The Southern Ocean diatom *Pseudo-nitzschia subcurvata* flourished better under simulated glacial than interglacial ocean conditions: Combined effects of CO$_2$ and iron

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Abstract

The ‘Iron Hypothesis’ suggests a fertilization of the Southern Ocean by increased dust deposition in glacial times. This promoted high primary productivity and contributed to lower atmospheric pCO$_2$. In this study, the diatom *Pseudo-nitzschia subcurvata*, known to form prominent blooms in the Southern Ocean, was grown under simulated glacial and interglacial climatic conditions to understand how iron (Fe) availability (no Fe or Fe addition) in conjunction with different pCO$_2$ levels (190 and 290 μatm) influences growth, particulate organic carbon (POC) production and photophysiology. Under both glacial and interglacial conditions, the diatom grew with similar rates. In comparison, glacial conditions (190 μatm pCO$_2$ and Fe input) favored POC production by *P. subcurvata* while under interglacial conditions (290 μatm pCO$_2$ and Fe deficiency) POC production was reduced, indicating a negative effect caused by higher pCO$_2$ and low Fe availability. Under interglacial conditions, the diatom had, however, thicker silica shells. Overall, our results show that the combination of higher Fe availability with low pCO$_2$, present during the glacial ocean, was beneficial for the diatom *P. subcurvata*, thus contributing more to primary production during glacial compared to interglacial times. Under the interglacial ocean conditions, on the other hand, the diatom could have contributed to higher carbon export due to its higher degree of silicification.

Introduction

The Southern Ocean (SO) is the world’s largest high-nutrient low-chlorophyll region (HNLC) and an area where physical forcing, atmospheric pCO$_2$, biological production and marine biogeochemical cycles are tightly linked. In this region, primary production is restricted by the bioavailability of the trace metal (TM) iron (Fe) [1–3]. Fe is an essential trace element, which is needed by phytoplankton to transfer electrons in key cellular and metabolic processes including photosynthesis, respiration, chlorophyll production, carbon (C) and nitrogen (N) fixation [4]. The availability of Fe strongly influences phytoplankton species composition and growth [5–9], and impacts the biological carbon pump and thus the global carbon cycle.
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Consequently, changes in Fe availability have caused feedback effects on climate over geological timescales. Furthermore, the SO is a region of high CO$_2$ exchange between ocean and atmosphere [10, 11]. The SO has been reported to be a major sink of atmospheric CO$_2$ during glacial periods, while it was on the other hand a source of CO$_2$ during glacial-interglacial transitions. At present day, the SO is the major sink of anthropogenic CO$_2$ [12, 13].

During the Last Glacial Maximum (LGM), the SO experienced changes in oceanic circulation and carbon storage. For instance, increased sea ice extent strengthened surface water stratification, thus limiting ocean ventilation and trapping more carbon in the deep ocean [14–16]. Additionally, the northward displacement of the westerly winds prevented the upwelling of CO$_2$-rich deep water [17]. Besides physical mechanisms, the strength of the biological pump might explain 25–50% of the roughly 100 μatm pCO$_2$ discrepancy between glacial (180 μatm pCO$_2$) and interglacial (280 μatm pCO$_2$) times as argued in several studies [18–21]. This supports John Martin’s ‘Iron Hypothesis’, which suggests that an increase in dust deposition during glacial times would fertilize the ocean, stimulate marine productivity, and enhance C export [1, 2, 22–24]. Indeed, analysis of sediment cores revealed a positive correlation between aeolian Fe supply and primary production during ice ages [20, 25]. Proxy data as well as model simulations showed a doubling of the global dust deposition during the last glacial climate condition, when 826 Tg/yr dust were deposited in the global ocean, compared to the 440 Tg/yr dust in pre-industrial times [26, 27]. The difference was mainly due to a dryer atmosphere and reduced vegetation cover [22]. Used a biogeochemical model to estimate the impact of Fe deposition on the global ocean. Under current conditions, 33% of the world’s oceans water masses have Fe concentrations, which limit the growth of phytoplankton. The model simulations revealed that the percentage of Fe-poor water masses decline to 25% and 13% with pre-industrial and LGM dust input, respectively. Along with the North Pacific Ocean, the SO showed the most significant difference in soluble Fe deposition during glacial and interglacial times, accordingly having the largest impact on marine biogeochemistry [22]. The dust deposition in the SO during glacial times was roughly ten times higher (0.04–0.17 Tg/yr) than in pre-industrial times (0.005–0.018 Tg/yr) [27]. However, the SO is geographically isolated from arid, dust-producing regions and is thus overall characterized by low aeolian Fe deposition [28]. Other sources of Fe include upwelling of deep nutrient-rich water, entrainment of sedimentary Fe from continental shelves and resuspension, island-wake effects, seasonal sea ice extent and melt, as well as iceberg drift and melt [3 and references therein].

The phytoplankton community in the current SO is dominated by different diatom species and the pycnocystophyte Phaeocystis antarctica [29]. Diatoms account for 40% of the ocean’s total primary production [30–33] and dominate the export of particulate organic matter to the seafloor [34, 35]. In other words, diatoms are crucial for the ocean’s ability to sequester C to the ocean’s interior. Diatoms also have an extensive impact on the oceanic silica inventory, as they produce frustules containing silica. Some frustules are resistant to remineralization and dissolution, are well preserved in the sediment, and thus provide precious information about past oceanic biogeochemistry. Pseudo-nitzschia species have been frequently observed in today’s phytoplankton assemblages in Antarctic waters [36]. Mesoscale Fe fertilization experiments in the SO triggered massive phytoplankton blooms dominated by large diatoms like the pennate Pseudo-nitzschia sp. [7, 37]. Large diatoms in the SO appear to have a higher Fe requirement compared to smaller phytoplankton because of physical constrains in the Fe uptake process [38]. To compensate for this, they have evolved various strategies to acquire bioavailable Fe. They generally reduce their biogeochemical Fe requirement through metal or protein substitution [39] and reduce Fe-rich components of the photosystem apparatus [9]. [8] suggested that Pseudo-nitzschia is able to accumulate intracellular Fe when ambient concentrations of this TM are high, while maintaining a low Fe demand. This luxury uptake and
subsequent storage of Fe supports growth in subsequent low Fe environments and enables *Pseudo-nitzschia* to dominate phytoplankton assemblages across a wide range of oceanic Fe concentrations.

In SO diatoms, Fe limitation often results in slower growth and reduced C fixation. The photochemical quantum efficiency, which indicates how efficiently excitation energy is transferred to the reaction centers, is usually lowered [8, 40–42]. In an Fe-poor environment, cells usually increase the functional absorption cross sectional area of their reaction centers, thereby enhancing the target area, which absorbs incoming photons [38, 41]. The absorbed photons can either drive photosynthesis, N reduction, C fixation, photorespiration or can be converted to heat (non-photochemical quenching). Fe deficiency induces changes in the photosystem II (PSII) reaction centers such as the reduction of the pigment content [43], causes less efficient electron transport [40] and increases non-photochemical quenching to dissipate the excess light energy [44].

Besides Fe limitation, phytoplankton cells have experienced variations in CO$_2$ concentration in the past. Previous studies on the effect of high CO$_2$ concentrations on phytoplankton reported changes in their elemental composition (e.g. [45, 46]), in cell size (e.g. [47]) and in the degree of silicification in diatoms (e.g. [48]). Furthermore, it was shown that low pCO$_2$ levels can influence the composition of Antarctic phytoplankton communities. For example, experiments with natural phytoplankton assemblages from different regions across the SO [24, 49, 50] concluded that *Pseudo-nitzschia* flourishes at low pCO$_2$ levels, while it does not do well in response to ocean acidification. Indeed, between ambient and future elevated pCO$_2$ levels, the growth of *P. subcurvata* in a laboratory experiment was not stimulated under enhanced Fe supply [51]. Under similar Fe conditions, a phytoplankton community from the Ross Sea, Antarctica, responded to CO$_2$ increase from 100 to 800 ppm with a dramatic reduction in cell abundance of *P. subcurvata*, being replaced by *Chaetoceros* species [49]. Similarly, a community from the Weddell Sea, Antarctica, shifted from *Pseudo-nitzschia* to *Fragilariopsis* after Fe addition between 390 to 800 μatm pCO$_2$ [50], while no difference in species composition was found between the glacial (190 μatm) and the present-day (390 μatm) pCO$_2$ levels. This implies that reduced CO$_2$ concentrations during glacial periods potentially favored pennate diatoms such as *Pseudo-nitzschia* while diatom species such as *Chaetoceros* and *Fragilariopsis* became most abundant under present-day and future pCO$_2$ levels [49]. A few studies investigated the SO phytoplankton assemblages and growth under low Fe supply in response to increasing pCO$_2$ [24, 50, 52]. [50] observed also a CO$_2$-dependent taxonomic shift in Fe-deplete conditions with increasing pCO$_2$ with *Pseudo-nitzschia* being replaced by the pennate diatom *Synechocystis* between 390 and 800 μatm pCO$_2$ levels. Similarly, when pCO$_2$ increased from 390 to 900 μatm another SO plankton community changed from being dominated by *P. proligeroides* to one, which was dominated by *P. antarctica* [24]. Hence, irrespective of Fe availability the genus *Pseudo-nitzschia* was found to be susceptible to ocean acidification pCO$_2$ levels.

Studies that assess the effects of low pCO$_2$ on phytoplankton often compare their results with high pCO$_2$ levels to understand ocean acidification. However, little is known about the smaller variation from 180 (glacial) to 280 μatm (interglacial/pre-industrial) pCO$_2$ under different Fe availability. Indeed, the potential interactive effect of low-pCO$_2$ (180 and 280 μatm) together with different Fe availability (deplete and replete) on net primary production and export production is currently often not considered, when developing models or designing laboratory experiments simulating glacial and interglacial ocean conditions. Studies looking at N-isotopes and Th-corrected sediment accumulation rates describe large fluxes of biogenic detritus out of surface waters in the glacial ocean due to a larger amount of lithogenic Fe transported by winds [26]. The latter study indicates that increased export production and thus
enhanced C storage potentially contributed to the observed lower atmospheric CO$_2$ concentrations during glacial times [53]. The above-mentioned studies offer first insights on how some phytoplankton species cope with glacial and interglacial climatic conditions. However, studies on the ecophysiology of Antarctic diatoms subject to glacial vs. interglacial ocean conditions under reduced Fe conditions, are yet lacking. In this study, the SO bloom-forming diatom _P. subcurvata_ was grown under Fe and CO$_2$ conditions representative of glacial (lower CO$_2$ and higher Fe) and interglacial (higher CO$_2$ and lower Fe) times to untangle the influence of these two environmental factors on growth, elemental stoichiometry, photosynthetic carbon production and photophysiology. This allowed to assess its role in the paleo carbon cycle.

