

1 **Zooplankton diel vertical migration and downward C flux into the Oxygen Minimum Zone in the**
2 **highly productive upwelling region off Northern Chile**

3 Pritha Tutasi^{1,3,4}, Ruben Escribano^{2,3}

4 ¹Doctoral Program of Oceanography, Universidad de Concepción, Chile

5 ²Department of Oceanography and Instituto Milenio de Oceanografía (IMO), Facultad de Ciencias
6 Naturales y Oceanográficas, ³Universidad de Concepción, Concepción, P.O. Box 160 C, Chile

7 ⁴Dirección Oceanografía Naval, Instituto Oceanográfico de la Armada (INOCAR), Guayaquil, Ecuador

8 **Correspondence to: Pritha Tutasi (prithatutasi@udec.cl)**

9 **Abstract.** Diel vertical migration (DVM), can enhance the vertical flux of carbon (C) and so contribute to the functioning of
10 the biological pump in the ocean. The magnitude and efficiency of this active transport of C may depend on the size and
11 taxonomic structure of the migrant zooplankton. However, the impact that a variable community structure can have on
12 zooplankton-mediated downward C flux has not been properly addressed. This taxonomic effect may become critically
13 important in highly productive eastern boundary upwelling systems (EBUS), where high levels of zooplankton biomass are
14 found in the coastal zone and composed by a diverse community with variable DVM behavior. In these systems, presence of
15 a subsurface oxygen minimum zone (OMZ) can impose an additional constraint to vertical migration and so influence the
16 downward C export. Here, we address these issues based on a vertically-stratified zooplankton sampling at three stations off
17 northern Chile (20°S-30°S) during November-December 2015. Automated analysis of zooplankton composition and taxa-
18 structured biomass allowed us to estimate daily migrant biomass by taxa and their amplitude of migration. We found that a
19 higher biomass aggregates above the oxycline, associated with more oxygenated surface waters and this was more evident
20 upon a more intense OMZ. Some taxonomic groups, however, were found closely associated with the OMZ. Most taxa were
21 able to perform DVM in the upwelling zone withstanding severe hypoxia. Also, strong migrants, such as eucalanid copepods
22 and euphausiids, can exhibit a large migration amplitude (~500 m), remaining either temporarily or permanently within the
23 core of the OMZ and thus contributing to the release of C below the thermocline. Our estimates of DVM-mediated C flux
24 suggested that a mean migrant biomass of ca. 958 mg C m⁻² d⁻¹ may contribute with about 71.3mg C m⁻² d⁻¹ to the OMZ
25 system through respiration, mortality and C excretion at depth, accounting for ca. 4% of the net primary production, and so
26 implying the existence of an efficient mechanism to incorporate freshly produced C into the OMZ. This downward C flux
27 mediated by zooplankton is however spatially variable and mostly dependent on the taxonomic structure due to variable
28 migration amplitude and DVM behavior.

29 **1 Introduction**

30 The Oxygen Minimum Zone (OMZ) in the Southeast Pacific, the fourth largest of the six permanent hypoxic regions in the
31 world oceans (Paulmier *et al.*, 2006), is a key component of the water column and a permanent feature intruding the coastal
32 zone of Ecuador, Peru and Chile (Fuenzalida *et al.*, 2009; Paulmier *et al.*, 2006). In the highly productive upwelling region of
33 northern Chile, the OMZ is closely linked to wind-driven upwelling in the coastal area and associated to the Equatorial
34 Subsurface Water (ESSW), which is transported southward along the continental shelf by the Peru-Chile Undercurrent (PUC),
35 as far south as 48°S (Fuenzalida *et al.*, 2009; Morales *et al.*, 1996a; Silva and Neshyba, 1979). Off Iquique (20°S) the OMZ is
36 characterized by being thick (500 m), very intense ($< 20 \mu\text{mol kg}^{-1}$) and with O_2 concentrations in the core of OMZ among the
37 lowest found in the global ocean reaching the current detection limit ($< 1 \mu\text{M}$) (Ulloa and Pantoja, 2009), although it becomes
38 thinner at about 30°S (Paulmier *et al.*, 2006).

39 During the last decades, the OMZ systems have attracted much scientific interest because of evidence showing that hypoxic
40 and anoxic conditions in coastal areas are expanding and becoming more intense (Ekau *et al.*, 2010; Stramma *et al.*, 2008). At
41 present, ongoing ocean deoxygenation is widely recognized as linked to global warming and it is rising much concern in
42 modern oceanography (Breitburg *et al.*, 2018).

43 The presence of oxygen-depleted water becomes a critical physiological constraint for pelagic and benthic organisms
44 inhabiting the upwelling zone, impacting their biomass and productivity, the species diversity, distribution, behaviour and
45 metabolic activity (Wishner *et al.*, 2018; Ekau *et al.*, 2010; Grantham *et al.*, 2004). For instance, diel vertical migration (DVM),
46 a common feature of the various size groups of zooplankton and also one of the most important movements of biomass in the
47 ocean, can also be affected by changes in intensity and distribution of the OMZ (Wishner *et al.*, 2018, 2013; Escribano *et al.*,
48 2009; Fernández-Álamo and Färber-Lorda, 2006; Hidalgo *et al.*, 2005; Morales *et al.*, 1996; Judkins, 1980). The OMZ can act
49 as an ecological barrier for vertical distribution of many organisms, constraining most zooplankton to a narrow (50 m) upper
50 layer, as shown in the coastal upwelling zone off Chile according to the works of Escribano (2006) and Donoso and Escribano
51 (2014). Zooplankton also become limited to the upper 150 or 300 m in the eastern tropical north Pacific (Wishner *et al.*, 2013).
52 However, the OMZ can also offer refuge for species adapted to live there, creating microhabitats of differing oxygen
53 concentration that are characterized by layers of high zooplankton biomass and abundance, with distinct species zonation
54 (Antezana, 2009; Wishner *et al.*, 2008; Fernández-Álamo and Färber-Lorda, 2006), which, in turn, may have important
55 consequences for carbon (C) cycling and its vertical flux. For example, it is known that zooplankton in the coastal upwelling
56 region off northern Chile may play a significant biogeochemical role by promoting carbon flux into the subsurface OMZ
57 (Escribano *et al.*, 2009). Therefore a significant proportion of the vertical material flux from the euphotic zone to the deep sea
58 (> 200 m) and within the food chain could be determined by DVM of zooplankton (Longhurst and Williams, 1992; Steinberg
59 and Landry, 2017).

60 As important contributors to the functioning of the biological pump, diel zooplankton migrants can actively increase the
61 magnitude of C export by transporting surface-ingested material in their guts to deep waters where it can be metabolized

62 (Steinberg and Landry, 2017). Zooplankton moults or mortality at depth can also contribute to the transportation of assimilated
63 organic biomass into the deep waters (Ducklow *et al.*, 2001). The biological pump process is also thought to be related to the
64 size structure of dominant zooplankton. This because some groups with large body sizes may exhibit a greater range of vertical
65 migration and sometimes higher levels of biomass, and so influencing the biogeochemical fluxes (Dai *et al.*, 2016; Ducklow
66 *et al.*, 2001). However, the effect of variable size structure on DVM performance and its consequence for active C transport
67 has not indeed been assessed. Size-structure is certainly related to zooplankton composition which has hardly been properly
68 addressed when examining the role of DVM on C flux. For instance, in areas with hypoxic subsurface layers some species are
69 more active migrants and thus more efficient C-transporters, because they have developed adaptations to low oxygen
70 conditions and can even use the OMZ as their habitat, either temporarily or permanently (Escribano *et al.*, 2009; Gonzalez and
71 Quiñones, 2002; Seibel, 2011). Adaptation may include in some cases reduction of aerobic metabolism by more than 50%
72 during exposure to hypoxia conditions, as a mechanism to facilitate low oxygen tolerance, and so reducing dramatically energy
73 expenditure during daytime within low oxygen waters, and therefore affecting the carbon C flux in areas subjected to low
74 concentrations of oxygen (Seibel *et al.*, 2016).

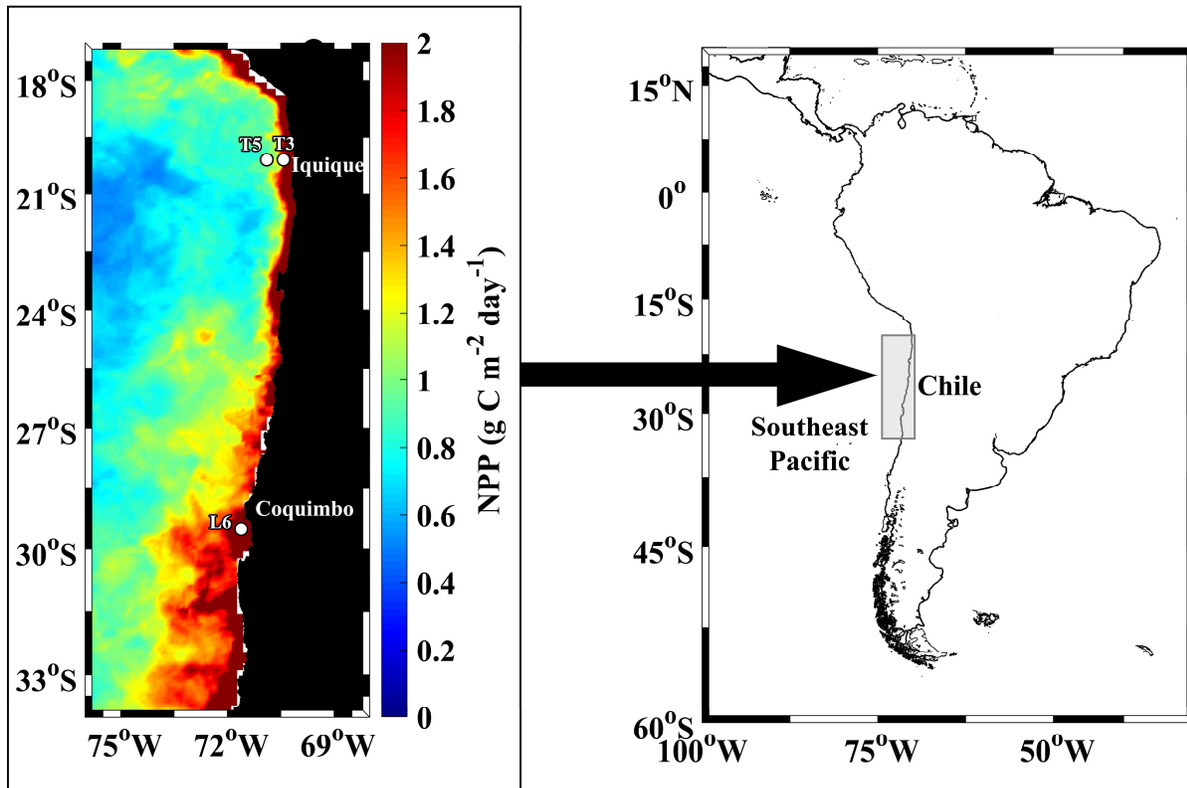
75 C-export to depth may also depend on the amount of biomass being produced in the photic zone. Primary production promotes
76 zooplankton feeding and growth and therefore determines C availability for both passive and active transport to depth. In this
77 context, highly productive upwelling zones can be assumed as systems where the C-flux mediated by zooplankton DVM can
78 be enhanced, although certainly depending on the size and taxonomic structure of zooplankton. In these regions, a shallow
79 OMZ might exert a further impact on the C-flux by affecting DVM or zooplankton metabolism at depth. In the present study,
80 based on vertically-resolved resolution sampling and automated analysis of mesozooplankton, we assessed zooplankton
81 vertical migration and downward C to the OMZ in the highly productive upwelling region of Northern Chile. We aimed at
82 understanding the role that taxonomic and size-structure can play in the magnitude and variability of the DVM behaviour
83 interacting with a shallow OMZ, and the implications this interaction can have on the magnitude of the downward C flux in a
84 highly productive coastal upwelling zone

85 **2 Methods**

86 **2.1 Study Area**

87 The study area was located in the Southeast Pacific Ocean and covered the coastal zone of the northern upwelling region of
88 Chile (21-29°S) (**Fig. 1**), which is a region known to be subjected to wind-driven upwelling throughout the year and containing
89 an intense and shallow OMZ (Ulloa *et al.*, 2012). The sampling design comprised three stations: Two stations (St. T3 and St.
90 T5) across a zonal section off Iquique (20°S) and a coastal station (St. L6) off Coquimbo (29°S). The study was carried out
91 during the LowpHox cruise conducted in November-December 2015 onboard the R/V *Cabo de Hornos*. At each station,
92 temperature, salinity and dissolved oxygen (DO) were recorded from 1000 m (St.T5 and St.T3) and only 356 m (St. L6) using

93 an oceanographic rosette with a CTD SeaBird 911 (SBE 911 plus) equipped with a SeaBird SBE-43 oxygen sensor and a
94 Seatech Fluorometer. Discrete water samples were also obtained for chemical measurements of oxygen.



