The suspended small-particles layer in the poorly-oxygenated Black 1

Sea: a proxy for delineating the effective N₂-yielding section 2

3 Rafael Rasse¹, Hervé Claustre¹, and Antoine Poteau¹

4 ¹Sorbonne Université and CNRS, Laboratoire d'Océanographie de Villefranche (LOV) UMR7093, Institut de la Mer de 5 6 Villefranche (IMEV), 06230, Villefranche-sur-Mer, France.

7 Correspondence to: rafael.rasse@obs-vlfr.fr; rjrasse@gmail.com

8 Abstract. The shallower poorly-oxygenated water masses of the ocean confine a majority of the microbial communities that 9 can produce up to 90% of oceanic N2. This effective N2-yielding section encloses a suspended small-particle layer, inferred 10 from particle backscattering (b_{bp}) measurements. It is thus hypothesized that this layer (hereafter, the b_{bp} -layer) is linked to 11 microbial communities involved in N₂-yielding such as nitrate-reducing SAR11 as well as sulphur-oxidizing, anammox and 12 denitrifying bacteria — a hypothesis yet to be evaluated. Here, data collected by three BGC-Argo floats deployed in the Black 13 Sea are used to investigate the origin of this b_{bp} -layer. To this end, we evaluate how the key drivers of N₂-yielding bacteria 14 dynamics impact on the vertical distribution of b_{bp} and the thickness of the b_{bp} -layer. In conjunction with published data on N₂ 15 excess, our results suggest that the b_{bp} -layer is at least partially composed of the bacteria driving N₂ yielding for three main 16 reasons: (1) strong correlations are recorded between b_{bp} and nitrate; (2) the top location of the b_{bp} -layer is driven by the 17 ventilation of oxygen-rich subsurface waters, while its thickness is modulated by the amount of nitrate available to produce 18 N_2 ; (3) the maxima of both b_{bp} and N_2 excess coincide at the same isopycnals where bacteria involved in N_2 yielding coexist. 19 We thus advance that b_{bp} and O_2 can be exploited as a combined proxy to delineate the N₂-yielding section of the Black Sea. 20 This proxy can potentially contribute to refining delineation of the effective N₂-yielding section of oxygen-deficient zones via 21 data from the growing BGC-Argo float network.

22 **1** Introduction

23 **Poorly-oxygenated** water masses ($O_2 < 3 \mu M$) host the microbial communities that produce between 20-40% of oceanic N_2 24 mainly via heterotrophic denitrification and anaerobic oxidation of ammonium (Gruber and Sarmiento, 1997; Devries et al. 25 2013; Ward 2013). The shallower poorly-oxygenated water masses (\sim 50-200 m) are the most effective N₂-producing section 26 because this is where the microbial communities that condition the process mainly develop and generate up to 90% of the N_2 27 (Ward et al., 2009; Dalsgaard et al., 2012; Babin et al., 2014). These microbial communities include nitrate-reducing SAR11, 28 and anammox, denitrifying, and sulphur-oxidizing bacteria (e.g. Canfield et al., 2010; Ulloa et al. 2012; Ward 2013; Callbeck 29 et al., 2018). It is thus important to unravel the biogeochemical parameters that trigger the accumulation of such bacteria in 30 the ocean's poorly-oxygenated water masses. This information is crucial for understanding and quantifying how bacterial 31 biomass and related N₂ yielding can respond to the ongoing expansion of oceanic regions with low oxygen (Keeling and 32 Garcia, 2002; Stramma et al., 2008; Helm et al., 2011; Schmidtko et al., 2017). Ultimately, greater accuracy in this domain 33 can contribute to improving mechanistic predictions on how such expansion affects the oceans' role in driving the Earth's 34 climate by sequestering atmospheric carbon dioxide (e.g. Oschlies et al., 2018).

35 In poorly-oxygenated water masses, the biogeochemical factors that can affect the abundance of denitrifying and anammox 36 bacteria are the levels of O_2 , organic matter (OM), nitrate (NO₃⁻), ammonium (NH₄⁺), and hydrogen sulfide (H₂S) (Murray et al., 1995; Ward et al., 2008; Dalsgaard et al., 2014; Bristow et al., 2016). Therefore, to elucidate what triggers the confinement
 of such bacteria, we need to investigate how the above biogeochemical factors drive their vertical distribution, with high
 temporal and vertical resolution. To this end, we should develop multidisciplinary approaches that allow us to permanently
 monitor the full range of biogeochemical variables of interest in poorly-oxygenated water masses.

41 Optical proxies of tiny particles can be applied as an alternative approach to assess the vertical distribution of N₂-yielding 42 microbial communities in poorly-oxygenated water masses (Naqvi et al., 1993). For instance, nitrate-reducing SAR11, and 43 anammox, denitrifying, and sulphur-oxidizing bacteria are found as free-living bacteria (0.2-2 μ m), and can be associated with 44 small suspended (> 2-30 μ m), and large sinking (> 30 μ m) particles (Fuchsman et al., 2011, 2012a, 2017; Ganesh et al., 2014, 45 2015). Therefore, particle backscattering (*b_{bp}*), a proxy for particles in the ~0.2-20 μ m size range (Stramski et al., 1999, 2004; 46 Organelli et al., 2018), can serve to detect the presence of these free-living bacteria and those associated with small suspended 47 particles.

