Spatial variations of biochemical content and stable isotope ratios of size-fractionated plankton in the Mediterranean Sea (MERITE-HIPPOCAMPE campaign)

Tesán-Onrubia Javier Angel ^{1, *}, Tedetti Marc ¹, Carlotti François ¹, Tenaille Melissa ¹, Guilloux Loïc ¹, Pagano Marc ¹, Lebreton Benoit ², Guillou Gaël ², Fierro-González Pamela ¹, Guigue Catherine ¹, Chifflet Sandrine ¹, Garcia Théo ¹, Boudriga Ismail ³, Belhassen Malika ³, Zouari Amel Bellaaj ³, Bănaru Daniela ^{1, *}

 ¹ Aix Marseille Univ., Université de Toulon, CNRS, IRD, MIO UM 110, 13288 Marseille, France
 ² UMR 7266 Littoral Environnement et Sociétés (CNRS - La Rochelle Université), La Rochelle, France
 ³ Institut National des Sciences et Technologies de la Mer (INSTM), 28, rue 2 mars 1934, 24, Salammbô 2025, Tunisia

* Corresponding authors : Javier Angel Tesan-Onrubia, email address : <u>javier.tesan@mio.osupytheas.fr</u>; Daniela Banaru, email address : <u>daniela.banaru@mio.osupytheas.fr</u>

Abstract :

Plankton represents the main source of carbon in marine ecosystems and is consequently an important gateway for contaminants into the marine food webs. During the MERITE– HIPPOCAMPE campaign in the Mediterranean Sea (April–May 2019), plankton was sampled from pumping and net tows at 10 from the French coast to the Gulf of Gabès (Tunisia) to obtain different size fractions in contrasted regions. This study combines various approaches, including biochemical analyses, analyses of stable isotope ratios (δ 13C, δ 15N), cytometry analyses and mixing models (MixSiar) on size-fractions of phyto- and zooplankton from 0.7 to >2000 µm. Pico- and nanoplankton represented a large energetic resource at the base of pelagic food webs. Proteins, lipids, and stable isotope ratios suggest different sources of carbon and nutrients at the base of the planktonic food webs depending on the coast and the offshore area. In addition, a link between productivity and trophic pathways was shown, with high trophic levels and low zooplankton biomass recorded in the offshore area. The results of our study highlight spatial variations of the trophic structure within the plankton size-fractions and will contribute to assess the role of the plankton as a biological pump of contaminants.

Keywords : Phytoplankton, Zooplankton, 813C, 815N, Plankton trophic structure, Contaminants

- 44 **1. Introduction**
- 45

46 Phyto- and zooplankton play a central role in the functioning of marine ecosystems by 47 producing, transforming and transferring the organic matter up to planktivorous species 48 (Bănaru et al., 2019; Chen et al., 2022). Plankton thus represents the main source of carbon 49 fueling the marine food webs (Bănaru et al., 2013; Cresson et al., 2020). Phyto- and 50 zooplankton are composed of a myriad of organisms of great diversity in terms of size, metabolism, physiology, and diet that are governed by different trophic pathways, and 51 52 complex ecological interactions. As taxonomic composition strongly differs between planktonic size-fractions, a size-based approach has become widely used to study the 53 structure and functioning of the planktonic compartment (Rau et al., 1990; Rolff, 2000; 54 55 Carlotti et al., 2008; Hunt et al., 2017). In pelagic food webs, body size determines rates of production (Banse and Mosher, 1980), energy requirements (Brown et al., 2004), mortality 56 57 rates (Hirst and Kiørboe, 2002) and predator-prev interactions (Cohen et al., 1993; Ljungström et al., 2020). The biochemical composition of the different plankton size-fractions 58 provides information on their energetic content, which may influence prey selection (Carlotti 59 et al., 2008, Harmelin-Vivien et al., 2019; Chen et al., 2019, 2021, 2022), while the carbon 60 and nitrogen isotopic ratios (δ^{13} C and δ^{15} N values) of the plankton size-fractions provide 61 information on the fluxes of organic matter within the planktonic food webs (Peterson et al., 62 1985; Cabana and Rasmussen, 1994; Vander Zanden and Rasmussen, 2001). 63 Besides its central role in the functioning of marine ecosystems, plankton is now recognized 64 as a key gateway of inorganic and organic contaminants into the marine food web (Tao et al., 65 2018; Chouvelon et al., 2019; Tedetti et al., 2023). Phytoplankton is exposed to contaminants 66 67 via water. Bioconcentration of contaminants in phytoplankton is driven mainly by partition equilibrium processes between the cells and the surrounding water (Frouin et al., 2013). For a 68 69 given species or cell size of phytoplankton, and a given physico-chemical habitat 70 (temperature, organic carbon content, etc.), the bioconcentration factors of organic contaminants may be directly correlated with their octanol-water partitioning coefficients (log 71 Kow), i.e., their degree of lipophilicity/hydrophobicity (Frouin et al., 2013). Correspondingly, 72 the bioconcentration of a given contaminant has been shown to increase with decreasing size 73 74 of phytoplankton (Fan and Reinfelder, 2003). Bioaccumulation processes in zooplankton are 75 highly complex due to the entry of contaminants by both the water aqueous phase 76 (bioconcentration) and diet, the trophic interactions and/or transfers between phytoplankton 77 and various zooplankton, and the contaminant removal processes used by these organisms

78 (Tiano et al., 2014; Alekseenko et al., 2018; Tao et al., 2018). Biomagnification may also

occur between the low and high trophic level organisms in the marine food webs (Chouvelonet al., 2019).

81 Therefore, the uptake, accumulation, and transfer of contaminants within the planktonic food 82 webs may be strongly influenced by its characteristics such as the size-fraction distribution, 83 the biochemical/energetic content (Hennig, 1986; Mason et al., 1995; Wu and Wang, 2011), 84 the trophic interactions and the fluxes of organic matter (Peterson et al., 1985; Cabana and Rasmussen, 1994; Vander Zanden and Rasmussen, 2001). Characterizing the content of 85 86 phyto-, and zooplankton in terms of size-fraction, biochemical/energetic content and stable isotope ratios (δ^{13} C and δ^{15} N) may thus provide key information on the structure, functioning, 87 and trophic interactions of this planktonic food webs, but also on the capacity of accumulation 88 and transfer of organic and inorganic contaminants within this planktonic network and 89 potentially their transfer to higher trophic levels. 90

91 The Mediterranean Sea has a high diversity of planktonic and exploited resources, and

92 contrasted biogeographical regions impacted by climate change and human activities (Durrieu

de Madron et al., 2011; Mayot et al., 2017). As the Mediterranean Sea is mainly oligotrophic,

94 its autotrophic biomass is dominated by small-size pico- and nanophytoplankton groups

95 (Leblanc et al., 2018; Boudriga et al., 2022). These groups play a key role in the biomass and

96 energy transfer to zooplankton (Bănaru et al., 2014; Hunt et al., 2017; Leblanc et al., 2018).

97 However, coastal areas impacted by riverine and urban inputs may locally be more enriched

98 and sometimes can even be characterized by higher phytoplankton biomass dominated by

99 larger cells such as dinobionts and diatoms (Harmelin-Vivien et al., 2008). Champalbert

100 (1996) highlighted spatial differences in the zooplankton diversity and biomass, with a lower

101 diversity and a higher biomass in coastal waters relative to offshore areas. The Mediterranean

102 Sea is also strongly exposed to chemical contamination due to the intensive human activities

in the bordering countries, its semi-closed status that limits the dilution of contaminants, and

the relatively short ventilation and residence times its waters (Durrieu de Madron et al., 2011;

105 Tedetti et al., 2023).

106 In this context, the aims of this work, carried out in the frame of the MERITE-HIPPOCAMPE

107 campaign, were: 1) to investigate the structure and functioning (in terms of trophic

108 interactions and organic matter fluxes) of the planktonic food webs of the Mediterranean Sea

109 through the characterization of the biomass, biochemical/energetic content and $\delta^{13}C/\delta^{15}N$

ratios of several size-fractions of phyto- and zooplankton ranging from 0.7 to $> 2000 \ \mu m$, 2)

111 to explore variations in size-spectra and spatial variations of these characteristics

112 (biochemical/energetic content and $\delta^{13}C/\delta^{15}N$ ratios), and relate these variations to their

113 composition (via cytometry and imagery analyses) and also to the specificities of the studied

areas, and 3) to discuss the implications of these findings regarding the accumulation and

transfer of contaminants in the planktonic food webs.

116 To our knowledge, this work represents the first study to use a wide variety of approaches,

and methodologies, such as biochemical analyses (proteins, carbohydrates, lipids and energy),

analyses of isotopic ratios (δ^{13} C, δ^{15} N), cytometry analyses and mixing models (MixSiar) on

fine size-fractions of phyto- and zooplankton ranging from 0.7 to $> 2000 \ \mu m$ for the purpose

of studying plankton food webs. The use of innovative sampling techniques, such as the

deployment of *in situ* pumps equipped with a sequential filtration system, or the towing of a

122 MultiNet device in chlorophyll maximum layer (CML), enabled us to obtain large amounts of

size fractionated phyto- and zooplankton, material rarely acquired together.

- 124
- 125

2. Material and methods

126

127 **2.1. Study area and sampling**

The MERITE-HIPPOCAMPE cruise was conducted in the Mediterranean Sea in spring 2019, 128 from April 13 to May 14, onboard the R/V Antea (Tedetti and Tronczynski, 2019; Tedetti et 129 al., 2023). Ten stations were sampled from the French coast (La Seyne-sur-Mer, Northwestern 130 Mediterranean) to the Gulf of Gabès in Tunisia (Southeastern Mediterranean) in coastal and 131 offshore locations, including the Gulf of Lion, the Ligurian and Algerian consensus regions, 132 the North Balearic Front and Sicilian Channel areas, and the gulfs of Hammamet and Gabès, 133 which presented different hydrological, biogeochemical and bloom conditions, and different 134 levels of anthropogenic pressures (Fig. 1, Table S1). Phytoplankton and zooplankton were 135 sampled at each station at the CML, during spring bloom (April to May), when maximum 136 137 primary and secondary production occurs (Liénart et al., 2018; Tedetti et al., 2023). Suspended particulate matter (SPM) was collected and filtered using McLane Large Volume 138 Water Transfer System Samplers (WTS6-142LV, 4–8 L min⁻¹), hereinafter referred to as 139 McLane *in situ* pumps. One pump was mounted with a regular 142-mm filter-holder holding 140 one 142-mm-diameter filter. This pump was equipped with a ~ 0.7 -µm-pore-size pre-141 combusted pre-weighed GF/F filter (Whatman) and covered with a 60-µm-pore-size 142 homemade sock-type pre-filter so that the filtered particle size-fraction was 0.7–60 µm. A 143 144 second McLane *in situ* pump was mounted with a mini-Multiple Unit Large Volume *in situ* Filtration System (MULVFS) filter holder composed of baffle tubes on the top followed by 145

- 146 successive baffle and filter support plates, for sequential filtration with three different filters
- 147 (142-mm diameter) (Bishop et al., 2012). The filter series used were one ~ 0.7-µm-pore-size
- 148 pre-combusted (450 °C, 6 h), pre-weighed GF/F filter, one 2.7-µm-pore-size pre-combusted,
- pre-weighed GF/D filter (Whatman), and one 20-µm-pore-size pre-cleaned (HCl 0.05% v/v)
- pre-weighed home-made Nylon filter so that the filtered particle size fractions were 0.7-2.7,
- 151 2.7-20, and > 20 μ m (hereafter labeled 20–60 μ m). The pumps were deployed in the CML
- between 40 and 60 min. The volumes of filtered seawater ranged between 169 and 300 L
- depending on the stations. After pump deployment, filters were 'dried' by connecting the filter
- holder to a vacuum pump, stored in pre-combusted (450 °C, 6 h) aluminum sheets, and
- 155 conserved at -20 °C and then freeze-dried. Then, punches with a known surface area were
- 156 made into filters to obtain subsamples for the different analyses.
- Zooplankton was sampled using a Multinet Plankton Sampler (Midi type with 0.25-m²) 157 aperture, Hydro-Bios), referred to hereafter as 'MultiNet', towed horizontally in the CML. The 158 159 MultiNet position was maintained stable at the defined layer by means of a V-fin deflector and controlled vessel speed (2.5 knots), and real-time control of its position from the onboard 160 161 desk-unit. The MultiNet was mounted with five individual (exchangeable) 2.5-m-long nylon nets with a mesh size of 60-um and cod ends of the same mesh size. The MultiNet frame was 162 163 equipped with various sensors: two Hydro-Bios flowmeters (one at the mouth and the other 164 on the side) to assess the volume of water filtered by the nets, a CTD sensor and a total
- 165 chlorophyll *a* (TChl*a*) fluorometer. Connected to the onboard desk unit *via* the electro-
- 166 mechanical cable, these captors inform on *in situ* depth, TChla concentration, filtered volume
- and flow rate, allowing the operator to decide to open and close the nets. Nets were closed
- 168 when the flow rate reached a threshold value to limit clogging. Filtered volumes for each net
- reached up to 185 m^3 . Once the five nets were filled, the MultiNet was hauled back on board,
- and the five cod ends linked by a helicoidal bucket connector were carefully recovered. The
- 171 fifth net remained open until the end of the operation, filtering the layer between the CML
- and the surface. The cod ends were rinsed out with local seawater, and their content
- transferred to pre-cleaned 10-L PFA bottles. The MultiNet was then returned to the water.
- 174 This operation was repeated several times to get sufficient amounts of plankton for all
- possible further contaminant content analyses (Tedetti et al., 2023). In the clean onboard
- 176 container lab, plankton collected in PFA bottles was then size-fractionated through a column
- of five stainless steel sieves (60, 200, 500, 1000 and 2000 µm mesh-size) by wet-sieving with
- seawater previously filtered onto GF/F filters and stored in stainless steel jerrycans to obtain
- the following size-fractions: 60-200, 200-500, 500-1000, 1000-2000 and $> 2000 \mu m$.

