- Slow particle remineralization, rather than
- suppressed disaggregation, drives efficient
- 3 flux transfer through the Eastern Tropical
- 4 North Pacific Oxygen Deficient Zone
- 5 30 October 2021
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17 1 Key Points

- 18 The upper mesopelagic of the oligotrophic Eastern Tropical North Pacific Oxygen Deficient Zone (ODZ)
- 19 has low flux attenuation.
- 20 Comparison of particle size observations to models suggests that the breakdown of particles of all sizes is
- 21 slow throughout the ODZ.
- 22 Zooplankton appear to transport organic matter into, and disaggregate particles within, the ODZ above
- 23 500 m.

24 **2** Abstract

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Models and observations suggest that particle flux attenuation is lower across the mesopelagic zone of anoxic environments compared to oxic environments. Flux attenuation is controlled by microbial metabolism as well as aggregation and disaggregation by zooplankton, all of which also shape the relative abundance of differently sized particles. Observing and modeling particle spectra can provide information about the contributions of these processes. We measured particle size spectrum profiles at one station in the oligotrophic Eastern Tropical North Pacific Oxygen Deficient Zone (ETNP ODZ) using an underwater vision profiler (UVP), a high-resolution camera that counts and sizes particles. Measurements were taken at different times of day, over the course of a week. Comparing these data to particle flux measurements from sediment traps collected over the same time-period allowed us to constrain the particle size to flux relationship, and to generate highly resolved depth and time estimates of particle flux rates. We found that particle flux attenuated very little throughout the anoxic water column, and at some time-points appeared to increase. Comparing our observations to model predictions suggested that particles of all sizes remineralize more slowly in the ODZ than in oxic waters, and that large particles disaggregate into smaller particles, primarily between the base of the photic zone and 500 m. Acoustic measurements of multiple size classes of organisms suggested that many organisms migrated, during the day, to the region with high particle disaggregation. Our data suggest that diel-migrating organisms both actively transport biomass and disaggregate particles in the ODZ core.

3 Plain Language Summary

- 43 Marine snow are microscopic particles that form in the surface of the ocean and sink into the deep ocean.
- 44 Most of these particles are the remains of dead algae and feces of tiny animals (zooplankton). The deeper
- 45 the particles sink into the ocean before microbes or animals consume them, the longer it takes before the
- 46 carbon in those particles can return to the atmosphere. In parts of the ocean where there is no oxygen,
- 47 more particles sink to greater depths, for reasons that are not well-understood. We used an underwater
- 48 camera to observe marine snow particles in the ocean just west of Mexico where there is little to no
- 49 oxygen at depth. We compared the observations to predictions from several computer simulations
- representing hypothesized mechanisms to explain why particles are consumed less in water without
- oxygen. Our measurements suggest that one reason that particles sink to deeper depths here is because
- 52 microbes consume the particles slowly when there is no oxygen. Meanwhile, zooplankton still break large
- 53 particles into smaller ones and produce fecal pellets in these low oxygen waters.

4 Introduction

- The biological pump, in which sinking microaggregate (< 500 μm) and marine snow (> 500 μm) particles
 (Simon et al., 2002) transport carbon from the surface into the deep ocean, is a key part of the global
 carbon cycle (Neuer et al., 2014; Turner, 2015). Organic matter flux into the deep ocean (>1000 m) is a
- function both of export from the photic zone into the mesopelagic (export flux), and the fraction of that
- flux that crosses through the mesopelagic (transfer efficiency) (François et al., 2002; Passow & Carlson,
- 60 2012; Siegel et al., 2016). While definitions vary between studies, we define "mesopelagic" as the region
- between the base of the photic zone, and 1000 m (following Francois et al., 2002; Cram et al., 2018). The
- transfer efficiency of the biological pump may affect global atmospheric carbon levels (Kwon & Primeau,
- 63 2008). Thus, understanding the processes that shape organic matter degradation in the mesopelagic is
- 64 critical.

- Oxygen concentrations, and especially the geographic and vertical extent of anoxic ocean regions, appear
- to modulate particle flux through the mesopelagic. Observations of particle flux in the Eastern Tropical
- North Pacific near the Mexican coast (Hartnett & Devol, 2003; Van Mooy et al., 2002; Weber & Bianchi,
- 68 2020), the Eastern Tropical South Pacific (Pavia et al., 2019), and Arabian Sea (Keil et al., 2016; Roullier
- 69 et al., 2014) have suggested lower flux attenuation in these ODZ systems. Models have shown that
- accounting for oxygen limitation in ODZs is necessary to fit global patterns of particle transfer (Cram et
- al., 2018; DeVries & Weber, 2017). Analysis of remineralization tracers has also shown evidence of slow
- 72 flux attention in the ODZs (Weber & Bianchi, 2020). Understanding the driving mechanisms of these
- patterns is important because the oxygen content of the ocean is decreasing (Ito et al., 2017; Schmidtko et
- 74 al., 2017), and the spatial extent and depth range of ODZs, including the Eastern Tropical North Pacific
- 75 (ETNP) Oxygen Deficient Zone (ODZ), are likely to change, though there is disagreement over whether
- they are expanding or undergoing natural fluctuation (Deutsch et al., 2014; Horak et al., 2016; Stramma et
- al., 2008). Recent data informed models suggest that ODZs may enhance carbon transport to the deep
- ocean, by inhibiting microbial degradation of sinking marine particles (Cram et al., 2018). However,
- 79 biological organic matter transport is also modulated by zooplankton whose interactions with particle flux
- 80 in pelagic ODZs are only beginning to be quantitatively explored (Kiko et al., 2020).
- 81 Zooplankton modulate carbon flux through the mesopelagic (Jackson & Burd, 2001; Steinberg & Landry,
- 82 2017; Turner, 2015), and by extension the transfer efficiency of the biological pump (Archibald et al.,
- 83 2019; Cavan et al., 2017), in three key ways that could be affected by ocean oxygen concentrations:
- 84 (1) Active transport: Zooplankton migrate between the surface and mesopelagic, consuming plankton and
- 85 particles in the surface and producing particulate organic carbon (POC), dissolved organic carbon (DOC),

- 86 respiratory CO₂, and zooplankton carcasses at depth (Archibald et al., 2019; Bianchi et al., 2013;
- Hannides et al., 2009; Steinberg et al., 2000; Stukel et al., 2018, 2019). This manuscript focuses on
- particles, so we only consider POC and carcass production, which cause particles to "appear" in the
- 89 midwater.
- 90 (2) Repackaging: Zooplankton fecal pellets have different physical properties than the parrticles and
- plankton that they ingest (Wilson et al., 2008). In this paper we define repackaging as zooplankton
- 92 feeding in the mesopelagic and producing fecal pellets, effectively aggregating POM.
- 93 (3) Disaggregation: Zooplankton break large particles into smaller ones in two ways by Coprorhexy
- 94 (also sometimes called sloppy feeding) in which they break particles apart while feeding on them
- 95 (Lampitt et al., 1990; Noji et al., 1991; Poulsen & Kiørboe, 2005), and by generating turbulence while
- 96 they swim (Dilling & Alldredge, 2000; Goldthwait et al., 2005). Disaggregation can reduce particle
- 97 transfer efficiency, because smaller particles sink more slowly and so reside longer in mesopelagic,
- allowing them to be consumed before reaching deep waters (Goldthwait et al., 2005; Lampitt et al., 1990;
- 99 Noji et al., 1991; Poulsen & Kiørboe, 2005). In some cases, disaggregation can explain around 50% of the
- particle flux attenuation over depth (Briggs et al., 2020).
- The migratory zooplankton that drive these mesopelagic processes spend the night in the surface layer
- and migrate into the core of the OMZ during the day (Bianchi et al., 2014). These organisms likely survive
- in ODZs by slowing their metabolic processes, but may supplement these with very efficient oxygen
- uptake and anaerobic metabolism (Seibel, 2011). Acoustic data suggest that zooplankton do not migrate
- as deeply into ODZs as they do into regions where ODZs are absent (Bianchi et al. 2011). New evidence
- suggests that in ODZ regions with shallower oxyclines, night-time migration depth remains the same but
- the depth where the organisms spend the day is compressed (Wishner et al., 2020). The rates at which
- zooplankton transport, repackage and disaggregate particles in ODZs are difficult to measure and
- therefore poorly constrained. Despite the importance of zooplankton mediated processes to global carbon
- flux, zooplankton are often missing from models of particle transfer.
- 111 Current models of particle transfer through the mesopelagic ocean predict that particle size, ocean
- temperature, and oxygen concentrations are the dominant factors controlling particle flux attenuation
- 113 (Cram et al., 2018; DeVries & Weber, 2017). These models, however, do not account for active transport
- or disaggregation by zooplankton. As a result of this assumption, the models predict that small particles
- preferentially attenuate with depth, which is often not borne out by observations (Durkin et al., 2015).
- Therefore, these models' predictions provide a useful null hypothesis of expected particle size
- distributions in the absence of zooplankton effects, which can be compared to observed distributions of
- particles to explore the magnitude of zooplankton effects.

119 Underwater vision profilers are cameras that can count and size many particles over large water volumes 120 (Picheral et al., 2010) and provide valuable information about particle distributions and transport. When 121 deployed in concert with particle traps in some regions, they can be used to predict flux in other regions 122 where traps have not been deployed (Guidi et al., 2008; Kiko et al., 2020). Connecting UVP and trap data 123 can furthermore inform about total particle flux variability across space and time, relationships between 124 particle size, biomass, composition, and sinking speed, as well as the contributions of the different 125 particle sizes to flux (Guidi et al., 2008; Kiko et al., 2017). Combined particle trap flux and UVP data 126 from the North Atlantic suggest active transport by zooplankton into hypoxic water (Kiko et al., 2020), 127 but the authors suggest that in more anoxic and larger ODZs, such as the modern day ETNP, there might 128 be reduced active transport into the mesopelagic, since many migratory organisms would presumably not 129 migrate into the anoxic water and would be less active. In this manuscript we provide the first combined 130 flux measurement and UVP data from such a fully anoxic region, the ETNP ODZ. 131 In addition to being fully anoxic, the ETNP ODZ is primarily oligotrophic: most of the volume of the 132 ETNP ODZ is below regions of very low surface productivity (Fuchsman et al., 2019; Pennington et al., 133 2006). Meanwhile most flux data have been measured in more coastal, higher productivity regions of the 134 ETNP (Hartnett & Devol, 2003; Van Mooy et al., 2002). 135 A recent modeling study posed three hypotheses to explain why particle flux attenuates slowly in ODZs 136 (Weber & Bianchi, 2020), which are susceptible to testing with UVP data. These are: H1: All particles in 137 ODZs remineralize more slowly than in oxic water, regardless of their size, due to slower carbon 138 oxidation during denitrification than aerobic respiration. H2: Breakdown of large particles into small 139 particles is suppressed in the ODZ because there is less disaggregation by zooplankton than elsewhere. 140 H3: Large particles remineralize more slowly in ODZs, but smaller ones do not, because carbon oxidation 141 in large particles can become limited by the diffusive supply of oxygen and nitrate. In this case, 142 respiration can only proceed by thermodynamically inefficient sulfate reduction (Bianchi et al., 2018; 143 Lam & Kuypers, 2011). Sulfide and organic matter sulfurization have been found on particles at this site 144 at nanomolar concentrations (Raven et al., 2021). Microbial analysis of particles found sulfate reducers 145 and S-oxidizing denitrifiers at low abundances (Fuchsman et al., 2017; Saunders et al., 2019). Each of the 146 hypotheses outlined above were predicted to leave distinct signatures in particle size distributions in the 147 core of ODZ regions (Weber & Bianchi, 2020). The model with slow remineralization of all particles, 148 predicts an increase in the abundance of small particles in the ODZ core relative both to overlying waters 149 and to similar, oxygenated environments (H1). The model with suppressed disaggregation predicts a large 150 decrease in small particle biomass in the ODZ, both relative to the surface and to oxygenated water (H2). 151 The model in which remineralization is depressed only in large particles predicts a small decrease with

- depth in small particle abundance, similar to that seen in oxygenated environments (H3). However, the
- necessary particle size data from an ODZ was not previously available to support any hypothesis at the
- exclusion of the others. In this manuscript we present a new dataset that is sufficient to test these three
- 155 hypotheses.
- To provide the data to test hypotheses **H1-H3** and illuminate zooplankton particle interactions in
- oligotrophic ODZs, we collected particle size data at high temporal resolution over the course of a week
- in an anoxic site typical of the oligotrophic ETNP ODZ, well away from the high productivity zone in the
- 159 coast. We integrated this size data with observed flux measurements, and acoustic data. We quantified,
- throughout the water column, how changes in size distribution deviate from changes that would be
- predicted by remineralization and sinking only models.
- We ask the following four questions:
- 163 **Question A:** How does the particle size distribution at one location in the oligotrophic Eastern Tropical
- North Pacific vary with respect to depth and time?
- Question B: Do our data support any of the three Weber and Bianchi (2020) models (H1-H3)?
- 166 Question C: Do our data suggest that regions of the oxygen deficient zone harbor disaggregation-like
- processes, and if so, do these co-occur with migratory zooplankton?
- 168 **Question D**: How do particle size distribution spectra in the ODZ compare to those seen in the oxic
- 169 ocean?

