Research Paper

Microbial communities associated with sinking particles across an environmental gradient from coastal upwelling to the oligotrophic ocean

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\textbf{ABSTRACT}

Complex interactions between microbial communities in the epipelagic and mesopelagic ocean drive variability in sinking carbon flux as part of the biological carbon pump. To investigate the relative contribution of dominant prokaryotic, protistan, and phytoplankton taxa to sinking particles, we used 16S/18S rRNA gene sequencing of particles collected in sediment traps and water column communities. Comparisons were done across a coastal upwelling-to-oligotrophic environmental gradient in the southern California Current Ecosystem to assess effects of natural environmental variability on taxon-specific contributions to sinking particle export. Particle-associated microbial assemblages differ from ambient mixed-layer communities. Gammaproteobacteria, dinoflagellates, and rhizarians were dominant microbes associated with sinking particles at all sampling locations, with diatoms increasing significantly at the inshore mesotrophic site. Parasitic groups, syndiniales and apicomplexans, were also major particle-associated taxa. Relative sequence abundance on exported particles from the mesotrophic inshore site more closely resembled the water-column dominants (especially diatoms) found above, while oligotrophic communities exhibited greater dissimilarity between the microbial communities on sinking particles and in the mixed layer. Protists generally showed greater similarity between mixed-layer and sinking particle communities than prokaryotes. Synechococcus was over-represented on sinking particles (relative to eukaryotic phytoplankton and other cyanobacteria) only in oligotrophic areas. Diatoms were the only phytoplankton group consistently over-represented on sinking particles across the region. Our results highlight important variability in microbial contributions to export that seem to align with differences in dominant grazing pathways. At the upwelling influenced site where export was primarily due to the fecal pellets of large copepods, diatoms were dominant contributors to export and protistan communities on sinking particles resembled those in the water column above. When doliolids were abundant, dinoflagellates contributed substantially to export, while the dominant picoeukaryote (Ostreococcus) was only a minor contributor. In oligotrophic areas, where grazing pathways were dominated by protists and sinking particles were primarily amorphous aged aggregates, there was little similarity between mixed layer and sinking particle communities.

1. Introduction

Predicting changes in marine carbon sequestration requires a detailed understanding of the ocean’s biological pump (BP) that transports organic carbon fixed by phytoplankton into the deep ocean (Buesseler and Boyd, 2009; Ducklow et al., 2001; Martin et al., 1987). The BP transforms organic particles physically and chemically through multiple food-web pathways and processes spanning all domains of life and spatiotemporal scales (Burd et al., 2016; Frischknecht et al., 2018). Export and sequestration efficiency are shaped not only by surface primary production, but also by particle sinking speeds and remineralization rates (Cram et al., 2018; McDonnell et al., 2015), the microbial ecology of sinking particles (Pelve et al., 2017; Piontek et al., 2009), and flux-feeding taxa in the mesopelagic (Jackson et al., 1993; Stukel et al.,...
2019b). These interacting processes, with varying importance in different regions and seasons, complicate simple predictions of export flux from primary production and necessitate integrative studies that link ecology and biogeochemistry (Burd et al., 2016; Siegel et al., 2016).

Advances in genetic characterization of communities, especially when combined with existing sediment trap methodologies, have recently allowed new investigations of microbial communities that create, modify, colonize, and remineralize sinking particles. These studies have shown, for instance, that Synechococcus is overrepresented on sinking particles relative to Prochlorococcus (Amacher et al., 2013) and uncovered an unexpectedly large role for Rhizaria in sinking flux (Guidi et al., 2016; Gutierrez-Rodriguez et al., 2019). They have also provided evidence of significant community shifts on sinking particles (Fontanez et al., 2015; Gutierrez-Rodriguez et al., 2019; Pelve et al., 2017) and vertical connectivity between the euphotic zone and bathypelagic realm (Mestre et al., 2018; Ruiz-Gonzalez et al., 2020). Time-series studies at abyssal depths have further documented relatively consistent dominance by piezophilic groups, variable contributions of eukaryotic taxa (Stoeo et al., 2019), and even major episodic flux events attributed to single diatom species (Preston et al., 2019). To date, however, most such studies have focused on a single ecological condition (e.g., the Sargasso Sea. Amacher et al., 2013) or have not quantified sinking particles and attached microbes (e.g., Guidi et al., 2016; Mestre et al., 2018).

In the present study, we investigate spatial patterns in suspended and sinking eukaryotic and prokaryotic communities in the California Current Ecosystem (CCE). The CCE is an excellent natural laboratory for such studies, because it includes an environmental gradient ranging from coastal upwelling to the edge of the oligotrophic North Pacific Subtropical Gyre within a compact spatial region (Ohman et al., 2013; Stukel and Barbeau, 2020). The coastal upwelling zone is characterized by high biomass and dominance of larger primary producers (Venrick, 2002; Taylor et al., 2015), and exported particles are mostly composed of zooplankton fecal pellets (Knauer et al., 1979; Stukel et al., 2013; Morrow et al., 2018). In contrast, the offshore open-ocean region has low biomass and dominance of small cells (Taylor and Landry, 2018), and exported particles are mainly amorphous aggregates (Knauer et al., 1979; Stukel et al., 2013; Morrow et al., 2018). Using Lagrangian drifters to follow marked water parcels on process cruise P1604 (April–May 2016) of the CCE Long-Term Ecological Research (CCE LTER) Program, we combine traditional biogeochemical and standing stock measurements with 16S and 18S rRNA gene sequencing of microbial communities for particles collected in sediment traps and sampled from the mixed layer, base of the euphotic zone, and 150 m. We focus on two hypotheses: (1) sinking material more closely resembles the water-column dominants in productive coastal waters than offshore oligotrophic waters due to larger taxa (e.g., diatoms, larger mesozooplankton), shallower euphotic depth, and more rapid, direct sinking of POM in coastal waters; and (2) the composition of eukaryotic taxa is more similar between water-column and sinking particles than prokaryotic taxa because the larger eukaryotes are more likely to sink rapidly as aggregates or in zooplankton fecal pellets.

