



# Photochemical vs. Bacterial Control of H<sub>2</sub>O<sub>2</sub> Concentration Across a pCO<sub>2</sub> Gradient Mesocosm Experiment in the Subtropical North Atlantic

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In the surface ocean, microorganisms are both a source of extracellular H<sub>2</sub>O<sub>2</sub> and, via the production of H<sub>2</sub>O<sub>2</sub> destroying enzymes, also one of the main H<sub>2</sub>O<sub>2</sub> sinks. Within microbial communities, H<sub>2</sub>O<sub>2</sub> sources and sinks may be unevenly distributed and thus microbial community structure could influence ambient extracellular H<sub>2</sub>O<sub>2</sub> concentrations. Yet the biogeochemical cycling of H<sub>2</sub>O<sub>2</sub> and other reactive oxygen species (ROS) is rarely investigated at the community level. Here, we present a time series of H<sub>2</sub>O<sub>2</sub> concentrations during a 28-day mesocosm experiment where a pCO<sub>2</sub> gradient (400–1,450 μatm) was applied to subtropical North Atlantic waters. Pronounced changes in H<sub>2</sub>O<sub>2</sub> concentration were observed over the duration of the experiment. Initially H<sub>2</sub>O<sub>2</sub> concentrations in all mesocosms were strongly correlated with surface H<sub>2</sub>O<sub>2</sub> concentrations in ambient seawaters outside the mesocosms which ranged from 20 to 92 nM over the experiment duration (Spearman Rank Coefficients 0.79–0.93, *p*-values < 0.001–0.015). After approximately 9 days of incubation however, H<sub>2</sub>O<sub>2</sub> concentrations had increased across all mesocosms, later reaching >300 nM in some mesocosms (2–6 fold higher than ambient seawaters). The correlation with ambient H<sub>2</sub>O<sub>2</sub> was then no longer significant (*p* > 0.05) in all treatments. Furthermore, changes in H<sub>2</sub>O<sub>2</sub> could not be correlated with inter-day changes in integrated irradiance. Yet H<sub>2</sub>O<sub>2</sub> concentrations in most mesocosms were inversely correlated with bacterial abundance (negative Spearman Rank Coefficients ranging 0.59–0.94, *p*-values < 0.001–0.03). Our results therefore suggest that ambient H<sub>2</sub>O<sub>2</sub> concentration can be influenced by microbial community structure with shifts toward high bacterial abundance correlated with low extracellular H<sub>2</sub>O<sub>2</sub> concentrations. We also infer that the nature of mesocosm experiment design, i.e., the enclosure of water within open containers at the ocean surface, can strongly influence extracellular H<sub>2</sub>O<sub>2</sub> concentrations. This has potential chemical and biological implications during incubation experiments due to the role of H<sub>2</sub>O<sub>2</sub> as both a stressor to microbial functioning and a reactive component involved in the cycling of numerous chemical species including, for example, trace metals and haloalkanes.

**Keywords:** hydrogen peroxide, H<sub>2</sub>O<sub>2</sub>, mesocosm, Atlantic, pCO<sub>2</sub>

## INTRODUCTION

Reactive oxygen species (ROS) are ubiquitous in sunlit natural surface waters (Van Baalen and Marler, 1966; Moore et al., 1993; Miller and Kester, 1994). The most extensively measured ROS in the marine environment, H<sub>2</sub>O<sub>2</sub>, is present in the surface mixed layer at concentrations on the order of 10–100 nM (Price et al., 1998; Yuan and Shiller, 2001; Gerringa et al., 2004). In the surface ocean, H<sub>2</sub>O<sub>2</sub> is known to be mainly produced by photochemistry (Fujiwara et al., 1993; Micinski et al., 1993) with a poorly quantified fraction produced via biochemical processes (Palenik et al., 1987; Croot et al., 2005; Milne et al., 2009). Biochemical processes are also used to explain H<sub>2</sub>O<sub>2</sub> production in the dark (Palenik and Morel, 1988; Moffett and Zafiriou, 1990; Vermilyea et al., 2010).

H<sub>2</sub>O<sub>2</sub> can cross cell membranes and cause a wide range of cellular damage, a process generically referred to as oxidative stress (Seaver and Imlay, 2001; Lesser, 2006; Imlay, 2008). Extracellular H<sub>2</sub>O<sub>2</sub> is however not generally considered to be a major constraint on cellular growth under natural conditions in the marine environment because most microorganisms are thought to produce catalase and peroxidase enzymes which control H<sub>2</sub>O<sub>2</sub> decomposition rates in the surface ocean (Moffett and Zafiriou, 1990; Petasne and Zika, 1997). However, recent work has challenged the assumption that extracellular H<sub>2</sub>O<sub>2</sub> at nanomolar concentrations does not negatively influence cellular metabolism in surface seawater. The susceptibility of a range of marine microorganisms to H<sub>2</sub>O<sub>2</sub> (Bogosian et al., 2000; Morris et al., 2011) and measurable effects on primary metabolism at extracellular H<sub>2</sub>O<sub>2</sub> concentrations within the range of surface marine concentrations (Morris et al., 2011; Baltar et al., 2013) have been demonstrated. Furthermore, it has been suggested that microbes sensitive to H<sub>2</sub>O<sub>2</sub> may not be cultivable under normal laboratory conditions (Morris et al., 2008, 2011), which may have severely biased our historical understanding of ROS interactions with marine microorganisms.

H<sub>2</sub>O<sub>2</sub> production rates, decomposition rates, and effects on cellular functioning may vary widely at the species level (Palenik et al., 1987; Baltar et al., 2013). Furthermore, cross-group interactions may be important in regulating ambient extracellular H<sub>2</sub>O<sub>2</sub> concentrations (Morris et al., 2011). Investigations at the community level are therefore required in order to comprehensively understand the interaction between biological processes and ROS in seawater. Mesocosm studies are one approach by which this could be achieved, yet rapid decay rates mean that investigating ROS during an offshore mesocosm experiment, with a setup for example as per Taucher (2017, this Research Topic), would be logistically challenging. A mesocosm experiment on a smaller and more accessible scale, with a similar pCO<sub>2</sub> gradient and timespan, was therefore conducted in the subtropical waters of Gran Canaria in March 2016. Our objective was to compare changes in ROS and other short-lived reactive species over the timescale of an induced phytoplankton bloom across a broad pCO<sub>2</sub> gradient. The pCO<sub>2</sub> gradient was designed to encompass pCO<sub>2</sub> under all plausible future climate scenarios until 2100 (IPCC Working Group 1, 2014) with the addition of some higher end-members to investigate potential thresholds

with respect to CO<sub>2</sub>-sensitive ecological and biogeochemical processes.

