Spatiotemporal Distribution of Key Pelagic Microbes in a Seasonal Oxygen-Deficient Coastal Upwelling System of the Eastern South Pacific Ocean

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The strong seasonal variability in physical-chemical conditions of the Eastern South Pacific Ocean creates an ideal setting to study spatiotemporal distribution of key marine microbial communities. We herein report a nearly 4-year-long time series of the variability in amoA gene counts of ammonia oxidizing archaea (AOA) and bacteria (betaproteobacteria, bAOB) by quantitative PCR, GI.1a Thaumarcheota and MG-II Euryarcheota by CARD-FISH, and the picoplanktonic community by flow cytometry for this area. During spring-summer, non-photosynthetic picoplankton such as MG-II Euryarcheota and GI.1a Thaumarcheota peaked at the surface and deeper waters, respectively. General AOA and bAOB achieved higher abundances at the oxycline mainly in summer (up to $10^5-10^4$ amoA copies mL$^{-1}$). Generalized additive models for location, scale, and shape (GAMLSS) indicated that season and depth account for 19–46% of variations in the abundance of the groups studied, particularly GI.1a Thaumarchaeota and AOA. The oxygen and nitrite concentration were statistically meaningful predictors for the studied groups. GAMLSS models indicate that ammonia oxidizing assemblage’s variability is coupled with ammonia, nitrite, and nitrate variations. Our results indicate that microbial abundances fluctuation is associated with upwelling variability and oxygen-deficient water conditions that shape the substrates availability and metabolic response of marine microbes, including keystone ammonia oxidizing assemblages and their ecological interactions. Overall, our results support planktonic nitrification activity and its contribution to nitrous oxide excess production in the time series off Concepción and the ecological dynamics regarding AOA and bAOB in coastal waters.

Keywords: Thaumarchaeota, chemolithotrophic nitrification, environmental forcing, coastal upwelling time-series station, oxygen minimum zone, ecological niche, quantitative PCR, CARD-FISH
INTRODUCTION

Coastal marine areas hold a diverse microbial community but with the predominance of abundant taxa, such as Gamma- and Alpha-proteobacteria, Bacteroidetes among other bacterial phyla, and archaea from *Thaumarchaea* and *MGII Euryarchaeota* which has been reported to have a spatial-temporal dynamics, e.g., surface versus subsurface, and in response to climatic and oceanographic conditions, e.g., San Pedro Ocean Time series station (Cram et al., 2015; Parada and Fuhrman, 2017); Monterey Bay (Reji et al., 2019; Tolar et al., 2020), based on their ecological and metabolic traits (e.g., Northwest Mediterranean coast, Galand et al., 2018; Monterey Bay, Reji et al., 2019). Marine microbes from coastal zones are also subjected to anthropological pressure, such as, nutrient fertilization and other pollutants, influencing biogeochemical conditions and oxygen-deficiency (Cavicchioli et al., 2019).

Besides photosynthetic communities, chemoautotrophic assemblages associated with nitrification have been reported to play a central role in the functioning of coastal microbial assemblages (e.g., network analyses, Parada and Fuhrman, 2017; Reji et al., 2020). Nitrification is a chemoautotrophic process associated with a two-steps aerobic reaction that oxidizes ammonia into nitrite followed by the conversion of nitrite into nitrate. Two functional groups represented by ammonia- and nitrite-oxidizing microorganisms catalyze complete marine nitrification, with the rate-limited first step of this reaction performed by ammonia-oxidizing bacteria (AOB) and archaea (AOA) (Ward, 2008). *Nitrospira* bacteria able to catalyze one-step ammonia oxidation to nitrate (Comamnox) represents the only known exception to this rule (Daims et al., 2015; van Kessel et al., 2015). However, the biogeochemical impact of *Nitrospira*-like Comamnox bacteria on the marine nitrogen cycle is still unclear, since these microorganisms appear to be absent in marine ecosystems (Daims et al., 2016).

Phylogenetically complex natural assemblages of ammonia-oxidizing microorganisms include *Thaumarchaea*, groups 1.1a, 1.1b, and Hot Water Crenarchaeota Group (HWCGIII) and *Nitrosocaldus* group (Pester et al., 2011), with the group 1.1a representing one of the most ubiquitous and abundant microbial lineages on earth (Karner et al., 2001). In contrast, AOB are less abundant and often undetectable in marine ecosystems (Mincer et al., 2007; Beman et al., 2010; Tolar et al., 2013). Nonetheless, according to the quantification of the functional marker *amoA* genes (encoding the alpha subunit of the ammonia monooxygenase enzyme), AOB could be more abundant than AOA. For example, higher ratios of benthic *amoA* gene copies were detected in freshwater and eutrophic waters (Bouskill et al., 2012) or in association with increased amounts of substrate and salinity in estuarine sediments (Bernhard and Bollmann, 2010; Li et al., 2015). In addition, the abundance and potential activity of AOB (as *amoA* gene counts and transcript counts, respectively) may be important in other marine environments characterized by strong physical-chemical gradients such as nitrification zones in oxygen-deficient areas (Lam et al., 2007) and near-surface depths (Tolar et al., 2013), including diel variability in sunlit coastal waters (Levipan et al., 2016).

Ecophysiological studies indicate a niche differentiation among AOA and AOB strains associated to factors such as temperature (Bayer et al., 2016), pH (Pommerning-Röser and Koops, 2005; Qin et al., 2014; Bayer et al., 2016), light (Guerrero and Jones, 1996; Qin et al., 2014), substrate concentrations (Martens-Habbena et al., 2009), and substrates used for nitrification (Qin et al., 2014; Bayer et al., 2016). For instance, AOA isolates are more photosensitive and grow at lower ammonia concentrations than AOB isolates (Martens-Habbena et al., 2009; Merbt et al., 2012). This is consistent with field studies indicating that natural AOB and AOA populations are commonly associated with high and intermediate to low ammonia concentrations, respectively (Bouskill et al., 2012; Sintes et al., 2013). Moreover, natural AOB and AOA populations may use different substrates such as urea and cyanate to fuel nitrification (Alonso-Sáez et al., 2012; Bowen et al., 2013; Connelly et al., 2014; Palatinszky et al., 2015).

