Eco-Evolutionary Interaction in Competing Phytoplankton: Nutrient Driven Genotype Sorting Likely Explains Dominance Shift and Species Responses to CO₂

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How ecological and evolutionary processes interact and together determine species and community responses to climate change is poorly understood. We studied long-term dynamics (over approximately 200 asexual generations) in two phytoplankton species, a coccolithophore (Emiliania huxleyi), and a diatom (Chaetoceros affinis), to increased CO₂ growing alone, or competing with one another in co-occurrence. To allow for rapid evolutionary responses, the experiment started with a standing genetic variation of nine genotypes in each of the species. Under co-occurrence of both species, we observed a dominance shift from C. affinis to E. huxleyi after about 120 generations in both CO₂ treatments, but more pronounced under high CO₂. Associated with this shift, we only found weak adaptation to high CO₂ in the diatom and none in the coccolithophore in terms of species’ growth rates. In addition, no adaptation to interspecific competition could be observed by comparing the single to the two-species treatments in reciprocal assays, regardless of the CO₂ treatment. Nevertheless, highly reproducible genotype sorting left only one genotype remaining for each of the species among all treatments. This strong evolutionary selection coincided with the dominance shift from C. affinis to E. huxleyi. Since all other conditions were kept constant over time, the most parsimonious explanation for the dominance shift is that the strong evolutionary selection was driven by the experimental nutrient conditions, and in turn potentially altered competitive ability of the two species. Thus, observed changes in the simplest possible two-species phytoplankton “community” demonstrated that eco-evolutionary interactions can be critical for predicting community responses to climate change in rapidly dividing organisms such as phytoplankton.

Keywords: eco-evolutionary interaction, CO₂, ocean acidification, competition, phytoplankton, C. affinis, E. huxleyi, species interaction
INTRODUCTION

Recent studies have repeatedly shown that ecological and evolutionary processes happen on similar time scales (Carroll et al., 2007; Reznick, 2013). Understanding how both processes interact has sparked the emergence of the new field of eco-evolutionary dynamics (Fussmann et al., 2007) aiming to understand how rapid evolutionary adaptation influences ecological processes, and vice versa (Hendry, 2016). Strong ecoevolutionary coupling is particularly expected in phytoplankton species, photoautotrophic aquatic microbes that are abundant in freshwater and marine ecosystems. With one cell division per day, many species lend themselves for experimental studies allowing for appropriate replication and hundreds of generations of evolutionary change (Reusch and Boyd, 2013). Moreover, their ecological importance cannot be overemphasized, as marine phytoplankton species are responsible for half of all global photosynthesis (Falkowski et al., 2008).

Here we studied eco-evolutionary dynamics in response to increased seawater CO$_2$ concentration, one prominent aspect of ocean global change (Doney et al., 2009). As primary producers, all phytoplankton species depend on the availability of inorganic carbon. On time scales too short for evolutionary adaptation, non-calcifying species mostly respond by enhanced growth (Schaum et al., 2012; Li et al., 2017) leading to higher abundances (Sommer et al., 2015; “ecological winners,” e.g., diatoms), while most calcifying species react negatively (“ecological losers,” e.g., coccolithophores) to CO$_2$ enrichment (Riebesell, 2004; Doney et al., 2009). Nested within such broad functional categories of sensitivity is pronounced interspecific variation (Schaum et al., 2012; Hattich et al., 2017; Des Roches et al., 2018), on which selection can operate (Lohbeck et al., 2012).

Within phytoplankton, competition for abiotic resources is ubiquitous (Tilman, 1977). In recent theoretical studies (de Mazancourt et al., 2008; Lancaster et al., 2017) and a long term bacterial community experiment (Lawrence et al., 2012), competition was shown to constitute one major biotic driver of adaptation. At the same time experimental evolution studies using phytoplankton and addressing the presence and absence of competition as a selection factor are largely absent (Scheinin et al., 2015).

