Analyzing the Impacts of Elevated-CO\textsubscript{2} Levels on the Development of a Subtropical Zooplankton Community During Oligotrophic Conditions and Simulated Upwelling

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Ocean acidification (OA) is affecting marine ecosystems through changes in carbonate chemistry that may influence consumers of phytoplankton, often via trophic pathways. Using a mesocosm approach, we investigated OA effects on a subtropical zooplankton community during oligotrophic, bloom, and post-bloom phases under a range of different pCO\textsubscript{2} levels (from \(\sim 400\) to \(\sim 1480\) µatm). Furthermore, we simulated an upwelling event by adding 650 m-depth nutrient-rich water to the mesocosms, which initiated a phytoplankton bloom. No effects of pCO\textsubscript{2} on the zooplankton community were visible in the oligotrophic conditions before the bloom. The zooplankton community responded to phytoplankton bloom by increased abundances in all treatments, although the response was delayed under high-pCO\textsubscript{2} conditions. Microzooplankton was dominated by small dinoflagellates and aloricate ciliates, which were more abundant under medium- to high-pCO\textsubscript{2} conditions. The most abundant mesozooplankters were calanoid copepods, which did not respond to CO\textsubscript{2} treatments during the oligotrophic phase of the experiment but were found in higher abundance under medium- and high-pCO\textsubscript{2} conditions toward the end of the experiment, most likely as a response to increased phyto- and microzooplankton standing stocks. The second most abundant mesozooplankton taxon were appendicularians, which did not show a response to the different pCO\textsubscript{2} treatments. Overall, CO\textsubscript{2} effects on zooplankton seemed to be primarily transmitted through significant CO\textsubscript{2} effects on phytoplankton and therefore indirect pathways. We conclude that elevated pCO\textsubscript{2} can change trophic cascades with significant effects on zooplankton, what might ultimately affect higher trophic levels in the future.

Keywords: microzooplankton, mesozooplankton, mesocosms, ocean acidification, nutrients, Oncnaea, trophic transfer efficiency
INTRODUCTION

Anthropogenic emissions are increasing atmospheric CO$_2$ concentrations from pre-industrial levels of $\sim$280 µatm to current levels of over 400 µatm, and increases up to 1000 µatm by the end of the century are projected for the RCP8.5 emission scenario (IPCC, 2013). The oceans act as carbon sinks, absorbing about one third of the anthropogenic CO$_2$ emission (Sabine et al., 2004). This oceanic CO$_2$ uptake causes a shift in carbonate chemistry with a decrease in seawater pH, commonly known as ocean acidification (OA). Recent years of intense research have shown that OA may cause substantial changes to marine ecosystems (IPCC, 2013; Kroeker et al., 2013).

Despite the large body of literature related to biological responses to OA, most studies investigated single species responses, which may rarely provide a sufficient basis to understand long-term responses in complex ecological environments (Harley, 2011; Queirós et al., 2015). Moreover, changes in pCO$_2$ may promote changes in trophic interactions, leading to the dampening or amplification of single species effects and hence promoting shifts in community composition (Lischka et al., 2011; Rossoll et al., 2012, 2013). Consequently, in situ community studies are important in order to evaluate OA effects at the level of assemblages and ecosystems (Guinotte and Fabry, 2008; Riebesell andGattuso, 2015).

Focusing on marine plankton, nutrient conditions can determine how communities respond to OA (Alvarez-Fernandez et al., 2018), being the most noticeable pCO$_2$ effects often observed under limiting inorganic nutrient conditions (Paul et al., 2015; Sala et al., 2015; Bach et al., 2016b). This is because elevated CO$_2$ levels cause an increase in phytoplankton standing stocks —more pronounced in smaller-sized taxa— and this effect on primary producers may be transferred differently into heterotrophic primary consumers depending on the inorganic nutrient availability (Alvarez-Fernandez et al., 2018). The present study focussed on an oligotrophic system around the island of Gran Canaria within the Canary Archipelago, located in the subtropical North Atlantic Ocean. Despite its overall oligotrophic character, this region experiences a short-term period of deep-water nutrient inputs in later winter (February–March) (de León and Braun, 1973; Cianca et al., 2007) as well as recurrent mesoscale upwelling events that act as an offshore pump of organic matter and carbon (Sangrà et al., 2009). The so-called late winter bloom usually causes an increase in primary production and chlorophyll $a$ concentration in the euphotic zone (Menzel and Ryther, 1961; Aristegui et al., 2001). Typically, mesozooplankton grazing pressure exerted on phytoplankton is low in the study area (Aristegui et al., 2001; Hernández-León et al., 2004), and mesozooplankters are considered to feed on microzooplankton which, in turn, control primary production (Hernández-León et al., 2001; Quevedo and Anadón, 2001; Calbet and Alcaraz, 2007). The microzooplankton community is usually dominated by small dinoflagellates and aloricate ciliates (Quevedo and Anadón, 2001), while the most important mesozooplankton during the annual cycle are copepods (Hernández-León et al., 2007). However, the plankton community typically changes during bloom conditions (Aristegui et al., 2001; Hernández-León et al., 2004; Schmoker et al., 2012). An increase in copepod abundances follows the increase in primary production, and a trophic cascade caused by the consumption of microzooplankton by mesozooplankton allows a further increase in autotrophic biomass by the combined effect of top-down control and nutrient remineralization (Hernández-León, 2009; Schmoker et al., 2012). This bloom situation may cause a reduction in the efficiency of the food web, considering that trophic transfer efficiency (i.e., zooplankton growth per unit phytoplankton production) tends to be diminished under nutrient enrichment conditions due to the limited capacity of grazers to use the boosted algae production (Calbet et al., 1996; Kemp et al., 2001; Calbet et al., 2014).

In order to assess the impacts of OA on zooplankton communities we must consider not only direct effects on zooplankton caused by pH reductions, but also effects that reach consumers indirectly, through trophic pathways (Boersma et al., 2008; Rossoll et al., 2012; Cripps et al., 2016). Detrimental indirect pCO$_2$ effects have been described in single species of herbivores (Schoo et al., 2013; Meunier et al., 2016) and secondary consumers (Lesniowski et al., 2015). In the case of copepods, bottom-up influences of OA seem to be largely associated with interspecific differences among prey items with regard to their sensitivity to elevated pCO$_2$ levels, as observed when analyzing the cellular stoichiometry of copepods’ photosynthetic preys (Isari et al., 2015; Meunier et al., 2016). In turn, microzooplankton may be affected by the effect of high pCO$_2$ levels on phytoplankton availability or quality such as an increase in picophytoplankton standing stock or changes in their cellular carbon-to-nutrient ratios (Bach et al., 2016b; Meunier et al., 2016). In addition, a high pCO$_2$ scenario is likely to favor harmful algal blooms (Wells et al., 2015) with substantial consequences for energy transfer from primary producers to consumers within marine communities.

