# Annual boom-bust cycles of polar phytoplankton biomass revealed by space-based lidar

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Polar plankton communities are among the most productive, seasonally dynamic and rapidly changing ecosystems in the global ocean. However, persistent cloud cover, periods of constant night and prevailing low solar elevations in polar regions severely limit traditional passive satellite ocean colour measurements and leave vast areas unobserved for many consecutive months each year. Consequently, our understanding of the annual cycles of polar plankton and their interannual variations is incomplete. Here we use space-borne lidar observations to overcome the limitations of historical passive sensors and report a decade of uninterrupted polar phytoplankton biomass cycles. We find that polar phytoplankton dynamics are categorized by 'boom-bust' cycles resulting from slight imbalances in plankton predator-prey equilibria. The observed seasonal-to-interannual variations in biomass are predicted by mathematically modelled rates of change in phytoplankton division. Furthermore, we find that changes in ice cover dominated variability in Antarctic phytoplankton stocks over the past decade, whereas ecological processes were the predominant drivers of change in the Arctic. We conclude that subtle and environmentally driven imbalances in polar food webs underlie annual phytoplankton boom-bust cycles, which vary interannually at each pole.

hip- and aircraft-mounted Light Detection And Ranging (lidar) systems have been employed for decades to characterize ocean optical, phytoplankton, zooplankton, and fish properties<sup>1-4</sup>. However, global-scale plankton retrievals from a space-based lidar were first demonstrated only in 2013<sup>5</sup> with measurements from the Cloud-Aerosol Lidar with Orthogonal Polarization (CALIOP) sensor<sup>6</sup>. These active lidar measurements retrieve ocean plankton properties both day and night, penetrate between and through significant cloud cover, and have minimal atmospheric correction errors, thus addressing many limitations of passive ocean colour measurements particularly in polar regions<sup>7</sup>. Here we use 1°-latitude binned ocean particulate backscattering coefficients  $(b_{bp})$  and diffuse attenuation coefficients  $(K_{D532})$ determined from CALIOP's 532 nm polarization channels<sup>5,8,9</sup> to quantify phytoplankton biomass<sup>10-12</sup> in the surface mixed layer over the period 2006 to 2015 (Methods). CALIOP is a nadirviewing sensor providing measurements along its orbit track to 81.5° latitude, with spatial coverage increasing with increasing latitude (Fig. 1c,g and Supplementary Fig. 1). Ocean properties retrieved by CALIOP are from its first 22.5 m vertical-resolution sampling bin below the surface<sup>5</sup>. Over its 16-day orbit repeat cycle, CALIOP views roughly 70% of the total 1°-resolution ocean bins poleward of 45° latitude (Supplementary Fig. 2).

#### Annual cycles in polar phytoplankton biomass

Before evaluating polar plankton properties, a 'comparison zone' of 45° to 55° latitude was defined where ocean observations are available throughout the year from both CALIOP and the Moderate-resolution Imagining Spectrometer (MODIS) passive ocean colour

sensor (Fig. 1). For this comparison zone, we find an excellent agreement (north  $r^2 = 0.84$ , n = 111, p < 0.001; south  $r^2 = 0.79$ , n = 111, p < 0.001) between phytoplankton biomass cycles from the two sensors (Fig. 1d,h), which lends confidence to the CALIOP retrievals. During periods when polar data ( $\geq 60^{\circ}$  latitude) are also available from both sensors, we again find an excellent agreement (north  $r^2 = 0.63$ , n = 84, p < 0.001; south  $r^2 = 0.81$ , n = 72, p < 0.001) between MODIS and CALIOP (Supplementary Figs 3 and 4). In these polar zones, however, CALIOP continues to characterize phytoplankton biomass distributions right up to the ice edge (Fig. 1b,f) throughout the extended periods when MODIS data are entirely absent (for example, Fig. 1a,e). This advantage of active lidar sensing allows CALIOP to 'fill in' the late autumn to earlyspring seasons hidden from ocean colour sensors and thus construct the first complete record of polar phytoplankton annual biomass cycles (Fig. 2a,c).

Phytoplankton concentrations retrieved by CALIOP in the northern and southern polar zones follow repeated winterminimum and summer-maximum annual cycles, with secondary interannual variations in cycle amplitude (black symbols in Fig. 2a,c; range in amplitude of 24 to 45 mg C m<sup>-3</sup> in the north and 11 to 21 mg C m<sup>-3</sup> in the south) but no long-term trend (p > 0.05) over the CALIOP record. Throughout these time series, biomass changes show a clear correspondence ( $r^2 = 0.78$ ) with mixed-layer phytoplankton division rates ( $\mu$ ; blue lines in Fig. 2a,c), calculated as:

$$\mu = \frac{\int_{t=\text{sunrise}}^{\text{sunset}} \int_{z=0}^{\text{MLD}} P_{\text{max}}^{C} \tanh\left(\frac{\alpha^{C} \text{PAR}_{t,z}}{P_{\text{max}}^{C}}\right) dz dt}{\text{MLD}}$$
(1)