95
96 **Fig.1.** Study area at the northern upwelling region of Chile, indicating sampling stations during the LowpHox cruise conducted
97 in November-December 2015 onboard the R/V *Cabo de Hornos*. Satellite estimated Net Primary Production (NPP), averaged
98 for November-December 2015 is shown. NPP was obtained from <http://science.oregonstate.edu.ocean.productivity>.
99

100 2.2 Zooplankton sampling

101 Zooplankton samples were collected during daytime and night-time conditions in two consecutive days at the three stations
102 off northern Chile (T5-T3-L6) (**Fig. 1**), also as indicated in **Table S1 (Supplemental Material)**. Vertical hauls of zooplankton
103 were performed from 600 to 0 m depth with a Multi Plankton Sampler Hydro-Bios Multinet system with a 0.25 m² opening
104 area and equipped with 200- μ m mesh-size nets. The Multinet towing speed was 1 m s⁻¹ and the flowmeter in the mouth of the
105 Multinet was to estimate the volume of filtered water. Once onboard the collected zooplankton samples were preserved
106 immediately in 5% buffered formalin-seawater solution. At T3 and T5, four replicate day and night hauls were conducted
107 (resulting in a total of eight hauls and 40 discrete samples at these stations). At L6, two replicate day and night hauls were

108 conducted (4 hauls and 20 samples total). from 600 to 0 m depth. Each sample corresponded to a different depth strata (30-0,
109 90-30, 150-90, 400-150 and 600–400m depth). These strata were defined in according to distribution of oxygen concentration
110 and localization of the OMZ (**Fig. 2**). Then, from the vertical profiles of oxygen and coinciding with the sampled layers of the
111 Multinet, strata were defined as: A well oxygenated stratum with oxygen approaching air saturation ($>250 \mu\text{mol O}_2 \text{ kg}^{-1}$) was
112 defined as the oxic mixed layer (OX-ML), an upper O_2 gradient (oxycline), defined by the level where O_2 reaches 4% of the
113 surface O_2 (Paulmier *et al.*, 2006), at whose base is located the upper boundary of the OMZ ($45 \mu\text{mol O}_2$ isoline, OMZ-UB)
114 (Escribano *et al.*, 2009; Hidalgo *et al.*, 2005; Morales *et al.*, 1999); the OMZ core defined by an upper boundary (OMZ-UC)
115 with the lowest concentration of O_2 ($<20 \mu\text{mol O}_2 \text{ kg}^{-1}$) and a lower boundary (OMZ-LC) (1 to $<20 \mu\text{mol O}_2 \text{ kg}^{-1}$), and finally
116 a lower O_2 gradient (OMZ-LW). Depth ranges and oxygen levels for these strata are detailed in **Table S2 (Supplemental**
117 **Material)**.

118

119 **2.3 Taxonomic and size measurements**

120 Taxonomic identification and enumeration of taxa were carried out by analysis of digitized images obtained with the Hydroptic
121 ZooScan digital imaging system (Gorsky *et al.*, 2010). Each sample was wet-sieved through a $1000 \mu\text{m}$ mesh into two size
122 fractions, although a few samples were not split into two fraction, because contained too few animals. Then, each size-fraction
123 was fractionated again separately with a Motoda splitter until the zooplankton concentration was sufficiently diluted to avoid
124 contact between organisms in the ZooScan scanning frame. Fractioning into small and large organisms and consequent separate
125 image acquisition of the two size classes prevented underestimates of large-rare objects which may need less fractioning
126 (Gorsky *et al.*, 2010). 179 subsamples were scanned and digitized at a resolution of 2400 dpi after manual separation of objects
127 on the scanning tray. After processing the samples with ZooProcess software, each of the objects was automatically sorted
128 with the help of a learning set, and then the sorting was visually validated by an expert (for details, see Chang *et al.*, 2012;
129 Gorsky *et al.*, 2010). Organisms making up the ZooScan datasets were enumerated, measured, biomass-estimated and
130 classified into 27 taxonomic groups, such as Copepods, Chaetognaths, Euphausiid, Gelatinous and other zooplankton. The
131 volume-specific abundance (ind. m^{-3}) of total zooplankton or of each taxonomic group was calculated following Eq. (1):

$$132 \text{ Abundance}/\text{m}^3 = \frac{N * S}{\text{Vol}} \quad (1)$$

133 were, N is the number of individuals with same identification (e.g. in last column written “copepod”), and Vol is net volume
134 Stratum-integrated abundance (ind. m^{-2}) was obtained after multiplying by width (m) of a given stratum.

135

136 **2.4 Patterns of vertical distribution of migrating zooplankton**

137 For the analysis of vertical distribution of organisms, the density estimates of the organisms were standardized to number of
138 individuals per m^3 (for each stratum) or per m^2 (for integrated values). In order to quantify the presence and extent of DVM of

139 various taxa at each station, we calculated weighted mean depth (WMD) for zooplankton abundance, as a measure of the center
140 of gravity of a population's vertical distribution for each taxon and haul, according to Andersen *et al.*(2004) following Eq. (2):

$$141 \quad WMD = \frac{\sum(ni*zi*di)}{\sum(ni*zi)}, \quad (2)$$

142 where d is the mean depth of the strata (m), z the width (m) of the strata and ni the abundance (ind. m⁻³) of a given i taxonomic
143 group.

144 We calculated the amplitude of vertical migration (Δ DVM) as the difference between the WMD of the organisms during the
145 day and the night and therefore this Δ DVM was considered as the criterion to assess the DVM behaviour for each taxonomic
146 group. Positive values indicated normal DVM (pattern of nocturnal ascent by individuals that reside at depth by day) and
147 negative values indicated reverse DVM (pattern of nocturnal descent by individuals that reside near the surface by day). The
148 individuals that occupied the same depth stratum by day and by night, whether near the surface or at depth were considered as
149 non migrant in according to Ohman (1990).

150

151 **2.5 Biomass estimates and Carbon fluxes**

152 The ZooScan Integrated System also provided zooplankton body size in terms of area (mm²) or volume (mm³) for each
153 organism. We used these organisms' area or volume to estimate dry weight of each individual of different taxonomic groups
154 using published regression equations relating organism size, area or volume to individual weight as detailed in **Table S3**
155 **(Supplemental Material)**. Mass unit conversions between dry weight (DW) and carbon content (C) were performed using
156 averaged conversion factors obtained for different zooplankton groups (Kjørboe, 2013) and ichthyoplankton (Childress and
157 Nygaard, 1973) (Table S3 Supplemental Material). Added biomasses (μ g C ind.⁻¹) of individuals within taxonomic categories
158 identified by ZooScan allowed us to estimate total biomass per taxa (mg C m⁻³) for each sample by station, daytime vs night-
159 time condition, and depth strata. Integrated values of biomass per depth strata (mg C m⁻²) and taxa were calculated multiplying
160 by strata width (m).

161 To calculate the migrant biomass, we integrated biomass in the upper 90 m layer from our two sampled strata 0-30 m and 30-
162 90 m. This 0-90 m stratum was considered the approximate above-oxyclyne layer after examining the vertical profiles of
163 oxygen. Biomass at night was thus subtracted from the corresponding day biomass in this layer to assess daily changes
164 involving migrants as in Putzeys *et al.*(2011). Thus, the negative values of the day-minus-night biomass corresponded to
165 migrant biomass that reached the epipelagic layer at night including organisms inhabiting above and below the oxycline.

166 The proportion of migrant biomass with respect to observed biomass in the upper 90 m of a given taxonomic group was defined
167 at the rate of migration on a daily basis. This rate of migration could thus be used as an index of DMV behaviour for a
168 taxonomic group. We additionally estimated daily migrant biomass from the difference between day and night samples in the
169 deeper 90-600 m layer (integrated data) and compared these estimates with those from upper 0-90 m layer.

170 To estimate the active C flux at each sampling station, we considered three processes contributing to C at depth mediated by
171 migrant zooplankton: respiration (R), excretion (E) and mortality (M) at depth. Respiration at depth (>90 m) was estimated
172 using the equation provided by Ikeda (1985) which relates individual respiration rate with body mass and temperature (**Table**
173 **S2 (Supplemental Material)**), independently of taxonomic category which may have a minor effect on R in according to a
174 more recent assessment (Ikeda, 2014). Mean body mass ($\mu\text{g C}$) for each taxonomic group from ZooScan estimates and mean
175 in situ temperature were used to obtain integrated R at each depth strata. Estimates of R for each taxonomic group are shown
176 in **Table S6 (Supplemental Material)**. Integrated R per station for the 0-600 depth strata along with the corresponding
177 integrated biomass was then used to estimate the fraction of C being respired at depth by zooplankton. The contribution of C
178 by excretion (E) at depth was assumed to be 31% of R as suggested by (Steinberg *et al.*, 2000) and daily mortality at depth
179 (M) was considered to be in the range of 0.03 and 0.05 as suggested by Edvardsen *et al.* (2002), so that a median value of 0.04
180 as a fraction of migrant biomass was assumed. Vertical C flux mediated by zooplankton was thus estimated as,

181

$$182 \text{ C Flux} = \text{MB} \times [(\text{R} + \text{M} + \text{E})/2] \quad (3)$$

183

184 where MB is the migrant biomass (mg C m^{-2}), R and M are daily Respiration and Mortality (expressed as a fraction of migrant
185 biomass), and E is the C excretion expressed as 0.31R . The three processes are divided by 2 assuming a 12 h incursion at
186 depth. We did not include the contribution by egestion at depth, because of lack of reliable estimates of ingestion rates in the
187 photic zone during our study.

188

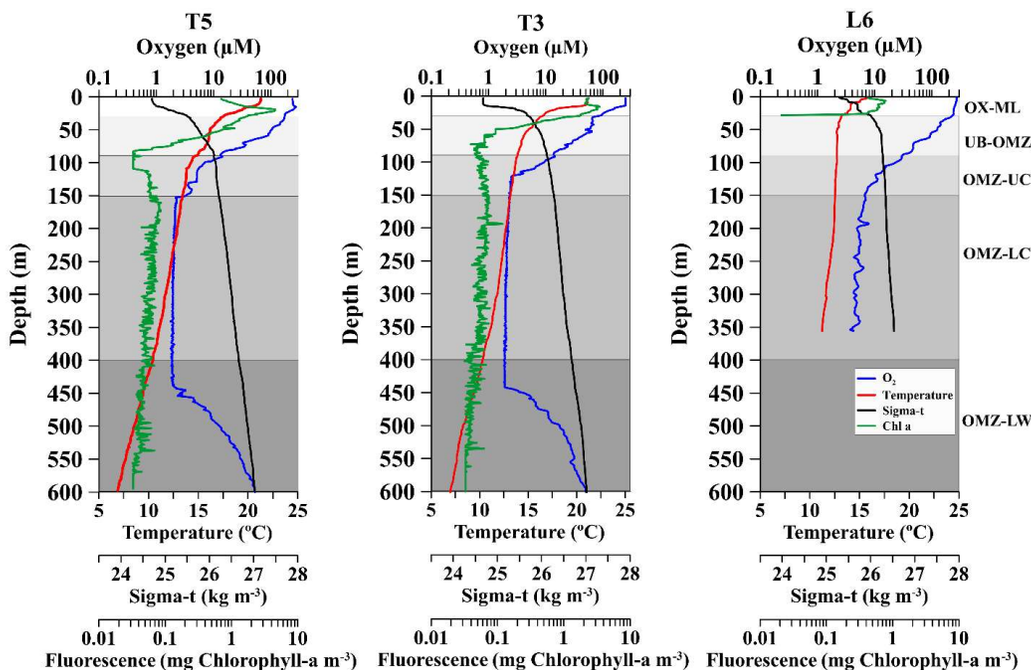
189 **2.6 Statistical analysis**

190 For statistical analysis, as a criterion for determining if the DVM was significant, we tested for differences in the WMD mean
191 between day and night using a two-tailed *t* test. We considered the occurrence of DVM when the difference in the WMD mean
192 between day and night was significant ($p < 0.05$). In order to evaluate the similarity/dissimilarity in the abundance and biomass
193 among stations, strata, and day–night conditions, the multivariate grouping techniques were applied ("cluster analysis"),
194 ANOSIM (Two-Way Crossed Analysis) tests and multidimensional scaling (MDS) with the data transformed in PRIMER v
195 6.1.16 (2013), prior to the application of the Bray-Curtis similarity index (Bray and Curtis, 1957). In general, WMD for
196 taxonomic groups did not exhibit a pronounced bimodal vertical distribution.

197 **3 Results**

198 **3.1 Hydrographic conditions**

199 Across the zonal section off Iquique the offshore station (St. T5) and onshore station (St. T3) showed two contrasting
 200 hydrographic regimes regarding the OMZ. Station T5 had a less pronounced and thicker OMZ than station T3. At both stations
 201 the 5 strata were well defined in the water column (**Fig. 2**). The OX-ML ($>250 \mu\text{mol O}_2 \text{ kg}^{-1}$) was present at 18 m (St. T5) and
 202 15 m (St. T3). The oxycline gradually decreased from oxic ($\sim 250 \mu\text{mol O}_2 \text{ kg}^{-1}$) to suboxic ($<20 \mu\text{mol O}_2 \text{ kg}^{-1}$)
 203 conditions associated with a strong stratification in the upper 80 m depth. The $45 \mu\text{mol O}_2$ isoline (OMZ-UB) was at the base
 204 of the oxycline at 70 m (St. T5) and 59 m (St. T3). The OMZ core ($<20 \mu\text{mol O}_2 \text{ kg}^{-1}$) was below the thermocline and
 205 below the 26.5 kg m^{-3} isopycnal following description of Paulmier *et al.* (2006). In the oceanic station (St. T5) the OMZ core
 206 was between 80 to 514 m, while in the coastal station (St. T3) it was between 80 to 507 m with 423 m thickness. The O_2
 207 concentration in the OMZ core was ca. $1 \mu\text{mol O}_2 \text{ kg}^{-1}$. The OMZ-LW at both stations was delimited above the core and below
 208 the depth where the O_2 slope changed significantly (slope break $>20 \mu\text{mol / m}$) (**Fig. 2**).
 209



210
 211 **Fig. 2.** Vertical profiles of dissolved oxygen (O_2), temperature, Chlorophyll-a and water density (sigma-t), at 3 stations off
 212 northern Chile (see **Fig. 1**) during the LowpHox Cruise in the austral spring 2015. Shaded areas represent different layers
 213 sampled with the mulinet and defined according to oxygen concentration. OX-M= Oxic mixed layer, OMZ-UB= Upper
 214 boundary of the oxygen minimum zone, OMZ-UC= Upper core of the oxygen minimum zone, OMZ-LC= Lower core of the
 215 OMZ and OMZ-LW=lower oxygen gradient.