Plankton community OA studies to date have been mostly carried out in relatively eutrophic environments (but see Sala et al., 2015; Gazeau et al., 2017), and led to varying conclusions. Some studies showed the lack of major effects of elevated pCO$_2$ levels in micro- (Aberle et al., 2013; Horn et al., 2016) and mesozooplankton abundances (Niehoff et al., 2013), while others detected both changes in community size distributions and biomass (Lischka et al., 2017; Taucher et al., 2017b) as well as positive bottom-up pCO$_2$ responses on mesozooplankton abundances (Algueró-Muñiz et al., 2017). Inorganic nutrient availability can control these different responses to OA in planktonic communities, thereby the nutrient-deplete phases could determine the transfer of the pCO$_2$ effects on primary producers to primary consumers (Alvarez-Fernandez et al., 2018). Taking this into account, studying OA effect in oligotrophic systems —which represent most of the global surface ocean— becomes of paramount importance. To accomplish this goal, we performed an experiment that allowed us to study the contrast between nutrient-depleted and nutrient-replete periods. Our aim was to analyze the effects of OA on the development of an autumn zooplankton community from the subtropical North Atlantic, including a simulated bloom situation. We assessed the effects of pCO$_2$ on (1) the abundance of subtropical...
micro- and mesozooplankton under oligotrophic and upwelling conditions, (2) the size and reproductive output of an important copepod species and (3) the trophic efficiency within the plankton community under different conditions.

**MATERIALS AND METHODS**

**Mesocosms Setup and Experimental Design**

The experiment was conducted from 27th September (t-4) until 26th November 2014 (t56) within the framework of the BIOACID II project (Biological Impacts of Ocean ACIDification) and was hosted by the Plataforma Oceánica de Canarias (PLOCAN, Spain). In order to study the effects of changing carbonate chemistry conditions on the plankton community succession, nine mesocosms (KOSMOS, M1-M9: “Kiel Off-Shore Mesocosms for Ocean Simulation”), were deployed in Gando Bay (27°55′41″ N, 15°21′55″ W), on the west coast of Gran Canaria (Canary Islands, Spain) (Taucher et al., 2017a). The nine cylindrical mesocosm units (13 m deep, 2 m diameter) enclosed water volumes (∼35 m$^3$) sealed by sediment traps installed at the bottom of each mesocosm bag (Boxhammer et al., 2016). Target $p$CO$_2$ was reached at the beginning of the experiment by adding CO$_2$ saturated seawater to the mesocosms following the protocol described in Riebesell et al. (2013). The carbonate chemistry of the enclosed seawater was manipulated by stepwise additions of CO$_2$-saturated seawater in four steps over 7 days. Two further CO$_2$ additions were conducted on days 21 and 38 to compensate for the loss of CO$_2$ through air-sea gas exchange. As $p$CO$_2$ treatments we established a gradient from current levels to end-of-century scenarios, representing IPCC predictions for mitigation scenarios (RCP 2.6) as well as medium (RCP 6.0) and high (RCP 8.5) $p$CO$_2$ levels (IPCC, 2013). The mean $p$CO$_2$ values per mesocosms between t1 and t55 were M1 = 369, M2 = 887, M3 = 563, M4 = 716, M5 = 448, M7 = 668, M8 = 1025 and M9 = 352 µatm. Analyzing the oligotrophic phase of the experiment, we could differentiate three $p$CO$_2$ groups by a k-means cluster analysis (Jain, 2010). The outcome showed three distinguishable clusters: low-$p$CO$_2$ (M1, M9, M5; $k = 460$ µatm) medium-$p$CO$_2$ (M3, M7, M4; $k = 721$ µatm) and high-$p$CO$_2$ levels (M2, M8; $k = 1111$ µatm) (Figure 1A) which were used for the analyses presented throughout this paper. Unfortunately, we detected a hole in the enclosure bag of the third high-$p$CO$_2$ mesocosm (M6 = 976 µatm) on t27, so M6 was excluded from sampling and analyses after that date.

To simulate a natural upwelling event, we collected deep water (∼84 m$^3$) from 650 m depth on t22, as described by Taucher et al. (2017a). From each mesocosm, a defined volume of water was removed from 5 m depth with a submersible pump (Grundfos SP-17-5R). Consequently, in a process of ∼9 h duration during the night of t24, deep water was pumped into the mesocosms,
were obtained every 4 days and fixed with Lugol’s solution. Phytoplankton samples for microscopy were carefully mixed to avoid a bias due to particle sedimentation. All carboys were protected from sunlight during sampling and deep-water addition ensured homogenous vertical distribution of deep water inside the mesocosms.

All sampling methods and analyses are described in detail in the overview paper provided by Taucher et al. (2017a). Briefly, regular sampling — conducted every 2nd day before deep water addition, daily after t25 — included CTD casts, water column sampling, and sediment sampling. CTD casts were carried out with a hand-held CTD probe (CTD60M, Sea and Sun Technologies) in each mesocosm and in the surrounding water. Thereby we obtained vertical profiles of temperature, salinity (Figure 1B), pH, dissolved oxygen, chlorophyll a, and photosynthetically active radiation (PAR). Vertical profiles of temperature and salinity showed a uniform distribution of both variables, indicating that there was no stratification and that the water columns in the mesocosms were well-mixed throughout the entire study period (Taucher et al., 2017a). Water column samples were collected with integrating water samplers (IWS, Hydrobios, Kiel), in which a total volume of 5 L from 0 to 13 m depth was collected evenly through the water column. This water was either used for samples sensitive to contamination such as nutrient analyses, which were directly filled into separate containers on board, or stored in carboys for later subsampling for parameters such as phytoplankton and microzooplankton. Some analyses required larger volumes of water than could be sampled with the IWS in a reasonable time frame, e.g., pigment samples for reverse-phase high-performance liquid chromatography (HPLC) analysis. To enable a faster water collection, we used a custom-built pump system connected to a 20 L carboy. By creating a gentle vacuum and moving the inlet of the tube evenly up and down in the mesocosm during pumping, samples similar to those from the IWS were obtained. All carboys were protected from sunlight during sampling and stored in a temperature-controlled room at 16°C upon arrival on shore. Before taking subsamples from the carboys, they were carefully mixed to avoid a bias due to particle sedimentation.

Pigments such as Chlorophyll a (Chla in the following) were analyzed using HPLC (Figure 1C). Nutrients (nitrate + nitrite (NOx), Figure 1D) were measured using an autoanalyzer (SEAL Analytical, QuAAstro) coupled to an autosampler (SEAL Analytical, XY2). NOx are presented here as a proxy for inorganic nutrients spec [see PO$_4^{3-}$, Si(OH)$_4$ and NH$_4^+$ dynamics in Taucher et al. (2017a)]. Phytoplankton samples for microscopy were obtained every 4 days and fixed with Lugol’s solution. They were analyzed using the Utermöhl (1958) technique and classified to the lowest possible taxonomical level. Biomass of phytoplankton was estimated by using conversion factors, as detailed in Supplementary Table S1 (Tomas and Hasle, 1997; Ojeda, 1998; Leblanc et al., 2012).