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**Figure 1** | **Phytoplankton biomass observations from CALIOP and MODIS. a**, MODIS phytoplankton biomass retrievals poleward of 45° latitude (white: no data) for northern latitudes (December 2010). **b**, CALIOP phytoplankton biomass retrievals for December 2010. **c**, Black lines: CALIOP orbit tracks poleward of 45° latitude. The red dashed ring demarks the 45°-55° 'comparison zone'. Yellow rings demark the north polar zone (60°-81.5° latitude). **d**, 2006-2015 monthly mean phytoplankton biomass for the north 'comparison zone'. **e-h**, As in **a-d**, respectively, but for southern latitudes during June 2010 (south polar zone in **f**: 60°-75°). Grey shading: ice cover in panels **a,b** and **e,f**.

where, MLD is mixed-layer depth (m),  $PAR_{t,z}$  is photosynthetically active radiation at time *t* and depth *z*, and  $\alpha^{C}$  and  $P_{max}^{C}$ are the carbon-specific light-limited and light-saturated rates of photosynthesis (Methods). This correlation between biomass and division rate, however, does not imply that rapid division causes high biomass.

Temporal changes in phytoplankton concentration (*C*, mg C m<sup>-3</sup>) reflect imbalances between division and loss (grazing, viral lysis, and so on) rates. This disequilibrium is quantified for a given time interval,  $t_0$  to  $t_1$ , through the specific rate of biomass change (r; d<sup>-1</sup>) (refs 13,14):

$$r = \frac{\ln\left(\frac{C_{t_1}}{C_{t_0}}\right)}{t_1 - t_0} \tag{2}$$

For both the north and south polar zones, the annual cycle in r shows that the spring increase in phytoplankton concentration begins (that is, r first becomes positive) when division rates are

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**Figure 2** | **Polar phytoplankton cycles. a,c**, Black symbols: CALIOP monthly mean phytoplankton biomass (*C*). Blue line: phytoplankton division rates ( $\mu$ ). Green line: phytoplankton loss rates (*I*, which are indistinguishable from  $\mu$ ). Red line: calculated *C* time series from model predictions of  $d\mu/dt$  (Methods). **b,d**, Open symbols, black line: biomass rates of change (*r*) calculated from CALIOP observations (equation (2)). Red line: modelled  $d\mu/dt$  (equations (1) and (3)). Statistics on relationships between *r* and  $d\mu/dt$  are:  $r^2 = 0.63$ , n = 110, p < 0.001 (**b**) and  $r^2 = 0.71$ , n = 110, p < 0.001 (**d**). Vertical tan bars: months with no MODIS data.

near their annual minimum (Fig. 3; purple dashed arrows). At the other extreme, bloom termination (that is, r first becomes negative) occurs when division rates are maximal (Fig. 3; green dashed arrows). Furthermore, the most rapid rate of biomass increase in spring (Fig. 3; black dashed arrows) coincides with division rates equivalent to those in autumn when biomass is most rapidly declining (Fig. 3; brown dashed arrows). These findings clearly demonstrate that the absolute rate of division defines neither the rate nor sign of biomass change.

The seemingly inconsistent annual cycles in polar biomass and division rate (Fig. 3) are reconciled if, as is the case for planktonic food webs<sup>13-15</sup>, phytoplankton loss rates  $(l; d^{-1})$  are closely coupled to, but slightly time-lagged behind, phytoplankton division rates  $(\mu)$  (ref. 15) (Fig. 2a,c). For a time lag of 1 day (that is,  $l_{t_1} = \mu_{t_0}$  and  $t_1 - t_0 = \Delta t = 1$  day), the specific rate of change in biomass (r) is then determined by the rate of change in division  $(d\mu/dt; d^{-2})$ :

$$r_{t1} = \mu_{t1} - l_{t1} = \mu_{t1} - \mu_{t_0} = \frac{\mathrm{d}\mu}{\mathrm{d}t} \Delta t \tag{3}$$

This relationship clearly emerges from the CALIOP record, where we find highly correlated and comparable magnitude annual cycles in  $d\mu/dt$  and *r* for both polar zones (Figs 2b,c and 3 top panels,

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**Figure 3** | **Climatological annual phytoplankton cycles.** Top panels: open symbols, black line: CALIOP-observed biomass change rate (r, grey shading indicates ±1 s.d.). Red line: modelled  $d\mu/dt$  (equations (1) and (3)). The horizontal dashed line demarks r=0. Below this line biomass is decreasing, while above this line biomass is increasing. Bottom panels: annual cycles in division rate. Top and bottom panels are connected by four key events: purple arrow: initial rise in biomass; black arrow: maximum biomass increase rate; green arrow: bloom termination; brown arrow: maximum rate of biomass decline. Upper brackets indicate month range of event. Lower boxes indicate range in  $\mu$  for each event (Methods).

Methods). Thus, the blooming phase of the annual cycle begins when division rates are minimal but accelerating (that is,  $d\mu/dt$  is positive) and then ends when division reaches its maximum rate (that is, acceleration stops)<sup>15</sup> (Fig. 3). Conversely, the declining phase of the annual cycle results from a deceleration in division rate (that is,  $d\mu/dt$  is negative)<sup>15</sup>, despite absolute rates of division being comparable to those during the blooming phase (Fig. 3).