Marine microbial time series (oceanic and coastal) could help to identify long-term trends in the biogeochemical cycles and predict future environmental sceneries, based on their inherent physical-chemical variability. However, there are still relatively few marine time series studies related to the variability of major marine archaeal lineages, besides ammonia oxidizing assemblages, especially in highly productive, oxygen-deficient and dynamic coastal upwelling areas. For example, exhaustive studies have been carried out along the California coast at San Pedro Ocean time series (SPOT) and Monterey Bay Aquarium Research Institute (MBARI) time series (e.g., Murray et al., 1999; Mincer et al., 2007; Beman et al., 2010, 2011; Steele et al., 2011; Robidart et al., 2012; Parada and Fuhrman, 2017; Tolar et al., 2020), at the station ALOHA in Hawaii (Karner et al., 2001), at Devil’s Hole in Bermuda (Parsons et al., 2015), and in marine ecosystems such as Chesapeake Bay (Bouskill et al., 2011), the Mediterranean Sea (Galand et al., 2010, 2018; Lambert et al., 2019), and the North Sea (Wuchter et al., 2006). These studies concluded that AOA are the main responsible for ammonia oxidation in oceanic and coastal areas in general.

The coastal area off central-southern Chile (~36°S) in the eastern South Pacific Ocean is an ideal location to determine the effects of oceanographic dynamics on microbial communities and their biogeochemical activities, since this is a highly productive seasonal upwelling system. Upwelling period occurs during the austral spring-summer modulated by southwesterly winds (Sobarzo et al., 2007). The upwelled water is influenced by the Equatorial Subsurface Waters (Euvw), characterized by cold (~9–12°C), high-salinity (>34.5 psu), nutrient-rich (reaching > 30 μM of nitrate), and oxygen-depleted conditions (~< 1 mL O2 L−1) (Ahumada and Chuecas, 1979). In contrast, strong water mixing events caused by intense northerly winds occur during the austral fall-winter, with water column being characterized by warmer temperatures (10–13°C) and low-nutrient concentrations (e.g., <20 μM of nitrate). Moreover, winter mixing events ventilate the surface
and middle depths (>1–6 mL O₂ L⁻¹), although low oxygen concentrations (<0.5 mL O₂ L⁻¹) persist at bottom depths almost the entire year (Sobarzo et al., 2007). During fall-winter, high precipitation and river-runoff reduce salinity (between ~32.5 to <34.1 psu) at the surface generating strong stratification in the first 10–20 m depth (Sobarzo et al., 2007; Saldías et al., 2012).

The oceanographic dynamic in central-southern Chile along with the high solar radiation in spring-summer promotes a high primary productivity (Montero et al., 2007; Hernández et al., 2012) and faster remineralization rates (Daneri et al., 2004) compared with winter, c.a. 21 nmol L⁻¹ d⁻¹ (Fernandez and Farías, 2012). Recently, ammonia and nitrite oxidation has been detected in this area at nanomolar concentrations of oxygen (5–33 nM, Bristow et al., 2016). Furthermore, molecular studies indicate the presence of an abundant and rich AOA community (Molina et al., 2010; Bertagnolli and Unloa, 2017), as well as of *Nitrospina* spp., as the main drivers of nitrification under oxygen-deficient conditions at bottom waters during the spring-summer time (Levipan et al., 2014). In general, the water column is characterized by higher bacterioplankton abundances in summer (Galán et al., 2012), and *Nitrosopumilus maritimus* predominance (an important AOA ecotype at the Sta.18) in winter months based on metagenomic data (Murillo et al., 2014). Additionally, annual variability of light intensity can favor transient co-variability between bAOB and certain AOA ecotypes to catalyze the first step of nitrification (Levipan et al., 2016).

Here, we investigate physical-chemical variability contribution in shaping the spatiotemporal distribution of marine microbial assemblages including ammonia oxidizers at a coastal upwelling station (Sta.18) using data from a nearly 4-year-long time series. We hypothesize that vertical and seasonal environmental fluctuations will significantly modulate the abundance variability of the microbial community in the study area, whereas oxygen and ammonium will be explanatory variables shaping the distribution and abundances of keystone ammonia oxidizing assemblages.

**MATERIALS AND METHODS**

**Study Area and Environmental Data Collection**

The Center for Oceanographic Research in the Eastern South Pacific (COPAS¹; now transformed to other initiatives http://www.sur-austral.cl/). Time-series station 18 is located over the continental shelf off central-southern Chile (36°30′ S; 73°08′ W) and has a maximum depth of approximately 90 m. Station 18 (hereafter, Sta.18) extends approximately 18 nautical miles offshore, and was visited monthly since 2005–2009 on board the R/V Kay Kay-I and -II (University of Concepcion). The wind speed and direction were measured every 10 min at a meteorological station (HOBO) located at Coliumo Bay (36° 31′ 41.2″ S; 72° 57′ 57.3″ W) between August 2004 and August 2009. Now different open observatory programs are available further south off Concepción².

Temperature, salinity, and dissolved oxygen concentrations were obtained using a CTD-O probe (Seabird 23B Electronics, Bellevue, United States). Water samples for chemical (NH₄⁺, NO₃⁻, NO₂⁻ and Chlorophyll-a) and biological analyses (picoplankton, DNA extractions, and CARD-FISH) were collected using a rosette sampler equipped with 10 L Niskin bottles. Triplicated NH₄⁺ samples (40 mL) were fixed on board and fluorometrically analyzed (Turner Designs 10AU fluorometer) using the method proposed by Holmes et al. (1999). Seawater samples for NO₃⁻ and NO₂⁻ analyses (20 mL) were filtered through glass filter (0.7 μm of pore-size, Millipore) and stored frozen (−20°C) until further processing (Parsons et al., 1984). Chlorophyll-a was determined in discrete seawater samples according to the method described by Holm-Hansen et al. (1965). In addition, discrete seawater samples were fixed in duplicate with glutaraldehyde (final concentration, 0.1% v/v) and stored at −80°C for flow cytometry analysis. Non-fluorescent picoplankton was stained with SYBR-Green I (Molecular Probes) and processed following Marie et al. (1997) on a FACSCalibur™ flow cytometer (Becton Dickinson Biosciences, CA, United States) equipped with a 15-mW argon-ion laser tuned at 488 nm.