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- By addressing these four questions, we demonstrate that our dataset from the ETNP supports Weber and
- Bianchi's first hypothesis, that microbial remineralization of all particles slows in the ODZ, while
- disaggregation continues unabated. Additionally, disaggregation-like processes do appear to co-occur
- with acoustic measurements of migratory zooplankton, suggesting that exclusion of zooplankton is not a
- major contributor to slow flux attenuation.

5 Methods

- Unless specified otherwise, measurements were taken on board the R/V Sikuliaq, cruise number
- 177 SKQ201617S, from 07 January 2017 through 13 January 2017 at a single station 16.5°N 106.9°W, which
- 178 was located in an oligotrophic region of the Eastern Tropical North Pacific Oxygen Deficient Zone
- 179 (ETNP Station P2; Figure 1A). Data are compared against measurements taken at 16.5°N 152.0°W on 08
- 180 May 2015, collected on the GO-SHIP CLIVAR/CARBON P16N Leg 1 Cruise (CCHDO Hydrographic
- 181 Cruise 33RO20150410). This station was at the same latitude as ETNP Station P2, west of the ODZ, but
- was not anoxic (P16 Transect Station 100; Figure S1).

5.1 Water property measurements

- We measured water properties of temperature, salinity, fluorescence, oxygen concentration and turbidity
- using the shipboard SeaBird 911 CTD. Auxiliary sensors included a WetLabs C-Star (beam attenuation
- and transmission) and a Seapoint fluorometer. Data were processed with Seabird Software, (programs-
- data conversion, align, thermal mass, derive, bin average and bottle summary) using factory supplied
- calibrations. Data was analyzed and visualized in R (Team 2011). Processed data are available under
- NCEI Accession number 1064968 (Rocap et al., 2017).

5.2 Water mass analysis

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- 191 Evans et al. (2020) previously employed optimum multiparameter analysis to map the percent identity of
- the water observed at each depth to three water masses: the 13 Degree Celsius Water (13CW), North
- 193 Equatorial Pacific Intermediate Water (NEPIW), and Antarctic Intermediate Water (AAIW). We subset
- and examine only the portion of these data that correspond to our site.

195 **5.3 Acoustic Measurements**

- Backscattering signals from the ship-board EK-60 were collected and archived by UNOLS as raw data
- files. We used Echopype software (Lee et al., 2021) to convert these raw files to netcdf files, which were
- down-sampled to five minute time-step resolution, saved as a text file, and later visualized in R. The
- acoustic data appeared to be off by one hour and so one hour was subtracted from all time measurements.
- 200 This correction resulted in zooplankton vertical migrations being synchronized with the diel light cycle as
- was recorded on board the ship by JAC.

5.4 Particle size measurements

- 203 Particle size data were collected by an Underwater Vision Profiler 5 (UVP) that was mounted below the
- 204 CTD-rosette and deployed for all CTD casts shallower than 2500 m. A UVP is a combination camera and
- 205 light source that quantifies the abundance and size of particles from 100 µm to several centimeters in size
- 206 (Picheral et al., 2010). UVP data were processed using the Zooprocess software, which prepares the data
- for upload to the Ecotaxa database (Picheral et al., 2017); data from all UVP instruments are located on
- this online database for ease of access. Detailed descriptions for installation of the Zooprocess software
- 209 can be found on the PIQv website (https://sites.google.com/view/piqv/zooprocess-uvpapp). Zooprocess
- uses the first and last image number selected by the user in metadata to isolate the downcast and process
- this subset for both particle size distribution and image data. The processed files and metadata are then
- 212 uploaded to a shared FTP database where it is available for upload to Ecotaxa. This project required the
- 213 extra step of filtering out images due to the discovery of an issue with the lighting system, where only one
- of the two LEDs would illuminate, resulting in an incomplete sample. The filtering procedure is

215 documented in a link available at the same location as the Zooprocess download. Images where only a 216 single light illuminated were removed from the dataset before it was uploaded on to Ecotaxa. Once 217 uploaded to Ecotaxa, data were downloaded from EcoPart (the particle section of the database) in detailed 218 TSV format, and analyzed in R. The UVP provided estimates of abundances of particles in different size-219 bins, as well as information about the volumes over which those particle numbers had been calculated. 220 The instrument is capable of observing smaller particles (down to 60 µm), but these tend to be 221 underestimated and so are excluded from the analysis. The instrument can in principle also measure larger 222 particles (up to the field of view of the camera), though these tend to be scarce enough to be not detected. 223 In this paper, we do not have an upper size cut-off for our analysis and rather implement statistics that are 224 robust to non-detection of scarce large particles (section 5.5.1). Visual inspection of images larger than 225 1 mm suggests that these large particles are primarily "marine snow" but about 5% are zooplankton. We 226 did not quantify the size distribution of these images.

5.5 Flux measurements

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Arrays, consisting of a surface float and two traps, were deployed and allowed to float freely during which time they collected particles. Trap deployments began on 07 January, concurrently with the beginning of the UVP sampling, and continued through 12 January. Trap recovery began on 08 January and continued through 13 January. Trap depths spanned the photic zone and mesopelagic, with the shallowest at 60 m and the deepest at 965 m. Trap deployments lasted between 21 and 91 hours, with deeper traps left out for longer, to collect more biomass. Two types of traps were deployed. One set of traps, generally deployed in shallower water, had a solid cone opening with area 0.46 m². The second set had larger conical net with opening of 1.24 m² area made of 53 µm nylon mesh similar to the description in Peterson et al. (2005). All equipment were combination incubators and particle traps, but in this study we only use trap data. No poisons were used, and both living and dead zooplankton, or 'swimmers', were manually removed prior to POC analysis. Sediment trap material was filtered immediately upon trap recovery onto pre-combusted GF-75 45 mm filters (nominal pore size of 0.3 µm) and preserved until further analysis at -80°C. These filters were split into several fractions for other analyses not discussed here. Total carbon content of particles in each trap were measured by isotope ratio mass spectrometry. Elemental analyses for particulate carbon and nitrogen quantities as well as ¹³C and ¹⁵N isotopic compositions were conducted at the U.C. Davis Stable Isotope Facility (http://stableisotopefacility.ucdavis.edu) on acidified freeze-dried trap samples to capture organic elemental contributions. Carbon was below mass spectrometry detection limits in four traps – these traps were excluded from further analysis. Traps at similar depths did detect carbon, lending confidence to the

Free floating, surface tethered particle traps were used to quantify carbon fluxes from sinking particles.

- idea that these non-detections were technical in nature, due to splitting of samples for multiple analyses,
- rather than reflecting environmental conditions.

5.6 Analysis

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- 251 Particles were binned by depth with 20 m resolution between the surface and 100 m, 25 m resolution
- between 100 m and 200 m depths and 50 m resolution below 200 m. This increasing coarseness of the
- depth bins helped account for more scarce particles deeper in the water column, while maintaining higher
- depth resolution near the surface. To perform this binning, particle numbers, and volumes of water
- sampled of all observations within each depth bin were summed prior to other analyses. Most analyses
- focused on the mesopelagic, defined here as the region between the base of the secondary chlorophyll
- 257 maximum layer (160 m hereafter the base of the photic zone), which is within the ODZ, and 1000 m.
- 258 Two normalized values of particle numbers were calculated. In the first, particle numbers were divided by
- volume sampled, to generate values in $particles/m^3$. In the second, particles were divided by both volume
- sampled and the width of the particle size-bins to generate values in *particles/m³/mm*.

261 **5.6.1 Particle size distribution**

- We determined the slope and intercept of the particle size distribution spectrum by fitting a power law to
- 263 the data, which is a common function for fitting particle size distributions (Buonassissi & Dierssen,
- 264 2010). Because large particles were infrequently detected, we used a general linear model that assumed
- residuals of the data followed a negative-binomial (rather than normal) distribution. We fit the equation

$$266 ln\left(\frac{E(Total\,Particles)}{Volume*Binsize}\right) = b_0 + b_1 \ln(Size) (Eqn 1).$$

- to solve for the Intercept (b_0) and particle size distribution slope (PSD = b_1). On the left-hand side of
- 268 Eqn 1. E(Total Particles) refers to the expected number of particles in a given depth and particle size bin
- assuming a negative binomial distribution of residuals (Date, 2020; Ooi, 2013). Volume indicates the
- volume of water sampled by the UVP, or in the case of depth-binned data, the sum of the volumes of all
- UVP images in that depth interval. *Binsize* indicates the width of the particle-size bin captured by the
- UVP. Thus, if particles between 0.1 and 0.12 mm are in a particle size bin, the *Binsize* is 0.02 mm. On the
- 273 right-hand side of Eqn 1, Size corresponds to the lower bound of the particle size-bin. We use the lower
- bound of a particle size-bin, rather than its midpoint, because, due to the power-law particle size
- distribution slopes, the average size of particles in each size-bin is closer to the size-bin's lower bound
- 276 rather than its midpoint.

277 5.6.2 Estimating particle flux

- We estimated particle flux throughout the water column, by fitting particle data to trap measurements. We
- assumed that particle flux in each size bin (j) followed the equation

280 Flux =
$$\sum_{j} \left[\frac{Total\ Particles_{j}}{Volume*Binsize_{i}} * C_{f} * (Size_{j})^{A} \right]$$
 (Eqn. 2)

- Such that flux at a given depth is the sum of all size-bin specific values.
- We used the *optimize()* function in R's stats package to identify values for the C_f and A coefficients in
- Eqn 2. that yielded closest fits of the UVP estimated flux to each particle trap.
- We also estimated the exponent of the particle size to biomass exponent α and size to sinking speed
- exponent γ per the equations Biomass_i ~ Size_i^{α} and Speed_i ~ Size_i^{γ}. This is done by assuming a
- spherical drag profile, in which case $A = \alpha + \gamma$ and $\gamma = \alpha 1$ (Guidi et al., 2008); with "A" referring to
- the exponent in Eqn 2.

288 5.6.3 Size specific information

- We separately analyzed total particle numbers, particle size distribution, and particle flux for particles
- larger than or equal to 500 μm, and those smaller than 500 μm, to determine the relative contributions of
- these two particle classes to particle properties. 500 µm was chosen as it has been previously defined as
- 292 the cutoff point between microscopic "microaggregates" and macroscopic "marine snow" (Simon et al.,
- 293 2002).