Samples were stored in pre-combusted (450 °C, 6 h) Pyrex bottles, conserved at -20 °C and freeze-dried. Large fractions (> 2000 µm) were most likely under-sampled compared to smaller fractions. These fractions should be carefully treated because they represent a relatively low – and difficult to estimate – biomass, form colonies, and may actively evade capture by swimming. The detailed methods to determine zooplankton group composition analysis presented in this paper and related to our results are detailed in Fierro-González et al. (2023).

187

188 **2.2. Analyses**

189

190

2.2.1. Particulate organic carbon and nitrogen

Samples (i.e., 22-mm diameter filter or 1 mg dry weight DW zooplankton) were leached with
100 µL of sulfuric acid (H₂SO₄ 0.5 mol L⁻¹) to remove any inorganic carbon. Samples were
then stored in 25 mL Schott[®] glass bottles for subsequent analyses. Filter blanks were
conditioned with 1 mL of ethanol and with 600 mL of 0.2-µm filtered seawater.
Determination of particulate organic carbon (POC) and nitrogen (PON) concentrations were
carried out simultaneously on the same sample using the persulfate wet-oxidation procedure
according to Raimbault et al. (1999).

198

199 2.2.2. Suspended particulate matter and plankton dry weight

Suspended particulate matter dry weight (SPM_{DW}) was measured in the 0.7–60 and 20–60-µm 200 filters gravimetrically by weighing the tare, wet and dry filters with a precision balance (d = 201 0.01 mg). Briefly, we subtracted from the dry filter the tare and the salts associated with the 202 203 evaporated water, considering a salinity of 38. Tests conducted with filtered seawater evidenced a higher mass after drying than the salt present in water, and most likely related to 204 205 the highly hygroscopic characteristics of salts. A ratio of 1.2 between the observed mass and 206 the theoretical mass associated with salt was applied to correct the water bonded to salt. When biomasses were low, as for 0.7–2.7 and 2.7–20-µm filters, high uncertainty in weighing 207 increased the margin of error. SPM_{DW} in the 0.7-2.7 and 2.7-20-µm filters were thus 208 estimated by multiplying their relative fraction of POC by the biomass measured 209 gravimetrically in the 0.7–60-µm size-fraction. POC can be employed as a proxy of SPM_{DW}, 210 based on the correlations assessed by Trimble and Baskaran (2005). SPM_{DW} at the CML is 211 212 mainly composed of POC (around 50%) but also contains biogenic silica, calcium carbonate and lithogenic matter (Bishop et al., 1977; Krasakopoulou and Karageorgis, 2005). On the 213

basis of these observations, relative POC was used to assess the contribution of the different

size-fractions to the total biomass. Plankton dry weight (Plankton _{DW}) of size fractions 60–

216 200, 200–500, 500–1000, 1000–2000, and > 2000 μ m were determined after drying samples

217 on pre-weighed GF/F filters (60 °C, 24 h) and then re-weighed with a microbalance, taking

into consideration the volumes of water filtered by the MultiNet (Fierro-González et al.,

- 219 2023).
- 220
- 221

2.2.3. Biochemical and energy content

222 Biochemical compounds (proteins, carbohydrates, and lipids) were extracted in triplicate from freeze-dried filters and zooplankton samples and expressed in µg mg⁻¹ DW. Absorbance of 223 the extracts was then measured at different wavelengths using a spectrophotometer 224 (Shimadzu, UV-1280). Briefly, proteins were extracted using a Folin phenol reagent 225 following the Lowry et al. (1951) method and measured at 700 nm. Carbohydrates were 226 extracted using the phenol-sulfuric acid reaction described in Dubois et al. (1956) and 227 measured at 490 nm. Finally, lipids were extracted with a monophasic methanol-228 229 dichloromethane solution (Bligh and Dyer, 1959) and measured at 360 nm. Ashes and residual organic compounds not recovered by these standard biochemical assays (chitin for 230 231 example) may be estimated by subtracting the weight of the biochemical compounds from the total dry weight, but were not presented in this study. The energy content (E_i) was estimated 232 by summing the three biochemical compounds after converting them into energetic units 233 (21.4 kJ g⁻¹ for proteins, 17.2 kJ g⁻¹ for carbohydrates and 35.6 kJ g⁻¹ for lipids) (Postel et al., 234 235 2000). The plankton energy amount (E_T) provided by plankton per cubic meter per station at CML (kJ m⁻³) was calculated for all the plankton size-fractions by multiplying the SPM_{DW} or 236 the biomass (B_i in mg DW m⁻³) of each size-fraction at a given station by its energy content 237 (E_i in kJ mg⁻¹ DW) and the sum of E_T of all size-fractions represents the plankton total energy 238 239 amount, E_{TS} : $E_T = B_i \times E_i$.

240

241

2.2.4. Stable isotope analyses (δ^{13} C and δ^{15} N)

The SPM_{DW} collected on filters was scraped off with a scalpel. Zooplankton samples were ground to a fine powder with an agate mortar and pestle. Presence of carbonates on samples can bias the measurement of δ^{13} C values as they are enriched in ¹³C compared to organic matter (Pinnegar and Polunin, 1999). Therefore, samples were acidified prior to the measurement of δ^{13} C values of the particulate organic matter (POM). Samples were acidified using HCl 1% and immediately rinsed with MilliQ-water. Measurements of δ^{15} N values of the

POM were carried out on raw samples. For zooplankton, approximately 0.5 mg of powder 248 was weighed into a tin cup (8×5 mm) using a precision balance (d = 0.01 mg). Carbon and 249 nitrogen stable isotope analyses were performed using a continuous-flow isotope-ratio mass 250 spectrometer (Delta V Plus, Thermo Scientific) with a Conflo IV interface coupled to an 251 252 elemental analyzer (EA Isolink, Thermo Scientific). Analyses were conducted at the Littoral, Environment and Societies Joint Research Unit stable isotope facility (CNRS - University of 253 254 La Rochelle, France). The δ^{13} C and δ^{15} N values are expressed in δ notation as deviations from standards (Vienna Pee Dee Belemnite for δ^{13} C and N₂ in air for δ^{15} N), in ‰, according to the 255 256 formula:

$$\delta X_{\text{sample}} = \left[\left(\frac{R_{\text{sample}}}{R_{\text{standard}}} \right) - 1 \right] x \ 1000$$

257

where X is 13 C or 15 N, R_{sample} is the isotopic ratio of the sample and R_{standard} is the isotopic ratio of the standard. Calibration was carried out using reference materials (USGS-61, -62, -63 for both carbon and nitrogen). The analytical precision of the measurements was < 0.10‰ for carbon and nitrogen based on analyses of USGS-61 and USGS-63 used as laboratory internal standards.

263

264

2.2.5. Cytometry and imagery analyses

For flow cytometry analyses, seawater was sampled with Niskin and Go-Flo bottles at the 265 CML. Seawater was filtered through a silk mesh filter with 100-µm pore size and then 266 267 immediately fixed with a 2% paraformaldehyde solution. Samples were incubated for 15 min at 4 °C in the dark and then stored in liquid nitrogen onboard and at -80 °C in the laboratory 268 269 until analysis. The autotrophic pico- and nanoplankton were analyzed using a CyFlow[®]Space flow cytometer (SysmexPartec) equipped with a blue diode pumped solid-state laser (20 mW; 270 488 nm) and a red diode laser (25 mW; 638 nm). Seawater samples were thawed in the dark, 271 filtered through a nylon mesh filter with a porosity of 30 µm (CellTrics[®], SysmexPartec), and 272 mixed with flow check high-intensity beads of 2-µm diameter (Polysciences, USA). Cells 273 were characterized based on their scatter and fluorescence signals (Khammari et al., 2018) 274 and data were analyzed with the FloMax software (Sysmex Partec), which directly calculates 275 the cell concentration (cells cm⁻³) of the resolved cell groups. Carbon biomass of each cell 276 group was estimated according to Khammari et al., (2020). The contributions of zooplankton, 277 phytoplankton (microalgae), and detritus components to the total Plankton _{DW} were estimated 278

using imagery methods (FlowCAM and ZOOSCAN) according to Fierro-González et al.
(2023).

281

282 **2.3. Bayesian mixing model**

283 The contributions of the different size-fractions of phytoplankton (0.7–2.7, 2.7–20 and 20–60 µm) as food resources to zooplankton size-fractions and their confidence intervals were 284 estimated using a Bayesian mixing model using the R package MIXSIAR (Parnell et al., 285 2013; https://github.com/brianstock/MixSIAR). This modeling approach incorporates 286 variability of potential food source isotopic compositions for consumers and of trophic 287 fractionation factors (TFF) and generates probability distributions of food source 288 contributions. The consumers (i.e., zooplankton) were grouped in the 60-500, 500-2000, and 289 > 2000-µm size-fractions based on statistical differences in their isotopic compositions. 290 Zooplankton size-fractions with similar isotopic compositions were thus considered to 291 consume the same food resources. Contributions were computed using δ^{13} C and δ^{15} N values. 292 As there is no well-established set of TFFs for plankton, we computed mixing model 293 estimates using the mean TFF values of 1.70 ± 0.38 % for δ^{13} C and 2.40 ± 0.26 % for δ^{15} N 294 295 (mean \pm standard error (SE)), as established by Tiselius and Fransson (2016). 296

297 2.4. Trophic level

The trophic level (TL) of consumers was calculated following the equation proposed by Post (2002):

$$TL = \left(\frac{\delta^{15} N_{consumer} - \delta^{15} N_{baseline}}{TFF \, \delta^{15} N}\right) + 1$$

300

301 where $\delta^{15}N_{\text{consumer}}$ is the isotopic composition of the consumer, $\delta^{15}N_{\text{baseline}}$ is the isotopic 302 composition of the size-fraction 0.7–2.7 µm considered as baseline with a trophic level of 1 303 and TFF $\delta^{15}N$ was fixed to 2.40 ± 0.26 ‰ (Tiselius and Fransson, 2016). TL of each size 304 fraction was weighted to its respective biomass per station (TL_B):

$$TL_{B} = \frac{\sum_{i}^{n} (TL_{i} * \% \text{ Biomass}_{i})}{100}$$

306 where *I* is represented by each size fraction from 60-200 to > 2000 μ m and n is the number of 307 size-fractions.

309 **2.5. Data treatment**

The effect of size-fractions and geographical area on the SPM_{DW}, Plankton_{DW}, POC, PON, C/N, carbon biomass of phytoplankton, biochemical composition, energy content, isotopic composition and trophic levels were tested by means of one-way ANOVA or non-parametric Kruskal-Wallis tests after testing for normality and homogeneity of variances, followed by

appropriate paired comparison tests, using the software Statistica 12. Spearman's rank order
correlation tests were used to assess the significance of the correlations between our results
and some environmental variables (Tedetti et al., 2023).

- 317
- 318

3. Results

319 320

321 **3.1.** Plankton dry weight, TChla, POC, PON and C:N

Significant differences in SPM_{DW} were observed between the different size-fractions (H = 322 23.2, p < 0.0001) (Table 1). When considering all stations, the highest mean SPM_{DW} was 323 measured in the 2.7–20 μ m size-fraction (0.18 ± 0.04 mg DW L⁻¹, n = 10) and the lowest in 324 the 20–60 μ m fraction (0.04 \pm 0.02 mg DW L⁻¹, n = 10). Generally, these concentrations 325 decreased from coastal stations to offshore stations, except for St9, which exhibited the 326 highest SPM_{DW} in the 2.7–20 µm size-fraction (Table 1). The size-fractions 0.7–2.7 and 2.7– 327 20 µm presented the lowest values at St15. The highest SPM_{DW} in the 20–60-µm size-fraction 328 were observed at St17, St19 and St4. TChla concentrations in the > 0.7-µm fraction ranged 329 from 0.21 (St17) to 1.54 μ g L⁻¹ (St9) with a mean value of 0.76 \pm 0.14 μ g L⁻¹ (n = 10) (Table 330 1). The 60–200 and 200–500 µm zooplankton size-fractions showed the highest Plankton_{DW} 331 (Table 1). POC and PON concentrations increased with phytoplankton size with the lowest 332 values in the 0.7–2.7 µm size-fraction (11.4 and respectively 2.1 µg C L⁻¹) and the highest 333 ones in the 0.7–60 μ m size-fraction (42.8 and respectively 6.9 μ g C L⁻¹) (Table S2). The 334 lowest POC and PON values were observed at St11 for the 0.7-2.7 µm size-fraction and at 335 St2 for the 2.7–20 µm size-fraction (Table S2). The C:N ratios in the 0.7 to 60 µm size-336 fractions ranged from 5.7 at St19 to 8.5 at St9. The 0.7–2.7 µm size-fraction presented 337 significantly lower C:N ratios than the 2.7–20-µm and the 20–60 µm size-fractions (Table 338 S2). 339

340

341 **3.2.** Planktonic group composition

Different plankton groups contributed to the biomass of the different size-fractions. The size-342 fraction 0.7–2.7 µm was on average dominated in biomass by Synechococcus spp. (4.6 µg C 343 L^{-1} or ~ 52%) and picoeukaryotes (4.0 µg C L^{-1} or ~ 45%) (Table S3). The size-fraction 2.7– 344 20 μ m was dominated in biomass by the nanoeukaryotes (20.0 μ g C L⁻¹ ~ 88%) and 345 represented the highest biomass among size-fractions. The highest cumulated biomass in the 346 range 0.7–20 µm was found at St1, St9 and St19 (Table S3). POC concentration and the 347 carbon biomass of phytoplankton size-fractions between 0.7 and 20 µm showed a significant 348 linear correlation ($R^2 = 0.66$; p = 0.018) (Fig. S1). Carbon biomass estimated by flow 349 cytometry represented around 86% of the POC measured on filters (y = 0.86x + 11.86). Total 350 biomass was dominated by detritus and phytoplankton in the 60-200 and 200-500 µm size-351 fractions (68.8 and 59.2%, respectively), which decreased with the increasing size-fractions 352 $(10.5\% \text{ in the} > 2000 \text{-}\mu\text{m} \text{ size-fraction})$ (Table S4). Copepods dominated in the zooplankton 353 biomass in the 60 to 2000 µm size-fractions (between 54.3% and 91.9%) while crustaceans 354 and gelatinous dominated the $> 2000 \,\mu m$ size-fraction (Table S4). 355

- 356
- 357

3.3. Biochemical and energy content

The Mediterranean plankton sampled had for all size-fractions and stations combined, except the 0.7–60- μ m fraction, mean dry weight concentrations (± SE) of 222.6 ± 6.9 μ g mg⁻¹ DW (n = 204) for proteins, 96.9 ± 6.5 μ g mg⁻¹ DW (n = 203) for carbohydrates, 35.1 ± 1.5 μ g mg⁻¹ DW (n = 197) for lipids and an E_i of 7.7 ± 0.2 kJ g⁻¹ DW (n = 196).