**Zooplankton: Sampling and Analysis**

For the analysis of the microzooplankton community (microZP) —the size class of 20–200 μm— samples from the IWS were taken every 8 days until day 50. 250 mL of mesocosm water was transferred into brown glass bottles, fixed with acidic Lugol’s solution (1–2% final concentration), and stored in the dark. MicroZP was counted and identified with an inverted microscope (Axiovert 25, Carl Zeiss) using the Utermöhl (1958). 50 mL of each sample was transferred into a sedimentation chamber and allowed to settle for 24 h prior to counting. Depending on plankton abundances, the whole or half of the chamber was counted at 100-fold magnification to achieve a count of at least 300–400 individuals for the most common taxa. MicroZP was identified to the lowest possible level (genus or species level) and otherwise grouped into size classes according to their distinct morphology. MicroZP were grouped into ciliates (aloricate and loricate) and dinoflagellates (athecate and thecate, size classes: small (<25 μm) and large (>25 μm)). As most dinoflagellates are capable of heterotrophic feeding (Calbet and Alcaraz, 2007), they can be considered as mixotrophic and were thus included in the microZP. Only few dinoflagellate taxa such as Ceratium or Dinophysis are considered to be predominantly autotrophic and were thus included in the phytoplankton (Tomas and Hasle, 1997). MicroZP biovolumes were estimated using geometric proxies obtained from literature (Ojeda, 1998; Hillebrand et al., 1999; Montagnes et al., 2001; Schmoker et al., 2014), and transformed to carbon biomass using conversion factors provided by Putt and Stoecker (1989) and Menden-Deuer and Lessard (2000) for ciliates and dinoflagellates, respectively (see Supplementary Table S1).

The mesozooplankton community (mesoZP) was sampled in the mesocosms by vertical net hauls with an Apstein net (55 μm mesh size, 17 cm diameter) equipped with a closed cod end. Sampling depth was restricted to 13 m to avoid resuspension of the material accumulated in the sediment traps at 15 m depth. Every net haul consisted in total filtered volume of 295 L. One net haul per mesocosm was carried out once every 8 days, always during the same time of the day (2–4 pm local time) to avoid diel differences in community composition. Samples were rinsed on board with filtered sea water, collected in containers and brought to the on-shore laboratory (PLOCAN, ~4 nm distance), where samples were preserved in denatured ethanol. For transportation, the samples were placed in cooling boxes until fixation of the organisms.

During analysis, organisms were sorted using a stereomicroscope (Olympus SZX9) and classified to the lowest possible taxonomical level. Copepodites and adults were classified together on a species/genus level, with the exception of the genus Oncceae, for which adults and copepodites were considered separately for a more in-depth study of this copepod. Nauplii from different species were pooled together. Taxonomical analysis was carried out focusing on copepods as the most

| TABLE 1 | Generalized additive mixed model (GAMM) structures. |
| Models | Meaning |
| s(DoE) | Temporal trend |
| s(DoE) : pCO$_2$ | Effect of pCO$_2$ on the temporal trend |
| s(DoE) + pCO$_2$ | Temporal trend and an independent pCO$_2$ effect on abundances |

DoE, day of experiment.
an abundant group (Boltovskoy, 1999). Every sample was sieved using a 50 μm mesh, rinsed with fresh water and divided with a Folsom plankton splitter (1:2, 1:4). Abundant species/taxa (>200 individuals in an aliquot) were only counted from subsamples, while less abundant species/taxa were counted from the whole sample. An in depth analysis of the succession of calcifying zooplankton is provided by Lischka et al. (2018).

As a proxy to explore the system’s energy transfer efficiency from producers to consumers (i.e., trophic transfer efficiency, TTE), we used the quotient autotrophy:heterotrophy (A:H) adapted from Calbet et al. (1996, 2014). This proxy was based on phytoplankton (Guan, 2018), heterotrophic microZP and mesoZP abundances transformed into biomass (see Supplementary Table S1 for further details). Low TTE is indicated by a higher biomass A:H ratio, hence TTE and A:H are inversely correlated.

Oncaea spp. Development

*Oncaea* is a common genus in the Canary Current System, where it has been typically recorded during the upwelling season (Hernández-León, 1998; Huskin et al., 2001; Hernández-León et al., 2007). *Oncaea* spp. is of special interest for this study because of (1) its trophic interaction with appendicularians (Go et al., 1998), which in turn may benefit from increased *pCO₂* levels and nutrient enrichment conditions (Troedsson et al., 2013) and (2) to our knowledge, poecilostomatoid copepods had not been studied in an OA context before. Hence, despite being not the most abundant mesoZP taxon within the mesocosms (Poecilostomatoida; 8% total mesoZP catch) we focused on the condition of *Oncaea* to investigate direct and/or indirect *pCO₂* effects on the female copepod length and reproductive output. Females were sorted from the same samples used for species determination, i.e., one sample per mesocosms (M1–M9) every 8 days during the whole study period (see section “Zooplankton: Sampling and Analysis”). The whole sample was scanned under the stereomicroscope (Olympus SZX9) and the first 20 adult females per sample were selected. Prosome length of every individual was measured, and females were classified regarding sexual development (mature/immature) and presence or absence of egg sacks. Females with developing egg sacks were classified as mature, while females which did not present any egg sack or eggs inside the prosome were rated as immature individuals.

Statistical Analyses

We used non-metric multidimensional scaling (NMDS) as exploratory analysis to describe the zooplankton community development in the mesocosms throughout the experiment. In our case the data matrix comprised abundances of each phytoplankton, microZP and mesoZP taxon in each mesocosm and on each sampling day (69 MK_timestep × 96 taxa). The treatment effect was assessed by using permutation tests on the community position in the NMDS space. These permutations check if the area of clusters formed by the treatment in the NMDS are smaller than randomized samples of the same size (Legendre and Anderson, 1999). In a complementary approach, we applied an ANalysis Of SIMilarity (ANOSIM) test (Clarke, 1993) as a post-analysis to compare the mean of ranked dissimilarities between *pCO₂* treatments to the mean of ranked dissimilarities within treatments. This analysis tests the assumption that ranges of (ranked) dissimilarities within groups are equal, or at least very similar (Buttigieg and Ramette, 2014).

To describe the temporal trends of each taxon during this experiment we used generalized additive mixed models (GAMMs) (Wood, 2006; Zuur et al., 2009) with a Gamma distribution and a logarithmic link. Three different kinds of models were fitted to each abundance group (Table 1).