294 5.7 Variability

- To explore the timescales of temporal variability in the POC flux, we determined how well the flux at
- each depth horizon can be described by the sum of daily and hourly temporal modes. This was achieved
- by fitting the general additive model of form
- 298 Flux^{1/5} ~ s(Depth) + s(Day) + s(Hour) (Eqn. 3)
- 299 This model explored whether estimated flux levels appeared to vary by decimal day and decimal hour,
- 300 holding the effects of depth constant, in the 250 m to 500 m region. The smooth terms s for *Depth* and
- 301 Day were thin plate splines, while the s term for Hour was a cyclic spline of 24-hour period.

5.8 Smoothing for Comparison to Model Results

- Normalized particle abundance data, from the only UVP cast that traversed the top 2000 m of the water
- 304 column, taken on January 13 at 06:13, was smoothed with respect to depth, time, and particle size using a
- 305 general additive model of the form

306 $\ln \left(\frac{E(Total \, Particles)}{Volume*Binsize} \right) \sim s(Depth, \ln(Size))$ (Eqn. 4)

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In this case, there is a single, two-dimensional, smooth term, rather than additive one-dimensional terms as in Eqn. 3 so that the smooth term can consider interactions between the two parameters, rather than assuming that the terms are additive. The predicted particle numbers at each particle size and depth, as well as particle size distribution spectra, and estimated particle masses of all particles smaller than 500 µm and all particles larger than or equal to 500 µm were then compared to each of Weber and Bianchi's (2020) models, corresponding to our **H1-H3**.

5.9 Modeling remineralization and sinking

To quantify disaggregation, our goal was to compare the particle size-abundance spectrum at each depth to a prediction of the null hypothesis, that it is simply governed by the effects of sinking and remineralization reshaping the spectrum observed shallower in the water column. This prediction is generated using the particle remineralization and sinking model (PRiSM), modified from DeVries et al. (2014), which we applied to the shallower spectrum as an initial condition. The difference between the null hypotheses prediction and observation indicates the role of processes not accounted for in PRiSM, such as disaggregation, aggregation, and active or advective transport of particles with a different size spectrum than the ones seen at the deeper depth. In practice we expanded the previous numerical implementation of PRiSM to allow for particle size distribution spectra with particle-size bins that match those obtained by the UVP, and to return estimates of the number of particles in those same size bins (Text S1). The model accepts inputs of particle size distributions at each depth, and changes in particle flux between each depth-bin and the next, deeper, depth-bin. The model optimizes a particle remineralization rate that would result in that observed flux loss. It finally returns a "predicted" particle size distribution spectrum that has total flux equal to the flux of the observed deeper spectrum that would be expected if the shallower spectrum only sank and remineralized. In cases where flux increased with depth, particles are assumed to put on mass rather than lose mass following a negative remineralization rate. Here, "negative remineralization" stands in for chemoautotrophy, active transport, and other processes that result in flux increases with depth. While these processes likely have more complex effect on the particle size distribution than is accounted for in our model, we note that flux increases with depth are very rare, and that allowing for negative remineralization allows our null model to be robust in those rare cases.

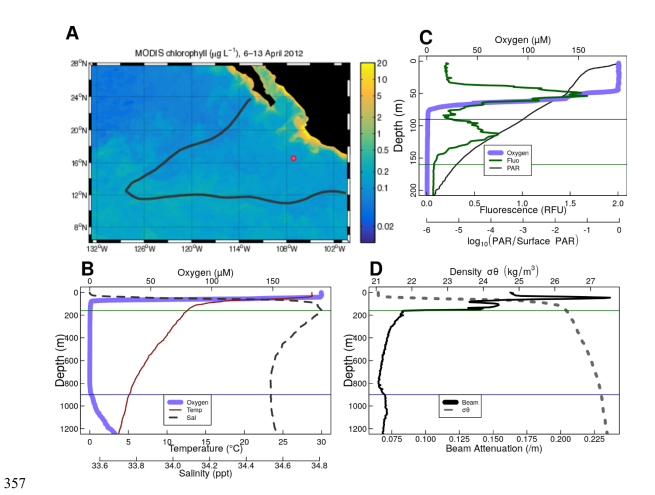
6 Results

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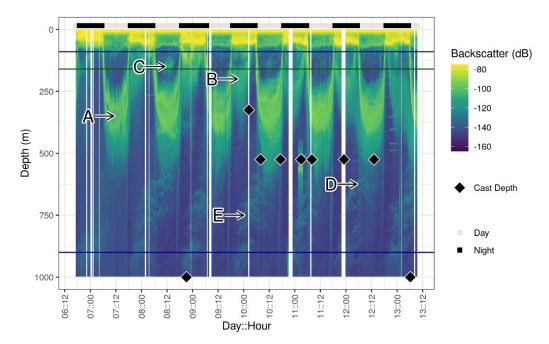
6.1 Physical and Chemical Data

337	The ODZ, characterized in this study by oxygen levels less than 1 μM , as measured by the CTD, extends
338	from 90 m to 900 m depth, with a sharp upper oxycline and a gradual lower oxycline (Figure 1B-C). This
339	station has been previously proven to be anoxic with a STOX sensor (Tiano et al., 2014). The upper
340	oxycline tracks a sharp pycnocline (Figure 1B-1D), set by the high salinity of the 13CW water mass
341	(Figure S2), and is characterized by an abrupt drop in temperature below the mixed layer and an increase
342	in salinity (Figure 1B). Water mass analysis indicated that water in the top part of the ODZ is dominated
343	by the 13CW water mass, while water between 275 and 500 m is primarily from the NEPIW, with water
344	from the AAIW dominating in the lower 500 m (Figure S2) (Evans et al., 2020). The site is characterized
345	by two fluorescence maxima (Figure 1C). The larger, shallower fluorescence peak is positioned just
346	above the oxycline, with fluorescence from this peak and oxygen attenuating together. The smaller, lower
347	peak is inside of the ODZ. Turbidity tracks the two chlorophyll peaks in the surface and has a tertiary
348	maximum at the lower oxycline (Figure 1D). The cyanobacteria at the secondary chlorophyll maximum
349	are known to be photosynthesizing and producing organic matter in the ODZ (Fuchsman et al., 2019;
350	Garcia-Robledo et al., 2017). To avoid complication due to this source of organic matter production, we
351	focus our further analysis below 160 m.
352	For the purposes of this study, we define the photic zone as ending at the base of this deeper fluorescence
353	layer (160m). This photic zone base corresponds with photosynthetically active radiation (PAR) $\leq 10^{-5}$ of
354	surface PAR levels (Figure 1C). We note that this photic zone depth is deeper than conventional
355	definitions, in which the base of the photic zone corresponds with 10 ⁻² (90 m) or 10 ⁻³ (120 m) of surface
356	PAR.



6.2 Acoustic data reveal diel migration patterns

Acoustic data, produced by the shipboard EK60 (Andersen, 2001), at ETNP Station P2, suggest the presence of multiple cohorts of migratory organisms. We focus initially on backscattering measurements from the EK60's lowest frequency 18000 Hz signal, corresponding to organisms the size of small fish, because it travels furthest into the water column and has the best resolution of the channels. Most migratory organisms appeared to leave the surface at dawn and return at dusk, spending the day between 250 m and 500 m (Figure 2A). There appeared to be two local maxima in backscattering intensity at midday, one at ~300 m and one at ~375 m (Figure 2A). There also appeared to be organisms that reverse migrated downward at dusk and upward at dawn, spending the night at ~300 m (Figure 2B). Just above the base of the photic zone, there was a peak of organisms that appeared, at mid-day, on some but not all days, without any visible dawn or dusk migration. (Figure 2C). Some diel migrators appeared to cross the ODZ and spend the day below the detection range of the EK60 (Figure 2D), as well as organisms that appeared between 500 m and 1000 m but did not appear to migrate to or from that depth at our site, but rather traveled through the EK60's field of view (Figure 2E). Similar patterns were evident in each of the other measured frequencies, with better resolution by the lower frequencies (Figure S3).



of the photic zone. Times are local Mexico General Standard time, which is the local time for the nearest port of call in Manzanillo and is equivalent to United States Central Standard Time. The black and white bar at the top indicates day and night periods, with day defined as times when the center of the sun is above the horizon, per the OCE R package. Diamonds indicate the depths and times of UVP casts, with casts deeper than 1000 m shown as diamonds on the 1000 m line. Several patterns are evident **A.** Two bands of organisms can be seen leaving the surface at dawn, spending the day between 250 m and 500 m and returning to the surface at dusk. **B.** Another group of nocturnally migrating organisms can be seen leaving the surface at dusk, spending the night near 250 m and returning at dawn. **C.** Some organisms appear at the base of the photic zone, during some, but not all mid days, and then disappear in the evening. **D.** A group of very deep migrating organisms appears to leave the surface with the diel migrators and pass all the way through the ODZ and out of the EK60's field of view. It returns at dusk. **E.** Swarms of organisms appear between 500 m and 1000 m disappearing later in the day. Swarms appear in the deepest layers at night and appear progressively shallower as the day progresses.

6.3 Flux data from traps

Flux measurements at Station P2 were consistent between the different particle trap types, showing a profile that broadly followed a power law with respect to depth, with the exception that flux appeared to increase in one trap at 500 m (Figure 3).

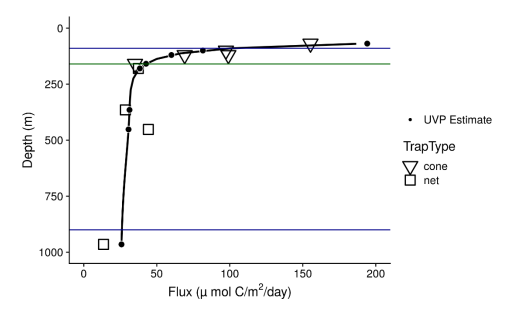


Figure 3. Sinking particle flux, measured from surface-tethered sediment traps (large symbols), at ETNP Station P2. Trap types are shown by the shape of the large points. Superimposed are

estimates of particle flux from the UVP generated by fitting the sum of particle numbers all four profiles to the trap observed flux. The black line indicates flux predictions made by fitting UVP observations to the trap data. Black circles indicate regions on the black line corresponding to the trap observation depths. Horizontal blue lines indicate the top and bottom of the ODZ, while the horizontal green line indicates the base of the photic zone.

418 6.4 Particle abundance measurements vary with size and depth 419 420 In all profiles, particle abundances were highest at the surface, and highest among the smallest particles 421 (Figure S4). Visual examination of the relationship between particle number and size suggested a power 422 law relationship where the log of volume and bin-size normalized particle abundance was proportional to 423 the log of the particles' size (Figure S5). The exception to this pattern were particles larger than 10 mm 424 (Figure S4, S5), which are rare enough that they are usually not detected by the UVP. Generalized linear 425 models that assume a negative-binomial distribution of the data accounted for this under-sampling of 426 large particles to estimate power law slopes, while considering rare occurrences of the large particles at 427 each depth (Figure S5). 428 Total particle numbers were generally similar between different casts, regardless of which day or hour 429 they were collected (Figure 4A). Particle numbers were highest in the surface and decreased within the 430 oxic region, then remained relatively constant from 160 m to 500 m, and gradually decreased between 431 500 m and the lower oxycline (Figure 4A). 432 The particle size distribution slope generally steepened (became more negative) between the base of the 433 photic zone (160 m) and 500 m, flattened (became less negative) between 500 m and 1000 m, and then 434 steepened again below 1000 m (Figure 4B). Steeper, more negative, slopes indicate a higher proportion of 435 small particles relative to large particles, while flatter, less negative, slopes indicate a more even particle 436 size distribution. Flatter distributions still have a higher absolute number of smaller particles than larger 437 particles; however, they have a higher proportion of larger particles relative to other samples with steeper 438 distributions.

