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## Global marine plankton functional type biomass distributions: Phaeocystis sp.

M. Vogt<sup>1</sup>, C. O'Brien<sup>1</sup>, J. Peloquin<sup>1</sup>, V. Schoemann<sup>2</sup>, E. Breton<sup>3</sup>, M. Estrada<sup>4</sup>, J. Gibson<sup>5</sup>, D. Karentz<sup>6</sup>, M. A. Van Leeuwe<sup>7</sup>, J. Stefels<sup>7</sup>, C. Widdicombe<sup>8</sup>, and L. Peperzak<sup>2</sup>

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<sup>&</sup>lt;sup>1</sup>Institute for Biogeochemistry and Pollutant Dynamics, Universitätsstrasse 16, 8092 Zürich, Switzerland

<sup>&</sup>lt;sup>2</sup>Royal Netherlands Institute for Sea Research, P.O. Box 59, 1790 AB Den Burg (Texel), The Netherlands

<sup>&</sup>lt;sup>3</sup>Université Lille Nord de France, ULCO, CNRS, LOG UMR8187, 32 Avenue Foch, 62930 Wimereux. France

<sup>&</sup>lt;sup>4</sup>Institut de Ciències del MAR (CSIC), Passeig Maritim de la Barceloneta, 3749, 08003 Barcelona, Catalunya, Spain

<sup>&</sup>lt;sup>5</sup>Tasmanian Aquaculture and Fisheries Institute, University of Tasmania, Private Bag 50, Hobart Tasmania 7001, Australia

<sup>&</sup>lt;sup>6</sup>University of San Francisco, College of Arts and Sciences, 2130 Fulton Street, San Francisco, CA 94117, USA

<sup>&</sup>lt;sup>7</sup>University of Groningen, Centre for Ecological and Evolutionary Studies, Department of Plant Ecophysiology, P.O. Box 14, 9750AA Haren, The Netherlands <sup>8</sup>Plymouth Marine Laboratory, Prospect Place, The Hoe, Plymouth PL1 3DH, UK

Received: 24 April 2012 - Accepted: 25 April 2012 - Published: 16 May 2012

Correspondence to: M. Vogt (meike.vogt@env.ethz.ch)

Published by Copernicus Publications.

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The planktonic haptophyte Phaeocystis has been suggested to play a fundamental role in the global biogeochemical cycling of carbon and sulphur, but little is known about its global biomass distribution. We have collected global microscopy data of the genus Phaeocystis and converted abundance data to carbon biomass using speciesspecific carbon conversion factors. Microscopic counts of single-celled and colonial Phaeocystis were obtained both through the mining of online databases and by accepting direct submissions (both published and unpublished) from *Phaeocystis* specialists. We recorded abundance data from a total of 1595 depth-resolved stations sampled between 1955-2009. The quality-controlled dataset includes 5057 counts of individual *Phaeocystis* cells resolved to species level and information regarding life-stages from 3526 samples. 83% of stations were located in the Northern Hemisphere while 17% were located in the Southern Hemisphere. Most data were located in the latitude range of 50-70° N. While the seasonal distribution of Northern Hemisphere data was well-balanced, Southern Hemisphere data was biased towards summer months. Mean species- and form-specific cell diameters were determined from previously published studies. Cell diameters were used to calculate the cellular biovolume of *Phaeocystis* cells, assuming spherical geometry. Cell biomass was calculated using a carbon conversion factor for Prymnesiophytes (Menden-Deuer and Lessard, 2000). For colonies, the number of cells per colony was derived from the colony volume. Cell numbers were then converted to carbon concentrations. An estimation of colonial mucus carbon was included a posteriori, assuming a mean colony size for each species. Carbon content per cell ranged from 9 pg (single-celled *Phaeocystis antarc*tica) to 29 pg (colonial Phaeocystis globosa). Non-zero Phaeocystis cell biomasses (without mucus carbon) range from  $2.9 \times 10^{-5} \,\mu g \, l^{-1}$  to  $5.4 \times 10^3 \,\mu g \, l^{-1}$ , with a mean of  $45.7\,\mu\text{g}\,\text{l}^{-1}$  and a median of  $3.0\,\mu\text{g}\,\text{l}^{-1}$ . Highest biomasses occur in the Southern Ocean below 70° S (up to 783.9 µg l<sup>-1</sup>), and in the North Atlantic around 50° N (up to

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http://doi.pangaea.de/10.1594/PANGAEA.779101.

#### Introduction

Plankton functional types (PFTs; Le Quéré et al., 2005) and marine ecosystem composition are important for the biogeochemical cycling of many abundant elements on Earth, such as carbon, nitrogen, and sulphur (e.g. Weber and Deutsch, 2010). In recent decades, changes have been observed in marine plankton communities (Chavez et al., 2003; Reid et al., 2007; Hatun et al., 2009; Beaugrand and Reid, 2003), and these changes are likely to affect local and global biodiversity, fisheries and biogeochemical cycling. Marine ecosystem models based on PFTs (Dynamic Green Ocean models: DGOMs) have been developed in order to study the lower trophic levels of marine ecosystems and the potential impact of changes in their structure and distribution (Le Quéré et al., 2005). DGOMs have been applied to a wide range of biological and biogeochemical questions (Aumont and Bopp, 2006; Hashioka and Yamanaka, 2007; Moore and Doney, 2007; Vogt et al., 2010; Weber and Deutsch, 2010). However, the validation of these models has proven difficult due to the scarcity of observational abundance and biomass data for individual PFTs.

The MARine Ecosystem DATa (MAREDAT) initiative is a community effort to provide marine ecosystem modellers with global biomass distributions for the major PFTs currently represented in marine ecosystem models (silicifiers, calcifiers, nitrogen fixers, DMS-producers, picophytoplankton, bacteria, microzooplankton, mesozooplankton and macrozooplankton). MAREDAT is part of the MARine Ecosystem Model Intercomparison Project (MAREMIP). All MAREDAT biomass fields are publicly available for use in model evaluation and development, and for other applications in biological oceanography.

The haptophyte Phaeocystis has been suggested to play a fundamental role in the global biogeochemical cycling of carbon and sulphur (Le Quéré et al., 2005).