**Material and methods**

**Experimental setup**

Prior to the execution of the experiment, the oceanic diatom _P. subcurvata_ (isolated by Philipp Assmy at 49° S, 2° E, R/V Polarstern cruise ANT-XXI/4, April 2004) was grown for more than one year in Antarctic seawater with a low total dissolved Fe (dFe) concentration of 0.5 nmol L$^{-1}$. Preacclimation and the main experiment were carried out in Fe-poor (0.4 nmol L$^{-1}$) Antarctic seawater collected at 60°32'S, 26°29'W (salinity of 33.8 ± 0.2), filtered through a sterilized, acid-cleaned 0.2 μm filter (Sartobran, Sartorius). This water was spiked with chelexed (Chelex 110, Sigma Aldrich, Merck) macronutrients (100 μmol L$^{-1}$ Si, 100 μmol L$^{-1}$ NO$_3$ and 6.25 μmol L$^{-1}$ PO$_4$) and vitamins (30 nmol L$^{-1}$ B$_1$, 23 nmol L$^{-1}$ B$_7$ and 0.228 nmol L$^{-1}$ B$_{12}$) according to the F/2$_R$ medium [54]. In addition, a TM mix containing Zn (0.16 nmol L$^{-1}$), Cu (0.08 nmol L$^{-1}$), Co (0.09 nmol L$^{-1}$), Mn (1.9 nmol L$^{-1}$), Mo (0.05 nmol L$^{-1}$) in the ratio of the original F/2 recipe adjusted to 4 nmol L$^{-1}$ Fe was added. As suggested by [55], in order to minimize the alteration of the natural seawater TM chemistry and ligands, no ethylenediaminetetraacetic acid (EDTA) was added. The Fe-deplete treatments (henceforth referred to as Control) contained 0.4 nmol L$^{-1}$ dFe while for the Fe-enriched treatments (henceforth referred to as +Fe), 4 nmol L$^{-1}$ FeCl$_3$ were added.

To avoid Fe contamination, TM clean techniques were used according to the GEOTRACES cookbook [56]. The sampling and handling of the incubations was conducted under a laminar flow hood (Class 100, Opta, Bensheim, Germany). All equipment was soaked for one week in 1% Citranox, followed by two weeks in 1 N HCl for polycarbonate and 5 N HCl for polyethylene materials. In between and after the cleaning process, the equipment was rinsed seven times with Milli-Q (MQ, Millipore). Finally, everything was air dried under a clean bench (U. S. class 100, Opta, Bensheim, Germany) and packed in three polyethylene bags.

All Control and +Fe incubations were bubbled with humidified air containing pCO$_2$ levels of 190 and 290 μatm, henceforth referred to as 190 and 290, respectively. Using a gas flow controller (CGM 2000, MCZ Umwelttechnik, Bad Nauheim, Germany), both CO$_2$ gas mixtures were generated by combining CO$_2$ free air (< 1 ppmv CO$_2$, Dominick Hunter, Kaarst, Germany) with pure CO$_2$ (Air Liquide Deutschland Ltd., Düsseldorf, Germany) in the respective ratios. They were regularly monitored with a Li-Cor (LI6252 Biosciences, Lincoln, NE) calibrated with CO$_2$ free air and purchased gas mixtures of 150 ± 10 and 1000 ± 20 ppmv CO$_2$ (Air Liquide Deutschland Ltd., Düsseldorf, Deutschland). Low pCO$_2$ and Fe input characterized the glacial ocean, which was here simulated in the +Fe 190 treatment. Vice versa, the interglacial ocean was characterized by higher pCO$_2$ and no Fe input and mimicked by the Control 290 treatment. In addition to the incubation bottles, Fe and carbonate chemistry were determined in the culture medium which was incubated in the same way as the respective
incubation bottles (pCO₂ and Fe availability), to check if the different pCO₂ and Fe mani-
pulations were successful. All incubations were placed in front of LED (light-emitting diode) lamps at 100 μmol photons m⁻² s⁻¹ under a light:dark cycle of 16:8 h. The light intensity was adjusted with a LI-1400 datalogger (Li-Cor Biosciences, Lincoln, NE, USA) with a 4π-sensor (Walz, Effeltrich, Germany). For this experiment, the long-term low Fe acclimated *P. subcurvata* stock culture was inoculated to the different CO₂-Fe conditions and was acclimated to each experimental condition at 2°C for at least two weeks. The main experiment was carried out in triplicate 4 L acid-cleaned polycarbonate bottles for each experimental treatment. The main experiment started with initial cell densities of ~1000 cells mL⁻¹, lasted between 8 and 9 days and reached final cell densities between 67000 and 107000 cells mL⁻¹.

**Trace metal chemistry**

At the end of the experiment, total dissolved Fe (dFe) samples were taken from the culture medium by filtering 100 mL from each bottle through 0.2 μm HCl-cleaned polycarbonate filters (47 mm, Nuclepore, Whatman, GE Healthcare, Chicago, IL, USA) using a trace metal clean filtration system under a clean laminar flow hood (Class 100, Opta, Bensheim, Germany). The filtrate was then filled into a 125 ml HCl-cleaned PE bottle and stored triple-bagged at 2°C until analysis. Between each filtration, the filtration manifold was cleaned in an acid bath consisting of 1 M HCl and rinsed seven times with Milli-Q. Prior to the dFe analysis, 0.2 μm pre-filtered seawater samples were acidified to pH 1.75 with double distilled HNO₃, minimizing the formation of Fe and Mn hydroxides. Next, samples were UV (ultraviolet) oxidized for 1.5 h using a 450 W photochemical UV power supply (photochemical lamp 7825; Power Supply 7830, ACE GLASS Inc., Vineland N.J., USA). Total dFe concentration of the seawater samples and the processed blanks were measured with a seaFAST system (Elemental Scientific, Omaha, NE, USA) [57] coupled to a sector field inductively coupled plasma mass spectrometer (ICP-MS; Element 2, Thermo Fisher Scientific; resolution of R = 4000; oxide forming rates below 0.3%). To minimize matrix effects, the seawater dFe concentrations were analyzed by standard addition. The accuracy of the dFe data was assessed by measuring NASS-6 (National Research Council of Canada) reference standards, with a recovery rate for Fe of 110%.

**Carbonate chemistry**

From the culture medium as well as from the incubation bottles at the end of the experiment, dissolved inorganic carbon (DIC) was filtered through 0.2 μm filters (Nalgene, Thermo Scientific) and was stored at 4°C in 5 mL borosilicate glass bottles without headspace. The colori-
metric analysis was performed with a QuAAtro autoanalyzer (Seal Analytical, [58]). Again, from the culture medium as well as from the incubation bottles at the end of the experiment, samples for the total alkalinity (TA) were filtered through 0.6 μm GF/F filters (Whatman) and stored at 4°C in 150 mL borosilicate glass bottles. TA was measured via potentiometric titra-
tion [59] and the concentrations were calculated using a linear Gran Plot [60]. The pCO₂ was calculated using the CO2Sys program [61] with the equilibrium constants of [62] as refitted by [63] using TA and DIC measurements, concentrations of phosphate and silicate, temperature and salinity.

**Growth**

Cell count samples of *P. subcurvata* were fixed with 10% acid lugol solution and stored at 2°C in the dark until counting. Cell numbers of *P. subcurvata* were enumerated according to the
method by [64] using 3 ml sedimentation chambers (Hydrobios, Kiel, Germany) on an inverted microscope (Zeiss Axiovert 200) counting at least 400 cells.

The growth rates $\mu (d^{-1})$ were determined with

$$\mu = \frac{\ln (N_t)}{\Delta t}$$

where $N_0$ and $N_t$ denote the initial and the final cell concentrations of the experiments, respectively and $\Delta t$ is the incubation time in days. Final harvest took place when the cells were in exponential growth and reached densities between 67 000 and 107 000 cells mL$^{-1}$.

The cell volume was computed using the volume formula of a prism on parallelogram base provided by [65]. The apical and transapical axes were measured via microscopy, while the pervalvar axis was estimated to be half of the transapical axis with an average value of 1.2 $\mu$m.

### Elemental composition

At the end of the experiment, particulate organic carbon (POC) and particulate organic nitrogen (PON) were measured after filtering onto pre-combusted (15 h, 500°C) GF/F filters (pore size ~ 0.6 $\mu$m, Whatman). The amount of seawater filtered ranged between 200–300 mL and was dependent on the biomass in the treatments. Filters were stored at -20°C and dried for > 12 h at 60°C. Analysis was performed using a Euro Elemental Analyzer 3000 CHNS-O (HEKAtech GmbH, Wegberg, Germany). At the end of the experiment, samples to determine biogenic silica (BSi) were filtered through a cellulose acetate filter (Sartorius, 0.6 $\mu$m) and stored at -20°C. The dried filters were submerged in 0.2 M NaOH at 95°C for 45 minutes, cooled in an ice bath for 15 minutes, neutralized with 1 M HCl according to [66] and analyzed colorimetrically for silicate using standard spectrophotometric techniques [67]. Contents of POC, PON and BSi were corrected for blank measurements and normalized to filtered volume and cell densities to obtain cellular quotas. Production rates of POC, PON and BSi were calculated by multiplying the cellular quotas with the respective growth rate.