216 The structure of the OMZ at the coastal station (St. L6) off Coquimbo (29°S) (**Fig.1**), was similar to St. T3 (21°S), but in this
217 area the OMZ was deeper and thinner. The OX-ML was shallower. The OMZ -UB (45 $\mu\text{mol O}_2$) in the base of the oxycline
218 was down to 80 m. The low O_2 concentrations in the core were less intense than at 21°S (4 to 20 $\mu\text{mol O}_2\text{kg}^{-1}$) and it was
219 located below 100 m (**Fig. 2**). The OMZ-LW could not be assessed because of lack of CTD data below 350 m.
220 Additional oceanographic variables showed a surface warming ($>20^\circ\text{C}$) and strongly stratified conditions at the three stations
221 with a sharp thermocline in the upper 100 m, coinciding with the oxycline, whereas Chlorophyll-a maximum ($>5\text{ mg}$
222 Chlorophyll-a m^{-3} was in the upper 20 m (**Fig. 2**).

223 **3.2 Zooplankton composition and abundance**

224 A total of 27 zooplankton taxa were identified by the ZooScan and ZooProcess (**Table S4 Supplemental Material**). The
225 number of taxa varied among stations and strata. Across the zonal section off Iquique the number of taxonomic groups
226 fluctuated between 23 (St. T3) and 26 (St. T5), whereas 25 taxa were off 29°S (St. L6). The most dominant taxa both daytime
227 and night conditions, were: Copepods 87% (in St. T5), 79% (in St. T3) and 69% (in St. L6). This group was constituted by
228 small Copepods, large Copepods, the eucalanid Copepods and the *Acartia* spp Copepods; fish eggs constituting 2% (in St. T5),
229 5% (in St. T3) and 6% (in St. L6); Nauplii being $<1\%$ (in St. T5), $<1\%$ (in St. T3) and 7% (in St. L6); Appendicularia 5% (in
230 St. T5), 4% (in St. T3) and 3% (in St. L6) (**Table S5 Supplemental Material**). The remaining 19 pooled groups only
231 constituted $<6\%$ (in St. T5), 11% (in St. T3) and 15% (in St. L6). The total integrated abundances of zooplankton (0-600 m) by
232 sampling station are in **Table S4 (Supplemental Material)**. As based on a Two-Way Crossed Analysis ANOSIM test, this
233 water-column integrated abundance did not show significant differences between day and night samples ($p>0.05$). However,
234 the abundance of these zooplankton groups regarding stations was significantly different (Two-Way Crossed Analysis
235 ANOSIM $p<0.05$), so that the stations were treated independently. Off Iquique the abundance was the lowest at the onshore
236 station (St. T3 with 18%), which was characterized by the strongest and most extensive OMZ in the study area. These values
237 increased at the offshore station (St. T5 with 31%), where the OMZ was less pronounced and thicker. Unlike stations T3 and
238 T5, the onshore station off 29°S (St. L6) had a weaker and less extensive OMZ showing the highest zooplankton abundance
239 (51% greater).

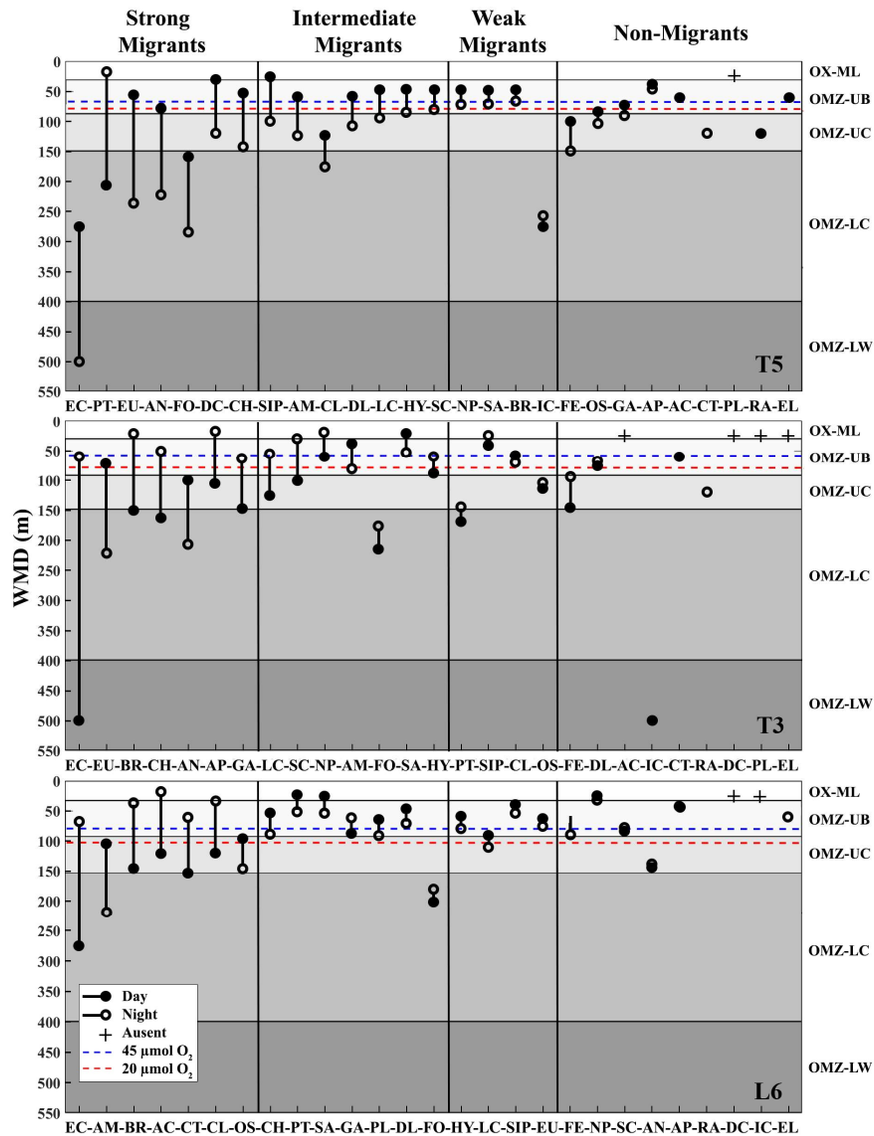
240

241 **3.3 Diel vertical migration (DVM) and vertical distribution**

242 **3.3.1 Main migrant groups of zooplankton**

243 The diel vertical migration of 27 zooplankton taxa in the 0–600 m water column is in **Fig. 3**. These taxa were classified into
244 four groups according to their amplitude of migration (ΔDVM) (**Table 1**): **1) Strong migrants**, represented by taxa with a
245 strong DVM and a broad range of ΔDVM from 225 to 99 m (in St. T5), 440 to 84 m (in St. T3) and 208 to 87 m (in St. L6).
246 This group constituted 70% of taxa with higher ΔDVM . The composition of taxa in this group was variable at each station

247 **(Table 1)**, but in general was well represented by Eucalanid copepods (EC), Euphausiids (EU), *Acartia* Copepods (AC),
248 Ctenophores (CT), Decapods (DC), Annelids (AN), Bryozoan L (BR), Pteropods (PT) and Chaetognaths (CH). These taxa
249 were mostly concentrated in the oxic surface stratum (**OX-ML**), and the **OMZ core** showing a strong interaction with both
250 the **OMZ-UC** and the **OMZ-LC**, and so changing from normoxia to hypoxia condition and vice versa between 0 to 550 m
251 **(Fig. 3), 2) Intermediate migrants**, represented by taxa with a moderate DVM and a range of Δ DVM from 73 to 34 m (in
252 St. T5), 70 to 27 m (in St. T3) and 49 to 22 m (in St. L6). This group constituted 23% of taxa with moderate Δ DVM. The
253 composition of taxa in this group was also variable at each station **(Table 1)**, but it was mostly represented by small (SC) and
254 large Copepods (LC), Amphipods (AM), Cirripedia Larvae (CL), Gastropods (GA), Siphonophores (SIP) and Appendicularia
255 (AP). These taxa were mostly concentrated in the oxic surface strata (**OX-ML**) and in the **OMZ-UC** showing some interaction
256 with the **OMZ core** and vertically changing from normoxia to hypoxia condition, and vice versa between 0 to 200 m, **3) Weak**
257 **migrants**, represented by taxa that with a weak DVM and a range of Δ DVM of 24 to 18 m (in St.T5), 23 to 12 m (in St.T3)
258 and 21 to 11 m (in St. L6). This group constituted 5% of taxa of low range of Δ DVM. The composition of taxa in this group
259 was also variable at each station **(Table 1)**, but in general it was represented by Hydrozoa (HY), Salps (SA), Platyhelminthes
260 (PT), Decapoda Larvae (DL), Ostracods (OS), Nauplii (NL) and Ichthyoplankton (IC). These taxa were concentrated mainly
261 in the oxic surface strata (**OX-ML**) and in the **OMZ-UP**, but also in the **OMZ-UC** at the onshore stations (Station T3 and
262 Station L6), showing much less interaction with the **OMZ core**, while spatially moving from normoxia to hypoxia condition
263 and vice versa between 0 to 100 m, and **4) Non-migrants**, represented by taxa which did not exhibit a significant DVM and
264 had a range of Δ DVM from 16 to 0 m (in St. T5), 7 to 0 m (in St. T3) and 6 to 0 m (in St. L6). This group constituted 1% of
265 taxa with not significant Δ DVM. The composition of taxa in this group was also variable at each station **(Table 1)**, but in
266 general it was represented by fish eggs (FE), Radiolarian (RA) and Echinoderm larvae (EL).
267



268

269 **Fig.3.** Weighed mean depth distribution (WMD) of the zooplankton community interacting with the OMZ off Iquique (Stations
 270 T5 and T3) and off Coquimbo (Station L6) at the northern upwelling area of Chile during the austral spring 2015. Shaded gray
 271 areas represent different layers defined by their oxygen levels (defined in Methods). The taxonomic groups were classified by
 272 automated analysis (ZooScan): EC= Eucalanid copepods, AM= amphipods, BR=Bryzoa larvae, AC= *Acartia* Copepods, CT=
 273 Ctenophores, CL=Cirripedia larvae, OS=Ostracods, CH=Chaetognaths, PT= Pteropods, SA=salps, GA= Gastropods, PL=
 274 Platyhelminthes, DL= Decapoda larvae, FO=Foraminifera HY=Hydrozoa, LC=Large Copepods, SIP=Siphonophores,
 275 EU=Euphausiids, FE=fish eggs, NP=Nauplii, SC=Small copepods, AN= Annelids, AP=Appendicularia, RA=Radiolarian,
 276 DC=Decapods, IC=ichthyoplankton, EL=Echinoderm larvae.

278 **Table 1-** Diel vertical migration indices for 27 taxonomic groups (TAXA) identified and sorted by ZooScan at 3 stations off
 279 northern Chile (see **Fig. 1** for acronyms), during the austral spring 2015. Amplitude of migration (Δ DVM) is in meters. Positive
 280 values indicate normal DVM and negative values indicate reverse DVM (see Methods). Four groups are defined in according
 281 to DVM behaviour. Relative abundances are shown in %.

T5			T3			L6			MIGRANTS
TAXA	Δ DVM	%	TAXA	Δ DVM	%	TAXA	Δ DVM	%	
EC.	225	14	EC.	-440	28	EC.	-208	20	STRONG MIGRANTS > 5%
PT.	-188	12	EU.	149	10	AM.	115	11	
EU.	181	11	BR.	-129	8	BR.	-107	10	
AN.	145	9	CH.	-114	7	AC.	-103	10	
FO.	126	8	AN.	105	7	CT.	-94	9	
DC.	90	6	AP.	-88	6	CL.	-87	8	
CH.	88	6	GA.	-84	5	OS.	49	5	
SIP.	73	5	LC.	-70	5	CH.	35	3	INTERMEDIATE MIGRANTS > 2%
AM.	64	4	SC	-70	5	PT.	29	3	
CL.	51	3	NP.	-42	3	SA.	29	3	
DL.	50	3	AM.	42	3	GA.	-27	3	
LC.	47	3	FO	-38	2	PL.	26	2	
HY.	39	2	SA.	32	2	DL.	22	2	
SC	34	2	HY.	-27	2	FO	-22	2	
NP.	24	2	PT.	-23	1	HY.	21	2	WEAK MIGRANTS > 1%
SA.	23	1	SIP.	-15	1	LC.	19	2	
BR.	18	1	CL.	12	1	SIP.	15	1	
IC.	-18	1	OS.	-9	1	EU.	11	1	
FE	49	3	FE	-52	3	FE	25	2	NON- MIGRANTS <1%
OS.	16	1	DL.	-7	0	NP.	6	1	
GA.	15	1	AC.	-	-	SC	-4	0	
AP.	7	0	IC.	-	-	AN.	-4	0	
AC.	-	-	CT.	-	-	AP.	2	0	
CT.	-	-	RA.	-	-	RA.	-	-	
PL.	-	-	DC.	-	-	DC.	-	-	
RA.	-	-	PL.	-	-	IC.	-	-	
EL.	-	-	EL.	-	-	EL.	-	-	

282 **3.2 Vertical distribution and DVM of dominant groups.**

283 Vertical distributions of zooplankton were assessed for 5 taxonomic groups, which represented 80% of total abundance in
284 average: Copepods represented by small Copepods, Large Copepods, Eucalanid copepods and *Acartia* Copepods;
285 Euphausiids; Decapods Larvae; Chaetognaths and Annelids, as well as their patterns of strata–station–abundance relationships
286 are detailed in **Table S4 (Supplemental Material)**. The abundance of these zooplankton groups regarding depth strata was
287 significantly different (ANOSIM, $p < 0.05$) at each station, and therefore representing distinctive microhabitats characterized
288 by specific depth and oxygen concentration. In general, the higher abundance (>80%) was found in the shallower strata and
289 well oxygenated layers (**OX-ML and OMZ UB**) ($>250 \mu\text{mol O}_2 \text{ kg}^{-1}$), and then it decreased rapidly in the strata associated
290 with the OMZ core (**OMZ-UC and OMZ-LC**). Below this stratum a second slight peak in abundance was in the **OMZ-LW**
291 in special at Sts. T5 and L6, occurring between 400-600 m, both daytime and night conditions.