**DNA Extraction**

Seawater samples (3 L) were collected monthly for DNA extraction at different depths (0, 5, 10, 15, 20, 30, 40, 50, and 80 m) from January 2005 to January 2009. The samples were pre-filtered serially through a 20-μm-mesh (Nitex® nylon) and 3-μm-filters (MCE, mixed cellulose ester, Millipore membrane filter), and then filtered onto cellulose ester membrane filters (47 mm diameter, 0.22 μm pore-size GPWP04700) through peristaltic pumping. Each membrane filter was soaked with 1 mL of DNA buffer (50 mM Tris-HCl [pH 9.0], 0.75 M sucrose, 400 mM NaCl), flash-frozen in liquid nitrogen, and stored at −80°C. The DNA was isolated from thawed filters using the phenol-chloroform extraction method as previously described (Molina et al., 2007). In order to estimate the contribution of different ecotypes, seawater samples (3 L) collected at 0, 5, 10, 20, 30, 50, and 80 m depth from two oceanographically contrasting months (August versus December 2011) were extracted by using the PowerSoil™ DNA Isolation Kit (MoBio Laboratories, Solana Beach, CA, United States) in accordance to the manufacturer’s specifications, with minor modifications described by Levipan et al. (2014). All DNA extracts were resuspended in 100 μL of nuclease-free water and their concentration and quality (A₂₆₀/A₂₈₀ ratio) were determined by optical absorption spectroscopy on a Synergy MX Microplate Reader (BioTek Instruments, Winooski, VT, http://www.cdom.cl/).

¹http://www.copas.cl

²http://www.wm.jh.edu/
United States). The DNA extracts were stored at −80°C until further analysis.

**Quantification of Ammonia-Oxidizing Assemblages**

The quantification of AOA and bAOB amoA genes was carried out with the primers Arch-amoAF/Arch-amoAR (Francis et al., 2005) and amoA-1F/amoA-2R (Rotthauwe et al., 1997), respectively. amoA gene copies were quantified for these two groups using standard curves of 10-fold serial dilutions made from amoA gene clones (dilution range, 2 × 10^7 to 20 copies) as previously described (Molina et al., 2010). The copy number of the amoA clones was calculated by dividing its DNA concentration (in ng μL⁻¹) by its mass (in ng) calculated using the following formula: mass = ([n] × (M/NA)) × 10^9 where n is the clone size (vector plus insert, in bp); M, is the average molecular weight of a base pair (660 g mol⁻¹); NA, is the Avogadro’s number (6,0221 × 10^23 bp mol⁻¹); and 109 is the factor to convert grams to nanograms. Efficiencies and correlation coefficients of standard curves were computed for AOA (E = 91 ± 6%, r² = 0.99 ± 0.012) and bAOB (E = 84 ± 6%, r² = 0.994 ± 0.004).

Profiling of amoA gene counts for AOA belonging to the shallow and deep water column clades named as WCA and WCB, and *N. maritimus* (Group 1.1a) were determined in two oceanographically contrasting months (August vs. December 2011). qPCR assays for these ecotypes were performed with the primer Arch-amoAR along with primers Arch-amoAFA and Arch-amoAFB for WCA and WCB amoA, respectively (Beman et al., 2008). The ecotype *N. maritimus-like* was quantified with the primer Nmar423F (Levipan et al., 2016) used in combination with the primer Arch-amoAR. These primers combination could also amplify other species such as *Nitrosopumilus oxyclinae* and *Ca. Nitrososarchaeum* but not the WCA cultivated representative *Nitrosopelagicus brevis* (Santoro et al., 2015, see alignment in Supplementary Table 2). qPCR standard curves were generated from 10-fold dilutions of amoA gene clones (2 × 10^7 to 20 copies) available in our laboratory for *N. maritimus* (GenBank accession number KJ555107) and WCA and WCB (Molina et al., 2010). The copy number of these amoA clones was calculated using the mentioned formula. PCR efficiencies and correlation coefficients of standard curves for WCA and WCB (E = 91 ± 6%, r² = 0.989 ± 0.012) and *N. maritimus* (E = 95.492%, r² = 0.986) were determined.

All qPCR reactions were performed in triplicate using a StepOne™ Real-Time PCR System (Applied Biosystems, Lincoln Centre Drive, Foster City, CA, United States); data were analyzed with the StepOne™ software package (v.2.2.2; Foster City, CA, United States). Each reaction was conducted on a volume of 20 μL containing 5–10 ng of template DNA, forward and reverse primers (0.4 μM, final concentration), and 2X Fast SYBR® Green Master Mix (1X; Applied Biosystems, 850 Lincoln Centre Drive, Foster City, CA, United States). The qPCR program consisted of an initial denaturation for 20 s at 95°C followed by 40 amplification cycles consisting of 95°C for 3 s, 20 s annealing at a temperature that varied depending on the primer pair used (Supplementary Table 1), and an extension of 20 s at 72°C. A melting curve was performed at the end of each qPCR experiment. The detection limit for standards was always observed at CT values lower than 30. The specificity of qPCR assays was verified by checking dissociation curves and standard electrophoresis of resulting amplicons.