362 363

3.3.1. Differences between plankton size-fractions

Considering all stations combined, the mean concentration of proteins increased with size and 364 365 was significantly higher in large fractions of zooplankton (500–1000 and 1000–2000 µm) than in small fractions of phytoplankton (0.7–2.7 and 2.7–20 µm). The highest mean protein 366 concentration was recorded in the 1000–2000 μ m size-fraction and the lowest in the largest (> 367 2000 µm) size-fraction (Fig. 2A). The mean concentration in carbohydrates decreased with 368 size. The concentrations measured in phytoplankton (0.7-60 and 20-60 µm) were 369 significantly higher than those of the large zooplankton (1000–2000 and $> 2000 \,\mu$ m) (Fig. 370 2B). The mean concentration in lipids was quite homogeneous between size-fractions, with 371 the highest values measured in the size-fractions 20-60 and 1000-2000 µm, and lowest in the 372 size-fractions 0.7-2.7 and $> 2000 \,\mu\text{m}$ (Fig. 2C). Energy content showed an increasing trend 373 with size for both phyto- and zooplankton size-fractions. The highest values were measured in 374

375

the 20-60 µm and 1000-2000 µm size-fractions, respectively, and the lowest in the sizefractions 0.7-2.7 and $> 2000 \,\mu\text{m}$. (Fig. 2D).

377

378

3.3.2. Differences between stations

To investigate the differences between stations, the size-fractions were grouped in two 379 380 fractions: the phytoplankton fraction (fractions between 0.7 and 60 μ m) and the zooplankton fraction (fractions between 60 and 500 µm that were sampled at all the stations). The highest 381 mean protein concentrations in phyto- and zooplankton fractions were recorded at St9, St11 382 383 and St15, and the lowest at St17 (Fig. 3A, B). The highest mean carbohydrate concentrations were observed at St9 for the phytoplankton (Fig. 3C). The highest mean lipid concentrations 384 and energy values for phyto- and zooplankton were measured at St1, St2, St9 and St11, and 385 the lowest at St3, St4 and St17 (Fig. 3E-H). Overall, greater differences appeared between 386 stations than between size-fractions for most of these biochemical compounds (Table S5). 387 E_{TS} ranged from 1.17 kJ m⁻³ at St3 to 6.30 kJ m⁻³ at St9 with a mean value (\pm SE) of 2.37 \pm 388 0.49 kJ m⁻³ (n = 10) (Fig. 4). High E_{TS} were also observed at coastal stations St1, St4, St17 389 and St19. The 0.7–2.7 and 2.7–20 µm size-fractions represented the largest energetic 390 reservoirs with mean values of 0.65 ± 0.09 kJ m⁻³ (n = 10) and 1.16 ± 0.37 kJ m⁻³ (n = 10), 391 392 respectively (Fig. 4). The 200–500-µm fraction represented the largest energy reservoir for the zooplankton with a mean of 0.164 kJ m⁻³. Although the 0.7–2.7 and 2.7–20 µm fractions 393 had lower E_i (6.2 and 6.3 kJ g⁻¹ DW, respectively) than those measured in the > 60 μ m size-394 fractions (between 7.0 and 9.5 kJ g⁻¹ DW) (Fig. 2), their dry weight was about one order of 395 396 magnitude higher (Tables 1 and S2), thus enhancing their major contribution of phytoplankton to the total plankton energy amount available in the system (Fig. 4). 397 398

3.4. Stable isotope compositions (δ^{13} C and δ^{15} N) 399

For all size fractions and stations combined, except the 0.7–60 μ m fraction, the plankton δ^{13} C 400 values ranged from -26.6 to -17.4 %, with a mean of -23.0 ± 0.1 % (n = 136), whereas the 401 δ^{15} N values ranged from -0.6 to 5.7‰, with a mean of 2.6 ± 0.1‰ (n = 136). 402

- 403
- 404

3.4.1. Differences between plankton size-fractions

Considering all stations combined, the mean δ^{13} C and δ^{15} N values increased with size, 405

especially in the fractions ranging from 0.7 to $60 \,\mu\text{m}$ (Fig. 5, Table S6). Significantly higher 406

mean δ^{13} C values were reported in the 20–60 µm fraction than in the 0.7–2.7 µm fraction. The 407

mean δ^{15} N values of the size-fractions from 0.7 to 60 µm were all significantly different. Size-408

fractions between 60 and 2000 μ m showed slightly higher δ^{13} C values than those of smaller fractions. The highest mean δ^{13} C and δ^{15} N values were measured in the 1000 to > 2000 μ m size-fractions (Fig. 5, Table S6).

412

413

3.4.2. Differences between stations

414 As for the biochemical composition, the size-fractions were grouped into two fractions: the phytoplankton fraction (fractions between 0.7 and 60 μ m) and the zooplankton fraction 415 (fractions between 60 and 500 μ m that were sampled at all the stations). The mean δ^{13} C and 416 417 δ^{15} N values followed the same spatial variations between phyto- and zooplankton fractions (Fig. 6A-D). For both phyto- and zooplankton, the highest mean δ^{13} C values were measured 418 at St4 and St17. The highest mean δ^{15} N values were detected in phytoplankton at St4, while 419 the lowest ones were observed at St9. In zooplankton, the highest mean δ^{15} N values were 420 measured at St10, and the lowest ones at St9, St17, and St19 (Fig. 6A-D). Overall, in contrast 421 to the biochemical composition, greater differences were highlighted between size-fractions 422 than between stations for both δ^{13} C and δ^{15} N values (Table S5). 423

424

425

3.5. Trophic levels

426 For all zooplankton size-fractions (60 to $> 2000 \,\mu$ m) and stations combined, the TL ranged from 1.5 to 3.6 with a mean of 2.2 ± 0.1 (n = 76). When considering all stations combined, 427 mean TL significatively increased with size (F = 5.2, p = 0.001) (Fig. 7). The highest mean 428 TL (2.6 ± 0.2 , n = 10) was measured for the 1000–2000 µm size-fraction and the lowest TL 429 430 $(1.9 \pm 0.1, n = 20)$ in the 60–200 µm fraction. For the 60–200, 200–500, and 500–1000 µm size-fractions, differences in mean TLs were overall significant between stations (H = 18.7, p 431 432 = 0.027; H = 18.9, p = 0.026 and H = 16.3, p = 0.037, respectively) with the highest mean trophic levels by size-fraction at St10. Mean TL weighted by the plankton biomass (TL_B) had 433 434 values larger than 2 at St2, St3 in northern coastal waters, and at St9, St10 and St11 in offshore waters (Fig. 8). The cumulated biomass of the zooplankton groups (60 to $> 2000 \,\mu$ m) 435 of the sampled stations was significantly and negatively correlated with TL_B ($R^2 = 0.61$, p =436 437 0.020) (Fig. S2).

438

439

3.6. Trophic flows in the plankton food web

440 The Mediterranean plankton food web analysis highlighted size predation (Fig. 9, Fig. S3). 441 For all stations combined, the $0.7-2.7 \,\mu m$ fraction was mainly consumed by the $60-500 - \mu m$ 442 fraction, while the consumption of the $2.7-20 \,\mu m$ fraction was similar between the 60-500 and 500–2000- μ m fractions. The > 2000 μ m size-fraction mainly consumed the 20–60 μ m size fraction (Fig. 9, Fig. S3).

445

446

4. Discussion

- 447
- 448

4.1. Plankton biochemical content

The present description of the plankton biochemical content along a north-south transect in
the Mediterranean Sea appears to be unique to our knowledge. The biochemistry of phytoand zooplankton has been extensively documented in many marine environments but remains
little observed in the Mediterranean Sea (Morris and Hopkins, 1983; Danovaro et al., 2000;
Yilmaz and Besiktepe, 2010; Chen et al., 2019).

454 Jónasdóttir (2019) summarizes the biochemistry of phytoplankton by a protein / carbohydrate / lipid average ratio of 5/3/2 (with a range of 40–60, 17–40, and 16–26%), similar to Ríos et 455 al. (1998) who found a ratio of 4.9/3.2/1.9. In our study, the mean ratio of the size-fractions 456 between 0.7 and 60 μ m was on average 5.6 / 3.6 / 0.9, therefore with relatively higher proteins 457 and lower lipids compared to previous ratios. The biochemical ratio in the $> 60 \ \mu m$ size-458 fractions was 6.9 / 2.0 / 1.1, reflecting higher protein and lower carbohydrate contents 459 compared to phytoplankton. The higher proportion of protein to carbohydrate is a good 460 indicator of the nitrogen availability in the environment whereas the inverse pattern may 461 reflect nitrogen-limited environments for both phyto- (Fabiano et al., 1999; Danovaro et al., 462 2000; Yilmaz and Besiktepe, 2010; Kim et al., 2019) and zooplankton (Bhat et al., 1993). 463 Chen et al. (2019) found in spring time in the Bay of Marseille (in the same location as our 464 station St4), for the size-fraction 200–500 µm, carbohydrate concentrations (62.4 to 66.7 mg 465 g^{-1}) similar to ours, while their concentrations in lipids (98.3 to 102.8 mg g^{-1}) and proteins 466

467 (292.7 to 335.7 mg g^{-1}) were higher.

468 The protein and lipid concentrations were higher in zooplankton than in phytoplankton.

469 Within the zooplankton, an increase in proteins with size was observed, in agreement with

470 Guisande (2006). Most of the organisms in our samples between 60 and 2000 μ m were

471 composed of copepods, and, consequently, the different fractions corresponding to different

- 472 life-stages of the same group (Fierro-González et al., 2023) may explain the size influence on
- 473 their biochemical content. The highest concentrations of carbohydrates and lipids reported in
- 474 the 20–60 μ m size-fraction can be related to a mixed composition of microphyto- and
- 475 microzooplanktonic organisms, such as diatoms and metazoan eggs (Danovaro et al., 2000;
- 476 Chen et al., 2019). However, an underestimation of the SPM_{DW} in the 20–60 µm fraction with

- 477 respect to that in the size-fractions from 0.7 to $60 \,\mu m$ due to different estimation methods
- 478 (see section 2.2.2) may explain the higher biochemical concentrations measured in the 20–
- 479 60 μm fractions. Finally, the lowest protein, carbohydrate, and lipid concentrations observed
- 480 in the > 2000- μ m fraction were probably related to the dominance in this fraction of filter
- 481 feeder organisms, such as salps and siphonophores (Bănaru et al., 2014; Hunt et al., 2017;
- 482 Chen et al., 2019; Fierro-González et al., 2023).
- 483 Organic contaminants have been shown to have a strong chemical affinity for lipids or
- 484 proteins (Mason et al., 1995; Wu and Wang, 2011, Frouin et al., 2013). Therefore, without
- 485 considering the size of the organisms (which is also an important factor in the accumulation
- 486 of contaminants in biota, in particular, phytoplankton), we can assume that the concentrations
- 487 of organic contaminants in zooplankton of stations St1, St2, St9 and St11 (which contains
- more lipids and proteins) may be higher than those in the zooplankton of stations St3, St4 and
 St17 (which have less lipids and proteins). The biochemical content of the plankton size
 fractions can change due to variations of the composition of the plankton community and the
 nutrient inputs linked to the physical and chemical environment, thus affecting the entire
- trophodynamics of the ecosystem (Chen et al., 2019, 2021; Tedetti et al., 2023).
- 493

4.2. Plankton total energy amount (E_{TS})

The E_T decreased with size due to plankton biomass reduction with size. In our study, the E_T 495 in SPM_{DW} were lower than previously measured in other areas (Mayzaud et al., 1989; Fabiano 496 et al., 1999; Kim et al., 2019). This is most likely due to the lower phytoplankton biomass 497 498 reported in the oligotrophic Mediterranean Sea, highlighting their low qualitative values even 499 if spatial and seasonal variations may occur. In the eastern Mediterranean basin, the E_i 500 measured in zooplankton by Danovaro et al., (2000) were even lower than in our study, probably related to higher oligotrophy in their study area. In the Bay of Marseille, the E_T 501 502 measured at St4 were higher for us than those recorded in similar zooplankton size-fractions during the same season in 2017 by Chen et al. (2019). Plankton showed higher E_{TS} at St1, St4, 503 504 St9 and St19. At these stations, where bloom conditions prevailed, trophic regimes were probably responsible for enhanced biomass (Mayot et al., 2017; Chen et al., 2019; Tedetti et 505 al., 2023) (Table S1), which may have impacted their E_{TS}. The 0.7–2.7, 2.7–20, and 200–500-506 µm size-fractions represented the largest energetic reservoir in the Mediterranean phyto-, and 507 zooplankton compartments. Our study highlights their importance as major potential sources 508 of contaminants for their consumers. 509

511 **4.3.** Plankton stable isotopes ratios and composition

The fractions comprised between 0.7 and 60 µm corresponded to SPM_{DW} made up of a 512 complex mixture of living, detrital and lithogenic material, difficult to separate (Lam et al., 513 514 2015; Tedetti et al., 2023). The main limitation of food web studies relying of stable isotope ratios is to establish an acceptable baseline, using samples mainly composed of primary 515 516 producers (Harmelin-Vivien et al., 2008; Tamelander et al., 2009). In our study, pico- and nanoplankton dominated the SPM_{DW} biomass which was composed of around 86% of 517 photosynthetic organisms, as confirmed by cytometry results (Boudriga et al., 2022). 518 519 Moreover, their low C:N ratios suggest a composition dominated by living photosynthetic

520 organisms.