292 As expected, Copepods dominated numerically the zooplankton community both within and outside the OMZ. Small Copepods
293 (SC) were the most abundant (70%), followed by large Copepods (LC) (6%), whereas the Copepods *Acartia* (AC) and
294 eucalanid copepods (EC) showed the lowest abundances among Copepods. The largest aggregation of Copepods (pooled data)
295 altogether during the entire study period was at the offshore station St. T5 (87%), where abundances reached $192088 \text{ ind. m}^{-2}$.
296 At the onshore station (St. T3) the percentage of contribution of copepods was 79% and 69% at the St L6 (**Table S5**
297 **Supplemental Material**). Off Iquique, the highest abundances were in the shallower strata (**OX-ML**) at St.T5 (46%) and at
298 St. T3 (47%), and reduced in the core of the OMZ at St. T5 (4 to 1%) and at T3 (8 to 1%) between 90-400 m, where oxygen
299 had the lowest concentrations ($< 20 \mu\text{M}$ to $1 \mu\text{M}$). At the St. T5 the second peak of abundances was in the **OMZ-LW** stratum
300 during daytime condition, where oxygen levels increased after the extremely low levels within the OMZ, while at the onshore
301 station St T3 it was much dimmer and during night condition. At the onshore station off 29° S (St. L6), having a weaker and
302 less extensive OMZ, the vertical distribution of abundance was similar. However, the abundance of copepods was lowest in
303 this station (in about 69%) in comparison with stations off Iquique, in the core of the OMZ the percentage was between 5 to
304 3%.

305 DVM of Copepods was pronounced at onshore stations (Stations T3 and L6), but the strength of migration was higher overall
306 at St. T3 off Iquique, as reflected by the migration indices (WMD and ΔDVM) (**Table 1**). The WMD of these taxa had a broad
307 range (17–500 m), which varied significantly among copepods groups and stations, both in day and night samples ($p < 0.05$)
308 (**Fig. 3**). During the night, at the offshore station (St. T5) most Copepods exhibited normal DVM and they were concentrated
309 mainly in the oxic surface strata (**OX-M**) and **OMZ-UB** (40-60 m) without interacting with the OMZ; except for eucalanid
310 copepods which concentrated deeper in the **OMZ-LC** stratum, associated with the lower core of the OMZ and showing a high
311 ΔDVM (225m). During the day these four groups of Copepods tended to remain deeper in the stratum associated with the
312 lower core of the OMZ (**OMZ-UC**) and lower O_2 gradient (**OMZ-LW**), except for small Copepods that remained at the **OMZ-**
313 **UB** stratum with a smaller ΔWMD (34 m). At the offshore stations (Stations T3 and L6) the DVM was reverse in most
314 Copepods, except for large Copepods (LC) which showed slightly normal DVM at St. L6 off 29° S . At night Copepods were

315 concentrated deeper in the stratum associated with the lower core of OMZ (**OMZ-UC**) and lower O₂ gradient (**OMZ-LW**),
316 particularly Eucalanidae with a strong DVM and high Δ WMD of 440 m (St. T3) and 208 m (St. L6) and *Acartia* Copepods
317 with 103 m (St. L6) (**Table 1**), whereas at St. L6 small Copepods (SC) were caught in abundance at the **OMZ-UB** stratum
318 down to 82–90 m depth, respectively (**Fig. 3**). During the day, Copepods remained shallower than at night, although they
319 concentrated at different depths. Small Copepods were in the oxic surface strata **OX-ML** (St. T3) and remained in the upper
320 boundary of the OMZ (St. L6) without detectable DVM, as judging by the small difference between their day- and night-time
321 distributions (ADV_M ca. 4 m). Large Copepods (LC), as expected, showed a normal migration and stayed inside the OMZ,
322 concentrated in the **OMZ-UC** stratum (St. L6) and **OMZ-UB** (St. T3). Finally, Eucalanidae, with a strong DVM tended to
323 distribute in the **OMZ-UC** (St. T3) and the **OMZ UB** (St. L6) (**Fig. 3**).

324 Unlike copepods, the Euphausiids were more abundant at the onshore station L6 (<1%), where they reached up to
325 $1683 \pm 473 \text{ ind. m}^{-2} \text{ d}^{-1}$. The OMZ-UB stratum was the most abundant in this station, with a peak of abundance during the
326 daytime, however no DVM was detectable, judging by the small difference between their day- and night-time distributions
327 (**Fig. 3**). Off Iquique, also the highest abundance was in OMZ-UB stratum at night, but with a second peak in OMZ-LC stratum
328 during daytime in both station (Sts. T3 and T5) (**Table S4 Supplemental Material**). The euphausiids appeared to perform a
329 strong DVM in these stations (**Fig. 3**), with a vertical range between 236 and 56 m and a mean Δ DV_M of 181 m at T5 and at
330 St. T3 between 222 and 73 m with a mean Δ DV_M of 149 m (**Table 1**).

331 Decapods larvae were more abundant at StT5 ($428 \pm 132 \text{ ind. m}^{-2} \text{ d}^{-1}$) and were associated with the **OMZ-UB** stratum, where
332 they performed a strong normal DVM with a vertical range between 120 and 30 m and a mean Δ DV_M of 90 m (**Table 1**). At
333 the offshore station (St.T3), the surface peak of abundance was in the **OX-ML** stratum during the day and in the **OMZ-UB**
334 layer at night, where they reached up to $292 \pm 62 \text{ ind. m}^{-2} \text{ d}^{-1}$, with a weak reverse DVM (Δ DV_M-7 m). Off Coquimbo (St. L6)
335 they reached up to $400 \pm 88 \text{ ind. m}^{-2} \text{ d}^{-1}$, the **OMZ-UB** stratum was the most abundant, with a slight second peak in the OMZ-
336 LW stratum during daytime, at this station the vertical range was between 70 and 48 m with a mean Δ DV_M of 22 m (**Fig. 3**).
337 The largest aggregation of Chaetognaths was at the onshore station St. L6 (~2%), where their abundances reached up to
338 $4755 \pm 1038 \text{ ind. m}^{-2} \text{ d}^{-1}$. The abundance and biomass of this group increased in the upper boundary of the OMZ (**OMZ-UB**)
339 during day and night. No DVM was discernible for this group in this station, because of the slight difference between their
340 day- and night-time distributions. By contrast, off Iquique they appeared to perform a strong DVM between the **OMZ-UB** and
341 the **OMZ-UC** strata, as indicated by the migration indices (WMD- Δ DV_M) (**Table 1**). However, at the onshore station (St. T3)
342 they showed a reverse DVM.

343 The other main taxon, Annelida was more abundant at the onshore station St. L6, where their abundances reached up to
344 $7395 \pm 847 \text{ ind. m}^{-2} \text{ d}^{-1}$ (**Table S4 Supplemental Material**). In the whole area, the highest of abundance was in the OMZ-UB,
345 however a second peak of abundance was in the OMZ-LC during daytime at St T5 and T3 and during the night at St. L6. The
346 DVM of this group was high off Iquique with Δ DV_M of 145 m (St. T5) and 105 m (St. T3), while at St. L6 off Coquimbo no
347 DVM was discernible for this group.

348 3.3.3 Others groups with vertical distribution associated to OMZ UC

349 The remaining 19 groups constituted 11% (in St. T5) 17% (in St. T3) and 27% (in St. L6) in abundance. The DVM behaviour
 350 was variable at each station, but in general it was Normal at St. T5 and reverse at Sts.T3-L6 (**Table 1**). These groups clearly
 351 exhibited different daytime and night depths associated with the **OMZ core (OMZ UC-LC)**. Overall, they tended to reside
 352 deeper by day and shallower by night in St. T5 than at the other sites (**Fig.3**).

353 3.4 Vertical distribution of zooplankton biomass

354 Estimates (mean \pm SD) of biomasses of the taxonomic groups integrated by depth strata are summarized in **Table 2**. These
 355 data, averaged from day and night measurements, contrast with the numerical abundances, which were dominated by copepods.
 356 In this case, the bulk of zooplankton biomass was dominated by different groups depending on Stations. In terms of biomass,
 357 Copepods, Euphausiids, Decapods L. Chaetognaths, and Annelids accounted, more or less equally, for >84% in the whole area
 358 (**Table 2**). At the stations less affected by the effect of OMZ the bulk of biomass was dominated by copepods ~50% at St. T5
 359 and ~40% at St. L6, while Decapods largely dominated the bulk of biomass at Station T3 (~40%), followed by Copepods(19%)
 360 and Euphausiids (16 %). (**Table 2**).

361

362 **Table2.** Mean and Standard Deviation (\pm) of integrated biomass (mg C m⁻²) by taxonomic groups identified and sorted by
 363 ZooScan during daytime/ night-time conditions at three stations (T5, T3 and L6) sampled off northern Chile, during the austral
 364 spring 2015. Mean \pm SD are from n=8 for Stations T5 and T3, and n=4 for Station L6.

	T5		T3		L6	
TAXA	Integrated Biomass		Integrated Biomass		Integrated Biomass	
Copepods						
Large C	383.73	\pm 89.3	286.75	\pm 50.6	1727.49	\pm 340.8
Small C	467.55	\pm 115.9	194.51	\pm 41.4	416.14	\pm 66.6
Eucalanid C	0.22	\pm 0.1	2.42	\pm 0.8	11.78	\pm 4.1
Acartia C	0.02	\pm 0.0	0.00	\pm 0.0	0.34	\pm 0.1
Euphausiids	349.48	\pm 77.6	412.38	\pm 103.0	1060.58	\pm 305.8
Decapoda L	20.04	\pm 4.7	1015.61	\pm 294.6	537.17	\pm 176.8
Chaetognaths	63.97	\pm 15.7	215.20	\pm 79.3	641.85	\pm 172.9
Annelida	50.76	\pm 7.4	79.59	\pm 11.8	734.67	\pm 196.4
Decapods	442.38	\pm 187.4	0.00	\pm 0.0	0.00	\pm 0.0
Ostracods	20.83	\pm 2.9	25.14	\pm 1.9	171.01	\pm 15.2
Ctenophores	0.45	\pm 0.2	101.44	\pm 45.4	100.01	\pm 29.9
Hydrozoan	31.48	\pm 9.9	28.39	\pm 8.3	132.34	\pm 39.0
Salps	15.96	\pm 6.0	129.33	\pm 33.5	35.14	\pm 9.9
Siphonophores	71.53	\pm 25.4	55.30	\pm 16.3	39.17	\pm 9.8

Amphipods	8.78	±	1.7	43.32	±	13.2	28.34	±	5.3
Platyhelminthes	0.00	±	0.0	0.00	±	0.0	48.83	±	21.0
Ichthyoplankton	7.62	±	2.2	28.64	±	12.8	0.00	±	0.0
Nauplius L	2.15	±	0.6	1.02	±	0.3	19.78	±	5.6
Pteropods	4.20	±	1.8	0.62	±	0.2	5.02	±	2.2
Foraminiphers	0.45	±	0.1	0.27	±	0.1	7.55	±	0.9
Gastropods	0.35	±	0.1	0.21	±	0.0	1.32	±	0.2
Cirripedia L	0.17	±	0.0	0.09	±	0.0	1.25	±	0.4
Bryozoa L	0.59	±	0.2	0.16	±	0.0	0.70	±	0.3
Radiolarian	0.02	±	0.0	0.05	±	0.0	0.60	±	0.2
Echinoderm L	0.01	±	0.0	0.00	±	0.0	0.04	±	0.0
Appendicularian	0.00	±	0.0	0.00	±	0.0	0.00	±	0.0
Fish Egg	0.00	±	0.0	0.00	±	0.0	0.00	±	0.0
Total	1943	±	435	2620	±	572	5721	±	1019