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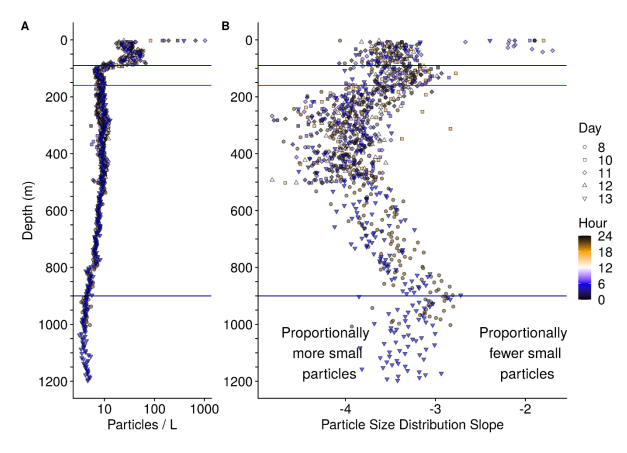


Figure 4. A. Observed, volume normalized total particle numbers from 9 casts taken at different times of the day at ETNP station P2. **B.** Calculated particle size distribution slopes of those particles. These data have not been binned by depth in order to better show sample to sample variability. Horizontal blue lines indicate the top and bottom of the ODZ, while the horizontal green line indicates the base of the photic zone. Hour corresponds to local, Mexican General Standard, time.

6.5 Estimated particle flux sometimes increases with depth in the ODZ core

- Optimization found best agreement between particle flux measured by traps, and UVP estimated particle flux when per particle flux is fit by the equation
- 449 Flux = $(133 \mu \text{ mol C} / \text{m}^2/\text{day}) = 133 * \text{Size (mm)} ^ 2.00$ (Eqn 5)
- This equation represents an empirical relationship between particle flux from traps and particle size measured by UVP. Applying this fit to the UVP data resulted in a UVP predicted flux profile that broadly fit the expected trap observed flux profiles (Figure 3).
- Particle flux profiles, predicted from the above particle size abundances and fit, varied between casts between the base of the photic zone and 500 m (Figure 5A-5B). To examine the rate of change of flux and

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455 to identify regions and time points where flux appeared to increase with depth, we examined the rate of 456 change of flux. This rate of change was fifth root transformed to normalize the data and to allow us to 457 focus on the cases where flux attenuation varied about zero, since we were interested in identifying 458 factors that related to whether flux was positive or negative. Between 250 m and 500 m, particle flux 459 appeared to increase on some, but not all, casts, while attenuating slowly on the other casts (Figure 5C). 460 Below 500 m, there were not enough casts to measure variability between casts. 461 The general additive model that quantified how the of change of flux between 250 m and 500 m varied 462 with depth, decimal study day and decimal hour found that depth (p = 0.061) and hour of the day (p =463 0.196) did not statistically associate with the fifth root transformed rate of change of flux while day of study did (p = 0.019, $R^2 = 0.264$, Figure S6). There were generally increases in flux over this region 464 towards the beginning and end of the sampling period and decreases in flux nearer to day 10 (Figure 465 466 S6B). A general additive model that looked only at the relationship between study day and rate of change 467 of flux (fifth root transformed) in this region suggested that day accounted for 14% of the variance in this 468 value, as determined by adjusted R^2 (p = 0.040). If the fifth root transformation was not applied to the rate 469 of change of flux, there was a statistically significant relationship between depth and rate of change (p = 470 0.001), but not study day (p = 0.062) or hour (p = 0.719, $R^2 = 0.341$). This pattern indicated that, without 471 the transformation, any temporal signal is swamped by the substantial changes in rate of change in depth, 472 with shallower depths losing flux faster than deeper ones.

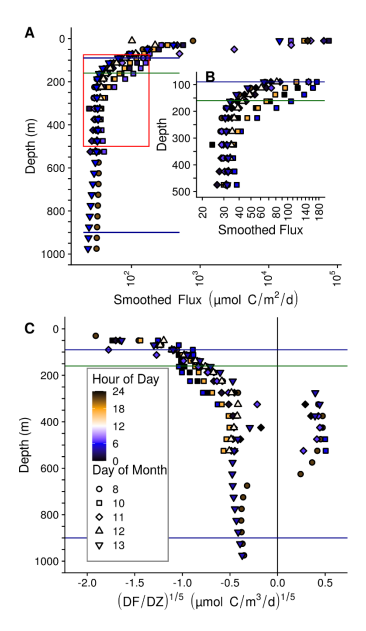


Figure 5. Within and between day variability in UVP predicted particle flux at ETNP Station P2. All profiles are depth binned with higher resolution towards the surface (methods). Horizontal blue lines indicate the top and bottom of the ODZ, while the horizontal green line indicates the base of the photic zone. **A.** Flux profiles in the top 1000 m of the water column. **B.** A more detailed depiction of the area enclosed by the rectangle in **A.** -- **C.** The rate of change of flux, divided by the rate in change in depth. The fifth root of these values are shown to highlight differences between values close to zero. Hour corresponds to local, Mexican General Standard, time.

6.6 Smoothed and averaged data

At the ETNP ODZ site, highly smoothed particle abundance data suggested that particle size, averaged across all casts, followed a pattern in which the abundance of $<500 \, \mu m$ particles increased between the oxycline and 350 m (Figure 6A), which corresponded with steepening of the particle size distribution slope (Figure 6B), and an increase in microaggregate ($<500 \, \mu m$) particle biomass (Figure 6C), but not of $>500 \, \mu m$ particle biomass (Figure 6D). Deeper in the ODZ, the microaggregate ($<500 \, \mu m$) particle number and biomass, and the particle size distribution slope declined.

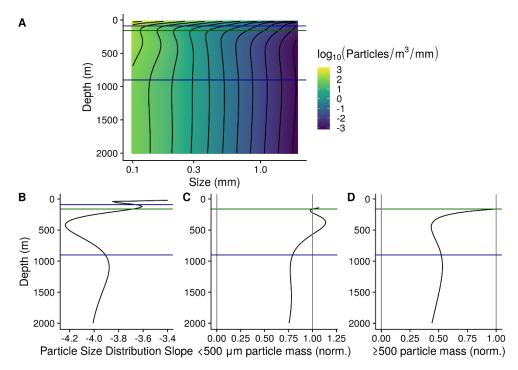


Figure 6. A. GAM smoothed, bin-size and volume normalized particle numbers across the particle size spectrum, at ETNP Station P2. Data are from the only cast that traversed the top 2000 m of the water column, collected on January 13 beginning at 06:13. Horizontal blue lines indicate the top and bottom of the ODZ, while the horizontal green line indicates the base of the photic zone. **B.** Particle size distribution slope. **C-D.** Estimated biomass of **(C)** microaggregate (<500 um) and **(D)** marine snow (≥500 um) particles, normalized to biomass at the base of the photic zone. In these two biomass panels, data from above the base of the photic zone are not shown.

At the oxic site, particle size distributions generally steepened with depth, while both microaggregates (<500 μm) and ≥500 μm estimated particle biomass followed a power law decrease with depth (Figure S10).

6.7 Particle number dynamics differ from model expectations

- The modified particle remineralization and sinking model predicted particle size distributions at each depth from the particle size distribution one depth-bin shallower and the calculated flux attenuation between the two depths. At the ETNP ODZ site, we found that the observed particle size distributions usually deviated from model expectations (Figure S11). In the model, remineralization rates are optimized, to ensure that the total predicted flux at each depth matches the observed total flux. However,
- 507 the predicted size spectrum will diverge from the observed spectrum if the assumptions of the model (i.e.,
- sinking and remineralization are the only particle transformations) are violated. The difference between
- 509 the observed and predicted flux of microaggregate particles (100 500 μm), normalized to depth,
- 510 therefore serves as a metric of observed deviations from the size distribution expected from sinking and
- remineralization alone. We call this value *Deviation from Model* (DFM).

DFM =
$$\frac{(<500 \,\mu m \,Flux \,Observed - <500 \,\mu m \,Flux \,Modeled)}{\Delta Z}$$
 (Eqn. 6)

- In the above equation ΔZ is the distance, in meters, between the current depth bin and the previous depth
- bin, whose particle size distribution is fed into the predictive model.
- DFM was positive between the base of the photic zone (160 m) and 500 m, meaning that less <500 μm
- particle flux attenuated than would be expected from the *PRiSM* model in this region (Figure 7). There
- was some variability in the DFM parameter between casts. A general additive model (GAM) that showed
- that the variability in DFM was statistically significantly related to depth ($p < 10^{-5}$), day of the study
- (p = 0.002), but not to hour of the day (p = 0.051), with these factors together explaining 41.6% of the
- variance, as measured by R². DFM was highest shallower in the water column (Figure S12A), highest
- near day 10 and lower at the beginning and end of the study (Figure S12B). A GAM that only explored
- 522 the effect of depth accounted for 27.4% of the variance. Comparing a GAM that accounted for study day
- and depth to one that only accounted for depth effects showed an increase in \mathbb{R}^2 value of 10.4%,
- suggesting that study day accounts for an additional 10.4% of the variance, after accounting for depth.
- 525 Comparing the model that accounts for depth, day and hour to one that only accounts for depth and day,
- suggests that hour of the day, while not statistically significant, could explain an additional 3.4% of the
- 527 variance. Below 500 m, DFM was negative. There were only two casts that reached below 500 m at this
- station, and so an analysis of the dynamics of DFM in this region is not possible. At P16 Station 100,
- 529 DFM was positive between the base of the photic zone and 350 m and negative below 350 m (Figure
- 530 S9C).

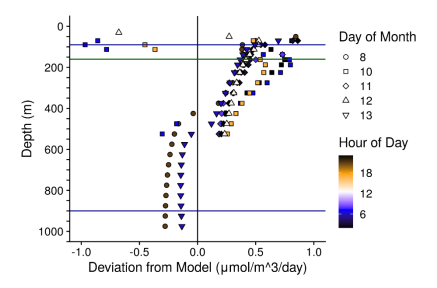


Figure 7. Deviation from Model (DFM) indicates the difference between the observed flux of small particles (< 500 μm), and the flux of small particles that would be estimated by a model, which assumes that particles in the depth bin above only remineralized and sank, following the PRiSM model. Values are normalized to the change in depth and are in units of μ mol Carbon m⁻³d⁻¹. This value serves as a metric of processes that cannot be captured by a null model, which assumes that particles only sink and remineralize. Positive values suggest an excess of <500 μm particles, suggesting disaggregation or advection of small particles, while negative values suggest a dearth of small particles, suggesting repackaging or aggregation. DFM is only reported for <500 μm particles (<500 μm), because it is the inverse of the deviation from expected flux of ≥500 μm particles. DFM is reported for all casts at ETNP Station P2. Horizontal blue lines indicate the top and bottom of the ODZ, while the horizontal green line indicates the base of the photic zone.

6.8 ETNP particle dynamics differ from those seen at an oxic site

The ODZ data were compared to an oxic water column in order to identify the spectral signatures that are particular to oxygen deficient waters. The oxic site, P16 Station 100, was characterized by a more gradually sloping pycnocline, and an oxygen minimum at 500 m of 19.7 µM, which is hypoxic (Figure S1B). There was no working fluorescence sensor on that cruise, but data from World Ocean Atlas (Boyer et al., 2018) suggest that the photic zone is characterized by a single fluorescence peak with a maximum at 110 m and which disappeared at 200 m (Figure S1C). Thus, we define the mesopelagic as beginning at 200 m at the oxic site. Turbidity followed chlorophyll concentration and did not have a peak in the mesopelagic (Figure S1D), unlike the ODZ site. There was a salinity peak at 150 m (Figure S1B).