Our results indicate that the large-amplitude biomass cycles of polar systems (Fig. 2a,c) are a consequence of division and loss rates being perpetually decoupled through environmental changes in upper-ocean growth conditions. By extension, it may be anticipated that interannual variations in biomass cycles will likewise reflect environmentally controlled changes in annual minimum-maximum ranges for division rate (that is, the ability for division to accelerate and decelerate). This prediction is confirmed in the CALIOP record for the north and south polar regions, where 75% (n = 34, p < 0.001) of the interannual differences in minimum-to-maximum and maximum-to-minimum biomass change rates (equation (2)) are accounted for by our modelled ranges in  $d\mu/dt$  (equations (1) and (3)) (Fig. 2b,c). The fidelity of this finding is further verified when modelled  $d\mu/dt$  values are applied to initial phytoplankton concentrations for the north and south polar zones and the entire decade of CALIOP-observed biomass changes are effectively recaptured (north  $r^2 = 0.78$ , n = 111, p < 0.001; south  $r^2 = 0.81$ , n = 111, p < 0.001) without any additional 'restoring' of the modelled time series to measured values (red line in Fig. 2a,c).

#### Interannual variations in polar phytoplankton biomass

Variations in phytoplankton concentration are only one factor impacting the broader biogeochemistry of the polar oceans. A second critical property is the extent of ice-free ocean area (IFA)<sup>16-19</sup>, with long-term trends in IFA (Supplementary Fig. 4) serving as harbingers of climate change. During the recent era of CALIOP measurements, seasonal anomalies in polar-zone integrated phytoplankton stocks ( $\Sigma C_{polar}$ ; Fig. 4 black symbols) have spanned similar ranges in the north (3.2 Tg C) and south (2.4 Tg C), but for different reasons. In the north, seasonal anomalies in mean phytoplankton concentration (mg C m<sup>-3</sup>) have dominated  $\Sigma C_{\text{polar}}$  variability (Fig. 4c;  $r^2 = 0.80$ , n = 37, p < 0.001), whereas IFA anomalies have driven variability in the south polar zone (Fig. 4b;  $r^2 = 0.72$ , n = 37, p < 0.001) (Supplementary Table 1). One reason for this difference is that surface nutrients in the polar north are often not fully depleted at the bloom climax, implying a potential for enhanced division rates during favourable years. In the south, the prevalence of iron-limiting conditions<sup>20,21</sup> is likely to play a key role in constraining anomalies in phytoplankton concentration (Fig. 4d). The second reason for the north-south difference is that IFA anomalies in the polar north were small over the CALIOP era (Fig. 4a), whereas changes in the polar south were the largest since the beginning of satellite ice-cover record in 1978 (Supplementary Fig. 5). These contrasting results for the two polar biomes highlight the importance of both ecological processes (for example, light, nutrients, grazing controls) and physical properties (for example, ice cover) in regulating plankton stocks and raise questions on what these changes observed today imply about polar ecosystem trajectories into the future.

#### Extending knowledge on planktonic ecosystems

Advantages of active satellite lidar sensing have enabled uninterrupted monitoring of polar plankton populations since 2006. Our CALIOP-based analysis provides the strongest observational



**Figure 4** | Interannual changes in polar-zone phytoplankton biomass. Black symbols, black line: seasonal anomalies in polar-zone integrated photic-layer phytoplankton biomass ( $\Sigma C_{polar} =$  phytoplankton concentration (mg C m<sup>-3</sup>) × photic-zone depth (m) × IFA (m<sup>2</sup>)/10<sup>12</sup> mg Tg<sup>-1</sup>). **a,b**, Blue symbols, blue line: ice-free-area (IFA) seasonal anomalies. **c,d**, Red symbols, red line: phytoplankton concentration seasonal anomalies. Anomalies are calculated as the difference between a property value for a given season and the average value for that season over the 2006 to 2015 CALIOP era. Seasons: December-February, March-May, June-August and September-November.

evidence to date that large-scale variability in polar phytoplankton concentrations is governed by the extent to which mixed-layer growth conditions allow division rates to accelerate or decelerate in time<sup>15</sup>. The temporal lag between phytoplankton division and loss rates implied by our data is on the order of days (Methods), but further studies are necessary to evaluate whether such tight predator-prey coupling operates at finer space and time scales. The finding that annual polar blooms terminate when division rates are maximal (Figs 2 and 3) is consistent with an earlier proposal regarding global phytoplankton biomass dynamics<sup>15</sup> and may provide additional insight into phytoplankton community structure during a bloom. Specifically, if our current results for total phytoplankton biomass can be extended to the species level, then an implication is that species with high light-use efficiencies and low maximum growth rate potentials will have an initial advantage under low-light conditions, but will saturate in division early during the blooming phase and thus soon be overcome by losses. On the other hand, species with high potential growth rates can continue to accelerate division ahead of loss rates and, accordingly, may ultimately come to dominate biomass at the bloom climax. Consistent with this view, diatoms are often noted for their exceptionally high division potentials<sup>22-24</sup> and their frequent dominance of climax bloom populations<sup>25-27</sup>. Extending this thought further, it might also be speculated that the dominance of diatoms is more likely when rapid division rates are permitted, but under less favourable bloom conditions other species may emerge as dominants in the climax community.