365

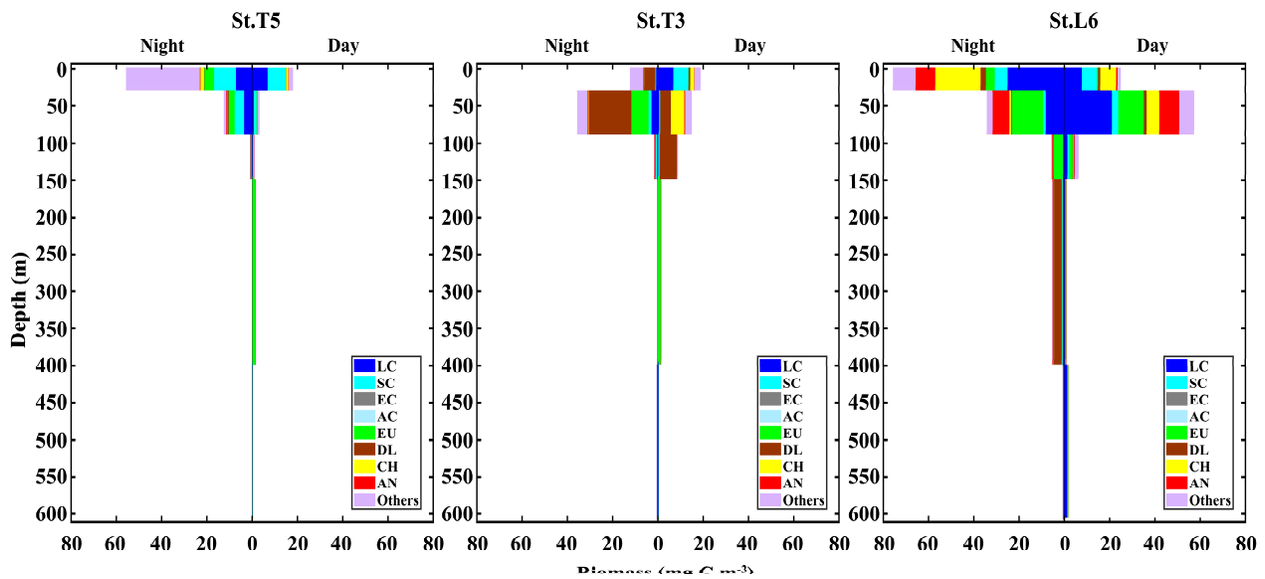
366 When assessing the day vs. night vertical distribution of taxonomic groups in terms of their contribution to biomass, different
367 patterns arise compared to numerical abundance. In this case, we used 9 taxonomic categories to examine vertical distribution
368 and DVM in terms of biomass: small Copepods (SC), large Copepods (LC), *Acartia* Copepods (AC), eucalanid copepods (EC),
369 Euphausiids (EU), Decapod larve (DL), Chaetognaths (CH), Annelids and all the other taxa (**Fig. 4**). Contrasting with
370 numerical abundance, the vertical distribution of biomass was more heterogeneously divided among taxonomic groups and
371 DVM patterns vary strongly between stations. Small Copepods continue to dominate at the St. T5 (24%), with two peaks of
372 biomass, a surface peak associated to the upper oxic layer (**OX-ML**) and OMZ-UB stratum during night condition, and a
373 second peak associated to deeper stratum (OMZ-LW) during daytime. At the onshore Stations T3 and L6 the biomass had a
374 similar vertical distribution but lower (~7%), at the Station T3 the peak of biomass was in the upper oxic layer (**OX-ML**)
375 during daytime condition and then it decreases sharply within the **OMZ-UB** and within the OMZ core (**OMZ-UC** and **OMZ-**
376 **LC**). This abrupt decrease in biomass coincides with the intense OMZ present at this station T3. The second peak of biomass
377 during daytime was in deeper stratum (OMZ-LW), where oxygen conditions seem to be restored. Large Copepods dominate
378 at the onshore St L6(30 %) where their biomass reached up to $1727.49 \pm 340.8 \text{ mg C. m}^{-2} \text{ d}^{-1}$ (**Table 2**). A surface peak of
379 biomass was associated to OMZ-UB stratum during daytime condition, and a second peak associated to deeper stratum (OMZ-
380 LW) also during daytime. Off Iquique they were the second dominant group, with a surface peak in OX-ML stratum during
381 night at St. T5 and daytime at St. T3, and a second peak in deeper stratum (OMZ-LW), during daytime in both stations (**Fig.**
382 **4**).The biomass of Eucalanidae and *Acartia* copepods were lower than the other copepods in the whole area, but in general
383 Eucalanidae was associated to the deeper stratum.

384 Following Copepods, Euphausiids were the second dominant group in term of biomass in the whole area. In general, their
385 ascent from deep layers to the upper ones at night was also evidenced by increasing proportions of these group in the OMZ-
386 UB stratum at night. The highest biomass was in St L6(19%) where reached up to $1060.58 \pm 305.8 \text{ mg C. m}^{-2} \text{ d}^{-1}$ (**Table 2**). A

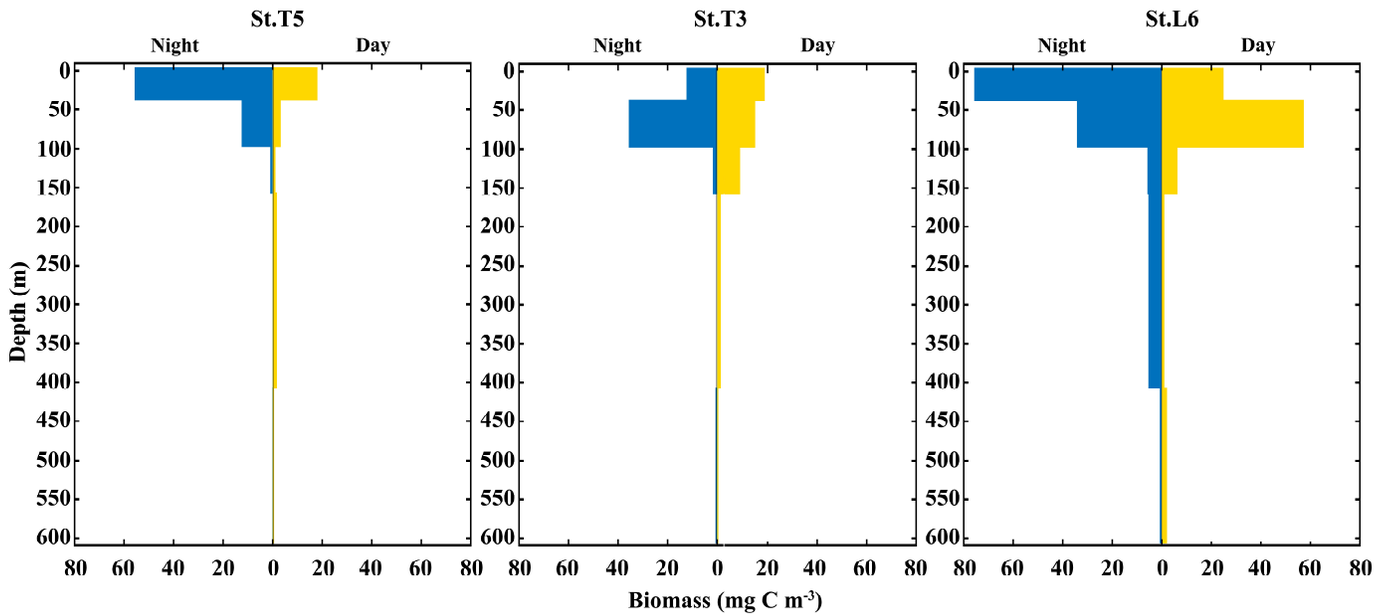
387 surface peak of biomass was associated to OMZ-UB stratum during night condition, decreasing in the deep strata (Fig. 4).
 388 Across of the zonal section off Iquique two peak of biomass was in both stations. A surface peak was in OMZ-UB stratum
 389 during night condition followed by lower biomass within the OMZ core, then a second peak was in the OMZ-LC during
 390 daytime.

391 Decapod Larvae clearly dominated over copepods in the St. T3 (39%). The high biomass was in OMZ-UB stratum during
 392 night condition followed by lower biomass within the OMZ Core. At the St L6 also was a second peak of biomass in the OMZ-
 393 LC during night. Chaetognaths and Annelids were other groups with an important vertical movement of biomass between day
 394 and night across strata, and like other groups with two peaks of biomass. The high biomass was at St. L6 in both groups (Fig.
 395 4).

396 Total added biomass of zooplankton revealed more clearly DVM behaviour of the whole zooplankton community (Fig. 5).
 397 The vertical distribution and daytime vs. Night-time variability of zooplankton biomass showed distinctive features associated
 398 with the OMZ structure, with significant differences ($p < 0.05$) between strata for both daytime and night-time samplings, as
 399 based on the ANOSIM test ($p < 0.05$). In the whole area most of the biomass was concentrated in a narrow band within the OX-
 400 ML and OMZ UB strata, associated with more oxygenated surface waters, with reduced values in deeper waters associated
 401 with the OMZ core, in special at the onshore station off Iquique (St. T3) (Fig.5). Overall, we observed that highest values of
 402 biomass were during the night at the shallower sampling stratum (Ox-ML) and in the subsurface during the day. There was
 403 also an important increase in biomass at the deepest stratum (OMZ-LW) during the daytime and night condition.



404
 405 **Fig.4.** Daytime vs. Night-time vertical distribution of biomass of dominant taxonomic groups at 3 stations off northern Chile:
 406 off Iquique (Stations T5 and T3) and off Coquimbo (Station L6). Data are from night and day replicated samples during two
 407 consecutive days in the austral spring 2015. Values represent means from sampling size $n=4$ for St. T5 and St. T3, and $n=2$ for
 408 St. L6.



409

410 **Fig. 5.** Vertical distribution of total zooplankton biomass during daytime and night-time conditions at 3 stations off northern
 411 Chile: off Iquique (Stations T5 and T3) and off Coquimbo (Station L6) during two consecutive days in the austral spring 2015.).

412 3.4.1 Migrant biomass of the zooplankton taxa

413 The migrant biomass of the zooplankton taxa and the rate of migration (RM), represented by the proportion of biomass (%)
 414 being vertically moved daily from the upper 90 m are shown in **Table 3**. Most dominant groups showed a high rate of migration
 415 as reflected in the RM. In terms of migrant biomass, Decapod larvae, Euphausiids, Decapods, Copepods and Chaetognaths
 416 accounted for a large proportion of total migrant biomass (81%), although high estimates of migrant biomasses were also
 417 associated with high standard deviations, indicating a strong variation among replicated samples (**Table 3**). Presence of zero
 418 values in **Table 3** represents absence of a given taxonomic group in the upper 90 m layer or extremely low values of biomass
 419 under both conditions' day and night, so that such groups did not contribute, or they had a non-significant contribution to total
 420 migrant biomass.

421

422

423

424

425

426

427

428 **Table 3.** Migrant biomass (mg C m⁻²) and rate of migration (RM) (%) for taxonomic groups of zooplankton sampled at northern
 429 Chile at 3 stations: off Iquique (Stations T5 and T3) and off Coquimbo (Station L6) during the austral spring 2015. RM
 430 represents the proportion (%) of migrant biomass with respect to total biomass found at night in the 0-90 m for a given
 431 taxonomic group. SD= Standard deviation of the migrant biomass estimated from n=4 (St. T5 and St. T3) and n=2 (St. L6).
 432

TAXA	ST-T5			ST-T3			ST-L6		
	MB	SD	RM %	MB	SD	RM %	MB	SD	RM %
Decapoda L	24.50	17.32	1	968.95	685.15	40	12.19	8.62	1
Euphausiids	269.50	190.56	11	455.14	321.83	19	247.96	175.34	10
Decapods	876.93	620.08	36	0.00	0.00	0	0.00	0.00	0
Chaetognaths	20.31	14.36	1	390.72	276.28	16	105.39	74.52	4
Copepods									
Small C.	223.31	157.90	9	139.02	98.30	6	128.19	90.64	5
Large C.	183.07	129.45	8	43.78	30.96	2	245.28	173.44	10
<i>Eucalanid C.</i>	0.00	0.00	0	1.10	0.78	0	0.12	0.09	0
<i>Acartia C.</i>	0.04	0.03	0	0.00	0.00	0	0.62	0.44	0
Ctenophores	0.00	0.00	0	202.89	143.46	8	81.61	57.71	3
Annelids	32.50	22.98	1	4.03	2.85	0	156.14	110.41	6
Hydrozoa	35.83	25.34	1	7.80	5.52	0	79.04	55.89	3
Siphonophores	34.89	24.67	1	58.18	41.14	2	0.33	0.23	0
Amphipods	5.62	3.98	0	32.50	22.98	1	32.93	23.29	1
Salps	27.36	19.34	1	38.56	27.27	2	4.57	3.23	0
Ostracods	17.80	12.59	1	12.92	9.14	1	36.37	25.72	2
Platyhelminthes	0.00	0.00	0	0.00	0.00	0	63.08	44.61	3
Pteropods	3.97	2.81	0	0.07	0.05	0	9.85	6.96	0
NaupliusL	3.06	2.17	0	0.36	0.25	0	2.84	2.01	0
Foraminifers	0.09	0.07	0	0.03	0.02	0	3.11	2.20	0
Cirripedia L	0.26	0.18	0	0.01	0.01	0	1.87	1.32	0
Bryozoa L	1.06	0.75	0	0.18	0.13	0	0.83	0.59	0
Gastropods	0.22	0.16	0	0.12	0.09	0	0.11	0.08	0
<i>Echinoderm larvae</i>	0.02	0.01	0	0.00	0.00	0	0.07	0.05	0
Ichthyoplankton	0.06	0.04	0	0.00	0.00	0	0.00	0.00	0
Appendicularia	0.00	0.00	0	0.00	0.00	0	0.00	0.00	0
Fish Eggs	0.00	0.00	0	0.00	0.00	0	0.00	0.00	0
Radiolarian	0.00	0.00	0	0.00	0.00	0	0.00	0.00	0

433

434 4 Discussion

435 Studies on zooplankton DVM and the active transport of C mediated by zooplankton have been documented previously for
 436 the Pacific Ocean and for other areas of the world's ocean, as summarized in **Table** . However, downward C flux due to DVM
 437 in highly productive upwelling regions, such as northern Chile, which is also characterized by severe subsurface hypoxic
 438 conditions upon presence of a shallow OMZ, is still poorly understood. Some studies have shown that hypoxic conditions can
 439 interfere with DMV of many meso- and macrozooplankton species (Wishner *et al.*, 2013; Ekau *et al.*, 2010; Escribano *et al.*,
 440 2009; Apablaza and Palma, 2006; Antezana, 2002; Escribano, 1998). These studies have shown that small differences in
 441 oxygen concentration can make a large difference for zooplankton behavior, physiology and adaptation (Wishner *et al.*, 2018;
 442 Kiko *et al.*, 2016; Seibel, 2011; Gonzalez and Quiñones, 2002; Escribano and McLaren, 1999). Therefore, it seems that the
 443 OMZ can play a very significant role influencing vertical distribution, DVM and ultimately the downward C flux mediated by
 444 zooplankton.