### Pigments

The amount of seawater filtered to collect pigment ranged between 200–300 mL on the GF/F filter and was dependent on the biomass in the treatments. Each pigment sample was flash frozen in liquid nitrogen and stored at -80°C until analysis. First, the pigments were homogenized and extracted for 24 h in 90% acetone at 4°C in the dark. Second, they were centrifuged for five minutes (4°C, 13000 rpm) and filtered through a 0.45 $\mu$m pore size nylon syringe filter (Nalgene, Nalge Nunc International, Rochester, NY, USA). The pigments were analyzed by reversed phase High Performance Liquid Chromatography (HPLC) on a LaChromElite system equipped with a chilled autosampler L-2200 and a DAD detector L- 2450 (VWR-Hitachi International GmbH, Darmstadt, Germany). A SpherisorbODS-2 column (25 cm × 4.6 mm, 5 $\mu$m particle size; Waters, Milford, MA, USA) with a LiChropher100-RP-18 guard cartridge was used for the separation of pigments, applying a gradient according to [68]. Peaks of light harvesting (LH) pigments chlorophyll a (Chl a) and c$_2$ (Chl c$_2$), fucoxanthin (Fuco), as well as of the light protective (LP) pigments diatoxanthin (Dt) and diadinoxanthin (Dd) were detected, identified and quantified by co-chromatography with the corresponding pigment standards (DHI Lab Products, Horsholm, Denmark) using the software EZChrom Elite ver. 3.1.3. (Agilent Technologies, Santa Clara, CA, USA). Pigment contents were normalized to filtered volume and cell densities to obtain cellular quotas.
Photophysiological parameters

The efficiency of photochemistry in the PSII of *P. subcurvata* was assessed regularly during and at the end of the experiment by means of a Fast Repetition Rate fluorometer (FRRf, FastOcean PTX) and a FastAct Laboratory system (both from Chelsea Technologies Group Ltd., West Molesey, United Kingdom). Values were obtained using the FastPro8 software (Version 1.0.50), [69]. Measurements were performed at least 2 hours after begin of the light period at 2˚C after 10 minutes of dark-adaptation to ensure that all PSII reaction centers were fully oxidized and non-photochemical quenching (NPQ) was relaxed [70]. For each treatment, a 0.2 μm filtered blank was collected, measured and subtracted.

The fluorometer’s LED (wavelength 450 nm) was automatically adjusted to a light intensity of 1.2 \(10^{22}\) photons m\(^{-2}\) s\(^{-1}\). A single turnover flashlet was applied to cumulatively saturate PSII, thus to close all PSII reaction centers, and consisted of 100 flashlets on a 2 μs pitch, followed by a relaxation phase made of 40 flashlets on a 50 μs pitch to reopen the PSII reaction centers. The saturation phase of the single turnover acquisition, comprised 24 sequences and was fitted according to [71]. The minimum \((F_0)\) and maximum \((F_m)\) Chl a fluorescence were determined and the apparent maximum PSII quantum yield \((F_v/F_m)\) was calculated according to the equation:

\[
F_v/F_m = (F_m - F_0)/F_m
\]

Further outputs of the FastPro8 software from the single turnover measurements of dark-adapted cells were the connectivity between PSII \((P, \text{dimensionless})\), thus the energy transfer between PSII units, the time constant for electron transport at the acceptor side of PSII \((\tau, \mu s)\), the functional absorption cross section of PSII photochemistry \((\sigma_{PSII}, \text{nm}^{-2})\) and the cellular concentration of functional PSII reaction centers \((RCII, \text{zmol cell}^{-1})\).

During the photosynthesis-irradiance-curve (PE-curve), cells were exposed to eight light levels ranging from 0 to 1868 μmol photons m\(^{-2}\) s\(^{-1}\) for five minutes each. At each light level, six measurements of the light-adapted minimum \((F_0')\) and maximum \((F_m')\) Chl a fluorescence were taken and the effective PSII quantum yield \((F_q'/F_m')\) was calculated [72].

Cellular electron transport rates \((cETR)\) were calculated following [73, 74] and normalized by RCII [75] using:

\[
cETR = RCII \cdot \sigma_{PSII} \cdot E \cdot \frac{F_q'/F_m'}{F_q/F_m}
\]

where E (photons m\(^{-2}\) s\(^{-1}\)) is the applied instantaneous irradiance, which was measured separately for each light level in seawater.

The cETR versus E curve was fitted according to [76] allowing to derive the maximum cETR \((cETR_{max})\), the minimum saturating irradiance \((I_K)\) determined by the interception of the light-limited region with the maximum photosynthetic rate, and the maximum light utilization efficiency \((\alpha)\).

NPQ of Chl a fluorescence was calculated using the Stern-Volmer equation [77] at each light level:

\[
NPQ = \frac{F_m}{F'_m} - 1
\]
Statistical assessment

To assess the effect of Fe concentration (Control and +Fe) and pCO₂ (190 and 290) on all experimental parameters among the different treatments of \textit{P. subcurvata}, we used a two-way analysis of variance (2-way ANOVA) followed by a pairwise multiple comparison test (post hoc) using the Holm-Sidak method. All statistical analyses and the curve fittings were performed using the program SigmaPlot (Version 13.0 from Systat Software, Inc., San Jose California USA, www.systatsoftware.com). Statistical significance was defined when p < 0.05.

Results

Trace metal and carbonate chemistry

The total dFe concentrations of the different culture medium showed a significant difference between the +Fe and the Control treatments (2-way ANOVA: p < 0.001, Table 1), with the +Fe treatments having higher dFe concentrations than the Control treatments. The parameters of the carbonate system are given in Table 1. TA remained constant in all culture media and incubation bottles. As expected, increasing pCO₂ significantly enhanced the DIC concentration in all culture media and incubation bottles (2-way ANOVA: p < 0.001; post hoc +Fe: p < 0.001; post hoc Control: p = 0.005). While Fe availability did not alter DIC of the different culture media bottles, a significant Fe effect was found for the \textit{P. subcurvata} incubations, but only for the 190 treatments (post hoc: p < 0.04). The interaction of CO₂ and Fe also led to significant effects in DIC of the \textit{P. subcurvata} incubations (2-way ANOVA: p < 0.02). As expected, the pCO₂ and DIC in all of the 290 treatments were significantly higher than in the 190 treatments (2-way ANOVA; p < 0.001; Table 1). Biologically driven changes to the carbonate chemistry were ruled out since TA, DIC, and pCO₂ values did not differ between the abiotic culture medium and the corresponding \textit{P. subcurvata} incubations for each treatment at the end of the experiment (Table 1).

Growth and elemental composition

The growth rates of \textit{P. subcurvata} were unaffected by Fe deficiency and changes in pCO₂ (Fig 1A). Similarly, cell volumes remained constant across all treatments (Table 2).

Table 1. Total dissolved iron (dFe) concentrations and carbonate chemistry determined at the end of the experiment in the culture medium (filtered seawater without cells) and the \textit{P. subcurvata} incubations of the four treatments (+Fe 190, Control 190, +Fe 290 and Control 290). The pCO₂ was calculated from measured dissolved inorganic carbon (DIC) and total alkalinity (TA). For the culture medium, dFe, TA, DIC and pCO₂ values represent the range of duplicate abiotic controls. TA, DIC and pCO₂ values of the \textit{P. subcurvata} incubations represent the means ± SD (n = 3). Differences between the individual treatments of the \textit{P. subcurvata} incubations were determined with post hoc tests, where significant statistical (p < 0.05) differences are denoted by different letters.

| Parameter | Culture medium | \(190\) | \(290\) |
|-----------|----------------|--------|--------|
| dFe (nmol L\(^{-1}\)) | +Fe | 2.92–3.10 | 0.94–1.07 | 1.36–1.41 | 0.37–0.50 |
| TA (μmol kg\(^{-1}\)) | Control | 2308–2318 | 2304–2319 | 2304–2323 | 2302–2311 |
| DIC (μmol kg\(^{-1}\)) | +Fe | 2077–2101 | 2058–2077 | 2125–2131 | 2131–2132 |
| pCO₂ (μatm) | Control | 208–249 | 201–208 | 269–308 | 296–309 |

| Parameter | \(P. subcurvata\) incubations | \(190\) | \(290\) |
|-----------|----------------|--------|--------|
| dFe (nmol L\(^{-1}\)) | +Fe | - | - | +Fe | - |
| TA (μmol kg\(^{-1}\)) | Control | 2317 ± 11 \(^a\) | 2326 ± 9 \(^a\) | 2327 ± 13 \(^a\) | 2320 ± 9 \(^a\) |
| DIC (μmol kg\(^{-1}\)) | +Fe | 2046 ± 17 \(^a\) | 2071 ± 11 \(^b\) | 2138 ± 14 \(^c\) | 2118 ± 2 \(^c\) |
| pCO₂ (μatm) | Control | 181 ± 15 \(^a\) | 202 ± 24 \(^a\) | 287 ± 31 \(^b\) | 283 ± 29 \(^b\) |

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Cellular POC quotas (Table 2) and POC production rates (Fig 1B) in both pCO$_2$ treatments were not affected by Fe deficiency. On the other hand, the increase of CO$_2$ concentration resulted in a 20–30% decrease of cellular POC quotas (2-way ANOVA: p < 0.001; Table 2) and POC production (2-way ANOVA: p < 0.001; Fig 1B) in both Control and +Fe treatments.

At 190, lowered Fe concentration led to a decrease of cellular PON concentrations by 19% (post hoc: p < 0.03), while no Fe effect was observed at 290. In response to increasing pCO$_2$, the cellular PON concentration was strongly reduced (2-way ANOVA: p = 0.005; Table 2) in the +Fe (post hoc: p < 0.004), but not in the Control treatments (Table 2). The PON production (Fig 1C) followed the same pattern as cellular PON quotas, showing a significant decrease of 15% with reduced Fe availability in the 190 treatments (post hoc: p < 0.03), while remaining constant in the 290 treatments. With increasing pCO$_2$, a loss of 26% in PON production in the +Fe (post hoc: p < 0.02), but not in the Control treatments was observed, resulting from an interactive effect of Fe and CO$_2$ availability (2-way ANOVA: p < 0.02; Fig 1C).

Molar C:N ratios ranged between 6.9 ± 0.1 and 8.1 ± 0.2 mol mol$^{-1}$. Fe deficiency led to a 13% increase in the C:N ratio in the 190 treatments (post hoc: p < 0.04), while no such Fe effect was observed in the 290 treatments. Furthermore, the increase of CO$_2$ concentration resulted in a decline of C:N by 15% in the Control (post hoc: p < 0.02), but not in the +Fe treatments. The interaction of Fe and CO$_2$ altered C:N ratios significantly (2-way ANOVA: p < 0.03; Table 2).
Neither low Fe concentrations nor increased pCO$_2$ changed the cellular BSi quota (Table 2). However, as a result of Fe deficiency the BSi production in 290 significantly increased by 35% (2-way ANOVA: p = 0.007; post hoc: p = 0.006; Fig 1D), but not in 190. A response to higher pCO$_2$ resulted in higher BSi production only in the Control treatments (post hoc: p < 0.04).