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Phaeocystis is a globally distributed genus of marine phytoplankton with a polymorphic life cycle, alternating between flagellated, free-living cells of 3–9 μm in diameter and colonial stages which form colonies reaching several mm–cm (Rousseau et al., 1990; Peperzak et al., 2000; Peperzak and Gäbler-Schwarz, 2012; Chen et al., 2002; Schoemann et al., 2005). Three of the six recognised *Phaeocystis* species are known to form massive blooms of gelatinous colonies, which may contribute significantly to carbon export (DiTullio et al., 2000). In addition, *Phaeocystis* cells are important producers of dimethylsulphoniopropionate (DMSP), which is the marine precursor of the trace gas dimethylsulphide (DMS). DMS has been suggested to play an important role in cloud formation, and DMS production is the main recycling pathway of sulphur from the ocean to the land. In their review, Schoemann et al. (2005) conclude that it should be possible to derive a single unique parameterisation of *Phaeocystis* growth for global modelling. Hence, *Phaeocystis* has recently been included in a number of regional and global DGOMs (e.g. Wang and Moore, 2011).

Here, we present biomass data that were estimated from direct cell counts of colonial and single-celled *Phaeocystis*. We show the spatial and temporal distribution of *Phaeocystis* biomass, with a particular emphasis on the seasonal and vertical patterns. We discuss in detail our method for converting abundance to carbon biomass and note the uncertainties in the carbon conversions. Our biomass estimates are tailored to suit the needs of the modelling community for marine ecosystem model validation and model development, but they are also intended to aid biological oceanographers in the exploration of the relative abundances of different PFTs in the modern ocean and their respective biogeochemical roles, for the study of ecological niches in marine ecosystems and the assessment of marine biodiversity.

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#### 2.1 Origin of data

Our data consists of abundance measurements from several databases (BODC, OBIS, OCB DMO, Pangaea, WOD09, US JGOFS<sup>1</sup>), and published and unpublished data from several contributing authors (E. Breton, M. Estrada, J. Gibson, D. Karentz, M. A. Van Leeuwe, J. Peloquin, L. Peperzak, V. Schoemann, J. Stefels, C. Widdicombe). Often, the online databases did not denote the method used for the quantitative analysis of *Phaeocystis* abundances. However, most known counts have been made using the common inverted microscopy and epifluorescence methods (Karlson et al., 2010). Both methods require the sampling of *Phaeocystis* colonies in Niskin bottles and the subsequent preservation of cells in Lugol's solution or another preservative. After storage of the sample prior to analysis, many scientists concentrate the sample through settling in counting chambers or filtration onto a polycarbonate filter.

Most conventional preservation agents cause the disintegration of the colonial matrix, such that colonial and single cells can no longer be distinguished. One preservation method based on a mixture of Lugol's, glutaraldehyde and iodine (Guiselin et al., 2009; Sherr and Sherr, 1993; Rousseau et al., 1990) is able to maintain colony structure (e.g. Karentz and Spero, 1995; Riebesell et al., 1995; Brown et al., 2008; Wassmann et al., 2005), but this is not widely used. Due to these difficulties, only few measurements resolve *Phaeocystis* life stages or morphotypes.

Table 1 summarizes the origin of all our data, sorted by database, principal investigator and the project during which measurements were taken. At present, the database

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<sup>&</sup>lt;sup>1</sup>BODC: British Oceanographic Data Centre; OBIS: The Ocean Biogeographic Information System, OCB DMO: Ocean Carbon and Biogeochemistry Coordination and Data Management Office, Pangaea: Data Publisher for Earth and Environmental Science, WOD: World Ocean Database, US JGOFS: US Joint Global Ocean Flux Study

contains 5057 individual data points from 3526 samples of 1595 depth-resolved stations.

### 2.2 Quality control

Given the low numbers of data points and the fact that *Phaeocystis* is a blooming species with a wide range of biomass concentration, the identification and rejection of outliers in our dataset is challenging. We use Chauvenet's criterion to identify statistical outliers in the log-normalized biomass data (Glover et al., 2011; Buitenhuis et al., 2012). Based on the analysis, none of the stations was identified to yield biomasses with a probability of deviation from the mean greater than 1/2n, with n = 2547 being number of non-zero data summed up for each station (two-sided z-score: |zc| = 3.72). In addition to the statistical testing of the biomass distribution, we also quality controlled the range of our cell abundances. We found that our maximum reported abundance of  $19 \times 10^7$  cells  $\Gamma^{-1}$  is within the range of previously reported abundances: Schoemann et al. (2005) report maximum cell abundances of the order of ca. 10<sup>7</sup> cells l<sup>-1</sup> in areas of colony occurrence (http://www.nioz.nl/projects/ironages). The largest bloom of P. antarctica was observed in Prydz Bay (http://www.nioz.nl/projects/ironages), with cell abundances measured up to  $6 \times 10^7$  cells I<sup>-1</sup>. Eilertsen et al. (1989) reported a maximum of  $1.2 \times 10^7$  cells I<sup>-1</sup> of *P. pouchetii* in the Konsfjord. For *P. globosa*, a maximal abundance of  $20 \times 10^7$  cells  $I^{-1}$  has been observed, corresponding to a total biomass of ca.  $10 \,\mathrm{mg} \,\mathrm{CI}^{-1}$  including mucus (Cadée and Hegeman, 1986; Schoemann et al., 2005). The latter biomass value is 20 times larger than the maximal biomass we report  $(5.4 \times 10^3 \, \mu g \, l^{-1})$ . Thus, based on statistical and observational evidence, none of the data were flagged.

#### 2.3 Biomass conversion

We distinguish between single, colonial and unspecified *Phaeocystis* cells. While *Phaeocystis* is generally observed and counted under bloom conditions, a significant

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fraction of cells is non-colonial even during bloom conditions (V. Schoemann, auxillary data). Hence, in order to calculate the lower limit biomass, we have assumed unspecified cells to be single cells. To first order, this choice does not affect the order of magnitude of our cell biomass estimates, since cell carbon is of the same order of magnitude for both colonial and single cells (see below). We define total *Phaeocystis* biomass to consist of cell biomass and biomass contained in the mucus surrounding *Phaeocystis* colonies. For our calculation of total biomass, we chose unidentified cells to be in the colonial stage. Hence, our cell biomass estimates represent a lower limit, and our total biomass estimates including colonial mucus represent an upper limit for global *Phaeocystis* biomass.

Biomass was determined from cell abundance using species- and form-specific conversion factors (Fig. 1). Similar conversion schemes have been previously described (e.g. Schoemann et al. 2005, and references therein). Total cell abundances were divided into single cells, colonial cells and undefined cell types. For each species, the mid-point of the range of reported cell diameters from the literature was used for single and colonial cells (Table 2; *P. globosa*: Rousseau et al., 2007; Schoemann et al., 2005; *P. antarctica*: Mathot et al., 2000; Rousseau et al., 2007).