Despite being designed for atmospheric (not oceanographic) research, CALIOP measurements have here revealed previously unobserved early stages of polar plankton blooms, provided new insights into mechanisms driving annual boom–bust cycles and interannual variability in plankton stocks, and established a comparative baseline for future regionally focused analyses. The full potential of satellite lidar technology for polar research is, however, yet to be realized. With today's technology, airborne lidar systems retrieve vertically resolved (3–5 m resolution) plankton properties to nearly 3 optical depths<sup>28</sup>, measure plankton absorption and

scattering separately<sup>29</sup>, and detect physiological properties of the phytoplankton<sup>30</sup>. By extending such capabilities to space and combining these measurements with passive satellite ocean colour data and *in situ* depth-profiling autonomous measurements, an observational infrastructure is foreseeable that enables three-dimensional reconstructions of polar-to-global ocean ecosystems and provides new insights into ocean ecology, biogeochemistry, and physical-biological interactions relevant to climate change research.

#### Methods

Methods, including statements of data availability and any associated accession codes and references, are available in the online version of this paper.

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#### Author contributions

M.J.B. designed the study; M.J.B., Y.H. and R.T.O'M. processed satellite data and analysed results; M.J.B. and R.T.O'M. prepared display items; M.J.B. wrote the manuscript with contributions from all authors.

#### Additional information

Supplementary information is available in the online version of the paper. Reprints and permissions information is available online at www.nature.com/reprints. Correspondence and requests for materials should be addressed to M.J.B.

#### **Competing financial interests**

The authors declare no competing financial interests.

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#### Methods

**Global data sets.** Particulate backscatter coefficients  $(b_{bp})$  and photosynthetically active radiation (PAR) data from the MODIS Aqua ocean colour sensor for the period June 2006 to July 2015 were acquired from http://oceancolor.gsfc.nasa.gov/cms. MODIS Aqua bbp values were based on the Generalized Inherent Optical Properties inversion algorithm (GIOP)<sup>31</sup>. CALIOP data for the June 2006 to July 2015 period were analysed as described in Behrenfeld et al.5 except for the following modifications. First, the CALIPSO satellite has been tilted 30° off-nadir for one orbit every two weeks since September 2015 to collect ocean measurements with minimal surface backscatter. These observations have allowed for an improved description of depolarization ratios among ocean subsurface backscatter, diffuse attenuation coefficients, and  $b_{bp}$ compared with the earlier study5. Second, collocated MODIS and CALIOP data were used to develop a relationship between diffuse attenuation coefficients at 532 nm ( $K_{D532}$ ) and  $b_{bp}$  to allow retrieval of  $b_{bp}$  at high latitudes from CALIOP measurements alone, whereas in Behrenfeld et al.5 collocated MODIS KD532 data were required to retrieve  $b_{bp}$ .

CALIOP  $b_{bp}$  and  $K_{D532}$  measurements are collected at nadir along the satellite orbit track, with an ocean surface lidar single-pulse footprint of ~100 m and an along-track pulse-to-pulse distance of 330 m. Ocean properties are retrieved along CALIOP orbit tracks only for cloud layers <1 optical depth. Over a 16-day repeat cycle, CALIOP measurement tracks (day and night) are separated by 0 to 1.5° Given these differences in along-track and between-track coverage, CALIOP data were aggregated into monthly 1° latitude bins for comparison with MODIS spatial coverage (Supplementary Fig. 2). For all other analyses, CALIOP data were aggregated into broader latitude zones, such as the 'comparison zones' (45° to  $55^\circ$  latitude, Fig. 1c,g) and the polar zones of the Northern (60°–81.5° latitude) and Southern (60°–75° latitude) hemispheres. CALIOP  $b_{bp}$  values were multiplied by 1.3 to scale with MODIS  $b_{bp}$  values, thus illustrating the close correspondence between annual cycles retrieved by these two sensors. In the analysis of Behrenfeld et al.<sup>5</sup>, an excellent agreement was found between in situ measured  $b_{bp}$ and unscaled CALIOP  $b_{\rm bp}$  values, whereas  $b_{\rm bp}$  values retrieved with the Garver-Siegel-Maritorena inversion algorithm<sup>32</sup> (GIOP products were not available at the time of the earlier study) were a factor of 1.3 higher than both the in situ and CALIOP data. The reason for this discrepancy is not yet resolved, but for the current study derived biomass accumulation rates (r) are little impacted by whether the CALIOP data are scaled to MODIS data or vice versa. Future analyses identifying the basis for this bias between sensor retrievals will be important for improving quantitative assessments of polar phytoplankton carbon standing stocks. Finally, CALIOP, MODIS, and all other satellite ocean colour sensors fail to detect phytoplankton populations under ice, so the contribution of these populations is not represented in any satellite assessment of polar phytoplankton stocks.