521 Few studies have measured isotopic compositions of the smallest size-fractions of plankton

- 522 (Wainright and Fry, 1994; Tamelander et al., 2009), and even fewer have separated them by
- size (Rau et al., 1990; Rolff, 2000; Im et al., 2015; Hunt et al., 2017; Décima, 2022). The
- size-fractions from 0.7 to $60 \,\mu m$ (without considering the 0.7–60 μm fraction) have similar
- 525 δ^{13} C values but their δ^{15} N values are rather low (-23.9 ± 0.2 ‰ and 1.4 ± 0.16 ‰,
- respectively) compared to those measured in other oceanic basins (from -28 to -17 ‰ and
- from 1 to 12 ‰, respectively) (Wainright and Fry, 1994; Rolff, 2000; Tamelander et al.,
- 528 2009) and in the Mediterranean Sea (from -25 to -22 ‰ and from 1 to 6 ‰, respectively)
- 529 (Rau et al., 1990; Harmelin-Vivien et al., 2008; Hunt et al., 2017; Liénart et al., 2017).
- 530 Several processes may explain the low δ^{15} N values of the plankton in our study: 1) in the
- convection areas subjected to algae blooms (such as St9), nutrients made of light isotopes are
- preferentially uptaken by phytoplankton (Wainright and Fry, 1994; Rolff, 2000; Tamelander
- et al., 2009), 2) in water with poor nutrient content, ammonium, the main nitrogen source
- emitted through excretion and recycling, has lower δ^{15} N values (Checkely and Miller, 1989),
- and 3) the high proportion of diazotrophs (atmospheric nitrogen fixer) in oligotrophic waters
- may lower δ^{15} N values (Pantoja et al., 2002; Montoya et al., 2002; Koppelmann et al., 2003).
- 537 Within the 0.7 and 60 μ m size-fractions, δ^{13} C and δ^{15} N values significantly increased with
- size from pico-, nano- to microplankton, similar to previous studies (Im et al., 2015; Hunt et
- al., 2017). Differences between primary producers can be related to differences in the isotopic
- 540 composition of inorganic nitrogen sources and/or fractionation between molecules made of
- heavy and light isotopes during physiological processes (Ostrom and Fry, 1993). A large
- 542 contribution of cyanobacteria, mainly composed of *Synechococcus* spp. may lower δ^{13} C and
- 543 δ^{15} N values of SPM in the 0.7–2.7 µm size-fraction (Rau et al., 1990; Rolff, 2000; Hunt et al.,
- 544 2017). These differences in the composition may also suggest predation within the $0.7-60 \,\mu m$

- 545 fraction (Onodera et al., 2018; Armengol et al., 2019). However, an increase in δ^{15} N values
- 546 with the size has already been observed in autotrophs (Karsh et al., 2003; Hunt et al., 2017).
- 547 The group composed of size-fractions larger than 60 µm and dominated by zooplankton,
- 548 mainly copepods (Fierro-González et al., 2023), presented higher δ^{13} C and δ^{15} N values than
- 549 phytoplankton.
- 550 The isotopic compositions reported here for the 200–500 μ m size-fractions (-22.6 ± 0.4 ‰
- and 3.2 ± 0.2 ‰, for δ^{13} C and δ^{15} N, respectively) were rather in the lower range of previous
- estimates in different oceanic basins (from -22 to -19% and from 2 to 8% for $\delta^{13}C$ and $\delta^{15}N$
- values, respectively) (Fry and Quiñones, 1994; Bode et al., 2007; Yang et al., 2017). Overall,
- the isotopic compositions measured in the different zooplankton size-fractions were within
- the same range as those reported in the ultra-oligotrophic eastern Mediterranean basin
- (Koppelmann et al., 2009; Denda and Christiansen, 2010) and in the more productive area of
- the Gulf of Lion (Espinasse et al., 2014; Bănaru et al., 2014; Hunt et al., 2017).
- 558 Within the zooplankton size-fractions, δ^{15} N values increased with size, up to the 1000–2000
- ⁵⁵⁹ μm fraction, underlining an enhancement of the predation with size (Fry and Quiñones, 1994;
- 560 Rolff, 2000; Bănaru et al., 2014; Espinasse et al., 2014; Hunt et al., 2017). The δ^{15} N values
- 561 increased with the size-fractions, which may be due to the succession of the different life
- stages of copepods, coinciding with the pattern observed in their biochemical composition
- 563 (Espinasse et al., 2014; Chen et al., 2019) related to metabolic changes during the lifespan
- (Guisande, 2006), adaptative foraging (Kozak et al., 2020), ontogenetics (Mauchline, 1998;
- 565 Im et al., 2015) or all these processes combined. Fierro-González et al. (2023) reported an
- 566 increase in the proportion of carnivores with size. However, in the largest size-fraction (>
- $2000 \,\mu\text{m}$), gelatinous organisms were generally overrepresented with respect to other
- 568 fractions, leading to higher δ^{13} C and lower δ^{15} N values due probably to distinct food sources
- from smaller fractions (Bănaru et al., 2014; Espinasse et al., 2014).
- 570
- 571

4.4. Spatial variation in plankton isotopic compositions

572 The highest δ^{15} N values in phytoplankton were recorded at St4, located in the Bay of 573 Marseille, which is highly impacted by anthropogenic and terrestrial inputs. These inputs may 574 induce, in case of declining nitrates related to intense bloom events, an increase of the δ^{15} N 575 values (Raimbault et al., 2008; Fey et al., 2021). Moreover, the effluents of the Marseille 576 sewage treatment plant may be ¹⁵N-enriched in the inorganic nitrogen pool due to bacterial 577 denitrification (Wainright and Fry, 1994; Cabana and Rasmussen, 1996). The high δ^{13} C 578 values measured in phyto- and zooplankton in the Bay of Marseille (St3 and St4) and in the

Gulf of Gabès (St17 and St19) may be due to higher growth rates and nutrient availability in 579 these productive areas (Laws et al., 1995; Bidigare et al., 1997), as well as to inputs of 580 sedimentary organic matter in the water column during sediment resuspension events (Table 581 S1). Moreover, the low $\delta^{15}N$ values measured on the Tunisian coasts support this hypothesis 582 and may be related to seagrass detritus (Cresson et al., 2012). St9, located at the border of the 583 584 convective area of the Northwestern Mediterranean Sea, was characterized by a post-bloom event during the campaign (Tedetti et al., 2023) and displayed the lowest δ^{15} N values in 585 phyto- and zooplanktonic organisms. The highest C:N measured at this station may suggest a 586 587 higher detrital organic matter content and a higher microbial production of regenerated isotopically lighter ammonium compared to the other stations (Maguer et al., 2000). 588 The relatively high δ^{13} C (and δ^{15} N) values measured in phyto- and zooplankton in the Bay of 589 Marseille and the Gulf of Gabès may also be related to various inputs (sediment resuspension, 590 effluents, etc.), and be the sign of higher levels of contaminants which could derive from the 591 same sources. To support this hypothesis, higher trace metal concentrations have been 592 reported in these coastal stations compared to other MERITE-HIPPOCAMPE stations 593 594 (Chifflet et al., 2023).

595

596

4.5. Trophic levels and food web implications

597 The TLs determined in this study, ranging from 1.5 to 3.6, increased with size and spread out 598 over 2 TLs. TLs lower than 2 may be related to a mixture of primary producers and 599 consumers mainly found at some stations in the 60–200 µm size-fraction, related to clogging 600 of the sampling device caused by phytoplankton. Further improvements in sampling methods 601 by using larger mesh-size nets for zooplankton size-fractions collection may avoid potential 602 clogging during bloom events and the presence of phytoplankton within the 60–200 µm size-603 fraction.

604 When biomass is averaged to the TL of zooplankton (TL_B), higher values are obtained at the coastal (St2, and St3), and offshore (St9, St10, and St11) stations. High adaptability in food 605 606 habits of zooplankton is well known and omnivory seems to be the most common trophic 607 pathway (Fierro-González et al., 2023). Higher TLs are generally reached in microbial food webs, often observed in oligotrophic regimes (Søreide et al., 2006; Kürten et al., 2013). St10 608 and St11 contained carnivorous plankton, such as chaetognaths which may explain the higher 609 TLs of the zooplankton community observed in these areas (Fierro-González et al., 2023). It 610 has been suggested that the TFF between phytoplankton and protozoans is equal to 0, 611 contributing to an underestimation of the number of TLs in planktonic food webs (Gutiérrez-612

Rodríguez et al., 2014). However, in this work, we suggest that the higher TLs of planktonwere probably related to the microbial food web.

615 Coastal stations (St1, St4, St15, St17 and St19) were characterized by lower TL_B. The 616 lower TL of plankton in productive systems is classic in herbivore food webs, dominated by 617 larger phytoplankton organisms readily available to zooplankton (Sommer et al., 2002; 618 Fileman et al., 2007). Food chain length decreases in productive ecosystems due to an 619 adaptive foraging of consumers which selects the most abundant low TL resources (Kondoh 620 and Ninomiya, 2009). Detritus and phytoplankton were dominant in these stations in the 60– 621 200 μ m size-fraction contributing to lowering TL_B values (Fierro-González et al., 2023).

In addition, the zooplankton biomass was related to the trophic pathways. A negative 622 relationship between the TL_B and the zooplankton biomass was observed during this study. 623 Higher biomasses of zooplankton have been measured at coastal stations relative to offshore 624 areas in relation to their primary production (Champalbert, 1996). Where TLs were high (St9, 625 St10 and St11), large size fractions (500–1000 and 1000–2000 µm) were also well 626 represented, accounting for 27 to 54% of the total zooplankton biomass. Espinasse et al. 627 628 (2014) have already highlighted the overrepresentation of large plankton organisms in moderately productive areas, while Décima (2022) have reported an inverse relationship 629 630 between food chain length and productivity in zooplankton. Although located in the offshore area, St9 was characterized by a still productive post-bloom situation (Tedetti et al., 2023) 631 with high TL_B. Usually, during post-bloom, diatoms give way to inedible algae not consumed 632 by zooplankton organisms and the source of organic matter is obtained through the cell lysis, 633 634 fuelling microbial loop and enhancing food steps (Sommer et al., 2002). High C:N ratios observed at St9 may indicate heterotrophy within the 0.7–60 µm fractions (Harmelin-Vivien 635 636 et al., 2008). The post-bloom situation may induce a decrease of the resources and thus mesoplankton preying on microplankton (Levinsen et al., 2000; Basedow et al., 2016). Hence, 637 638 the results reported here highlight different trophodynamic scenarios in the Mediterranean plankton food webs and their control on planktonic biomasses, as hypothesized by Fry and 639 640 Quiñones (1994). Contrasting carbon fluxes can impact contaminant transfer. The ability of each contaminant to 641

bioaccumulate can lead to different scenarios. In low-productive ecosystems, organic matter

fluxes go through more trophic steps, which can favor a higher transfer of contaminants that

644 will bioaccumulate such as methylmercury, PCBs or Cd (Tiano et al., 2014; Schartup et al.,

645 2018; Chouvelon et al., 2019). Conversely, in productive ecosystems, direct access to organic

646 matter sources and reduced food steps favors higher transfer of contaminants with little or no

capacity to bioaccumulate, such as the numerous trace elements Co, Ni, Cu, Ag, Pb, and Zn(Chouvelon et al., 2019).

649

650 **4.6. Plankton food web flows**

651 The difference in stable isotope composition between phytoplankton size-fractions made them 652 relevant candidates for mixing model analysis of their relative contributions to zooplankton biomass (Phillips et al., 2014; Hunt et al., 2017). Overall, the contribution of the 0.7–2.7-µm 653 size-fraction to the diet of the $60-500 \,\mu\text{m}$ size fraction underlines the role of the 654 655 microzooplankton (60–200 μ m) and of the small mesozooplankton (200–500 μ m) in the transfer of organic matter from the baseline to higher trophic level consumers. The 656 picoplankton size fraction was mainly composed of Synechococcus spp., the most abundant 657 phytoplankton organism in the Mediterranean Sea that sustains primary production (Boudriga 658 et al., 2022). In fact, during the MERITE-HIPPOCAMPE campaign, picoplankton accounted 659 660 for, on average, 27% of the total biomass of phytoplankton. Classically, we consider that nanoflagellates and microzooplankton (from 2 to 200 µm size-661 fractions) may represent intermediate food steps (Ryther, 1969; Calbet and Landry, 1999) 662 ensuring the link between the 0.7–2.7 μ m fraction and mesozooplankton (200–2000 μ m) 663 664 (Sommer et al., 2000). The 60 to 500 µm size-fractions were constituted of a large percentage of phytoplankton and detritus that would have contributed to lowering their stable isotope 665 ratios. This may have consequences with regard to the mixing model results and may explain 666 the high percentage of picoplankton estimated in the diet of 60–500 µm size-fraction. 667 668 However, in agreement with our results, and despite their small size, recent studies

- demonstrated that small copepods may feed on picoplankton (Motwani and Gorokhova, 2013;
- 670 Im et al., 2015; Major et al., 2017).
- 671 Primary producers were also dominant in the $2.7-20 \,\mu m$ fractions, representing 69% of the

672 phytoplanktonic biomass and about half of the total food supply to both the 60–500 and the

- 673 500–2000 μm size-fractions. Nanoplankton was mainly composed of nanoeukaryotes and its
- 674 contribution to the planktonic food web seems essential, in agreement with the results of Hunt
- et al. (2017). Finally, the 20–60 μ m fraction has low biomass and represents an important
- 676 contribution only to the diet of the $> 2000 \,\mu m$ size-fractions. TFFs for plankton should be
- 677 experimentally determined and a higher number of replicates should be analyzed to improve
- 678 model results and to reduce their variability.
- 679 Differences in contaminant accumulation have been reported in phytoplankton size-fractions.
- 680 Trace metals reflected, for each element, contrasted accumulation patterns between small and

large phytoplankton fractions (Chifflet et al., 2023). Higher methylmercury concentrations