2. Methods

2.1. Sample collection and environmental measurements

Data are derived from the P1604 Process cruise of the CCE LTER Program (19 Apr - 12 May 2016; Kelly et al., 2018; Nickels and Ohman, 2018). The cruise took place during the end of the 2015–2016 El Nino, which followed the North Pacific marine heat wave of 2014–2015 (Bond et al., 2015; Jacob et al., 2016; Kahrn et al., 2018). The region had thus experienced excessively warm conditions with reduced upwelling for some time, although upwelling resumed near the coast during our cruise. The cruise sampling plan featured four Lagrangian experiments of ~3–5 day-duration, during which two drifting arrays with 3 × 1-m drogues were used to track mixed-layer communities (Landry et al., 2009). One array held sediment traps (see below), while the other contained attachment points for daily in situ incubations conducted to measure primary productivity using H14CO3− uptake (6 depths per day), among other measurements (see Morrow et al., 2018). Daily CTD casts were conducted at 02:00 (to collect water for daily incubations and standing stock measurements at 6 depths in the euphotic zone) and at approximately local noon (to determine photosynthetically active radiation profiles). Nutrients were measured using an autoanalyzer. Chlo- rophyll (Chl) a (total and size-fractionated) was measured using the acidification method. Heterotrophic bacteria, Prochlorococcus, Synecho- coccus, and picoeukaryotes were enumerated by flow cytometry. Mes- zozooplankton biomass and community composition was determined from paired day-night 202-μm mesh bongo net samples (Morrow et al., 2018; Nickels and Ohman, 2018). Samples for DNA analyses were collected from Niskin bottles at the mixed layer, the base of the euphotic zone, and 150 m (Table S1). Seawater samples (280 mL, 200 μm Nitex screen or 650 mL, 500 μm Nitex screen) were filtered onto 0.2-μm Supor membrane filters (Pall), flash-frozen, and stored at ~80°C until analysis. VERTEX-style cylindrical sediment traps (12 tubes per depth) were deployed on the trap array at fixed depths of 100 and 150 m, with additional traps placed at the base of the euphotic zone at the two sites closest to shore (Knauer et al., 1979; Morrow et al., 2018). Traps with top baffles were deployed with a formaldehyde brine (filtered seawater + 50 g L−1 NaCl + formaldehyde (0.4% final concentration); for most analyses), an unpoisoned brine (for sequencing), or RNA later (also for sequencing, prepared following methods in Fontanez et al., 2015). Independent 234Th deficiency measurements confirmed that the traps had no consistent over- or under-collection bias (Stukel et al., 2019a). After recovery, all tubes were filtered through a 200-μm filter, which was then carefully inspected by stereomicroscope and mesozooplankton ‘swimmers’ were removed, before the remainder of the >200-μm material were frozen at ~80°C. Tubes were then split and filtered for analyses including POC and PN, Chl a, pheopigments, fecal pellet mass flux, 234Th, and sequencing. Samples from the unpreserved (live) and RNA later traps were immediately filtered through Sterivex filters (Millipore, 0.22 μm) and frozen at ~80°C for 18S/16S rRNA gene sequencing analyses. Samples for POC and PN were filtered through pre-Combusted GF/F filters, frozen, acidified on land, and analyzed by an elemental analyzer coupled to an isotope ratio mass spectrometer. Samples for Chl a and pheopigments were filtered through GF/F filters and analyzed by the acidification method. Samples for fecal pellet mass flux were analyzed using a stereomicroscope (20X magnification). Fecal pellets were categorized by shape and morphometric measurements were determined by image analysis and qualitative characteristics of sinking material (e.g., dominance by fecal pellets or aggregates) were noted. Samples for 234Th were filtered through QMA filters and analyzed using a RISO low-level beta multi-counter. For additional details on other analyses and general sediment trap processing, see Morrow et al. (2018) and Stukel et al. (2019a).

The Lagrangian experiments sampled four regions of differing trophic conditions in the CCE: 1) offshore oligotrophic (OO) waters on the edge of the subtropical gyre, 2) core water of the California Current (CC), 3) the transition zone (TZ) region where inshore mesotrophic waters were mixing with the edge of the California Current, and 4) inshore (IN) nutrient-rich water showing evidence of recent upwelling. Additional environmental data from these experiments are available on the CCE LTER Datazoo website.

2.2. Library construction and sequencing

DNA from water-column and trap samples was extracted using the NucleoMag 96 Plant kit (Macherey Nagel) following the manufacturer’s instructions. Although DNA was extracted separately for the sediment trap particles <200 μm and >200 μm, these were pooled for subsequent analysis. DNA was amplified by Polymerase Chain Reaction using the Q5
high-fidelity PCR kit (New England Biolabs). Prokaryotes were characterized by amplification of the V4–V5 regions of the 16S small subunit ribosomal RNA gene (SSU-rRNA) using primers 515F and 926R (Quince et al., 2011) (Table S2). Eukaryotes were characterized by amplifying the V9 region of the 18S rRNA gene using primers 1389F and 1510R (Amaral-Zettler et al., 2009) (Table S2). For sediment trap samples that did not amplify during PCR, we used the OneStep™ PCR Inhibitor Removal Kit (Zymo Research) to remove inhibitory substances. The PCR products were pooled in equimolar amounts (~10 ng µL⁻¹) and sequenced using a dual-barcode index on an Illumina MiSeq platform at the Institute for Genomic Medicine (IGM, University of California, San Diego). Initial quality control of the raw sequence reads was done using the workflow for read filtering, swarm Operational Taxonomic Unit (OTU) clustering, and taxonomic classification of the SSU-rRNA written by JP McCrow (https://github.com/allenlab/rRNA_pipel ine). Demultiplexed raw reads were provided by IGM and have been deposited in NCBI under BioProject PRJNA445287 and Biosample accession numbers SAMN08784582–SAMN08784552 for 18SrRNA gene sequences, and under BioProject PRJNA422420 and Biosample accession numbers SAMN08784494–SAMN08784464 for 16S rRNA gene sequences. Additional processing details are in the Supplementary Information. The total number of sequence reads and OTUs are summarized for each primer set in Table S3 and by sample types in Table S4 (prokaryotes) and Table S5 (eukaryotes). These results show relative contributions of different taxa in each sample, but should not be interpreted as differences in absolute abundance of taxa across samples (Gong and Marchetti, 2019; Vandeputte et al., 2017). Indeed, the abundance of heterotrophic bacteria at the surface (determined by flow cytometry), varied from 3.2 × 10⁶ cells mL⁻¹ at the OO site to 4.8 × 10⁶ cells mL⁻¹ at the IN site, while surface Chl a (proxy for phytoplankton biomass) varied by a factor of 40.