CALIOP and MODIS  $b_{pp}$  values were converted to  $C_{phyto}$  (mg C m<sup>-3</sup>) following (Supplementary Discussion) (refs 10,12):

$$C_{\rm phyto} = 12,128 \,{\rm mg}\,{\rm C}\,{\rm m}^{-2}(b_{\rm bp} - 0.00035 \,{\rm m}^{-1})$$
 (4)

CALIOP K<sub>D532</sub> values were converted to diffuse attenuation coefficients for PAR  $(K_{\text{PAR}}, \text{m}^{-1})$  (ref. 33) and photic-layer depth  $(Z_{eu}, \text{m})$  was taken as the depth of the 0.415 mol quanta m<sup>-2</sup> d<sup>-1</sup> isolume<sup>34,35</sup> and calculated as  $Z_{eu} = \ln(0.415/\text{PAR})/K_{\text{PAR}}$ , except during polar night when  $Z_{eu}$  was assigned a value of 1 m. For calculations of integrated photic-zone biomass ( $\Sigma C_{polar}$ ), we assume that biomass is distributed uniformly with depth and equal to the value observed at the surface. Field studies have shown that subsurface chlorophyll maxima are common in polar regions1 These features are not detected by satellite sensors and accordingly will cause our estimates of  $\Sigma C_{\rm polar}$  to be conservative. Day length values were calculated as a function of date and latitude following Kirk<sup>36</sup>. Mixed-layer depths (MLDs) were calculated from salinity, temperature, and pressure data converted to density (sigma-theta) and based on the Fleet Numerical Meteorology and Oceanography Center (FNMOC) high-resolution ocean analysis model<sup>37</sup> for the period 2006-2007 and the HYbrid Coordinate Ocean Model (HYCOM) (https://hycom.org) for the period 2008-2015. The FNMOC and HYCOM models are both tuned to in situ data (that is, they are 'data-assimilating models'). The FNMOC model does not provide MLD data at latitudes >70° so monthly climatological average MLD data from the HYCOM model were applied at >70° latitude for the years 2006 and 2007. Results presented for the current manuscript are based on mixed-layer depths defined as a density change of  $0.030\,kg\,m^{-3}$  from the value at 10 m (ref. 38). Alternative definitions of MLD were evaluated and resulted in small decreases, but similar annual cycles, in modelled values of  $\mu$  as the MLD criterion was increased from 0.005 to 0.125 kg m<sup>-3</sup>, with small impacts on values of  $d\mu/dt$  and its relationship with biomass accumulation rates, r. Ice-free ocean area for latitudes >60° was based on monthly Sea Ice Index extent data obtained from the National Snow and Ice Data Center (NSIDC) in Boulder, Colorado.

Assessment of phytoplankton division and accumulation rates. Daily net primary production ( $\Sigma$ NPP, mg C m<sup>-2</sup> d<sup>-1</sup>) was calculated for the mixed layer as a

function of time and depth following:

$$\Sigma NPP = \int_{t=\text{sunrise}}^{\text{sunset}} \int_{z=0}^{\text{MLD}} CP_{\text{max}}^{C} \tanh\left(\frac{\alpha^{C} PAR_{t,z}}{P_{\text{max}}^{C}}\right) dz \, dt \tag{5}$$

where C is CALIOP-retrieved phytoplankton concentration (mg C  $m^{-3}$ ) and is assumed to be uniform through the mixed layer, MLD is mixed-layer depth (m), and PAR<sub>t,z</sub> is photosynthetically active radiation (mol photon  $m^{-2} h^{-1}$ ) at time t and depth z. Equation (5) was operated at 1 m depth increments and 64 time increments over the photoperiod. Temporal changes in incident PAR at the sea surface were modelled as a cosine function of time such that the integral of PAR, for the photoperiod equalled the MODIS cloud-corrected PAR product. PAR<sub>t.2</sub> was calculated as  $PAR_{t,z} = PAR_t e^{-K_{PAR,z}}$ . The carbon-specific light-saturated rate of photosynthesis,  $P_{\max}^C$  (mg C (mg C h)<sup>-1</sup>), was calculated as the product of the chlorophyll-specific light-saturated rate of photosynthesis,  $P_{max}^{b}$  (mg C  $(mg Chl h)^{-1}$ ), and the phytoplankton Chl/C ratio  $(mg mg^{-1})$ .  $P_{max}^{b}$  was calculated as a function of sea surface temperature (SST) following an exponential relationship<sup>39</sup> with a  $Q_{10}$  of 2 (ref. 40). Chl/C values were calculated following the model of Behrenfeld and colleagues<sup>41</sup>. The light-saturation parameter,  $E_k$  (mol photon m<sup>-2</sup> h<sup>-1</sup>), was then calculated from Chl/C (ref. 41) and the carbon-specific light-limited slope of the photosynthesis-irradiance relationship,  $\alpha^C \pmod{\operatorname{cmg} C \operatorname{m}^2 (\operatorname{mg} C \operatorname{mol} \operatorname{photon})^{-1}}$ , calculated as  $P_{\operatorname{max}}^C / E_k$ . Phytoplankton division rates ( $\mu$ ) were calculated as  $\Sigma NPP / \Sigma C_{ML}$ , where  $\Sigma C_{ML}$  is the mixed-layer integrated phytoplankton biomass,  $C \times MLD$ . Phytoplankton concentration, C, cancels from the numerator and denominator when equation (5) is divided by  $\Sigma C_{\text{ML}}$ , thus leaving the simplified expression for  $\mu$  given as equation (1). Thus, variability in  $\mu$  and its temporal derivative,  $d\mu/dt$ , is determined by time-dependent changes in MLD, PAR, KPAR, and SST (owing to the dependence of  $P_{\text{max}}^{C}$  on SST). 2006 through 2015 time series of MLD, PAR, SST and  $Z_{eu}$  are provided in Supplementary Fig. 6.