**Pigment composition**

All quantified pigments, except for Chl $c_2$, were significantly affected by Fe deficiency in either the 190 or the 290 treatments (2-way ANOVA: Chl $a$ p < 0.001; Fuco p < 0.02; Dd p < 0.02; Dt p < 0.02; Fig 2A and Table 3). At 190, reduced Fe availability resulted in a decrease of Chl $a$
### Table 3. Pigment concentrations determined at the end of the experiment in the four treatments of *P. subcurvata* (+Fe 190, Control 190, +Fe 290 and Control 290). The values represent the means ± SD (n = 3). Different letters indicate significant differences between treatments.

| Parameter                  | *P. subcurvata* incubations |                     |                     |
|----------------------------|----------------------------|---------------------|---------------------|
|                            | +Fe | Control | +Fe | Control |
| Chlorophyll c₂ (fg cell⁻¹) | 28.9 ± 6.9ᵃ | 19.4 ± 5.7ᵃ | 22.3 ± 5.1ᵃ | 19.1 ± 5.0ᵃ |
| Fucoxanthin (fg cell⁻¹)    | 140 ± 10ᵃ | 93 ± 24ᵇ | 110 ± 16ᵃ | 86 ± 24ᵃᵇ |
| Diadinoxanthin (fg cell⁻¹)  | 28.3 ± 3.8ᵃ | 19.4 ± 4.5ᵇ | 24.1 ± 4.3ᵃ | 17.7 ± 3.9ᵃᵇ |
| Diatoxanthin (fg cell⁻¹)   | 1.27 ± 0.24ᵃ | 1.08 ± 0.12ᵃᵇ | 1.47 ± 0.44ᵃ | 0.64 ± 0.21ᵇ |
| Chl a:C (mol mol⁻¹)         | 0.21 ± 0.03ᵃ | 0.14 ± 0.02ᵇ | 0.22 ± 0.04ᵃ | 0.16 ± 0.03ᵇ |

by 37% (post hoc: p = 0.002), of Fuco by 34% (post hoc: p < 0.02) and of Dd by 29% (post hoc: p = 0.03), while Dt was not affected. At 290, the reduction of Fe significantly reduced the Chl a concentration by 23% (post hoc: p = 0.03) and Dt by 60% (post hoc: p = 0.007), whereas Fuco and Dd remained constant. In response to elevated pCO₂, cellular Chl a quotas of *P. subcurvata* were significantly reduced in the +Fe (251 ± 17) by 192 ± 19 fg cell⁻¹ for 190 and 290, respectively; 2-way ANOVA: p < 0.03; post hoc: p = 0.02; Fig 2A), while this trend was absent in the Control. No other pigments (Fuco, Chl c₂, Dd or Dt) responded to changes in the pCO₂.

The Chl a:C ratio in *P. subcurvata* was significantly affected by Fe deficiency (2-way ANOVA: p = 0.005; Table 3) leading to a decrease of 33% (post hoc: p < 0.03) and 27% (post hoc: p < 0.04) in the 190 and 290 treatments, respectively. Conversely, increased pCO₂ had no effect on the Chl a:C ratio.

### Maximum quantum yield and changes to PSII

The photosynthetic yield of *P. subcurvata* (F₀/F₉) showed a significant Fe effect (2-way ANOVA: p < 0.001; Fig 2B). At 190, F₀/F₉ decreased significantly by 21% in response to Fe deficiency (from 0.52 ± 0.01 to 0.41 ± 0.02 in the +Fe and Control, respectively, post hoc: p < 0.001), while no Fe effect was observed in 290. Interestingly, CO₂ enhancement differently affected the photosynthetic yield of the two Fe treatments. While increasing pCO₂ enhanced the F₀/F₉ in the Control treatment by 15% (from 0.41 ± 0.02 to 0.47 ± 0.01, post hoc: p = 0.005), it reduced F₀/F₉ in the +Fe treatments by 8% (from 0.52 ± 0.01 to 0.48 ± 0.01, post hoc: p < 0.04). Hence, there was a significant interactive effect of CO₂ and Fe availability on F₀/F₉ (2-way ANOVA: p = 0.002; Fig 2B).

The connectivity (P) was significantly affected by Fe deficiency (2-way ANOVA: p = 0.002; Table 4), with the Control treatment having an 11% smaller energy transfer between PSII units

### Table 4. Connectivity (P), cellular concentration of functional PSII reaction centers (RCII), light utilization efficiency at low irradiance (α), maximum cellular electron transport rate (cETRₘₐₓ) and minimum saturating irradiance (Iₛ) of *P. subcurvata* in the four treatments (+Fe 190, Control 190, +Fe 290 and Control 290) at the end of the experiment. The values represent the means ± SD (n = 3). Different letters indicate significant differences between treatments (p < 0.05).

| Parameter                  | *P. subcurvata* incubations |                     |                     |
|----------------------------|----------------------------|---------------------|---------------------|
|                            | +Fe | Control | +Fe | Control |
| P (rel. unit)               | 0.44 ± 0.01ᵃ | 0.39 ± 0.02ᵇ | 0.43 ± 0.01ᵃ | 0.40 ± 0.01ᵃᵇ |
| RCII (zmol cell⁻¹)          | 515 ± 58ᵃ | 525 ± 42ᵃ | 370 ± 38ᵇ | 519 ± 47ᵃ |
| α (amol e⁻ cell⁻¹ s⁻¹ μmol photons m⁻² s⁻¹) | 0.75 ± 0.13ᵃ | 0.97 ± 0.14ᵇ | 0.58 ± 0.08ᵃ | 0.82 ± 0.07ᵇ |
| cETRₘₐₓ (amol e⁻ cell⁻¹ s⁻²) | 119 ± 21ᵃ | 165 ± 26ᵇ | 85 ± 5ᵃ | 139 ± 19ᵇ |
| Iₛ (μmol photons m⁻² s⁻¹)   | 155 ± 9ᵃ | 171 ± 11ᵃᵇ | 143 ± 15ᵃ | 169 ± 9ᵇ |

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than the +Fe at 190 (post hoc: p = 0.002). In the 290 treatments, a similar, however, not significant, decreasing trend was seen. In contrast, no response of P to increased CO₂ was observed.

The functional absorption cross section of PSII (σPSII) showed a significant effect to Fe deficiency (2-way ANOVA: p < 0.001; Fig 2C). While σPSII increased by 26% with reduced Fe availability in 190 (from 2.47 ± 0.03 to 3.11 ± 0.18 nm⁻², respectively, post hoc: p < 0.001), this Fe effect was not seen in the 290 treatments. Furthermore, only in the Control treatments σPSII was reduced by 10% from 3.11 ± 0.21 to 2.79 ± 0.09 nm⁻² between 190 and 290, respectively (post hoc: p = 0.01). Moreover, there was a synergistic effect between Fe and CO₂ on σPSII (2-way ANOVA: p = 0.009; Fig 2C).

The cellular concentration of functional PSII reaction centers (RCII) was significantly altered by Fe deficiency (2-way ANOVA: p < 0.04; Table 4). This effect was only seen in 290, where RCII increased by 29% (post hoc: p < 0.02). Increasing CO₂ significantly reduced the RCII concentration (2-way ANOVA: p < 0.05), but only in the +Fe treatments (post hoc: p < 0.02).

Fe deficiency differently influenced the time constant for electron transport at the acceptor of PSII (τ) in the two CO₂ treatments. While lower Fe concentration reduced τ when grown at 190 μatm pCO₂ (post hoc: p < 0.001), it was enhanced at 290 μatm pCO₂ (post hoc: p = 0.006; Fig 2D). The effect of increased CO₂ on τ was significant (2-way ANOVA: p < 0.004). In the Control treatments, τ increased from 548 ± 21 to 659 ± 23 μs from 190 to 290 μatm pCO₂ (post hoc: p < 0.001) while it remained constant in the +Fe treatments. Hence, there was a strong interactive effect of Fe and CO₂ on τ apparent (2-way ANOVA: p < 0.001).

**PE-curve**

The cellular electron transport rates (cETR) of all treatments followed the shape of a typical PE-curve (Fig 3A). The light utilization efficiency of *P. subcurvata* at low irradiance (α) was significantly affected by Fe deficiency (2-way ANOVA: p = 0.005; Table 4), with α increasing by 29% at 190 (post hoc: p < 0.04) and by 41% at 290 (post hoc: p < 0.02). A CO₂ effect was also observed (2-way ANOVA: p = 0.02), where increased CO₂ reduced α, but due to large uncertainties, the individual post hoc tests of the +Fe and Control treatments were not significant. In response to Fe deficiency, cETRmax (Table 4 and Fig 3A) was significantly enhanced (2-way ANOVA: p < 0.006) by 39% at 190 and by 64% at 290 (both post hoc: p < 0.03). The increase in CO₂, however, did not lead to significant changes in cETRmax. The minimum saturating irradiance (Iₙ) displayed a significant Fe effect (2-way ANOVA: p < 0.02; Table 4), where Iₙ increased by 10% in the 290 treatments (post hoc: p < 0.04). Although not significant (p>0.05), in the 190 treatments a similar trend was observed. Iₙ remained unchanged by increasing CO₂ irrespective of Fe availability.

The non-photochemical quenching of all treatments was similarly low at low irradiances (Fig 3B). Exposed to irradiances higher than 350 μmol photons m⁻² s⁻¹, the NPQ in *P. subcurvata* increased nearly linearly and then leveled off between ~1.5 and 2.5 for all treatments. No Fe or CO₂ effect on NPQ was observed in any treatment.

**Discussion**

The ‘Iron Hypothesis’ suggests that the fertilization of the SO by increased dust deposition in glacial times promoted growth and productivity of phytoplankton. The biological pump in the SO was thus hypothesized to have reduced atmospheric pCO₂. In this study, we assessed the ecophysiological response of *P. subcurvata* simulating glacial and interglacial climate scenarios in terms of changes in Fe and CO₂ availability. It is important to note that while we manipulated two of the main environmental parameters, Fe concentrations and pCO₂, other
parameters (macronutrient concentrations, temperature etc.) likely also differed between the glacial and interglacial ocean. For this study, however, the focus was on the interactive effects of Fe and pCO$_2$.