2.3. Statistical analyses

We evaluated differences in prokaryotic (16S) and eukaryotic (18S) assemblages at the genus level by multivariate analyses based on the Bray-Curtis dissimilarity index. Prior to analyses, the sequence reads were merged by genera (prokaryotes: 379; eukaryotes: 510), and only those contributing >1% to the relative abundances in any sample (prokaryotes: 79; eukaryotes: 97) were included in the analyses. For the genera selected, relative abundances per sample were recalculated and square-root transformed to reduce the impact of the most abundant microbes. Community differences were assessed by hierarchical clustering and ordination using non-metric multidimensional scaling (nMDS). Cluster significance was established by similarity profile analysis (SIMPROF). Multivariate analyses were done in R using the packages ‘vegan’ (Oksanen et al., 2017) and ‘clustsig’ (Whitaker and Christman, 2014).

3. Results

3.1. Environmental conditions and export fluxes

Our drifter experiments covered a seven-fold range in productivity (229–1659 mg C m⁻² d⁻¹) and a six-fold range in sediment-trap productivity fluxes at 150 m (32–202 mg C m⁻² d⁻¹) (Table 1, Figs. 1 and 2). Environmental conditions in offshore oligotrophic (OO) and California Current waters (CC) were marked by deep mixed layers (50–80 m), low nutrients (0.02–0.08 µM nitrate), low chlorophyll (0.1–0.2 µg Chl a L⁻¹), and low primary production (PP, 229–276 mg C m⁻² d⁻¹) (Table 1, Fig. 1b–d, Fig. 2a–b). In contrast, conditions in the transition zone (TZ) and inshore (IN) areas had shallower mixed layers (30 and 12 m, respectively) with elevated nutrients, Chl a, and PP compared to the offshore region. Mixed-layer nitrate concentrations were highest in the TZ region (4.6 vs 2.1 µM), but Chl a and PP were greatest at the IN site (4.2 vs 1.2 µg L⁻¹ and 1659 vs 868 mg C m⁻² d⁻¹, respectively). There was also a shift towards larger cells in the upwelling region. The <1-µm size fraction accounted for 59% of chlorophyll at the OO site, while the median (chlorophyll-weighted) cell size was between 3 and 8 µm at the IN site. The transition from offshore (OO) to coastal (IN) regions was also associated with order-of-magnitude increases in mesozooplankton community biomass and grazing (Morrow et al., 2018; Nickels and Olman, 2018). Mesozooplankton communities were typically dominated by crustaceans (especially copepods), except for a high biomass of doliolids (pelagic tunicates) in the TZ region.

The composition of sinking particles collected in sediment traps also shifted across the productivity gradient (Fig. 2c–g). In the upwelling region, export was dominated by recognizable mesozooplankton fecal pellets, primarily cylindrical pellets believed to be crustacean egesta. Recognizable pellets contributed nearly all of the carbon flux at the IN site and approximately 50% in the TZ area (Fig. 2d and f). In contrast, recognizable pellets were a negligible component of sinking material in most traps in the offshore region, where most of the carbon was unrecognized detrital material and aggregates. IN and TZ traps also had substantially higher pigment:carbon ratios than OO or CC traps, reflecting rapid export of pigmented material from the euphotic zone (Fig. 2e). The pigments were predominantly phaeopigments (degradation products of Chl a) formed during digestion in mesozooplankton guts, suggesting that phytoplankton were consumed by grazers prior to export (Fig. 2g). These ecological and biogeochemical differences are typical of the gradient in ecosystem characteristics across the CCE and underlie the >6-fold difference in POC export fluxes into the 150-m

| Table 1 |
| Summary of environmental data during sediment trap drift deployments across a gradient of ecosystem conditions in the California Current Ecosystem. PP: primary production (mg C m⁻² d⁻¹), POC flux: export of particulate organic carbon (mg C m⁻² d⁻¹), Temperature (°C), Chl: chlorophyll (µg L⁻¹), NO₃: nitrate (µg L⁻¹). Number of chlorophyll-nitrate measurements per depth and CTD casts were as follows: OO: (n: 3, 4), CC: (n: 7, 10), TZ: (n: 6, 11), and IN: (n: 7, 11). Mean ± standard error. |

| Inshore (IN) |
| PP Depth (m) | POC flux | Temperature | Chl | NO₃ |
| 1658.7 ± 195.6 | 0–5 | 14.42 ± 0.05 | 4.20 ± 0.15 | 1.13 ± 0.04 |
| 47 | 251.0 ± 8.3 | 10.39 ± 0.04 | 0.42 ± 0.06 | 24.27 ± 2.07 |
| 147 | 202.3 ± 14.0 | 8.82 ± 0.04 | – | – |

| Transition Zone (TZ) |
| PP Depth (m) | POC flux | Temperature | Chl | NO₃ |
| 868.0 ± 133.8 | 0–5 | 13.63 ± 0.05 | 1.01 ± 0.03 | 3.45 ± 0.28 |
| 57 | 120.4 ± 9.7 | 11.24 ± 0.06 | 0.25 ± 0.06 | 13.21 ± 0.71 |
| 147 | 77.4 ± 4.4 | 8.78 ± 0.01 | – | – |