Figure 2b,c compares for the north and south polar zones observed rates of change in phytoplankton biomass (*r*) and values of  $d\mu/dt$  calculated as:

$$d\mu/dt = \frac{\mu_{t_1} - \mu_{t_0}}{t_1 - t_0} \tag{6}$$

where  $t_1 - t_0 = 1$  month (see Supplementary Discussion for analysis of uncertainty). We show that *r* and  $d\mu/dt$  have correlated annual cycles of comparable magnitude, implying that phytoplankton loss rates are time-lagged behind division rates by  $\sim 1$  day when averaged over a monthly period. This result does not mean that a 1 day lag will necessarily be observed between division and loss rates when measured on a daily timescale.

Figure 2a,b compares observed and modelled time series of phytoplankton biomass. Modelled phytoplankton biomass in this comparison (red line) was calculated by assigning an initial biomass of 45 and 20 mg C m<sup>-3</sup> for the polar north and south, respectively, and then applying the model predictions of  $d\mu/dt$  (that is, the derivative of equation (1)) over the 10-year time series.

While an excellent correspondence is observed between the CALIOP time series of *r* and  $d\mu/dt$  for both polar zones, there are also secondary discrepancies. For example, observed values of *r* were notably higher than calculated values of  $d\mu/dt$  during May and June of 2013 for the north polar zone. Such discrepancies can arise for multiple reasons, including errors in the observational data, comparison of monthly satellite biomass data with differences in spatial coverage, uncertainties in modelled values of  $d\mu/dt$  associated with errors in input data fields (for example, MLD, PAR,  $K_{PAR}$ , and SST), or unaccounted for temporal variations in phytoplankton physiological properties and loss processes. Differences between observed *r* and modelled  $d\mu/dt$  are responsible for mismatches between measured and modelled biomass values (Fig. 2a,c).

**Phytoplankton climatological annual cycles.** Figure 3 shows climatological average monthly cycles of (top panels) phytoplankton biomass accumulation rates (r), modelled  $d\mu/dt$ , and (bottom panels) phytoplankton division rates ( $\mu$ ). For these data, linear regression statistics for the r versus  $d\mu/dt$  relationships are  $r^2 = 0.72$ , n = 12, p < 0.001 for the north polar zone and  $r^2 = 0.85$ , n = 12, p < 0.001 for the south polar zone. In the figure, the top and bottom panels are connected for four key events. Here, the initial rise in biomass (purple dashed arrows) was taken as the point where the biomass rate of change first becomes positive and thereafter remains positive to the bloom climax. For each arrow, the ranges in r (coloured brackets in top panels) and  $\mu$  (boxes in the bottom panels) are taken from the year-by-year occurrences of these events shown in Fig. 2.

Data availability. Remote sensing data on ocean plankton properties and surface mixed-layer depth used in this study are available at http://oceancolor.gsfc.nasa.gov/cms and http://www.science.oregonstate.edu/ocean.productivity and from the corresponding author on request. Ice-cover data are available at https://nsidc.org.



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Supplementary Information for Behrenfeld et al. manuscript, "Decade of polar phytoplankton biomass cycles characterized using a space-based lidar"

#### SUPPLEMENTARY DISCUSSION

(1) **Phytoplankton Carbon Retrievals:** For the current study, phytoplankton biomass  $(C_{phyto})$ is estimated from satellite retrievals of particulate backscatter coefficients  $(b_{bp})$  following the relationship given in equation 2 of the Methods section. The scalar in this relationship (i.e., 12,128 mg C m<sup>-2</sup>) was determined using co-located field measurements of  $b_{bp}$  and analytically measured phytoplankton carbon<sup>1</sup>. These measurements were conducted over a wide range of ecosystem conditions (e.g., equatorial upwelling, open ocean gyres, temperate bloom forming regions), but none of the measurements included polar samples. We have no *apriori* reason to suspect that a different mean relationship exists for polar regions, but if it does then its primary impact is expected to be an adjustment in derived absolute phytoplankton biomass values, with little (if any) impact on derived biomass rates of change. Consequently, the main conclusions of our study are expected to be robust to this uncertainty. Other factors can also contribute to variability in the  $b_{bp}$  to  $C_{phyto}$  relationship, such as suspended particulate inorganic carbon, glacial flour, and suspended sediments. However, for the current analysis where data are integrated over vast polar regions, these more local phenomena should have only secondary impacts on derived annual plankton properties and are not expected to compromise our conclusions. Finally, it should be noted that, while the method for analytically measuring phytoplankton carbon is relatively new<sup>2</sup>, a diversity of studies have been conducted supporting the use of  $b_{bp}$  data as a metric of phytoplankton biomass<sup>3-9</sup>. Together, these studies significantly expand the range of conditions under which relationships between phytoplankton biomass and light scattering properties have been evaluated, albeit again without polar samples. Nevertheless, future studies

