Active mesopelagic prokaryotes support high respiration in the subtropical northeast Atlantic Ocean

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[1] Here we provide evidence, based on prokaryote metabolic proxies and direct estimates of oxygen consumption, that the mesopelagic prokaryote assemblage in the subtropical Northeast Atlantic is an active one. It supports a high respiration (0.22 ± 0.05 μmol O₂ l⁻¹ d⁻¹, corresponding to 68 ± 8 mmol CO₂ m⁻² d⁻¹), comparable to that of the epipelagic zone during the same period (64–97 mmol C m⁻² d⁻¹). Our findings suggest that mesopelagic prokaryotes in the NE subtropical Ocean, as well as in other eastern boundary regions, are important carbon sinks for organic matter advected from the highly productive coastal systems, and would play a key role in the global carbon cycle of the oceans. Citation: Arístegui, J., C. M. Duarte, J. M. Gasol, and L. Alonso-Sáez (2005), Active mesopelagic prokaryotes support high respiration in the subtropical northeast Atlantic Ocean, Geophys. Res. Lett., 32, L03608, doi:10.1029/2004GL021863.

1. Introduction

[2] The mesopelagic (200–1000 m) open ocean has been suggested as a major site for biological oxygen consumption and CO₂ production [del Giorgio and Duarte, 2002]. Yet, oxygen consumption in the mesopelagic zone has only been examined directly in a few regions, thereby precluding verification of the presumed role of its biota on the carbon budget of the ocean. The pioneering work of Jenkins in the Sargasso Sea [Jenkins, 1977] led for the first time to the estimation of oxygen consumption in the mesopelagic zone of the ocean, from changes in the apparent oxygen utilization (AOU) and the use of tracers to calculate the apparent age of a water mass. His results were surprising in that he showed that oxygen consumption in the mesopelagic zone of the Sargasso Sea was considerably higher than the estimates of new production in the same region [Jenkins, 1982]. Unfortunately, this biogeochemical approach is only useful for the few oceanic regions where mixing below the surface thermocline is assumed to be unimportant, and consequently the age of the water mass can be calculated with relative confidence. This handicap has impeded the derivation of global estimates of respiration in the dark ocean through AOU/tracers.

[3] The first assessment of global respiration in the dark ocean was recently produced by Arístegui et al. [2003]. These authors compiled available estimates of enzymatic ETS (electron transport system) respiratory activity in the dark ocean, and converted them to actual respiration rates (R), by using a constant R/ETS ratio of 0.09. The ratio was derived from empirical R/ETS relationships determined in vitro from monospecific cultures of bacteria at their senescent phase [Christensen et al., 1980], suggested to be representative of the dark ocean [Packard et al., 1988]. The calculated average global R in the dark ocean (5 mol C m⁻² yr⁻¹) was in close agreement with the oxygen utilization rates inferred from large-scale tracer balances by Jenkins for the Sargasso Sea, building confidence to the R/ETS calculations. However, Arístegui et al. [2003] cautioned about the uncertainty in the general use of a constant R/ETS ratio derived from bacteria cultures in a low-activity physiological state. Since, prokaryotic organisms in the deep ocean could maintain a rather high activity [e.g., Cho and Azam, 1988], the ETS to R conversion factor would need to be validated in each region through concomitant oxygen consumption and enzymatic measurements of natural mesopelagic communities.

[4] Here, we have addressed this challenge by combining direct measurements of in vitro oxygen consumption at selected depths and detailed vertical profiles of ETS determinations, in the mesopelagic zone of the subtropical Northeast Atlantic Ocean. Additionally, we examined different metabolic proxies for the degree of activity of the prokaryote assemblages in the mesopelagic zone, and compared them with the activity of surface-water assemblages. We aimed to test the hypothesis that eastern boundary regions, which are supposed to receive high lateral inputs of organic matter from coastal upwelling ecosystems, would maintain highly active mesopelagic microbial assemblages, supporting a high respiratory activity.

2. Methods

2.1. Data Collection

[5] The study was conducted along two zonal sections (21°N and 26°N) extending from the NW African shelf to the open Atlantic Ocean at 26°W, during May–June 2003.
A total of 10, roughly equidistant, stations (5 in each section) were occupied. Samples for determination of prokaryote abundance and their physiological state were collected each 100 m down to 1000 m. Samples for respiration experiments were obtained from 600 m ($10 \pm 0.1^\circ C$) and 1,000 m ($6.7 \pm 0.1^\circ C$) depth, and immediately taken to chambers set at the in situ temperature ($\pm 0.2^\circ C$) conditions.