682 were reported during a picoplanktonic bloom, probably due to a higher surface to volume

ratio, favoring uptake (Heimbürger et al., 2010). In addition, for PCBs, a higher accumulation

- 684 was observed in smaller cells (Axelman et al., 1997). The different carbon size-fractions of
- food sources can thus influence the transfer of contaminants and concentrations in
- 686 zooplankton consumers.
- 687

688 5. Conclusions

During the MERITE-HIPPOCAMPE campaign, plankton size-fractions revealed contrasted 689 biochemical and isotopic compositions related to their size, composition and location. 690 Carbohydrate concentrations were the highest in phytoplankton (between 0.7 and 60 µm size-691 fractions) with respect to zooplankton (> $60 \mu m$ size-fractions), which displayed high protein 692 and lipid contents increasing with size. The affinity of proteins and lipids for contaminants 693 makes zooplankton more sensitive to their accumulation (regardless of the size of organisms). 694 Due to the high amount of energy contained in their biomass, the pico- and nanoplankton 695 represents a major food resource in Mediterranean ecosystems fueling the zooplankton food 696 webs and an essential pathway for contaminants due to the combination of small size and high 697 698 energetic content. In the zooplankton community, the 200-500 µm size-fraction, dominated by copepods, showed the highest energy amount, which explains their role as essential food 699 700 resource for many planktivorous fishes. Spatial variations of isotopic compositions of plankton size-fractions were also observed, revealing different carbon sources between the 701 702 coast and the offshore areas, and between the Bay of Gabès (southern coast) and the Bay of 703 Toulon (northern coast), and resulting in a different exposure to contaminants. Trophic levels 704 revealed increasing predation with size, which can result in the biomagnification of contaminants. Trophic pathways in planktonic food webs displayed spatial variations 705 706 influenced by the availability of phytoplankton resources. Higher predation occurred in low productive areas where the zooplankton community reached the highest mean trophic levels, 707 708 increasing the potential exposure of planktonic food webs to contaminants. Our findings on 709 the spatial variations of the biomass, biochemical composition, and the role of the different plankton size-fractions in the food web, as well as the different flows of organic matter in the 710 pelagic food webs, are an essential step for the comprehension of the transfer of contaminants 711 in the Mediterranean pelagic food webs. 712

- 713
- 714

715 Author contribution statement

- 716 Conception and design of study: DB, MT, FC, MP,
- 717 Acquisition of data: JATO, MeTe, LG, PF, BL, GG, IB, MB, AZ, MT, SC
- 718 Analysis and/or interpretation of data: JATO, DB, MT, TG, CC, SC
- 719 Drafting of the manuscript: JATO, DB, MT
- 720 Revising/editing of the manuscript: JATO, DB, MT, FC, BL
- 721 Project administration and funding acquisition: DB, MT, FC, MP
- 722

723 Acknowledgements

- The authors wish to thank the crew of the R/V Antea and the various platforms of the
- 725 Mediterranean Institute of Oceanography having contributed to the data acquisition:
- 726 Plateforme Analytique de Chimie des Environnements Marins (PACEM) for the POC/PON
- 727 measurements, Plateforme Régionale de Cytométrie pour la Microbiologie (PRECYM) for
- 728 cytometric analyses, Microscopie et Imagerie numérique (MIM) for identification expertise
- and the Service Atmosphère-Mer (SAM) for the technical and operational tasks. The authors
- thank the LIENSs joint research unit (CNRS La Rochelle University) for the measurement
- of δ^{13} C and δ^{15} N values. The MERITE-HIPPOCAMPE project has been funded by the cross-
- 732 disciplinary *Pollution & Contaminants* axis of the CNRS-INSU MISTRALS program (joint
- action of the MERMEX-MERITE and CHARMEX subprograms) and received support from
- the IRD French-Tunisian International Joint Laboratory (LMI) COSYS-Med. We are grateful
- for the additional funding received from IFREMER, the MIO Action Sud and Transverse
- Axis programs (CONTAM Transverse Axis), and from the IRD Ocean Department. This
- study received funding by the CONTAMPUMP project (ANR JCJC #19-CE34-0001-01).
- Finally, we warmly thank three anonymous Reviewers for their very relevant and useful
- comments and corrections on the manuscript, as well as Michael Paul for the English
- 740 corrections.
- 741

742 Supplementary information

- 743 Supplementary material related to this article is available online at: xxx
- 744

745 **References**

- Alekseenko, E., Thouvenin, B., Tronczyński, J., Carlotti, F., Garreau, P., Tixier, C., Baklouti,
- 747 M., 2018. Modeling of PCB trophic transfer in the Gulf of Lions; 3D coupled model

- application. Marine Pollution Bulletin 128, 140–155.
- 749 https://doi.org/10.1016/j.marpolbul.2018.01.008
- 750 Armengol, L., Calbet, A., Franchy, G., Rodríguez-Santos, A., Hernández-León, S., 2019.
- 751 Planktonic food web structure and trophic transfer efficiency along a productivity
- gradient in the tropical and subtropical Atlantic Ocean. Scientific Reports 9, 2044.
- 753 https://doi.org/10.1038/s41598-019-38507-9
- Axelman, J., Broman, D., Näf, C., 1997. Field Measurements of PCB Partitioning between
- 755 Water and Planktonic Organisms: Influence of Growth, Particle Size, and
- Solute–Solvent Interactions. Environmental Science & Technology 31, 665–669.
 https://doi.org/10.1021/es960088+
- 758 Bănaru, D., Carlotti, F., Barani, A., Grégori, G., Neffati, N., Harmelin-Vivien, M., 2014.
- 759 Seasonal variation of stable isotope ratios of size-fractionated zooplankton in the Bay of
- 760 Marseille (NW Mediterranean Sea). Journal of Plankton Research 36, 145–156.
- 761 https://doi.org/10.1093/plankt/fbt083
- 762 Bănaru, D., Diaz, F., Verley, P., Campbell, R., Navarro, J., Yohia, C., Oliveros-Ramos, R.,
- Mellon-Duval, C., Shin, Y.-J., 2019. Implementation of an end-to-end model of the Gulf
 of Lions ecosystem (NW Mediterranean Sea). I. Parameterization, calibration and
- revaluation. Ecological Modelling 401, 1–19.
- 766 https://doi.org/10.1016/j.ecolmodel.2019.03.005
- 767 Bănaru, D., Mellon-Duval, C., Roos, D., Bigot, J.-L., Souplet, A., Jadaud, A., Beaubrun, P.,
- Fromentin, J.-M., 2013. Trophic structure in the Gulf of Lions marine ecosystem (north-
- western Mediterranean Sea) and fishing impacts. Journal of Marine Systems 111–112,

770 45–68. https://doi.org/10.1016/j.jmarsys.2012.09.010

- Banse, K., Mosher, S., 1980. Adult Body Mass and Annual Production/Biomass Relationships
 of Field Populations. Ecological Monographs 50, 355–379.
- 773 https://doi.org/10.2307/2937256
- Basedow, S.L., de Silva, N.A.L., Bode, A., van Beusekorn, J., 2016. Trophic positions of
- mesozooplankton across the North Atlantic: estimates derived from biovolume spectrum
- theories and stable isotope analyses. Journal of Plankton Research 38, 1364–1378.
- 777 https://doi.org/10.1093/plankt/fbw070
- Bhat, K.L., Rayadurga, S., Ansari, Z.A., 1993. Biochemical composition of zooplankton from
 the northern Arabian Sea. Pakistan Journal of Marine Sciences2, 17-22.
- 780 Bidigare, R.R., Fluegge, A., Freeman, K.H., Hanson, K.L., Hayes, J.M., Hollander, D.,
- Jasper, J.P., King, L.L., Laws, E.A., Milder, J., Millero, F.J., Pancost, R., Popp, B.N.,

- 782 Steinberg, P.A., Wakeham, S.G., 1997. Consistent fractionation of 13C in nature and in
- the laboratory: Growth-rate effects in some haptophyte algae. Global Biogeochemical
- 784 Cycles 11, 279–292. https://doi.org/10.1029/96GB03939
- Bishop, J.K.B., Edmond, J.M., Ketten, D.R., Bacon, M.P., Silker, W.B., 1977. The chemistry,
 biology, and vertical flux of particulate matter from the upper 400 m of the equatorial
- 787 Atlantic Ocean. Deep Sea Research 24, 511–548. https://doi.org/10.1016/0146788 6291(77)90526-4
- Bishop, J.K.B., Lam, P.J., Wood, T.J., 2012. Getting good particles: Accurate sampling of
 particles by large volume in-situ filtration: Getting good particles. Limnology and
 Oceanography: Methods 10, 681–710. https://doi.org/10.4319/lom.2012.10.681

Bligh, E.G., Dyer, W.J., 1959. A rapid method of total lipid extraction and purification.

Canadian Journal of Biochemistry and Physiology 37, 911–917.

- 794 https://doi.org/10.1139/o59-099
- Bode, A., Alvarez-Ossorio, M.T., Cunha, M.E., Garrido, S., Peleteiro, J.B., Porteiro, C.,
- Valdés, L., Varela, M., 2007. Stable nitrogen isotope studies of the pelagic food web on
 the Atlantic shelf of the Iberian Peninsula. Progress in Oceanography, 74, 115–131.
 https://doi.org/10.1016/j.pocean.2007.04.005
- Boudriga, I., Thyssen, M., Zouari, A., Garcia, N., Tedetti, M., Bel Hassen, M., 2022.
- 800 Ultraphytoplankton community structure in subsurface waters along a North-South
- 801 Mediterranean transect. Marine Pollution Bulletin 182, 113977.
- 802 https://doi.org/10.1016/j.marpolbul.2022.113977
- Brown, J.H., Gillooly, J.F., Allen, A.P., Savage, V.M., West, G.B., 2004. Toward a Metabolic
 Theory of Ecology. Ecology 85, 1771–1789. https://doi.org/10.1890/03-9000
- 805 Cabana, G., Rasmussen, J.B., 1994. Modelling food chain structure and contaminant

bioaccumulation using stable nitrogen isotopes. Nature 372, 255–257.

- 807 https://doi.org/10.1038/372255a0
- 808 Cabana, G., Rasmussen, J.B., 1996. Comparison of aquatic food chains using nitrogen
- isotopes. Proceedings of the National Academy of Sciences 93, 10844–10847.
- 810 https://doi.org/10.1073/pnas.93.20.10844
- 811 Calbet, A., Landry, M.R., 1999. Mesozooplankton influences on the microbial food web:
- 812 Direct and indirect trophic interactions in the oligotrophic open ocean. Limnology and
- 813 Oceanography 44, 1370–1380. https://doi.org/10.4319/lo.1999.44.6.1370
- 814 Carlotti, F., Thibault-Botha, D., Nowaczyk, A., Lefèvre, D., 2008. Zooplankton community
- structure, biomass and role in carbon fluxes during the second half of a phytoplankton

- bloom in the eastern sector of the Kerguelen Shelf (January–February 2005). Deep Sea
- 817 Research Part II: Topical Studies in Oceanography, 55, 720–733.
- 818 https://doi.org/10.1016/j.dsr2.2007.12.010
- Champalbert, G., 1996. Characteristics of zooplankton standing stock and communities in the
 western Mediterranean Sea: relations to hydrology. Scientia Marina 60, 97–113.
- 821 Checkley, D.M., Miller, C.A., 1989. Nitrogen isotope fractionation by oceanic zooplankton.
- Deep Sea Research Part A. Oceanographic Research Papers 36, 1449–1456.

823 https://doi.org/10.1016/0198-0149(89)90050-2

- Chen, C.-T., Bănaru, D., Carlotti, F., Faucheux, M., Harmelin-Vivien, M., 2019. Seasonal
 variation in biochemical and energy content of size-fractionated zooplankton in the Bay
- of Marseille (North-Western Mediterranean Sea). Journal of Marine Systems 199,

827 103223. https://doi.org/10.1016/j.jmarsys.2019.103223

- 828 Chen, C.-T., Carlotti, F., Harmelin-Vivien, M., Guilloux, L., Bănaru, D., 2021. Temporal
- variation in prey selection by adult European sardine (Sardina pilchardus) in the NW
 Mediterranean Sea. Progress in Oceanography 196, 102617.
- 831 https://doi.org/10.1016/j.pocean.2021.102617
- Chen, C.-T., Carlotti, F., Harmelin-Vivien, M., Lebreton, B., Guillou, G., Vassallo, L., Le
 Bihan, M., Bănaru, D., 2022. Diet and trophic interactions of Mediterranean
- planktivorous fishes. Marine Biology 169, 119. https://doi.org/10.1007/s00227-022-
- 835 04103-1
- 836 Chifflet, S., Briant, N., Tesán-Onrubia, J.A., Zaaboub, N., Amri, S., Radakovitch, O., Bănaru,
- B37 D., Tedetti, M., 2023. Distribution and accumulation of trace metal in the planktonic food
- web of the Mediterranean Sea (MERITE-HIPPOCAMPE campaign). Marine Pollution
 Bulletin 186, 114384. https://doi.org/10.1016/j.marpolbul.2022.114384.
- 840 Chouvelon, T., Strady, E., Harmelin-Vivien, M., Radakovitch, O., Brach-Papa, C., Crochet,
- 841 S., Knoery, J., Rozuel, E., Thomas, B., Tronczynski, J., Chiffoleau, J.-F., 2019. Patterns
- 842 of trace metal bioaccumulation and trophic transfer in a phytoplankton-zooplankton-small
- pelagic fish marine food web. Marine Pollution Bulletin 146, 1013–1030.
- 844 https://doi.org/10.1016/j.marpolbul.2019.07.047
- Cohen, J.E., Pimm, S.L., Yodzis, P., Saldana, J., 1993. Body Sizes of Animal Predators and
- Animal Prey in Food Webs. The Journal of Animal Ecology 62, 67.
- 847 https://doi.org/10.2307/5483
- 848 Cossa, D., Knoery, J., Bănaru, D., Harmelin-Vivien, M., Sonke, J.E., Hedgecock, I.M., Bravo,
- A.G., Rosati, G., Canu, D., Horvat, M., Sprovieri, F., Pirrone, N., Heimbürger-Boavida,