Each of these models allowed the abundance temporal trend to vary differently between *pCO₂* treatments, representing (a) an equal temporal trend for all mesocosms [s(DoE)], (b) an effect of *pCO₂* on the temporal trend [s(DoE):*pCO₂*], and (c) an equal temporal trend with an independent *pCO₂* effect [s(DoE) + *pCO₂*]. This way, potential differences between *pCO₂* treatments could be detected as either (b) changes in phenology or (c) an increase/decrease of overall abundance. If necessary, models were fitted with an autocorrelation structure of first order to account for temporal autocorrelation in the data (Zuur et al., 2009). Statistically significant models were compared by the coefficient of determination ($R^2$), which indicates the proportion of the variance in the dependent variable that is predictable from the independent variables. For each taxon, the model with the highest $R^2$ was considered to best represent the abundance data. Models presented here accounted from t1, whilst t-3 abundances have been included in the figures to illustrate conditions prior to *pCO₂* manipulations within the mesocosms.

Differences in the condition of *Oncaea* females were analyzed by generalized linear mixed models (GLMMs) comparing the potential effect of *pCO₂* and time on development, prosome length and reproductive output. The effect of the day of experiment (t1–t56) and *pCO₂* treatment (low-, medium-, and high-*pCO₂*) on the studied parameters as well as their interaction were considered in the models. A Poisson distribution with a log link was used for the GLMM of count data, while length data was analyzed with a Gamma distribution. Unfortunately, the relatively low zooplankton sampling frequency did not allow for testing *pCO₂* effects on a continuous manner. As an alternative, different *pCO₂* levels were grouped in low-, medium- and high-*pCO₂* by a k-means cluster analysis, as described in Section “Mesocosms Setup and Experimental Design.”

We used R [version 3.0.2, (R Core Team, 2012)] to fit abundance data with the GAMMs and GLMs. The significance level for all statistical analysis was set to $p < 0.05$.

RESULTS

Community Change

The 2-dimensional representation of the community (NMDS) showed a strong trend in time (plankton succession), and a divergence of this trend from ca. t25 between the high-*pCO₂* mesocosms and the low- and medium- ones (Figure 2). Treatments followed a similar trend from t-3 until t17, but tended to separate afterwards, matching the simulated upwelling caused by deep water addition (t24). Permutation tests (with 999 permutations) did not show the areas (i.e., clusters of samples)
representing the different pCO$_2$ treatments to be significantly smaller than randomized areas, indicating that the variation due to CO$_2$ is smaller than the changes over time (i.e., natural succession) (ANOSIM test, $p$-value = 0.246). Areas representing the sampling day were significantly different from randomized areas using the same test, indicating a temporal trend ($p$-value = 0.001). Moreover, results for the interaction between sampling day and pCO$_2$ treatment (ANOSIM test, $p$-value = 0.001)
matched with the NMDS, suggesting that there was a significant effect of pCO$_2$ on plankton succession, ultimately affecting the plankton community development after the simulated upwelling event. Consequently, the plankton community developed differently within the different pCO$_2$ treatments, the largest difference being in the high-pCO$_2$ mesocosms.

**Zooplankton Temporal Trends**

In view of zooplankton abundance and Chla levels we could define three experimental phases: pre-bloom (from t1 until deep water addition on t24), phytoplankton bloom phase (t25–35) and post-bloom phase (from t35 until the end of the experiment), as shown in Figure 1C.

The microzooplankton (microZP) community comprised 13 different taxonomic groups of heterotrophic dinoflagellates and ciliates. Temporal trends of total microZP were affected by pCO$_2$ [s(DoE):Treat, Table 2], resulting in higher abundances under the high-pCO$_2$ treatment on the last sampling day. Averaged microZP abundances at the beginning of the experiment (t1) were 4.5 × 10$^6$ ± 2.89 × 10$^6$ individuals per m$^3$ for the low-, 3.45 × 10$^6$ ± 8.03 · 10$^5$ for the medium-, and 4.07 · 10$^6$ ± 9.36 · 10$^5$ for the high-pCO$_2$ treatments, respectively. After deep water addition (t24), abundances increased in all treatments, especially in the medium-pCO$_2$ treatment. Maximum values were reached at the end of the experiment (t50) in the high-pCO$_2$ treatment with 2.14 · 10$^7$ ± 8.94 · 10$^6$ individuals per m$^3$ (1.44 · 10$^7$ ± 6.61 · 10$^6$ and 1.52 · 10$^7$ ± 1.08 · 10$^7$ individuals per m$^3$ in the low- and medium-pCO$_2$ treatments, respectively). MicroZP responded rapidly to phytoplankton bloom formation following the simulated upwelling (t123/24) and showed the strongest increase in abundance in the medium-pCO$_2$ treatment. On t50, however, abundances in the medium-pCO$_2$ treatment decreased again while a pronounced increase in the high-pCO$_2$ was observed (Figure 3G).

Aloricate ciliates, mainly represented by specimens <30 µm, accounted for ~26% on average of total microZP abundances. Ciliate abundance was lower in high-pCO$_2$ during the bloom phase and increased after t35, matching with Chla decrease (Figure 1). An effect of pCO$_2$ on the temporal trend was detected on these ciliate abundances [s(DoE):Treat], indicating a direct link between CO$_2$-enhanced phytoplankton growth and increases in ciliate abundance under high-pCO$_2$ conditions (Table 2 and Figure 3A). Aloricate ciliates were clearly dominant while loricate ciliates, mainly represented by tintinnids, accounted for only ~2.5% of total microZP abundance. No significant pCO$_2$ effect was detected on the temporal trend of loricate ciliates [s(DoE) + Treat], even though abundances were higher at lower pCO$_2$ during the oligotrophic phase of the experiment (Table 2 and Figure 3B). Most dinoflagellates in low- and medium-pCO$_2$ treatments responded to the deep-water addition and followed the Chla build-up and decrease (Figure 1) resulting in an increase in dinoflagellates abundance following the addition (t24), although only some (>25 µm thecate) responded to high-pCO$_2$ at the end of the experiment (Figures 3C–F). Small thecate dinoflagellate abundances (Figure 3C) were higher under high-pCO$_2$ conditions during most of the oligotrophic phase, although highest abundances were recorded under medium-pCO$_2$ treatment toward the end of the experiment [s(DoE):Treat]. The most abundant group within the dinoflagellates were small thecate dinoflagellates. The best fitting model was an interaction of pCO$_2$ and the temporal trend resulting in higher abundances at medium pCO$_2$ in the second half of the experiment [s(DoE):Treat]. Thus, higher abundances of this group were recorded at medium- and low-pCO$_2$ treatments during the bloom, followed by a subsequent decrease in the post-bloom phase (Table 2 and Figure 3D). Large thecate dinoflagellates (Figure 3E) showed a similar trend during the bloom phase, but abundance resulted to be ultimately higher under low-pCO$_2$ toward the end of the experiment [s(DoE):Treat]. Large thecate dinoflagellates (Figure 3F) responded differently than other dinoflagellates, reaching lowest abundance before deep water addition and increasing again when the phytoplankton bloom decayed, independent of the pCO$_2$ treatment [s(DoE) + Treat]. Large dinoflagellates were mainly represented by the genus *Gyrodinium*, comprising ~12% of the total microZP abundances. Small dinoflagellates from the genera *Proteroperidiun* and *Gymnodinium* accounted for ~22 and 20% total microZP abundances, respectively.