Here, we set out to study how the evolutionary response to increased CO$_2$ of two bloom forming and geographically co-occurring phytoplankton species, Emiliania huxleyi (a coccolithophore) and Chaetoceros affinis (a diatom), was altered by competition and assessed putative effects on ecological dynamics. Studying such eco-evolutionary processes requires that at least two species can be kept over long-term (i.e., more than 10 s of generations such that evolutionary change can happen) in co-occurring experimental settings, delaying, or preventing Gause’s principle of competitive exclusion (Hardin, 1960; Tilman, 1977). Hence, we established long-term coexistence in semi-continuous batch cycles by taking advantage of the species’ different nutrient uptake related strategies (Sommer, 1984; Riebesell, 2004; Doney et al., 2009). Diatoms have high nutrient uptake rates and are consequently characterized by high maximum growth rates (Litchman et al., 2007). At the same time, diatoms have high half saturation constants and low affinity for nutrient uptake (Litchman et al., 2007). As such they represent the “velocity-adapted” species (Sommer, 1984) and thrive in nutrient replete and fluctuating conditions, being able to rapidly monopolize nutrients. Coccolithophores in contrast, have lower maximum nutrient uptake rates. Hence, after a nutrient pulse they initially lose out against diatoms. Their lower half-saturation constants and higher affinity, however, make them the better competitors under nutrient poor conditions thriving at later successional stages (Sommer, 1984). In this system, we ran the experiment for ca. 200 asexual generations under fully factorial selection conditions of adaptation to ambient and increased CO$_2$, in combination with and without competition, i.e., the co-occurrence of the respective other species (Figure 1A). To allow for rapid evolutionary adaptation via genotype sorting (Lohbeck et al., 2012), populations of both species were assembled using nine genotypes in equal frequencies (Figure 1A) that could be traced via microsatellite genotyping. Reciprocal adaptation assays were conducted to test for adaptation of both species to enhanced CO$_2$ and the presence of the respective other species in the first and second half of the experiment (after ca. 50 and 200 generations; Figure 1B).

We hypothesize that on the short term, species contribution to the “community” would be diverging between CO$_2$ treatments due to varying CO$_2$ responses of the target species. Specifically, the relative contribution of E. huxleyi should be lower in the high CO$_2$ compared to the ambient treatment due to potential negative or neutral effects of CO$_2$. In contrast, the relative contribution of C. affinis should be higher in high CO$_2$ treatment due to a potential fertilizing effect of increased CO$_2$ in combination with the effect of CO$_2$ on E. huxleyi. In the long-term, we hypothesize that evolutionary dynamics (i.e., genotype sorting and eventually adaptation) diverge between both target species. Specifically, the potentially negatively affected coccolithophore should be selected toward more stress tolerance, while the possible fertilizing effect on the diatom could lead to selection for the diatom’s potential to efficiently utilize increased levels of CO$_2$ and in turn achieve higher growth rates. Moreover, any evolutionary change can feed back onto ecological processes by altering the abundance and thus contribution of the species to the community over a longer time (i.e., 50–200 generations). Overall, we hypothesize that the presence of the competitor can alter any evolutionary (genotype sorting to adaptive) response. For example, for E. huxleyi, the presence of the diatom assimilating CO$_2$ could alleviate CO$_2$ stress, and thus reduce selection toward tolerance, slowing down its adaptation in co-culture.

MATERIALS AND METHODS

We exposed two important phytoplankton species; the coccolithophore E. huxleyi and the diatom C. affinis to increased CO$_2$ (ambient, high) either alone or in co-culture (mono, mix) with the respective other species over 288 days.
FIGURE 1 | Set up of Selection and Assay experiments: Here the schematic setup of the selection phase of the experiment is depicted in the top row (A). The colors show the CO₂ treatments, whereas the frames of the species show mono or mix treatment. All species populations were started with 9 genotypes each. Each treatment consisted of 5 replicates. In the reciprocal assays, all cultures of both species were exposed to all possible treatments (four rows below) thus resulting in a fully crossed assay set-up (B). The 16 resulting assay treatments are shown for each species. The star shows which of the species was the focal species in the mix assay treatments.

(=36 batch cycles of 8 days), representing approximately 200 asexual generations. We followed absolute species biovolume at the end of each batch cycle over time to understand how the species respond to the different treatments. Each species started as a mixture of nine different genotypes to allow for rapid adaptation via genotype selection. All genotypes used were isolates originating from one geographical region (Gran Canary, 27°59′N 15°22′W, isolated 2012 and 2015) and are deposited in the culture collection in Roscoff (Table 1). In previous studies we have shown that the genotypes differed in growth, and to some extent in their response to CO₂ (Hattich et al., 2017) indicating ecological variability and as such standing genetic variation. After 64 days (ca. 50 generations) and 288 days (ca. 200 generations), we conducted reciprocal adaptation experiments to test for adaptation to CO₂ with and without competition. Ideally, to fully disentangle selection to inter- and intraspecific competition and the intended abiotic treatment single genotype cultures under both CO₂ conditions would be required. However, this was technically not feasible for the phytoplankton species in this study as it would have required to handle another 180 bottle’s each batch cycle.