48 Time series of b_{bp} acquired by biogeochemical Argo (BGC-Argo) floats highlight the presence of a permanent layer of 49 suspended small particles in shallower poorly-oxygenated water masses $(b_{bp}$ -layer) (Whitmire et al., 2009; Wojtasiewicz et 50 al., 2018). It has been hypothesized that this b_{bp} -layer is linked to N₂-yielding microbial communities such as nitrate-reducing 51 SAR11, and denitrifying, anammox, and sulphur-oxidizing bacteria. However, this hypothesis has not yet been clearly 52 demonstrated. To address this, the first step is to evaluate: (1) potential correlations between the biogeochemical factors that 53 control the presence of the *b_{bp}-layer* and such arrays of bacteria (O₂, NO₃⁻, OM, H₂S;, Murray et al., 1995; Ward et al., 2008; 54 Fuchsman et al., 2011; Ulloa et al., 2012; Dalsgaard et al., 2014; Bristow et al., 2016), and (2) the possible relationship between 55 the b_{bp} -layer and N₂ produced by microbial communities.

This first step is thus essential for identifying the origin of the b_{bp} -layer and, ultimately, determining if BGC-Argo observations of b_{bp} can be implemented to delineate the poorly-oxygenated water masses where such bacteria are confined. The Black Sea appears as a suitable area for probing into the origin of the b_{bp} -layer in low-oxygen waters in this way. It is indeed a semienclosed basin with permanently low O₂ levels where N₂ production and related nitrate-reducing SAR11, and denitrifying and anammox bacteria are mainly confined within a well-defined poorly-oxygenated zone (Kuypers et al., 2003; Konovalov et al., 2005; Kirkpatrick et al., 2012). In addition, a permanent b_{bp} -layer is a typical characteristic of this region, which is linked to such microbial communities and inorganic particles (Stanev et al., 2017, 2018, see details in section 2.0).

63 The goal of our study is therefore to investigate the origin of the b_{bp} -layer in the poorly-oxygenated waters of the Black Sea 64 using data collected by BGC-Argo floats. More specifically, we aim to evaluate, within the poorly-oxygenated zone, how: (1) 65 two of the main factors (O_2 and NO_3) that drive the dynamics of denitrifying and anammox bacteria, impact on the location 66 and thickness of the b_{bp} -layer, (2) NO₃ controls the vertical distribution of b_{bp} within this layer, (3) temperature drives the 67 formation of the b_{bp} -layer and consumption rates of NO₃, and (4) particle content inferred from b_{bp} and N₂ produced by 68 microbial communities can be at least qualitatively correlated. Ultimately, our findings allow us to infer that b_{bp} can potentially 69 be used to detect the presence of the microbial communities that drive N_2 production in poorly-oxygenated water masses – 70 including nitrate-reducing SAR11, and sulphur- oxidizing, denitrifying and anammox bacteria.

71 **2.0.** Background-nature of the small particles contributing to the b_{bp} -layer and their links with N₂ yielding

72 The poorly-oxygenated water masses of the Black Sea are characterized by a permanent layer of suspended small particles

- 73 constituted of organic and inorganic particles (Murray et al., 1995; Kuypers et al., 2003; Konovalov et al., 2005; Kirkpatrick
- ret al., 2012). Organic particles are mainly linked to microbial communities involved in the production of N₂, and these include

- nitrate-reducing SAR11, and anammox, denitrifying, and sulphur-oxidizing bacteria (Kuypers et al., 2003; Lam et al., 2007; Yakushev et al., 2007; Fuchsman et al. 2011; Kirkpatrick et al., 2012). The first group listed, SAR11, provides NO₂⁻ for N₂ yielding, and make the largest contribution (20-60%) to microbial biomass (Fuchsman et al., 2011, 2017; Tsementzi et al., 2016). Meanwhile, the second and third groups of bacteria make a smaller contribution to microbial biomass (~10%; e.g. Fuchsman et al., 2011, 2017) but *dominate* N₂ yielding via anammox (NO₂⁻ + NH₄⁺ \rightarrow N₂) and heterotrophic denitrification (NO₃⁻ \rightarrow NO₂⁻ \rightarrow N₂) (Murray et al., 2005; Kirkpatrick et al., 2012; Devries et al., 2013;Ward, 2013). Finally, the last group can potentially produce N₂ via autotrophic denitrification (e.g. 3H₂S + 4NO₃⁻ \rightarrow 3SO₄²⁻ + 2N₂ + 6H₂O; Sorokin, 2002;
- 82 Konovalov et al., 2003; Yakushev et al., 2007).
- 83 The inorganic component is mainly due to sinking particles of manganese oxides (Mn, III, IV) that are formed due to the 84 oxidation of dissolved Mn (II, III) pumped from the sulfidic zone (e.g. $2Mn^{2+}(l) + O_2 + H_2O \rightarrow 2MnO_2(s) + 4H^+$; Konovalov 85 et al., 2003; Clement et al, 2009; Dellwig et al., 2010). Ultimately, sinking particles of manganese oxides are dissolved back 86 to Mn (II, III), mainly via chemosynthetic bacteria that drive sulphur reduction (e.g. $HS^- + MnO_2(s) + 3H^+ \rightarrow S^0 + Mn^{2+}(l) + 3H^+$ 87 2H₂O; Jorgensen et al., 1991; Konovalov et al., 2003; Johnson, 2006; Yakushev et al., 2007; Fuschman et al., 2011; Stanev et 88 al., 2018). Overall, these arrays of bacteria mediate the reactions described above by using electron acceptors according to the 89 theoretical "electron tower" (e.g., $O_2 \rightarrow NO_3 \rightarrow Mn(IV) \rightarrow Fe(III) \rightarrow SO_4^2$; Stumm and Morgan, 1970; Murray et al., 1995; 90 Canfield and Thamdrup, 2009). Therefore, the vertical distributions of NO_3 , N_2 excess, and content of small particles are 91 driven by the reactions that occur in the chemical zones of poorly-oxygenated water masses (e.g. nitrogenous and manganous 92 zones, which correspond to the sections where NO_3 and Mn(IV), respectively, are predominantly used as electron acceptors; 93 Murray_et al., 1995; Konovalov et al., 2003; Yakushev et al., 2007; Canfield and Thamdrup, 2009; see also sections 4.2 and
- 94 4.3).