Where the species was not specified, Southern Ocean cell counts were assumed to be *Phaeocystis antarctica*. For cell counts in other regions, the mid-point of the range of cell diameters for *P. pouchetii* and *P. globosa* was taken (Table 2; flagellates:  $5.0\,\mu m$ , colonial cells:  $6.7\,\mu m$ ). From cell diameter we computed biovolume, assuming spherical geometry of all cell types. We then converted biovolume to carbon biomass using an empirical volume-carbon conversion formula for prymnesiophytes developed by Menden-Deuer and Lessard (2000) (Table 2).

Most colonial cells were reported in the form of cell abundances. However, one dataset (P. globosa; number of data points: n = 30) provided colony counts only, but additionally reported the corresponding colony diameters. We used the reported colony diameter to calculate colony volume (assuming spherical colonies), and from this

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estimated the number of cells per colony using published conversion factors (Table 2; *P. globosa*: Rousseau et al., 1990; *P. antarctica*: Mathot et al., 2000; no colony-only cell counts reported for *P. pouchetii*). Total cell counts per colony were then converted to carbon biomass using the method described above.

We show biomass estimates based on cell carbon excluding colonial mucus as our lower limit for *Phaeocystis* biomass. The range of uncertainty for the lower limit biomass estimates is given by the uncertainty in cell diameters. Additional uncertainty is introduced where cell life form is not specified. The uncertainty introduced by this assumption is addressed by calculating a minimum cell biomass estimate treating all undefined cell types as single cells.

Estimates for colonial mucus are included to provide an upper limit for *Phaeocystis* biomass. Estimating mucus carbon from cell counts alone is problematic, as the ratio of mucus carbon to cell number increases with colony size. Colony size therefore needs to be known in order to calculate accurate estimates of mucus carbon. Only one of the datasets (n = 30) included information on colony size. Consequentially, we have used a standard colony diameter of 200 µm for all three species, based on a review of previously reported colony sizes: Verity et al. (2007) find most P. pouchetii colonies in their study to range between 20-450 µm in diameter, Reigstad and Wassmann (2007) observe most of their P. pouchetii colonies in a size range between 65-115 µm, Mathot et al. (2000) observe P. antarctica colonies to range from 9.3–560 µm, and Rousseau et al. (1990) report colony sizes of P. globosa to range from 10 µm-2 mm. In all references, larger colonies occured, but were rarer than the smaller colonies. In our data, P. globosa colonies range from 11-594 µm in diameter, with a mean diameter of 197 µm. Given that the samples of Verity et al. (2007), Mathot et al. (2000) and Rousseau et al. (1990) cover a similar range of sizes for all three species, and that the dataset that reports colony sizes confirms a mean colony size of ca. 200 µm, these findings suggest that the chosen standard diameter is a realistic value for a typical *Phaeocystis* bloom. Maximum sizes are reported in Schoemann et al. (2005) and Baumann et al. (1994), and range between 9 mm-3 cm for P. globosa, between 1.5-2 mm for P. pouchetii, and around

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1.4–9 mm for *P. antarctica*, but given the lack of data on colony sizes, we are unable to quantify the impact of large colonies on average carbon biomass. However, we assess the uncertainty of our estimates by calculating mucus carbon for the minimum and maximum colony sizes reported for each species (Schoemann et al., 2005; Baumann et al., 1994).

Conversion factors have previously been published for estimating mucus biomass and number of cells from colony volume for *P. antarctica* (Mathot et al., 2000) and *P. globosa* (Rousseau et al., 1990). Using these estimates we calculated the expected mucus biomass per cell (Table 2). Unspecified cell types were assumed to be colonial cells when calculating these upper estimates of *Phaeocystis* biomass.

For *P. pouchetii*, no direct mucus carbon conversion factor has been developed, but Verity et al. (2007) provides a conversion factor for colony volume to total colony biomass (Table 2; cells and mucus). Following the same procedure as for the other two species, we used this to calculate total biomass per cell. We then subtracted our cell biomass estimate for colonial cells to obtain an estimate of mucus carbon per cell for comparison with *P. globosa* and *P. antarctica* estimates.

Unspecified species outside of the Southern Ocean were given a total biomass per cell of 224 pg, which corresponds to the mean total biomass estimate for *P. globosa* and *P. pouchetii* (Table 2).

#### 3 Results

#### 3.1 Global distribution of abundance data

Of the 1595 stations contained in the database (Fig. 2), 83% are located in the Northern Hemisphere (NH) and only 17% in the Southern Hemisphere (SH; Fig. 3). Out of the 3526 samples, 2547 were reported as non-zero biomass, with 2054 non-zero abundances out of 2862 samples for the NH, and 493 non-zero abundances out of 664 samples for the SH (Table 3). Most measurements (53%) were taken in the latitudinal

band of 50–70° N (Fig. 3). When only data points with non-zero abundances are taken into account, we find that most non-zero data were collected between 60–80° N (64 %; Table 3), with relatively few non-zero abundances recorded between 50–60° N (11 %). Several latitudinal bands are undersampled. We could not collect data for the 40–20° S, 0–10° N and 30–40° N latitudinal bands. All in all, we have few non-zero data in tropical and sub-tropical latitudes from 40° S to 40° N, where sampling is targeted at other phytoplankton groups.

While 60% of measurements were taken in the upper 10 m of the water column, the mean sampling depth of our dataset is 27 m, and the median sampling depth is 10 m. Reported cell abundances were maximal at depths between 0–80 m. Observations and laboratory experiments suggest that *Phaeocystis* is well-adapted to low light conditions (Arrigo et al., 1999; Moore et al., 2007; Shields and Smith, 2009). In our database, the deepest occurrence of *Phaeocystis* was at 292 m at 65° N/35° W (Barents Sea; OBIS dataset).

### 3.2 Temporal distribution of data

The data were collected from 1955–2009, with 79 % of measurements taken during the period of 1990–2009 (Fig. 4). 6 % (8 %) of (non-zero) measurements were taken in the 1950s, <1 % (<1 %) in the 1960s, <1 % (1 %) in the 1970s, 14 % (10 %) in the 1980s, 55 % (60 %) in the 1990s and 23 % (20 %) between 2000–2009.

Dividing the data into the four seasons for both hemispheres gives a first indication of the level of temporal bias (Table 4). In the Northern Hemisphere, 56% (64%) of all (non-zero) data were taken in spring, 29% (31%) in summer, 9% (5%) in autumn and 6% (<1%) in winter. For the For the Southern Hemisphere, 27% (32%) of data was collected in spring, 58% (52%) in summer, 13% (16%) in autumn and only 2% (<1%) in winter. Hence, NH data is biased towards spring values, and SH data towards summer values.