**Glacial conditions favored POC production by *P. subcurvata***

Between 190 and 290 μatm pCO$_2$, no change in growth rate was observed in the +Fe treatments of *P. subcurvata* (Fig 1A). Previous laboratory studies with cultures of the same *P.*
Contrary to other studies, we did not observe a decrease in cell volume of _P. subcurvata_ levels [24, 49, 50]. Overall, we can conclude that glacial conditions simulated by a low pCO\(_2\) [51]. Similarly, growth remained unaffected in the temperate strain also reported no changes in growth rate between 180 and 390 ppm pCO\(_2\). Between 200 and 380 μm pCO\(_2\) with decreasing Fe availability (Table 2) [42, 82]. This may have been masked by the fact that enhanced between 220 and 400 ppm pCO\(_2\) subcurvata possessed highest photochemical fitness under simulated glacial conditions. With \(F_c/F_m\) was highest in the +Fe 190 treatment (Fig 2B), indicating that _P. subcurvata_ possessed highest photochemical fitness under simulated glacial conditions. With increasing pCO\(_2\), however, \(F_c/F_m\) declined in the diatom, but only when Fe was added (Fig 2B). Such a negative CO\(_2\) effect in Fe-enriched conditions was also observed in the Chl a content (Fig 2A) and the number of functional RCII (Table 4). Indeed, _P. subcurvata_ cells grown in the +Fe 290 treatment had a lower Chl a content compared to ones in the +Fe 190 treatment (Fig 2A), although the Chl a:C ratios were similar. Moreover, cellular BSi quotas and production remained constant with increasing pCO\(_2\) in the +Fe treatments (Fig 1D, Table 2) while a decline in POC and PON quotas as well as in POC and PON production rates (Fig 1B and 1C and Table 2) was found. Reducing both POC and PON quotas, _P. subcurvata_ was able to maintain a constant C:N ratio (Table 2) in response to increasing pCO\(_2\) under Fe-enriched conditions. Considering, however, that cETRs remained similar between 190 and 290 (Fig 3A, Table 4), a reduction in POC and PON contents indicates that the contribution of linear electron transport was reduced while cycling of electron via alternative pathways was required to avoid excess light energy. These physiological characteristics resemble those observed in various field incubation experiments under ocean acidification conditions and indicate that _P. subcurvata_ struggles when exposed to high pCO\(_2\) levels [24, 49, 50]. Overall, we can conclude that glacial conditions simulated by a low pCO\(_2\) of 190 μatm together with Fe enrichment was neither limiting growth nor POC production of _P. subcurvata_. On the contrary, these conditions were beneficial for biomass production and photochemical fitness of the diatom.

_P. subcurvata_ adjusted its physiological machinery to cope with low Fe supply

Contrary to other studies, we did not observe a decrease in cell volume of _P. subcurvata_ grown with decreasing Fe availability (Table 2) [42, 82]. This may have been masked by the fact that
the *P. subcurvata* strain used in our experiment was acclimated to low Fe conditions for a long time. Indeed, it exhibited large and elongated cells compared to the much shorter cells of the stock culture grown in the Fe-rich F2 medium (12 μM Fe), thus increasing their surface area-to-volume ratio. Furthermore, this strain was isolated from open ocean waters in the Atlantic sector of the SO. It is well known that oceanic diatoms acclimate to Fe limitation by increasing their surface area-to-volume ratio in order to maximize the number of transporter sites and nutrient uptake kinetics [83, 84].

Many studies reported a decrease in growth rate with decreasing Fe availability [8, 39, 40, 42, 44, 85–89]. Nonetheless, some of them also observed that particular oceanic diatoms grew at comparable rates under high and low Fe conditions [8, 86], as they have evolved acclimation strategies to reduce their Fe requirement. In our experiment, the growth rate of the oceanic *P. subcurvata* also displayed no difference between +Fe and Control conditions at the two pCO2 levels tested (Fig 1A) [42]. Suggested that the response of physiological and biochemical parameters to Fe reduction precedes changes in growth rate. This may explain why we did not see a decrease in growth rate here, despite observing typical responses to Fe-limiting conditions as substantial reductions in photochemical quantum efficiency (Fig 2B), connectivity (Table 4) and Chl *a* content (Fig 2A) accompanied by large functional absorption cross sections (Fig 2C) [9, 38, 40, 41, 85, 87, 88, 90, 91].

Under Fe deficiency, lowered *F*<sub>v</sub>/*F*<sub>m</sub> values indicate that the excitation energy was less efficiently transferred in the antennae, due to damaged and altered parts of the photosynthetic apparatus [41]. A decrease in *F*<sub>v</sub>/*F*<sub>m</sub> was commonly observed in cells grown in Fe-poor environments [9, 40, 41, 44, 87, 88] and, as expected, we observed this trend in the Control 190 treatment of *P. subcurvata* (Fig 2B). In line with the tested *P. subcurvata* here, oceanic *Pseudo-nitzschia* species usually decouple *F*<sub>v</sub>/*F*<sub>m</sub> and growth rate, reducing the former while maintaining the latter [8]. This decoupling was suggested to be due to either a low energy requirement of the diatom, or a compensating mechanism that generates reducing power, thus supporting rapid growth [8].

The decrease in *F*<sub>v</sub>/*F*<sub>m</sub> (Fig 2B) and lowered connectivity (P, Table 4) at low pCO2 in the low Fe *P. subcurvata* cells indicate that the transfer of excitation energy to the reaction centers was compromised [9]. Because Fe deficiency affects the synthesis and thus cellular content of Chl *a*, as seen in our data (Fig 2A), light harvesting may become more difficult for the cell. While [85] held lowered pigment concentration during Fe starvation responsible for a decline in photosynthesis, we did not observe reduced POC production rates (Fig 1B). Rather *P. subcurvata* compensated for a low Chl *a* content by increasing the functional absorption cross section of PSII (*σ*<sub>PSII</sub>), which is a measure of the target area of the light harvesting antenna (Fig 2C). In response to Fe deficiency this strategy can reduce the Fe demand and keep up the same capacity of the cell to absorb light [92]. Our results agree with literature showing an increase in *σ*<sub>PSII</sub> with Fe reduction [9, 38, 40, 41, 75, 87–89].

These photophysiological adjustments, however, did not prevent changes in light absorption completely, as shown by the strongly impacted light use capacities of Fe-limited *P. subcurvata* (Table 4). Higher *α* values were found under Fe deficiency for both 190 and 290 treatments, indicating that cells were able to respond better to lower irradiances than Fe-replete cells. Surprisingly, this effect was not always observed for *I*<sub>L</sub> values of *P. subcurvata*, which remained similar at 190 and was slightly higher at 290 (Table 4). Thus, while Fe deficiency at 290 resulted in a more efficient light utilization at lower irradiances (higher *α*), the cells required more light (higher *I*<sub>L</sub>) in order to cover their photosynthetic requirement [40]. In other studies, *I*<sub>L</sub> either decreased [40, 93] or remained unchanged [44, 85] under Fe reduction.
Even though POC-fixation remained constant under Fe deficiency (Fig 1B), cETR_{max} (Fig 3A, Table 4) and RCII concentration (only seen at 190 μatm pCO\textsubscript{2}) were enhanced (Table 4), indicating similar linear electron transport, but also cycling of electrons into alternative pathways such as cyclic electron flow within PSII [94] or Mehler reaction [95]. Considering, however, that the latter pathways are Fe-expensive, other pathways such as activity of a putative plastid plastoquinol terminal oxidase (PTOX) seem more plausible [96]. In support for this, [9] also observed constant C assimilation, but enhanced electron transport with Fe limitation in open ocean phytoplankton. Furthermore, a quicker turnover time at the acceptor side of PSII (τ) was found at 190 μatm pCO\textsubscript{2} in the Fe deficient P. subcurvata cells (Fig 2D), supporting PTOX activity, as previously observed for the Fe-limited Antarctic diatom Chaetoceros debilis [44]. Interestingly, this was not reflected in higher NPQ activities (Fig 3B).

At low pCO\textsubscript{2}, BSi quotas and production rates of P. subcurvata remained unaltered in response to Fe deficiency (Fig 1D, Table 2), as previously observed in Chaetoceros debilis [44], Corethron pennatum [97] and Chaetoceros dichaeta [98]. Considering the importance of Fe in C and N assimilation pathways, many studies reported a decrease in C and N under Fe deficiency [40, 42]. In [99], the C quota per cell volume ranged between 0.02 and 0.03 pg μm\textsuperscript{3} and was similar between Fe-replete and Fe-deficient treatments in the oceanic Pseudo-nitzschia fraudulenta, P. heimi, P. inflataula and P. turgidula, as well as in the coastal species P. multiseries and P. pseudodelicatissima. This matches with our results for the two tested pCO\textsubscript{2} levels (POC per cell volume at 190 +Fe: 0.041±0.002 pg μm\textsuperscript{3}, Control: 0.035±0.005 pg μm\textsuperscript{3} and at 290 +Fe: 0.029±0.005 pg μm\textsuperscript{3}, Control: 0.027±0.003 pg μm\textsuperscript{3}). The C:N ratio of diatoms was reported to increase [78], decrease [100] or remain unchanged [42, 44] with reduced Fe availability. We observed an increase in the C:N ratio in response to Fe deficiency at 190 μatm pCO\textsubscript{2} (Table 2). In this case, POC quotas remained constant, whereas PON cell quotas decreased with Fe deficiency (Table 2). Literature showed that Fe limitation can affect the supply of ‘new nitrogen’ to the cell as Fe is needed in some N-rich enzymes [101, 102]. [75] observed less abundant transcripts for nitrite reductase under Fe limiting conditions in Phaeocystis antarctica. Considering this, our reduced PON-fixation in P. subcurvata, allowing it to reduce its Fe demand. In order to keep up the same POC production, P. subcurvata needed to rely on alternative electron pathways such as cyclic electron flow as well as PTOX activity to prevent over-excitation.

**Increased pCO\textsubscript{2} weakened the effects of low Fe supply, but did not promote biomass build up**

Previous experiments with Pseudo-nitzschia demonstrated on the one hand, that the cell volume of P. pseudodelicatissima increased significantly as pCO\textsubscript{2} decreased, while, on the other hand, cell volume was found to decrease with decreasing Fe availability [42, 82]. In our experiments, the cell volume of P. subcurvata did not decrease with reduced Fe availability and increased pCO\textsubscript{2} (Table 2), potentially due to a counteracting effect of both factors together. Moreover, Fv/Fm decreased in response to Fe reduction at 190 (Fig 2B), while such Fe-dependent decrease in Fv/Fm was not observed at 290. This indicates that increasing pCO\textsubscript{2} had a positive effect on the maximum photochemical efficiency of low Fe P. subcurvata cells. A similar effect by high CO\textsubscript{2} concentration was also found for σPSII in Fe-deplete cells, being much smaller (Fig 2C). Apparently, these positive CO\textsubscript{2} effects weakened the strong Fe reduction effects previously observed at 190. Such positive response did, however, not translate into
more efficient energy transfer from photochemistry to biomass production. In fact, re-oxidation of the primary electron acceptor Q_a of low Fe cells was strongly compromised at 290 (Fig 2D). This was associated with reduced POC fixation and enhanced cETRs at 290 (Fig 3A, Table 4), and as a consequence, alternative electron acceptors were required. Due to a synergistic effect of reduced Fe availability and increased pCO₂, in our experiment we observed the highest BSi production in low Fe high pCO₂ conditions (Fig 1D). This increase in BSi production with reduced Fe concentrations at 290 hints towards stronger silicification and the production of thicker shells by *P. subcurvata* [104–106].