95 3 Methods

96 3.1 Bio-optical and physicochemical data measured by BGC-Argo floats

- 97 We used data collected by three BGC-Argo floats that profiled at a temporal resolution of 5-10 days in the first 1000 m depth 98 of the Black Sea from December 2013 to July 2019 (Figure 1). These floats - allocated the World Meteorological 99 Organization (WMO) numbers 6900807, 6901866, and 7900591 — collected 239, 301, and 518 vertical profiles, respectively. 100 BGC-Argo float 6901866 was equipped with four sensors: (1) a SBE-41 CP conductivity-T-depth sensor (Sea-Bird Scientific), 101 (2) an Aanderaa 4330 optode (serial number:1411), (3) a WETLabs ECO Triplet Puck, and (4) a Satlantic Submersible 102 Ultraviolet Nitrate Analyzer (SUNA). These sensors measured upward profiles of: (1) temperature (T), conductivity, and 103 depth, (2) dissolved oxygen (O_2), (3) chlorophyll fluorescence, total optical backscattering (particles + pure seawater) at 700 104 nm and fluorescence by Colored Dissolved Organic Matter, and (4) nitrate (NO₃; detection limit of 0.5 μ M with T/salinity 105 correction processing) and bisulfide (HS⁻, detection limit of 0.5 μ M; Staney et al., 2018). Floats 6900807 and 7900591 were
- 106 equipped with only the first three sensors.
- 107 Raw data of fluorescence and total backscattering were converted into Chlorophyll concentration (chl) and particle
- backscattering (b_{bp}) following standard protocols, respectively (Schmechtig et al., 2014, 2015). Spike signals in vertical profiles of *chl* and b_{bp} and due to particle aggregates were removed by using a median filter with a window size of three data
- promos of our and v_{pp} and due to parallel aggregates were removed by using a median inter with a window side of an ee and 110 points (Briggs et al., 2011). NO₃⁻, HS⁻ and O₂ data were processed following BGC-Argo protocols (Johnson et al., 2018;
- 111 Thierry et al., 2018; Bittig and Körtzinger, 2015). Sampling regions covered by the three floats encompassed most of the
- 112 Black Sea area (Figure 1, and Appendix A). However, we only used data collected during periods without a clear injection of

- small particles derived from the productive layer and Bosporus plume (e.g. advection of water masses, Stanev et al., 2017).
- 114 This restriction allowed us to focus on the *in-situ* 1D processes driving local formation of the b_{bp} -layer, with minimal
- 115 interference from any possible external sources of small particles.
- 116 We only describe the time series of data collected by float 6901866 because this was the only float carrying a NO_3^{-}/HS^{-} sensor.
- 117 Data acquired by floats 6900807 and 7900591 are described in Appendix A, and nevertheless used as complementary data to
- 118 those of float 6901866 to corroborate: (1) qualitative correlations between O_2 levels and the location of the b_{bp} -layer, and (2)
- 119 consistency in the location of the b_{bp} maximum within the b_{bp} -layer.

120 **3.2 Defining the poorly-oxygenated zone**, mixed layer depth, and productive layer

- 121 We used O_2 and NO_3^- to respectively define the top and bottom isopycnals of the poorly-oxygenated zone where denitrifying 122 and anammox bacteria are expected to be found. To set the top isopycnal, we applied an O₂ threshold of $\sim 3 \mu M$ because 123 denitrifying and anammox bacteria seem to tolerate O_2 concentrations beneath this threshold (Jensen et al., 2008; Dalsgaard 124 et al., 2014; Babbin et al., 2014). The bottom isopycnal was defined as the deepest isopycnal at which NO_3^- was detected by 125 the SUNA sensor $(0.23 \pm 0.32 \mu M)$. NO₃ was used to set this isopycnal because heterotrophic denitrification and subsequent 126 reactions cannot occur without NO₃⁻ (Lam et al., 2009; Bristow et al., 2017). HS⁻ was not used to delimit the bottom of this 127 zone because the maximum concentration of HS that denitrifying and anammox bacteria tolerate is not well established 128 (Murray et al., 1995; Kirkpatrick et al., 2012; see also section 4.1).
- Mixed layer depth (MLD) was computed as the depth at which density differed from 0.03 kg m⁻³ with respect to the density recorded at 1m depth (de Boyer Montégut et al., 2004). We used *chl* to define the productive layer where living phytoplankton were present and producing particulate organic carbon. The base of this layer was set as the depth at which *chl* decreased below 0.25 mg m⁻³. This depth was used only as a reference to highlight the periods when surface-derived small particles were clearly injected into the poorly-oxygenated zone.