**Experimental Set up and Culturing Conditions**

The selection treatments were set-up in a fully orthogonal way with the single factors CO₂ (two levels, ambient and high) and co-occurrence or absence of a 2nd species (Mono and Mix) resulting in 4 treatment combinations per species (Figure 1A). Five replicate cultures of each of the four treatment combinations were kept for 9 months (36 batch cycles, resulting in 210–230 generations in *E. huxleyi* and 195–210 generations in *C. affinis*) in semi-continuous batch cultures. However, after ca. 28–30 batch cycles one replicate culture per treatment combination was lost in *E. huxleyi* and one in 3 out of 4 treatment combinations in *C. affinis.*
The ambient and high CO$_2$-treatments were manipulated by aerating the artificial-seawater [35 PSU salinity; after (Kester et al., 1967)] for 24 h with ambient or CO$_2$-enriched air (400 and 1250 ppm, respectively) prior to two batch cycles. The dissolved inorganic carbon (Hansen et al., 2013) varied between 2052.39 ± 34.49 and 2202 ± 8.19 µmol × kg$^{-1}$ with a total alkalinity [following (Matthiessen et al., 2012)] of 2272.56 ± 11.24 and 2273.12 ± 11.43 µmol × kg$^{-1}$ for ambient and high CO$_2$, respectively. Nutrients were added to the final concentrations of 19.61 ± 0.99 µmol × L$^{-1}$ nitrate, 0.99 ± 0.02 µmol × L$^{-1}$ phosphate, and 4.34 ± 0.14 µmol × L$^{-1}$ silicate. We chose these comparably (to other long-term selection experiments) low nutrient concentrations to mirror natural conditions in the ocean. In addition, to simulate a bloom situation with resource competition under depleted nutrients (Paulton, 1991), we ran the study as semi-continuous batch cultures with both species reaching the stationary phase (Supplementary Figure S1 growth graphs of the different species; Supplementary Figure S2). Here the natural nutrient concentrations allowed this in a feasible time frame of 8 days per batch cycle. At the onset of the experiment each experimental unit (500 mL Nalgene® bottle) was inoculated with an initial total biovolume of 8280 µm$^3$ × mL$^{-1}$ of non-axenic, exponentially growing cultures, resulting in an inoculation of the communities with both species (mix-culture) in a 1:1 ratio of E. huxleyi and C. affinis (4140 µm$^3$ × mL$^{-1}$ each). Genotypes in each species were assembled in equal abundance for the nine genotypes of each species. For every following batch cycle 8280 µm$^3$ × mL$^{-1}$ of each replicate was transferred into replete media (prepared as described above). In order to calculate the contribution of each species to the total biovolume/mL and the amount needed to transfer to the next batch cycle, cell numbers, and cell volume (after Hillebrand et al., 2002) were determined using microscopy (Zeiss Axiovert Observer) at the end of each batch cycle. The experiment was carried out under constant rotation (0.75 min$^{-1}$) at ~22°C and a 17:7 day:night cycle reaching a maximum light intensity of 350 µmol × m$^{-2}$ × s$^{-1}$ 3 h after dusk and dawn.

### Frequency Assessments of Genotypes

In order to follow genotype frequencies, we re-isolated cells of both species after 8, 32, 64, and 288 days. Genotypes of both species could be unambiguously identified via microsatellite genotyping [see (Hattich et al., 2017), for detailed information on E. huxleyi and for C. affinis genotype identification see Supplementary Material]. For the quantification of E. huxleyi and C. affinis genotypes a maximum of 20 cells per culture were re-isolated by dilution in 48 well plates. This provided a theoretical detection limit of 5% difference between the contributions of the genotypes to both species’ populations. A lower detection limit could in theory have been achieved via isolation and identification of more cells but was not the goal of this study. Details on the resolation and quantification via microsatellite analysis are given in Supplementary Material. For the genotype composition of C. affinis we only had enough data for analysis two time points in the mixed cultures.

### Reciprocal Adaptation Assays

We carried out reciprocal adaptation assays to test for adaptation in both species to enhanced CO$_2$, the competition of the respective other species, and both factors in combination (Figure 1B). The assays on day 64 and 288 assessed how adaptation played out over time. Those adaptation assays compared evolved populations in control and new environments rather than evolved and ancestral populations because methods such as cryopreservation are not readily available for our target study species (Collins, 2011b). Specifically, every evolved culture was tested in all four treatment combinations in a full factorial way. For example, E. huxleyi that were long-term treated with high CO$_2$ conditions (treatment combination Mono High), were in the assay exposed to both ambient and high CO$_2$ concentrations and then also to the co-occurring diatom (already long-term exposed to ambient and high CO$_2$ competition to avoid confounding responses) in ambient and high CO$_2$ concentrations. Thus, each treatment combination from the selection leads to 4 assay treatment combinations. This resulted in a total of 16 assay treatment combinations per species (Figure 1B; for the details of the reciprocal assay set up see Supplementary Material). Using a full factorial adaptation assay allowed us to test on the one hand for the adaptation of the single factor CO$_2$ [i.e., in the statistical analysis, this could be identified as significant interactions of the respective selection and assay treatment factors (e.g., selection CO$_2$x assay CO$_2$)]. On the other hand, we were able to additionally test for the adaption to increased CO$_2$ in combination with competition, but then also how competition itself affected adaptive responses. This allowed disentangling the single and combined treatment factors at the same time. Here we focused on growth rates because this response is directly related to Darwinian fitness in an asexual population (Elena and Lenski, 2003) and is independent of nutrient availability due to the presence of another species (Tilman, 1977). The cultures for the assay treatments were inoculated in the same way as the long-term experiment, i.e., with a biovolume of 8280 µm$^3$ × mL$^{-1}$ (see Supplementary Material for details) in 500 ml Nalgene® bottles. After one batch
cycle of acclimation to assay treatments the assay responses were measured. We took daily samples for microscopy to determine growth rates over the course of the assay batch cycle. The growth rates (μ) were determined using a linear-fit approach of the “growthrates” (Petzoldt, 2016) package on log transformed cell count data.