## MATERIALS AND METHODS

### Mesocosm Design

The mesocosm study, conducted in Taliarte Harbor, Gran Canaria in March 2016, used eight thermoplastic polyurethane bags with a 2 m diameter, a depth of ~3 m, a starting volume of ~8,000 L, and no lid or screen on top. The bags were mounted on a buoyant frame which was allowed to drift ~2–3 m away from a sampling jetty. After filling with ambient seawater (on 1 March, experiment day –4), pumped from outside Taliarte Harbor (depth 15 m), the mesocosms were allowed to function without nutrient addition for 21 days (**Supplementary Figure 1**). A pCO<sub>2</sub> gradient across the eight mesocosms was induced on day 0 by the addition of varying volumes of filtered, CO<sub>2</sub> saturated seawater using a custom made “spider” distribution device described by Riebesell et al. (2013). The pCO<sub>2</sub> gradient (400–1,450 μatm) was designed to be similar to that used during the offshore KOSMOS mesocosm experiment conducted in Gando Bay, Gran Canaria in September/October 2014 (Taucher, 2017, this Research Topic). A further top-up of pCO<sub>2</sub> saturated seawater was then made to the mesocosms as necessary to maintain the pCO<sub>2</sub> gradient (**Supplementary Figure 1**) following CO<sub>2</sub> outgassing. The precise volume of each mesocosm was determined (on day 18) by measuring salinity before, and after, the addition of 40 L freshwater to each mesocosm, similar to Czerny et al. (2013). A single macronutrient addition (3.1 μM nitrate, 1.5 μM silicate, and 0.2 μM phosphate) was then made (after day 18 sampling).

### Analysis

H<sub>2</sub>O<sub>2</sub> samples were collected in opaque 125 mL high density polyethylene (HDPE) bottles (Nalgene) which were pre-cleaned (1 day soak in detergent, 1 week soak in 1 M HCl, three rinses with de-ionized water; 18.2 MΩ·cm, Milli-Q, Millipore) and dried in a laminar flow hood prior to use. Sample bottles were rinsed once with seawater and filled with no headspace by gently submerging the bottles within the mesocosms. Chlorophyll a, bacterial abundance, chromophoric dissolved organic matter (CDOM) and macronutrient concentrations were determined from depth integrated water samples collected using 2.5 m long custom-made samplers with an internal volume of ~10 L. These samplers were constructed from polypropylene tubing with valves at both ends. After filling, by submerging into the mesocosms and closing the valves, the samplers were removed and gently inverted to facilitate mixing. Samplers were then slowly drained through 1 cm diameter silicone tubing into pre-rinsed (de-ionized water and then mesocosm water) 25 L transparent HDPE containers, which were then transferred to shaded boxes and moved to a dark, refrigerated room for subsampling. Analysis of chlorophyll a and macronutrients then began immediately.

Chlorophyll a was measured by fluorometry as per Welschmeyer (1994) and macronutrient concentrations (nitrate + nitrite, phosphate, silicate) were determined by colorimetry as per Hansen and Koroleff (1999). CDOM absorption spectra

were measured with a 100 cm, 250  $\mu$ L capillary (LPC100CM) connected via an optical fiber to a light source (DH2000BAC) and a USB2000+UV-VIS ES detector (Ocean Optics). The system was controlled using Spectra-suite software (Ocean Optics). Samples were injected into the capillary with a peristaltic pump at a flow rate of 1 mL min<sup>-1</sup>. Relative molecular weight was estimated from CDOM absorption by deriving the slope ratio ( $S_R$ ) as the ratio of the slope of the shorter wavelength region (275–295 nm) to that of the longer wavelength region (350–400 nm; Helms et al., 2008). The spectral slopes were calculated from the linear regression of the log-transformed absorption spectra.

H<sub>2</sub>O<sub>2</sub> was always analyzed within 1 h of collection via flow injection analysis (FIA) using the Co(II) catalyzed oxidation of luminol (Yuan and Shiller, 1999). A FIA system was assembled and operated exactly as per Hopwood et al. (2017) resulting in a detection limit of <1 nM. Calibrations were run daily, and with every new reagent batch, by at least six standard additions of diluted H<sub>2</sub>O<sub>2</sub> (TraceSelect, Fluka) to aged (stored at room temperature for >48 h) seawater (unfiltered). The stability of H<sub>2</sub>O<sub>2</sub> stock solution was checked by measuring absorbance at 240 nm using a quartz 10 cm cell and a USB4000 spectrometer (Ocean Optics). H<sub>2</sub>O<sub>2</sub> (TraceSelect, Fluka) was sequentially diluted weekly to create stock solutions of 100 mM and 100  $\mu$ M using de-ionized water. With respect to the measurement of chemical species other than H<sub>2</sub>O<sub>2</sub>, the H<sub>2</sub>O<sub>2</sub> luminol based FIA method is expected to be robust for measurements in oxygenated surface seawater (Yuan and Shiller, 1999, 2004).

Bacterial counts were obtained on depth integrated water samples. Two mL water samples were fixed with 1% paraformaldehyde (final concentration) and stored at -80°C until analysis. Samples were analyzed by flow cytometry (FACSCalibur, Becton Dickinson), with a 15 mW laser set to excite at 488 nm (Gasol and del Giorgio, 2000). Subsamples (400  $\mu$ L) for the determination of heterotrophic bacteria were stained with the fluorochrome SybrGreen-I (4  $\mu$ L) at room temperature for 20 min and run at a flow rate of 16  $\mu$ L min<sup>-1</sup>. Cells were enumerated in a bivariate plot of 90° light scatter and green fluorescence. Molecular Probes latex beads (1  $\mu$ m) were used as internal standards.

## Ancillary Measurements

A CTD cast (using a CTD60M, Sea and Sun Technology) was conducted every sample collection day in all mesocosms. Additionally, a CTD cast was conducted on the sampling jetty adjacent to the mesocosms where a H<sub>2</sub>O<sub>2</sub> sample in ambient seawater was also collected alongside every mesocosm sampling event. Photosynthetically active radiation (PAR) data was obtained from an ELDONET (Häder and Lebert, 2006) monitoring site located <200 m from the mesocosm jetty. Complete diurnal light profiles were available for experiment day 7 onwards, excluding days 12 and 13.

The diurnal change in H<sub>2</sub>O<sub>2</sub> concentration in ambient seawater, and inside two mesocosms, was monitored by setting up FIA equipment on the mesocosm sampling jetty with a PTFE sampling line weighted to float ~10 cm below the water surface. Seawater was pumped continuously using a peristaltic pump

(MiniPuls 3, Gilson) with a time delay between water inflow and analysis of ~60–120 s. The sample line was used without a filter and visually inspected regularly for blockage. Calibration was undertaken three times in every 24 h period by 6–8 standard additions of H<sub>2</sub>O<sub>2</sub> into aged (>48 h in the dark) seawater. A coiled 3 m PTFE sample line, which could be extended to the bottom of the mesocosms, was used to determine whether H<sub>2</sub>O<sub>2</sub> was vertically homogenous within the mesocosm bags. During the diurnal experiment monitoring ambient seawater, salinity was measured regularly (<3 h intervals) using a LF 325 conductivity meter (WTW) which was calibrated before use with a KCl solution.