134 **3.3** Complementary cruise data on N₂ excess and NO₃.

- 135 Published data on N₂:Ar ratios and NO₃⁻ collected at the southwest of the Black Sea in March 2005 (Fuchsman et al., 2008,
- 136 2019) were exploited to complement discussion of our results. N_2 produced by anaerobic microbial communities (N_2 excess,
- 137 µM) was estimated from N₂:Ar ratios and argon concentrations at atmospheric saturation (Hamme and Emerson, 2004). N₂
- 138 excess data were used to: (1) describe the poorly-oxygenated zone where N_2 is expected to be predominantly produced, and
- 139 (2) highlight qualitative correlations between N_2 excess, the location of the b_{bp} -layer, and vertical distribution of small particles
- 140 within the b_{bp} -layer.



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Figure 1: (a) Sampling locations of float 6901866 between May 2015 and July 2019. Colored circles indicate the date (color bar) for a given profile. The white star in (a) marks the sampling site of the cruise (March 2005). The white x in (a) highlights the float location on 6th April 2016. Float profiles of (b) $\log(O_2)$, (c) NO_3^- , (d) $\log(b_{bp})$, and (e) HS⁻ collected on 24th November 2018.

146 4 Results and discussion

147 **4.1 Description of the poorly-oxygenated zone**

148 The top and bottom of the poorly-oxygenated zone are located around the isopycnals (mean \pm standard deviation) 15.79 \pm 0.23 149 kg m⁻³ and 16.30 \pm 0.09 kg m⁻³, respectively. The two isopycnals therefore delimit the poorly-oxygenated water masses where 150 nitrate-reducing SAR11, and denitrifying, anammox, and sulphur-oxidizing bacteria are expected to be found (zone hereafter 151 called the PO_{D-A} , Figure 2; Kuypers et al., 2003; Lam et al., 2007; Yakushev et al., 2007; Fuschman et al., 2011; Kirkpatrick 152 et al., 2012). The top location of the PO_{DA} shows large spatial-temporal variability ranging between 80-180 m (or σ_{θ} between 153 15.5-15.9 kg m⁻³, Figure 2). Similarly, the PO_{D-A} thickness varies between 30-80 m, which corresponds to a σ_{θ} separation of 154 ~0.50 kg m⁻³. The bottom of the PO_{DA} is slightly sulfidic (HS⁻ = 11.4 ± 3.53 μ M, n = 86) and deeper than suggested (e.g. σ_{θ} = 155 16.20 kg m³, and $H_2S \le 10$ nM, Murray et al., 1995). However, our results coincide with the slightly sulfidic conditions of the

- 156 deepest isopycnal at which anammox bacteria can be still recorded ($\sigma_{\theta} = 16.30$ kg m⁻³, and H₂S $\geq 10 \mu$ M; Kirkpatrick et al.,
- 157 2012).



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Figure 2: Time series of: (a) Salinity (S), (b) O_2 , (c) NO_3^- , (d) $log(b_{bp})$, and (e) HS⁻. The blue lines in (a) and (b) indicate the mixed layer depth. The red lines in (a), (b) and (d) show the base of the productive region. The isopycnals 15.79 kg m⁻³ and 16.30 kg m⁻³ describe the top and bottom of the poorly-oxygenated zone (PO_{D-A}), respectively. SU, A, W, and SP stand for summer, autumn, winter, and spring, respectively. The colored horizontal line in (b) indicates the sampling site for a given date (Figure 1). The horizontal white lines in (d) are the profiles used to: (1) delimit the PO_{D-A} , and (2) compute correlations between b_{bp} , NO_3^- , and T within the PO_{D-A} .

165 **4.2** NO₃⁻, O₂, and MnO₂ as key drivers of the thickness and location of the suspended small-particle layer

166 The permanent b_{bp} -layer is always confined within the two isopycnals that delimit the PO_{D-A} (Figure 2). It follows that the

167 thickness and top location of this layer demonstrate the same spatial and temporal variability as the one described for the $P_{O_{D}}$.