| California Current (CC) |
| PP Depth (m) | POC flux | Temperature | Chl | NO₃ |
| 275.7 ± 41.1 | 0–5 | 15.20 ± 0.02 | 0.13 ± 0.01 | 0.07 ± 0.04 |
| 97 | 40.2 ± 3.8 | 13.18 ± 0.08 | 0.34 ± 0.06 | 4.94 ± 1.67 |
| 147 | 37.0 ± 5.0 | 9.54 ± 0.03 | – | – |

| Oligotrophic Offshore (OO) |
| PP Depth (m) | POC flux | Temperature | Chl | NO₃ |
| 229.0 | 0–5 | 17.26 ± 0.12 | 0.07 ± 0.00 | 0.03 ± 0.02 |
| 100 | 72.1 ± 7.3 | 12.22 ± 0.05 | 0.16 ± 0.05 | 11.16 ± 1.68 |
| 150 | 31.8 ± 4.9 | 9.51 ± 0.01 | – | – |
traps deployed at the OO and IN sites (Table 1).

3.2. Major groups from sequence analyses: Eukaryotes

We collected water-column samples from multiple depths (typically mixed layer, base of the euphotic zone, and 150 m) and analyzed them for 18S and 16S rRNA gene sequences (Fig. 3). In addition to the protistan reads determined from 18S, we identified numerous metazoan sequences (Fig. 3). Within our working OTU table, which excludes singletons and merges sequence reads of over-split OTUs, metazoans accounted for 1%–63% of total reads, with an arithmetic mean of 27% across all depths and sites. Most of the metazoan sequences are ascribed to copepods, especially the calanoids *Metridia* and *Calanus* spp., both important consumers and vertical migrators in CCE waters (Supp. Fig. 1). Doliolid sequences are also present due to a bloom of *Doliolum denticulatum* during the study period, with highest densities in the TZ. Percent metazoan reads typically increased with depth, except at the TZ site, which had high doliolid abundances in the mixed layer. In contrast to the water column, metazoan sequences were more abundant than protistan sequences (84%–16%) in the sediment traps (mean: 82%), despite the fact that all recognizable >200-μm mesozooplankton “swimmers” were removed from those samples during examination under a stereomicroscope (Fig. 3).

Protistan communities in mixed-layer samples are co-dominated by alveolates and stramenopiles at the IN site, chlorophytes (Archeaplastida) are more important in the TZ, and alveolates dominate the OO and CC sites (Fig. 4). Clear shifts were also evident with depth. At the IN and TZ sites, the contributions of stramenopiles and Archaeplastida, respectively, decreased consistently with depth (Fig. 4). In contrast, Rhizaria increased in proportional abundance at deeper depths at all sites. They accounted for ~30% of sequences at 150 m at the IN station and approximately half of the sequences at the other three sites (Fig. 4).

Protistan communities on sinking particles collected by sediment traps were typically dominated by parasitic alveolates (apicomplexan and syndiniales) and rhizarians, although their contributions varied among sampling locations (Figs. 4 and 5). In the IN traps, alveolates and stramenopiles are all important (Fig. 4). At the TZ site, the large contribution of chlorophytes (Archaeplastida) to the water-column...
assemblage is not seen in the trap samples. In the CC traps, alveolate sequences dominate (except for rhizarians in one live trap), while both alveolates and rhizarians are important in the OO trap samples (Fig. 4).

### 3.3. Water-column community composition: Eukaryotes

18S rRNA gene sequences from the mixed layer showed substantial variability in protistan communities across the environmental gradient (Fig. 5, Table S6). At the coastal upwelling-influenced IN site, stramenopiles (especially the diatoms *Thalassiosira* and another unclassified Bacillariophyceae) were particularly important, while the chlorophyte *Micromonas* was another important contributor (Fig. 5). TZ samples were characterized by very high abundances of Archaeplastida sequences, including the chlorophyte *Ostreococcus*, which was responsible for 44% of protistan reads. At both of the offshore sites (CC and OO), the protistan community was dominated by alveolates, with high sequence abundances of an uncultured dinoflagellate (16% of reads at OO; 18% at CC). The dinoflagellate *Glenodiniopsis* contributed 28% of mixed-layer sequences at OO, and a third dinoflagellate, *Karlodinium*, was common across the region (from IN to OO) and at multiple depths from the euphotic zone to 150 m. Besides alveolates, the Acantharian *Chaunacanthida* was particularly important at 150 m, comprising 26%, 55%, and 32% of total protistan reads at the IN, TZ, and OO sites, respectively (Fig. 5). At all sites, multiple groups of marine alveolates (including cultured and uncultured dinoflagellates as well as MALV-I, MALV-II, and MALV-III) comprised most of the remaining sequences.

We used non-metric multidimensional scaling (nMDS, Fig. 5a) and dendrogram (Fig. 5b) analyses based on the Bray-Curtis dissimilarity index, with broadly similar results, to compare assemblage differences across sites, depths, and sampling type (water column or sediment trap). For water-column samples, microbial community structure clusters mainly by depth, although mixed-layer assemblages also differ among locations (Fig. 5). The 150-m samples from all sites are characterized by high concentrations of acantharians and polycystine radiolarians (Fig. 5b, Table S6). Samples from the euphotic zone base in the TZ, CC, and OO regions cluster in a single group that is closely aligned with mixed-layer samples from the CC and OO sites. Euphotic zone samples (both mixed layer and euphotic zone base) from the IN site cluster more...
Fig. 3. Number of sequence reads of eukaryotes (18S rRNA, top panel) and prokaryotes (16S rRNA, bottom panel) in water-column and sediment-trap samples across an environmental gradient in the California Current Ecosystem. IN: inshore, TZ: transition zone, CC: California Current, OO: oligotrophic offshore.