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*Phaeocystis* biomass estimates based on cell carbon only, without mucus carbon included, constitute a lower boundary for carbon biomass of this PFT in the global ocean. Since mucus carbon biomass is difficult to quantify based on *Phaeocystis* cell counts, many marine ecosystem models do not include a parameterisation of mucus carbon for this PFT. Thus, in the following section, our estimates of cell biomass represent a lower limit of carbon biomass for model validation. *Phaeocystis* biomasses span a wide range of concentrations, which is why we show log transformed biomass concentrations in all subsequent figures. However, we report only non log-transformed biomass concentrations in this manuscript for better comparability with the original data submission.

#### 3.3.1 Global surface cell biomass characteristics

*Phaeocystis* biomass estimated from cell carbon alone is depicted in Fig. 5a for the surface layer of the ocean (0–5 m). The maximal biomass calculated from the reported cell abundances is  $5449.3 \,\mu g \, C \, I^{-1}$ , located at  $53^{\circ} \, N$  at a depth of 0 m during the spring bloom (month of May). The maximal cell biomass in the Southern Hemisphere is  $783.9 \,\mu g \, C \, I^{-1}$ , recorded in the Ross Sea in January ( $76.49^{\circ} \, S$ ,  $171.97^{\circ} \, E$ , depth 1 m). The mean of all reported non-zero cell biomass values is  $45.7 \,\mu g \, C \, I^{-1}$ , and the median is  $3.0 \,\mu g \, C \, I^{-1}$ . Of all calculated cell biomasses,  $40.1 \,\%$  are in the range of 0–0.1  $\,\mu g \, C \, I^{-1}$ ,  $55.6 \,\%$  in the range of 0–1  $\,\mu g \, C \, I^{-1}$ , and  $67.5 \,\%$  between 0 and  $5 \,\mu g \, C \, I^{-1}$ .

Figure 5b shows the range of uncertainty for cell biomass in % resulting from the uncertainty in cell diameters reported for each species and life stage. Biomasses calculated using the higher estimates of cell diameter are 246 to 355 % higher than estimates calculated using mean cell dimensions. Biomasses calculated using the lower cell diameter estimates are between 4 and 26 % of the mean values. Uncertainties are highest when species or life form is not reported. Biomass estimates are highly

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sensitive to changes in cell size, and reduced uncertainty is only possible if cell measurements are available in addition to abundance data.

#### 3.3.2 Latitudinal cell biomass distribution

Calculated cell biomasses do not follow a distinct latitudinal pattern (Fig. 6a). Highest cell biomasses occur at latitudes around 50° N and 80° S, lowest cell biomasses are calculated for latitudes around 20° S (Peruvian upwelling). Cell biomasses decrease from 50° N towards the pole in the Northern Hemisphere, but Southern Hemisphere concentrations increase polewards towards the Antarctic continent. Given that many of our data stem from coastal regions, we note that our latitudinal distributions are biased towards high coastal concentrations in some areas, as open ocean areas are still undersampled. However, cell biomass distributions confirm previous findings that *Phaeocystis* blooms occur in the temperate and high latitudes of both hemispheres, and that *Phaeocystis* is fairly ubiquitous, occurring in all major ocean basins.

### 3.3.3 Depth distribution of cell biomass

Figure 7 shows calculated cell biomass estimates for *Phaeocystis* in six different depth ranges (0–5 m, 5–25 m, 25–50 m, 50–75 m, 75–100 m and depths >100 m). All depth bands have not been sampled at each station, and many datasets contain only surface measurements. Where depth profiles are available, cell biomass concentrations are generally highest in the surface layer and decrease with depth to 100 m (Fig. 6b). Cell biomasses are low between 100–300 m (mean non-zero biomass concentrations of 7.3 μg C l<sup>-1</sup>), however, high *Phaeocystis* abundances are reported even at depths of close to 300 m in the Northern Hemisphere. The highest cell biomass reported below 100 m is 311.9 μg C l<sup>-1</sup> in the Arctic (66.42° N 34.36° E) in late May, at a depth of 270 m. In the Southern Ocean, *Phaeocystis* cells are reported to a maximum depth of 200 m in the Weddell Sea during February and March, but biomass values below 100 m never exceed 0.01 μg C l<sup>-1</sup>. Given the limited number of data points reported for this depth

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range, it is unclear how representative our data are of deep *Phaeocystis* cell biomasses

#### Seasonal distribution of cell biomass 3.3.4

in other sampling locations.

Cell biomass distributions for the Northern and Southern Hemisphere show that the calculated *Phaeocystis* biomasses reflect those of a typical blooming species (Fig. 8a and b). In the NH, Phaeocystis blooms during the spring months, with the spread of the biomass distribution being a combination of the temporal development of a bloom, and different blooms starting times at different latitudes. In the SH, cell biomasses are highest in December and January. The temporal development mostly reflects Southern Ocean dynamics, as few samples were taken at latitudes below 40°S (compare Fig. 6b).

### Total *Phaeocystis* biomass distribution (mucus included)

Biomass estimates including colonial mucus are given as an upper limit for our biomass estimates (Fig. 9a). Given that the ratio of mucus carbon to cell carbon is highly dependent on colony size, the addition of mucus carbon estimates introduces a high level of uncertainty to total biomass estimates where colony size data is unavailable. Calculating mucus carbon biomass based on the minimum and maximum reported colony sizes for each species (Schoemann et al., 2005) gives a huge range of values: percent colony carbon as mucus ranges from 0.2-99.6% for P. globosa, 1.4-94.3% for P. antarctica and 55.8-99.8% for P. pouchetii. Using a standard colony diameter of 200 µm increases biomass estimates by a factor of 1.2 for colonial P. globosa and P. antarctica cells, but 32.8 for P. pouchetii compared to estimates considering cell biomass alone. The contribution of (standard) mucus to total carbon per cell is 96.9 % for P. globosa, and 14.6% for P. pouchetii and P. antarctica (Table 2) for this standard colony size. The difference between the three species leads to a greater contribution of the Northern Hemisphere species to total *Phaeocystis* biomass (Fig. 9a and b).

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Total *Phaeocystis* biomass estimates including (standard) mucus range from  $2.9 \times 10^{-5} \, \mu g \, C \, I^{-1}$  to  $19\,823 \, \mu g \, C \, I^{-1}$ . The maximal total biomass ( $19\,823 \, \mu g \, C \, I^{-1}$ ) is 3.6 times higher than the corresponding data point with the maximal cell biomass of 5449.3  $\, \mu g \, C \, I^{-1}$ . This data point is associated with high cell numbers during a bloom of *P. pouchetii* off the coast of the Netherlands in the Wadden Sea. In contrast, the maximal total biomass in the Southern Hemisphere is only  $918 \, \mu g \, C \, I^{-1}$ , and thus one order of magnitude lower than maximal total biomasses in the Northern Hemisphere (Fig. 9). The global mean of all reported non-zero total biomass values is  $183.8 \, \mu g \, C \, I^{-1}$ , and the median is  $11.3 \, \mu g \, C \, I^{-1}$ . While our publicly available dataset also contains an estimate of maximal and minimal total carbon biomass based on maximal and minimal reported colony sizes (and thus maximal and minimal mucus), we do not visualize these results here. Uncertainties in the mucus contribution to total biomass due to these uncertainties in colony size range from hundreds to thousands of percent, and total carbon biomass estimates are far from certain at this point in time.

