solid lines represent primary production for the portion of the region where ice concentration is greater than 50%.

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<sup>372</sup> Figure 1. The atmosphere/land and ice/ocean domains of the Regional Arctic System Model



373	<b>Figure 2.</b> Modeled surface chl- <i>a</i> distribution in the northern Chukchi Sea during July 3-8
374	2011 (a) and Jun 20-24 2011 (b). Circles represent the locations and observed surface chl- $a$ con-
375	centrations for hydrographic stations sampled during the July 2011 ICESCAPE cruise (Arrigo
376	et al., 2012). Red lines indicate modeled ice concentration; green lines in both panels indicate
377	observed ice concentration from satellite during the ICESCAPE cruise.



Figure 3. Surface chl-a distributions for the hydrographic stations shown in Figure 2 and for the corresponding model grid cells.



Figure 4. Top row shows the modeled ocean surface chl-a distributions before, at peak chl-a, 380 and after the Western Arctic Bloom during 2011 in the region where ice fraction is greater than 381 50%. Red areas in the second row indicate the regions where PAR through the ice to the ocean 382 surface exceeds the critical value as determined in Section 3.2. Red areas in the third row in-383 dicate the regions where surface nitrate concentration exceeds the critical value as determined 384 in Section 3.1. Red areas in the bottom row indicate the regions where both PAR and nitrate 385 exceed the their critical values. Pink contour indicates the region of the 2011 ICESCAPE cruise 386 (Arrigo et al., 2012) 387



Figure 5. Same as in Figure 5, but for the Eastern Arctic Bloom.





Figure 6. Same as in Figure 4, but for the Western Arctic Bloom in 2001





Figure 7. Same as in Figure 4, but for the Eastern Arctic Bloom in 2001





Figure 8. Same as in Figure 4, but for the Western Arctic Bloom in 1991

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Figure 9. Same as in Figure 4, but for the Eastern Arctic Bloom in 1991.



Figure 10. Time series of spatially integrated surface chl-*a* (green lines) and percentage seaice area (black lines) for the eastern and western under sea-ice bloom areas for the years 1991, 2001 and 2001. Vertical green bars delineate the 5-day period surrounding the date of the chl-*a* maximum for each time series. Vertical gray bars delineate July 3-8 2011. Dashed lines indicate the pre-bloom and post-bloom periods as shown in the first and last columns of Figures 4-9



Figure 11. Spatially-integrated model primary production for the WB and EB regions for May, June and July of 1991, 2001 and 2001. Dashed lines represent primary production for the entire region. Solid lines represent primary production for the portion of the region where ice concentration is greater than 50%.