The mesozooplankton (mesoZP) community was dominated by copepods and comprised 28 different species or taxonomic groups (see Table 3). Nauplii were counted from the net hauls (>55 µm) and thus included in the mesoZP category, although we are aware that early copepod life stages would in principle belong to microZP when strictly following Sieburth et al.’s (1978).

### Table 2: Zooplankton GAMM analyses.

| Microzooplankton | Model             | Edf  | F   | R$^2$-adj. (%) |
|------------------|-------------------|------|-----|----------------|
| Aloricate ciliates| s(DoE):Treat      | 4.106| 11.26| *** 0.69 72.6  |
| Loricate ciliates | s(DoE) + Treat    | 6.779| 579.2| *** 0.753 79   |
| Athec. dinoflag. <25 µm| s(DoE) | 4.035| 2.327| 0.38 39.3   |
| Thec. dinoflag. <25 µm| s(DoE) | 5.219| 7.227| 0.438 55.1  |
| Athec. dinoflag. >25 µm| s(DoE) | 5.388| 13.191| *** 0.385 79.7 |
| Thec. dinoflag. >25 µm| s(DoE) + Treat | 6.886| 91.33| 0.113 32.2  |
| Total microZP     | s(DoE):Treat      | 3.568| 6.259| * 0.488 42.3  |
| Mesozooplankton   |                   |      |     |                |
| Calanoida         | s(DoE)            | 3.062| 37.07| *** 0.726 81.4 |
| Cyclopoidea       | s(DoE)            | 6.275| 19  | ** 0.289 36.7 |
| Harpacticoidia    | s(DoE)            | 1    | 87.91| *** 0.756 37.9 |
| Pectinostomatoida | s(DoE):Treat      | 5.96 | 7.664| *** 0.382 37.4 |
| Nauplii           | s(DoE):Treat      | 1.372| 5.912| ** 0.329 40.6 |
| O. dioica         | s(DoE)            | 5.739| 3.98 | ** 0.151 13.6 |
| mesoZP total catch| s(DoE):Treat      | 3.596| 5.786| *** 0.571 67.1 |
| Oncaea spp.       |                   |      |     |                |
| Adults            | s(DoE):Treat      | 2.144| 7.533| ** 0.204 9.37 |
| Copepodites       | s(DoE):Treat      | 2.062| 5.914| *** 0.148 17.2 |

Models defined the temporal trend of the abundances alone [s(DoE)], or within an interaction with the pCO$_2$ treatments [s(DoE):Treat]. Only significant values (p-value < 0.05) are presented. DoE, day of experiment; edf, estimated degrees of freedom; Dev. Expl., deviance explained. Significance codes: <0.001 *** 0.001 *** 0.01 *** 0.05.
FIGURE 3 | Microzooplankton abundances during the study period. (A) aloricate ciliates, (B) loricate ciliates, (C) small athecate dinoflagellates (<25 µm), (D) small thecate dinoflagellates (<25 µm), (E) large thecate dinoflagellates (>25 µm), (F) large athecate dinoflagellates (>25 µm), (G) total microZP. Color code: blue = low-PCO2 (M1, M5, and M9), gray = medium-PCO2 (M3, M4, and M7), red = high-PCO2 (M2, M6, and M8). DoE, day of experiment. Note that, for a better visibility of the data, y-axes have been adapted to abundances in each panel. Numbers represent abundances for the respective mesocosm (e.g., 9 for M9). Solid lines = prediction from Generalized Additive Mixed Models (GAMMs) (smoother trends p-value < 0.05); shaded area = confidence interval. Dashed line: t24, deep water addition.

size definition. Total mesoZP catch showed a different temporal trend for each pCO2 treatment [s(DoE):Treat, Table 2]. Averaged mesoZP abundances at the beginning of the experiment (t1) varied between 4730 ± 1202 (low-pCO2), 6023 ± 982 (medium-pCO2), and 5242 ± 369 (high-pCO2) individuals per m$^3$, respectively. Our results showed that mesoZP abundances increased after deep water addition (t24), although this increase was delayed in the high-pCO2 treatment. Highest averaged abundances were recorded for all three treatments on the last sampling day (Figure 4): 23038 ± 9230 individuals per m$^3$ in low-pCO2, 25295 ± 14196 in medium-pCO2 and 24403 ± 10928 in high-pCO2.

Different responses to pCO2 treatments were observed among the studied copepod orders. All copepods, including nauplii, represented ~90% of total mesozooplankton abundances. Calanoid copepods were mainly represented by Clausocalanus spp. and Paracalanus spp. (including e.g., Clausocalanus furcatus, C. arcuicornis, Paracalanus indicus), and accounted for 7–89% (average = 38%) of the total mesozooplankton abundances. An increase in calanoid abundances was detected after deep water addition (t24) in low- and medium-pCO2. Calanoida evolved similarly within the low- and the medium-pCO2 treatments until ~t40. Afterwards, abundances under medium-pCO2 and high-pCO2 treatments increased, resulting in abundances higher than those in low-pCO2 mesocosms on the last sampling day (Figure 4A). Hence, a significant interaction between pCO2 and temporal trend abundances was detected on calanoid abundances [s(DoE):Treat, Table 2] resulting in higher abundances under elevated pCO2 conditions (medium- and high-) during the last two sampling days.

Cyclopoid copepods abundance (Figure 4B) decreased throughout the experiment independently of the treatment.
high-
the experiment, even though they were completely absent in the
catch. We could not detect a
accounted for 0–40% (average = 8%) of total mesozooplankton
was mainly composed by juveniles and
day (p
abundances under the medium-
pabundances under low-
and medium-
pabundances under low-
abundances under high-
abundance was highest in high-
p
A similar trend was observed under medium-
developments, resulting in higher number of immature females under high-
CO
on the temporal trend was detected
on both adults and copepodites [s(DoE):Treat], although no reaction to deep water addition (t24) was observed. Elevated
CO
levels resulted in higher abundances for both adults (only under high-pCO2) and copepodites (under both medium- and high-pCO2 conditions) (Figure 6A and Table 2). There were no significant differences between the numbers of mature females without egg sacks across treatments (Figure 6B). In contrast, the number of females carrying eggs during the experiment was significantly different across treatments. At high-pCO2 there were no egg-carrying females after t24, and a clear increase in numbers could only be detected at medium-pCO2 (Figure 6C).