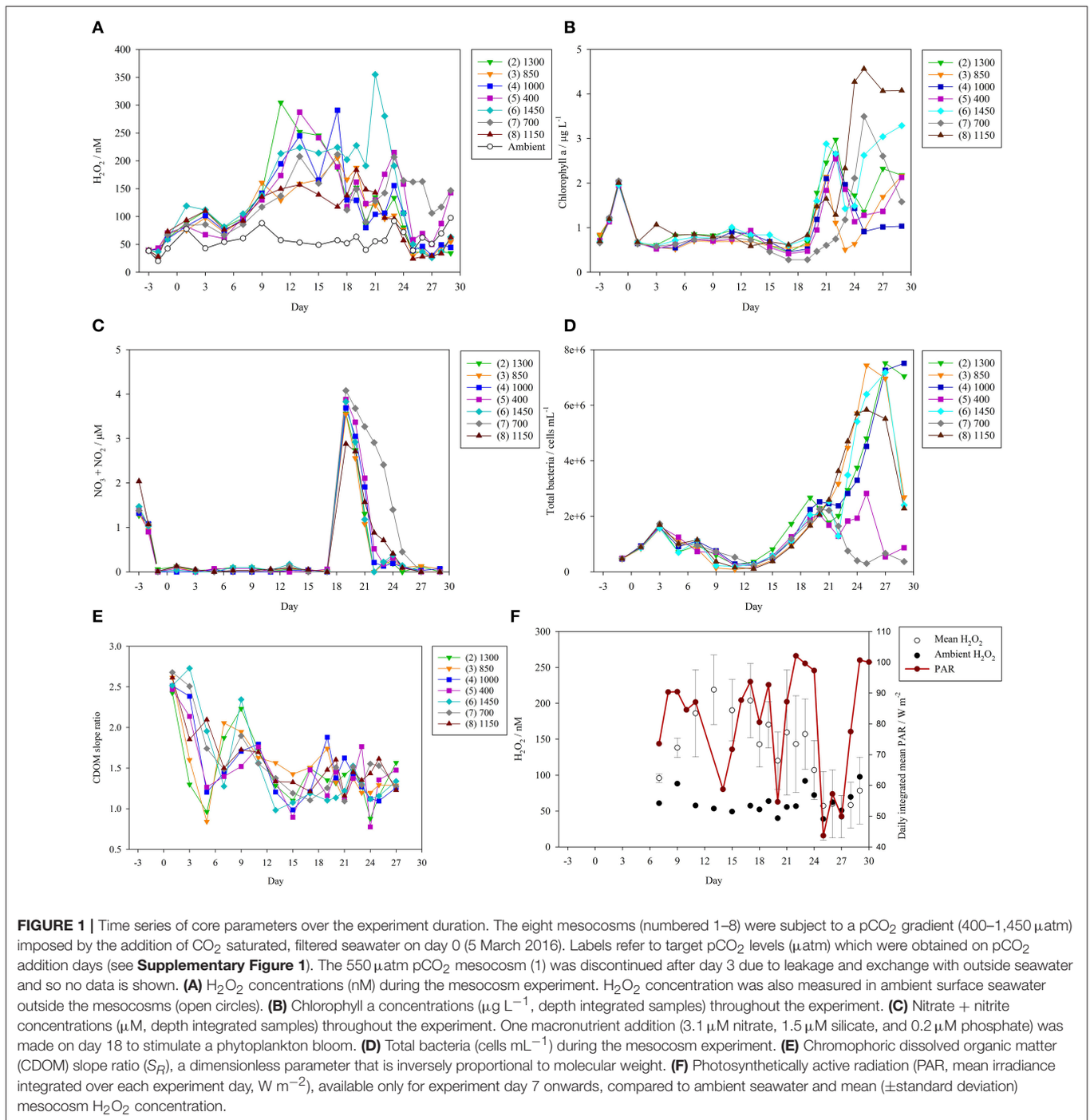
During rain events, rainwater was collected by deploying open low density polyethylene (LDPE) bags adjacent to the mesocosms. Rainwater was diluted prior to analysis by spiking 200  $\mu$ L unfiltered rainwater into 50 mL aged seawater with the H<sub>2</sub>O<sub>2</sub> concentration in seawater measured before, and after, the rainwater spike.

## RESULTS

### Mesocosm Time Series

Time series for the core parameters discussed herein are included in **Supplementary Datasheet 1**. Initial measurements in the mesocosms (2 March 2016, experiment day -3, after filling the mesocosms—but before any treatments were applied) verified that the enclosed waters were close to identical with respect to H<sub>2</sub>O<sub>2</sub> concentration. Differences between the eight enclosures (range 38.3–40.9 nM, mean 39.3  $\pm$  0.8 nM) were small compared to the detection limit of the analytical method (<1 nM; Croot et al., 2004; Hopwood et al., 2017) and the standard deviation of quadruple measurements of H<sub>2</sub>O<sub>2</sub> in seawater collected on the sample jetty (mean 3.3 nM over the experiment duration). Depth profiles within a mesocosm bag (~3 m depth) were conducted on day 25 mid-afternoon, when any stratification was expected to be maximal, and verified that H<sub>2</sub>O<sub>2</sub> concentration was relatively well-mixed with only a slight vertical gradient (increasing from 43 to 51 nM bottom-top, mean 47  $\pm$  3.5 nM). On day 12, surface seawater collected using a small inflatable boat showed a range of H<sub>2</sub>O<sub>2</sub> concentrations at different locations around Taliarte Harbor (39  $\pm$  9.9 nM,  $n$  = 4), outside the harbor within 150 m of the coastline (63  $\pm$  1.8 nM,  $n$  = 3) and outside the harbor >400 m offshore (38  $\pm$  5.6 nM,  $n$  = 4). Ambient seawater concentrations within the harbor (used to determine a background concentration for comparison with the mesocosms) were therefore similar to those found in near-shore waters.

The imposition of a large pCO<sub>2</sub> gradient across the mesocosm bags after sampling on day 0 had no clear prolonged effect on observed H<sub>2</sub>O<sub>2</sub> concentrations (**Figure 1A**). Temporal trends in H<sub>2</sub>O<sub>2</sub> concentration were relatively similar in all mesocosms until day 9 of the experiment with the standard deviation ranging from 0.7 to 13 nM. Data for mesocosm 1 are shown only until day 3, after which exchange with surrounding seawater occurred following leakage, and monitoring was thus discontinued. After day 9 the enclosed waters diverged and some mesocosms experienced swings to very high (>300 nM) H<sub>2</sub>O<sub>2</sub> concentrations compared to ambient water (**Figure 1A**).