**CARD-FISH**

Seawater samples were collected monthly since September 2006 to September 2008 (2-year-long time series), fixed after collection with paraformaldehyde (final concentration, 1% v/v), and stored at 4°C until processing. Subsamples (50 mL) obtained from different depths and dates were filtered by vacuum (<10 cm Hg) through 0.2-μm polycarbonate filters (Millipore; 45-mm diameter filters placed over 0.45-μm cellulose nitrate support filters) and then stored frozen (at −20°C) until further CARD-FISH assays. Specific oligonucleotide probes targeting 16S rRNA genes of *Thaumarchaeota* GL1a (CREN55: HRP-TTAGGCCCAATAATCMTCTCT-3’; Massana et al., 1997) and *Euryarchaeota* MG-II (EURY806: HRP-CAACGCGTTTACACTAG-3’; Teira et al., 2004) were used following procedures described by Sekar et al. (2004) and Woebken et al. (2007). A mix of three eubacterial probes (EUB338, HRP-GCTGGCTTCCCGTAGGAGT-3’; EURY806-II, HRP-GAGGGCAACCGTAGTT-3’; and EUB338-III, HRP-GCTGCCACCGTAGTTG-3’) was used to detect total bacterial cells (Amann et al., 1990; Daims et al., 1999). The NON338 probe (HRP-ACTCTTACGGAGGCAGC-3’; Wallner et al., 1993) was used to detect false positive in hybridization experiments. Briefly, probes were hybridized during 4 h (EUB338 mix and NON338) and 8 h (CREN55 and EURY806) at 35°C in a standard hybridization buffer (360 mM NaCl, 8 mM Tris-HCl pH 8, 0.008% w/v SDS, 0.4% w/v blocking reagent, 1% w/v dextran sulfate, and the respective probe). The wash temperature for removing non-specifically hybridized probes was set at 37°C for 15 min. Depending upon the probe, the stringency of the hybridization reaction was regulated with different formamide percentages (% v/v) in the hybridization buffer and different NaCl concentrations in a standard washing buffer (2.5 mM EDTA, 20 mM Tris-HCl). The amount of formamide/NaCl (%/mM) used for archaeal (CREN554 and EURY806) and non-archaeal probes (EUB338 mix and NON338) was 20%/3 mM and 55%/135 mM, respectively. Prokaryotic cell abundance was estimated using 4, 6-diamidino-2-phenylindole (DAPI) staining (Porter and Feig, 1980) and an epifluorescence microscope (Zeiss Axiosplan 2) equipped with a 100 W Hg lamp (HBO 103W/2, Osram, Germany) and an appropriate filter set for 5-(6-) FAM and DAPI dyes. Ten epifluorescence photomicrographs (1000×) were randomly taken from each sample, counting between 500–1000 DAPI-stained cells with the AxioVision software (Release 4.6.3.).

**Statistical Analyses**

The abundances of total non-photosynthetic picoplankton, *Thaumarchaeota* (formerly known as marine GI Crenarchaeota; Massana et al., 1997), AOA, and bAOB, were modeled in response to the spatiotemporal variability (season and depth) and
chemical variability (ammonium, nitrite, nitrate, and oxygen) using Generalized Additive Models for Location, Scale, and Shape (GAMLSS) from R ‘gamlss’ package (Rigby and Stasinopoulos, 2005). MG-II *Euryarchaeota* was not included in this analysis because there were temporal gaps in its abundance, but a Spearman correlation analysis indicates that it is significantly correlated with non-photosynthetic picoplankton in surface waters ($R = 0.4; P = 0.0013$). Predictive variables showed a high collinearity (up to Spearman's $R = -0.79, P < 0.001$) due to the strong seasonality in wind-driven upwelling conditions. To solve this problem, for each microbial group, we fitted a spatiotemporal model with season and depth as predictive variables, and independent models for every chemical parameter used as a predictive variable (i.e., nitrite, nitrate, ammonium, and oxygen). Before running the analysis, predictive variables were scaled (Z-transformed) to allow a direct comparison between parameters derived from resultant models (Schielzeth, 2010). In order to count both skewness and leptokurtosis of the response variables, microbial groups were modeled by using the follow distributions (Rigby and Stasinopoulos, 2005): (i) negative binomial type I, (ii) negative binomial type II, and (iii) Box-Cox $t$ (Table 1). Model diagnostic procedures were performed (to choose the most appropriate distribution) using (i) the plot.gamlss function which provided a series of plots for checking normalized quantile residuals of the fitted models, and (ii) the Generalized Akaike Information Criterion (GAIC). Microbial groups were modeled as a linear function of variables seasonality, ammonium, nitrite, nitrate, and oxygen. The depth was adjusted using a non-parametric cubic smoothing spline function available in R ‘gamlss’ package.

The data that support the findings of this study are available from the corresponding author upon reasonable request.

**RESULTS**

**Seasonal Regime of Upwelling-Favorable Winds and Physical-Chemical Variables in the Study Area**

Upwelling-favorable winds (i.e., southwesterly) usually extended from September (early austral spring) to April (early austral fall) in the study area, however, inter-annual differences were detected during our temporal framework (Supplementary Figure 1). The 2004–2005 annual wind cycle showed the most intense southerly winds and the highest accumulation of wind-stress, whereas the lowest accumulated wind-stress was observed during the 2005–2006 annual wind cycle. Upwelling-favorable winds in the 2007–2008 annual wind cycle were similar to those from 2005 to 2006 cycle in terms of their respective starting points-spring transition (early October), but the former was more intense and prolonged. Moreover, upwelling-favorable winds from the 2006 to 2007 and 2008 to 2009 annual wind cycles showed the earliest start (mid-July to early August) and strongest intensities (Supplementary Figure 1).