Fig. 4. Major groups of protists in the water column and sinking particles across an environmental gradient in the California Current Ecosystem, assessed by 18S V9 rRNA. Sinking particles were collected at two depths (base of the euphotic zone and 150 m) and trap tubes were filled with either a brine (live traps) or a RNA later solution (fixed traps).
closely with sediment traps than other water-column samples, while the TZ mixed-layer assemblage is not similar to any other samples.

3.4. Sediment trap community composition: Eukaryotes

Alveolates and stramenopiles were all important in the IN traps, with high relative abundances of sequences assigned to the diatom *Thalassiosira* and syndiniales MALV III (Fig. 5b, Table S6). Alveolate sequences assigned to the apicomplexan Cephalodophoroidea (likely parasitic protists) dominated protistan reads on sinking particles, especially at the TZ and CC sites (Fig. 5b). Alveolates and rhizarians were particularly important in OO samples. An uncultured dinoflagellate comprised ~10% of the total protistan sequence reads in this region, while the rhizarians included Chaunacanthia (class Acantharia) and Spumellaria (class Polycystina) (Fig. 5b). *Karolidinium*, an important dinoflagellate in water-column samples, contributed substantially to all trap samples (1–10% of reads), although it was only over-represented on sinking particles relative to water-column samples at the OO site (5–10% of reads) (Fig. 5b).

For sinking particles, the protist communities cluster mainly by site, with little difference between the base of the euphotic zone and 150 m depth (Fig. 5). OO samples form a distinct cluster with one of the shallow sediment traps from the TZ site. This cluster is more compositionally similar to water-column samples than to other trap samples, in part because apicomplexans comprise only a small component of the Alveolata in these samples (Fig. 5b). Trap sample sequences at the IN site (and one shallow TZ trap sample) cluster close to samples from the overlying mixed layer and the base of the euphotic zone at IN, all with high relative abundances of *Thalassiosira* (15–29% of reads). In contrast, CC and deep TZ trap samples with high contributions from apicomplexans (5–91% of reads) form distinct groups with low similarity to overlying water-column samples (Fig. 5b).

3.5. Major groups from sequence analyses: Prokaryotes

Bacteria sequences dominated water-column (92.6%) and trap (99.9%) 16S samples over Archaea (Fig. 3). In the mixed layer, Bacteroidetes and Proteobacteria dominate prokaryote sequences at the inshore sites (TZ and IN), and Proteobacteria and Cyanobacteria dominate at the offshore sites (OO and CC) (Fig. 6). Archaea, Bacteroidetes, and Proteobacteria are the main contributors at all sampling locations in the base of the euphotic zone and 150-m samples (Fig. 6). In contrast, sediment-trap sequences were mostly dominated by Proteobacteria at all sampling locations (Fig. 6), although relative abundances of Bacteroidetes are high in the IN, CC, and OO samples from the brine-filled traps, in conjunction with Cyanobacteria at the offshore sites (Fig. 6).

3.6. Water-column community composition: Prokaryotes

Similar to protists, the prokaryote assemblages show clear differences with depth (Fig. 7, Table S7). The mixed-layer communities have high relative abundances of cyanobacteria (*Prochlorococcus* and *Synechococcus*) and Alphaproteobacteria (SAR11 surface clade), with these three groups comprising 38% and 52% of the 16S rRNA gene sequences, respectively, at the offshore sampling locations (OO and CC) (Fig. 7b). *Synechococcus* was the most represented group, comprising 16% of reads, at the TZ site, and *Roseobacter* was conspicuously abundant at the IN site. Flavobacteria (multiple groups) were also proportionally more important at the TZ and IN sites (Fig. 7b).

At the base of the euphotic zone, Flavobacteria were fairly consistent contributors to prokaryotic assemblages. The SAR11 surface clade was also important in CC and TZ waters. Archaea (*Thaumarchaeota*) increased substantially with depth (Fig. 7b). Bacteroidetes and Alphaproteobacteria were the main contributors in samples from the base of the euphotic zone and 150 m at all locations (Fig. 7b). SAR11 surface clade was also important at 150 m at all sites (Fig. 7b).

Bray-Curtis dissimilarity analysis shows a clear distinction between water-column (which cluster in one large group with ~50% similarity) and sediment trap samples (Fig. 7b). Within the water-column super group, 150-m samples comprise a distinct group with >80% similarity, identified by high proportional abundances of *Thaumarcheota* and SAR11. This group is most closely associated with TZ, CC, and OO.
samples from the base of euphotic zone, which have lower proportions of Thaumarchaeota and SAR11 and a greater influence of Flavobacteria. Mixed-layer samples fall into two subgroups based on relative system productivity, with TZ and IN samples separate from OO and CC samples (Fig. 7b).

3.7 Sediment trap community composition: Prokaryotes

Prokaryotic communities were distinctly different in trap samples compared to the water column (Fig. 7, Table S7). Gammaproteobacteria (Pseudoalteromonas and Vibrio) comprised the majority of reads at all locations (Figs. 6 and 7, Table S7), whereas variability in other groups drive cross-site differences. Unlike water-column or eukaryotic samples; however, there is not a clear pattern. Relative abundances of Bacteroidetes, Alphaproteobacteria, and Cyanobacteria were high in OO and CC samples from the brine-filled traps (Fig. 7b). Prochlorococcus and Gammaproteobacteria clade BD1-7 were major components of two OO samples, while the Flavobacteria Fluviicola was important in one IN sample (Fig. 7b). These relatively minor differences determined the clustering of samples in the dissimilarity analysis. However, the dissimilarity analysis did not elucidate clear patterns in the data, with most groups including samples from multiple sites and multiple depths. Across all sites, the general similarity and over-representation of Gammaproteobacteria on sinking particles is clear, in striking contrast to water-column assemblages and the variability of protistan communities on sinking particles.