- 850 L.-E., 2022. Mediterranean Mercury Assessment 2022: An Updated Budget, Health
- 851 Consequences, and Research Perspectives. Environmental Science & Technology 56,
- 852 3840–3862. https://doi.org/10.1021/acs.est.1c03044
- 853 Cresson, P., Ruitton, S., Fontaine, M.-F., Harmelin-Vivien, M., 2012. Spatio-temporal
 854 variation of suspended and sedimentary organic matter quality in the Bay of Marseilles
- 855 (NW Mediterranean) assessed by biochemical and isotopic analyses. Marine Pollution
- 856 Bulletin 64, 1112–1121. https://doi.org/10.1016/j.marpolbul.2012.04.003
- 857 Cresson, P., Chouvelon, T., Bustamante, P., Bănaru, D., Baudrier, J., Le Loc'h, F., Mauffret,
- A., Mialet, B., Spitz, J., Wessel, N., Briand, M.J., Denamiel, M., Doray, M., Guillou, G.,

Jadaud, A., Lazard, C., Prieur, S., Rouquette, M., Saraux, C., Serre, S., Timmerman, C.-

A., Verin, Y., Harmelin-Vivien, M., 2020. Primary production and depth drive different

- trophic structure and functioning of fish assemblages in French marine ecosystems.
- Progress in Oceanography 186, 102343. https://doi.org/10.1016/j.pocean.2020.102343

B63 Danovaro, R., Dell'Anno, A., Pusceddu, A., Marrale, D., Della Croce, N., Fabiano, M.,

- Tselepides, A., 2000. Biochemical composition of pico-, nano- and micro-particulate organic matter and bacterioplankton biomass in the oligotrophic Cretan Sea (NE
- Mediterranean). Progress in Oceanography 46, 279–310. https://doi.org/10.1016/S00796611(00)00023-9
- Décima, M., 2022. Zooplankton trophic structure and ecosystem productivity. Marine
 Ecology Progress Series 692, 23–42. https://doi.org/10.3354/meps14077
- 870 Denda, A., Christiansen, B., 2010. Zooplankton at a seamount in the eastern Mediterranean:
- distribution and trophic interactions. Journal of the Marine Biological Association of the
 United Kingdom 91, 33–49. https://doi.org/10.1017/S0025315410001153
- 873 DuBois, Michel., Gilles, K.A., Hamilton, J.K., Rebers, P.A., Smith, Fred., 1956. Colorimetric
- Method for Determination of Sugars and Related Substances. Analytical Chemistry 28,
 350–356. https://doi.org/10.1021/ac60111a017
- 876 Durrieu de Madron, X., Guieu, C., Sempéré, R., Conan, P., Cossa, D., D'Ortenzio, F.,
- 877 Estournel, C., Gazeau, F., Rabouille, C., Stemmann, L., Bonnet, S., Diaz, F., Koubbi, P.,
- 878 Radakovitch, O., Babin, M., Baklouti, M., Bancon-Montigny, C., Belviso, S.,
- 879 Bensoussan, N., Bonsang, B., Bouloubassi, I., Brunet, C., Cadiou, J.-F., Carlotti, F.,
- 880 Chami, M., Charmasson, S., Charrière, B., Dachs, J., Doxaran, D., Dutay, J.-C., Elbaz-
- 881 Poulichet, F., Eléaume, M., Eyrolles, F., Fernandez, C., Fowler, S., Francour, P.,
- 882 Gaertner, J.C., Galzin, R., Gasparini, S., Ghiglione, J.-F., Gonzalez, J.-L., Goyet, C.,
- 883 Guidi, L., Guizien, K., Heimbürger, L.-E., Jacquet, S.H.M., Jeffrey, W.H., Joux, F., Le

- Hir, P., Leblanc, K., Lefèvre, D., Lejeusne, C., Lemé, R., Loÿe-Pilot, M.-D., Mallet, M., 884 Méjanelle, L., Mélin, F., Mellon, C., Mérigot, B., Merle, P.-L., Migon, C., Miller, W.L., 885 Mortier, L., Mostajir, B., Mousseau, L., Moutin, T., Para, J., Pérez, T., Petrenko, A., 886 Poggiale, J.-C., Prieur, L., Pujo-Pay, M., Pulido-Villena, Raimbault, P., Rees, A.P., 887 888 Ridame, C., Rontani, J.-F., Ruiz Pino, D., Sicre, M.A., Taillandier, V., Tamburini, C., Tanaka, T., Taupier-Letage, I., Tedetti, M., Testor, P., Thébault, H., Thouvenin, B., 889 Touratier, F., Tronczynski, J., Ulses, C., Van Wambeke, F., Vantrepotte, V., Vaz, S., 890 Verney, R., 2011. Marine ecosystems' responses to climatic and anthropogenic forcings 891 892 in the Mediterranean. Progress in Oceanography 91, 97–166. https://doi.org/10.1016/j.pocean.2011.02.003 893 Espinasse, B., Harmelin-Vivien, M., Tiano, M., Guilloux, L., Carlotti, F., 2014. Patterns of 894 variations in C and N stable isotope ratios in size-fractionated zooplankton in the Gulf of 895 Lion, NW Mediterranean Sea. Journal of Plankton Research 36, 1204–1215. 896 897 https://doi.org/10.1093/plankt/fbu043 Fabiano, M., Danovaro, R., Povero, P., 1999. Vertical Distribution and Biochemical 898 Composition of Pico- and Microparticulate Organic Matter in the Ross Sea (Antarctica), 899 in: Spezie, G., Manzella, G.M.R. (Eds.), Oceanography of the Ross Sea Antarctica. 900 901 Springer Milan, Milano, pp. 233–246. https://doi.org/10.1007/978-88-470-2250-8_16 Fan, C.-W., Reinfelder, J.R., 2003. Phenanthrene Accumulation Kinetics in Marine Diatoms. 902 903 Environmental Science & Technology 37, 3405–3412. https://doi.org/10.1021/es026367g Fey, P., Parravicini, V., Bănaru, D., Dierking, J., Galzin, R., Lebreton, B., Meziane, T., 904 905 Polunin, N.V.C., Zubia, M., Letourneur, Y., 2021. Multi-trophic markers illuminate the understanding of the functioning of a remote, low coral cover Marquesan coral reef food 906 907 web. Scientific Reports 11, 20950. https://doi.org/10.1038/s41598-021-00348-w Fierro-González, P., Pagano, M., Guilloux, L., Makhlouf, N., Tedetti, M., Carlotti, F., 2023. 908 909 Zooplankton biomass, size structure, and associated metabolic fluxes with focus on its roles at the chlorophyll maximum layer during the plankton-contaminant MERITE-910 HIPPOCAMPE cruise. Submitted to this special issue. 911 912 Fileman, E., Smith, T., Harris, R., 2007. Grazing by Calanus helgolandicus and Para-
- 913 Pseudocalanus spp. on phytoplankton and protozooplankton during the spring bloom in
- the Celtic Sea. Journal of Experimental Marine Biology and Ecology 348, 70–84.
- 915 https://doi.org/10.1016/j.jembe.2007.04.003
- 916 Frouin, H., Dangerfield, N., Macdonald, R.W., Galbraith, M., Crewe, N., Shaw, P., Mackas,
- 917 D., Ross, P.S., 2013. Partitioning and bioaccumulation of PCBs and PBDEs in marine

- 918 plankton from the Strait of Georgia, British Columbia, Canada. Progress in
- 919 Oceanography, Strait of Georgia Ecosystem Research Initiative (ERI) 115, 65–75.
- 920 https://doi.org/10.1016/j.pocean.2013.05.023
- Fry, B., Quiñones, R.B., 1994. Biomass spectra and stable isotope indicators of trophic level
 in zooplankton of the northwest Atlantic. Marine Ecology Progress Series 112, 201–204.
- 923 Guisande, C., 2006. Biochemical fingerprints in zooplankton. Limnetica 25, 369–376.
- 924 Gutiérrez-Rodríguez, A., Décima, M., Popp, B.N., Landry, M.R., 2014. Isotopic invisibility of
- protozoan trophic steps in marine food webs. Limnology and Oceanography 59, 1590–
 1598. https://doi.org/10.4319/lo.2014.59.5.1590
- 927 Harmelin-Vivien, M., Bănaru, D., Dromard, C.R., Ourgaud, M., Carlotti, F., 2019.
- 928 Biochemical composition and energy content of size-fractionated zooplankton east of the
- 929 Kerguelen Islands. Polar Biology 42, 603–617. https://doi.org/10.1007/s00300-019930 02458-8
- Harmelin-Vivien, M., Loizeau, V., Mellon, C., Beker, B., Arlhac, D., Bodiguel, X., Ferraton,
 F., Hermand, R., Philippon, X., Salen-Picard, C., 2008. Comparison of C and N stable
 isotope ratios between surface particulate organic matter and microphytoplankton in the
 Gulf of Lions (NW Mediterranean). Continental Shelf Research, 28, 1911–1919.
 https://doi.org/10.1016/j.csr.2008.03.002
- Heimbürger, L.-E., Cossa, D., Marty, J.-C., Migon, C., Averty, B., Dufour, A., Ras, J., 2010.
- 937 Methyl mercury distributions in relation to the presence of nano- and picophytoplankton
 938 in an oceanic water column (Ligurian Sea, North-western Mediterranean). Geochimica et
- 939 Cosmochimica Acta 74, 5549–5559. https://doi.org/10.1016/j.gca.2010.06.036
- Hirst, A.G., Kiørboe, T., 2002. Mortality of marine planktonic copepods: global rates and
 patterns. Marine Ecology Progress Series 230, 195–209.
- 942 https://doi.org/10.3354/meps230195
- 943 Hunt, B.P.V., Carlotti, F., Donoso, K., Pagano, M., D'Ortenzio, F., Taillandier, V., Conan, P.,
- 2017. Trophic pathways of phytoplankton size classes through the zooplankton food webover the spring transition period in the north-west Mediterranean Sea. Journal of
- 946 Geophysical Research: Oceans 122, 6309–6324. https://doi.org/10.1002/2016JC012658
- 947 Im, D.-H., Wi, J.H., Suh, H.-L., 2015. Evidence for ontogenetic feeding strategies in four
- 948 calanoid copepods in the East Sea (Japan Sea) in summer, revealed by stable isotope
- 949 analysis. Ocean Science Journal 50, 481–490. https://doi.org/10.1007/s12601-015-0044-y
- 950 Jónasdóttir, S.H., 2019. Fatty Acid Profiles and Production in Marine Phytoplankton. Marine
- 951 Drugs 17, 151. https://doi.org/10.3390/md17030151

- 952 Karsh, K.L., Trull, T.W., Lourey, M.J., Sigman, D.M., 2003. Relationship of nitrogen isotope
- 953 fractionation to phytoplankton size and iron availability during the Southern Ocean Iron
 954 RElease Experiment (SOIREE). Limnology and Oceanography 48, 1058–1068.

955 https://doi.org/10.4319/lo.2003.48.3.1058

956 Khammeri, Y., Bellaaj-Zouari, A., Hamza, A., Medhioub, W., Sahli, E., Akrout, F., Barraj,

957 N., Ben Kacem, M.Y., Bel Hassen, M., 2020. Ultraphytoplankton community

- 958 composition in Southwestern and Eastern Mediterranean Basin: Relationships to water
- mass properties and nutrients. Journal of Sea Research 158, 101875.

960 https://doi.org/10.1016/j.seares.2020.101875

961 Khammeri, Y., Hamza, I.S., Zouari, A.B., Hamza, A., Sahli, E., Akrout, F., Ben Kacem,

962 M.Y., Messaoudi, S., Hassen, M.B., 2018. Atmospheric bulk deposition of dissolved

963 nitrogen, phosphorus and silicate in the Gulf of Gabès (South Ionian Basin); implications

- 964 for marine heterotrophic prokaryotes and ultraphytoplankton. Continental Shelf Research
- 965 159, 1–11. https://doi.org/10.1016/j.csr.2018.03.003
- Kim, Y., Lee, Jang Han, Kang, J.J., Lee, Jae Hyung, Lee, H.W., Kang, C.K., Lee, S.H., 2019.
 River discharge effects on the contribution of small-sized phytoplankton to the total
 biochemical composition of POM in the Gwangyang Bay, Korea. Estuarine, Coastal and

969 Shelf Science 226, 106293. https://doi.org/10.1016/j.ecss.2019.106293

970 Kondoh, M., Ninomiya, K., 2009. Food-chain length and adaptive foraging. Proceedings of

the Royal Society B: Biological Sciences 276, 3113–3121.