### 2.2. Prokaryote Respiration (R)

[6] Two approaches were used to estimate R: (1) Water samples drawn into carboys were siphoned into 5 replicate “time-zero” and “dark” (incubated for 2–4 days) 125 mL-BOD bottles. A time-series experiment was carried out at the most oceanic station; with 2 extra sets of “dark” replicate bottles added at intermediate times. R was estimated from the difference in oxygen concentration between the “zero” and “dark” bottles following incubation. Dissolved oxygen was measured by the Winkler technique, using an automated precise titration system, with colorimetric end-point detection. The mean standard error (SE) among replicated bottles was 0.08 $\mu$mol O$_2$ L$^{-1}$. (2) Because R might often lie beyond the detection limit for the procedure described, R was also assessed by pre-concentrating (on-average, a factor of 2.2 fold) the prokaryote assemblage by inverse filtration under gentle pressure, using a temperature-controlled cell concentration chamber fitted with 76 mm, 0.1 $\mu$m polycarbonate filters. The oxygen concentration was calculated spectrophotometrically, with a mean SE among replicated bottles of 0.43 $\mu$mol O$_2$ L$^{-1}$ and a mean SE for replicated measurements from the same bottle of 0.19 $\mu$mol O$_2$ L$^{-1}$. R was calculated as above. Additionally, we monitored changes in abundance and metabolism of the microbial assemblage during the incubations, allowing the back-scaling of R to that corresponding to the assemblage prior to incubation (see results for details).

### 2.3. ETS Measurements

[7] Seawater (10 to 20 l), collected every 50 m from 200 m to 1000 m depth, was filtered through 47 mm Whatman GF/F filters, at a low vacuum pressure (<0.3 atm). The filters were stored in liquid nitrogen until being assayed in the laboratory. ETS determinations were carried out as described by Arístegui and Montero [1995]. An incubation time of 20 min at 18°C was used. ETS activities measured at 18°C were converted to activities at in situ temperatures by using the Arrhenius equation. A calculated activation energy of 16 kcal mole$^{-1}$ was used.

### 2.4. Prokaryote Abundance and Biomass

[8] Cells were counted by flow cytometry, using a FACScalibur (Becton & Dickinson) instrument [Gasol and del Giorgio, 2000]. Samples were fixed with 1% of paraformaldehyde + 0.05% glutaraldehyde (final concentrations) and stored deep-frozen until analyzed. Samples were stained with Syto13 (Molecular Probes Inc) at 2.5 $\mu$M final concentration. Prokaryotes were detected by their signature in a plot of side scatter vs. green fluorescence. The division of total prokaryotes into high-NA and low-NA prokaryotes served both as an estimate of assemblage structure and of relative activity [Gasol and del Giorgio, 2000]. Green fluorescence was used as an estimate of prokaryote cell size [Gasol and del Giorgio, 2000] and converted to biomass using the alometric equation of Norland [1993].

### 2.5. Prokaryote Physiological State

[9] The CTC (5 cyano-2, 3 diotyl tetrazolium chloride) reduction technique was used to estimate the physiological state of the prokaryote assemblages. We added 5 mM CTC of a daily-prepared batch to water samples that were incubated (from 3 to 12 hours) at in situ temperatures. After incubation, the samples were analyzed on board the ship with the cytometer, to measure the red fluorescence of the formazan granules and the relative size of each formazan particle.

### 2.6. Prokaryote Production (P)

[10] P was estimated from the incorporation of tritiated leucine (Leu) and thymidine (TdR), following standard procedures. We used 4 replicates plus two TCA-killed blanks and incubated with 40 nM TdR and 40 nM Leu for 3 to 8 hours. Precipitation was done with ice-cold TCA. Incorporation rates were transformed to carbon production rates with conversion factors of 1.5 kg C mol$^{-1}$ Leu$^{-1}$, which assumes no intracellular isotope dilution, and 20 kg C mol TdR$^{-1}$, derived from a TdR conversion factor of $1.6 \times 10^{18}$ cells mol$^{-1}$ and 13 fg C cell$^{-1}$. The average value of TdR and Leu production for each sample was used as the final P estimate.