**Conclusion: Glacial vs. interglacial**

In our study, in a simulated Fe-fertilized glacial ocean (+Fe 190), *P. subcurvata* displayed similar growth rates as in interglacial ocean conditions (Control 290), despite lower Fe availability, hinting towards an efficient acclimation strategy to reduce the Fe requirement. Under glacial conditions, electrons were more efficiently channeled, leading to higher cellular POC and PON concentrations and production rates. In comparison, the interglacial conditions with higher pCO₂ and reduced Fe availability resulted in reduced POC buildup of the diatom. Thus, we observed that both higher Fe availability and lower CO₂ concentration as in the glacial ocean, promoted POC production by *P. subcurvata*. Assuming that *P. subcurvata* dominated phytoplankton blooms in the SO during glacial and interglacial times, we can conclude that *P. subcurvata* contributed more to primary production in the glacial than interglacial ocean. The higher POC production rates by the diatom under glacial conditions facilitated higher CO₂ uptake from the atmosphere and potentially higher C export. This matches the ‘Iron Hypothesis’ of [1], which states that in the last glacial maximum higher Fe input from dust fertilized the SO, thus stimulating higher primary production and reducing thereby the atmospheric CO₂ concentration. On the other hand, however, the thicker shells of *P. subcurvata* under the simulated interglacial conditions hint towards reduced grazing and thus its higher contribution to C export [107]. Biogeochemical cycles changed in the past and will change in response to future global climate change. Thus, understanding the dynamic interactions of the ocean’s biogeochemistry and phytoplankton is important in order to better simulate past and future climatic scenarios.

**Supporting information**

S1 File. Cellular trace metal quotas. Trace metal (TM) quotas without oxalate (Total TM content) and with oxalate wash (Intracellular TM content) determined at the end of the experiment in the four treatments of *P. subcurvata* (+Fe 190, Control 190, +Fe 290 and Control 290). The values represent the means ± SD (n = 3). Different letters indicate significant (p < 0.05) differences between treatments.

(DOCX)

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References

1. Martin JH. Glacial-interglacial CO$_2$ change: The Iron Hypothesis. Paleoceanography. 1990. https://doi.org/10.1029/PA005i001p00001

2. de Baar HJW, de Jong JTM, Bakker DCE, Löscher BM, Veth C, Bathmann U, et al. Importance of iron for plankton blooms and carbon dioxide drawdown in the Southern Ocean. Nature. 1995. https://doi.org/10.1038/373412a0

3. Boyd PW, Ellwood MJ. The biogeochemical cycle of iron in the ocean. Nature Geoscience. 2010. https://doi.org/10.1038/ngeo892 PMID: 20634995

4. Raven JA, Evans MCW, Korb RE. The role of trace metals in photosynthetic electron transport in O$_2$-evolving organisms. Photosynthesis Research. 1999. https://doi.org/10.1023/A:1006282714942

5. Martin JH, Coale KH, Johnson KS, Fitzwater SE, Gordon RM, Tanner SJ, et al. Testing the iron hypothesis in ecosystems of the equatorial Pacific Ocean. Nature. 1994. https://doi.org/10.1038/371123a0

6. Boyd PW, Watson AJ, Law CS, Abraham ER, Trull T, Murdoch R, et al. A mesoscale phytoplankton bloom in the polar Southern Ocean stimulated by iron fertilization. Nature. 2000. https://doi.org/10.1038/36037500 PMID: 11048709

7. de Baar HJW, Boyd PW, Coale KH, Landry MR, Tsuda A, Assmy P. Synthesis of iron fertilization experiments: From the Iron Age in the Age of Enlightenment. Journal of Geophysical Research: Oceans. 2005. https://doi.org/10.1029/2004JC002901

8. Marchetti A, Maldonado MT, Lane ES, Harrison PJ. Iron requirements of the pennate diatom _Pseudo-nitzschia_: Comparison of oceanic (high-nitrate, low-chlorophyll waters) and coastal species. Limnology and Oceanography. 2006. https://doi.org/10.4319/lo.2006.51.3.1530 PMID: 21258624

9. Schuback N, Schallenberg C, Duckham C, Maldonado MT, Tortell PD. Interacting effects of light and iron availability on the coupling of photosynthetic electron transport and CO2-assimilation in marine phytoplankton. PLOS ONE. 2015. https://doi.org/10.1371/journal.pone.0133235 PMID: 26171963

10. Takahashi T, Sutherland SC, Wanninkhof R, Sweeney C, Feely RA, Chipman DW. Climatological mean and decadal change in surface ocean pCO$_2$, and net sea-air CO$_2$ flux over the global oceans. Deep Sea Research Part II: Topical Studies in Oceanography. 2009. https://doi.org/10.1016/j.dsr2.2008.12.009

11. Hauck J, Lenton A, Langlais C, Matear R. The fate of carbon and nutrients exported out of the Southern Ocean. Global Biogeochemical Cycles. 2018. https://doi.org/10.1029/2018GB005977

12. Raven JA, Falkowski PG. Oceanic sinks for atmospheric CO$_2$. Plant, Cell & Environment. 1999. https://doi.org/10.1046/j.1365-3040.1999.00419.x

13. Gruber N, Gloor M, Mikoloff Fletcher SE, Doney SC, Dutkiewicz S, Follows MJ, et al. Oceanic sources, sinks, and transport of atmospheric CO$_2$. Global Biogeochemical Cycles. 2009. https://doi.org/10.1029/2008GB003349
14. Francois R, Altabat MA, Yu E-F, Sigman DM, Bacon MP, Frank M, et al. Contribution of Southern Ocean surface-water stratification to low atmospheric CO\textsubscript{2} concentrations during the last glacial period. Nature. 1997. https://doi.org/10.1038/40073

15. Sigman DM, Boyle EA. Glacial-interglacial variations in atmospheric carbon dioxide. Nature. 2000. https://doi.org/10.1038/35038000 PMID: 11057657

16. Stephens BB, Keeling RF. The influence of Antarctic sea ice on glacial-interglacial CO\textsubscript{2} variations. Nature. 2000. https://doi.org/10.1038/35004556 PMID: 10724166

17. Mayr C, Lücke A, Wagner S, Wissel H, Ohlendorf C, Haberzettl T, et al. Intensified Southern Hemisphere Westerlies regulated atmospheric CO\textsubscript{2} during the last deglaciation. Geology. 2013. https://doi.org/10.1130/G34335.1

18. Kohfeld KE, Quéré CL, Harrison SP, Anderson RF. Role of marine biology in glacial-interglacial CO\textsubscript{2} cycles. Science. 2005. https://doi.org/10.1126/science.1105375 PMID: 15802597

19. Martinez-Garcia A, Rosell-Mel A, Gersonde R, Masqu P, Gaspari V, et al. Links between iron supply, marine productivity, sea surface temperature, and CO\textsubscript{2} over the last 1.1 Ma. Paleoceanography. 2009. https://doi.org/10.1029/2008PA001657

20. Martinez-Garcia A, Sigman DM, Ren H, Anderson RF, Straub M, Hodell DA, et al. (2014). Iron fertilization of the subtropical ocean during the Last Ice Age. Science. 2014. https://doi.org/10.1126/science.1246848 PMID: 24653031

21. Lambert F, Tagliabue A, Shaffer G, Lamy F, Winckler G, Farias L, et al. Dust fluxes and iron fertilization in Holocene and Last Glacial Maximum climates. Geophys. Res. Lett. 2015. https://doi.org/10.1002/2015GL064250

22. Moore JK, Doney SC, Lindsay K, Mahowald N, Michaels AF. Nitrogen fixation amplifies the ocean biogeochemical response to decadal timescale variations in mineral dust deposition. Tellus B: Chemical and Physical Meteorology. 2006. https://doi.org/10.1111/j.1600-0889.2006.00209.x

23. Sunda W. Feedback interactions between trace metal nutrients and phytoplankton in the ocean. Frontiers in Microbiology. 2012. https://doi.org/10.3389/fmicb.2012.00204 PMID: 22701115

24. Trimborn S, Brenneis T, Hoppe CJM, Laglera LM, Norman L, Santos-Echeandía J, et al. Iron sources alter the response of Southern Ocean phytoplankton to ocean acidification. Mar Ecol Prog Ser. 2017; 578: 35–50.