3.8 Plastid sequences in sinking particles

Relative abundances of 16S and 18S rRNA gene sequences do not allow direct comparison of relative abundances of prokaryotes and eukaryotes. Hence, we analyzed plastid sequences to compare cyanobacteria and eukaryotic phytoplankton. Eighteen eukaryotic phytoplankton taxa were obtained from 16S rRNA gene sequencing of the water column and trap samples, with Mamiellophyceae, Prymnesiophyceae, and Bacillariophyceae exhibiting the greatest relative abundances in mixed-layer samples at the IN site (Fig. 8, Table S8). At this site, Bacillariophyceae was the only group that was over-represented in sediment traps (35% of plastid sequences) relative to the mixed layer (12.0%). Also, at the IN site, the relative abundances of Mamiellophyceae and Prymnesiophyceae were similar in traps and the water column, while cyanobacteria (Synechococcus and Prochlorococcus) were under-
represented in sinking particles relative to the water column. At the TZ site, mixed-layer plastid sequences were dominated by Mamielophyceae (64.5%), which had lower proportional representation in shallow sediment traps, but roughly similar proportional representation in deep sediment traps (relative to the mixed layer). Meanwhile, Synechococcus (30.0% of plastid sequences at the TZ site) had a similar contribution to water column and the sediment traps. At the two offshore sites (CC and OO), the two cyanobacterial groups dominated the water-column plastid samples compared to eukaryotic phytoplankton, and Synechococcus was over-represented on sinking particles in the CC (but not OO) relative to the mixed layer, while Prochlorococcus was never over-represented on sinking particles. Across all sites, Bacillariophyceae proportions in the sediment trap samples were typically similar to or greater than its contribution to the mixed layer water column community (Fig. 8, Table S8).

4. Discussion

4.1. Environmental conditions during the 2015–2016 El Niño

The prolonged anomalous warm-water conditions that prevailed in the North Pacific from late 2013 to 2016 likely influenced the microbes contributing to sinking particles during our study. In the northern region of the California Current Ecosystem (CCE), considerable changes in the plankton communities (protists and zooplankton) were documented by Peterson et al. (2017), who found an increase in taxa with tropical-subtropical affinities, increased species richness of dinoflagellates, and reduced plankton biomass. During our study in the southern CCE, plankton with subtropical affinities were also evident (e.g., Doliolum denticulatum). However, the inshore-offshore distributions of the major groups of microbes were generally consistent with their normal ranges in the region (Freibott et al., unpublished), despite warmer than normal conditions ( McClatchie et al., 2016 ). For example, the higher relative abundances of Prochlorococcus at the offshore sites (OO and CC) and of Synechococcus at the CC and TZ sites are consistent with their inshore-offshore biomass distributions (Taylor and Landry, 2018). Likewise, among protists primarily present in the mixed layer (Fig. 5), the dominance of dinoflagellates offshore and the importance of both diatoms and dinoflagellates inshore (assuming that syndiniales are mostly within their hosts) agree with the inshore-offshore contributions of these groups to protistan abundance and biomass in the CCE (Taylor et al., 2015; Venrick, 2002). The high dominance of rhizarians deep in the water column is consistent with their increased importance to plankton biomass with depth (Biard et al., 2018). In addition, particulate organic carbon fluxes during this study at the OO, CC, and TZ sites were within the ranges previously documented in the CCE, whereas at the IN site, fluxes were actually higher than prior measurements (Stukel et al., 2015). Although the effects of the 2015–2016 El Niño on the specific microbes exported are uncertain due to the lack of a basis of comparison, they are likely representative of conditions in the region because our sampling was done when El Niño was in decline.

4.2. Microbial communities and sinking particle flux: Ecosystem variability

Foundational studies relating food webs and the biological pump have highlighted the importance of large, mineral-ballasted phytoplankton, metazoan fecal pellets, and aggregates in sinking particle flux (Michaels and Silver, 1988; Passow et al., 1994; Smayda, 1971). Additional studies have documented heterogeneity of sinking particles and aggregates, including variability in the taxonomic composition and size of sinking particles (Durkin et al., 2015; Fender et al., 2019; McDonnell and Buesseler, 2010; Richardson, 2019), though without substantially changing the expectation for higher export efficiency in diatom- and mesozooplankton-dominated regions (Steinberg and Landry, 2017; Turner, 2015). Until recently, however, such studies have been severely limited by the poor taxonomic resolution of microscopy and pigment-based approaches to investigating microbial communities.

Next generation sequencing techniques now allow more detailed interrogation of microbial communities on sinking particles and in the mesopelagic realm generally (Amacher et al., 2009; Swan et al., 2011). These sequencing techniques have identified unexpected taxa that either correlate with export flux or are commonly associated with sinking material, including rhizarians, parasitic Syndiniales, and Flavobacteria (Boeuf et al., 2019; Fontanez et al., 2015; Guildi et al., 2016; Gutierrez-Rodriguez et al., 2019). Such studies have additionally raised new questions. For instance, Boeuf et al. (2019) found specialized piezophilic communities living on sinking particles in the bathypelagic, suggesting community evolution on sinking particles. In contrast, Mestre et al. (2018) observed similarities in microbial communities associated with large particles collected in surface, mesopelagic, and bathypelagic waters suggesting strong vertical connectivity. Sequencing has also given differing results with respect to the relative importance of cyano bacteria and eukaryotic phytoplankton (especially diatoms) on sinking material. Amacher et al. (2013) found that Synechococcus was often over-represented on sinking material in the Sargasso Sea relative to other cyano bacteria and eukaryotic plastids, and only rarely found genetic evidence of diatom plastids. In contrast, Preston et al. (2019)
analyzed sequences from a deep-sea time-series sediment trap in the northeast Pacific and found that a single diatom species (*Thalassiosira aestivalis*) accounted for 64% of all 18S rRNA gene sequences during the only high flux event in their dataset.