Statistical Analyses

In all treatments with co-occurring species, the relative species contribution of *E. huxleyi* to total biovolume (Mix) over time was analyzed using a generalized least squares (GLS) model \[ \text{m0 < -gls (Biovolume ~ Selection CO}_2 \times \text{Time)} \]. Differences in variance structure were adjusted for the factor “Selection CO\(_2\)” and accounted for autocorrelation over time. For the statistical analysis, we only considered the relative contribution of one of the two species, as their respective contribution to the community was complementary. For the analysis of the absolute biovolume of each species in all cultures we used the same GLS model. The analysis was done for each species separately and started with the following full model: m0 < -gls (Biovolume ~ Selection CO\(_2\) × Selection Culture × Time). After reduction and analysis of the model structure we accounted for autocorrelation over time and differences in variance structure (varIDEN) for the “Selection Culture” in *E. huxleyi* and “Selection CO\(_2\)” and “Selection Culture” in *C. affinis*. There was a strong change around 160 days of experiment and we found significant interactions with “Time” for all factors in the GLS model. In order to investigate the effects of the different treatments on both species before and after this time point, we divided the time series data into two parts – BC1-BC20 or BC21-BC36 on which we repeated the described GLS model \[ \text{BC1-20 or BC21-36, respectively, -gls (Biovolume~Selection CO}_2 \times \text{Selection Culture)} \]. A permutational multivariate analysis of variance (permanova, with 999 permutations (using the package “vegan”) was used to test for the differences in genotype compositional change between the treatments.

To analyze the reciprocal assay data we used repeated measure analysis of variance (rmANOVA). Since during the experiment, 4 and 3 cultures were lost, we had to omit the data of the “lost” replicates from all statistical analyses of the assays. Before starting the analyses we tested for the homogeneity of variance using a Fligner Killeen Test (Fligner and Killeen, 1976) and for the normality of the residuals using a Shapiro–Wilk Normality Test (Shapiro and Wilk, 1965). The assumption of sphericity was not violated because we only had one repeated measure (Field, 2018). All assumptions were met such that we could continue with the analysis without data transformation. The growth rate was first assessed for the overall effects of “Time” (between the two assays), “CO\(_2\)” and “Culture” in both selection and assay environment and thus we started with a complete data set analysis \[ \text{aov((muExp)~(Selection CO}_2 \times \text{Selection Culture} \times \text{Assay CO}_2 \times \text{Assay Culture} \times \text{Time} + \text{Error(Replicate/Time))]} \] followed by a separate analysis of each assay. Here, interactions of selection treatment × assay treatment would indicate evolutionary adaptation over the course of the experiment.

All modeling and statistical analyses were done using the software R (R Core Team, 2016). The following packages were used for the analyses, plotting, and modeling: “ggplot2,” “deSolve,” “vegan,” and “ez” (Oksanen et al., 2007; Wickham, 2009; Soetaert et al., 2010; Lawrence, 2016).

RESULTS

Changes in Species Composition Over Time

The outcome of interspecific competition in ambient and high CO\(_2\) was mirrored in relative biovolume shifts: In the first half of the experiment the diatom dominated the two-species community with a relative biovolume (mean over first 20 batch cycles) of ca. 95 ± 7% (mean ± SD) in high compared to ambient CO\(_2\) condition with 80 ± 13% (mean ± SD; Figure 2; and Supplementary Table S4 BC1-20 “Selection CO\(_2\)” \[ F_{1,186} = 164.039, p < 0.0001 \]. From ca. 160 days onward there was not only a dominance reversal from *C. affinis* to *E. huxleyi* (Figure 2 and Supplementary Table S4 Full Model “Time” \[ F_{1,337} = 613.093, p < 0.0001, “Selection CO}_2 \times \text{Time} \] \[ F_{1,337} = 26.036, p < 0.0001 \], but also a different reaction of both species to CO\(_2\), with *E. huxleyi* being favored by high CO\(_2\) in the second phase of the experiment. The reversal of dominance was reflected in decreased final relative contribution of *C. affinis* of 43 ± 24% and 37 ± 26% in ambient and high CO\(_2\) conditions, respectively, without a statistically significant difference between the CO\(_2\) treatments (Figure 2, and Supplementary Table S4 BC21-36 “Selection CO\(_2\)” \[ F_{1,142} = 3.266, p = 0.072 \].