#### 4 Discussion

We have estimated the carbon biomass of the haptophyte *Phaeocystis* from microscopic determinations of cell abundances. This approach is associated with several uncertainties:

First, since the data included in this database are sparse, we may have biases that we cannot account for. Whether the biomass estimates truly represent global averages is unclear. Free-living cells of *Phaeocystis* are often ignored in experimental studies, while colonies are counted, despite the fact that there is always a background concentration of *Phaeocystis* cells when this genus is present in colonial form. Furthermore, even though *Phaeocystis* is ubiquitous (Schoemann et al., 2005), our data show a poor spatial resolution and data coverage outside the high-latitude coastal regions. Our biomass estimates for the coastal seas may not be representative of open ocean concentrations. Some areas such as the Pacific Ocean are clearly under-represented

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and we were not able to acquire any Phaeocystis measurements from the North Pacific. Our data is also seasonally biased in the Southern Hemisphere, with 58 % of the data acquired during the summer months. In addition, we note that *Phaeocystis* is only accurately counted at times when it is expected to form large blooms, when there is a strong likelihood that its abundance is high and when scientists are specifically looking for this group. Hence, low background concentrations of single-celled *Phaeocystis* will often be overlooked. Since the single-celled life stages of *Phaeocystis* lack a clear morphological distinction, this gap in our current knowledge is unlikely to be resolved using microscopic methods, but will require genetic identification methods.

Second, there are methodological issues with the determination of abundance data that will influence our biomass calculations. Several data contributors do not report the life stage cells were in at the time of sampling, most likely due to the disruption of colony structure during cell fixation. This fact results in difficulties in distinguishing single and colonial cells. Hence, in order to obtain a lower limit on Phaeocystis cell biomass, we chose to assume undefined cells to be in the form of flagellates, which will bias the resulting biomass calculations. The ratio of free-living to colonial cells is highly variable, but a significant background concentration of free-living cells is present even during bloom conditions. Our assumption that all unspecified cells are flagellates is therefore likely to lead to an underestimation of *Phaeocystis* cell biomass.

Furthermore, non-blooming species such as P. cordata, P. jahnii or P. scrobiculata are not recorded explicitly in our abundance data, but may constitute a non-negligible fraction of total global *Phaeocystis* biomass in some oceanic regions.

Third, there are large uncertainties associated with the conversion of cell abundances to biomass. Cell measurements were only provided for very few datasets; for the majority of the database biovolumes were calculated using mean published cell dimensions. Cell size is highly variable for all *Phaeocystis* species (Schoemann et al., 2005) and using a constant biovolume estimate for each species will underestimate the spatial and temporal variability that occurs in *Phaeocystis* biomass. Due to the differences in the reported size range, our estimates of cell carbon content are different from

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some previously reported figures. For example, our estimates of cell carbon content for P. globosa (Table 2; flagellates: 13 pg C cell<sup>-1</sup>; colonial cells: 29 pg C cell<sup>-1</sup>) are higher than estimates by Rousseau et al. (1990; flagellates: 11 pg C cell<sup>-1</sup>; colonial cells: 14 pg C cell<sup>-1</sup>), and our estimates for *P. antarctica* (Table 2; flagellates: 9 pg C cell<sup>-1</sup>; 5 colonial cells: 21 pg C cell<sup>-1</sup>) are higher than those reported in Mathot (2000; flagellates: 3 pg C cell<sup>-1</sup>; colonial cells: 14 pg C cell<sup>-1</sup>) due to these differences in the reported mean cell diameters that were used to calculate the carbon estimates. Furthermore, literature values for the carbon conversion factor are only given for Prymnesiophytes in general, but we lack information on the individual species of *Phaeocystis*, which may have a species-dependent, spatially and temporally varying cell carbon content.

Last, there is a large uncertainty associated with the addition of mucus carbon biomass due to the lack of data on cell forms, colony size and the amount of mucus per colonial cell. Greater use of preservation methods that maintain colony structure, along with routine colony size measurements, would allow for more reliable estimates of colonial mucus carbon. Further data on *Phaeocystis* colony sizes are clearly needed if mucus carbon is to be included in global biomass estimates and model validation. Moreover, there are uncertainties related to the structure of the mucilaginous carbon surrounding colonies. For example, an alternative method for estimating the total carbon biomass of P. globosa has been suggested by van Rijssel et al. (1997), based on the observed hollow structure of the colonies. van Rijssel et al. (1997) compute total biomass per cell based on a linear relationship between colony surface area and carbon content. A comparison of the estimated mean total carbon per P. globosa cell leads to significant differences: For our standard colonies of 200 µm diameter, we find total *P. globosa* carbon per cell to be 33.6 pg C cell<sup>-1</sup> following Rousseau et al. (1990) (Table 2), we compute an amount of 202.5 pg C cell<sup>-1</sup> using van Rijssel et al. (1997). The Rousseau relationship results in 9.6 ng C colony<sup>-1</sup>, whereas the Van Rijssel relationship would lead to 58 ng C colony<sup>-1</sup> for this species. Prior to the publication of Verity et al. (2007), the contribution of mucus carbon to total carbon per cell for P. pouchetii

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was done using the Rousseau et al. (1990) and Mathot et al. (2000) or the Van Rijssel et al. (1997) formulations (Reigstad and Wassmann, 2007). Using these relationships, Reigstad and Wassmann (2007) find a much lower contribution of mucus (10%) to total carbon per cell than what we find using Verity et al. (2007) (96.9%), earlier estimates of *P. pouchetii* mucus carbon may not be compatible with our estimations. Clearly, future studies are needed to address this uncertainty in colony structure and mucus distribution, and the corresponding volume to biomass conversion factors.

#### 5 Conclusions

This is the first attempt at creating a global *Phaeocystis* biomass database. At present, however, we are still far from being able to give a global estimate of *Phaeocystis* biomass concentration. Data are limited by lack of spatial and temporal resolution and at most sampling sites we lack a seasonal cycle that would be necessary to determine reasonable estimates for annual mean biomass concentration. Annual and monthly mean biomasses are of particular interest for the modelling community, but these will only be meaningful if further microscopic data can be added to the database. Targeted explorations of marine ecosystems with the aim to determine phytoplankton biomass would be desirable, but such endeavours tend to be expensive and laborious. A marine census of species biomass would shed light on the relative importance of key marine plankton groups, and their respective importance for global biogeochemical cycling.