- 972 https://doi.org/10.1098/rspb.2009.0482
- Weikert, H., Lahajnar, N., 2003. Vertical distribution of mesozooplankton
 and its δ15N signature at a deep-sea site in the Levantine Sea (eastern Mediterranean) in

975 April 1999. Journal of Geophysical Research: Oceans 108.

976 https://doi.org/10.1029/2002JC001351

977 Koppelmann, R., Böttger-Schnack, R., Möbius, J., Weikert, H., 2009. Trophic relationships of

- 200 zooplankton in the eastern Mediterranean based on stable isotope measurements. Journal
 of Plankton Research 31, 669–686. https://doi.org/10.1093/plankt/fbp013
- 980 Kozak, E.R., Franco-Gordo, C., Godínez-Domínguez, E., Suárez-Morales, E., Ambriz-
- 981 Arreola, I., 2020. Seasonal variability of stable isotope values and niche size in tropical
- calanoid copepods and zooplankton size fractions. Marine Biology 167, 37.
- 983 https://doi.org/10.1007/s00227-020-3653-7
- Krasakopoulou, E., Karageorgis, A.P., 2005. Spatial and temporal distribution patterns of
 suspended particulate matter and particulate organic carbon in the Saronikos Gulf

- 986 (eastern Mediterranean, Greece). Geo-Marine Letters 25, 343–359.
- 987 https://doi.org/10.1007/s00367-005-0007-x
- Kürten, B., Painting, S.J., Struck, U., Polunin, N.V.C., Middelburg, J.J., 2013. Tracking
 seasonal changes in North Sea zooplankton trophic dynamics using stable isotopes.
- 990 Biogeochemistry 113, 167–187. https://doi.org/10.1007/s10533-011-9630-y
- 291 Lam, P.J., Ohnemus, D.C., Auro, M.E., 2015. Size-fractionated major particle composition
- and concentrations from the US GEOTRACES North Atlantic Zonal Transect. Deep Sea
 Research Part II: Topical Studies in Oceanography, 116, 303–320.
- 994 https://doi.org/10.1016/j.dsr2.2014.11.020
- Laws, E.A., Popp, B.N., Bidigare, R.R., Kennicutt, M.C., Macko, S.A., 1995. Dependence of
- 996 phytoplankton carbon isotopic composition on growth rate and [CO2]aq: Theoretical
- 997 considerations and experimental results. Geochimica et Cosmochimica Acta 59, 1131–

998 1138. https://doi.org/10.1016/0016-7037(95)00030-4

- 999 Leblanc, K., Quéguiner, B., Diaz, F., Cornet, V., Michel-Rodriguez, M., Durrieu de Madron,
- 1000 X., Bowler, C., Malviya, S., Thyssen, M., Grégori, G., Rembauville, M., Grosso, O.,
- Poulain, J., de Vargas, C., Pujo-Pay, M., Conan, P., 2018. Nanoplanktonic diatoms are
 globally overlooked but play a role in spring blooms and carbon export. Nature
 Communications 9, 953. https://doi.org/10.1038/s41467-018-03376-9

1004 Levinsen, H., Turner, J.T., Nielsen, T.G., Hansen, B.W., 2000. On the trophic coupling

- between protists and copepods in arctic marine ecosystems. Marine Ecology Progress
 Series 204, 65–77. https://doi.org/10.3354/meps204065
- Liénart, C., Savoye, N., Bozec, Y., Breton, E., Conan, P., David, V., Feunteun, E., Grangeré,
 K., Kerhervé, P., Lebreton, B., Lefebvre, S., L'Helguen, S., Mousseau, L., Raimbault, P.,
- 1009 Richard, P., Riera, P., Sauriau, P.-G., Schaal, G., Aubert, F., Aubin, S., Bichon, S.,
- 1010 Boinet, C., Bourasseau, L., Bréret, M., Caparros, J., Cariou, T., Charlier, K., Claquin, P.,
- 1011 Cornille, V., Corre, A.-M., Costes, L., Crispi, O., Crouvoisier, M., Czamanski, M., Del
- 1012 Amo, Y., Derriennic, H., Dindinaud, F., Durozier, M., Hanquiez, V., Nowaczyk, A.,
- 1013 Devesa, J., Ferreira, S., Fornier, M., Garcia, F., Garcia, N., Geslin, S., Grossteffan, E.,
- 1014 Gueux, A., Guillaudeau, J., Guillou, G., Joly, O., Lachaussée, N., Lafont, M.,
- 1015 Lamoureux, J., Lecuyer, E., Lehodey, J.-P., Lemeille, D., Leroux, C., Macé, E., Maria,
- 1016 E., Pineau, P., Petit, F., Pujo-Pay, M., Rimelin-Maury, P., Sultan, E., 2017. Dynamics of
- 1017 particulate organic matter composition in coastal systems: A spatio-temporal study at
- 1018 multi-systems scale. Progress in Oceanography 156, 221–239.
- 1019 https://doi.org/10.1016/j.pocean.2017.03.001

1020 Liénart, C., Savoye, N., David, V., Ramond, P., Rodriguez Tress, P., Hanquiez, V., Marieu, 1021 V., Aubert, F., Aubin, S., Bichon, S., Boinet, C., Bourasseau, L., Bozec, Y., Bréret, M., 1022 Breton, E., Caparros, J., Cariou, T., Claquin, P., Conan, P., Corre, A.-M., Costes, L., 1023 Crouvoisier, M., Del Amo, Y., Derriennic, H., Dindinaud, F., Duran, R., Durozier, M., 1024 Devesa, J., Ferreira, S., Feunteun, E., Garcia, N., Geslin, S., Grossteffan, E., Gueux, A., 1025 Guillaudeau, J., Guillou, G., Jolly, O., Lachaussée, N., Lafont, M., Lagadec, V., 1026 Lamoureux, J., Lauga, B., Lebreton, B., Lecuyer, E., Lehodey, J.-P., Leroux, C., L'Helguen, S., Macé, E., Maria, E., Mousseau, L., Nowaczyk, A., Pineau, P., Petit, F., 1027 1028 Pujo-Pay, M., Raimbault, P., Rimmelin-Maury, P., Rouaud, V., Sauriau, P.-G., Sultan, E., Susperregui, N., 2018. Dynamics of particulate organic matter composition in coastal 1029 systems: Forcing of spatio-temporal variability at multi-systems scale. Progress in 1030 Oceanography 162, 271–289. https://doi.org/10.1016/j.pocean.2018.02.026 1031 Ljungström, G., Claireaux, M., Fiksen, Ø., Jørgensen, C., 2020. Body size adaptions under 1032 climate change: zooplankton community more important than temperature or food 1033 abundance in model of a zooplanktivorous fish. Marine Ecology Progress Series 636, 1-1034 1035 18. https://doi.org/10.3354/meps13241 1036 Lowry, O.H., Rosebrough, N.J., Farr, A.L., Randall, R.J., 1951. Protein measurement with the 1037 Folin phenol reagent. Journal of Biological Chemistry 193, 265-275. 1038 Maguer, J.-F., L'helguen, S., Le Corre, P., 2000. Nitrogen Uptake by Phytoplankton in a 1039 Shallow Water Tidal Front. Estuarine, Coastal and Shelf Science 51, 349-357. https://doi.org/10.1006/ecss.2000.0678 1040 1041 Major, Y., Kifle, D., Niedrist, G.H., Sommaruga, R., 2017. An isotopic analysis of the 1042 phytoplankton-zooplankton link in a highly eutrophic tropical reservoir dominated by 1043 cyanobacteria. Journal of Plankton Research 39, 220-231. 1044 https://doi.org/10.1093/plankt/fbx007 1045 Mason, R.P., Reinfelder, J.R., Morel, F.M.M., 1995. Bioaccumulation of mercury and methylmercury. Water Air Soil Pollut 80, 915-921. https://doi.org/10.1007/BF01189744 1046 1047 Mauchline, J., 1998. The biology of Calanoid copepods. In: Blaxter, J.H.S., Southward, A.J., Tyler, P.A.(Eds.), Advances in Marine Biology, 245–292. Academic Press, San Diego, 1048 1049 USA. Mayot, N., D'Ortenzio, F., Uitz, J., Gentili, B., Ras, J., Vellucci, V., Golbol, M., Antoine, D., 1050 1051 Claustre, H., 2017. Influence of the Phytoplankton Community Structure on the Spring and Annual Primary Production in the Northwestern Mediterranean Sea. Journal of 1052 Geophysical Research: Oceans 122, 9918–9936. https://doi.org/10.1002/2016JC012668 1053

- Mayzaud, P., Chanut, J., Ackman, R., 1989. Seasonal changes of the biochemical composition
 of marine particulate matter with special reference to fatty acids and sterols. Marine
- 1056 Ecology Progress Series 56, 189–204. https://doi.org/10.3354/meps056189
- Montoya, J.P., Carpenter, E.J., Capone, D.G., 2002. Nitrogen fixation and nitrogen isotope
 abundances in zooplankton of the oligotrophic North Atlantic. Limnology and
 Oceanography 47, 1617–1628. https://doi.org/10.4319/lo.2002.47.6.1617
- 1060 Morris, M.J., Hopkins, T.L., 1983. Biochemical composition of crustacean zooplankton from
- the eastern Gulf of Mexico. Journal of Experimental Marine Biology and Ecology 69, 1–
 1062 19.
- Motwani, N.H., Gorokhova, E., 2013. Mesozooplankton Grazing on Picocyanobacteria in the
 Baltic Sea as Inferred from Molecular Diet Analysis. PLOS ONE 8, e79230.
- 1065 https://doi.org/10.1371/journal.pone.0079230
- 1066 Onodera, T., Kanaya, G., Hatamoto, M., Kohzu, A., Iguchi, A., Takimoto, Y., Yamaguchi, T.,
 1067 Mizuochi, M., Syutsubo, K., 2018. Evaluation of trophic transfer in the microbial food
 1068 web during sludge degradation based on 13C and 15N natural abundance. Water
 1069 Research 146, 30–36. https://doi.org/10.1016/j.watres.2018.09.016
- Ostrom, P. H., Fry, B., 1993. Sources and cycling of organic matter within modern and
 prehistoric food webs. Organic geochemistry, 785-798. Springer, Boston, MA.
- 1072 Pantoja, S., Repeta, D.J., Sachs, J.P., Sigman, D.M., 2002. Stable isotope constraints on the
- 1073 nitrogen cycle of the Mediterranean Sea water column. Deep Sea Research Part I:
- 1074 Oceanographic Research Papers 49, 1609–1621. https://doi.org/10.1016/S09671075 0637(02)00066-3
- Parnell, A.C., Phillips, D.L., Bearhop, S., Semmens, B.X., Ward, E.J., Moore, J.W., Jackson,
 A.L., Grey, J., Kelly, D.J., Inger, R., 2013. Bayesian stable isotope mixing models.
- 1078 Environmetrics 24, 387–399. https://doi.org/10.1002/env.2221
- Peterson, B.J., Howarth, R.W., Garritt, R.H., 1985. Multiple stable isotopes used to trace the
 flow of organic matter in estuarine food webs. Science 227, 1361–1363.
- 1081 https://doi.org/10.1126/science.227.4692.1361
- 1082 Phillips, D.L., Inger, R., Bearhop, S., Jackson, A.L., Moore, J.W., Parnell, A.C., Semmens,
- 1083 B.X., Ward, E.J., 2014. Best practices for use of stable isotope mixing models in food-
- 1084 web studies. Canadian Journal of Zoology 92, 823–835. https://doi.org/10.1139/cjz-2014-
- 1085 0127

- Pinnegar, J.K., Polunin, N.V.C., 1999. Differential fractionation of δ13C and δ15N among
 fish tissues: implications for the study of trophic interactions. Functional Ecology 13,
- 1088 225–231. https://doi.org/10.1046/j.1365-2435.1999.00301.x
- Post, D.M., 2002. Using Stable Isotopes to Estimate Trophic Position: Models, Methods, and
 Assumptions. Ecology 83, 703–718. https://doi.org/10.1890/0012-
- 1091 9658(2002)083[0703:USITET]2.0.CO;2
- 1092 Postel, L., Fock, H., Hagen, W., 2000. 4 Biomass and abundance, In: Harris, R., Wiebe, P.,
- 1093 Lenz, J., Skjoldal, H.R., Huntley, M. (Eds.), ICES Zooplankton Methodology Manual.
- Academic Press, London, pp. 83–192. https://doi.org/10.1016/B978-012327645-2/50005 0
- 1096 Raimbault, P., Diaz, F., Pouvesle, W., Boudjellal, B., 1999. Simultaneous determination of
 1097 particulate organic carbon, nitrogen and phosphorus collected on filters, using a semi-

automatic wet-oxidation method. Marine Ecology Progress Series 180, 289.

1099 Raimbault, P., Garcia, N., Cerutti, F., 2008. Distribution of inorganic and organic nutrients in

- the South Pacific Ocean; evidence for long-term accumulation of organic matter in
 nitrogen-depleted waters. Biogeosciences 5, 281–298. https://doi.org/10.5194/bg-5-2812008
- 1103 Rau, G., Teyssie, J., Rassoulzadegan, F., Fowler, S., 1990. C-13/C-12 and N-15/N-14
- variations among size-fractionated marine particles implications for their origin and
 trophic relationships. Marine Ecology Progress Series 59(1/2), 33–38.