Temporal Dynamics of Absolute Species Biovolume

Similar to the dynamics of relative species contributions, the time course of absolute biovolume of both species was characterized by two distinct phases that changed around 160 days of the experiment (Figures 3A–D and Supplementary Table S5 *E. huxleyi* and *C. affinis* single and interactive effects of “Time” in Full model).

Overall *E. huxleyi* biovolume increased over time about one order of magnitude with an average 30-fold increase across all treatments (Figures 3A, B, and Supplementary Table S5 Full model, “Time” \[ F_{1,660} = 122.726, p < 0.0001 \]. During the first half of the experiment high CO\(_2\) reduced the biovolume of *E. huxleyi* by a third in the mono-cultures (Figures 3A–E, Supplementary Table S5 *E. huxleyi* BC1-20 “Selection CO\(_2\)” \[ F_{1,372} = 113.818, p < 0.0001 \]. The effect of high CO\(_2\) was, with a biovolume reduction of almost 4 times compared to the ambient treatment, even more negative in *E. huxleyi* in mix-cultures (Figures 3A–E, Supplementary Table S5 *E. huxleyi* BC1-20 “Selection CO\(_2\)” \[ Selection CO\(_2\) \times \text{Selection Culture} \] \[ F_{1,372} = 8.313, p = 0.0042 \]. This suggested that interspecific competition had an additive negative effect at first. In the second half of the experiment the effect of competition reversed and *E. huxleyi* biovolume was only slightly reduced in high CO\(_2\) in the mono-cultures whereas in mix-culture increased CO\(_2\) even had...
a slightly positive effect (Figures 3A–E, and Supplementary Table S5 E. huxleyi BC21-36 “Selection CO$_2$ × Selection Culture” $F_{1,280} = 7.156$, $p = 0.0079$).

In contrast to E. huxleyi the biovolume of C. affinis varied markedly between batch cycles with only a slight decrease of biovolume over time (Figures 3C,D, and Supplementary Table S5 C. affinis Full model, “Time” $F_{1,671} = 25.538$, and $p < 0.0001$). Specifically, the biovolume of C. affinis was not affected by increased CO$_2$ in the first half of the experiment, neither in mix- nor mono-culture (Figures 3C,D, and Supplementary Table S5 C. affinis BC1-20, “Selection CO$_2$” $F_{1,372} = 0.534$, and $p = 0.465$). This pattern changed toward the end of the experiment where C. affinis biovolume declined to half in the high CO$_2$ treatment in the mix-cultures (Figures 3D–F) while it doubled in the mono-cultures (Figures 3C–F, and Supplementary Table S5 C. affinis BC21-36, “Selection CO$_2$ × Selection Culture” $F_{1,352} = 15.986$, and $p < 0.0001$). This suggests that, over time, the diatoms in mix-culture were affected by the competing cocolithophores: in high CO$_2$ treatments biovolume of E. huxleyi increased more than in ambient CO$_2$ conditions leading to potentially stronger depletion of nutrient and consequently reduced growth of competing C. affinis (Supplementary Table S5 C. affinis Full model, three-way interaction $F_{1,671} = 12.225$, and $p = 0.0005$).

Genotype Sorting

The genotype composition changed uniformly over time in both species (Figure 4, and Supplementary Table S6, E. huxleyi “Time” $R^2 = 0.999$, $p = 0.001$, permutations = 999, C. affinis “Time” $R^2 = 0.999$, $p = 0.001$, and permutations = 999) with only a single genotype left for all cultures of E. huxleyi and C. affinis (C41 and B57, respectively). In E. huxleyi there was no effect of CO$_2$ on the genotype sorting (Figure 4, and Supplementary Table S6, “Selection CO$_2$” $R^2 = 0$, $p = 0.401$, and permutations = 999) and only a small difference between the single and co-occurring cultures (Figure 4, and Supplementary Table S6 “Selection culture” $R^2 = 0$, $p = 0.085$, and permutations = 999). The small difference likely came about due to slightly slower sorting to the dominant genotype C41 in the mix-culture (Figures 4A,B). For C. affinis we could not analyze the effect of either treatment on the genotype composition.