### Appendix A

#### A1 Data table

A full data table containing all biomass data points can be downloaded from the data archive PANGAEA, http://doi.pangaea.de/10.1594/PANGAEA.779101. The data

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file contains longitude, latitude, depth, sampling time, abundance counts and biomass concentrations, as well as the full data references.

### A2 Gridded netcdf biomass product

Monthly mean biomass data has been gridded onto a 360 x 180° grid, with a vertical resolution of 33 depth levels (equivalent to World Ocean Atlas depths) and a temporal resolution of 12 months (climatological monthly means). Data has been converted to netcdf format for easy use in model evaluation exercises. The netcdf file can be downloaded from PANGAEA, http://doi.pangaea.de/10.1594/PANGAEA.779101. This file contains total and non-zero abundances, cell biomasses and total biomass estimates. For all fields, the means, medians and standard deviations resulting from multiple observations in each of the 1° pixels are given. The ranges in cell and total biomasses due to uncertainties in cell size and life form are not included as variables in the netcdf product, but are given as ranges (minimum cell biomass, maximum cell biomass; minimum total biomass, maximum total biomass) in the data table.

Acknowledgements. We thank P. Assmy, G. C. Cadée, D. A. Caron, G. R. DiTullio, B. Hansen, I. R. Jenkinson, I. Joint, S.-H. Kang, B. Karlson, D. J. Lonsdale, S. Mathot, R. Riegman, M. W. Silver, W. O. Smith, P. Tett, P. Tréguer, R. Uncles, F. C. Van Duyl, E. L. Venrick, T. Weisse, G. V. Wolfe, and P. Wassmann for the permission to use and redistribute *Phaeocystis* data, and the BODC, JGOFS, OBIS OCB, PANGAEA and WOD databases for providing and archiving data. We also thank E. Buitenhuis for producing the gridded netcdf product, S. Doney for fruitful discussions on quality control, and S. Pésant for archiving the data. M. V. acknowledges funding from ETH Zürich. C. O'B.'s contribution to the research leading to these results has received funding from the European Community's Seventh Framework Programme (FP7 2007-2013) under grant agreement no [238366].

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**Table 1.** List of data contributors, in temporal order; Databases: BODC: British Oceanographic Data Centre, OBIS: Ocean Biogeographic Information System, US JGOFS: US Joint Global Ocean Flux Study OCB: Ocean Carbon and Biogeochemistry, WOD09: World Ocean Database 2009, Institutes: AWI: Alfred-Wegener-Institute, Bremerhaven, Germany, IMARPE: Institut del Mar del Peru, Paita, Peru, IOS: Institute of Ocean Sciences, Sidney, Canada, MMBI: Murmansk Marine Biological Institute, Murmansk, Russia.

Entry No.	Database	Investigator/Institute	Project	Year(s)	Region	No of data points	Reference(s)
1	WOD09	AWI	IAPP	1991	Arctic	6	
2	WOD09	IOS	-	1980	US coast (Oregon)	4	-
3	WOD09	IMARPE	-	1966-1977	Peruvian coastal zone	8	-
4	WOD09	MMBI	-	1955-1997	Kola Bay (Barents Sea)	395	-
5	WOD09	University of Alaska	OCSEAP	1975-1977	Prince William Sound (Gulf of Alaska)	20	-
6	-	M. van Leeuwe & J. Stefels	Ant 16/3 R/V Polarstern	1999	Southern Ocean	33	Koeman (1999)
7	-	M. Estrada	Antarctic 85	1985	Weddell Sea, Antarctica	126	Estrada and Delgado (1990)
8	OBIS	P. Wassmann & T. Ratkova	ArcOD	1993-2003	Arctic	1815	
9	BODC	D. Harbour	BOFS	1989-1991	North Atlantic	13	BODC (1994)
10	-	J. Gibson		1993-1995	East Antarctica	136	J. Gibson (unpublished data)
11	Pangaea	P. Assmy	EIFEX	2004	Southern Ocean	28	Assmy and Philipp (2007)
12	_	M. Estrada	Fronts	1985	Mediterranean Sea	156	Estrada (1991)
13	_	D. Karentz	GRINCHES	2004-2005	Ross Sea, Antarctica	14	
14	_	D. Karentz	Icecolors	1986	Southern Ocean	74	Karentz and Spero (1995)
15	BODC	D. Harbour	JGOFS	1994	Arabian Sea	25	BODC (1998);
16	US JGOFS Data System	W. Smith, D. Caron & D. Lonsdale	AESOPS	1996-1997	Southern Ocean	184	_ ` ` ` `
17	BODC	R. Uncles	LOIS	1994-1995	North Sea	19	BODC (2001);
18	BODC	P. Tett	North Sea Project	1988-1989	North Sea	18	BODC (1992);
19	BODC	I. Joint	OMEX	1994-1995	North Atlantic	7	BODC (1997);
20	BODC	P. Wassmann	OMEX	1994	North Atlantic	186	BODC (1997);
21		J. Peloguin	Boss Sea	2001-2005	Ross Sea. Antarctica	84	J. Peloquin (unpublished data)
22	_	E. Breton	SOMLIT-MONITO	2006-2009	English Channel	216	E. Breton (unpublished data)
23	OCB DMO	M. Silver	VERTIGO	2004	Hawaii	1	
24	-	C. Widdicombe	Western Channel Observatory	1992-2008	English Channel	1248	Widdicombe et al. (2010)
25	Schoemann et al. (2005)	Palmisano	Woodom onamio oboorvatory	1984	McMurdo Sound, Antarctica	10	Palmisano et al. (1986)
26	Schoemann et al. (2005)	S. Mathot		1994-1995	Ross Sea, Antarctica	35	Mathot et al. (2000)
27	Schoemann et al. (2005)	C. Robinson		1993	East Antarctica	1	Robinson et al. (1999)
28	Schoemann et al. (2005)	G. DiTullio		1996	Ross Sea. Antarctica	i	DiTullio et al. (2000)
39	Schoemann et al. (2005)	F. Scott		1992	Fast Antarctica	i	Scott et al. (2000)
30	Schoemann et al. (2005)	S. Kang		1986	Weddell Sea. Antarctica	3	Kang and Fryxell (1993)
31	Schoemann et al. (2005)	P. Tréquer		1988	Scotia Sea, Antarctica	1	Tréguer et al. (1991)
32	Schoemann et al. (2005)	G. Cadée	Marsdiep	1976–1985	Dutch coastal zone	2	Cadée and Hegeman (1986)
33	Schoemann et al. (2005)		Marsdiep	1991	Dutch coastal zone	2	Fransz et al. (1992)
33	Scribernarin et al. (2005)	n. Flansz & G. Cauee	Maisulep	1991	Dutcii coastai zone	2	Cadée and Hegeman (1993)
34	Schoemann et al. (2005)	G. Cadée	Marsdiep	1989-1992	Dutch coastal zone	3	Cadee (1991)
35	Schoemann et al. (2005)	G. Cadée	Marsdiep	1989-1992	Dutch coastal zone	2	Cadee (1991) Cadée (1991)
36	Schoemann et al. (2005)	H. Pieters	Marsdiep	1990	Dutch coastal zone  Dutch coastal zone	1	Pieters et al. (1980)
37				1976	Dutch coastal zone  Dutch coastal zone	4	
38	Schoemann et al. (2005)	R. Riegman F. Van Duyl	Marsdiep	1991	Dutch coastal zone Dutch coastal zone	2	Riegman et al. (1993)
38	Schoemann et al. (2005)		Marsdiep	1995		12	van Duyl et al. (1998)
	Schoemann et al. (2005)	I. Jenkinson			German Bight		Jenkinson and Biddanda (1995)
40	Schoemann et al. (2005)	K. Kennington		1996	Irish Sea	1	Kennington et al. (1999)
41	Schoemann et al. (2005)	T. Weisse		1975-1976	German Bight, North Sea	2	Weisse et al. (1986)
42	Schoemann et al. (2005)	B. Karlson		1993	Skagerrak Strait, North Sea	5	Karlson et al. (1996)
43	Schoemann et al. (2005)	G. Wolfe		1997	Labrador Sea	2	Wolfe et al. (2000)
44	Schoemann et al. (2005)	B. Hansen		1988-1989	Barents Sea	6	Hansen et al. (1990)
45	Schoemann et al. (2005)	A. Luchetta		1991	Barents Sea	1	Luchetta et al. (2000)
46	-	V. Schoemann	BGC of Phaeocystis colonies, EC-FP4	1994	Dutch coastal zone	80	Schoemann et al. (1998)