- 168 $_{A}$ (Figure 2 and Appendix A). This correlation indicates that variations in the thickness and top location of the b_{bp} -layer are 169 partially driven, respectively, by: (1) the amount of NO₃⁻ available to produce N₂ inside the PO_{D-A} via the set of bacteria 170 communities involved, and (2) downward ventilation of oxygen-rich subsurface waters (Figure 2 and Appendix A).
- 171 NO_3 and O_2 are two of the key factors that modulate the presence of: (1) denitrifying and anammox bacteria working in 172 conjunction with nitrate-reducing SAR11 (Fuschman et al., 2011; Ulloa et al., 2012; Tsementezi et al., 2016; Bristow et al., 173 2017), and probably with chemoautotrophic ammonia-oxidizing bacteria (in this case, only with anammox, e.g. yAOB; Ward 174 and Kilpatrick, 1991; Lam et al., 2007), and (2) sulphur-oxidizing bacteria (e.g. SUP05; Canfield et al., 2010; Fuschman et al., 175 2011; Ulloa et al., 2012). Therefore, the results described above highlight that at least a fraction of the b_{bp} -layer should be due 176 to this array of bacteria. This notion is supported by three main observations. Firstly, the top location of the b_{bp} -layer is driven 177 by the intrusion of subsurface water masses (S $\leq 20.36 \pm 0.18$ psu) with O₂ concentrations above the levels tolerated by 178 denitrifying and anammox bacteria ($O_2 \ge 3 \mu M$, Jensen et al., 2008; Babbin et al., 2014; Figure 2). As a result, in regions where 179 O_2 is ventilated to deeper water masses, the top location of the b_{bp} -layer is also deeper. The contrary is observed when O_2 180 ventilation is shallower (Figure 2 and Appendix A). Secondly, nitrate-reducing SAR11, and denitrifying, anammox, and 181 sulphur-oxidizing bacteria reside between the isopycnals 15.60-16.30 kg m⁻³ (Fuchsman et al., 2011; 2012a; Kirkpatrick et al., 182 2012), while the b_{bp} -layer is formed between isopycnals ~15.79-16.30 kg m⁻³. We can thus infer coexistence of such bacteria 183 between the coincident isopycnals where the b_{bv} -layer is generated. Thirdly, NO₃ declines from around isopycnal 15.79 kg m 184 ³ to the isopycnal 16.30 kg m⁻³ due to the expected N_2 production via the microbial communities involved (Figures 2-3, and 185 Kirkpatrick et al., 2012).
- 186 The ventilation of subsurface O_2 is also key in driving the depth at which MnO_2 is formed ($O_2 \le 3-5 \mu M$;, Clement et al., 2009), 187 and can thus contribute to setting the characteristics of the b_{bp} -layer via its subsequent accumulation and dissolution 188 (Konovalov et al., 2003; Clement et al., 2009; Dellwig et al., 2010). Thus, in regions where subsurface O_2 (e.g. $O_2 \ge 3-5 \mu M$, 189 and $S \le 20.36 \pm 0.18$ psu) is ventilated to deeper water masses, both the formation of MnO_2 and top location of the b_{bp} -layer 190 can be expected to be deeper, and vice versa (Figure 2). Finally, the dissolution of MnO_2 should also influence the thickness 191 of the b_{bp} -layer because it occurs just beneath the maxima of the optical particles inside this *layer* (Konovalov et al., 2006; see 192 the explanation in section 4.3).
- 193 Overall, the qualitative evidence presented above points out that particles of MnO_2 as well as nitrate-reducing SAR11, and 194 denitrifying, anammox, and sulphur-oxidizing bacteria, appear to define the characteristics of the b_{bp} -layer (Johnson, 2006; 195 Konovalov et al., 2003; Fuchsman et al., 2011, 2012b; Stanev et al., 2018). This observation leads us to argue, in the next 196 section, that the b_{bp} -layer is partially composed of the main group of microbial communities involved in N₂ yielding, as well 197 as of MnO_2 .

198 **4.3** Role of the removal rate of NO_3 , MnO₂, and temperature in the vertical distribution of small particles

We propose that the removal rate of NO₃⁻ is a key driver of the vertical distribution of small particles and N₂ excess within the PO_{D-A} . This is because the vertical profiles of small particles and of N₂ excess are qualitatively similar, and both profiles are clearly related to the rate at which NO₃⁻ is removed from the PO_{D-A} (Figures 3-4). For instance, maxima of N₂ excess and b_{bp} coincide around the isopycnal 16.11 ± 0.11 kg m⁻³ (Figure 3; Konovalov et al., 2005; Fuchsman et al., 2008, 2019). At this isopycnal, the mean concentration of NO₃⁻ is 1.19 ± 0.53 µM. We thus propose that this NO₃⁻ threshold value splits the PO_{D-A} in two sub-zones with distinctive biogeochemical conditions (e.g. nitrogenous and manganous zones; Canfield and Thamdrup,