During the austral spring-summer, upwelled waters were characterized by low temperatures (~9–12°C), and high salinities (>34.5) and densities (~26 kg m$^{-3}$), influencing surface layer; whereas during non-upwelling periods the densest water was just restricted to bottom depths (Supplementary Figure 2). The upwelled water was also characterized by suboxic conditions (~22 µM), and its influence was perceptible up to almost 15 m depth, mostly from November to April (Figure 1). Oxic conditions (>200 µM) were restricted in the first 5–10 m depth over the entire study period and at intermediate depths during the austral fall-winter (i.e., upwelling-unsuitable period) (Figure 1). In general, during the upwelling season, high concentrations of ammonium (>1 µM) were detected at the base of the mixed layer (~10 m depth), except during the 2005–2006 upwelling period, when elevated ammonium concentrations (up to ~3 µM) were found at almost the entire water column (Figure 1). During this period, also high nitrite concentrations (~7 µM) were normally found in the water layer below 40 m depth (Figure 1), except during the 2005–2006 upwelling period, when higher nitrite concentrations were registered both in bottom waters and oxycline. Surface waters showed lower concentrations of nitrate (~10 µM) during the entire study period, while the higher concentrations of this nutrient (up to ~35 µM) were detected at mid-depth waters (between ~20 and 50 m) but not just associated with the seasonal upwelling signature (Figure 1).

**Spatiotemporal Variability of Chlorophyll-a and Microbial Assemblages**

During fall-winter months (upwelling-unsuitable seasons), chlorophyll-a concentrations were always ≤ 5 µg m$^{-3}$ throughout the water column (Figure 2). In contrast, during spring-summer months, the highest chlorophyll-a concentration (i.e., >10 mg m$^{-3}$) was tightly associated with surface waters (between 0 and ~20 m depth), except by the 2006–2007 spring-summer period, when the lower chlorophyll-a concentrations were detected and the span of maximal chlorophyll-a concentration finished earlier than the other years (Figure 2). Cell abundances of the non-photosynthetic picoplankton ranged between 0.23–6.50 × 10$^6$ cells mL$^{-1}$ in the water column, reaching their maxima at surface and bottom waters during late summer and fall, and minima in winter months (Figure 2).

Cell abundances of GI.1a *Thaumarchaeota* and MG-II *Euryarchaeota* (referred to as *Thaumarchaeota* and *Euryarchaeota* in the following sections) are shown in Supplementary Figure 3. The abundance of *Euryarchaeota* range between zero and 78 × 10$^4$ cells mL$^{-1}$, accounting for 9.6 ± 9.0% of total picoplankton community (determined by counting DAPI-stained cells) and showing four maxima in the first approximately 30 m depth (Supplementary Figure 3). Moreover, its abundance decreased with depth, with only one *Euryarchaeota* maxima detected at depth during late summer 2008 (see march-associated peak). In contrast, cell abundances of *Thaumarchaeota* were usually between ~0.2 and 85 × 10$^4$ cells mL$^{-1}$ and more abundant during late summer and winter. This pattern was observed in deeper waters, where this group was accounted for 25.1 ± 13.8% of DAPI counts detected below the oxycline (Supplementary
In addition, the contribution of *Thaumarchaeota* to the whole picoplankton community (as % of the total number of DAPI-stained cells) was as high as 50% below 50 m depth (Supplementary Figure 3).

In total, ammonia-oxidizing assemblages amoA gene copies quantification indicated that AOA amoA outnumbered bAOB amoA by up to an order of magnitude (Figure 2). Moreover, if it is considered that one amoA gene copy corresponds to one cell, AOA and bAOB were accounted for 1.2 ± 2.9% and 0.008 ± 0.015% of total abundance of the non-photosynthetic picoplankton (determined from flow cytometry) in the entire water column over time, respectively. In addition, AOA amoA was detected in the entire water column, but especially at the oxycline (up to 10^5 gene copies mL^-1 between ~20 and 40 m), during the summer 2006–2007. In 2005, 2006, and 2008, high AOA amoA abundance was determined below the oxycline (up to 10^5), during the summer 2006–2007. In 2005, 2006, and 2008, high AOA amoA abundance was determined below the oxycline (up to 10^5).
oxycline (40 m depth) during fall and winter (between April and August, Figure 2). The AOA amoA - to - Thaumarchaeota cell ratios ranged between $2.75 \times 10^{-7}$ and 0.72, showing higher numbers in spring followed by fall, summer and winter (Supplementary Figure 4). The number of bAOB amoA ranged from undetectable (i.e., < 2 copies mL$^{-1}$) during the 2005–2006 upwelling-favorable period up to 10$^4$ copies mL$^{-1}$ in the oxycline from mid-2006 onward, following the variability of AOA amoA, but usually centered at the oxycline and intermediate depths, when detected (Figure 2).

The comparison of bAOB and AOA amoA gene copies and with the AOA amoA/Thaumarchaeota cell ratios showed that the relative contribution of bAOB amoA was separated in spatial and seasonal scale; bAOB were favored at surface and intermediate waters but mainly during summer (Supplementary Figure 5A). Summertime was characterized by low ratios of archaeal amoA/Thaumarchaeota cells ratios (Supplementary Figure 5B). In fact, a plateau of ~181 bAOB amoA (i.e., ~2.26 as log-transformed value) was determined when AOA amoA/Thaumarchaeota cells ratios were >0.2, usually associated with surface and intermediate depths during summer (Supplementary Figure 5B).

The spatiotemporal differences in gene number or cell abundance between the studied microbial groups (except for Euryarchaeota) are shown in Figure 3. Data derived from Generalized Additive Models for Location, Scale, and Shape (GAMLSS) statistical analyses are shown in Table 1. In general, a significant decrease in microbial abundance was observed during winter, more evident for ammonia oxidizing assemblages and spring for Thaumarchaeota. GAMLSS analyses indicated that spatiotemporal variations (i.e., season and depth) account for >19.1% of the fluctuations, i.e., in non-photosynthetic picoplankton (24.3%), Thaumarchaeota, (45.6%), AOA amoA (34.8%) and bAOB amoA (19.1%) in the water column (see model 1 in Table 1).