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**Table 2.** Literature values for conversion factors from abundance to biomass. Cell diameters, biovolumes, carbon content and colony number conversions for *P. globosa, P. pouchetii, P. antarctica*. Reported means, and ranges are given in brackets.

	P. globosa Flagellate	Colonial	P. pouchetii Flagellate	Colonial	P. antarctica Flagellate	Colonial
Diameter (μm)	5.5 <sup>1</sup> (3–8)	7.5 <sup>2</sup> (4.5–10.4)	5.0 <sup>3</sup> (2–8)	5.5 <sup>3</sup> (3–8)	4.8 <sup>4</sup> (2–7.5)	6.6 <sup>5</sup> (3.2–10)
Biovolume <sup>6</sup> (μm <sup>3</sup> )	87 (14–268)	217 (48–589)	65 (4–268)	87 (14–268)	56 (4–221)	151 (17–524)
Carbon per cell (pg)	13 <sup>7</sup> (3–35)	29 <sup>8</sup> (7–71)	10 <sup>7</sup> (1–35)	13 <sup>9</sup> (3–35)	9 <sup>7</sup> (1–29)	21 <sup>10</sup> (3–63)
Colony diameter (µm)		200 (10–30 000)		200 (20–2000)		200 (25–9000)
Colony volume – cell number conversion <sup>8,10</sup> (V: [mm <sup>-3</sup> ])		$\log N_c = 0.51 \log V + 3.67$		$\log N_c = 0.537 \log V + 3.409$		$N_c = \left(\frac{V}{417}\right)^{0.60}$
Colony volume – mucus carbon conversion		335 ng mm <sup>-3 8</sup>				213 ng mm <sup>-3</sup> 10
Colony volume – total carbon conversion				$\log C = 0.924 \cdot V + 3.947^{9}$ $C: [\mu g]; V: [mm^{-3}]$		
Total carbon per cell including mucus (pg)		34 (29–7768)		415 (29–6008)		24 (21–362)
Percent of total carbon associated with mucus contribution (pg)		14.6 (0.2–99.6)	96.9 (1.4–94.3)		14.6 (55.8–99.8	

References for the cell diameters: <sup>1</sup> Rousseau et al. (2007); Schoemann et al. (2005); Rousseau et al. (2007); <sup>3</sup> Wassmann et al. (2005); Rousseau et al. (2007); <sup>4</sup> Schoemann et al. (2005); Mathot et al. (2000); Rousseau et al. (2007); Mathot et al. (2000); Rousseau et al. (2007). References for biovolume conversion, assuming spherical geometry of cells: <sup>6</sup> Hillebrand et al. (1999). Reference for the biovolume-carbon conversion: <sup>7</sup> Menden-Deuer and Lessard (2000). References for colony volume-cell number conversion and for colony volume-mucus biomass conversion: <sup>8</sup> Rousseau et al. (1990); <sup>9</sup> Verity et al. (2007) (colony volume – total biomass conversion); <sup>10</sup> Mathot et al. (2000).

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**Table 3.** Latitudinal distribution of abundance data in ten degree latitudinal bands (-90 to 90°). Number of data points for each latitudinal band. All: all measurements, non-zero: data with non-zero carbon biomass.

Latitudinal band	All data	non-zero data
-9080°	0	0
−80− −70°	334	284
-7060°	283	162
-6050°	1	1
-5040°	37	37
-4030°	0	0
-3020°	0	0
-2010°	6	6
-10-0°	1	1
0–10°	0	0
10–20°	17	17
20–30°	10	10
30–40°	0	0
40–50°	152	30
50–60°	852	284
60–70°	1010	1010
70–80°	727	609
80–90°	94	94

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**Table 4.** Seasonal distribution of abundance data for the Northern and Southern Hemisphere. Number of data points for each month. All: all data, non-zero: Data with non-zero carbon biomass. 27 observations did not include the month, where measurements were taken.