- 205 2009). Ultimately, these two different sets of conditions drive the rates at which $NO_{3^{-}}$ and small particles are removed and 206 formed within the *PO*_{*D-A*}, respectively (Figure 3, and explanation below).
- 207 The first sub-zone is thus located between the top of the PO_{D-A} ($\sigma_{\theta} = 15.79$ kg m⁻³) and around the isopycnal 16.11 kg m⁻³. 208 Here, removal rates of NO₃⁻ (-0.16 \pm 0.10 μ M m⁻¹, Figure 4) are likely to be boosted by: (1) high content of organic matter 209 (dissolved organic carbon = $122 \pm 9 \mu$ M, Margolin et al., 2016) and NO₃ ($\geq 1.19 \pm 0.53 \mu$ M), and (2) O₂ levels staying between 210 a range that maintain the yielding of N₂ ($0.24 \pm 0.04 \mu M \ge O_2 \le 2.8 \pm 0.14 \mu M$, n = 100, the means of the minima and maxima 211 of O_2 , respectively, in the first sub-zone) and promote the formation of MnO_2 (e.g. maximum of Mn(II) oxidation is at O_2 levels 212 \sim 0.2 μ M; Clement et al., 2009). Consequently, the formation of biogenic and inorganic small particles (and related N₂ excess) 213 increases from the top of the PO_{DA} to around the isopycnal 16.11 kg m⁻³ (Figure 3). This hypothesis is: (1) in part confirmed 214 by significant and negative power-law correlations between the suspended small-particle content and NO₃⁻ in this sub-zone 215 (Figure 3), and (2) in agreement with the progressive accumulation of MnO_2 from around isopycnal 15.8 kg m⁻³ to the isopycnal
- 216 16.10 kg m^{-3} (e.g. Konovalov et al., 2006).
- 217 The second sub-zone is located between isopycnal 16.11 kg m⁻³ and the bottom of the PO_{DA} ($\sigma_{\theta} = 16.30$ kg m⁻³, Figure 3). 218 Here, NO₃⁻ is low ($\leq 1.19 \pm 0.53 \mu$ M) and O₂ is relatively constant (0.23 ± 0.02 μ M, n= 2284, mean of O₂ calculated in the 219 second sub-zone for all profiles), or lower than the minimum of O₂ recorded by this sensor (0.22 \pm 0.02 μ M, n = 89). These 220 constant (or lower) levels of O₂ roughly correspond to those at which anammox and heterotrophic denitrification are inhibited 221 by $\sim 50\%$ (0.21 µM, and 0.81 µM, respectively; Dalsgaard et al., 2014). In addition, low levels of NO₃ necessarily promotes 222 the microbial use of Mn(IV) as an electron acceptor, ultimately dissolving the particles of MnO₂ into Mn(II) (e.g. manganous 223 zone; Konovalov et al., 2006; Yakushev et al., 2007; Canfield and Thamdrup, 2009). As a result, this sub-zone exhibits a 224 decline in removal rates of NO₃⁻ (-0.04 \pm 0.01 μ M m⁻¹, Figure 4) along with inhibited formation of biogenic small particles and 225 dissolution of MnO₂. Ultimately, both the content of small particles and related N_2 excess decrease from around isopycnal 226 16.11 kg m⁻³ to the bottom of the PO_{D-A} (Figure 3). These results are in agreement with: (1) significant and positive exponential 227 correlations computed between the small-particle content inferred from b_{bp} and NO₃⁻ within this sub-zone (Figure 3), and (2) 228 the overlap of nitrogenous and manganous zones in this sub-zone because the content of MnO₂ particles and dissolved Mn(II) 229 concurrently declines and increases just beneath the isopycnal 16.11 kg m⁻³, respectively (e.g. Murray et al., 1995; Konovalov 230 et al., 2003, 2005, 2006; Yakushev et al., 2007; Canfield and Thamdrup, 2009).
- 231 Strong-positive linear correlations are also recorded between b_{hp} and T in the first sub-zone of the PO_{D-A} (Figure 4). This is 232 likely to indicate that the formation of small particles is sensitive to very tiny increments in T (0.003 ± 0.001 °C m⁻¹, n = 133). 233 We thus infer a tendency for the decline rates of NO_3^{-1} and related production of N_2 to increase with T. This hypothesis is at 234 least partially supported by the significant correlation between NO3⁻ decline rates and T increase rates in this sub-zone (Figure 235 4). Within the second sub-zone, T continues increasing while b_{bp} decreases, likely due to inhibition of the formation of small 236 particles for the reasons described above (Figure 4). These observations suggest that the production of small particles is likely 237 to have first- and second-order covariations, with NO_3^{-1} and T, respectively – a likelihood backed up by a lack of correlation 238 between NO₃⁻ decline rates and T increase rates in this sub-zone (Figure 4). Finally, more information is needed to investigate 239 the physical and/or biogeochemical processes driving the correlation between the increase rates of T, and declines rates of 240 NO_{3} in the first sub-zone. This is however out of the scope of our study.



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242 Figure 3: (a) Cruise profiles of NO₃, and N₂ excess, collected in March 2005 (Fuchsman et al., 2019). (b) Float profiles 243 of NO₃, *b_{bp}*, and log(O₂) measured on 6th April 2016. Profiles in (a) and (b) were conducted at the northwest of the basin 244 (see Figure 1). The top and bottom of the PO_{D-A} are described in (a) and (b) as horizontal blue and red lines, respectively. 245 The b_{bp} maximum is the horizontal black line in (b). The first and second sub-zone of the PO_{D-A} are respectively 246 highlighted in (b) as blue and red squares. NO₃ vs b_{bp} in (c) the first, and (d) the second sub-zone, of the float profile in 247 (b). The number of data points visualized in (c) is lower than in (b) for the first sub-zone because b_{bp} and NO₃⁻ are not 248 always recorded at the same depths. (e) Frequency distributions of correlation coefficients (R, blue bars), and root 249 mean square errors (RMSE, white bars) for NO₃ vs b_{bp} in the first sub-zone. (f) Same as (e) but for the second sub-zone. 250 (g) Frequency distributions of the isopycnals at which b_{bp} maxima are found within the PO_{D-A} . Dotted, dashed, and solid 251 black lines in (g) are data collected by floats 7900591, 6901866, and 6900807, respectively. Gray bars include all data.