25. Anderson RF, Barker S, Fleisher M, Gersonde R, Goldstein SL. Biological response to millennial variability of dust and nutrient supply in the Subantarctic South Atlantic Ocean. Philosophical Transactions of the Royal Society A: Mathematical, Physical and Engineering Sciences. 2014. https://doi.org/10.1098/rsta.2013.0054 PMID: 24891398

26. Kumar N, Anderson RF, Mortlock RA, Froelich PN, Kubik P, Dittrich-Hannen B, et al. Increased biological productivity and export production in the glacial Southern Ocean. Nature. 1995. https://doi.org/10.1038/378675a0

27. Albani S, Mahowald NM, Murphy LN, Raiswell R, Moore JK, Anderson RF, et al. Paleodust variability since the Last Glacial Maximum and implications for iron inputs to the ocean. Geophysical Research Letters. 2016. https://doi.org/10.1002/2016GL067911

28. Wagener T, Guieu C, Losno R, Bonnet S, Mahowald N. Revisiting atmospheric dust export to the Southern Hemisphere ocean: Biogeochemical implications. Global Biogeochemistry. 2008. https://doi.org/10.1029/2007GB002984

29. Arrigo KR, Robinson DH, Worthen DL, Dunbar RB, DiTulio GR, VanWoert M, et al. Phytoplankton Community Structure and the Drawdown of Nutrients and CO\textsubscript{2} in the Southern Ocean. Science. 1999. https://doi.org/10.1126/science.283.5400.365 PMID: 9888847

30. Nelson DM, Trügler P, Brzezinski MA, Leynaert A, Quagheber B. Production and dissolution of biogenic silica in the ocean: Revised global estimates, comparison with regional data and relationship to biogenic sedimentation. Global Biogeochemical Cycles. 1995. https://doi.org/10.1029/95GB01070

31. Trügler P, Nelson DM, Van Bennekum AJ, DeMaster DJ, Leynaert A, Quagheber B. (1995). The silica balance in the world ocean: A reestimate. Science. 1995. https://doi.org/10.1126/science.268.5209.375 PMID: 17746543

32. Diatoms Smetacek V. and the ocean carbon cycle. Protist. 1999. https://doi.org/10.1016/S1434-4610(99)70006-4 PMID: 10724516

33. Smetacek V. Seeing is believing: Diatoms and the ocean carbon cycle revisited. Protist. 2018. https://doi.org/10.1016/j.protis.2018.08.004 PMID: 30342384

34. Smetacek VS. Role of sinking in diatom life-history cycles: ecological, evolutionary and geological significance. Marine Biology. 1985. https://doi.org/10.1007/BF00392493
35. Buesseler K, Ball L, Andrews J, Cochran J, Hirschberg D, Bacon M, et al. Upper ocean export of particulate organic carbon and biogenic silica in the Southern Ocean along 170°W. Deep Sea Research Part II: Topical Studies in Oceanography. 2001. https://doi.org/10.1016/S0967-0645(01)00089-3

36. Hasle GR. *Nitzschia* and *Fragilariopsis* species studied in the light and electron microscopes: 1. Some marine species of the groups *Nitzschia* and *Lanceolatae*. Skrifter utgitt av Det Norske Videnskaps-Akademi i Oslo. 1964.

37. Smetacek V, Klaas C, Strass VH, Assmy P, Montresor M, Cisewski B. et al. Deep carbon export from a Southern Ocean iron-fertilized diatom bloom. Nature. 2012. https://doi.org/10.1038/nature11229 PMID: 22810695

38. Strzepek RF, Maldonado MT, Hunter KA, Frew RD, Boyd PW. Adaptive strategies by Southern Ocean phytoplankton to lessen iron limitation: Uptake of organically complexed iron and reduced cellular iron requirements. Limnology and Oceanography. 2011. https://doi.org/10.4319/lo.2011.56.6.2411 PMID: 28966398

39. Allen AE, LaRoche J, Maheswari U, Lommer M, Schauer N, Lopez PJ. Whole-cell response of the pennate diatom *Phaeodactylum tricornutum* to iron starvation. Proceedings of the National Academy of Sciences. 2008. https://doi.org/10.1073/pnas.0711370105 PMID: 18653757

40. Greene RM, Geider RJ, Falkowski PG. Effect of iron limitation on photosynthesis in a marine diatom. Limnology and Oceanography. 1991. https://doi.org/10.4319/lo.1991.36.8.1772

41. Greene RM, Geider RJ, Kolber Z, Falkowski PG. Iron-induced changes in light harvesting and photochemical energy conversion processes in eukaryotic marine algae. Plant Physiology. 1992. https://doi.org/10.1104/pp.100.2.565 PMID: 16653030

42. Marchetti A, Harrison PJ. Coupled changes in the cell morphology and elemental (C, N, and Si) composition of the pennate diatom *Pseudo-nitzschia* due to iron deficiency. Limnology and Oceanography. 2007. https://doi.org/10.4319/lo.2007.52.5.2270

43. Van Leeuwe M, Visser RJW, Stefels J. The pigment composition of *Phaeocystis antarctica* (Haptophyceae) under various conditions of light, temperature, salinity, and iron. Journal of Phycology. 2014; 50 (6): 1070–1080. https://doi.org/10.1111/j.py.12238 PMID: 26988788

44. Trimborn S, Thorns S, Bischof K, Beszteri S. Susceptibility of two Southern Ocean phytoplankton key species to iron limitation and high light. Frontiers in Marine Science. 2019. https://doi.org/10.3389/fmars.2019.00167

45. Reinfelder JR. Carbon dioxide regulation of nitrogen and phosphorus in four species of marine phytoplankton. Marine Ecology Progress Series. 2012. https://doi.org/10.3354/meps09905

46. Zhu Z, Qu P, Gale J, Fu F, Hutchins DA. Individual and interactive effects of warming and CO$_2$ on *Pseudo-nitzschia subcurvata* and *Phaeocystis antarctica*, two dominant phytoplankton from the Ross Sea, Antarctica. Biogeosciences. 2017. https://doi.org/10.5194/bg-14-5281-2017

47. Hoogstraten A, Timmermans KR, de Baar HJW. Morphological and physiological effects in *Proboscia alata* (bacillariophyceae) grown under different light and CO$_2$ conditions of the modern southern ocean.1. Journal of Phycology.2012. https://doi.org/10.1111/j.1529-8817.2012.01148.x PMID: 27011071

48. Sun J, Hutchins DA, Feng Y, Seubert EL, Caron DA, Fu F-X. Effects of changing pCO$_2$ and phosphate availability on domoic acid production and physiology of the marine harmful bloom diatom *Pseudo-nitzschia multiseries*. Limnology and Oceanography. 2011. https://doi.org/10.4319/lo.2011.56.6.2411 PMID: 28966398

49. Tortell PD, Payne CD, Li Y, Trimborn S, Rost B, Smith WO, et al. CO$_2$ sensitivity of Southern Ocean phytoplankton. Geophysical Research Letters. 2008. https://doi.org/10.1029/2008gl035090 PMID: 24347740

50. Hoppe CJM, Hassler CS, Payne CD, Tortell PD, Rost B, Trimborn S. Iron limitation modulates ocean acidification effects on Southern Ocean phytoplankton communities. PLOS ONE. 2013. https://doi.org/10.1371/journal.pone.0079890 PMID: 24278207

51. Trimborn S, Brenneis T, Sweet E, Rost B. Sensitivity of Antarctic phytoplankton species to ocean acidification: Growth, carbon acquisition, and species interaction. Limnology and Oceanography. 2013. https://doi.org/10.4319/lo.2013.58.3.0997

52. Feng Y, Hare C, Rose J, Handy S, DiTulio G, Lee P, et al. (2010). Interactive effects of iron, irradiance and CO$_2$ on Ross Sea phytoplankton. Deep Sea Research Part I: Oceanographic Research Papers. 2010. https://doi.org/10.1016/j.dsr.2009.10.013

53. Muglia J, Skinner LC, Schmittner A. Weak overturning circulation and high Southern Ocean nutrient utilization maximized glacial ocean carbon. Earth and Planetary Science Letters. 2018. https://doi.org/10.1016/j.epsl.2018.05.038
54. Guillard RRL, Ryther JH. Studies of marine planktonic diatoms: I. *cyclotella nana* Hustedt, and *detonula confervacea* (Cleve) Gran. Canadian Journal of Microbiology. 1962. https://doi.org/10.1139/m62-029 PMID: 13902807

55. Gerringa LJA, de Baar HJW, Timmermans KR. A comparison of iron limitation of phytoplankton in natural oceanic waters and laboratory media conditioned with EDTA. Marine Chemistry. 2000. https://doi.org/10.1016/S0304-4203(99)00092-4

56. Cutter G, Casciotti K, Croot P, Geibert W, Heimburger L.-E., Lohan M, et al. Sampling and sample-handling protocols for GEOTRACES cruises. Version 3, August 2017. Toulouse, France, GEOTRACES International Project Office, 2017. pp 139. https://doi.org/10.25607/OPB-2

57. Hather CE, Haley B, Stichel T, Grasse P, Zieringer M, Frank, M. Online preconcentration ICP-MS analysis of rare earth elements in seawater. Geochimistry, Geophysics, Geosystems. 2012. https://doi.org/10.1029/2011GC003907

58. Stoll MHC, Bakker K, Nobbe GH, Haese RR. Continuous-flow analysis of dissolved inorganic carbon content in seawater. Analytical Chemistry. 2001. https://doi.org/10.1021/ac010303r PMID: 11569799

59. Brewer PG, Bradshaw AL, Williams RT. Measurements of total carbon dioxide and alkalinity in the North Atlantic Ocean in 1981. In: Trabalka JR, Reichle DE, editors. The changing carbon cycle: A global analysis. New York: Springer; 1986. pp. 348–370. https://doi.org/10.1007/978-1-4757-1915-4-18

60. Gran G. Determination of the equivalence point in potentiometric titrations. Part ii. Analyst. 1952. https://doi.org/10.1039/AN9527700661

61. van Heuven S, Pierrot D, Rae JWB, Lewis E, Wallace DWR. MATLAB Program Developed for CO2 System Calculations. ORNL/CDIAC-105b. Carbon Dioxide Information Analysis Center, Oak Ridge National Laboratory, U.S. Department of Energy, Oak Ridge, Tennessee. https://doi.org/10.3334/CDIAC/co2sys-matlab-v1.1

62. Mehrbach C, Culberson CH, Hawley JE, Pytkowicz RM. Measurement of the apparent dissociation constants of carbonic acid in seawater at atmospheric pressure. Limnology and Oceanography. 1973. https://doi.org/10.4319/lo.1973.18.6.0897

63. Dickson AG, Millero FJ. A comparison of the equilibrium constants for the dissociation of carbonic acid in seawater media. Deep Sea Research Part A. Oceanographic Research Papers. 1987. https://doi.org/10.1016/0198-0149(87)90021-5

64. Utermöhl H. Methods of collecting plankton for various purposes are discussed. SIL Communications, 1953–1996. 1958. https://doi.org/10.1080/05384680.1958.11904091

65. Hillebrand H, Drselen C-D, Kirschtel D, Pollingher U, Zohary T. Biovolume calculation for pelagic and benthic microalgae. Journal of Phycology. 1999. https://doi.org/10.1046/j.1529-8817.1999.3520403.x

66. Brzezinski MA, Nelson DM. The annual silica cycle in the Sargasso Sea near Bermuda. Deep Sea Research Part I: Oceanographic Research Papers. 1995. https://doi.org/10.1016/0967-0637(95)93592-3

67. Koroleff F. (1983). Determination of silicon. In: Grasshoff K, Kremling M, editors. Methods of Seawater Analysis. Weinheim: Wiley-VCH; 1983. pp. 174–183.