The appendicularia population — represented by the species Oikopleura dioica— was mainly composed by juveniles and accounted for 0–40% (average = 8%) of total mesozooplankton catch. We could not detect a pCO2 effect on O. dioica during the experiment, even though they were completely absent in the high-pCO2 treatment after deep water addition [s(DoE), Table 2]. This order of copepods was mainly represented by Oithona spp. Harpacticoid copepod abundances (Figure 4C) decreased from the start of the experiment, and no pCO2 effect was detected [s(DoE), Table 2]. This order of copepods was only represented by Microsetella spp. during this experiment. A significant effect of pCO2 on the temporal trend was detected on poecilostomatoid copepods (Figure 4D), mainly represented by Oncaea spp. [s(DoE):Treat, Table 2]. Poecilostomatoids abundance was highest in high-pCO2, increasing until t25 and decreasing gradually afterwards until the end of the experiment. A similar trend was observed under medium-pCO2 while abundances under low-pCO2 conditions did not vary much during the experiment. pCO2 had an effect on the temporal trend of nauplii abundances [s(DoE):Treat, Table 2], which accounted for ~33% of total mesozooplankton abundances. An increase in nauplii abundances under low- and medium-pCO2 conditions was detected after the deep-water addition (t24), with maximum abundances under the medium-pCO2 treatment, while at high-
pCO2 abundances did not increase until the last sampling day (Figure 4E).

Genus Oncaea
A significant effect of pCO2 on the temporal trend was detected on both adults and copepodites [s(DoE):Treat], although no reaction to deep water addition (t24) was observed. Elevated
CO
levels resulted in higher abundances for both adults (only under high-pCO2) and copepodites (under both medium- and high-pCO2 conditions) (Figure 6A and Table 2). A GLMM detected a negative pCO2 effect on females’ sexual development, resulting in higher number of immature females under high-
CO
conditions [s(DoE):Treat, Table 2 and Figure 6]. Approximately 60% of the females in the high-
CO
mesocosms were classified as immature, versus ~30% in medium- and ~36% low-pCO2 treatments over the whole duration of the experiment. The number of immature females at high- and low-pCO2 increased during the experiment while it decreased under medium-pCO2 (Figure 6A). There were no significant differences between the numbers of mature females without egg sacks across treatments (Figure 6B). In contrast, the number of females carrying eggs during the experiment was significantly different across treatments. At high-pCO2 there were no egg-carrying females after t24, and a clear increase in numbers could only be detected at medium-pCO2 (Figure 6C). Thus, we observed a clear negative effect at high-pCO2 on Oncaea
potential offspring (Table 4 and Figure 6), represented by females carrying an egg-sack.

The model revealed a negative effect of the \( pCO_2 \) treatment on the prosome length of mature and immature *Oncaea* females (Table 4 and Figure 7), although this result must be taken with caution due to the relatively weak fit of our models (pseudo-\( R^2 \) ~0.1, Table 4). Pooling together mature and immature individuals, females’ prosome length was slightly shorter under high-\( pCO_2 \) conditions (0.45 ± 0.058 mm) when compared to medium-\( pCO_2 \) (0.56 ± 0.085 mm) and low-\( pCO_2 \) (0.52 ± 0.082 mm).

**Trophic Transfer Efficiency (TTE)**

The simulated upwelling induced a phytoplankton bloom (t25–t35) and amplified differences in succession patterns and food-web structure under high-\( pCO_2 \) conditions (Figure 8).

In the high-\( pCO_2 \) mesocosms the phytoplankton bloom lasted for longer than in the other two treatments, and zooplankton responses were not detected until the bloom decayed (~t48). MicroZP abundance built up only in high-\( pCO_2 \) treatment, while we observed an increase in mesoZP abundances in both medium- and high-\( pCO_2 \) conditions toward the end of the experiment.
Generalized additive mixed models revealed a significant pCO$_2$ effect on the temporal trend of the A:H ratio [s(DoE):Treat, p-value < 0.05, Figure 9]. The model detected highest A:H ratio at the end of the phytoplankton bloom (~t35) in the high-pCO$_2$ treatment. During the post-bloom phase (i.e., after t35), the A:H ratio responded to the differential increase in microZP and mesoZP abundances (see Figures 3G, 4G). Hence, A:H in high-pCO$_2$ decreased faster than in the other two treatments, overlapping low-pCO$_2$ A:H on t50, when highest values corresponded to medium-pCO$_2$ treatment.

**DISCUSSION**

The main objective of this study was to analyze the effect of OA on the zooplankton community from subtropical waters during pre-bloom, bloom and post-bloom conditions. During the oligotrophic phase of this experiment we could not detect important differences in total zooplankton abundances between the treatments (Figures 3G, 4G). However, after the simulated upwelling, the zooplankton community under high-pCO$_2$ conditions evolved significantly differently compared to the low- and medium-pCO$_2$ conditions (Figure 2). Overall, higher zooplankton abundances were observed at elevated pCO$_2$ conditions (medium- and high-) in the post-bloom phase. This result matches with a previous KOSMOS study in coastal mesotrophic conditions (Bach et al., 2016b) where certain groups of consumers capitalized on CO$_2$-enhanced phytoplankton biomass, resulting in higher zooplankton abundances under moderate IPCC end-of-century pCO$_2$ scenarios (RCP6.0) (Horn et al., 2016; Algueró-Muñiz et al., 2017). However, unexpectedly, both microZP and mesoZP abundances (Figures 3G, 4G) increased much later in the experiment under high-pCO$_2$ than under medium- and low-pCO$_2$. In the following, we will discuss the differences in zooplankton densities as well as the timing of bloom development.

**pCO$_2$ Effects on Zooplankton Densities**

As reported by other authors (e.g., Isari et al., 2015), responses to OA are not only dependent on species-specific sensitivities, but, much more importantly, depend on CO$_2$ effects on the community and the trophic interactions taking place in a species’ natural habitat. In fact, most of the reported effects of OA on planktonic communities need to be attributed to these community effects, as many indicate a positive effect of OA (or rather carbon availability) on the plankton (Algueró-Muñiz et al., 2017; Taucher et al., 2017b). The temporal trends in major microZP groups (aloricate ciliates, small dinoflagellates) and Calanoida (Figures 3, 4, respectively) were most likely triggered by the food supply for microZP combined with the preference of most copepods for heterotrophic protists (Suzuki et al., 1999; Turner, 2004). As expected, picoplanktonic phytoplankton were a dominant component during the oligotrophic phase and large chain-forming diatoms dominated during the nutrient induced bloom (Taucher et al., 2017a). Diatoms are an ideal food source for larger mesoZP and this direct consumption of mesoZP on phytoplankton might have caused a release of microZP from grazing pressure after the deep-water addition.