445 Our approach to assess downward C flux into the Oxygen Minimum Zone, based on estimates of the migrant biomass and our
 446 proposed migration indices, allowed us on one hand to examine the contribution that different zooplankton can have for vertical
 447 flux of C and hence export production, and in the other hand to assess zooplankton responses (e.g. vertical distribution and
 448 DVM performance) to changes in environmental conditions over the vertical gradient, such as temperature, water density and
 449 the abrupt changes in oxygenation levels. In this subtropical upwelling region, vertical gradients are much stronger than in
 450 temperate upwelling zones. For example, the coastal zone in this region is more stratified and with a very shallow OMZ (<50
 451 m) with a weak seasonal signal and moderate upwelling throughout the year (Paulmier and Ruiz-Pino, 2009; Fuenzalida *et al.*,
 452 2009; Escribano *et al.*, 2004). This means that zooplankton must cope with hypoxic conditions during their entire life cycle,
 453 except for some species that may reside in near surface water (<30 m), such as *C. chilensis* and *C. brachiatus* which have been
 454 reported as mostly restricted to the upper layer without performing any substantial DVM (Escribano *et al.*, 2012, 2009;
 455 Escribano and Hidalgo, 2000; Escribano, 1998).

456

457 **Table 4.** Comparison of active transport of carbon (AC) ($\text{mg C m}^{-2} \text{d}^{-1}$) by vertically migratory taxa in Pacific Ocean. Diel
 458 vertically migratory taxa (DVM), productivity primary (PP) ($\text{mg C m}^{-2} \text{d}^{-1}$), migrant biomass (MB) (mg C m^{-2}), respiratory
 459 loss (R) ($\text{mg C m}^{-2} \text{d}^{-1}$), faecal pellets production (F) ($\text{mg C m}^{-2} \text{d}^{-1}$) and mortality (M) ($\text{mg C m}^{-2} \text{d}^{-1}$). Where provided by
 460 authors, estimated passive export (POC) is listed. Fluxes refer to carbon export beneath the epipelagic zone (150–200 m depth,
 461 depending on the study) in $\text{mg C m}^{-2} \text{d}^{-1}$

Location	Taxa	PP	MB	AC	R	F	M	% POC	References
N.Hawaii ALOHA	DVM Zooplankton		108–216	7.1	2.6–4.8			12–18	Al-Mutairi and Landry (2001)
N. ALOHA	Hawaii		157.9	3.2–13.6	3.7			18	Steinberg <i>et al.</i> (2008)

N.W. Pac.	DVM Metridia	418	144	9	3	5	1	23.1– 61.8	Kobari <i>et al.</i> (2008)
N.W. Pac.	DVM copepods			8				22.3	Takahashi <i>et al.</i> (2009)
N.E. Pac.	Mesopelagic fishes	170			23.9				Davison <i>et al.</i> (2013)
Eastern Equator			96.0± 25.2		4.2± 1.2		2.9±0. 8	18.4	Zhang and Dam (1997)
Eastern Equator	DVM zooplankton		154.8± 32.4		7.3± 1.4		5.4±1. 1	25.4	Zhang and Dam (1997)
Central Equator (HNLC)			52.9		6		–	4	Rodier and Le Borgne (1997)
Western Equator			46.9		3		–	6	Rodier and Le Borgne (1997)
E. Eq. Pac.	DVM Zooplankton		1214	7.1	7.1			204	Rodier and Le Borgne (1997)
Western Equator	DVM Zooplankton		144–447	23.53– 9.97	7.3–19.1		2.6– 4.4	13–35	Hidaka <i>et al.</i> (2002)
Equator divergence			2.8–21.8		0.9–1.2			<1-2	Roman <i>et al.</i> (2002)
Oligotrophic area			30.2–33.8		1.3–1.7			4	Roman <i>et al.</i> (2002)
E.S.Pac. N.Chile		5503							Gonzalez <i>et al.</i> (1998)
E.S.Pac. N.Chile		10000							Daneri <i>et al.</i> (2000)
E.S.Pac. N.Chile	DVM Eucalanus		8.0 -34	14.1					Hidalgo <i>et al.</i> , (2005)
E.S.Pac. N.Chile	DVM zooplankton		37810	7200		670 0			Escribano <i>et al.</i> (2009)
E.S.Pac. N.Chile	DVM zooplankton	2833±1155	958 ±778	71±64					This study

462 The vertical distribution and diurnal variability of zooplankton biomass seem to be disturbed by the OMZ, such that high
463 biomass aggregates above the oxycline in a narrow band within the OX-ML and OMZ-UB layers, associated with more
464 oxygenated surface waters, whereas extremely low biomass reside in deeper waters, in particular within the OMZ core. This
465 condition was more evident in the coastal station off Iquique (St. T3), characterized by the most intense OMZ in the whole
466 study area. In the eastern tropical north Pacific, biomass distribution seemed different, exhibiting a secondary peak at depth
467 during the daytime within the upper oxycline or OMZ core (Wishner *et al.*, 2013).

468 Regarding the estimates of biomass for each of the taxonomic groups, our approaches can certainly introduce variation,
469 depending on selected regressions and conversion factors from highly diverse body shapes and body densities of the
470 zooplankton taxa affecting the estimates of body area and volume, dry weight and C content. Various approaches have been
471 adopted for converting sizes to body masses. For example, Lehette and Hernández-León (2009) provided some general
472 regression equations for subtropical and Antarctic zooplankton describing the relationship between scanned area and body
473 mass (C content). These authors also proposed two separate regressions for crustacean and gelatinous zooplankton, because of
474 different body densities. In our study, we adopted more direct estimates of body masses by converting individual areas or
475 volumes (from ZooScan) using published regressions for separate taxonomic groups. Also, in our samples there was a high

476 diversity of taxonomic groups as identified by ZooScan, such that unique regressions for crustacean and gelatinous organisms
477 may lead to strong biases in body mass estimates, because of high variability in C content, which is the key component of body
478 mass needed to estimate C flux. Therefore, the use of taxa-specific conversion factors, as those detailed in our **Table S3**
479 **(Supplemental Material)** is strongly recommended.

480 Despite the apparently hostile oxygen-deficient habitat, associated with the OMZ, we found that most taxa were able to perform
481 DVM in the upwelling zone withstanding severe hypoxia. Even, several zooplankton groups are strong migrants, exhibiting
482 large DVM amplitude (~500 m). Among them, an important migrant group is comprised by the eucalanid copepods which
483 have been described as even being able to enter the core of the OMZ, and then migrate downward to the lower limit of the
484 OMZ, which is slightly more oxygenated (Hidalgo *et al.*, 2005). In our study however, their contribution to total migrant
485 biomass was too small (ca. 0.4 mg C m⁻² d⁻¹), as compared to the estimate made by Hidalgo *et al.*(2005). In fact, the migrant
486 biomass and rate of migration of this group was non-significant when considering DVM between the upper 90 m and below,
487 suggesting a little or no contribution to downward flux of C for this group of copepods. However it seems that eucalanid
488 copepods remain below the oxycline or nearby the base of the oxycline day and night, as shown by their weighted mean depth
489 (WMD) and therefore suggesting that they may still contribute to vertical flux by feeding at the base of the oxycline at night
490 and then migrating into the OMZ during the day.

491 Other taxa, such as Euphausiids, *Acartia* spp., other copepods, Ctenophores, Decapods, Annelidae, Bryozoa L, Pteropods and
492 Chaetognaths tended to concentrate their populations inside the OMZ core showing a strong link to the OMZ with important
493 movement throughout the water column. Antezana (2010) showed that *E. mucronata*, an endemic and abundant euphausiid in
494 the coastal upwelling zone off Chile, is a well-adapted species to vertically migrate into the core of the OMZ. In fact, the
495 euphausiids studied here showed a large DVM amplitude (~250 m), descending into the core of the OMZ and below 250 m
496 each day. In general, all strong migrants' taxa showed a strong interaction with the core of OMZ, remaining there either
497 temporarily or permanently during the day or night condition, contributing in this way to the release of C below the
498 thermocline, despite presence of hypoxic conditions.

499
500 Our estimates of DVM-mediated C flux showed that migrant biomass (958±778 mg C m⁻² d⁻¹) and C flux estimates (71±64mg
501 C m⁻² d⁻¹) of the major taxa performing DVM, were greater than those reported for the Pacific Ocean, both in oligotrophic,
502 such as Hawaii, and mesotrophic waters, as the subarctic North Pacific(Steinberg *et al.*, 2008), and even greater than that
503 informed by Yebra *et al.* (2005) within eddies with enhanced biological production. Most of these previous estimates however
504 have not been done in regions with severe hypoxia or anoxia at mid water depths (e.g. Kiko *et al.*, 2016), such as the highly
505 productive upwelling region of the coastal zone off northern of Chile, where the oxygen concentrations may fall below <1
506 μmol in the core of OMZ (Paulmier and Ruiz-Pino, 2009). Moreover, only few works have considered the whole zooplankton
507 community (**Table 4**). High productivity and strong aggregation of zooplankton in coastal areas of this region (Escribano *et*
508 *al.*, 2000; Escribano and Hidalgo, 2000) may promote greater amounts of migrant biomass. This requires however that DVM
509 should not be majorly constrained by presence of the OMZ and that most migrant taxa are tolerant to low-oxygen. On the other

510 hand, our estimates of downward C flux were substantially lower than previous ones reported off northern Chile by Hidalgo
511 *et al.* (2005) for *Eucalanus inermis* alone (14.1 mg C m⁻² d⁻¹). Although, such previous estimates may be too high, considering
512 the level of primary production in the upwelling zone of Chile (~10000 mg C m⁻² d⁻¹, the maximum estimated value) (Daneri
513 *et al.*, 2000). It should be noted that potential contribution to C at depth by faecal pellet production (egestion) was not
514 considered in our estimate of active transport. The lack of an estimate of ingestion rates at the upper layer (nominally 0-90 m)
515 precludes us to make reliable calculations of egestion at depth. We also consider that in situ production of faecal pellets at
516 depth (below the thermocline) and its actual contribution to active transport of C need further study and it should be estimated
517 for particular feeding conditions.