By sampling across an environmental gradient while simultaneously assessing suspended and sinking prokaryotic and eukaryotic communities, we are able to elucidate broad patterns in sinking microbial communities and reconcile some of these disparate results. We hypothesized that variability in the relative contributions of eukaryotic phytoplankton and microbial communities assessed in previous studies would be driven by variability in trophic structure. Specifically, we hypothesized that in productive ecosystems diatoms and rapidly sinking fecal pellets would promote vertical connectivity and hence greater similarity between euphotic zone microbial communities and microbial communities on sinking particles below the euphotic zone. In contrast, slower sinking rates and high recycling in oligotrophic regions would promote community succession on sinking particles, leading to communities on sinking particles that are largely distinct from the mixed layer. These relationships and mechanisms are generally supported by our results.

At the high-productivity, upwelling-influenced IN site, surface Chl a was 40 times higher than at the oligotrophic (OO) site and vertically-integrated primary production was nearly an order of magnitude higher (Figs. 1 and 2). This high productivity was associated with substantially higher export and a diatom-dominated phytoplankton community that was over-represented in sinking particles (Fig. 4). The IN site was also the only site for which Bray-Curtis dissimilarity analysis of protistan communities showed sediment trap and mixed layer community clustering together (Fig. 5). For the remaining three sites, water-column communities clustered by depth, and sediment traps samples clustered separately. Particle-associated microbes at the richer upwelling IN site had high relative abundances of dinoflagellates (syndiniales MALV I, MALV III, and *Karolodinium*) and, notably, diatoms (an unclassified Bacillariophyceae and *Thalassiosira*). Microscopical analysis indicate that *Thalassiosira* cells are numerically important in mesotrophic CCE waters (Du et al., 2015), and that diatoms in general often dominate biomass in the coastal upwelling region (Taylor and Landry, 2018; Taylor et al., 2015). Although two diatom taxa were important in the water-column community at the IN site (Fig. 5), only *Thalassiosira* was strongly represented in the sediment trap material, likely transported to depth within copepod fecal pellets. Notably, this diatom species, responsible for 24% of protistan sequences in the deepest trap at the IN site, was closely related to the diatom that contributed 64% of all 18S rRNA gene sequences during the only high flux event sampled by Preston et al. (2019) in the bathypelagic. Overall, the strong similarity between euphotic zone and trap particle communities at the mesotrophic IN site suggests efficient export of “fresh” carbon from the ocean’s surface, consistent with higher export expected for diatom-rich waters.

In stark contrast to our results at the IN site, the CC and OO sites showed strong dissimilarity between sediment trap and water-column communities (Fig. 5). Water-column communities clustered by depth, while the trap communities formed separate and distinct clusters. The protists on sinking particles at the OO site were characterized by high relative abundances of alveolates (mostly dinoflagellates) and rhi- zarians, consistent with their recognized importance on sinking particles (Amacher et al., 2009, 2013; Fontanet et al., 2015) and carbon export in oligotrophic systems (Guidi et al., 2016; Gutierrez-Rodriguez et al., 2019; Lampitt et al., 2009). Likewise, these groups are major components of protistan biomass in oligotrophic waters of the CCE (Biard et al., 2018; Taylor et al., 2015). Notably, the sinking material analyzed by microscopy at the OO site was mostly comprised of unrecognizable particles and aggregates, with very low pigment content (Fig. 2c–g). There were very few recognizable fecal pellets, although metazoaan were actually a higher proportion of total reads than at other sites. Taken together, this suggests that sinking material in offshore oligotrophic waters is likely derived from fecal material, but aged and degraded during sinking, allowing substantial microbial colonization and community evolution. We also note that, although the coastal area was dominated by epipelagic taxa (e.g., diatoms) that contributed to export, export efficiency (i.e., carbon export/NPP) was not elevated in this region relative to offshore areas. This discrepancy likely arises from spatiotemporal decoupling of production and export, which has been described for the CCE region (Plattner et al., 2005; Kelly et al., 2018; Kahru et al., 2020), and leads to a supplement of organic matter input to offshore regions (Stephens et al. 2018). This organic matter subsidy increases export efficiency in the offshore regions relative to what would be expected in a one-dimensional system with oligotrophic communities. Disentangling these processes to quantify the relationship between epipelagic communities and export efficiency will require further sampling in coordination with mechanistic models.

The protists on sinking particles in the CC and TZ were dominated by the apicomplexan Cephalodinophoroidea and/or the acantharian Chaunacanthida. Acantharians are known for reproductive strategies that involve forming cysts, which sink rapidly due to their dense mineral (strontium sulfate) composition (Bernstein et al., 1987; Martin et al., 2010; Decelle et al., 2013). Because Chaunacanthida were important in the deeper water column at 150 m except at the CC, the high relative abundances of Chaunacanthida sequences in traps at this site could also reflect cyst sinking. We observed round structures similar to acantharian cysts in trap samples in the CC; however, these were considerably smaller (22–28 μm) than cysts collected in traps in the North Pacific (80–165 μm, Bernstein et al., 1987) and North Atlantic (200–1000 μm, Martin et al., 2010; Decelle et al., 2013). Cephalodinophoroidea is a poorly known group of gregarine apicomplexans that parasitize the intestines of crustaceans, including copepods (Rueckert et al., 2011). The high relative abundances of apicomplexans in CC and TZ trap-collected particles coincide with the bloom of *D. denticulatum*. Due to the preferred particle sizes filtered by these dioloids (2.5–15 μm, Katechakis et al., 2004), it is unlikely that the parasites were acquired by feeding directly on copepods. Considering that other gregarine apicomplexans have been identified in guts of salps (Claxton, 2002; Wallis et al., 2017), Cephalodinophoroidea possibly also parasitize dioloids. Because dioloids are passive filter-feeders, the proportions of appropriately sized microbes in their diet should be similar to those in the water column (Allredge and Madin, 1982). Nonetheless, at the sites where dioloids were most abundant, microbial communities in the water column and on sinking particles showed little similarity, suggesting substantial post-consumption transformations before particles were exported. The most notable contrast in the TZ was the virtual absence of *Ostreococcus*, a dominant mixed-layer picophytoplankton, on the sinking particles. *Ostreococcus* and *Micromonas* are both important components of the water-column communities in nutrient-rich regions of the CCE (Allen et al., 2012). It is possible, however, that the very tiny cells of *Ostreococcus* (~1 μm) were not efficiently grazed by *D. denticulatum* or rapidly degraded in the doliolid guts. In contrast to *Ostreococcus*, the relative abundances of two dinoflagellates (an uncultured taxon and *Karlodinium*) with lower contributions to the water-column assemblages were higher on sinking particles. These dinoflagellates were likely grazed and transported in doliolid fecal pellets.