1106 Ríos, A.F., Fraga, F., Pérez, F.F., Figueiras, F.G., 1998. Chemical composition of

- phytoplankton and Particulate Organic Matter in the Ría de Vigo (NW Spain). Scientia
 Marina 62 (3), 257–271 https://doi.org/10.3989/scimar.1998.62n3257
- 1109 Rolff, C., 2000. Seasonal variation in δ 13C and δ 15N of size-fractionated plankton at a
- 1110 coastal station in the northern Baltic proper. Marine Ecology Progress Series 203, 47–65.
 1111 https://doi.org/10.3354/meps203047
- 1112 Ryther, J.H., 1969. Photosynthesis and Fish Production in the Sea. Science 166, 72–76.
- 1113 https://doi.org/10.1126/science.166.3901.72
- 1114 Schartup, A.T., Qureshi, A., Dassuncao, C., Thackray, C.P., Harding, G., Sunderland, E.M.,
- 1115 2018. A Model for Methylmercury Uptake and Trophic Transfer by Marine Plankton.
- 1116 Environmental Science & Technology 52, 654–662.
- 1117 https://doi.org/10.1021/acs.est.7b03821

- 1118 Sommer, F., Stibor, H., Sommer, U., Velimirov, B., 2000. Grazing by mesozooplankton from
- Kiel Bight, Baltic Sea, on different sized algae and natural seston size fractions. Marine
 Ecology Progress Series 199, 43–53. https://doi.org/10.3354/meps199043
- 1121 Sommer, U., Stibor, H., Katechakis, A., Sommer, F., Hansen, T., 2002. Pelagic food web
- 1122 configurations at different levels of nutrient richness and their implications for the ratio
- fish production:primary production, in: Vadstein, O., Olsen, Y. (Eds.), Sustainable
- 1124Increase of Marine Harvesting: Fundamental Mechanisms and New Concepts. Springer
- 1125 Netherlands, Dordrecht, pp. 11–20. https://doi.org/10.1007/978-94-017-3190-4_2
- Søreide, J.E., Hop, H., Carroll, M.L., Falk-Petersen, S., Hegseth, E.N., 2006. Seasonal food
 web structures and sympagic–pelagic coupling in the European Arctic revealed by stable
 isotopes and a two-source food web model. Progress in Oceanography 71, 59–87.
- 1129 https://doi.org/10.1016/j.pocean.2006.06.001
- 1130 Tamelander, T., Kivimäe, C., Bellerby, R.G.J., Renaud, P.E., Kristiansen, S., 2009. Base-line
- variations in stable isotope values in an Arctic marine ecosystem: effects of carbon and
 nitrogen uptake by phytoplankton. Hydrobiologia 630, 63–73.
- 1133 https://doi.org/10.1007/s10750-009-9780-2
- 1134 Tao, Y., Yu, J., Liu, X., Xue, B., Wang, S., 2018. Factors affecting annual occurrence,
- bioaccumulation, and biomagnification of polycyclic aromatic hydrocarbons in plankton
- food webs of subtropical eutrophic lakes. Water Research 132, 1–11.
- 1137 https://doi.org/10.1016/j.watres.2017.12.053
- 1138 Tedetti, M., Tronczynski, J., 2019. HIPPOCAMPE cruise, RV Antea.
- 1139 https://doi.org/10.17600/18000900
- 1140 Tedetti, M., Tronczynski, J., Carlotti, J.F., Pagano, M., Ben Ismail, S., Sammari, C., Bel
- 1141 Hassen, M., Desboeufs, K., Poindron, C., Chifflet, S., Bellaaj Zouari, A., Abdennadher,
- 1142 M., Amri, S., Bănaru, D., Ben Abdallah, L., Bhairy, N., Boudriga, I., Bourin, A., Brach-
- 1143 Papa, C., Briant, N., Cabrol, L., Chevalier, C., Chouba, L., Coudray, S., Daly Yahia, M.
- 1144 N., de Garidel Thoron, T., Dufour, A., Dutay, J-C., Espinasse, B., Fierro-González, P.,
- 1145 Fornier, M., Garcia, N., Giner, F., Guigue, C., Guilloux, L., Hamza, A., Heimbürger-
- 1146 Boavida, L-E., Jacquet, S., Knoery, J., Lajnef, R., Makhlouf Belkahia, N., Malengros, D.,
- 1147 Martinot, P.L., Bosse, A., Mazur, J-C., Meddeb, M., Misson, B., Pringault, O.,
- 1148 Quéméneur, M., Radakovitch, O., Raimbault, P., Ravel, R., Rossi, R., Rwawi, C., Sakka
- 1149 Hlaili, A., Tesán-Onrubia, J.A., Thomas, B., Thyssen, M., Zaaboub, N., Garnier, C.,
- 1150 2023. Contamination of planktonic food webs in the Mediterranean Sea: Setting the

- 1151 frame for the MERITE-HIPPOCAMPE oceanographic cruise (spring 2019). Marine
- 1152 Pollution Bulletin. In press in this special issue.
- 1153 Tiano, M., Tronczyński, J., Harmelin-Vivien, M., Tixier, C., Carlotti, F., 2014. PCB
- 1154 concentrations in plankton size classes, a temporal study in Marseille Bay, Western
- 1155 Mediterranean Sea. Marine Pollution Bulletin 89, 331–339.
- 1156 https://doi.org/10.1016/j.marpolbul.2014.09.040
- 1157 Tiselius, P., Fransson, K., 2016. Daily changes in δ 15N and δ 13C stable isotopes in copepods:
- equilibrium dynamics and variations of trophic level in the field. Journal of Plankton
 Research 38, 751–761. https://doi.org/10.1093/plankt/fbv048
- 1160 Trimble, S.M., Baskaran, M., 2005. The role of suspended particulate matter in 234Th
- scavenging and 234Th-derived export fluxes of POC in the Canada Basin of the Arctic

1162 Ocean. Marine Chemistry 96, 1–19. https://doi.org/10.1016/j.marchem.2004.10.003

- 1163 Vander Zanden, M.J.V., Rasmussen, J.B., 2001. Variation in δ 15N and δ 13C trophic
- 1164 fractionation: Implications for aquatic food web studies. Limnology and Oceanography
 1165 46, 2061–2066. https://doi.org/10.4319/lo.2001.46.8.2061
- Wainright, S.C., Fry, B., 1994. Seasonal variation of the stable isotopic compositions of
 coastal marine plankton from Woods Hole, Massachusetts and Georges Bank. Estuaries
 17, 552–560. https://doi.org/10.2307/1352403
- 1169 Wu, Y., Wang, W.-X., 2011. Accumulation, subcellular distribution and toxicity of inorganic
- mercury and methylmercury in marine phytoplankton. Environmental Pollution, Nitrogen
 Deposition, Critical Loads and Biodiversity 159, 3097–3105.

1172 https://doi.org/10.1016/j.envpol.2011.04.012

- Yang, G., Li, C., Guilini, K., Wang, X., Wang, Y., 2017. Regional patterns of δ13C and δ15N
 stable isotopes of size-fractionated zooplankton in the western tropical North Pacific
- 1175 Ocean. Deep Sea Research Part I: Oceanographic Research Papers 120, 39–47.
 1176 https://doi.org/10.1016/j.dsr.2016.12.007
- 1177 Yılmaz, A.Z., Besiktepe, S., 2010. Annual variations in biochemical composition of size
- 1178 fractionated particulate matter and zooplankton abundance and biomass in Mersin Bay,
- 1179 NE Mediterranean Sea. Journal of Marine Systems 81, 260–271.
- 1180 https://doi.org/10.1016/j.jmarsys.2010.01.002

1182 Figure captions

1183

Figure 1. Location of the ten sampling stations of the MERITE-HIPPOCAMPE campaign in the Mediterranean
Sea (April-Mai 2019).

1186

1187 **Figure 2.** Boxplot of the concentrations of A) proteins, B) carbohydrates, C) lipids (µg mg⁻¹ DW) and D) energy 1188 content (E_i) (kJ g⁻¹ DW) in the different plankton size-fractions (fractions between 0.7 and 60 μ m in green and > 1189 $60 \,\mu\text{m}$ in orange) for all stations combined. H = Kruskal–Wallis non-parametric test and the associated p-value 1190 for the respective biochemical compounds: H = 43.5, p < 0.0001; H = 110.6, p < 0.001; H = 41.8, p < 0.0001 and 1191 H = 51.7, p < 0.0001. Mean values with different letters are significantly different (p < 0.05). The mean and 1192 median values are represented by a cross and a horizontal line, respectively, and the box length is defined as the 1193 interquartile range. The minimum and maximum values are represented by whiskers. Mean values with different 1194 post-hoc letters are significantly different (p < 0.05).

1195

1196 Figure 3. Boxplot of the concentrations of A, B) proteins, C, D) carbohydrates, E, F) lipids ($\mu g m g^{-1} DW$) and 1197 G, H) energy content (E_i) (kJ g⁻¹ DW) in the phyto-(fractions between 0.7 and 60 μ m – in green) and 1198 zooplankton (fractions between 60 and 500 μ m – in orange) fractions for each station. H = Kruskal–Wallis non-1199 parametric test and the associated p-value for the respective biochemical compounds for phytoplankton (H =43.4, p < 0.0001; H = 30.6, p < 0.0001; H = 34.9, p < 0.0001 and H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H = 42.9, p < 0.0001) and for zooplankton (H1200 1201 = 43.8, p < 0.0001; H = 22.9, p < 0.006; H = 38.8, p < 0.0001 and H = 38.8, p < 0.0001). Stations are grouped 1202 by geographical area: St1, St2, St3 and St4 for the Northern coast, St9, St10 and St11 for the offshore area, and 1203 St15, St17 and St19 for the Southern coast. The mean and median values were represented by a cross and a 1204 horizontal line, respectively, and the box length is defined as the interquartile range. The minimum and 1205 maximum values are represented by whiskers. Mean values with different post-hoc letters are significantly 1206 different (p < 0.05).

1207

Figure 4. Total amount of energy provided by plankton (E_T in kJ m⁻³) for each of the size-fractions by station.
Stations are grouped by geographical area: St1, St2, St3 and St4 for the Northern coast, St9, St10 and St11 for the offshore area, and St15, St17 and St19 for the Southern coast.

1211

1212 Figure 5. Mean (\pm standard error, SE) stable isotope compositions (δ^{13} C and δ^{15} N values, ‰) in the different 1213 plankton size-fractions for all stations combined. Green dots correspond to the phytoplankton size-fractions

- 1214 (from 0.7 to 60 μ m), while the orange dots correspond to zooplankton size-fractions (> 60 μ m).
- 1215

1216 Figures 6. Boxplot of the A, B) δ^{13} C and C, D) δ^{15} N values (‰) in the phyto-(fractions between 0.7 and 60 µm

- $1217 \qquad \text{ in green) and zooplankton (fractions between 60 and 500\,\mu\text{m} \text{ in orange) fractions for each station. Stations}$
- 1218 are grouped by geographical area: St1, St2, St3 and St4 for the Northern coast, St9, St10 and St11 for the
- 1219 offshore area, and St15, St17 and St19 for the Southern coast. Kruskall-Wallis non-parametric test were
- 1220 performed for A) (H = 25.5, p < 0.05), C) (H = 24.1, p < 0.05) and D) (H = 33.5, p < 0.0001), and one-way

1221ANOVA test was performed for B) (F = 4.4, p < 0.0001). The mean and median values are represented by a</td>1222cross and a horizontal line, respectively, and the box length is defined as the interquartile range. The minimum1223and maximum values are represented by whiskers. Mean values with different post-hoc letters are significantly1224different (p < 0.05).</td>

1225

1226Figure 7. Boxplot of the trophic levels (TL) of the zooplankton size-fractions (60 to > 2000 μ m). The median1227value is represented by a horizontal line and the box length is defined as the interquartile range. The minimum1228and maximum values are represented by whiskers. One-way ANOVA test (F = 5.17, p = 0.001). Superscript1229letters represent Newman-Keuls post-hoc groups and values with the different letters are significantly different1230(p < 0.05).</td>

1231

1232 Figure 8. Trophic levels (TL) of the zooplankton size-fractions ($60 \text{ to} > 2000 \,\mu\text{m}$) and mean TL weighted by

 $\label{eq:their biomass} 1233 \qquad \text{their biomass} (TL_B) \text{ for each station. Stations are gathered by geographical area: } St1, St2, St3 and St4 \text{ for the}$

1234 Northern coast, St9, St10 and St11 for the offshore area, and St15, St17 and St19 for the Southern coast.

1235

1236 Figure 9. Relative contributions of $0.7-2.7 \ \mu m$ (pico-), $2.7-20 \ \mu m$ (nano-), and $20-60 \ \mu m$ (microplankton) as

1237 food resources for three zooplankton size-fractions. Arrows indicate the trophic transfer between pico-POM

- 1238 nano-POM, and micro-POM fractions and zooplankton size fractions: $60-500 \,\mu$ m (in green), $500-2000 \,\mu$ m (in 1239 blue) and > 2000 μ m (in orange). Their thickness is proportional to relative contribution in percentage, indicated 1240 close to the corresponding arrow.
- 1241

1242 **Table 1.** Suspended particulate matter and plankton dry weight (SPM $_{DW}$ and Plankton $_{DW}$, in mg DW L^{-1}) for the

1243 different plankton size-fractions and stations, as well as total chlorophyll a concentration (TChla, µg L⁻¹) for the

1244 > 0.7- μ m fraction. Kruskal-Wallis test and the associated p-value for SPM_{DW} (H = 23.15, p < 0.0001) and

1245 Plankton_{DW} (H = 28.51, p < 0.0001). Superscript letters represent rank comparison groups and values with

 $1246 \qquad different \, letters \, are \, significantly \, different \, (p < 0.05). \, St = station, SE = standard \, -error, n = number \, of \, samples.$

1248 Figures

















Figure 6







Figure 9

1277 Tables

1278 Table 1.

Stations	St1	St2	St3	St4	St9	St10	St11	St15	St17	St19	n	$Mean \pm SE$				
Size-fractions	SPM _{DW} (mg DW L ⁻¹)															
$0.7-2.7\mu m$	0.11	0.23	0.09	0.11	0.14	0.12	0.06	0.05	0.07	0.16	10	0.11	±	0.02	ab	
$2.7-20\mu m$	0.24	0.09	0.12	0.11	0.53	0.13	0.09	0.07	0.23	0.17	10	0.18	±	0.04	bc	
$20-60\mum$	0.03	0.01	0.00	0.11	0.01	0.01	0.01	0.01	0.11	0.11	10	0.04	±	0.02	a	
$0.7-60\mu m$	0.35	0.32	0.21	0.22	0.67	0.25	0.15	0.12	0.29	0.34	10	0.29	±	0.05	с	
TChla (> 0.7	0.77	0.29	0.69	0.08	154	0.55	0.28	0.67	0.21	1 45	10	0.76		0.14		
µm) (µg L ⁻¹)	0.77	0.38	0.08	0.98	1.54	0.55	0.38	0.07	0.21	1.45	10	0.70	Ξ	0.14		
		Plankton _{DW} (mg DW L ⁻¹)														
$60-200\mu m$	0.020	0.007	0.002	0.068	0.017	0.003	0.004	0.006	0.083	0.029	10	0.024	±	0.027	ab	
$200500\mu\text{m}$	0.037	0.007	0.002	0.028	0.023	0.004	0.005	0.010	0.072	0.038	10	0.023	±	0.021	ab	
$5001000\mu\text{m}$	0.005	0.007	0.004	0.007	0.004	0.002	0.009	0.009	0.002	0.011	10	0.006	±	0.003	b	
$1000-2000\mum$	0.002	0.005	0.007	0.002	0.0002	0.001	0.002	0.001		0.0000	10	0.002	±	0.002	а	
$>2000\mum$	0.0004	0.005	0.013			0.0001	0.0004	0.001		0.0002	10	0.003	±	0.005	a	
Sum																
$60 -> 2000 \mum$	0.064	0.031	0.028	0.105	0.044	0.011	0.020	0.026	0.157	0.078	50	0.057	±	0.019		
1279																