Over the duration of the mesocosm experiment, ambient surface seawater H<sub>2</sub>O<sub>2</sub> concentrations ranged from only 20–92 nM.

The initial nitrate concentration present in ambient seawater was depleted rapidly after filling of the mesocosms (**Figure 1C**). Nitrate + nitrite fell to <0.1 µM in all mesocosms by day -1. Nitrate concentration then remained depleted until the macronutrient addition on day 18 (macronutrient addition occurred after sampling on day 18). Correspondingly, a small

peak in chlorophyll a was observed in all mesocosms on day -1 (**Figure 1B**). Chlorophyll a then declined to low concentrations until a larger peak following the macronutrient addition on day 18. Maximum chlorophyll a was then observed in most mesocosms on days 21–22.

A notable exception to the general trend in mesocosm H<sub>2</sub>O<sub>2</sub> under post-bloom conditions was mesocosm 7 (700 µatm pCO<sub>2</sub>). On days 25–27, H<sub>2</sub>O<sub>2</sub> concentrations in the majority of mesocosms dropped below those measured in ambient

seawater, but H<sub>2</sub>O<sub>2</sub> in mesocosm 7 always remained >100 nM (**Figure 1A**). Mesocosm 7 was also anomalous with respect to the bloom development. In mesocosm 7 only it appeared that grazing may have impeded bloom development after nutrient addition (this was consistent with higher meso-zooplankton abundances in mesocosm 7, data not shown), as evidenced by a noticeably slower decline in nitrate concentration (**Figure 1C**) and a late peak in chlorophyll a (**Figure 1B**).

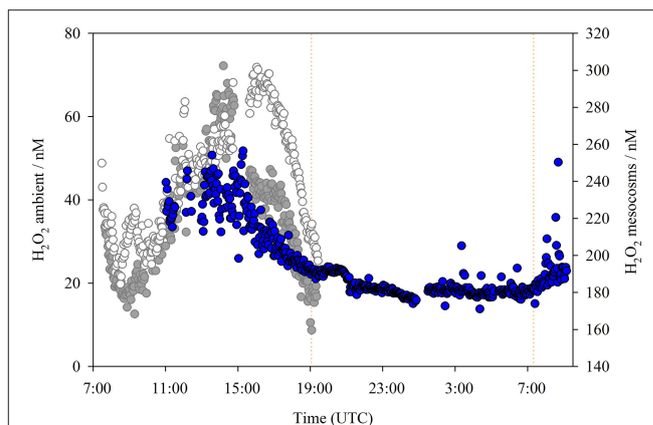
Bacterial abundance was similar in all mesocosms until after nutrient addition (**Figure 1D**). Unlike chlorophyll a however, an increase in bacterial abundance was evident in all treatments prior to the nutrient addition on day 18. A small dip in bacterial abundance was then evident in most mesocosms between days 19 and 22. Under post-bloom conditions bacterial abundance was lowest in the anomalous treatment (700 μatm) and the 400 μatm pCO<sub>2</sub> mesocosm with notably elevated abundances in all pCO<sub>2</sub> enriched treatments. The observed trend in bacterial abundance can be interpreted as resulting from grazing pressure, and enhanced growth rates post-nutrient addition. Bacterial abundance pre-nutrient addition was inversely related to nanoeukaryotes' abundance due to grazing pressure. The increase in bacterial abundance from day 13 reflected a decline in the abundance of nanoeukaryotes. After nutrient addition (day 18), grazing on bacteria was probably considerable, but bacterial growth was enhanced sufficiently to overcome grazing pressure (except in the anomalous treatment). An in depth discussion of phytoplankton community structure over the experiment duration will be presented in a companion text.

The trend in CDOM over the experiment duration was similar across all pCO<sub>2</sub> treatments. The generally higher S<sub>R</sub> at the start of the experiment (**Figure 1E**) corresponds to lower mean molecular weight and suggests an overall increase in CDOM molecular weight over the experiment duration. Photochemical bleaching would be expected to have had the opposite effect; to have produced low molecular weight CDOM from high molecular weight CDOM (i.e., to have increased S<sub>R</sub>). Thus, the overall trend suggests that bacterial production of high molecular weight CDOM exceeded the rate of photochemical bleaching. Only between experiment days 7 and 13 was a sustained increase in S<sub>R</sub> evident across most mesocosms and this corresponded to a temporary decline in bacterial abundance (**Figure 1D**).

Photosynthetically active radiation (PAR) data was available from day 7 until the end of the mesocosm experiment (**Figure 1F**) from a recording site close to the mesocosm jetty (<200 m displacement). Integrated daily PAR was subject to notable variations over the duration of the experiment ranging from a mean irradiance of 44 (day 25) to 102 (day 22) W m<sup>-2</sup>.

## Diurnal Cycling of H<sub>2</sub>O<sub>2</sub>

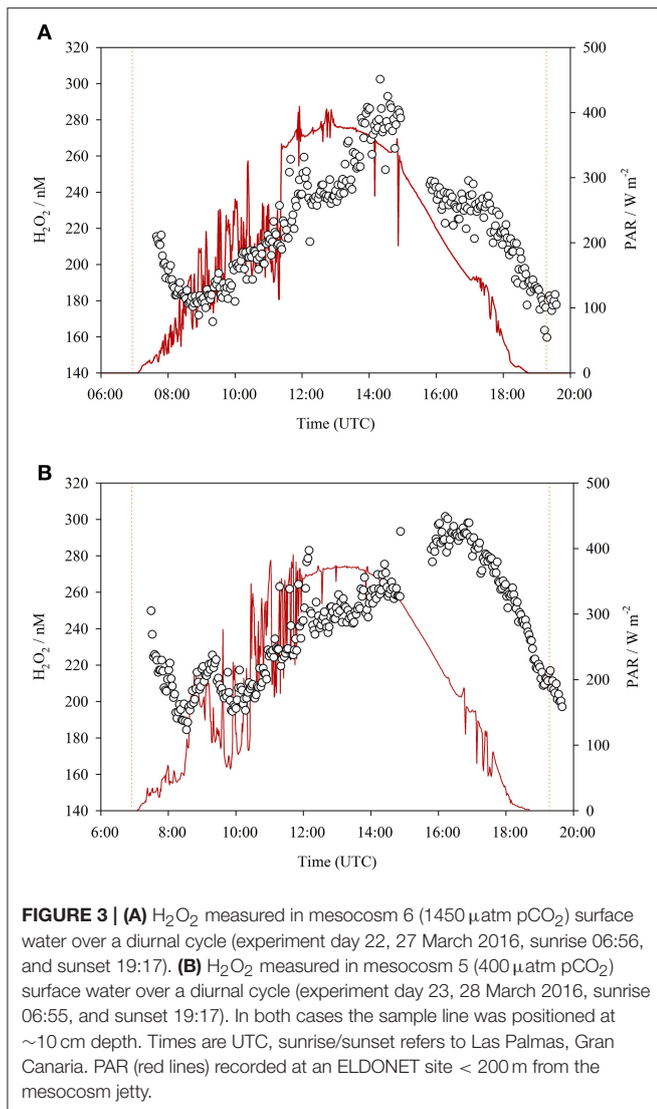
The diurnal cycling of H<sub>2</sub>O<sub>2</sub> was followed both in ambient seawater (**Figure 2**) and inside mesocosm numbers 6 (1,450 μatm pCO<sub>2</sub>, **Figure 3A**) and 5 (400 μatm pCO<sub>2</sub>, **Figure 3B**) on days 22 and 23, respectively. Gaps in the data series corresponded to periods when standard additions were analyzed, the FIA instrument was cleaned, or malfunctions occurred such as air bubbles which occasionally resulted in missed data collection points. Generally, a clear increase in H<sub>2</sub>O<sub>2</sub> associated with