Reciprocal Adaptation Experiments

In E. huxleyi we found no significant interaction of selection with the assay treatment in neither of the two assays which would indicate evolutionary adaptation (Supplementary Table S7, E. huxleyi). Specifically, long-term selection to high CO$_2$ did not result in increased growth in the high CO$_2$ assay condition compared to ambient selected populations (Figures 5A,B). However, there were non-adaptive effects of CO$_2$ and culture selection and assay treatments that varied between the two assay experiments: After 64 days, selection under high CO$_2$ had a negative effect on the growth rates of E. huxleyi when measured in the presence of the diatom (“mix” assay treatment), while “mix” assay conditions led to a decline in growth rates in general (Figures 5A,B and Supplementary Table S7 E. huxleyi Assay 64 days “Selection CO$_2$ × Assay Culture” $F_{1,47} = 10.511$, $p = 0.002$, “Assay Culture” $F_{1,47} = 254.708$, and $p < 0.0001$). These effects were largely absent in the second assay after 288 days where growth rates were only affected by the assay CO$_2$ treatment (Supplementary Table S7 E. huxleyi Assay 64 days, $F_{1,47} = 8.066$, and $p = 0.007$). There was a significant increase in growth rates of ca. 10–45% over time in all selection treatments (Figure 5A and Supplementary Table S7 E. huxleyi, “Time” $F_{1,47} = 391.511$, and $p < 0.0001$), with the strongest increase in cultures that had been selected in high CO$_2$ in the presence of the diatom.
FIGURE 3 | Biovolume changes over approx. 200 generations. The biovolume of *E. huxleyi* (A,B) and *C. affinis* (C,D) measured at the end of each batch cycle over 36 batch cycles in the four selection treatments is shown here (mean ± SE; *n* = 5 (5 replicates/treatment) for days 1–200; and *n* = 4 (4 replicates/treatment) for days 200–288 per time point). The light gray and black lines show the time points where samples were taken for genotype reisolation. The black lines show when the cultures were additionally tested for adaptation in the reciprocal assay test. Panels (E,F) show the relative difference between the ambient and high CO$_2$ treatments for *E. huxleyi* (E) and *C. affinis* (F) when grown alone (black points) and with the respective other species (gray points) for better understanding how the responses to increased CO$_2$ concentrations changed over time.

No selection for higher growth rates to exposure to increased CO$_2$ could be observed in *C. affinis*. In contrast to this expectation the growth rates decreased by a tenth over time throughout all treatments (Figure 5C and Supplementary Table S7 *C. affinis* "Time" *F*$_{1,46}$ = 14.843, *p* < 0.001). Nevertheless, we found different responses to CO$_2$ with and without competition that even resulted in an seemingly adaptive effect to increased CO$_2$ in the second assay after 288 days, evident as a significant interaction of selection and assay treatment (Figure 5D "u-shape" pattern in pooled data, Supplementary Table S7 *C. affinis*).
Assay 288 days “Selection CO\textsubscript{2}” \times “Assay CO\textsubscript{2}” \(F_{1,47} = 17.807,\) and \(p = 0.0001\). However, this “adaptive” response was mainly driven by very low growth rates of \textit{C. affinis} that have been selected in mix-cultures to high CO\textsubscript{2} as well as low growth rates of \textit{C. affinis} selected under ambient CO\textsubscript{2} in mono-cultures. This result showed that disentangling the effects of both abiotic and biotic factors is crucial to avoid overestimation of adaptive responses to abiotic factors alone.

**DISCUSSION**

Our results show that long-term selection in response to abiotic factors in communities reaching stationary phase can drastically diverge from the predicted outcomes of (i) short term experiments with the same abiotic driver and (ii) selection experiments where species are held under exponential growth (Riebesell, 2004; Lohbeck et al., 2012; Schlüter et al., 2014; Hattich et al., 2017). We here show that the divergence in the community response potentially arises from the effect of evolutionary on ecological processes, as previously shown in other systems (Hairston et al., 2005; Schoener, 2011). During the first half of the experiment our initial expectations (for biovolume) with respect to species being an ecological “loser” and “winner” in response to enhanced CO\textsubscript{2} were met for \textit{E. huxleyi} but not \textit{C. affinis}. This led to the expected dominance of \textit{C. affinis}, which was more pronounced under increased CO\textsubscript{2} conditions. However, over time, the observed effects of increased CO\textsubscript{2} on biovolume changed for both species so as to revert our predicted ecological “loser” and “winner” outcome. Consequently, the relative contributions in the communities flipped and \textit{E. huxleyi} became dominant under both CO\textsubscript{2} conditions. This change in community composition was reflected in a strong increase in biovolume of \textit{E. huxleyi} and a weak decrease in biovolume of \textit{C. affinis}. However, these changes in species’ biovolume over time were the result of different drivers: While the \textit{E. huxleyi} biovolume correlated mainly with cell numbers (Supplementary Figures S3A, S4A), \textit{C. affinis} biovolume correlated mainly with cell volume (Supplementary Figures S3B, S4B). We note that the “dominance reversal” coincided with strong genotype selection in both species, which was unexpectedly not driven by CO\textsubscript{2} or species competition treatment. Likewise, neither species showed
adaptation to CO$_2$ or competition in the reciprocal assays. We can only speculate why this "dominance flip" occurred at the specific point in the experiment: Coinciding with the flip the winning genotype C41 in E. huxleyi became dominant, suggesting a feedback from evolutionary changes on ecology. Potentially a critical abundance threshold was reached by E. huxleyi that was necessary to become competitively superior over C. affinis. Characteristics and traits of this winning genotype likely have played a role when competing with C. affinis, but could not be characterized herein.