553 Particle numbers were higher between the base of the photic zone through 1000 m at the ETNP ODZ site, 554 than at the same-latitude, oxygenic, P16 Station 100 (Figure S7A). Particle size distributions were similar 555 between the two sites above 500 m, being characterized by overlapping confidence intervals generated by 556 a general additive model. From 500 m to 1000 m, particle size distributions were flatter at the ETNP site, 557 being characterized by a smaller proportion of smaller particles, relative to larger ones (Figure S7B). 558 Microaggregate particles (100 μm - 500 μm) at the ETNP ODZ site were about two orders of magnitude 559 more common than marine snow particles (≥ 500 μm) (Figure S8). ≥500 μm particle numbers appeared 560 to attenuate more quickly than <500 µm particles, and more generally follow a power law decrease, while 561 <500 µm particles appeared to increase around 500 m depth. Flux was predicted to be predominantly 562 from <500 um, rather than ≥500 μm particles, at all depths except the shallowest depth bin in the surface 563 of the photic zone. The particle size distribution, calculated only on ≥500 µm particles, was more variable 564 between depths than calculated for <500 µm particles. Data from the oxic P16 Station 100 suggested 565 more particles, steeper particle size distribution, and more flux at this station than at the ETNP station. 566 They also suggested that differences between $<500 \mu m$ and $\ge 500 \mu m$ particles, with respect to number, 567 flux and size distribution that were broadly similar to the ones seen at ETNP Station P2. In contrast to the 568 anoxic station, at the oxic station flux always decreased with depth (Figure S9A+B).

7 Discussion

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7.1 Diel migrators spend time in the ODZ core

571 Organisms of all sizes appear to migrate into the core of the ODZ at our site. Most migrators appear to 572 leave the surface at dawn, spend the day in the top 500 m of the ODZ and return to the surface at dusk 573 (Figure 2A), while others show the opposite pattern, leaving the surface at dusk and returning at dawn 574 (Figure 2B). Diel migration is prevalent throughout the oceans (Cisewski et al., 2010; Hays, 2003; 575 Heywood, 1996; Jiang et al., 2007; Rabindranath et al., 2011; Sainmont et al., 2014; Yang et al., 2019), 576 including at other ODZ sites (Antezana, 2009; Kiko et al., 2020; Riquelme-Bugueño et al., 2020), 577 including highly anoxic sites with secondary, anoxic, deep chlorophyll maxima, like this one (Bianchi et 578 al., 2014; Hidalgo et al., 2005), and much of the ETNP ODZ (Herrera et al., 2019). Sampling efforts in 579 the Costa Rica Dome, a unique system in the ETNP, find that euphausiids and fish migrate into the ODZ 580 (Maas et al., 2014; Wishner et al., 2013), but that diel migrators are primarily 2 mm-5 mm in size 581 (Wishner et al., 2013). Krill in the Humboldt current OMZ similarly spend the day at depth and migrate to 582 the surface at night (Riquelme-Bugueño et al., 2020). The presence of organisms that appear and 583 disappear just above the base of the photic zone, in the region of the deeper anoxic fluorescence peak 584 region, but absence of a tell-tale signature of mass migration before or after they appear (Figure 2C) may

585 suggest that these organisms migrate at different times of the day to this deep region, rather than all at 586 once. Another possibility is that they pass through our station at this depth in mid-day, but migrate to 587 depth at another location. 588 The organisms that appear between 500 m and 1000 m (Figure 2E) have acoustic signatures that resemble 589 those of jellyfish (Kaartvedt et al., 2007), but could also represent other organisms such as salps (Maas et 590 al., 2014; Ressler, 2002). That they appear in horizontal bands that do not appear to trend upwards over 591 time suggests that these swarms are traveling through our site at progressively shallower depths over the 592 course of the day, but that the individual swarms are not themselves moving upward at this station. This pattern suggests that any vertical migration by these organisms happens elsewhere or occurs more slowly 593 594

than the advection seen at this site. That they appear at different depths at different times of the day

595 suggest that these organisms have some sort of vertical migration pattern. Future work may consider more

596 highly resolved spatial and temporal monitoring of this phenomenon. Indeed molecular surveys have

found evidence of both Cnidarians and Ctenophores both within and below the ETSP ODZ near Chile

598 (Parris et al., 2014).

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7.2 Flux is lower at this site than previous measurements in the ETNP

600 Flux at P2 was lower at all depths, ranging from 10–100 μmol/m²/day, than was seen in previous

601 measurements by traps at other, more productive, ODZ sites, which ranged from 1000-

602 10000 μmol/m²/day (Hartnett & Devol, 2003; Van Mooy et al., 2002).

The flux to size relationship is typical of other sites

The exponent of the particle size to flux relationship that we saw at our site (2.00) is of a similar magnitude to, but slightly smaller than, those seen by other studies that compare UVP flux to trap flux (Guidi et al., 2008; Kiko et al., 2020). Differences in the size-flux relationship could indicate that this relationship truly varies between sites, or that imprecision in flux measurements leads to differences in these values between studies. The single fit and relationship that we carried out does not account for variation in the size to flux ratio across time and depth, nor does it account for differences in particles of different origin. In practice, this value could change over depth and time. Setting up, deploying and retrieving each trap array is a large effort. However, coupled particle flux and size measurements that are more resolved with respect to depth, space or time might allow for further exploration of the spatiotemporal variability of this relationship. In other systems, combined image analysis and gel traps (McDonnell & Buesseler, 2010, 2012) have provided opportunities to explore particle size to flux

relationships and how they vary between particle types in more detail.

7.4 Remineralization rates of all particles decrease in the ODZ, but disaggregation does not

Particle size profiles, particle size distribution slopes, and estimated biovolume, averaged across all casts and smoothed, are all similar to the predictions made by Weber and Bianchi's (2020) "Model 1". (Figure 5), and therefore our hypothesis H1, that all particles are remineralized more slowly than in oxic sites. This suggests that the low oxygen at this site decreases the remineralization rate of all particles, including <500 µm microaggregates. It does not support the **H2** in which disaggregation is suppressed in the ODZ, nor H3 in which only the very large particles' remineralization is slowed due to sulfate reduction. The data at the oxic site generally conformed to Weber and Bianchi's null model, "Model 0", which was their prediction for particle distributions at oxic sites (2020). However, one difference was that the observed particle size distribution, while essentially constant from the base of the photic zone through 1000 m, appeared to steepen between 1000 m and 2000 m, suggesting an increase in the abundance of <500 µm particles, relative to Model 0. This could indicate increased disaggregation in this region or horizontal transport of small particles through advection in this region. One possible source of disaggregation are zooplankton communities that have been found to specialize in feeding in the lower oxycline (Saltzman & Wishner, 1997; Wishner et al., 1995). These communities actively seek out the lower oxycline and feed on particles that have escaped remineralization in the ODZ, potentially resulting in the increased disaggregation we observe in this depth interval. Such a community would likely be comprised primarily of small organisms which the EK60 is not able to measure at this depth. One possible source of horizontal transport is colloids in a deep iron plume (Homoky et al., 2021; Lam et al., 2020).

7.5 Zooplankton likely transport organic matter into the ODZ core

Predicted flux levels sometimes increase between 275 m and 625 m, and at all other times attenuate very slowly in this region. The EK60 data suggest the diel migration of all sizes of organisms to this region, agreeing with previous analysis of copepods collected with nets (Wishner et al., 2020). Taken together, the concurrent intermittent increases in flux with diel migration in the top 500 m suggests that zooplankton are transporting organic matter. The observation that the rate of change in flux changes with depth suggests some day-to-day variability in this transport. That this rate does not vary statistically significantly between day and night suggests that any diel release of particles is relatively small compared to the particles already present in situ. Indeed, it suggests that particle sinking is slow enough that any particles that are transported to depth during the day are retained at night. Furthermore, nocturnal migrators are likely playing a role in carbon transport which may smooth out any diel signal. Another possibility, given that the magnitude of the day-to-day variability in apparent particle flux is small, is that

the zooplankton themselves, which likely make up about 5% of what the UVP counts as particles, may be driving this apparent pattern and that particle flux itself does not vary. More likely, especially given the observation that this flux variability did not track well with the within day backscattering patterns seen by the EK60 and the small number of particles that are zooplankton, is that this factor accounts for some, but not all, of the observed variability in flux. An additional source of temporal variability in flux is variation in particle export from the photic zone. Zooplankton, if they are more common in large particle size bins, or even if they have a flatter size distribution spectrum than non-living particles, may flatten the particle size spectrum, where they are present. However, this effect, if present at our site, appears to be overpowered by the disaggregation effect, since the particle size spectra appear to be steeper where zooplankton are present. Zooplankton are also known to congregate at the lower boundaries of ODZs (Wishner et al., 2018, 2020) and high urea concentrations in the lower oxycline of the ETNP have been suggested to be due to these zooplankton (Widner et al., 2018). Beam attenuation indicates a third peak in the oxycline below the ODZ. We do not see this congregation in the EK60 data. However, it is possible that small organisms do congregate here, but are not detected by the EK60's 12000 and 20000 kHZ signals, which do not penetrate to 1000 m in our data. The EK60 data do however suggest that larger, krill to fish sized organisms are not abundant in the lower oxycline.

7.6 Zooplankton likely disaggregate particles in the ODZ core

by remineralization and sinking alone (Figure 7), between the photic zone and 500 m suggests that some process is disaggregating large particles into smaller ones. That this apparent disaggregation corresponds with the region where migratory organisms are found suggests that some of these organisms, likely small animals such as copepods and euphausiids (Herrera et al., 2019; Maas et al., 2014), may break down particles (Dilling & Alldredge, 2000; Goldthwait et al., 2005). While, in principle, other processes such as horizontal advection of water containing <500 μm particles (Inthorn, 2005) could be responsible for this increase in <500 μm particles, there is no reason to expect horizontal differences at this site, which is at the core of the ODZ and far from shore.

Other deviations from model assumptions could alternatively explain the increase in <500 μm particles relative to model predictions. In particular, smaller particles might break down more slowly than larger ones, or sink more quickly for their size than expected, as has been seen elsewhere (McDonnell & Buesseler, 2010). Our model assumes a spherical particle drag profile, such that the particle sinking speed fractal dimension (γ) is one less than the particle size fractal dimension (α) (Cram et al., 2018; Guidi et

The observation that there is greater flux by microaggregate particles (< 500 µm) than would be predicted

- al., 2008), and that these two values sum to the particle flux fractal dimension. If any of these assumptions
- do not hold, the magnitude of the values may differ.
- In contrast to the upper ODZ core, there is an apparent flattening of the particle size distribution below
- 500 m, beyond the expected effects generated by particle remineralization. This could suggest
- aggregation processes (Burd & Jackson, 2009). Indeed, aggregation could be occurring throughout the
- 686 ODZ core, but only exceed disaggregation in the lower ODZ region. Alternatively, in this region,
- processes resembling Weber and Bianchi's (2020) Model 3, corresponding to **H3**, in which large particles
- 688 remineralize more slowly than larger ones could also occur. Like aggregation, such processes could be
- occurring through the ODZ but are overwhelmed by the effects of disaggregation above 500 m.

7.7 Water mass changes may affect particle flux and size changes

- The observation that particle flux begins to attenuate below 500 m more quickly than it does between the
- base of the photic zone and 500 m could be explained in part by a shift in water mass at this depth where
- 693 AAIW begins to mix with NEPIW (Figure S2). The AAIW is suggested to have micromolar oxygen
- 694 concentrations, as compared to the NEPIW, such that a small contribution of AAIW can raise the oxygen
- 695 concentration (Evans et al., 2020). However, measurements taken at this station in 2012 observed zero
- oxygen though 800 m with the highly sensitive STOX electrode, suggesting that oxygen, if present, is
- below 4 nM (Tiano et al., 2014). It is conceivable that the AAIW has larger particle sizes and lower
- particle abundance characteristics due to its having advected from different geographic regions than the
- overlying water, but it is difficult to see why this would be the case as these water masses stay in the ODZ
- region for years (DeVries et al., 2012) and particles have a much shorter residence time. In any case, the
- NEPIW to AAIW transition coincides with the lower limit of the depth to which vertically migrating
- zooplankton travel (Figure 2), and so we are not able to deconvolve the effects of water mass changes
- from that of changes in zooplankton effects on particle characteristics.
- The change in water mass between 13CW and NEPIW, around 250 m, in contrast, does not appear to
- correspond to any apparent changes in particle flux or size. Thus, we would argue that any historical
- effects of these water mass differences are likely to be small, and that active transport differences above
- and below 500 m likely have a larger effect.