The initial microZP abundance, as well as their taxonomic composition, was in agreement with those reported previously for the same area (Ojeda, 1998; Schnöcker et al., 2014). During the post-bloom phase, microZP was dominated by dinoflagellates <25 $\mu$m and aloricate ciliates. Ciliates and dinoflagellates are the main grazers on phytoplankton in marine systems, especially oligotrophic ones and also contribute to a large part to the copepod diets (Calbet, 2008). This is attributed to their appropriate size and high nutritional quality of microZP relative to phytoplankton (Stoecker and Capuzzo, 1990) and the dominance of small-sized phytoplankton in oligotrophic systems which is outside the food spectrum of many mesozooplankters (Kleppel, 1993). Previous OA studies reported a tolerance of microZP communities toward high CO$_2$ concentrations, or only...
very subtle changes in the community (Suffrian et al., 2008; Aberle et al., 2013; Horn et al., 2016; Lischka et al., 2017) while other studies showed detrimental (Calbet et al., 2014) or even positive effects (Rose et al., 2009). In this study, an increase in aloricate ciliate abundances was observed in all treatments in response to the deep water-induced phytoplankton bloom, although the increase showed a considerable time-lag relative to the phytoplankton bloom, especially at high CO\textsubscript{2} conditions. In contrast to aloricate ciliates, loricate ciliates played a minor role in this experiment and showed only a very small peak during the oligotrophic phase. Loricate ciliates started to decline after t10 and were virtually absent after deep water addition (see Figure 3).

For dinoflagellates, especially small-sized athecates, we expected a positive effect of high-pCO\textsubscript{2} levels due to findings from previous OA studies conducted in oligotrophic (Sala et al., 2015) and eutrophic regions (Horn et al., 2016). During the oligotrophic phase of the experiment, this expectation was confirmed since higher abundances of small athecate dinoflagellates at high-pCO\textsubscript{2} were also found in our study. However, after deep water addition overall dinoflagellate abundances were higher at low- and medium-pCO\textsubscript{2} conditions. Unlike ciliates, heterotrophic dinoflagellates are known to feed on phytoplankton of various sizes up to several times larger than their body size and have been shown to prey on bloom-forming diatoms including taxa as, e.g., *Thalassiosira* (Sherr and Sherr, 2007). The abundance
of diatoms, however, was higher at high-pCO2 compared to the low- and medium-pCO2 conditions (Taucher et al., 2017a). Thus, the effect of high pCO2 on dinoflagellates was most likely an indirect one based on changes in the phytoplankton community and more precisely, on prey edibility (see section “pCO2 Effects on Zooplankton Bloom Timing”).

Calanoida were positively affected by medium- and high-pCO2, although the trend was only visible during the last two sampling days. These results match with previous outcomes described for copepodes and adult Pseudocalanus acusipes in eutrophic waters and pCO2 levels of ∼760 µatm (Algueró-Muñiz et al., 2017; Taucher et al., 2017b), suggesting a benefit of realistic end-of-century pCO2 levels on calanoid copepodes through higher food availability. Small planktonic copepodes are dominant in the plankton communities in many parts of the world’s oceans and therefore important members of pelagic food webs (Turner, 2004). Thus, a positive pCO2 effect on these major zooplankton components could have a crucial impact on the transfer of energy to higher trophic levels thus affecting, e.g., future fisheries (Moyano et al., 2009; Swat et al., 2018).

C copepod species that do not exhibit vertical migration behavior are considered as evolutionarily less exposed to high-pCO2 levels compared to other copepodes, and typically more sensitive to OA (Fitzter et al., 2012; Lewis et al., 2013). Accordingly, we expected cyclopoid (dominated by Oithona) and harpacticoid copepodes (dominated by Microsetella) to show lower abundances under elevated pCO2 conditions as neither species shows diel migration (Maar et al., 2006). However, during this experiment, elevated pCO2 did not cause a significant effect on Cyclopoida and Harpacticoida abundances, according to the GAMM analyses (Figures 4B,C). The reason for the decay in Cyclopoida and Harpacticoida abundances is not entirely clear but could be due to the distribution of the copepodes in the water column, closer to the mesocosm sediment traps. Such a loss through sedimentation was previously observed in other mesocosm experiments during periods of low food availability (Bach et al., 2016a; Algueró-Muñiz et al., 2017). Oithona and Microsetella have been reported to concentrate on marine snow (Ohtsuka et al., 1993; Koski et al., 2005) and during the present experiment, the cumulative flux of particulate organic matter to the sediment traps increased after deep water addition (Stange et al., 2018). This might have promoted a downward migration of Microsetella — already from the beginning of the experiment on — to enhance their feeding on sinking material, preventing us to sample them in the net hauls.

### pCO2 Effects on Zooplankton Bloom Timing

Where the observed response of the zooplankton densities is in line with previously published results, the differences in timing of the blooms were rather unexpected. In fact, zooplankton density increases after the simulated upwelling under high-pCO2 treatment were much slower than under the low and medium treatments. The most probable explanation for this observation lies in the differences in taxonomy of the phytoplankton responding to the nutrient addition of the upwelling. The phytoplankton bloom in the high-pCO2 mesocosms (M2 and M8) was dominated by Vicicitus globosus (Dictyotephyceae) which bloomed only in the high-pCO2 mesocosms from t25 until t47 (Riebesell et al., 2018). Harmful or non-edible for zooplankton, it seems likely that V. globosus caused adverse effects on the plankton community. MicroZP as potential grazers were most likely affected by the inadequacy of the available phytoplankton food (Chang, 2015), thus preventing the subsequent increase in mesoZP abundances. This is even more likely considering that once the phytoplankton bloom ceased in the high-pCO2 treatments, microZP started to increase in numbers at a time point when they were already decreasing at low and medium-pCO2. The tolerance to harmful algae has previously been described for copepod species closely related to those recorded in the mesocosms such as Paracalanus parvus (tolerant to Chattonella antiqua) and Oncaea venusta (tolerant to Karenia brevis) (Turner and Tester, 1989). Although Paracalanus sp. nauplii may exhibit adverse effects from feeding upon Alexandrium tamiaiyavanichii (Silva et al., 2013), we did not detect negative effects on nauplii abundances when relating them to the harmful algae abundance, but a delay in the reaction time likewise in aloricate ciliates, dinoflagellates and calanoid copepods. Accordingly, we based our conclusions for copepods on temporal trends and pCO2 treatments rather than on possible effects of inedible/harmful food items. Our results suggest that copepods reacted to the different pCO2 levels only after their preferred prey [i.e., heterotrophic protists (Turner, 2004)] reacted...
to the stimulated bloom, thus highlighting the importance of microZP in bloom situations within oligotrophic ecosystems (Calbet and Alcaraz, 2007; Calbet, 2008).