providing new data from polar regions will contribute to improved assessments of phytoplankton biomass

#### (2) Analysis of uncertainty in derived $d\mu/dt$ associated with parameters describing

**phytoplankton physiology:** A central finding of the current study is that the rate of change in phytoplankton biomass (*r*) is linked to the rate of change in phytoplankton division ( $d\mu/dt$ ). Temporal variations in modeled  $d\mu/dt$  values reflect variations in the time-dependent variables, *MLD*, *PAR*,  $P^{C}_{max}$ , and  $\alpha^{C}$  (main manuscript Eq. 1; Methods Eq. 4), with the two latter terms characterizing variations in phytoplankton physiology. To evaluate the sensitivity of derived  $d\mu/dt$  estimates on the two physiological terms, we operated our *NPP* model with  $P^{C}_{max}$  decreased (Alternative Model #1) and increased (Alternative Model #2) by a factor of 1.5 from





**Figure Caption:** Comparison of (top) North Polar Zone and (bottom) South Polar Zone time-series in Standard Model  $d\mu/dt$  and values calculated using alternative values for  $P^{c}_{max}$  and  $\alpha^{c}$  (see description in text).

the values calculated as described in the Methods section (i.e., the 'Standard Model') and with  $\alpha^{C}$  values based on an earlier description of *Chl:C* variability<sup>8</sup> (Alternative Model #3 in the figure on the left). Regarding this latter adjustment, our 'standard model' assigns a value to  $\alpha^{C}$  based on the recent *Chl:C* model of Behrenfeld et al.<sup>10</sup>, which represents an 'upper end' model of the photoacclimation response. By contrast, the model of Westberry et al.<sup>8</sup> assigns a very conservative fraction of *Chl:C* variability to photoacclimation and thus represents a 'lower end' response model. Results from these reparameterizations of our *NPP* model (figure on previous page) indicate only minor changes in the amplitude of calculated annual cycles in  $d\mu/dt$  associated with these rather significant changes in  $P^{C}_{max}$ , and  $\alpha^{C}$ . Furthermore, the regression analyses of  $d\mu/dt$  from the Standard Model and values from any of the Alternative Models give coefficients of determination of  $r^2 = 0.99$  for both the North and South Polar Zones, indicating that the model variations slightly impact the amplitude of  $d\mu/dt$  and *r*.

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Supplementary Table 1. Linear regression analysis of relationships between seasonal mean ice-free ocean area (*IFA*; km<sup>2</sup> × 10<sup>6</sup>), polar zone average phytoplankton concentration (*C*; mg C m<sup>-3</sup>), and polar zone integrated photic layer phytoplankton biomass ( $\mathcal{IC}_{polar}$ ; Tg C). (Column 1) Regression analyses were conducted separately for all data from the CALIOP record (Fig. 4) and for winter and summer months only. (Column 2) For each time period, relationships were evaluated separately for northern (60° to 81.5° N) and southern (60° to 75° S) polar zones. (Columns 3 – 6) Regression result for (Row 1) *IFA* versus  $\mathcal{IC}_{polar}$ , (Row 2)  $C_{phyto}$  versus  $\mathcal{IC}_{polar}$ , and (Row 3, shaded) Cross-correlation between *IFA* and  $C_{phyto}$ .