68. Wright SW. Improved HPLC method for the analysis of chlorophylls and carotenoids from marine phytoplankton. Mar. Ecol. Prog. Ser. 1991. https://doi.org/10.3354/meps07718

69. Oxborough K, Moore CM, Suggett DJ, Lawson T, Chan HG, Geider RJ. Direct estimation of functional PSII reaction center concentration and PSII electron flux on a volume basis: a new approach to the analysis of Fast Repetition Rate fluorometry (FRRf) data. Limnology and Oceanography: Methods. 2012. https://doi.org/10.4319/lom.2012.10.142

70. Trimborn S, Thoms S, Petrou K, Kranz SA, Rost B. Photophysiological responses of Southern Ocean phytoplankton to changes in CO2 concentrations: Short-term versus acclimation effects. Journal of Experimental Marine Biology and Ecology. 2014. https://doi.org/10.1016/j.jembe.2013.11.001

71. Kolber ZS, Prasil O, Falkowski PG. Measurements of variable chlorophyll fluorescence using fast repetition rate techniques: defining methodology and experimental protocols. Biochimica et Biophysica Acta (BBA)—Bioenergetics. 1999. https://doi.org/10.1016/s0005-2728(98)00135-2 PMID: 9784616

72. Genty B, Briantais J-M, Baker NR. The relationship between the quantum yield of photosynthetic electron transport and quenching of chlorophyll fluorescence. Biochimica et Biophysica Acta (BBA)—General Subjects. 1989. https://doi.org/10.1016/S0304-4165(89)80016-9

73. Suggett DJ, MacIntyre HL, Geider RJ. Evaluation of biophysical and optical determinations of light absorption by photosystem II in phytoplankton. Limnology and Oceanography: Methods. 2004. https://doi.org/10.4319/lom.2004.2.316
74. Suggett DJ, Moore CM, Hickman AE, Geider RJ. (2009). Interpretation of fast repetition rate (FRR) fluorescence: signatures of phytoplankton community structure versus physiological state. Marine Ecology Progress Series. 2009. https://doi.org/10.3354/meps07830

75. Koch F, Beszteri S, Harms L, Trimborn S. The impacts of iron limitation and ocean acidification on the cellular stoichiometry, photophysiology, and transcriptome of Phaeocystis antarctica. Limnology and Oceanography. 2018. https://doi.org/10.1002/lno.11045

76. Ralph PJ, Gademann R. Rapid light curves: A powerful tool to assess photosynthetic activity. Aquatic Botany. 2005. https://doi.org/10.1016/j.aquabot.2005.02.006

77. Bilger W, Bjorkman O. Temperature dependence of violaxanthin de-epoxidation and non-photochemical fluorescence quenching in intact leaves of Gossypium hirsutum L. and Malva parviflora L. Planta. 1991. https://doi.org/10.1007/BF00197951 PMID: 24194074

78. Sugie K, Yoshihara T. Effects of pCO2 and iron on the elemental composition and cell geometry of the marine diatom Pseudo-nitzschia pseudodelicatissima (Bacillariophyceae). Journal of Phycology. 2013. https://doi.org/10.1111/j.1529-8817.2011.00612.x

79. King AL, Jenkins B, Wallace J, Liu Y, Wikfors G, Milke L, et al. Effects of CO2 on growth rate, C:N:P, and fatty acid composition of seven marine phytoplankton species. Mar. Ecol. Prog. Ser. 2015. https://doi.org/10.3354/meps11458

80. Boelen P, van de Poll WH, van der Strate HJ, Neven IA, Beardall J, Buma AG. Neither elevated nor reduced CO2 affects the photophysiological performance of the marine Antarctic diatom Chaetoceros brevis. Journal of Experimental Marine Biology and Ecology. 2011. https://doi.org/10.1016/j.jembe.2011.06.012

81. Trimborn S, Lundholm N, Thoms S, Richter K-U, Krock B, Hansen PJ, et al. Inorganic carbon acquisition in potentially toxic and non-toxic diatoms: the effect of pH-induced changes in seawater carbonate chemistry. Physiologia Plantarum. 2008. https://doi.org/10.1111/j.1399-3047.2007.01038.x PMID: 18405335

82. Sugie K, Kuma K. Resting spore formation in the marine diatom Thalassiosira nordenskioldii under iron- and nitrogen-limited conditions. Journal of Plankton Research. 2008. https://doi.org/10.1093/plankt/fbn080

83. Hudson RJM, Morel FMM. Iron transport in marine phytoplankton: Kinetics of cellular and medium coordination reactions. Limnology and Oceanography. 1990. https://doi.org/10.4319/lo.1990.35.6.1343 PMID: 11538698

84. Sunda WG, Huntsman SA. Iron uptake and growth limitation in oceanic and coastal phytoplankton. Marine Chemistry. 1995. https://doi.org/10.1016/0304-4203(95)00035-P

85. Davey M., Geider RJ. Impact of iron limitation on the photosynthetic apparatus of the diatom Chaetoceros muelleri (Bacillariophyceae). Journal of Phycology. 2001. https://doi.org/10.1046/j.1529-8817.2001.99169.x

86. Strzepek RF, Harrison PJ. Photosynthetic architecture differs in coastal and oceanic diatoms. Nature. 2004. https://doi.org/10.1038/nature02954 PMID: 15470428

87. Strzepek RF, Hunter KA, Frew RD, Harrison PJ, Boyd PW. Iron-light interactions differ in Southern Ocean phytoplankton. Limnology and Oceanography.2012. https://doi.org/10.4319/lo.2012.57.4.1182

88. Petrou K, Trimborn S, Rost B, Ralph PJ, Hassler CS. The impact of iron limitation on the physiology of the Antarctic diatom Chaetoceros simplex. Marine Biology. 2014. https://doi.org/10.1007/s00227-014-2392-z PMID: 24719494

89. Koch F, Trimborn S. Limitation by Fe, Zn, Co and B results in similar physiological responses in two Antarctic phytoplankton species. Front. Mar. Sci. 2019. https://doi.org/10.3389/fmars.2019.00514

90. Trimborn S, Hoppe CJM, Taylor BB, Bracher A, Hassler C. Physiological characteristics of open ocean and coastal phytoplankton communities of Western Antarctic Peninsula and Drake Passage waters. Deep Sea Research Part I: Oceanographic Research Papers. 2015. https://doi.org/10.1016/j.dsr.2014.12.010

91. Strzepek RF, Boyd PW, Sunda WG. Photosynthetic adaptation to low iron, light, and temperature in Southern Ocean phytoplankton. Proceedings of the National Academy of Sciences of the United States of America. 2019. https://doi.org/10.1073/pnas.1810886116 PMID: 30787187

92. Ryan-Keogh TJ, Macey AI, Cockshutt AM, Moore CM, Bibby TS. The cyanobacterial chlorophyll-binding-protein isf acts to increase the in vivo effective absorption cross-section of psi under iron limitation1. Journal of Phycology. 2012. https://doi.org/10.1111/j.1529-8817.2011.01092.x PMID: 27009659

93. McKay RML, Geider RJ, LaRoche J. Physiological and biochemical response of the photosynthetic apparatus of two marine diatoms to Fe stress. Plant Physiology. 1997. https://doi.org/10.1104/pp.114.2.615 PMID: 12223732
94. Prasıl O, Kolber Z, Berry JA, Falkowski PG. Cyclic electron flow around Photosystem II in vivo. Photosynthesis Research. 1996. https://doi.org/10.1007/BF00029472 PMID: 24271480

95. Mehler AH. Studies on reactions of illuminated chloroplasts: I. mechanism of the reduction of oxygen and other hill reagents. Archives of Biochemistry and Biophysics. 1951. https://doi.org/10.1016/0003-9861(51)90082-3 PMID: 14857775

96. Mackey KRM, Paytan A, Grossman AR, Bailey S. A photosynthetic strategy for coping in a high-light, low-nutrient environment. Limnology and Oceanography. 2008. https://doi.org/10.4319/lo.2008.53.3.0900

97. Timmermans KR, van der Wagt B, de Baar HJW. Growth rates, half-saturation constants, and silicate, nitrate, and phosphate depletion in relation to iron availability of four large, open-ocean diatoms from the Southern Ocean. Limnology and Oceanography. 2004. https://doi.org/10.4319/lo.2004.49.6.2141

98. Hoffmann L, Peek I, Lochte K. Effects of iron on the elemental stoichiometry during EIFEX and in the diatoms Fragilariopsis kerguelensis and Chaetoceros dichaeta. Biogeosciences (BG) 2007. https://doi.org/10.5194/bg-4-569-2007

99. Marchetti A. Ecophysiological aspects of iron nutrition and domoic acid production in oceanic and coastal diatoms of the genus Pseudo-nitzschia. PhD thesis, University of British Columbia. 2005.

100. Bucciarelli E, Pondaven P, Sarthou G. Effects of an iron-light colimitation on the elemental composition (Si, C, N) of the marine diatoms Thalassiosira oceanica and Ditylum brightwellii. Biogeosciences. 2010. https://hal.univ-brest.fr/hal-00472043.

101. Morel FMM, Hudson RJM, Price NM. Limitation of productivity by trace metals in the sea. Limnology and Oceanography. 1991. https://doi.org/10.4319/lo.1991.36.8.1742

102. Milligan AJ, Harrison PJ. Effects of non-steady-state iron limitation on nitrogen assimilatory enzymes in the marine diatom Thalassiosira weissflogii (Bacillariophyceae). Journal of Phycology. 2000. https://doi.org/10.1046/j.1529-8817.2000.99013.x

103. Nunn BL, Faux JF, Hippmann AA, Maldonado MT, Harvey HR, Goodlett DR, et al. Diatom proteomics reveals unique acclimation strategies to mitigate Fe limitation. PLOS ONE. 2013. https://doi.org/10.1371/journal.pone.0075653 PMID: 24146769

104. Hutchins DA, Bruland KW. Iron-limited diatom growth and Si:N uptake ratios in a coastal upwelling regime. Nature. 1998. https://doi.org/10.1038/31203

105. Takeda S. Influence of iron availability on nutrient consumption ratio of diatoms in oceanic waters. Nature. 1998. https://doi.org/10.1038/31674

106. Boyle E. Pumping iron makes thinner diatoms. Nature. 1998. https://doi.org/10.1038/31585

107. Hamm CE, Merkel R, Springer O, Jurkojc P, Maier C, Prechtel K, et al. Architecture and material properties of diatom shells provide effective mechanical protection. Nature. 2003. https://doi.org/10.1038/nature01416 PMID: 12594512