The distribution of different archaeal ecotypes (N. maritimus, WCA and WCB) was studied only for selected months during winter and summer of 2011 in order to decipher the relevance of specific groups within the AOA amoA (Figure 4). The ammonia-oxidizing ecotypes presented similar trends with depth than the average values obtain at the time series distribution, showing a weak peak at the subsurface (~10 m depth) and higher values below 30 m depth for AOA amoA during both summer and winter months. N. maritimus amoA presented higher magnitudes followed by WCA, WCB, and bAOB amoA. The most relevant difference observed was that during winter, N. maritimus, WCA, and bAOB amoA peaked at intermediate depths (50 m depth), whereas during summer, WCB amoA maximum was determined at intermediate to bottom depths (50–90 m) (Figure 4).

Environmental Forcing of Microbial Communities, Including Ammonia-Oxidizers

The distribution of total non-photosynthetic picoplankton (flow cytometry) was significantly and positively associated with oxygen and nitrite, and negatively associated with nitrate and depth (Figure 5). Nitrite account for 15.1% of non-photosynthetic picoplankton spatiotemporal variability (Table 1). In contrast, Thaumarchaeota cell abundances were negatively correlated with oxygen, ammonium, and nitrite, and positively associated with nitrate and depth (Figure 5). Oxygen (45.9%), ammonium (25.2%) and nitrate (26.8%) were the explanatory variables for the Thaumarchaeota fluctuations (Table 1). A similar tendency was found for AOA amoA (Figure 5), but the most explicative variable was nitrate (15.6%) (Table 1). Except for depth, bAOB amoA were significantly correlated with ammonium, nitrite, and nitrate (Figure 5). Nitrate (11.3%) was the variable with the higher explanatory percentage to the bAOB amoA distribution in the time series (Table 1).

DISCUSSION

The biological activity in the study area is mainly influenced by seasonal upwelling events and its remote-forcing variability; see examples in the special volume for the COPAS time series (Escribano and Morales, 2012; Farias et al., 2015). In the present study, upwelling-favorable conditions were associated with higher phytoplanktonic biomass (chlorophyll) and abundances of total non-photosynthetic picoplanktonic communities, except during the 2005–2006 upwelling-favorable period for the non-photosynthetic picoplanktonic component (Figure 2). In fact, upwelling favorable winds cumulative intensity denoted an evident interannual upwelling variability during our study timeframe, characterized by a delayed and less intense events triggered during 2005–2006 (Supplementary Figure 1). The influence of upwelling variability on non-photosynthetic picoplankton has been reported in other seasonal upwelling ecosystems, such as, northwestern Indian Ocean (Wiebinga et al., 1997) and Benguela (Bergen et al., 2015) and agrees with previous results from the study area (Cuevas et al., 2004; Daneri et al., 2012).

In the study area, upwelling not only fertilizes the surface with high-nutrient content waters fueling primary and secondary productivity (e.g., Montero et al., 2007), but also develops a seasonal oxygen deficiency in the subsurface associated with Equatorial Subsurface Waters (ESSW) which was evident during our study (Ahumada and Chuecas, 1979; Sobarzo et al., 2007). Oxygen has been reported as a controlling factor of microbial community abundance (Eissler et al., 2010), structure (Aldunate et al., 2018), and biogeochemical active processes (Galán et al., 2014, 2017; Srain et al., 2020) in the study area. During our study, GAMLSS analyses indicated that the spatiotemporal variability in non-photosynthetic picoplanktonic cell abundance (studied by flow cytometry) was significant in the study area, and among the variables studied, oxygen and nitrite largely account for the changes observed (Table 1). Both, oxygen and nitrite are associated variables in subsurface waters, since nitrite accumulation depends on the redox conditions in the water column triggered by oxygen deficiency (e.g., Farias et al., 2015). Nitrite is a key intermediary of many biogeochemical processes, i.e., as a product of aerobic ammonia oxidation and
FIGURE 1 | Spatiotemporal variability of dissolved oxygen, ammonium, nitrite, and nitrate at the time series station (St.18). Dots indicate specific date and depth samplings.
FIGURE 2 | Spatiotemporal variability of chlorophyll concentration, non-photosynthetic picoplankton abundance, and amoA gene counts for general AOA and bAOB at the time series station (St.18). Dots indicate specific date and depth samplings.
nitrate reduction, and as a substrate for aerobic nitrite oxidation, anammox and nitrite reduction related with heterotrophic and chemoautotrophic denitrification (Ward, 2008). In fact, coupling of keystone OMZ microbial communities such as SUP05 clade with anammox and denitrifying sulfur-oxidizers were determined to be associated to nitrite based in a biogeochemical model in the Saanich Inlet seasonally anoxic fjord (Louca et al., 2016) and in bottom waters during upwelling periods in the study area (Galán et al., 2014).

Variability of Euryarchaea Cells Using CARD-FISH

Euryarchaea and Thaumarchaea presented a differential temporal and spatial distributions in the water column. Euryarchaea was characterized by a higher contribution at surface seawater at fall or spring. This result agrees with other findings on a temporal rather than a spatial scale since Euryarchaeota showed a narrower distribution than previously reported (Levitan et al., 2007a; Quiñones et al., 2009). However, sampling methodological constraints could influence these spatial differences, that is, our study was focused on free-living fraction cell counts, whereas the mentioned reports involved bulk DNA analyses (7 L samples) that favor the presence of different particle sizes. Particle aggregates seem to enhance the contribution of Euryarchaea off northern Chile (Ganesh et al., 2014) and in the study area (Levitan et al., 2007a), where methylotrophic methanogens were abundant and viable at depth in a large-particle fraction (between 0.22 and 25 μm). Moreover, differences in vertical distribution patterns of Euryarchaea caused by the filter size fraction have also been reported in contrasting areas near ALOHA time series station (Lincoln et al., 2014) and Oregon coast (Smith et al., 2013). In addition, our results
could indicate that using CARD-FISH distinct phylotypes were detected, associated with euphotic MGII during 2006–2007, and also at subsurface during 2008, supported by previous findings of a wider distribution and diversity within *Euryarchaeota* as in other coastal areas, e.g., SPOT (Parada and Fuhrman, 2017).