4.3. Prokaryotic and eukaryotic communities associated with sinking particles

The above discussion highlights the impact of ecosystem trophic structure and food webs on the connectivity between water column and sinking microbial communities. However, many studies have found distinct microbial taxa, such as *Vibrio* and *Acrbacter*, associated with sinking particles (Boeuf et al., 2019; Guidi et al., 2016; Pelve et al., 2017). We hypothesize that eukaryotic organisms associated with sinking particles show greater compositional similarity to water-column communities than prokaryotic taxa, because they are more likely to sink individually, or within aggregates or fecal pellets. This hypothesis is
supported by our Bray-Curtis dissimilarity results showing that some protistan water-column communities cluster with trap-collected communities and vice versa (Fig. 5), while all prokaryotic water-column communities clustered within a single large group (Fig. 7).

These results agree with previous metabarcoding analyses from oligotrophic and mesotrophic systems that have shown significant differences in the community structure of water-column and trap-collected prokaryotes (Amacher et al., 2013; Fontanez et al., 2015; LeCleir et al., 2014). Among heterotrophic bacteria, mixed-layer sequences were mainly represented by members of Flavobacteria (NS4 and NS5 clades, *Flavicola*) and Alphaproteobacteria (SAR11, *Roseobacter*), some of which are important free-living (e.g., SAR11) or particle-associated microbes in pelagic ecosystems (e.g., *Flavicola, Roseobacter*) (DeLong et al., 1995). SAR11 relative abundances tended to be higher offshore (OO and CC) and were rarely major components of sinking particles. Allen et al. (2012) also documented that SAR11 sequences were rarely found in the largest size-fractions analyzed (3.0 μm). Likewise, the higher relative abundances of *Roseobacter* in nutrient-rich waters is consistent with Allen et al.’s (2012) findings in the CCE. Although *Roseobacter* are recognized as important colonizers of sinking particles (LeCleir et al., 2014), their relative abundances at the mesotrophic inshore sites (TZ and IN) during our study were similar in the water column and sinking particles. Also at the IN site, *Flavicola* was the only heterotrophic bacterium significantly enriched on sinking particles, likely reflecting growth of these particle-associated bacteria during sinking (Suzuki et al., 2017). The increased importance of Archaea with depth at all sites is consistent with their vertical distribution (Karner et al., 2001) and their general absence on sinking particles collected in traps (Fontanez et al., 2015). In contrast, the dominance of *Pseudodieteromonas* and *Vibrio* (Gammaproteobacteria) sequences in trap-collected samples is consistent with their association with sinking particles or eukaryote surfaces and their enzymatic capabilities for degrading high molecular weight compounds (DeLong et al., 1993; Fontanez et al., 2015). *Pseudodieteromonas* are also known to produce anti-bacterial compounds (Holmström and Kjelleberg, 1999), which may allow them to dominate on particles by inhibiting the growth of other bacteria. In contrast, we suspect that Synechococcus, which was enriched on sinking particles only in oligotrophic regions, is often found in sediment trap results not because its ecology enhances carbon export, but rather because digestion-resistance properties enable it to survive intact (and be recognized) on sinking particles (Johnson et al., 1982; Valencia et al., 2021).

Although protistan groups in sediment traps generally showed greater similarity to mixed-layer communities, syndiniales (MALV), Cephalodiformida and, Rhizaria were specific taxa associated with sinking particles. The importance of syndiniales and Cephalodiformida on sinking particles suggests that these parasites might be transported within their eukaryotic hosts (Guilhou et al., 2008). Although the mechanisms are not well understood, syndiniales, among parasites generally, appear to play important roles in modulating export flux (e.g., Amacher et al., 2013; Guidi et al., 2016). Rhizaria, meanwhile, are surmised to be important contributors to sinking particles both because they often reside in the mesopelagic, and because their siliceous tests promote sinking (Biard et al., 2016, 2018; Gutiérrez-Rodríguez et al., 2019). In Valencia et al. (2021), we consider specific properties of these prokaryotic and protistan communities that make them more likely to be detected on sinking particles. Here, however, we simply note that the greater influence of mixed-layer communities on protistan, rather than prokaryotic, sinking particle-associated communities helps explain the greater temporal variability in protistan communities collected in time-series sediment traps (Boeuf et al., 2019; Preston et al., 2019).

5. Conclusions

Metabarcoding analyses of water-column and sediment trap samples from the CCE confirm some general findings from other systems, such as the strong associations of Gammaproteobacteria and rhizarians with particle export. Additionally, they provide new perspectives on variability within taxa and across gradients of ecosystem trophic state. For the former, we document differential sequence retention of *Thalassiosira* relative to co-occurring diatoms in sinking particles. We also find that protistan communities associated with sinking particles more closely resemble mixed-layer communities than prokaryotic communities on the same particles. Comparison of sites across an environmental gradient show that sinking particles beneath high-biomass, coastal upwelling-influenced water more closely resembles the water-column dominants (especially diatoms) found above, while oligotrophic communities exhibit greater dissimilarity between the microbial communities on sinking particles and in the mixed layer. These differences highlight important variability in microbial contributions to export from the water column between inshore mesotrophic and offshore oligotrophic waters that seem to align with differences in dominant grazing pathways (large copepods versus doliolids versus protistan microzooplankton) and export delivery mechanisms (fecal pellets versus aged amorphous aggregates).

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

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