Time Period	Hemisphere	Regression	Slope	$r^2$	p value
All data	Northern	IFA vs $\mathcal{I}C_{polar}$	1.16	0.20	0.005
		$C_{phyto}$ vs $\mathcal{D}C_{polar}$	0.18	0.80	< 0.001
		<i>IFA</i> vs $C_{phyto}$ (cross correlation)	0.01	0.03	0.33
	Southern	IFA vs $\mathcal{Z}C_{polar}$	0.60	0.72	< 0.001
		$C_{phyto}$ vs $\mathcal{D}C_{polar}$	0.10	0.11	0.05
		<i>IFA</i> vs $C_{phyto}$ (cross correlation)	0.00	0.00	0.97
Summer	Northern	IFA vs $\mathcal{Z}C_{polar}$	2.33	0.32	0.09
	(June-Aug)	$C_{phyto}$ vs $\mathcal{D}C_{polar}$	0.18	0.84	< 0.001
		<i>IFA</i> vs $C_{phyto}$ (cross correlation)	0.01	0.08	0.44
	Southern	IFA vs $\mathcal{Z}C_{polar}$	0.71	0.75	0.003
	(Dec-Feb)	$C_{phyto}$ vs $\mathcal{D}C_{polar}$	0.15	0.12	0.37
		<i>IFA</i> vs $C_{phyto}$ (cross correlation)	-0.09	0.03	0.67
				-	
Winter	Northern	IFA vs $\mathcal{Z}C_{polar}$	-0.30	0.06	0.52
	(Dec-Feb)	$C_{phyto}$ vs $\mathcal{D}C_{polar}$	0.08	0.75	0.003
		<i>IFA</i> vs $C_{phyto}$ (cross correlation)	-0.04	0.20	0.22
	Southern	IFA vs $\Sigma C_{polar}$	0.35	0.69	0.003
	(June-Aug)	$C_{phyto}$ vs $\overline{Z}C_{polar}$	0.08	0.78	< 0.001
		<i>IFA</i> vs $C_{phyto}$ (cross correlation)	0.12	0.36	0.07



Supplementary Figure 1. CALIOP ground tracks achieved within a single 16-day repeat cycle. (a) Northern hemisphere  $\ge 45^{\circ}$  latitude. (b) Southern hemisphere  $\ge 45^{\circ}$  latitude. Red circles delineate 10° latitudinal integration bins used in supplemental figure 2 and 4.



Supplementary Figure 2: Comparison of CALIOP and MODIS pixel coverage per month per 10° latitudinal bin. White symbols = Total number of 1° latitude × 1° longitude ice-free ocean pixels per month with valid MODIS  $b_{bp}$  data. Black symbols = Total number of 1° latitude × 1° longitude ice-free ocean pixels per month with valid CALIOP  $b_{bp}$  data (left panels) Northern hemisphere latitudinal zones as shown in Supplemental figure 1. (right panels) Southern latitudinal zones as shown in Supplemental figure 1.

## **North Polar Zone**



South Polar Zone



Supplementary Figure 3: Comparison of CALIOP and MODIS phytoplankton biomass records for the Polar Zones. Monthly mean phytoplankton biomass ( $C_{phyto}$ ; units = mg C m<sup>-3</sup>) for (top panel) North Polar Zone and (bottom panel) South Polar Zone (i.e.,  $\ge 60^{\circ}$ ). White symbols = MODIS. Black symbols = CALIOP. Vertical yellow bars = months with no MODIS data for at least one of the 10° latitudinal zones shown in Supplemental figure 1. For multiple annual cycles, particularly in the South Polar Zone, MODIS biomass values are notably lower than CALIOP values for the first month of MODIS data. This bias is in-part due to MODIS data coverage being very poor during the first month (often <10% area coverage; Supplemental Fig. 2) and biased toward lower latitudes of the Polar Zone due to solar illumination conditions. Active measurements by CALIOP, in contrast, are more evenly distributed across the Polar Zone because they are not dependent on solar angle and can even be collected during polar night. In the above time series for the North Polar Zone, there are also occasions where MODIS biomass values are significantly higher than CALIOP values during the summer maximum (e.g., 2007 and 2013). As indicated in Supplemental figure 4, these discrepancies are largely attributable to the northernmost latitudes (75° - 81.5°) where both MODIS and CALIOP coverage is weakest and co-located retrievals are least likely.



Supplementary Figure 4: Comparison of CALIOP and MODIS phytoplankton biomass records for 10° latitudinal zones. Monthly mean phytoplankton biomass ( $C_{phyto}$ ; units = mg C m<sup>-3</sup>) for (left panels) northern hemisphere and (right panel) southern hemisphere latitudinal zones polarward of 45° as shown in Supplemental figure 1. White symbols = MODIS. Black symbols = CALIOP. Latitudinal zones are indicated in blue text for each panel. Correspondence between MODIS and CALIOP biomass values is indicated by the coefficient of determination (r<sup>2</sup>) shown in red text.



Supplementary Figure 5: 1978 to 2016 monthly anomalies in polar zone ice free ocean area. Left panel = North polar zone ( $60^{\circ}N - 81.5^{\circ}N$ ). Right panel = South polar zone ( $60^{\circ}S - 75^{\circ}S$ ). Gray shaded area = period of CALIOP measurements (2006 - 2015). Note that large spikes in the anomaly records result from temporal shifts in the timing of annual ice melt and ice formation.

# a. North Polar Zone



# b. South Polar Zone



Supplementary Figure 6: 2006 to 2016 time series of division rate of change ( $d\mu/dt$ ), incident photosynthetically active radiation (PAR; mole quanta m<sup>-2</sup> d<sup>-1</sup>), sea surface temperature (SST; °C), mixed layer depth (MLD; m), and photic layer depth (Zeu; m). (a) North Polar Zone. (b) South Polar Zone. Note that it is the strong annual cycles in MLD and PAR that primarily drive the annual cycle in  $d\mu/dt$ , whereas the highly constrained annual cycles in SST have a minor impact on modeled  $d\mu/dt$ .