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Figure 4: Float profiles of (a) NO₃⁻, and b_{bp} , and (b) T and log(O₂) collected on 10th September 2017. Horizontal blue and red lines in (a) and (b) are the top and bottom of the PO_{D-A} . The b_{bp} maximum is indicated in (a) and (b) as horizontal black lines. The first and second sub-zones of the PO_{D-A} are respectively highlighted in (a) and (b) as blue and red squares. (c) b_{bp} vs T for the first sub-zone of the profile in (b). (d) Frequency distributions of correlation coefficients (R, blue bars), and root mean square errors (RMSE, white bars), for b_{bp} vs T in the first sub-zone, including data collected by the three floats. Decrease rates of NO₃⁻ vs increase rates of T in (e) the first and (f) the second sub-zone.

259 To summarize, BGC-Argo float data combined with a proxy of N₂ production suggest that in regions without the Bosporus 260 plume influence, the b_{bp} -layer systematically tracks and delineates the effective N₂-yielding section independently of: (1) the 261 biogeochemical mechanisms driving N_2 yielding, and (2) the contribution that MnO_2 and other microorganisms can be 262 expected to make to the formation of the *b_{bp}-layer* (e.g. Lam et al., 2007; Fuchsman et al., 2011; 2012a; Kirkpatrick et al., 263 2018). It is thus finally inferred that this b_{bp} -layer is at least partially composed of the predominant anaerobic microbial 264 communities involved in the production of N₂, such as *nitrate-reducing SAR11*, and anammox, denitrifying, and sulphur-265 *oxidizing* bacteria. These results also suggest that N₂ production rates can be highly variable in the Black Sea because the 266 characteristics of the b_{bp} -layer show large spatial-temporal variations driven by changes in NO₃⁻ and O₂ (Figures 2 and 4). 267 Finally, we propose that b_{bp} and O_2 can be exploited as a combined proxy for defining the N₂-producing section of the poorly-268 oxygenated Black Sea. We consider that this combined proxy can delineate the top and base of this section, by applying an O_2 269 threshold of 3.0 μ M, and the bottom isopycnal of the b_{bp} -layer, respectively. This section should thus be linked to free-living 270 bacteria (0.2-2 μ m), and those associated with small suspended particles (> 2-20 μ m), as well as to small inorganic particles 271 (0.2-20 μm).

272 4.4 New perspectives for studying N₂ losses in ODZs

The conclusions and inferences of this study, especially those related to the origin and drivers of the b_{bp} -layer, primarily apply to the Black Sea. However, these findings may also have a wider application. In particular, the shallower water masses of

- 276 be optically detected by b_{bp} and the attenuation coefficients of particles (Spinrad et al., 1989; Naqvi et al., 1993; Whitmire et 277 al., 2009). This layer is mainly linked to N₂-yielding microbial communities because: (1) its location coincides with the maxima 278 of N_2 excess, microbial metabolic activity, and nitrite (NO_2^- , the intermediate product of denitrification-anammox that is mainly 279 accumulated in the N₂-yielding section, Spinrad et al., 1989; Naqvi et al., 1991, 1993; Devon et al., 2006; Chang et al., 2010, 280 2012; Ulloa et al., 2012; Wojtasiewicz et al., 2018), and (2) MnO₂ is not accumulated as in the Black Sea (Martin and Knauer, 281 1984; Johnson et al., 1996; Lewis and Luther, 2000). Therefore, our findings suggest that highly resolved vertical profiles of 282 b_{bp} and O₂ can potentially be used as a combined proxy to define the *effective* N₂-production section of ODZs. Such definition 283 can be key to better-constrained global estimates of N_2 loss rates because it can allow us to: (1) accurately predict the poorly-284 oxygenated water volume where around 90% of N_2 is produced in the ODZ core (Babin et al., 2014), and (2) evaluate how the 285 location and thickness of the N₂-yielding section vary due to changes in the biogeochemical factors that modulate anammox 286 and heterotrophy denitrification.
- 287 Global estimates of N₂ losses differ by 2-3 fold between studies (e.g. 50-150 Tg N yr¹, Codispoti et al., 2001; Bianchi et al., 288 2012, 2018; DeVries et al., 2012; Wang et al., 2019). These discrepancies are caused in part by inaccurate estimations of the 289 **poorly-oxygenated** volume of the N_2 -production section. Other sources of uncertainties arise from the methods applied to 290 estimate the amount of POC that fuels N₂ production. For instance, POC fluxes and their subsequent attenuation rates are not 291 well resolved because they are computed respectively from satellite-based primary-production algorithms and generic power-292 law functions (Bianchi et al., 2012, 2018; DeVries et al., 2012). POC-flux estimates based on these algorithms visibly exclude: 293 (1) POC supplied by zooplankton migration (Kiko et al., 2017; Tutasi and Escribano, 2020), (2) substantial events of POC 294 export decoupled from primary production (Karl et al., 2012), and (3) the role of small particles derived from the physical and 295 biological fragmentation of larger ones (Karl et al., 1988; Briggs et al., 2020), which are more efficiently remineralized by 296 bacteria in ODZs (Cavan et al., 2017). In addition, these estimates do not take into consideration the inhibition effect that O_2 297 intrusions may have on N₂-yield rates (Whitmire et al., 2009; Ulloa et al., 2012; Dalsgaard et al., 2014; Peters et al., 2016; 298 Margolskee et al., 2019).