### 3. Results and Discussion

[11] The abundance of prokaryotes in the mesopelagic zone of the subtropical NE Atlantic averaged $2 \times 10^8$ cells L$^{-1}$, with an average Leu and TdR-based production (P) of 33 mmol C L$^{-1}$ d$^{-1}$, about 7 and 12 fold lower, respectively, than in the epipelagic zone (Table 1). Highest P values were found in the area affected by the African coastal upwelling, while lower values were observed towards the central Atlantic. However, the average turnover of the mesopelagic prokaryote assemblage (0.13 ± 0.02 d$^{-1}$, Table 1) was only 3 times less than that of the epipelagic prokaryotes, indicative of an actively growing mesopelagic assemblage. This was consistent with the relatively high contribution of prokaryotes with high-nucleic acid (NA) content to the

### Table 1. Prokaryote Abundance and Activity in the Epipelagic and Mesopelagic Zones

<table>
<thead>
<tr>
<th></th>
<th>Epipelagic Zone</th>
<th></th>
<th>Mesopelagic Zone</th>
<th></th>
<th>Ratio E:M</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prokaryote abundance (10$^6$ Cells l$^{-1}$)</td>
<td>23</td>
<td>12.4 ± 0.3</td>
<td>31</td>
<td>1.8 ± 0.2</td>
<td>6.9</td>
</tr>
<tr>
<td>High-NA-content prokaryotes (% of total)</td>
<td>23</td>
<td>37.8 ± 2.2</td>
<td>27</td>
<td>57.9 ± 2.0</td>
<td>0.7</td>
</tr>
<tr>
<td>Leucine/Thymidine-based P (nmols C l$^{-1}$ d$^{-1}$)</td>
<td>27</td>
<td>391.6 ± 77.1</td>
<td>29</td>
<td>32.7 ± 6.5</td>
<td>12.4</td>
</tr>
<tr>
<td>Prokaryote P/B (d$^{-1}$)</td>
<td>27</td>
<td>0.40 ± 0.12</td>
<td>29</td>
<td>0.13 ± 0.02</td>
<td>2.9</td>
</tr>
<tr>
<td>CTC + prokaryote abundance (10$^7$ Cells l$^{-1}$)</td>
<td>23</td>
<td>8.4 ± 2.6</td>
<td>20</td>
<td>1.7 ± 0.3</td>
<td>4.8</td>
</tr>
<tr>
<td>CTC + prokaryotes (% of total)</td>
<td>23</td>
<td>6.8 ± 1.1</td>
<td>20</td>
<td>8.1 ± 1.1</td>
<td>0.8</td>
</tr>
</tbody>
</table>
assemblage (58 ± 2%, Table 1). Indeed, 8 ± 1% of the mesopelagic prokaryotes were actively respiring, as indicated by their reduction of the tetrazolium salt CTC (Table 1), a fraction similar to that in the upper ocean (7 ± 1%). Furthermore, the respiring cells at mesopelagic depths produced 28% larger formazan granules than the surface prokaryotes (t-test, P < 0.05), which suggests that the respiration rate per cell was higher in the mesopelagic zone.

Prokaryote abundance increased upon confinement of the samples for respiration experiments with rates averaging 0.44 ± 0.04 d⁻¹ (average doubling time: 2.8 days). The 3-days time-series experiment showed that the oxygen concentration decreased and the cumulative heterotrophic production increased gradually along the experiment (Figure 1). The integrated prokaryote respiration (μmol O₂ l⁻¹) throughout all the experiments was closely related to the integrated prokaryote production (μmol C l⁻¹), measured both as the change in biomass (r = 0.81, n = 30) and from the average Leu and TdR uptake (r = 0.79, n = 30) (Figure 2). The measured daily rate of prokaryote respiration (R) was also closely related to the average abundance of prokaryotes (PA), during each experiment (Figure 3), in spite that the average cell-specific respiration rate apparently decreased towards the end of the time-series experiment. R was therefore corrected using the relationship with PA obtained (R = 7.2 × 10⁻¹¹ PA¹.₈₇±₀.₁₇; Figure 3) to account for the observed increase in prokaryote abundance. Using this equation, we derived a value of R corresponding to the prokaryote assemblage representative of conditions in situ (i.e., the prokaryote abundance inside the Niskin bottles). The resulting corrected estimates of R averaged 0.22 ± 0.05 μmol O₂ l⁻¹ d⁻¹ across the study area, being about 4 fold higher at 600 m (0.35 ± 0.08 μmol O₂ l⁻¹ d⁻¹) than at 1000 m depth (0.08 ± 0.03 μmol O₂ l⁻¹ d⁻¹). These rates are about two thirds of those directly derived from the incubation of unconfined samples (0.33 ± 0.03 μmol O₂ l⁻¹ d⁻¹), showing that the re-scaling procedure probably accounted for potential artifacts derived from growth after confinement.