518 Differences of our estimates with previous works may also be accounted by strong variability of zooplankton abundance in
519 the upwelling zone. In fact, our estimates of migrant biomasses of the different taxonomic groups based on 2 days of sampling
520 and two replicates for each condition (day and night) are strongly variables, as shown by the standard errors in **Table 3** which
521 can be as much as 100% from the mean value. Therefore, comparisons must take caution upon strong time-space variation
522 when assessing zooplankton abundance. Nevertheless, a strong spatial variation in migrant biomass was also evident when
523 comparing the three sampling stations. For instance, St. L6 had more biomass than the other stations, but much less migrant
524 biomass in the upper layer (**Table 5**), and thus a very low contribution to vertical flux of C by DVM. At station L6, large
525 copepods, Euphausiids, Annelids and Chaetognaths largely contributed to biomass, although they did not show significant
526 DVM. Therefore, species composition and their DVM behaviour appear as a key factor to determine the downward flux of C
527 mediated by active transport. Even although the OMZ did not greatly prevent DVM migration, zooplankton behaviour
528 appeared disrupted or exhibited reversed patterns, depending on vertical distribution of OMZ and on the taxonomic group
529 being considered. This behaviour was more evident in the onshore stations (Stations T3 and L6), but in particular in the station
530 off Iquique (St. T3) that also showed a higher migration rate (60%). According to Ekau *et al.* (2010), other indirect effects
531 could also be caused by the hypoxia conditions, such as changes in prey availability, prey size or predation risk, as well as
532 changes in species composition, the strength of which depends on the duration and intensity of the hypoxic events. This could
533 explain why individuals within a single population can perform reverse, normal, or non DVM, apparently depending on the
534 more important source of mortality: predation by nocturnal feeding, normally migrating carnivorous zooplankton, or visually
535 hunting planktivorous fish (Ohman, 1990). These kind of DVM behaviors can only be better assessed and understood when
536 looking at the population level, although again time-space variation in zooplankton abundance in a highly heterogeneous
537 upwelling zone should be kept in mind. Important to consider that our automated analysis of the zooplankton community may
538 not account for differences in species composition between stations or strata, and therefore changing taxa within assigned
539 groups between stations, such as strong migrants, or non-migrants may obey to variable species compositions. although the
540 possibility that same populations change their DVM performance depending on changing environmental conditions cannot be
541 discarded, in particular referring to vertical distribution of oxygen. Such effects may provide explanations to observed variation
542 in migrant biomass between stations, but also between strata. In fact, we noted strong differences in estimates of migrant
543 biomass when comparing the upper 0-90 m stratum and the deeper 90-600 m stratum (**Table 5**, also by taxa in **Table S7**

544 **(Supplemental Material)**). Furthermore, sampling biases should also be considered, especially when using a vertically towed
545 multinet which may not properly sample large-sized zooplankton at daytime condition in the 0-90 m, because of net avoidance,
546 so that introducing a source of variation when comparing surface vs deeper layers under daytime and nighttime conditions.
547 Concerning C fluxes, our estimates of active transport of carbon by zooplankton were about half the estimates of passive C
548 sinking obtained off northern Chile at 60 m depth off Antofagasta (23°S) by Gonzalez *et al.*(1998) based on sediment traps
549 (125 to 176 mg C m⁻² d⁻¹). Regarding the question on how efficient is active C transport mediated by DVM, we obtained
550 satellite-based (<http://science.oregonstate.edu.ocean.productivity>) estimates of net primary production (monthly means for
551 November-December 2015) for the coastal area (Stations T3 and L6) and the coastal transition zone (Station T5), averaged for
552 the months of November and December 2015. Our estimates of downward C flux represented a mean of ca. 4 % of export of
553 Carbon resulting from net primary production in the upwelling region, estimated in the range of 1500-3500 mg C m⁻² d⁻¹
554 **(Table 5)**. If we consider this is accounted only by mesozooplankton, then an important fraction of freshly produced C might
555 be taken downward by zooplankton, and this DVM-mediated C flux ought to be taking into account when analysing and
556 modelling the C budget in the upwelling zone.
557

558 **Table 5.** Mean net primary production rate and estimates of daily downward C flux due to passive sinking and mediated by
559 diel vertical migration (DVM) of mesozooplankton at three stations (T5, T3 and L6) in the coastal upwelling region off
560 northern Chile during the austral spring 2015. Primary production represents satellite-based estimates of monthly mean
561 (November–December 2015) at the 3 sampling stations. Passive C flux is a mean value estimated from sediment traps by
562 González et al. (2000) off Antofagasta (northern Chile, 23 °S) for January 1997. Total biomass and epipelagic biomass are
563 mean observed values from day-night conditions after 2 consecutive days of sampling.

564

Station	T5	T3	L6	Mean ± SD	
Primary Production (C m ⁻² d ⁻¹)	1500	3500	3500	2833	± 1155
Passive Carbon Flux (C m ⁻² d ⁻¹)				151	± 36
Integrated Abundance ind. m ⁻² (0–600 m)	221735	127085	371235	240018	± 123097
S Total biomass mg (C m ⁻² d ⁻¹) (0–600m)	1943	2620	5721	3428	± 2015
Epipelagic biomass (mg C m⁻² d⁻¹) (0–90m)	1581	1988	4259	2609	± 1443
Migrant biomass mg (C m ⁻² d ⁻¹) (0–90m)	1686	1048	139	957.7	± 778
Migrant biomass mg (C m ⁻² d ⁻¹) (90–600m)	503	756	810	689.9	± 164
Rate of Biomass migration (%)	70	42	3	38	± 34
Active Carbon Flux mg (C m ⁻² d ⁻¹)	139	63	11	71.13	± 64
Total Active Carbon exported (%)	9.3	1.8	0.3	3.8	± 4.8

565

566 **5. Conclusions**

567 In the coastal upwelling zone off northern Chile the presence of a subsurface oxygen minimum zone (OMZ) can impose an
568 important constraint for diel vertical migration of zooplankton and so influencing the downward C export mediated by
569 zooplankton. We found that most of the zooplankton biomass aggregates above the oxycline, associated with more oxygenated
570 surface waters and this was evident upon presence of a more intense OMZ. Some taxonomic groups, however, were found
571 closely associated with the OMZ and several taxa were able to perform DVM in the upwelling zone withstanding severe
572 hypoxia. Also strong migrants, such as large sized copepods and copepod of the group Eucalanidae and Euphausiids, can
573 exhibit a large migration amplitude (~500 m), remaining either temporarily or permanently during the day or night condition
574 within the core of the OMZ, and so contributing to the release of C below the oxycline (and thermocline). Our estimates of
575 DVM-mediated C flux suggested that a mean migrant biomass of 957.7 mg C m⁻² d⁻¹ may contribute with about 71.1 mg C m⁻²
576 d⁻¹ to the OMZ system through respiration, mortality, and C excretion at depth, accounting for ca, 4% of the net primary
577 production, and so implying the existence of a efficient mechanism to incorporate freshly produced C into the OMZ. This

578 downward C flux mediated by zooplankton DVM is however strongly depending on the taxonomic structure due to variable
579 migration amplitude and DVM behavior. These estimates should also consider the strong temporal-spatial variation in
580 zooplankton abundance in the upwelling zone for comparison purposes.

581

582 **Acknowledgements**

583 This work has been funded the Millennium Institute of Oceanography (IMO) (Grant IC 120019) and the CONICYT Project
584 Grant No PCHA 21160038 through which the LowpHox I cruise was conducted. We are thankful to two anonymous reviewers
585 who greatly contributed to improve the work. We are also grateful to D. Toledo for assistance during field work. The work is
586 a contribution to IMBeR Program and SCOR EBUS WG 155.

587 **References**

- 588 Al-Mutairi, H. and Landry, M. R.: Active export of carbon and nitrogen at station ALOHA by diel migrant zooplankton, *Deep.*
589 *Res. Part II Top. Stud. Oceanogr.*, 48(8–9), 2083–2103, doi:10.1016/S0967-0645(00)00174-0, 2001.
- 590 Andersen, V., Devey, C., Gubanova, A., Picheral, M., Melnikov, V., Tsarin, S. and Prieur, L.: Vertical distributions of
591 zooplankton across the Almeria-Oran frontal zone (Mediterranean Sea), *J. Plankton Res.*, 26(3), 275–293,
592 doi:10.1093/plankt/fbh036, 2004.
- 593 Antezana, T.: Vertical distribution and diel migration of *Euphausia mucronata* in the oxygen minimum layer of the Humboldt
594 Current, *Oceanogr. East. Pacific II*, 13–28, 2002.
- 595 Antezana, T.: Species-specific patterns of diel migration into the Oxygen Minimum Zone by euphausiids in the Humboldt
596 Current Ecosystem, *Prog. Oceanogr.*, 83(1–4), 228–236, doi:10.1016/j.pocean.2009.07.039, 2009.
- 597 Antezana, T.: *Euphausia mucronata*: A keystone herbivore and prey of the Humboldt Current System, *Deep. Res. Part II Top.*
598 *Stud. Oceanogr.*, 57(7–8), 652–662, doi:10.1016/j.dsr2.2009.10.014, 2010.
- 599 Apablaza, P. and Palma, S.: Efecto de la zona de mínimo oxígeno sobre la migración vertical de zooplancton gelatinoso en la
600 bahía de Mejillones, *Investig. Mar.*, 34(2), 81–95, doi:10.4067/s0717-71782006000200009, 2006.
- 601 Bray, J. R. and Curtis, J. T.: An ordination of the upland forest communities of southern Wisconsin, *Ecol. Monogr.*, 27(4),
602 325–349, 1957.
- 603 Breitburg, D., A Levin, L., Oschlies, A., Grégoire, M., P Chavez, F., J Conley, D., Garcon, V., Gilbert, D., Gutiérrez, D.,
604 Isensee, K., S Jacinto, G., Limburg, K., Montes, I., Naqvi, S. W. A., Pitcher, G., Rabalais, N., R Roman, M., Rose, K., Seibel,
605 B. and Zhang, J.: Declining oxygen in the global ocean and coastal waters, *Science*, 359(6371), doi:10.1126/science.aam7240,
606 2018.
- 607 Chang, C. Y., Ho, P. C., Sastri, A. R., Lee, Y. C., Gong, G. C. and Hsieh, C. hao: Methods of training set construction: Towards
608 improving performance for automated mesozooplankton image classification systems, *Cont. Shelf Res.*, 36, 19–28,
609 doi:10.1016/j.csr.2012.01.005, 2012.

610 Childress, J. J. and Nygaard, M. H.: The chemical composition of midwater fishes as a function of depth of occurrence off
611 Southern California, *Deep. Res.*, 20, 1091–1093, 1973.

612 Dai, L., Li, C., Yang, G. and Sun, X.: Zooplankton abundance, biovolume and size spectra at western boundary currents in the
613 subtropical North Pacific during winter 2012, *J. Mar. Syst.*, 155, 73–83, doi:10.1016/j.jmarsys.2015.11.004, 2016.

614 Daneri, G., Dellarossa, V., Quiñones, R., Jacob, B., Montero, P. and Ulloa, O.: Primary production and community respiration
615 in the Humboldt Current System off Chile and associated oceanic areas, *Mar. Ecol. Prog. Ser.*, 197, 41–49,
616 doi:10.3354/meps197041, 2000.

617 Davison, P. C., Checkley, D. M., Koslow, J. A. and Barlow, J.: Carbon export mediated by mesopelagic fishes in the northeast
618 Pacific Ocean, *Prog. Oceanogr.*, 116, 14–30, doi:10.1016/j.pocean.2013.05.013, 2013.

619 Donoso, K. and Escribano, R.: Mass-specific respiration of mesozooplankton and its role in the maintenance of an oxygen-
620 deficient ecological barrier (BEDOX) in the upwelling zone off Chile upon presence of a shallow oxygen minimum zone, *J.*
621 *Mar. Syst.*, 129, 166–177, 2014.

622 Ducklow, H. W., Eborah, D., Steinberg, K. and Buesseler, K. O.: Upper ocean carbon export and the biological pump,
623 *Oceanography*, 14(4), 50–58, doi:10.5670/oceanog.2001.06, 2001.

624 Edvardsen, A., Zhou, M., Tande, K. S. and Zhu, Y.: Zooplankton population dynamics: measuring in situ growth and mortality
625 rates using an Optical Plankton Counter, *Mar. Ecol. Prog. Ser.*, 227, 205–219, 2002.

626 Ekau, W., Auel, H., Pörtner, H. O. and Gilbert, D.: Impacts of hypoxia on the structure and processes in pelagic communities
627 (zooplankton, macro-invertebrates and fish), *Biogeosciences*, 7(5), 1669–1699, doi:10.5194/bg-7-1669-2010, 2010.

628 Escribano, R.: Population dynamics of *Calanus chilensis* in the Chilean Eastern Boundary Humboldt Current, *Fish. Oceanogr.*,
629 7(3–4), 245–251, doi:10.1046/j.1365-2419.1998.00078.x, 1998.

630 Escribano, R.: Zooplankton interactions with the oxygen minimum zone in the eastern South Pacific, *Supl. Gayana*, 70, 19–
631 21, 2006.

632 Escribano, R. and McLaren, I.: Production of *Calanus chilensis* in the upwelling area of Antofagasta, Northern Chile, *Mar.*
633 *Ecol. Prog. Ser.*, 177, 147–156, doi:10.3354/meps177147, 1999.

634 Escribano, R. and Hidalgo, P.: Spatial distribution of copepods in the north of the Humboldt Current region off Chile during
635 coastal upwelling, *J. Mar. Biol. Assoc. United Kingdom*, 80(2), 283–290, 2000.

636 Escribano, R., Iribarren, C. and Marín, V.: Distribution of *Euphausia mucronata*; at the upwelling area of Peninsula Mejillones,
637 northern Chile: the influence of the oxygen minimum layer, *Sci. Mar. ISSN 0214-8358*, Vol. 64, N°. 1, 2000, págs. 69-77,
638 64(1), 69–77, doi:10.3989/scimar.2000.64n169, 2000.

639 Escribano, R., Daneri, G., Farías, L., Gallardo, V. A., González, H. E., Gutiérrez, D., Lange, C. B., Morales, C. E., Pizarro,
640 O., Ulloa, O. and Braun, M.: Biological and chemical consequences of the 1997-1998 El Niño in the Chilean coastal upwelling
641 system: A synthesis, *Deep. Res. Part II Top. Stud. Oceanogr.*, 51(20–21), 2389–2411, doi:10.1016/j.dsr2.2004.08.011, 2004.

642 Escribano, R., Hidalgo, P. and Krautz, C.: Zooplankton associated with the oxygen minimum zone system in the northern
643 upwelling region of Chile during March 2000, *Deep Sea Res. Part II Top. Stud. Oceanogr.*, 56(16), 1083–1094, 2009.

644 Escribano, R., Hidalgo, P., Fuentes, M. and Donoso, K.: Zooplankton time series in the coastal zone off Chile: Variation in
645 upwelling and responses of the copepod community, *Prog. Oceanogr.*, 97–100, 174–186, doi:10.1016/j.pocean.2011.11.006,
646 2012.

647 Fernández-Álamo, M. A. and Färber-Lorda, J.: Zooplankton and the oceanography of the eastern tropical Pacific: a review,
648 *Prog. Oceanogr.*, 69(2–4), 318–359, 2006.