7.8 Oxic site differences

- The oxic site provides validation that the patterns that we see at the ETNP are unique to the ODZ region,
- and do not apply to a same latitude ODZ site. The particle size distribution slope varied little and there
- was not an increase in particle mass in the oxic site, consistent with Weber and Bianchi's (2020) null
- model (Figure S10), in which oxygen is not limiting and particle sizes are not affected by anoxia. In this

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- case, small particles break down more quickly in the oxic site than our site and so there is no small
 particle excess in this region. Similarly, the higher flux attenuation in the oxic site (Figure S9A) suggests
 that the differences in attenuation of all particle sizes by microbes at both sites do indeed drive differences
 in flux profiles, and by extension transfer efficiency, between oxic and anoxic regions. The lack of
 increases in flux at the oxic site (Figure S9B) suggest that active transport may play a greater role in the
 anoxic region than elsewhere. The lack in apparent excess of small particles over model prediction
 (Figure S9C) could either indicate less activity by zooplankton in this region, or perhaps that
- remineralization of small particles quickly removes any small particles produced by zooplankton in this

721 region.

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7.9 Future directions

- We advocate exploring the relationships between particle size distribution, flux and acoustic signatures in
- other parts of the ETNP and other ODZ regions. Expanded spatial analysis of particle size spectra in
- ODZs would allow the community to confirm whether Weber and Bianchi's (2020) model (H1), that
- particles of all sizes break down more slowly in ODZs, applies elsewhere. Similarly, a clear next step is to
- apply our disaggregation model to other ocean regions, perhaps using particle size data already collected
- 728 by other groups (Guidi et al., 2008; Kiko et al., 2017, 2020).
- While the UVP characterizes dynamics of particles $>100 \mu m$, particles smaller than this range contribute
- dramatically to carbon flux (Durkin et al., 2015), and so their size distribution matters as well. However,
- at some point particles become small enough that they likely do not sink, and so exploring
- remineralization and disaggregation of <500 μm microaggregate particles into non-sinking size classes
- would provide valuable context to these measurements. In-situ pumped POC data from the GEOTRACES
- program have been used to describe the dynamics of smaller particle size classes (Lam et al., 2011; Lam
- 735 & Marchal, 2015). Other sensors, such as coulter counters (Sheldon et al., 1972) and Laser In-Situ
- 736 Scattering transmissometers (Ahn & Grant, 2007) provide size resolved distribution information about
- these smaller size classes of particles. Comparison between UVP data and past and ongoing (Siegel et al.,
- 738 2016) studies of the characteristics of <100 µm particles provide opportunities to better understand the
- dynamics of the full range of particle sizes.
- 740 The image data collected by the UVP offers opportunities to quantify the abundance and taxonomic
- distribution of the zooplankton that migrate into the mesopelagic, as well as the particle types within this
- 742 region. Identifying this visual data would have the added benefit of allowing researchers analyze particle
- size spectra, rather than the sum of particles and zooplankton as we do here.

8 Conclusions

744

745 If ODZs expand in response to the changing climate, larger areas of the ocean are likely to resemble this 746 environment, which is oligotrophic and has an oxygen deficient zone spanning most of the mesopelagic 747 zone. Previous models and observations have suggested that ODZs are sites of efficient carbon transfer to 748 the deep ocean (Cram et al., 2018; Hartnett & Devol, 2003; Van Mooy et al., 2002; Weber & Bianchi, 749 2020), and our data appear to support this contention. Indeed, the mechanism of efficient transfer appears 750 to be slowing of particle remineralization, presumably from decreased microbial metabolism, with 751 zooplankton playing an important role in both active particle transport and particle disaggregation. 752 Our data could potentially be used in conjunction with mechanistic models (e.g. Weber & Bianchi, 2020) 753 to constrain the relative carbon oxidation rate by nitrate reduction, denitrification and sulfate reduction 754 processes, which is currently poorly understood (Bristow, 2018). Furthermore, it appears that diel 755 migratory organisms both disaggregate particles and transport carbon throughout the top 500 m of the 756 water column. Day-to-day and within day variability in organic matter transport was evident, though 757 overall patterns in particle size, flux and disaggregation appeared to be consistent over the course of the 758 time-series. The change in particle abundance and size between 500 m and the bottom of the ODZ has 759 implications for the free-living microbes living in this region. These microbes are likely particularly 760 organic matter starved, and so these decaying particles are likely an important energy source for them. 761 Our data highlights the heterogeneous nature of the ETNP ODZ with depth and indicates that more 762 detailed sampling should be performed for rate and microbial measurements to properly extrapolate to the 763 entire ODZ.

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10 References

776	Ahn, J. H., & Grant, S. B. (2007). Size Distribution, Sources, and Seasonality of Suspended Particles in
777	Southern California Marine Bathing Waters. Environmental Science & Technology, 41(3), 695-
778	702. https://doi.org/10.1021/es061960+
779	Andersen, L. N. (2001). The new Simrad EK60 scientific echo sounder system. The Journal of the
780	Acoustical Society of America, 109(5), 2336–2336. https://doi.org/10.1121/1.4744207
781	Antezana, T. (2009). Species-specific patterns of diel migration into the Oxygen Minimum Zone by
782	euphausiids in the Humboldt Current Ecosystem. Progress in Oceanography, 83(1), 228–236.
783	https://doi.org/10.1016/j.pocean.2009.07.039
784	Archibald, K. M., Siegel, D. A., & Doney, S. C. (2019). Modeling the Impact of Zooplankton Diel
785	Vertical Migration on the Carbon Export Flux of the Biological Pump. Global Biogeochemical
786	Cycles, 33(2), 181–199. https://doi.org/10.1029/2018GB005983
787	Bianchi, D., Stock, C., Galbraith, E. D., & Sarmiento, J. L. (2013). Diel vertical migration: Ecological
788	controls and impacts on the biological pump in a one-dimensional ocean model. Global
789	Biogeochemical Cycles, 27(2), 478–491. https://doi.org/10.1002/gbc.20031
790	Bianchi, D., Babbin, A. R., & Galbraith, E. D. (2014). Enhancement of anammox by the excretion of diel
791	vertical migrators. Proceedings of the National Academy of Sciences, 111(44), 15653–15658.
792	https://doi.org/10.1073/pnas.1410790111
793	Bianchi, D., Weber, T. S., Kiko, R., & Deutsch, C. (2018). Global niche of marine anaerobic metabolism
794	expanded by particle microenvironments. Nature Geoscience, 11(4), 263.
795	https://doi.org/10.1038/s41561-018-0081-0
796	Boyer, T., Garcia, H. E., Locarini, R. A., Ricardo, A., Zweng, M. M., Mishonov, A. V., et al. (2018).
797	World Ocean Atlas 2018. NOAA National Centers for Environmntal Information.

798	Briggs, N., Dall'Olmo, G., & Claustre, H. (2020). Major role of particle fragmentation in regulating
799	biological sequestration of CO2 by the oceans. Science, 367(6479), 791–793.
800	https://doi.org/10.1126/science.aay1790
801	Bristow, L. A. (2018). Anoxia in the snow. Nature Geoscience, 11(4), 226-227.
802	https://doi.org/10.1038/s41561-018-0088-6
803	Buonassissi, C. J., & Dierssen, H. M. (2010). A regional comparison of particle size distributions and the
804	power law approximation in oceanic and estuarine surface waters. Journal of Geophysical
805	Research: Oceans, 115(C10). https://doi.org/10.1029/2010JC006256
806	Burd, A. B., & Jackson, G. A. (2009). Particle Aggregation. Annual Review of Marine Science, 1(1), 65-
807	90. https://doi.org/10.1146/annurev.marine.010908.163904
808	Cavan, E. L., Trimmer, M., Shelley, F., & Sanders, R. (2017). Remineralization of particulate organic
809	carbon in an ocean oxygen minimum zone. Nature Communications, 8, 14847.
810	https://doi.org/10.1038/ncomms14847
811	Cisewski, B., Strass, V. H., Rhein, M., & Krägefsky, S. (2010). Seasonal variation of diel vertical
812	migration of zooplankton from ADCP backscatter time series data in the Lazarev Sea, Antarctica.
813	Deep Sea Research Part I: Oceanographic Research Papers, 57(1), 78–94.
814	https://doi.org/10.1016/j.dsr.2009.10.005
815	Cram, J. A., Weber, T., Leung, S. W., McDonnell, A. M. P., Liang, JH., & Deutsch, C. (2018). The Role
816	of Particle Size, Ballast, Temperature, and Oxygen in the Sinking Flux to the Deep Sea. Global
817	Biogeochemical Cycles, 32(5), 858–876. https://doi.org/10.1029/2017GB005710
818	Date, S. (2020, November 21). Generalized Linear Models. Retrieved May 2, 2021, from
819	https://towardsdatascience.com/generalized-linear-models-9ec4dfe3dc3f
820	Deutsch, C., Berelson, W., Thunell, R., Weber, T., Tems, C., McManus, J., et al. (2014). Centennial
821	changes in North Pacific anoxia linked to tropical trade winds. Science, 345(6197), 665-668.
822	https://doi.org/10.1126/science.1252332

823	DeVries, T., & Weber, T. (2017). The export and fate of organic matter in the ocean: New constraints
824	from combining satellite and oceanographic tracer observations. Global Biogeochemical Cycles,
825	2016GB005551. https://doi.org/10.1002/2016GB005551
826	DeVries, T., Deutsch, C., Primeau, F., Chang, B., & Devol, A. (2012). Global rates of water-column
827	denitrification derived from nitrogen gas measurements. Nature Geoscience, 5(8), 547-550.
828	https://doi.org/10.1038/ngeo1515
829	DeVries, T., Liang, JH., & Deutsch, C. (2014). A mechanistic particle flux model applied to the oceanic
830	phosphorus cycle. Biogeosciences Discuss., 11(3), 3653–3699. https://doi.org/10.5194/bgd-11-
831	3653-2014
832	Dilling, L., & Alldredge, A. L. (2000). Fragmentation of marine snow by swimming macrozooplankton:
833	A new process impacting carbon cycling in the sea. Deep Sea Research Part I: Oceanographic
834	Research Papers, 47(7), 1227–1245. https://doi.org/10.1016/S0967-0637(99)00105-3
835	Durkin, C. A., Estapa, M. L., & Buesseler, K. O. (2015). Observations of carbon export by small sinking
836	particles in the upper mesopelagic. Marine Chemistry, 175, 72-81.
837	https://doi.org/10.1016/j.marchem.2015.02.011
838	Evans, N., Boles, E., Kwiecinski, J. V., Mullen, S., Wolf, M., Devol, A. H., et al. (2020). The role of
839	water masses in shaping the distribution of redox active compounds in the Eastern Tropical North
840	Pacific oxygen deficient zone and influencing low oxygen concentrations in the eastern Pacific
841	Ocean. Limnology and Oceanography, 65(8), 1688–1705. https://doi.org/10.1002/lno.11412
842	Francois, R., Honjo, S., Krishfield, R., & Manganini, S. (2002). Factors controlling the flux of organic
843	carbon to the bathypelagic zone of the ocean. Global Biogeochemical Cycles, 16(4), 34-1-34-20.
844	https://doi.org/10.1029/2001GB001722
845	Fuchsman, C. A., Devol, A. H., Saunders, J. K., McKay, C., & Rocap, G. (2017). Niche Partitioning of
846	the N Cycling Microbial Community of an Offshore Oxygen Deficient Zone. Frontiers in
847	Microbiology, 8. https://doi.org/10.3389/fmicb.2017.02384