**FIGURE 2** | H<sub>2</sub>O<sub>2</sub> concentrations (nM) in ambient waters adjacent to the mesocosm sampling jetty over a complete diurnal cycle at ~2 min resolution (blue circles), superimposed on H<sub>2</sub>O<sub>2</sub> concentrations for different days (as per **Figure 3**) from within the 1,450 μatm pCO<sub>2</sub> (dark gray) and 400 μatm pCO<sub>2</sub> (light gray) mesocosms. Ambient H<sub>2</sub>O<sub>2</sub> concentration data collected 5–6 March 2016. All times are UTC, sunrise/sunset (illustrated) refers to Las Palmas: sunrise (5 March) 7:21, sunset (5 March) 19:04, sunrise (6 March) 07:20. The sample line was positioned at ~10 cm depth. There was no large change in surface salinity over the experiment duration (range 0.2 practical salinity units).

daylight was evident in all three diurnal datasets (**Figures 2, 3**). Yet H<sub>2</sub>O<sub>2</sub> concentrations were much higher throughout the diurnal cycle within the mesocosms compared to ambient waters. In all three diurnal cycles, peak H<sub>2</sub>O<sub>2</sub> concentration occurred mid-late afternoon (times refer to UTC). In the two mesocosms where diurnal H<sub>2</sub>O<sub>2</sub> concentration was followed, the range between peak and minimum H<sub>2</sub>O<sub>2</sub> was similar (180–300 nM, mesocosm 5, day 23; 160–300 nM, mesocosm 6, day 22), despite the pronounced difference in pCO<sub>2</sub> (400 μatm [5] compared 1,450 μatm [6]) and the 2 h offset between the timing of peak daytime H<sub>2</sub>O<sub>2</sub> concentration (**Figure 3**). In ambient seawater, the amplitude of diurnal variation in H<sub>2</sub>O<sub>2</sub> (~20–30 nM, **Figure 2**) was very similar to that reported previously in the central Atlantic Ocean (25 nM, Yuan and Shiller, 2001).

Two curious features were notable in both mesocosm diurnal cycles (**Figure 3**). These features could only be observed by virtue of the very high data resolution and, to some extent, also the relatively high H<sub>2</sub>O<sub>2</sub> concentrations within the mesocosms compared to ambient waters. First, peak H<sub>2</sub>O<sub>2</sub> concentrations occurred >1 h after peak irradiance. PAR data from ELDONET was not available for the seawater diurnal study date (**Figure 2**), but noting the timing of peak H<sub>2</sub>O<sub>2</sub>, there was likely approximately a 1 h offset between peak irradiance and peak H<sub>2</sub>O<sub>2</sub> here also. Second, the net decline in H<sub>2</sub>O<sub>2</sub> concentration which occurred in the dark continued until sometime after sunrise in both mesocosm studies (**Figure 3**). An increase in H<sub>2</sub>O<sub>2</sub> was not apparent until after PAR increased above ~100 W m<sup>-2</sup>. This was not apparent in the ambient seawater study (**Figure 2**) where a relatively stable concentration of 18.1 ± 1.3 nM was maintained from 21:11 to 07:20 (sunrise) with a sustained rise in H<sub>2</sub>O<sub>2</sub> thereafter. This time offset could



simply therefore have related to the incident solar angle as the mesocosm walls may have reduced the rate of light induced H<sub>2</sub>O<sub>2</sub> formation at low incident solar angles.

### Minor H<sub>2</sub>O<sub>2</sub> Additions From Rainwater

The open design of the mesocosms created the potential for significant atmospheric deposition of H<sub>2</sub>O<sub>2</sub>. Rain water H<sub>2</sub>O<sub>2</sub> concentrations were quantified for the two rainfall events during the mesocosm experiment with sufficient amounts of deposition (>1 mm) to facilitate sampling (Table 1) and were similar to values reported elsewhere over the Atlantic (Zika et al., 1982; Kieber et al., 2001). H<sub>2</sub>O<sub>2</sub> concentrations in rainwater are thought to be sufficiently high to offset the expected beneficial effect of rainwater derived nutrient and micro-nutrient addition on primary production when rainwater is mixed with seawater (Willey et al., 2004) unless the dilution factor is high. The calculated H<sub>2</sub>O<sub>2</sub> addition to mesocosms resulting from rainfall in Gran Canaria was however modest relative to the mesocosm

**TABLE 1 |** Contribution of rainwater events to H<sub>2</sub>O<sub>2</sub> concentrations in the mesocosms.

Location	Date and time	Rainfall (mm)	Rainwater H <sub>2</sub> O <sub>2</sub> (μM)	Calculated H <sub>2</sub> O <sub>2</sub> increase in mesocosms (nM)
Taliarte, Gran Canaria	21/03/16 07:00 (day 16)	1	20.5	6
	30/03/16 17:30 (day 25)	3	44.4	40

H<sub>2</sub>O<sub>2</sub> concentrations due to the low ratio of rain:mesocosm volume (Table 1).

## DISCUSSION

### Diurnal Cycling of H<sub>2</sub>O<sub>2</sub>

Photochemical processes are thought to be the dominant influence on ambient H<sub>2</sub>O<sub>2</sub> concentrations in the surface ocean (O'Sullivan et al., 2005; Steigenberger and Croot, 2008), with large scale spatial variations typically explained in terms of latitudinal changes in light and ocean temperature (Yocis et al., 2000; Yuan and Shiller, 2001). In the high-resolution temporal experiments a clear diurnal cycle in H<sub>2</sub>O<sub>2</sub> concentration was evident, both inside and outside the mesocosm bags (Figures 2, 3). The amplitude of H<sub>2</sub>O<sub>2</sub> concentration over a diurnal cycle was however much greater inside the mesocosms (100 nM compared to 20–30 nM in ambient seawater). H<sub>2</sub>O<sub>2</sub> concentrations over the 28 day duration of the mesocosm experiment (Figure 1A) were always measured at 14:00–15:00 (UTC) daily to ensure that the month long data series was not affected unduly by diurnal variation.