For the coccolithophore, the absence of CO$_2$ adaptation found here is in contrast to several previous studies (Lohbeck et al., 2012; Schlüter et al., 2014), while there is conflicting prior evidence on the diatom species (Scheinin et al., 2015; Li et al., 2017). The contrasting finding of no adaption or differential genotype sorting to increased CO$_2$ for E. huxleyi could potentially be explained by differences in the experimental setups concerning the CO$_2$ manipulation, batch cycle length, experimental duration, light, temperature, and nutrient availability (Rost et al., 2008; Gao and Campbell, 2014; Meyer and Riebesell, 2015). A marked difference of our experimental setup compared to other experimental evolution studies was that our species were not kept under constant exponential growth (Lohbeck et al., 2012; Li et al., 2017; Pardew et al., 2018) but ran into stationary phase. The resulting recurrent nutrient repletion and limitation at the start and end of each batch cycle, respectively, could likely have overridden CO$_2$ effects on growth rate as observed herein (or f.e. in Aranguren-Gassis et al., 2019), because selection on other traits, e.g., important for nutrient acquisition, might have been more important. For C. affinis, however, it remains uncertain if nutrient limitation has likewise overwritten the expected adaptive response to CO$_2$ or if the species was really unaffected by this driver. There is no directly comparable study with the same species, which does not run into nutrient limitation. Moreover the two long-term studies existing for diatoms do show diverging adaptive responses, which might depend on different experimental set-ups as well as species identity, as largely diverging short-term responses have already been described between diatom species (Gao and Campbell, 2014; Bach and Taucher, 2019).

It is well accepted that a more complex ecological context allowing for competition can affect evolutionary dynamics (de Mazancourt et al., 2008; Barralough, 2015; Lancaster et al., 2017) but we did not find a significant effect in this study. Although not demonstrated to date, it is quite likely that interspecific competition has the potential to affect evolution in phytoplankton, as already intraspecific competition was shown to alter adaptive responses to CO$_2$ (Collins, 2011a). In this study, the dominant C. affinis strongly reduced the biovolume (throughout the experiment) and growth rate (in both reciprocal assays) of E. huxleyi, demonstrating the strong effect of competition, especially in the first half of the experiment. In addition, we found a further reduction of growth rates to increased CO$_2$ with the co-occurrence of the diatom indicating that the ecological context did play a role albeit not affecting the evolutionary change. That no effect of interspecific competition on species' evolution, or adaptation could be observed potentially depends on how coexistence was allowed in our experiment. Owing to the species' different nutrient uptake kinetics (Sommer, 1984) niche partitioning likely appeared temporary over the course of a batch cycle. Whereas the diatom had an advantage during the first batch cycle days with replete nutrients, the coccolithophore with higher affinity was favored and could grow longer toward the end of each batch cycle when nutrients became limiting (Sommer, 1984; Supplementary Figures S1-1, S1-2). As such selection on standing genotype composition likely took place within these two niche spaces of both coexisting species, unaffected by the presence of the interspecific competitor.

Although adaptive responses to either increased CO$_2$ or competition were absent, we still observed strong evolutionary change as very reproducible, directional sorting of genotypes across all treatments. Interestingly, these evolutionary dynamics selected for one single genotype of each species that became dominant already after approximately 64–160 days of the experiment. While we know that evolution was independent of the intended CO$_2$ and species competition treatments, non-intended laboratory selection was the most likely driver in our study. Laboratory selection is inevitable and often strong and has been previously demonstrated over very long time scales by comparing ancestral bacterial populations to laboratory evolved populations (Lenski and Travisano, 1994). Here, selection became apparent when looking at different responses in both species over time: We found an overall change in E. huxleyi and C. affinis biovolume and growth over time (Supplementary Figure S5). Interestingly, changes in traits of E. huxleyi (e.g., growth rate, cell size) were also observed in another long-term laboratory selection study (Schlüter et al., 2016). In these experiments, however, laboratory adaptation was only a “background” signal, whereas in our experiments the chosen treatments were too weak to impose a selection force to overcome laboratory selection. Schlüter et al. (2016) started their experiment with one single genotype, implying that observed changes in growth rate were a result of novel mutations, while in our study, observed changes in biovolume and growth rate could be the result of both selection on genotypic diversity and mutations. Among the single remaining genotype it is
possible that within the 200 generations of experimentation new mutations occurred (Elena and Lenski, 2003) that may contribute to phenotypic divergence in genotypes solely characterized by microsatellite marker alleles. With the caveat of not having ancestral populations of genotypes as a comparison we cannot say how laboratory selection played out in our species. Certain is, however, that changes in species’ characteristic as observed in changed biovolume growth rate responses to the treatments throughout the experiment were strong.