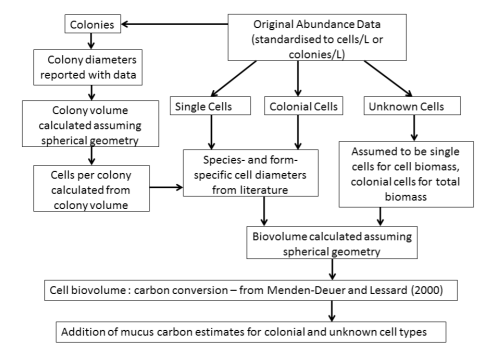
Month	Globe	Globe	NH	NH	SH	SH
	all	non-zero	all	non-zero	all	non-zero
January	164	82	59	4	105	78
February	213	56	59	4	154	52
March	379	187	347	157	32	30
April	687	641	638	593	49	48
May	618	561	612	560	6	1
June	384	318	380	318	4	0
July	263	185	258	183	5	2
August	202	131	198	131	4	0
September	119	56	114	56	5	0
October	169	94	90	27	79	67
November	164	94	67	15	97	91
December	164	130	40	6	124	124
Spring	_	_	1597	1310	181	158
Summer	_	_	836	632	383	254
Autumn	_	_	271	98	87	79
Winter	-	-	158	14	13	2
Total	3526	2547	2862	2054	664	493

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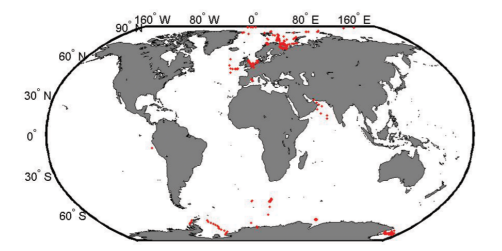


**Fig. 1.** Flow diagram of methodology used to derive mean *Phaeocystis* biomass estimates from abundance data for single cells, colonial cells and unidentified cells. Abundance data was converted to biovolume, and a biovolume to carbon ratio was applied to derive biomass. Finally, an estimate of mucus carbon was added for colonial cell types.

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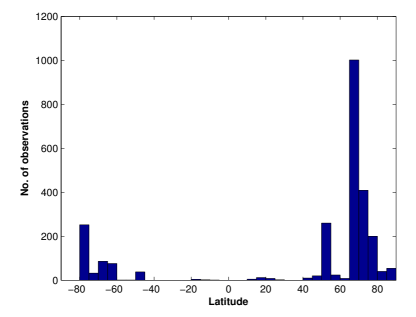


**Fig. 2.** Global distribution of stations where *Phaeocystis* abundance counts were made available for this study. Most stations are located at temperate latitudes and in coastal areas.

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**Fig. 3.** Number of *Phaeocystis* observations as a function of latitude for the period of 1950–2009. Most observations are located in the temperate and high latitudes of the Northern Hemisphere.

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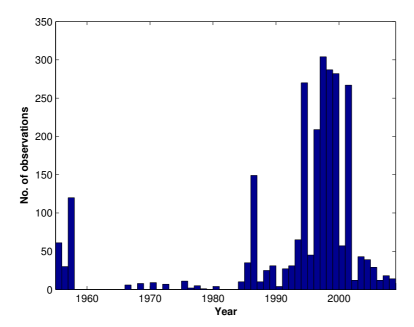
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**Fig. 4.** Number of observations for *Phaeocystis* species per year, for the years 1950–2009. Most counts were made after 1990.

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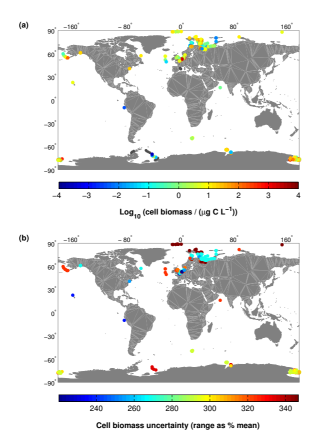


Fig. 5. (a) Surface mean log-normalized Phaeocystis cell biomass concentration in units of carbon ( $\mu g C I^{-1}$ ) and (b) range of uncertainty in cell biomass in % of the mean, due to uncertainty in cell size. Black dots represent zero biomass values. Data has been log-transformed for a better visualization of the wide range of concentrations.

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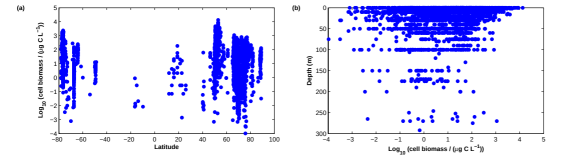
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**Fig. 6.** Distribution of non-zero log-normalized *Phaeocystis* cell biomass ( $\mu g C I^{-1}$ ) (a) as a function of latitude and (b) as a function of depth.

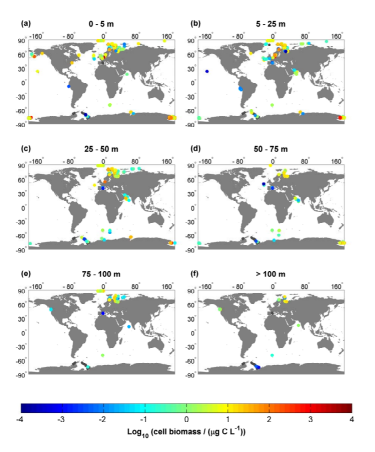
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**Fig. 7.** Log-normalized *Phaeocystis* biomass in units of carbon ( $\mu$ g C I<sup>-1</sup>) at different depths **(a)** surface measurements (0–5 m) **(b)** measurements between 5–25 m **(c)** 25–50 m **(d)** 50–75 m **(e)** 75–100 m and **(f)** >100 m depth. Black dots represent zero biomass values.

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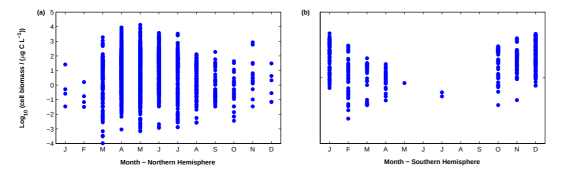
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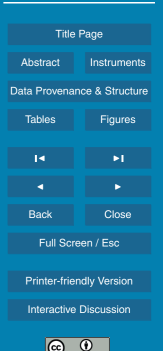




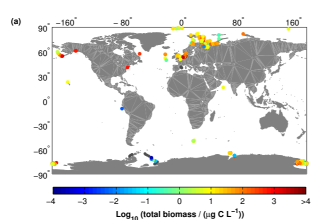
**Fig. 8.** Seasonal distribution of log-normalized non-zero *Phaeocystis* cell biomass data for **(a)** the Northern and **(b)** the Southern Hemisphere.

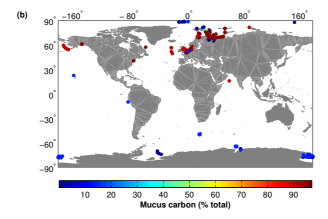
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**Fig. 9.** Estimates of **(a)** log-normalized total mean *Phaeocystis* biomass including colonial mucus for the surface layer (0–5 m) and **(b)** fraction of total mean surface biomass composed of mucus carbon. Zero values are not represented.

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