During overnight monitoring of ambient H<sub>2</sub>O<sub>2</sub> it appeared that an equilibrium concentration of 18 nM was maintained in the dark (Figure 2). This was indicative of a dark production mechanism, such as that highlighted in prior work (Palenik and Morel, 1988; Moffett and Zafriou, 1990), sufficient to offset H<sub>2</sub>O<sub>2</sub> decay. Whilst photochemical processes are no doubt a major source of H<sub>2</sub>O<sub>2</sub>, inter-day changes in integrated PAR were only significantly correlated with the H<sub>2</sub>O<sub>2</sub> trend in ambient seawater (Spearman rank correlation 0.73, *p* value 0.002, *n* = 14) and in the 400 μatm mesocosm (Table 2). In all other mesocosms there was no apparent correlation between integrated daily PAR and H<sub>2</sub>O<sub>2</sub> concentration from days 7–29.

The two high-resolution diurnal cycles within mesocosms (Figure 3) revealed some interesting features that would not be apparent at lower H<sub>2</sub>O<sub>2</sub> concentrations (due to the increased signal:noise ratio) or reduced sampling resolution. An offset between irradiance and H<sub>2</sub>O<sub>2</sub> concentration was reflected in the later delayed peak in H<sub>2</sub>O<sub>2</sub> compared to maximum irradiance. Partial shading of the mesocosms in the early morning and late afternoon could create a local PAR exposure that was reduced compared to that reported by a mounted sensor. However, as the displacement of the sensor was <200 m from the mesocosms, this could not explain the offset between peak H<sub>2</sub>O<sub>2</sub> and peak irradiance in the mid-afternoon. Instead, we suggest that net

**TABLE 2** | Spearman rank correlation coefficients and *p* values for H<sub>2</sub>O<sub>2</sub> concentration time series in each mesocosm compared to: baseline H<sub>2</sub>O<sub>2</sub> concentration in ambient seawater (nM), chlorophyll *a* (μg L<sup>-1</sup>), total bacteria (cells mL<sup>-1</sup>), colored dissolved organic matter slope ratio (S<sub>R</sub>), and daily integrated mean photosynthetically active radiation (PAR, W m<sup>-2</sup>).

Mesocosm (number) pCO <sub>2</sub> (μatm)	Spearman rank correlation coefficient									
	Baseline H <sub>2</sub> O <sub>2</sub>			Chlorophyll <i>a</i>		Bacteria		CDOM SR		PAR
	Day -3 to 9	Day 11+	All days	Day 11+	All days	Day 11+	All days	Day 11+	All days	Days 7+
(5) 400	0.86	NSR	0.48	NSR	NSR	NSR	NSR	NSR	NSR	0.58
(7) 700	0.79	NSR	0.42	NSR	NSR	NSR	NSR	NSR	-0.51	NSR
(3) 850	0.81	NSR	NSR	-0.58	-0.47	-0.85	-0.52	0.70	NSR	NSR
(4) 1000	0.81	NSR	NSR	NSR	NSR	-0.87	-0.50	NSR	NSR	NSR
(8) 1150	0.86	NSR	NSR	-0.78	-0.53	-0.83	-0.56	NSR	NSR	NSR
(2) 1300	0.83	NSR	NSR	-0.61	-0.46	-0.94	-0.58	NSR	NSR	NSR
(6) 1450	0.93	NSR	NSR	NSR	NSR	-0.59	NSR	NSR	NSR	NSR
<b>P VALUES</b>										
(5) 400	0.0018	0.44	0.019	0.29	0.87	0.15	0.59	0.92	0.50	0.028
(7) 700	0.015	0.23	0.040	0.74	0.91	0.081	0.16	0.85	0.035	0.13
(3) 850	0.0096	0.76	0.33	0.035	0.030	<0.001	0.023	0.010	0.17	0.66
(4) 1000	0.0096	0.82	0.12	0.098	0.36	<0.001	0.028	0.97	0.99	0.20
(8) 1150	0.0018	0.67	0.57	<0.001	0.013	<0.001	0.013	0.72	0.54	0.92
(2) 1300	0.0053	0.73	0.38	0.025	0.034	<0.001	0.010	0.60	0.54	0.70
(6) 1450	<0.001	0.70	0.46	0.17	0.60	0.031	0.28	0.85	0.20	0.22

NSR, no significant relationship (*P* > 0.05). Highly significant correlations (*P* 0.010 or less) are highlighted. Mesocosm 1 (550 μatm pCO<sub>2</sub>), was excluded due to the lack of data after day 3.

biological production of H<sub>2</sub>O<sub>2</sub> is slightly offset from irradiance. A critical factor, not measured herein, which likely affects the net rate of change in H<sub>2</sub>O<sub>2</sub> concentration during daylight hours is the production rate of peroxidase or catalase enzymes. This process is known to be diurnally variable, with oxidative defenses more active during daylight hours (Morris et al., 2016). Whilst it is not clear if this response under natural conditions is triggered by increasing H<sub>2</sub>O<sub>2</sub> induced stress or by a circadian rhythm, the rate and efficiency (which may also change with incident light) of enzymatic H<sub>2</sub>O<sub>2</sub> removal are likely to be key influences on the shape of the diurnal H<sub>2</sub>O<sub>2</sub> trend.

## Correlation of H<sub>2</sub>O<sub>2</sub> in Mesocosms With Core Parameters

Between days -3 and 9 the range of H<sub>2</sub>O<sub>2</sub> within the different mesocosms was small (<45 nM) and the behavior of chlorophyll *a* and nitrate very similar across all mesocosms (Figures 1A–C). During this early phase, H<sub>2</sub>O<sub>2</sub> inside and outside the mesocosms was relatively well correlated across all pCO<sub>2</sub> conditions (positive Spearman Rank Coefficients of 0.79–0.93). Combined with the comparability of the diurnal H<sub>2</sub>O<sub>2</sub> cycle in 400 and 1,450 μatm pCO<sub>2</sub> mesocosms (Figure 3), despite the extreme pCO<sub>2</sub> difference, this suggests that the direct effect of the pCO<sub>2</sub> gradient (applied after day 0) on H<sub>2</sub>O<sub>2</sub> concentrations was minimal.

In contrast to the early stage of the experiment, H<sub>2</sub>O<sub>2</sub> concentrations in the mesocosms diverged after day 9 and there was then no longer a statistically significant relationship between the paired H<sub>2</sub>O<sub>2</sub> concentrations in and outside the mesocosm (Spearman Rank Correlation *P* > 0.05, Table 2) for any pCO<sub>2</sub>

treatment. If irradiance was the dominant factor controlling extracellular H<sub>2</sub>O<sub>2</sub> concentrations throughout the experiment, we would expect H<sub>2</sub>O<sub>2</sub> concentrations to have remained similar across all mesocosms and to remain correlated with H<sub>2</sub>O<sub>2</sub> in ambient water outside the mesocosms. Yet this was not the case. Variations in daily integrated PAR, available from day 7 onwards, could explain some of the variation in background seawater H<sub>2</sub>O<sub>2</sub>, but inter-day changes in PAR were only significantly correlated with H<sub>2</sub>O<sub>2</sub> in the 400 μatm mesocosm (Table 2).