Overall, mechanistic predictions of N_2 losses misrepresent the strong dynamics of the biogeochemical and physical processes that regulate them. Consequently, it is still debated whether the oceanic nitrogen cycle is in balance or not (Codispoti, 2007; Gruber and Galloway, 2008; DeVries et al., 2012; Jayakumar et al., 2017; Bianchi et al., 2018; Wang et al., 2019). The subsiding uncertainty points to a compelling need for alternative methods that allow accurate refinement of oceanic estimations of N_2 losses.

- Our study supports the proposition that robotic observations of b_{bp} and O_2 can be used to better delineate the N₂-yielding section at the appropriate spatial (e.g. vertical and regional) and temporal (e.g. event, seasonal, interannual) resolutions. In addition, POC fluxes and N₂ can be simultaneously quantified using the same float technology (BGC-Argo, Bishop et al., 2009; Dall'Olmo and Mork, 2014; Reed et al., 2018; Boyd et al., 2019; Estapa et al., 2019; Rasse and Dall'Olmo, 2019). These robotic measurements can contribute to refining global estimates of N₂ losses by better constraining both the poorlyoxygenated section where N₂ is produced, and POC fluxes that fuel its loss. Ultimately, O₂ intrusions into the N₂-yielding section can potentially be quantified by BGC-Argo floats to assess their regulatory effect on N₂ losses.
- 311 Conclusions

Our results suggest that the b_{bp} -layer of the poorly-oxygenated Black Sea is at least partially composed of nitrate-reducing SAR11, and anammox, denitrifying, and sulphur-oxidizing bacteria. The location and thickness of this layer show strong

314 spatial-temporal variability, mainly driven by the ventilation of oxygen-rich subsurface waters, and nitrate available to generate

N₂, respectively. Such variations in the characteristics of the b_{bp} -layer highlight that N₂-production rates can be highly variable in the Black Sea. We therefore propose that high resolution measurements of O₂ and b_{bp} can potentially be exploited as a combined proxy to delineate the *effective* N₂-yielding section of ODZs. This proposition is in part supported by evidence that the b_{bp} -layer and a majority of N₂-yielding microbial communities are both confined in the shallower poorly-oxygenated water masses of ODZs. We however recommend investigation into the key biogeochemical drivers of the b_{bp} -layer for each ODZ. This information will be critical for validating the applicability of the b_{bp} -layer in assessing spatial-temporal changes in N₂ production.

Finally, it is evident that BGC-Argo float observations can acquire essential proxies of N_2 production and associated drivers at appropriate spatial and temporal resolutions. The development of observation-modeling synergies therefore holds the potential to deliver an unprecedented view of N_2 -yielding drivers if robotic observations become an integrated part of model validation. Ultimately, this approach could prove essential for reducing present uncertainties in the oceanic N_2 budget.







Figure A1: Sampling locations of floats (a) 7900591 and (b) 6900807 between December 2013 and July 2019. Colored squares and hexagons indicate the date (colorbar) for a given profile of floats 6900807 and 7900591, respectively.



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Figure A2: Time series of (a) S, (b) O_2 , (c) log(chl), and (d) $log(b_{bp})$ for float 7900591. The blue line in (c) indicates the mixed layer depth. The red lines in (c) and (d) show the base of the productive region. The isopycnals 15.79 kg m⁻³ and 16.30 kg m⁻³ describe the top and bottom of the poorly-oxygenated zone (PO_{D-A}), respectively. SU, A, W, and SP stand for summer, autumn, winter, and spring, respectively. The colored horizontal line at the bottom indicates the sampling site for a given date (Figure S1). The horizontal white lines in (d) are the profiles used to: (1) delimit the SO_{D-A} , and (2) find the isopycnals at which b_{bp} is maximum in the SO_{D-A} . *chl* is set to zero in the SO_{D-A} due to fluorescence contamination (Stanev et al., 2017).



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339 Figure A3: Same as Figure A2 but for float 6900807

340 Data availability. Data from Biogeochemical-Argo floats used in this study are freely available at ftp.ifremer.fr/ifremer/argo.
341 These data were collected and made freely available by the International Argo Program and the national programs that
342 contribute to it (http://www.argo.ucsd.edu; the Argo Program is part of the Global Ocean Observing System). Data on N₂:Ar
343 ratios are freely available at https://agupubs.onlinelibrary.wiley.com/doi/abs/10.1029/2018GB006032.

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