848	Fuchsman, C. A., Palevsky, H. I., Widner, B., Duffy, M., Carlson, M. C. G., Neibauer, J. A., et al. (2019)
849	Cyanobacteria and cyanophage contributions to carbon and nitrogen cycling in an oligotrophic
850	oxygen-deficient zone. The ISME Journal, 1. https://doi.org/10.1038/s41396-019-0452-6
851	Garcia-Robledo, E., Padilla, C. C., Aldunate, M., Stewart, F. J., Ulloa, O., Paulmier, A., et al. (2017).
852	Cryptic oxygen cycling in anoxic marine zones. Proceedings of the National Academy of
853	Sciences, 114(31), 8319-8324. https://doi.org/10.1073/pnas.1619844114
854	Goldthwait, S. A., Carlson, C. A., Henderson, G. K., & Alldredge, A. L. (2005). Effects of physical
855	fragmentation on remineralization of marine snow. Marine Ecology Progress Series, 305, 59-65.
856	Guidi, L., Jackson, G. A., Stemmann, L., Miquel, J. C., Picheral, M., & Gorsky, G. (2008). Relationship
857	between particle size distribution and flux in the mesopelagic zone. Deep Sea Research Part I:
858	Oceanographic Research Papers, 55(10), 1364–1374. https://doi.org/10.1016/j.dsr.2008.05.014
859	Hannides, C. C. S., Landry, M. R., Benitez-Nelson, C. R., Styles, R. M., Montoya, J. P., & Karl, D. M.
860	(2009). Export stoichiometry and migrant-mediated flux of phosphorus in the North Pacific
861	Subtropical Gyre. Deep Sea Research Part I: Oceanographic Research Papers, 56(1), 73–88.
862	https://doi.org/10.1016/j.dsr.2008.08.003
863	Hartnett, H. E., & Devol, A. H. (2003). Role of a strong oxygen-deficient zone in the preservation and
864	degradation of organic matter: a carbon budget for the continental margins of northwest Mexico
865	and Washington State. Geochimica et Cosmochimica Acta, 67(2), 247–264.
866	https://doi.org/10.1016/S0016-7037(02)01076-1
867	Hays, G. C. (2003). A review of the adaptive significance and ecosystem consequences of zooplankton
868	diel vertical migrations. In M. B. Jones, A. Ingólfsson, E. Ólafsson, G. V. Helgason, K.
869	Gunnarsson, & J. Svavarsson (Eds.), Migrations and Dispersal of Marine Organisms (pp. 163-
870	170). Dordrecht: Springer Netherlands. https://doi.org/10.1007/978-94-017-2276-6_18
871	Herrera, I., Yebra, L., Antezana, T., Giraldo, A., Färber-Lorda, J., & Hernández-León, S. (2019). Vertical
872	variability of Euphausia distinguenda metabolic rates during diel migration into the oxygen

873	minimum zone of the Eastern Tropical Pacific off Mexico. Journal of Plankton Research, 41(2),
874	165–176. https://doi.org/10.1093/plankt/fbz004
875	Heywood, K. J. (1996). Diel vertical migration of zooplankton in the Northeast Atlantic. Journal of
876	Plankton Research, 18(2), 163–184. https://doi.org/10.1093/plankt/18.2.163
877	Hidalgo, P., Escribano, R., & Morales, C. E. (2005). Ontogenetic vertical distribution and diel migration
878	of the copepod Eucalanus inermis in the oxygen minimum zone off northern Chile (20–21° S).
879	Journal of Plankton Research, 27(6), 519-529. https://doi.org/10.1093/plankt/fbi025
880	Homoky, W. B., Conway, T. M., John, S. G., König, D., Deng, F., Tagliabue, A., & Mills, R. A. (2021).
881	Iron colloids dominate sedimentary supply to the ocean interior. Proceedings of the National
882	Academy of Sciences, 118(13), e2016078118. https://doi.org/10.1073/pnas.2016078118
883	Horak, R. E. A., Ruef, W., Ward, B. B., & Devol, A. H. (2016). Expansion of denitrification and anoxia
884	in the eastern tropical North Pacific from 1972 to 2012. Geophysical Research Letters, 43(10),
885	2016GL068871. https://doi.org/10.1002/2016GL068871
886	Inthorn, M. (2005). Lateral particle transport in nepheloid layers - a key factor for organic matter
887	distribution and quality in the Benguela high-productivity area. Retrieved from
888	https://media.suub.uni-bremen.de/handle/elib/2212
889	Ito, T., Minobe, S., Long, M. C., & Deutsch, C. (2017). Upper ocean O2 trends: 1958-2015. Geophysical
890	Research Letters, 44(9), 4214-4223. https://doi.org/10.1002/2017GL073613
891	Jackson, G. A., & Burd, A. B. (2001). A model for the distribution of particle flux in the mid-water
892	column controlled by subsurface biotic interactions. Deep Sea Research Part II: Topical Studies
893	in Oceanography, 49(1), 193–217. https://doi.org/10.1016/S0967-0645(01)00100-X
894	Jiang, S., Dickey, T. D., Steinberg, D. K., & Madin, L. P. (2007). Temporal variability of zooplankton
895	biomass from ADCP backscatter time series data at the Bermuda Testbed Mooring site. Deep Sec
896	Research Part I: Oceanographic Research Papers, 54(4), 608–636.
897	https://doi.org/10.1016/j.dsr.2006.12.011

398	Kaartvedt, S., Klevjer, T. A., Torgersen, T., Sørnes, T. A., & Røstad, A. (2007). Diel vertical migration of
399	individual jellyfish (Periphylla periphylla). Limnology and Oceanography, 52(3), 975–983.
900	https://doi.org/10.4319/lo.2007.52.3.0975
901	Keil, R. G., Neibauer, J. A., & Devol, A. H. (2016). A multiproxy approach to understanding the
902	"enhanced" flux of organic matter through the oxygen-deficient waters of the Arabian Sea.
903	Biogeosciences, 13(7), 2077–2092. http://dx.doi.org/10.5194/bg-13-2077-2016
904	Kiko, R., Biastoch, A., Brandt, P., Cravatte, S., Hauss, H., Hummels, R., et al. (2017). Biological and
905	physical influences on marine snowfall at the equator. Nature Geoscience, 10(11), 852-858.
906	https://doi.org/10.1038/ngeo3042
907	Kiko, R., Brandt, P., Christiansen, S., Faustmann, J., Kriest, I., Rodrigues, E., et al. (2020). Zooplankton-
808	Mediated Fluxes in the Eastern Tropical North Atlantic. Frontiers in Marine Science, 7.
909	https://doi.org/10.3389/fmars.2020.00358
910	Kwon, E. Y., & Primeau, F. (2008). Optimization and sensitivity of a global biogeochemistry ocean
911	model using combined in situ DIC, alkalinity, and phosphate data. Journal of Geophysical
912	Research: Oceans, 113(C8), C08011. https://doi.org/10.1029/2007JC004520
913	Lam, P., & Kuypers, M. M. M. (2011). Microbial Nitrogen Cycling Processes in Oxygen Minimum
914	Zones. Annual Review of Marine Science, 3(1), 317–345. https://doi.org/10.1146/annurev-marine
915	120709-142814
916	Lam, P. J., & Marchal, O. (2015). Insights into Particle Cycling from Thorium and Particle Data. Annual
917	Review of Marine Science, 7(1), 159–184. https://doi.org/10.1146/annurev-marine-010814-
918	015623
919	Lam, P. J., Doney, S. C., & Bishop, J. K. B. (2011). The dynamic ocean biological pump: Insights from a
920	global compilation of particulate organic carbon, CaCO3, and opal concentration profiles from
921	the mesopelagic. Global Biogeochemical Cycles, 25(3), n/a-n/a.
22	https://doi.org/10.1029/2010GB003868

923 Lam, P. J., Heller, M. I., Lerner, P. E., Moffett, J. W., & Buck, K. N. (2020). Unexpected Source and 924 Transport of Iron from the Deep Peru Margin. ACS Earth and Space Chemistry, 4(7), 977–992. 925 https://doi.org/10.1021/acsearthspacechem.0c00066 926 Lampitt, R. S., Noji, T., & von Bodungen, B. (1990). What happens to zooplankton faecal pellets? 927 Implications for material flux. Marine Biology, 104(1), 15–23. 928 https://doi.org/10.1007/BF01313152 929 Lee, W.-J., Staneva, V., Mayorga, E., Nguyen, K., Satiewan, L., & Majeed, I. (2021). Echopype: 930 Enhancing the interoperability and scalability of ocean sonar data processing. The Journal of the Acoustical Society of America, 149(4), A63-A63. https://doi.org/10.1121/10.0004522 931 932 Maas, A. E., Frazar, S. L., Outram, D. M., Seibel, B. A., & Wishner, K. F. (2014). Fine-scale vertical 933 distribution of macroplankton and micronekton in the Eastern Tropical North Pacific in 934 association with an oxygen minimum zone. Journal of Plankton Research, 36(6), 1557–1575. 935 https://doi.org/10.1093/plankt/fbu077 936 McDonnell, A. M. P., & Buesseler, K. O. (2010). Variability in the average sinking velocity of marine 937 particles. Limnology and Oceanography, 55(5), 2085–2096. 938 https://doi.org/10.4319/lo.2010.55.5.2085 939 McDonnell, A. M. P., & Buesseler, K. O. (2012). A new method for the estimation of sinking particle 940 fluxes from measurements of the particle size distribution, average sinking velocity, and carbon 941 content. Limnology and Oceanography: Methods, 10(5), 329–346. 942 https://doi.org/10.4319/lom.2012.10.329 943 Neuer, S., Iversen, M., & Fischer, G. (2014). The Ocean's Biological Carbon pump as part of the global 944 Carbon Cycle. *Limnology and Oceanography E-Lectures*, 4(4), 1–51. 945 https://doi.org/10.4319/lol.2014.sneuer.miversen.gfischer.9 946 Noji, T. T., Estep, K. W., Macintyre, F., & Norrbin, F. (1991). Image Analysis of Faecal Material Grazed 947 Upon by Three Species Of Copepods: Evidence For Coprorhexy, Coprophagy and Coprochaly.