Formation of ROS generally increases with dissolved organic carbon concentration in estuarine waters and this is specifically linked to the presence of terrestrially derived humic material (Timko et al., 2014) which has a relatively large component of CDOM. However, CDOM properties are not clearly linked to increasing H<sub>2</sub>O<sub>2</sub> concentrations in offshore seawater (O'Sullivan et al., 2005). The general decline in S<sub>R</sub> throughout the mesocosms (from day 1 to day 27, Figure 1E) suggested that *in situ* production of higher molecular weight CDOM was sufficient to offset photo bleaching. However, this was not clearly related to any change in H<sub>2</sub>O<sub>2</sub> concentration as no clear correlation was found between S<sub>R</sub> and H<sub>2</sub>O<sub>2</sub> either over the whole experiment or specifically for days 11+ (Table 2).

Similarly, the overall trend in H<sub>2</sub>O<sub>2</sub> concentration could not be related directly to macronutrient depletion. The observed nitrate concentrations were similar across all mesocosms until after the addition of macronutrients on day 18 (Figure 3) and there was no consistent trend in H<sub>2</sub>O<sub>2</sub> concentration in the days immediately following this addition. Most mesocosms obtained the highest measured H<sub>2</sub>O<sub>2</sub> concentration between days 9–18. Yet mesocosm 6 was notable for obtaining a peak H<sub>2</sub>O<sub>2</sub>

concentration 3 days after the nutrient addition. A sustained drop in H<sub>2</sub>O<sub>2</sub> concentration across all mesocosms (apart from the anomalous mesocosm 7) was only observed between days 23–25. The lowest H<sub>2</sub>O<sub>2</sub> concentrations, both within the mesocosm experiment and relative to ambient seawater, were observed in this post-bloom phase when bacterial abundance peaked (Figure 1D) and when daily integrated PAR was persistently low for 3 days (Figure 1F).

## A Contribution to H<sub>2</sub>O<sub>2</sub> From Mesocosm Design?

H<sub>2</sub>O<sub>2</sub> concentrations might be expected to be generally higher inside the mesocosms due to the nature of their design. Sunlight is attenuated with depth, so in a natural surface mixed layer H<sub>2</sub>O<sub>2</sub> is formed predominantly at the surface and then physically mixed throughout the surface mixed layer (Johnson et al., 1989; Miller and Kester, 1994), although sub-surface H<sub>2</sub>O<sub>2</sub> peaks can occasionally be observed at the chlorophyll a maxima (Yuan and Shiller, 2001; Croot et al., 2005; Steigenberger and Croot, 2008). The confinement of the surface 3 m within the mesocosm bags used here thereby encloses the water with the highest light exposure and, throughout most of this experiment, a high level of biological activity relative to ambient seawater, whilst removing the physical processes that would constantly act to mix H<sub>2</sub>O<sub>2</sub> into deeper waters. The enclosure of seawater was thereby a likely factor contributing to the high H<sub>2</sub>O<sub>2</sub> concentrations occurring during the middle (days 9–18) of the mesocosm experiment. Yet this alone does not explain the divergence in H<sub>2</sub>O<sub>2</sub> levels between the various mesocosms after day 9 (Figure 1A). The timings of the initial mini-bloom after the mesocosms were filled, of the macronutrient addition and the subsequent bloom were (with the exception of mesocosm 7) relatively uniform (Figures 1B,C).

## A Link Between Community Structure and Extracellular H<sub>2</sub>O<sub>2</sub>?

Two possible explanations for why H<sub>2</sub>O<sub>2</sub> concentrations were so variable between mesocosms, which are not mutually exclusive, are that either the biological H<sub>2</sub>O<sub>2</sub> source(s) or the biological H<sub>2</sub>O<sub>2</sub> sink(s) varied because of different microbial communities across the pCO<sub>2</sub> gradient. An anti-correlation between bacterial abundance and extracellular H<sub>2</sub>O<sub>2</sub> concentrations is therefore intriguing. Microorganisms are simultaneously both a H<sub>2</sub>O<sub>2</sub> source and a H<sub>2</sub>O<sub>2</sub> sink. ROS are generated as waste products from aerobic cellular metabolism (Kruk, 1998; Apel and Hirt, 2004), with H<sub>2</sub>O<sub>2</sub> produced both directly and from the decay of other ROS (for example, the enzyme superoxide dismutase produces H<sub>2</sub>O<sub>2</sub> from O<sub>2</sub><sup>-</sup>; McCord and Fridovich, 1969). Conversely, biological H<sub>2</sub>O<sub>2</sub> sinks arise from both synthesis of H<sub>2</sub>O<sub>2</sub> destroying enzymes (catalase and peroxidase) and also, under some circumstances, serendipitously with increasing biomass. The quenching of extracellular H<sub>2</sub>O<sub>2</sub> with increasing biomass is referred to as “self-shading” and has been described both in monocultures of *Prochlorococcus*, which lacks catalase-peroxidase genes (Morris et al., 2011) and at the community level (Barros and Colepicolo, 2003). It

presumably arises from physical shading and/or non-enzymatic ROS sinks.

The anti-correlation between H<sub>2</sub>O<sub>2</sub> concentrations in each mesocosm both with chlorophyll a and with total bacterial abundance over the duration of the experiment (Table 2) was statistically significant in most mesocosms. Overall there was a significant ( $P < 0.05$ ) anti-correlation between chlorophyll a and H<sub>2</sub>O<sub>2</sub> concentrations with negative correlation coefficients (Table 2) in mesocosms 3, 4, 8, and 2 (850–1,300 μatm pCO<sub>2</sub>). The anti-correlation was slightly stronger when the initial phase of the experiment (Days –3 to 9, when the range of H<sub>2</sub>O<sub>2</sub> concentrations across all mesocosms was relatively narrow) was excluded. A negative relationship between chlorophyll a and H<sub>2</sub>O<sub>2</sub> concentrations could arise from an increasing biological H<sub>2</sub>O<sub>2</sub> sink simply due to self-shading associated with increasing biomass, yet no correlation with chlorophyll a was found either at low (400, 700 μatm) or at the highest (1,450 μatm) pCO<sub>2</sub> treatments. For bacteria, the correlation followed the same pattern as chlorophyll a with respect to pCO<sub>2</sub>, yet the effect was stronger and more significant. From day 11 onwards, the intermediate pCO<sub>2</sub> mesocosms (850–1,300 μatm) all exhibited a strong negative correlation (Spearman Rank Coefficients of –0.83 to –0.94) between bacteria cell counts and H<sub>2</sub>O<sub>2</sub> concentration ( $P$  values  $< 0.001$ ). This suggests that under post-bloom conditions, bacteria were an important net-sink for extracellular H<sub>2</sub>O<sub>2</sub>.