649 Fuenzalida, R., Schneider, W., Garcés-Vargas, J., Bravo, L. and Lange, C. B.: Vertical and horizontal extension of the oxygen
650 minimum zone in the eastern South Pacific Ocean, *Deep Sea Res. Part II Top. Stud. Oceanogr.*, 56(16), 992–1003,
651 doi:10.1016/j.dsr2.2008.11.001, 2009.

652 Gonzalez, H. E., Daneri, G., Figueroa, D., Iriarte, J. L., Lefevre, N., Pizarro, G. A., Quiñones, R., Sobarzo, M. and Troncoso,
653 A.: Producción primaria y su destino en la trama trófica pelágica y océano-atmósfera de CO₂ en la zona norte de la Corriente
654 de Humboldt (23°S): Posibles efectos del evento El Niño, 1997-98 en Chile, *Rev. Chil. Hist. Nat.*, 71, 429–458, 1998.

655 Gonzalez, R. R. and Quiñones, R. A.: Ldh activity in *Euphausia mucronata* and *Calanus chilensis*: Implications for vertical
656 migration behaviour, *J. Plankton Res.*, 24(12), 1349–1356, doi:10.1093/plankt/24.12.1349, 2002.

657 Gorsky, G., Ohman, M. D., Picheral, M., Gasparini, Stephane Stemann, L., Romagnan, Jean-Baptiste Cawood, A., Pesant,
658 S., García-Comas, C. and Prejger, F.: Digital zooplankton image analysis using the ZooScan integrated system, *J. Plankton
659 Res.*, 32(3), 285–303, doi:10.1093/plankt/fbp124, 2010.

660 Grantham, B. A., Chan, F., Nielsen, K. J., Fox, D. S., Barth, J. A., Huyer, A., Lubchenco, J. and Menge, B. A.: Upwelling-
661 driven nearshore hypoxia signals ecosystem and oceanographic changes in the northeast Pacific, *Nature*, 429(6993), 749–754,
662 doi:10.1038/nature02605, 2004.

663 Hidaka, K., Kawaguchi, K., Murakami, M. and Takahashi, M.: Downward transport of organic carbon by diel migratory
664 micronekton in the western equatorial Pacific:: its quantitative and qualitative importance, *Deep Sea Res. Part I Oceanogr.
665 Res. Pap.*, 48(8), 1923–1939, doi:10.1016/s0967-0637(01)00003-6, 2002.

666 Hidalgo, P., Escribano, R. and Morales, C. E.: Ontogenetic vertical distribution and diel migration of the copepod *Eucalanus
667 inermis* in the oxygen minimum zone off northern Chile (20-21° S), *J. Plankton Res.*, 27(6), 519–529,
668 doi:10.1093/plankt/fbi025, 2005.

669 Ikeda, T.: Metabolic rates of epipelagic marine zooplankton as a function of body mass and temperature, *Mar. Biol.*, 85, 1–11,
670 doi:10.1007/BF00396409, 1985.

671 Ikeda, T.: Respiration and ammonia excretion by marine metazooplankton taxa: synthesis toward a global-bathymetric model,
672 *Mar. Biol.*, 161(12), 2753–2766, 2014.

673 Judkins, D. C.: Vertical distribution of zooplankton in relation to the oxygen minimum off Peru, *Deep Sea Res. Part A,
674 Oceanogr. Res. Pap.*, 27(6), 475–487, doi:10.1016/0198-0149(80)90057-6, 1980.

675 Kiko, R., Hauss, H., Buchholz, F. and Melzner, F.: Ammonium excretion and oxygen respiration of tropical copepods and
676 euphausiids exposed to oxygen minimum zone conditions, *Biogeosciences (BG)*, 13(8), 2241–2255, 2016.

677 Kiørboe, T.: Zooplankton body composition, *Limnol. Oceanogr.*, 58(5), 1843–1850, 2013.

678 Kobari, T., Steinberg, D., Ueda, A., Tsuda, A., Silver, M. W. and Kitamura, M.: Impacts of ontogenetically migrating copepods
679 on downward carbon flux in the western subarctic Pacific Ocean.pdf, *Deep Sea Res. Part II Top. Stud. Oceanogr.*, 55, 1648–
680 1660 [online] Available from: doi: 10.1016/j.dsr2.2008.04.016, 2008.

681 Lehette, P. and Hernández-León, S.: Zooplankton biomass estimation from digitized images: a comparison between subtropical
682 and Antarctic organisms, *Limnol. Oceanogr. Methods*, 7(4), 304–308, doi:10.4319/lom.2009.7.304, 2009.

683 Longhurst, A. and Williams, R.: Carbon flux by seasonally migrating copepods is a small number, *J. Plankton Res.*, 14(11),
684 1495–1509, 1992.

685 Morales, C. E., Braun, M., Reyes, H., Blanco, J. L. and Davies, A. G.: Anchovy larval distribution in the coastal zone off
686 northern Chile: the effect of low dissolved oxygen concentrations and of a cold-warm sequence (1990-95), *Invest. Mar.*,
687 Valparaíso, 24, 77–96, doi:10.4067/s0717-71781996002400007, 1996a.

688 Morales, C. E., Davies, A. G., Braun, M., Reyes, H. and Blanco, J. L.: Anchovy larval distribution in the coastal zone off
689 northern Chile: the effect of low dissolved oxygen concentrations and of a cold-warm sequence (1990-95), *Investig. Mar.*, 24,
690 77–96, doi:10.4067/s0717-71781996002400007, 1996b.

691 Morales, C. E., Hormazábal, S. E. and Blanco, J. L. J.: Interannual variability in the mesoscale distribution of the depth of the
692 upper boundary of the oxygen minimum layer off northern Chile (18–24S): Implications for the pelagic system and
693 biogeochemical cycling, *J. Mar. Res.*, 57(6), 909–932, 1999.

694 Ohman, M. D.: The demographic benefits of diel vertical migration by zooplankton, *Ecol. Monogr.*, 60(3), 257–281, 1990.

695 Paulmier, A. and Ruiz-Pino, D.: Oxygen minimum zones (OMZs) in the modern ocean, *Prog. Oceanogr.*, 80(3–4), 113–128,
696 doi:10.1016/j.pocean.2008.08.001, 2009.

697 Paulmier, A., Ruiz-Pino, D., Garçon, V. and Fariás, L.: Maintaining of the Eastern South Pacific Oxygen Minimum Zone
698 (OMZ) off Chile, *Geophys. Res. Lett.*, 33(20), 2–7, doi:10.1029/2006GL026801, 2006.

699 Putzeys, S., Yebra, L., Almeida, C., Bécognée, P. and Hernández-León, S.: Influence of the late winter bloom on migrant
700 zooplankton metabolism and its implications on export fluxes, *J. Mar. Syst.*, 88(4), 553–562,
701 doi:10.1016/j.jmarsys.2011.07.005, 2011.

702 Rodier, M. and Le Borgne, R.: Export flux of particles at the equator in the western and central Pacific ocean, *Deep. Res. Part*
703 *II Top. Stud. Oceanogr.*, 44(9–10), 2085–2113, doi:10.1016/S0967-0645(97)00092-1, 1997.

704 Roman, M. R., Adolf, H. A., Landry, M. R., Madin, L. P., Steinberg, D. K. and Zhang, X.: Estimates of oceanic
705 mesozooplankton production: A comparison using the Bermuda and Hawaii time-series data, *Deep. Res. Part II Top. Stud.*
706 *Oceanogr.*, 49(1–3), 175–192, doi:10.1016/S0967-0645(01)00099-6, 2002.

707 Seibel, B. A.: Critical oxygen levels and metabolic suppression in oceanic oxygen minimum zones, *J. Exp. Biol.*, 214(2), 326–
708 336, doi:10.1242/jeb.049171, 2011.

709 Seibel, B. A., Schneider, J. L., Kaartvedt, S., Wishner, K. F. and Daly, K. L.: Hypoxia Tolerance and Metabolic Suppression
710 in Oxygen Minimum Zone Euphausiids: Implications for Ocean Deoxygenation and Biogeochemical Cycles, *Integr. Comp.*
711 *Biol.*, 56(4), 510–523, doi:10.1093/icb/icw091, 2016.

712 Silva, N. and Neshyba, S.: On the southernmost extension of the Peru-Chile undercurrent, *Deep Sea Res. Part A, Oceanogr.*
713 *Res. Pap.*, 26(12), 1387–1393, doi:10.1016/0198-0149(79)90006-2, 1979.

714 Steinberg, D. K. and Landry, M. R.: Zooplankton and the Ocean Carbon Cycle, *Ann. Rev. Mar. Sci.*, 9(1), 413–444,
715 doi:10.1146/annurev-marine-010814-015924, 2017.

716 Steinberg, D. K., Carlson, C. A., Bates, N. R., Goldthwait, S. A., Madin, L. P. and Michaels, A. F.: Zooplankton vertical
717 migration and the active transport of dissolved organic and inorganic carbon in the Sargasso Sea, *Deep. Res. Part I, Oceanogr.*
718 *Res. Pap.*, 47(1), 137, doi:10.1016/S0967-0637(99)00052-7, 2000.

719 Steinberg, D. K., Cope, J. S., Wilson, S. E. and Kobari, T.: A comparison of mesopelagic mesozooplankton community
720 structure in the subtropical and subarctic North Pacific Ocean, *Deep. Res. Part II Top. Stud. Oceanogr.*, 55(14–15), 1615–
721 1635, doi:10.1016/j.dsr2.2008.04.025, 2008.

722 Stramma, L., Johnson, G., Sprintall, J. and Mohrholz, V.: Expanding Oxygen-Minimum in the Tropical Oceans, *Science* (80-
723 .), 320, 655–658, 2008.

724 Takahashi, K., Kuwata, A., Sugisaki, H., Uchikawa, K. and Saito, H.: Downward carbon transport by diel vertical migration
725 of the copepods *Metridia pacifica* and *Metridia okhotensis* in the Oyashio region of the western subarctic Pacific Ocean, *Deep.*
726 *Res. Part I Oceanogr. Res. Pap.*, 56(10), 1777–1791, doi:10.1016/j.dsr.2009.05.006, 2009.

727 Ulloa, O. and Pantoja, S.: The oxygen minimum zone of the eastern South Pacific, *Deep. Res. Part II Top. Stud. Oceanogr.*,
728 56(16), 987–991, doi:10.1016/j.dsr2.2008.12.004, 2009.

729 Ulloa, O., Canfield, D. E., DeLong, E. F., Letelier, R. M. and Stewart, F. J.: Microbial oceanography of anoxic oxygen
730 minimum zones, *Proc. Natl. Acad. Sci.*, 109(40), 15996–16003, doi:10.1073/pnas.1205009109, 2012.

731 Wishner, K. F., Gelfman, C., Gowing, M. M., Outram, D. M., Rapien, M. and Williams, R. L.: Vertical zonation and
732 distributions of calanoid copepods through the lower oxycline of the Arabian Sea oxygen minimum zone, *Prog. Oceanogr.*,
733 78(2), 163–191, doi:10.1016/j.pocean.2008.03.001, 2008.

734 Wishner, K. F., Outram, D. M., Seibel, B. A., Daly, K. L. and Williams, R. L.: Zooplankton in the eastern tropical north
735 Pacific: Boundary effects of oxygen minimum zone expansion, *Deep. Res. Part I Oceanogr. Res. Pap.*, 79, 122–140,
736 doi:10.1016/j.dsr.2013.05.012, 2013.

737 Wishner, K. F., Seibel, B. A., Roman, C., Deutsch, C., Outram, D., Shaw, C. T., Birk, M. A., Mislán, K. A. S., Adams, T. J.,
738 Moore, D. and Riley, S.: Ocean deoxygenation and zooplankton: Very small oxygen differences matter, *Sci. Adv.*, 4(12),
739 eaau5180, doi:10.1126/sciadv.aau5180, 2018.

740 Yebra, L., Almeida, C. and Hernández-León, S.: Vertical distribution of zooplankton and active flux across an anticyclonic
741 eddy in the Canary Island waters, *Deep. Res. Part I Oceanogr. Res. Pap.*, 52(1), 69–83, doi:10.1016/j.dsr.2004.08.010, 2005.

742 Zhang, X. and Dam, H. G.: Downward export of carbon by diel migrant mesozooplankton in the central equatorial Pacific,
743 *Deep Sea Res. Part II Top. Stud. Oceanogr.*, 44, 2191–2202 [online] Available from: [https://ac.els-](https://ac.els-cdn.com/S096706459700060X/1-s2.0-S096706459700060X-main.pdf?_tid=d3d3eae5-5ca5-4ac5-ba8e-6e934a548139&acdnat=1535698698_4634563185704240d6eef101ba42f036)
744 [cdn.com/S096706459700060X/1-s2.0-S096706459700060X-main.pdf?_tid=d3d3eae5-5ca5-4ac5-ba8e-](https://ac.els-cdn.com/S096706459700060X/1-s2.0-S096706459700060X-main.pdf?_tid=d3d3eae5-5ca5-4ac5-ba8e-6e934a548139&acdnat=1535698698_4634563185704240d6eef101ba42f036)
745 [6e934a548139&acdnat=1535698698_4634563185704240d6eef101ba42f036](https://ac.els-cdn.com/S096706459700060X/1-s2.0-S096706459700060X-main.pdf?_tid=d3d3eae5-5ca5-4ac5-ba8e-6e934a548139&acdnat=1535698698_4634563185704240d6eef101ba42f036), 1997.