**FIGURE 4** | Vertical profiles of the main ammonia-oxidizer ecotypes (*amoA* gene counts in a log-transformed scale) at the St.18 during two contrasting oceanographic months, that is, a representative month for the unfavorable (August 2011, austral winter) and favorable conditions (December 2011, austral summer) for upwelling. For the sake of comparison, the seasonal average for βAOB and general AOA are shown in blue. Error bars are standard deviations.
The Spatiotemporal Distribution of *Thaumarchaeota* Cell Abundances

Unlike Euryarchaeota, thaumarchaeal cells were found to be favored in subsurface waters, characterized by peaks at the oxycline during fall and spring (2007) and maxima at the bottom during wintertime and late summer 2008. Thaumarchaeal cells made up the bulk fraction of the prokaryote community at deep waters and represented a large fraction of the archaea considering the sum of Euryarchaeota and Thaumarchaeota (Supplementary Figure 3). These results are supported by previous findings in the Eastern South Pacific (ESP) off Chile, based on rRNA dot blot hybridization (Levican et al., 2007b; Quiñones et al., 2009) and lipid-based approach (Rossi et al., 2012; Srain et al., 2015). GAMLSS analysis indicates that *Thaumarchaeota* was the microbial group with the greater percentage of predictability associated with its spatial and temporal variability (Table 1), and with all the other variables (oxygen, nitrate, ammonium and nitrate), associated with substrates and products of nitrification. In general, these results support a recent report considering the water column position (euphotic, 50 m, and deeper waters 60 ñ 500 m) and other factors such as hydrographic explanatory variables, including oxygen, nitrite and ammonium *Thaumarchaeota* contribution to total microbial community (iTAG-16S rRNA) variability in the Monterey Bay coastal time series (Tolar et al., 2020). However, in the case of the time series off Concepción the *Thaumarchaeota* depth distribution is resolved within a shallow water column depth (92 m depth).

Contribution of Ammonia Oxidizing Archaea in the Study Area

The AOA *amoA* abundance results indicate that AOA in the study area reached comparable abundances with those determined in other marine ecosystems, e.g., the northern Gulf of Mexico (up to $10^5$ copies mL$^{-1}$ at ≤100 m depth; Tolar et al., 2013), Sargasso Sea or ETNA (up to $10^5$ copies mL$^{-1}$, Löschter et al., 2012; Newell et al., 2013), North Sea in winter months (from $10^4$ up to $10^5$ copies mL$^{-1}$, Wuchter et al., 2006; Herfort et al., 2007; Pitcher et al., 2011b), and within the oxycline and just over the upper limit of the OMZ in the Arabian Sea (up to $10^5$ copies mL$^{-1}$, Newell et al., 2011; Pitcher et al., 2011a; Bouskill et al., 2012). However, variable ratios between *amoA* and 16S rRNA genes or cell counts have been observed in marine ecosystems (e.g., Wuchter et al., 2006; Mincer et al., 2007; Agogué et al., 2008). In our study, a variable contribution of AOA *amoA* versus *Thaumarchaeota* cell counts determined through CARD-FISH was observed, reaching higher ratios during spring and fall (0.2 ñ 0.7) compared with <0.2 ratios observed during summer and winter (Supplementary Figure 4). Similar values were obtained in surface waters from Monterey Bay (Mincer et al., 2007) and in subsurface (100 ñ 150 m depth) from subtropical and equatorial regions of the North Atlantic (Agogué et al., 2008). However, lower ratios of AOA *amoA* to *Thaumarchaeota* 16S rRNA gene ratios (<0.01) were associated with the presence of non-nitrifying Thaumarchaeota from bathypelagic waters (Agogué et al., 2008) or with methodological biases (Mincer et al., 2007). Alternatively, different ecotypes could account for a wide range of AOA *amoA* versus *Thaumarchaeota* 16S rRNA gene ratios (e.g., Sintes et al., 2013). GAMLSS modeling indicates that similar predictive variables account for both *Thaumarchaeota* cell and AOA *amoA* fluctuation (Table 1).

Contribution of Ammonia Oxidizing Archaea Ecotypes in a Representative Upwelling Versus Non-upwelling Season Profile

AOA in the study area were found to be associated to the candidate order *Nitrosoarchaeales* (Stieglmeier et al., 2014), and...
to other ecotypes, i.e., Water Column Surface clade (WCA) with one culture Candidate Nitrosopelagicus brevis and Water Column Deep clades (WCB), as in other coastal marine time series (e.g., Beman et al., 2008; Santoro et al., 2010, 2015; Reji et al., 2019; Tolar et al., 2020). The ammonia-oxidizing assemblages were predominantly characterized by *N. maritimus*-like followed by WCA *amoA* reaching higher abundances during winter, whereas the deep WCB *amoA* ecotype showing maxima during summer at the bottom. This result was supported by previous reports in the study area based on AOA *amoA* survey using clone libraries (Molina et al., 2010), and iTag sequencing (Bertagnolli and Ulloa, 2017) using primers designed by Pester et al. (2012). These previous studies revealed specific predominant contribution of the WCA *amoA* in both upwelling and non-upwelling season and of the deep WCB *amoA* during the upwelling period. The WCB was strongly coupled to oxygen deficient conditions, and was suggested to be advected by upwelling waters toward the continental shelf (Bertagnolli and Ulloa, 2017). Indeed, WCA and *N. maritimus* *amoA* ecotypes are the main AOA in the study area in terms of the abundances of *amoA* genes and transcripts, and related to ammonium oxidation (Molina et al., 2010; Bristow et al., 2016; Levipan et al., 2016). In addition, both archaeal ecotypes, but mainly WCA, appear to have an important role in the ammonium oxidation in other marine ecosystems; e.g., the Gulf of California (Beman et al., 2008), Monterey Bay (Smith et al., 2014; Tolar et al., 2020), from the equatorial Pacific to the Arctic (Shiozaki et al., 2016), and the suboxic zone of the central Baltic Sea (Labrenz et al., 2010). Nonetheless, *N. maritimus*-like ecotypes may be less abundant at lower latitudes of the ETSP (~12 and 18°S; Peng et al., 2013).