**pCO₂ Effects on *Oncaea* and *O. dioica* Interactions in Pre- and Post-bloom Conditions**

*Oncaea*’s feeding strategies are associated with surface materials, such as fine particles, bacteria, or the tegument fluid of gelatinous zooplankton (*Sagitta* spp., *Oikopleura* spp. and *Salpa* spp.) (Go et al., 1998). During this study, abundances of *Oncaea* spp. and *O. dioica* were inversely correlated, as previously observed at other study sites (Itoh et al., 2014). *Oncaea* was positively affected by pCO₂, recording higher abundances under medium- and high-pCO₂ treatments from (approximately) the beginning of the experiment until the end of the phytoplankton bloom, on t35 (Figure 4D). The *O. dioica* trends showed some similarities with other studies at elevated nutrient concentrations (Troedsson et al., 2013). We did not detect a significant pCO₂ effect on *O. dioica* when considering the whole experimental period (Figure 4F), what agrees with previous results from Troedsson et al. (2013) and Winder et al. (2017) on the tolerance of appendicularians to OA.

After deep water addition, we observed that *O. dioica* completely disappeared under high-pCO₂ while *Oncaea* abundances were higher than in the other two treatments, suggesting a top-down control of *Oncaea* on *O. dioica* abundances. Hence, the fact that during the last sampling days *Oncaea* spp. abundances decayed in the high-pCO₂ treatment might reflect the scarcity of *O. dioica* as food resource (Go et al., 1998).

Concerning the condition of *Oncaea* females (Figures 6, 7), we observed smaller individuals, as well as a higher number of immature females and a lower number of egg-carrying mature females in the high-pCO₂ treatment. These results are in line with previous studies in calanoid copepods which also observed a decrease in copepod size (Garzke et al., 2015) and fecundity loss (Thor and Dupont, 2015) caused by increased CO₂ levels. However, unlike the major sensitivities to OA previously described for early life stages of calanoid copepods (Pedersen et al., 2013; Algueró-Muníz et al., 2017), we did not observe a stronger pCO₂ effect on *Oncaea* copepodites than on adults (Figure 5), suggesting differences between poecilostomatoids and calanoids in their offspring responses to expected future OA levels. We conclude that the negative pCO₂ effect detected on *Oncaea* females’ reproductive output might affect food web interactions in the long term in those tropical and subtropical communities dominated by this species (e.g., Böttger-Schnack, 1994), especially in those where oncaeid copepods are the main prey for larvae and juvenile fish (Itoh et al., 2014). The lack of published OA research on *Oncaea* spp. (Poecilostomatoida) makes the analysis presented here of special relevance and calls for multigenerational OA studies on this species.

**Influence of OA on the Transfer of Energy Within the Plankton Community**

As discussed above, community effects and trophic interactions can alter sensitivities to OA (Rossoll et al., 2013), which in turn may have an effect on the efficiency of the food web (Calbet et al., 2014; Cripps et al., 2016; Algueró-Muníz et al., 2017). The autotrophic community was expected to experience an increase in biomass (Gismervik et al., 2002) responding to the nutrient input created by the deep water addition. Thus, a significant effect of CO₂ on plankton succession was observed after deep water addition (Taucher et al., 2017a), suggesting that phytoplankton blooms are boosted at elevated pCO₂. This situation could in turn cause a CO₂-dependant reduction in trophic efficiency after deep water addition, due to the limited capacity of micro- and mesoZP grazers to exploit the elevated phytoplankton productivity (Calbet et al., 2014). Accordingly, the A:H ratio (autotrophy/heterotrophy) proposed as a proxy for the trophic efficiency of the system was highest during the phytoplankton built-up at high-pCO₂. TTE decreased in all three pCO₂ treatments during the phytoplankton bloom (t25–t35), and lowest TTE was detected under high-pCO₂ conditions, likely because under these conditions microZP was limited by inadequate food items thus leading to a delayed response of microZP after phytoplankton bloom initiation, consequently affecting mesoZP production. These results are in line with previous studies (Calbet et al., 2014; Cripps et al., 2016) which point at a more-autotrophic and less efficient food web under higher pCO₂ conditions when the consumers mismatch the phytoplankton bloom (Edwards and Richardson, 2004; Calbet et al., 2014), as observed during this experiment until t~40. Similarly than Calbet et al. (2014) and Cripps et al. (2016), we did not account for nanoplankton to estimate TTE in our study, what might have incurred in an underestimation of the system efficiency when considering phytoplankton as the only carbon source for zooplankton. The increase in calanoid copepod abundance observed in both high- and medium-pCO₂ treatments toward the end of the experiment points at pCO₂-induced effects under nutrient-replete conditions, which could travel up the food web reaching secondary consumers, as previously observed in eutrophic systems (Algueró-Muníz, 2017; Swat et al., 2018). In case of the medium-pCO₂ treatment, an increased grazing pressure of copepods (Calanoida) on dinoflagellates could explain that TTE in medium-pCO₂ was lower than in the other two treatments after the phytoplankton bloom. Our results thus suggest that pCO₂ effects on plankton succession depend on the coupling of the phytoplankton bloom with microZP and mesoZP grazers, ultimately affecting the development of the plankton community and the energy transfer efficiency of the system.

Based on this study, end-of-century pCO₂ levels are not expected to cause major effects on subtropical zooplankton communities during oligotrophic phases. However, during bloom and post-bloom conditions, elevated pCO₂ might promote higher zooplankton abundances by bottom-up effects of CO₂-enhanced primary production. Hence, pCO₂-fertilized phytoplankton productivity would reach grazers through trophic cascades, which might in turn be disrupted when CO₂ benefits harmful algae. These pCO₂ effects on plankton communities could be specially relevant in oligotrophic environments with short bloom periods such as the Canary Islands, where zooplankton biomass has been shown to have direct implications.
on larval abundance in different fish species during late winter bloom (Moyano et al., 2009). Therefore, a positive effect of pCO₂ on zooplankton abundance after a bloom event might eventually benefit larval recruitment, and consequently have an effect on future fisheries.

DATA AVAILABILITY
All zooplankton abundance data are archived at the PANGEA data library, https://doi.pangaea.de/10.1594/PANGAEA.887283.

ETHICS STATEMENT
No specific permission was required for activities related to field sampling. The field location was not privately owned or protected, and neither regulated animals, endangered nor protected species were involved.

AUTHOR CONTRIBUTIONS
UR, MA-M, HH, NA, LB, WG, EA, and MB conceived and designed the experiments. MA-M, HH, CS, LB, WG, and UR performed the experiments. MA-M, HH, SA-F, LB, WG, and EA analyzed the data. MA-M and HH contributed equally to this work. MA-M wrote this paper with input from all co-authors.

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