As both cellular H<sub>2</sub>O<sub>2</sub> production and defense mechanism activity vary over diurnal cycles and at the species level, it is difficult to separate the source and sink terms within separate mesocosms and over the pCO<sub>2</sub> gradient. Yet in any case, the effects of a pCO<sub>2</sub> gradient on H<sub>2</sub>O<sub>2</sub> concentration were apparently indirect, resulting from changes to bacterial abundance and community composition rather than directly arising as a consequence of perturbing the carbonate system alone.

## Effects of Elevated H<sub>2</sub>O<sub>2</sub> Under Experimental Conditions?

Reported H<sub>2</sub>O<sub>2</sub> concentrations in seawater incubated under laboratory conditions are typically 100–300 nM (e.g., Coe et al., 2016), but can be much higher depending on the buffer composition and irradiance exposure (Morris and Zinser, 2013). In a spot-check of baseline H<sub>2</sub>O<sub>2</sub> concentrations in incubated seawater (with added nutrients) in our own laboratory, we found  $42 \pm 50$  nM H<sub>2</sub>O<sub>2</sub> in freshly made mixtures (made from seawater which was previously stored in the dark for >1 year) and  $180 \pm 130$  nM in the same water after 72 h of incubation subject to a diel light cycle. In all these cases, the H<sub>2</sub>O<sub>2</sub> concentrations in seawater incubated under laboratory conditions are thereby at the mid-high end of the range observed in the surface ocean (Price et al., 1998; Yuan and Shiller, 2001; Hopwood et al., 2017).

The unmodified concentrations of H<sub>2</sub>O<sub>2</sub> in growth media are sufficiently high to prevent the cultivation of some microorganisms, including strains of *Prochlorococcus*, under laboratory conditions (Morris et al., 2011). However, uncertainties remain about what the effects of elevated H<sub>2</sub>O<sub>2</sub> are

at the community level. Whilst H<sub>2</sub>O<sub>2</sub> is not widely investigated during mesocosm studies, field evidence suggests that increases in H<sub>2</sub>O<sub>2</sub> to concentrations less than the peaks observed here (**Figure 1A**) can affect a broad range of biogeochemical processes. Additions of 10–100 nM H<sub>2</sub>O<sub>2</sub> to water from 100 m in the North Atlantic were found to reduce the extracellular activity of  $\beta$ -glucosidase, alkaline phosphatase, and leucine aminopeptidase by 20–80% with the inhibition effect consistently strongest on  $\beta$ -glucosidase (Baltar et al., 2013). A similar small absolute increase of only 30 nM is reported to suppress rates of ammonia oxidation to below detection in surface (10 m depth) Antarctic seawater (Tolar et al., 2016) where the initial natural concentration of H<sub>2</sub>O<sub>2</sub> was low (14 nM). Additional specific effects of increasing extracellular H<sub>2</sub>O<sub>2</sub> concentrations in seawater can also be deduced from laboratory experiments. For example, an increase on the order of 100 nM H<sub>2</sub>O<sub>2</sub> is sufficient to measurably decrease the half-life of Fe(II), thereby theoretically decreasing dissolved Fe bioavailability (González-Davila et al., 2005), and to double brominating activity in some diatoms (Hughes and Sun, 2016).

A particularly curious phenomenon observed here was the extremely high H<sub>2</sub>O<sub>2</sub> concentrations observed during the pre-bloom phase of the experiment (**Figure 1A**). It is questionable to what extent this phenomenon would occur in a natural environment because small-scale mesocosms inevitably fail to mimic the mixing processes that occur within the natural water column. Nevertheless, the potential for oxidative stress from extracellular ROS to be increased under natural, or incubated, oligotrophic conditions should be investigated concurrently with other stressors to understand the interactive effect(s) on microbial functioning. As outlined above, the sensitivity of some enzymatic processes to increasing extracellular H<sub>2</sub>O<sub>2</sub> concentrations and the involvement of H<sub>2</sub>O<sub>2</sub> in the biogeochemical cycling of various chemicals including the micronutrient Fe, means that large perturbations to H<sub>2</sub>O<sub>2</sub> during mesocosm experiments are undesirable.

## CONCLUSIONS

During the progression of a 28 day mesocosm experiment in North Atlantic waters an applied pCO<sub>2</sub> gradient (400–1,450  $\mu$ atm) had no discernable direct effect upon extracellular H<sub>2</sub>O<sub>2</sub> concentrations. Whilst a clear diurnal trend in H<sub>2</sub>O<sub>2</sub> was observed, both within high/low pCO<sub>2</sub> mesocosms and in ambient waters, inter-day variation in PAR was not correlated with mesocosm H<sub>2</sub>O<sub>2</sub> concentrations.

The general elevation of H<sub>2</sub>O<sub>2</sub> above ambient North Atlantic concentrations was attributed to the effect of enclosing seawater in an open mesocosm at the ocean surface. Yet during the pre-bloom phase of the experiment, unexpected swings occurred to H<sub>2</sub>O<sub>2</sub> concentrations >300 nM. Across the majority of mesocosms, bacteria appeared to be a net-sink for H<sub>2</sub>O<sub>2</sub>, particularly under post-bloom conditions. Thus,

microbial community structure does appear to strongly influence extracellular H<sub>2</sub>O<sub>2</sub> concentrations.

Given the multitude of possible direct and indirect, chemical and biological effects of large changes to H<sub>2</sub>O<sub>2</sub> concentrations, it is important to consider to what extent ROS are a source of oxidative stress in the natural surface ocean and how well experiments manipulate the natural environment with respect to this stressor.

## AUTHOR CONTRIBUTIONS

All authors contributed to the design of the study. Fieldwork and analytical work was conducted by MH, UR, AL, JA, and NH. MH wrote the initial draft of the manuscript and then all authors contributed to its revision.

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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.2018.00105/full#supplementary-material>

**Supplementary Figure 1** | Mesocosm timeline labeled with experiment days (used throughout the text). Day -4, when the mesocosms were filled, was 1 March 2016. A pH gradient was imposed on day 0 by the addition of CO<sub>2</sub> saturated, filtered seawater. Further pCO<sub>2</sub> additions were then made as necessary to compensate for out-gassing. The volume of each mesocosm was determined on day 18 by measuring the change in salinity resulting from a freshwater addition. A single macronutrient spike was then added after sampling on day 18.

**Supplementary Datasheet 1** | Mesocosm time series data for core parameters used: chlorophyll a, nitrate + nitrite, hydrogen peroxide, bacterial abundance and CDOM SR.

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**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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