### Ammonia Oxidizing Bacteria and Its Interaction With Their Archaea Counterparts in the Study Area

The abundance of bAOB *amoA* ranged from the limit of detection (2 copies) to 10⁴ copies mL⁻¹ (*Figures 2, 4*), reaching higher numbers mainly during spring-summer at intermediate depths. This range was lower than that previously described in the ETSP (10³ to 10⁵ copies mL⁻¹, Bouskill et al., 2012; Lösch et al., 2012), similar to the range reported by Wuchter et al. (2006) in the North Sea, and wider than that found in locations such as Monterey Bay (undetectable to 10³ copies mL⁻¹, Mincer et al., 2007), shallow waters in the California Current (undetectable to 10² copies mL⁻¹, Santoro et al., 2010, 2013), Gulf of California or ETNP (undetectable to 10² copies mL⁻¹, Beman et al., 2008, 2012), and within the oxycline in the Arabian Sea (from ~10² to 10³ copies mL⁻¹, Newell et al., 2011; Bouskill et al., 2012).

Spatiotemporal followed by nitrate and ammonium were significant explanatory variables for bAOB *amoA*, distribution. In addition, the variability in bAOB *amoA* appears to be constrained by AOA distribution. Shifts in bAOB *amoA* were more obvious when plotted against the AOA *amoA* versus *Thaumarchaeota* cell ratios estimated from CARD-FISH counts (Supplementary Figure 5B). These results show that the number of bAOB *amoA* was maximum in surface waters during summer-fall when low AOA *amoA* versus *Thaumarchaeota* cell ratios were found, and reached a plateau at spring-time. This finding suggests that competitive interactions between AOA and bAOB *amoA* are more intense in surface waters during springtime and were potentially associated with lower ammonium availability and/or oxygen deficiency. The numerical dominance of AOA *amoA* in the study area could be attributed to the well-known kinetic differences in ammonium uptake between AOA and bAOB (Martens-Habbena et al., 2009). It is known that ammonium concentration promotes niche separation not only between phylogenetically distant species of AOA and AOB (Schleper, 2010), but also between different clusters of AOA (Sintes et al., 2016). In the present study, ammonia-oxidizing assemblages significantly varied with ammonium (model 5 in *Table 1*), suggesting niche overlapping between AOA and AOB and hence substrate competition for ammonium. Nevertheless, in contrast to Fernandez and Farias (2012), who reported that bacterial Ammonium oxidation might benefit from low oxygen concentrations, we found that the oxygen concentration was a statistically meaningful predictor for AOA *amoA* but not for bAOB *amoA* distribution (models 3 in *Table 1*), suggesting that oxygen concentration can be involved in niche differentiation between AOA and bAOB *amoA* (models 1 and 3 in *Table 1*).

Moreover, our results support the key role that oxygen concentration plays in AOA-A and B ecotypes partitioning in this and other areas (Beman et al., 2008; Bertagnolli and Ulloa, 2017), as well as in controlling (at low concentrations) the distribution and activity of marine AOA populations (Qin et al., 2017).

Ammonia oxidizing assemblages are significant contributors to biogeochemical cycling in the study area associated with high rates of nitrification that are correlated with the availability of ammonium mainly in the oxyclines in the study area (Fernandez and Farias, 2012). Moreover, in a 10-year (2002–2012) biogeochemical report associated with nitrous oxide dynamics the COPAS time series, nitrous oxide positive excess, mainly in the oxyclines and subsurface waters, was associated with nitrification (Farias et al., 2015; Galán et al., 2017), as in other oxygen deficient areas of the Pacific Ocean, e.g., Lösch et al. (2012) and Frey et al. (2020). In order to compare potential role of microbial communities studied here, the average monthly inventories of nitrous oxide reported by Farias et al. (2015) for the 2005–2009 upwelling periods were compared with nitrate, ammonium, AOA, bAOB and non-fluorescent picoplanktonic community estimated inventories in our study (Supplementary Figure 6 and *Table 2*). This comparison helps us identify the connection between ammonia oxidizing assemblages with Nitrous oxide and nitrate mainly during 2007–2009, but not for the previous upwelling periods. During 2005–2007, nitrous oxide and nitrate inventories could be associated with high ammonium accumulation and an enrichment of picoplanktonic communities with unknown identity in the bottom (Supplementary Figure 6). Therefore, changes in microbial assemblages are associated with upwelling variability, generating significant impact in the water column biogeochemistry in this natural oxygen-deficient upwelling ecosystem.
CONCLUSION

In total, our results indicate the physical-chemical oceanographic conditions associated with the seasonal upwelling systems modulate the distribution and dynamics of abundant marine microbial groups including key ammonia oxidizing assemblages. The statistical GAMlSS analyses indicate that spatiotemporal changes associated with biogeochemical conditions trigger by oxygen-deficiency, including substrate and products of nitrification, were significant predictors shaping the marine microbial community distribution. Biological interactions including AOA and bAOB, and possibly with other microbial groups in surface waters influences the temporal dynamics of ammonia oxidizing assemblages. Considering the key role that coastal marine microbial community plays in the cycle and the balance of nutrients and carbon at a global scale, an appropriate understanding of their natural variability will allow to better predict the metabolic and biogeochemical consequences associated with the expected expansion and intensification of oxygen deficient zones due to anthropogenic forcing.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article are given in Supplementary Table 2 and complementary data will be made available by the authors, without undue reservation.

AUTHOR CONTRIBUTIONS

VM, LB, and HL wrote the manuscript with the support from SR-F, CA, AG, IM, and OU. VM and LB conceived the original idea. VM and OU supervised the project. All the authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmars.2020.561597/full#supplementary-material

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