When studying eco-evolutionary dynamics, complex ecological conditions can evidently impinge upon evolutionary dynamics, while the complementary question asks how evolutionary change affects ecological processes (for example Fussmann et al., 2007; Ellner et al., 2011; Koch et al., 2014). In our study, the genotype compositional change (i.e., strong evolutionary change) coincided or rather preceded reversal in community composition (i.e., ecological change). That this dominance shift and underlying genotype selection also occurred in an earlier attempt of the experiment, suggests that the eco-evolutionary dynamics observed are reproducible (Supplementary Figures S6, S7). We, however, propose that selection by nutrient limitation on standing variability in genotype’s competitive abilities was the most likely driver resulting in the eco-evolutionary interaction. However, necessary experimental characterization of the genotypes’ nutrient uptake associated traits is (currently) missing. Other studies reported nutrient uptake related traits in marine protists to vary between genotypes to the same extent as growth rates (Brandenburg et al., 2018). Since we have shown that the variability in growth rates among the genotypes used in this study is even larger than that of the response to increased CO$_2$ (Hattich et al., 2017), we postulate that their nutrient uptake related traits vary as well on which selection can take place. Just recently evolution in nutrient related competitive traits in phytoplankton has been demonstrated (Bernhardt et al., 2020). Another indication that nutrients were driving the change in our system is the observed shift away from C. affinis toward E. huxleyi, the favored species by nutrient limitation (Tyrrell and Merico, 2004). While studies on natural phytoplankton communities (i.e., in mesocosm studies; see for example Peter and Sommer, 2015) and nutrient dependent seasonal successions within phytoplankton populations (Sommer, 1984) are well understood, the underlying eco-evolutionary mechanisms driving nutrient associated changes are less well studied. Most previous studies were not run over sufficient generations to permit genotypic selection, the simplest form of evolution to occur in phytoplankton communities. Consequently, future experimental eco-evolutionary studies should focus more on the consequences on how nutrients select to predict phytoplankton change (Thomas et al., 2017; Bernhardt et al., 2020). This will become particularly relevant as nutrient uptake related traits explain competitive ability under different nutrient conditions (Sommer, 1984; Litchman et al., 2007) and there will be more nutrient limitation on phytoplankton in the future ocean (Boyd et al., 2013; Moore et al., 2013).

To conclude, the most parsimonious explanation for the observed dominance flip among the two globally important phytoplankton species E. huxleyi and C. affinis, is rapid evolutionary feedback to the ecology. Rapid evolutionary change as genotype sorting driven by experimental selection (potentially via nutrient limitation) and not CO$_2$ environment turned an ecological “loser” (with respect to biovolume contribution to the community) into a “winner” and vice versa. As such, this flip demonstrates for the first time that eco-evolutionary interactions play out in competing phytoplankton communities. Such interactions can drastically alter the effect of environmental drivers and lead to diverging predictions of future changes compared to such resulting of short-term studies. Considering the role that phytoplankton play in biogeochemical cycles as primary producers, changes in biovolume composition and/or in size within populations owing to evolutionary change due to the combination of abiotic and biotic interactions identified here could have far reaching effects on ecosystem dynamics (Falkowski et al., 2008). Our results call for an inclusion of more realistic experimental evolution conditions in future studies, not only using realistic nutrient regimes, but more importantly also including multi-species settings and their underlying mechanisms allowing for stable coexistence to simultaneously investigate ecological and evolutionary processes in phytoplankton.

**DATA AVAILABILITY STATEMENT**

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: PANGAEA (https://doi.org/10.1594/PANGAEA.887780).

**AUTHOR CONTRIBUTIONS**

The basic grant proposal supporting this work was written by BM and TR. The experiment was developed by GH and then the experiments further designed by LL and GH. Lab work carried out by LL and GH and data analyses by LL, GH, BM, and TR drafted the manuscript. All authors revised the manuscript and gave final approval for publication.

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Supplementary Material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmars.2020.00634/full#supplementary-material

Further methodological explanations on reciprocal assay experiments and genotype identification

Supplementary Material

Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmars.2020.00634/full#supplementary-material

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