948	Journal of the Marine Biological Association of the United Kingdom, 71(2), 465–480.
949	https://doi.org/10.1017/S0025315400051717
950	Ooi, H. (2013, August 8). Where does the offset go in Poisson/negative binomial regression? Retrieved
951	May 2, 2021, from https://stats.stackexchange.com/questions/66791/where-does-the-offset-go-in-
952	poisson-negative-binomial-regression
953	Parris, D. J., Ganesh, S., Edgcomb, V. P., DeLong, E. F., & Stewart, F. J. (2014). Microbial eukaryote
954	diversity in the marine oxygen minimum zone off northern Chile. Frontiers in Microbiology, 5.
955	https://doi.org/10.3389/fmicb.2014.00543
956	Passow, U., & Carlson, C. (2012). The biological pump in a high CO2 world. Marine Ecology Progress
957	Series, 470, 249–271. https://doi.org/10.3354/meps09985
958	Pavia, F. J., Anderson, R. F., Lam, P. J., Cael, B. B., Vivancos, S. M., Fleisher, M. Q., et al. (2019).
959	Shallow particulate organic carbon regeneration in the South Pacific Ocean. Proceedings of the
960	National Academy of Sciences, 116(20), 9753–9758. https://doi.org/10.1073/pnas.1901863116
961	Pennington, J. T., Mahoney, K. L., Kuwahara, V. S., Kolber, D. D., Calienes, R., & Chavez, F. P. (2006).
962	Primary production in the eastern tropical Pacific: A review. Progress in Oceanography, 69(2-4),
963	285–317. https://doi.org/10.1016/j.pocean.2006.03.012
964	Peterson, M. L., Wakeham, S. G., Lee, C., Askea, M. A., & Miquel, J. C. (2005). Novel techniques for
965	collection of sinking particles in the ocean and determining their settling rates. Limnology and
966	Oceanography: Methods, 3(12), 520–532. https://doi.org/10.4319/lom.2005.3.520
967	Picheral, M., Guidi, L., Stemmann, L., Karl, D. M., Iddaoud, G., & Gorsky, G. (2010). The Underwater
968	Vision Profiler 5: An advanced instrument for high spatial resolution studies of particle size
969	spectra and zooplankton. Limnology and Oceanography: Methods, 8(9), 462-473.
970	https://doi.org/10.4319/lom.2010.8.462
971	Picheral, M., Colin, S., & Irisson, JO. (2017). EcoTaxa, a tool for the taxonomic classification of
972	images. Retrieved from http://ecotaxa.obs-vlfr.fr

973	Poulsen, L., & Kiørboe, T. (2005). Coprophagy and coprorhexy in the copepods Acartia tonsa and
974	Temora longicornis: clearance rates and feeding behaviour. Marine Ecology Progress Series, 299
975	217-227. https://doi.org/10.3354/meps299217
976	Rabindranath, A., Daase, M., Falk-Petersen, S., Wold, A., Wallace, M. I., Berge, J., & Brierley, A. S.
977	(2011). Seasonal and diel vertical migration of zooplankton in the High Arctic during the autumn
978	midnight sun of 2008. Marine Biodiversity, 41(3), 365–382. https://doi.org/10.1007/s12526-010-
979	0067-7
980	Raven, M. R., Keil, R. G., & Webb, S. M. (2021). Microbial sulfate reduction and organic sulfur
981	formation in sinking marine particles. Science, 371(6525), 178–181.
982	https://doi.org/10.1126/science.abc6035
983	Ressler, P. H. (2002). Acoustic backscatter measurements with a 153kHz ADCP in the northeastern Gulf
984	of Mexico: determination of dominant zooplankton and micronekton scatterers. Deep Sea
985	Research Part I: Oceanographic Research Papers, 49(11), 2035–2051.
986	https://doi.org/10.1016/S0967-0637(02)00117-6
987	Riquelme-Bugueño, R., Pérez-Santos, I., Alegría, N., Vargas, C. A., Urbina, M. A., & Escribano, R.
988	(2020). Diel vertical migration into anoxic and high- p CO 2 waters: acoustic and net-based krill
989	observations in the Humboldt Current. Scientific Reports, 10(1), 17181.
990	https://doi.org/10.1038/s41598-020-73702-z
991	Rocap, G., Keil, R., Devol, A., & Deutsch, C. (2017). Water temperature, salinity, and other data from
992	CTD taken from the RV Sikuliaq in the Pacific Ocean between San Diego, California and
993	Manzanillo, Mexico from 2016-12-21 to 2017-01-13 (NCEI Accession 0164968). [Temperature,
994	Salinity, Oxygen, Beam Attenuation, Fluorescence, PARJ. NOAA National Centers for
995	Environmental Information. Retrieved from https://accession.nodc.noaa.gov/0164968
996	Roullier, F., Berline, L., Guidi, L., Durrieu De Madron, X., Picheral, M., Sciandra, A., et al. (2014).
997	Particle size distribution and estimated carbon flux across the Arabian Sea oxygen minimum
998	zone. Biogeosciences, 11(16), 4541-4557. https://doi.org/10.5194/bg-11-4541-2014

999	Sainmont, J., Gislason, A., Heuschele, J., Webster, C. N., Sylvander, P., Wang, M., & Varpe, Ø. (2014).
1000	Inter- and intra-specific diurnal habitat selection of zooplankton during the spring bloom
1001	observed by Video Plankton Recorder. Marine Biology, 161(8), 1931–1941.
1002	https://doi.org/10.1007/s00227-014-2475-x
1003	Saltzman, J., & Wishner, K. F. (1997). Zooplankton ecology in the eastern tropical Pacific oxygen
1004	minimum zone above a seamount: 2. Vertical distribution of copepods. Deep Sea Research Part
1005	I: Oceanographic Research Papers, 44(6), 931–954. https://doi.org/10.1016/S0967-
1006	0637(97)00006-X
1007	Saunders, J. K., Fuchsman, C. A., McKay, C., & Rocap, G. (2019). Complete arsenic-based respiratory
1008	cycle in the marine microbial communities of pelagic oxygen-deficient zones. Proceedings of the
1009	National Academy of Sciences, 116(20), 9925–9930. https://doi.org/10.1073/pnas.1818349116
1010	Schmidtko, S., Stramma, L., & Visbeck, M. (2017). Decline in global oceanic oxygen content during the
1011	past five decades. Nature, 542(7641), 335–341. https://doi.org/10.1038/nature21399
1012	Seibel, B. A. (2011). Critical oxygen levels and metabolic suppression in oceanic oxygen minimum
1013	zones. <i>Journal of Experimental Biology</i> , 214(2), 326–336. https://doi.org/10.1242/jeb.049171
1014	Sheldon, R. W., Prakash, A., & Sutcliffe Jr., W. H. (1972). The Size Distribution of Particles in the
1015	Ocean. Limnology and Oceanography, 17(3), 327–340. https://doi.org/10.4319/lo.1972.17.3.0327
1016	Siegel, D. A., Buesseler, K. O., Behrenfeld, M. J., Benitez-Nelson, C. R., Boss, E., Brzezinski, M. A., et
1017	al. (2016). Prediction of the Export and Fate of Global Ocean Net Primary Production: The
1018	EXPORTS Science Plan. Frontiers in Marine Science, 3.
1019	https://doi.org/10.3389/fmars.2016.00022
1020	Simon, M., Grossart, H., Schweitzer, B., & Ploug, H. (2002). Microbial ecology of organic aggregates in
1021	aquatic ecosystems. Aquatic Microbial Ecology, 28(2), 175–211.
1022	https://doi.org/10.3354/ame028175
1023	Steinberg, D. K., & Landry, M. R. (2017). Zooplankton and the Ocean Carbon Cycle. Annual Review of
1024	Marine Science, 9, 413-444. https://doi.org/10.1146/annurev-marine-010814-015924

1025	Steinberg, D. K., Carlson, C. A., Bates, N. R., Goldthwait, S. A., Madin, L. P., & Michaels, A. F. (2000).
1026	Zooplankton vertical migration and the active transport of dissolved organic and inorganic carbon
1027	in the Sargasso Sea. Deep Sea Research Part I: Oceanographic Research Papers, 47(1), 137-
1028	158. https://doi.org/10.1016/S0967-0637(99)00052-7
1029	Stramma, L., Johnson, G. C., Sprintall, J., & Mohrholz, V. (2008). Expanding Oxygen-Minimum Zones
1030	in the Tropical Oceans. Science, 320(5876), 655-658. https://doi.org/10.1126/science.1153847
1031	Stukel, M. R., Décima, M., Landry, M. R., & Selph, K. E. (2018). Nitrogen and Isotope Flows Through
1032	the Costa Rica Dome Upwelling Ecosystem: The Crucial Mesozooplankton Role in Export Flux.
1033	Global Biogeochemical Cycles, 32(12), 1815–1832. https://doi.org/10.1029/2018GB005968
1034	Stukel, M. R., Ohman, M. D., Kelly, T. B., & Biard, T. (2019). The Roles of Suspension-Feeding and
1035	Flux-Feeding Zooplankton as Gatekeepers of Particle Flux Into the Mesopelagic Ocean in the
1036	Northeast Pacific. Frontiers in Marine Science, 6. https://doi.org/10.3389/fmars.2019.00397
1037	Tiano, L., Garcia-Robledo, E., Dalsgaard, T., Devol, A. H., Ward, B. B., Ulloa, O., et al. (2014). Oxygen
1038	distribution and aerobic respiration in the north and south eastern tropical Pacific oxygen
1039	minimum zones. Deep Sea Research Part I: Oceanographic Research Papers, 94, 173-183.
1040	https://doi.org/10.1016/j.dsr.2014.10.001
1041	Turner, J. T. (2015). Zooplankton fecal pellets, marine snow, phytodetritus and the ocean's biological
1042	pump. Progress in Oceanography, 130, 205–248. https://doi.org/10.1016/j.pocean.2014.08.005
1043	Van Mooy, B. A. S., Keil, R. G., & Devol, A. H. (2002). Impact of suboxia on sinking particulate organic
1044	carbon: Enhanced carbon flux and preferential degradation of amino acids via denitrification.
1045	Geochimica et Cosmochimica Acta, 66(3), 457–465. https://doi.org/10.1016/S0016-
1046	7037(01)00787-6
1047	Weber, T., & Bianchi, D. (2020). Efficient Particle Transfer to Depth in Oxygen Minimum Zones of the
1048	Pacific and Indian Oceans. Frontiers in Earth Science, 8.
1049	https://doi.org/10.3389/feart.2020.00376

1050	Widner, B., Fuchsman, C. A., Chang, B. X., Rocap, G., & Mulholland, M. R. (2018). Utilization of urea
1051	and cyanate in waters overlying and within the eastern tropical north Pacific oxygen deficient
1052	zone. FEMS Microbiology Ecology, 94(10). https://doi.org/10.1093/femsec/fiy138
1053	Wilson, S. E., Steinberg, D. K., & Buesseler, K. O. (2008). Changes in fecal pellet characteristics with
1054	depth as indicators of zooplankton repackaging of particles in the mesopelagic zone of the
1055	subtropical and subarctic North Pacific Ocean. Deep Sea Research Part II: Topical Studies in
1056	Oceanography, 55(14-15), 1636-1647. https://doi.org/10.1016/j.dsr2.2008.04.019
1057	Wishner, K. F., Ashjian, C. J., Gelfman, C., Gowing, M. M., Kann, L., Levin, L. A., et al. (1995). Pelagic
1058	and benthic ecology of the lower interface of the Eastern Tropical Pacific oxygen minimum zone.
1059	Deep Sea Research Part I: Oceanographic Research Papers, 42(1), 93–115.
1060	https://doi.org/10.1016/0967-0637(94)00021-J
1061	Wishner, K. F., Outram, D. M., Seibel, B. A., Daly, K. L., & Williams, R. L. (2013). Zooplankton in the
1062	eastern tropical north Pacific: Boundary effects of oxygen minimum zone expansion. Deep Sea
1063	Research Part I: Oceanographic Research Papers, 79, 122–140.
1064	https://doi.org/10.1016/j.dsr.2013.05.012
1065	Wishner, K. F., Seibel, B. A., Roman, C., Deutsch, C., Outram, D., Shaw, C. T., et al. (2018). Ocean
1066	deoxygenation and zooplankton: Very small oxygen differences matter. Science Advances, 4(12),
1067	eaau5180. https://doi.org/10.1126/sciadv.aau5180
1068	Wishner, K. F., Seibel, B., & Outram, D. (2020). Ocean deoxygenation and copepods: coping with
1069	oxygen minimum zone variability. Biogeosciences, 17(8), 2315–2339. https://doi.org/10.5194/bg-
1070	17-2315-2020
1071	Yang, C., Xu, D., Chen, Z., Wang, J., Xu, M., Yuan, Y., & Zhou, M. (2019). Diel vertical migration of
1072	zooplankton and micronekton on the northern slope of the South China Sea observed by a moored
1073	ADCP. Deep Sea Research Part II: Topical Studies in Oceanography, 167, 93–104.
1074	https://doi.org/10.1016/j.dsr2.2019.04.012
1075	