The derived respiration rates for the mesopelagic subtropical NE Atlantic are about one order of magnitude higher than the average global rate implied in previous assessments (0.02 μmol O₂ l⁻¹ d⁻¹) [Arístegui et al., 2003]. The reason for this discrepancy may be two-fold: (1) the subtropical NE Atlantic is an area with particularly high respiratory activity, as it receives substantial lateral organic inputs from the African upwelling which may fuel mesopelagic R; (2) previous estimates of the global average mesopelagic R may be underestimates as these were indirect ones, largely inferred from measured ETS activity and a R/ETS of 0.09, derived from senescent bacterial cultures [Packard et al., 1988; Arístegui et al., 2003]. However, the results presented here depict the mesopelagic prokaryote
assemblages as actively growing (Table 1) rather than senescent. Indeed, the average R/ETS ratio in our data set, after excluding two data points from two coastal stations in the Cape Blanc upwelling, was 0.68 (SE ± 0.11; 0.9 without excluding the two outlier points), 8 fold greater than that assumed in the past. The rather high R/ETS ratio in the mesopelagic subtropical NE Atlantic is similar to that of the mixed layer microbial assemblages [Arístegui and Montero, 1995]. More interestingly, the ratio is comparable to that (average R/ETS = 1.1, range = 0.6–1.7) observed during the exponential growth phase of the same bacterial cultures used to derive the R/ETS in their senescent state [Christensen et al., 1980]. Our findings suggest that the general use of a low ratio (<0.1) to derive mesopelagic R from ETS may lead to gross underestimation of this process in the global ocean, if prokaryote assemblages are actively growing as observed in our study. Indeed, the prokaryote growth efficiency [PGE = P/(P + R)], estimated with the back-scaled R and the Leu/Tdr-based P was, on average, 0.18 ± 0.03 at 600 m, and 0.13 ± 0.02 at 1000 m. These PGE values are comparable to the average PGE of sea surface prokaryote assemblages [del Giorgio and Cole, 2000], and suggest that mesopelagic R must be comparable to that of the epipelagic zone if similar integrated abundances of prokaryotes are found in both zones, as was the case in our region of study.

We further calculated integrated mesopelagic R in the subtropical NE Atlantic by combining 10 vertical profiles (200–1000 m) of ETS activity with the empirically-estimated R to ETS ratio of 0.68, yielding an average value of 68 ± 8 mmol C m⁻² d⁻¹ (assuming a respiratory quotient, RQ = 0.69). The mesopelagic R is comparable to the epipelagic R measured during the same cruise (64–97 mmol C m⁻² d⁻¹) (N. Navarro, personal communication, 2004) or in previous studies (89–136 mmol C m⁻² d⁻¹) in the same region [Duarte et al., 2001]. This indicates a very high carbon demand by mesopelagic prokaryotes that cannot possibly be fulfilled by vertical inputs of organic carbon alone. Estimates of vertical inputs of organic carbon into the mesopelagic layer at this region are well below these values (unpublished results), and the activity of the layer must be, therefore, largely fueled by lateral exports from the highly productive NW African upwelling system.

Several sets of evidence support this hypothesis. Recent studies in the NW African coast indicate that upwelling filaments may transport offshore up to 50% of coastal upwelling primary production, even during low to moderate upwelling pulses [Gabric et al., 1993; García-Muñoz et al., 2004]. Upwelling filaments are known to be a widespread and recurrent phenomena in the NW African coast [Kostianoy and Zatsepina, 1996]; hence their annual contribution to the coastal-offshore exchange of organic matter must be important. Additionally, high concentrations of particulate organic carbon (POC) might be also transport to the open ocean via intermediate or deep nepheloid layers. Indeed, vertical profiles of suspended POC from the upper 1000 m, more intense towards the coast, which could only be explained by lateral advection of organic matter from the African shelf.

In conclusion, our findings confirm that mesopelagic prokaryote assemblages are active nodes of organic carbon remineralisations and act as major sinks for organic carbon in the subtropical NE Atlantic Ocean. A large part of the mesopelagic respiration must be supported by coastal advection of organic matter from the highly productive upwelling system to the open ocean. The possibility that the high mesopelagic R reported here could be a feature of other eastern boundary regions, bordering high productive upwelling systems, awaits verification. If confirmed, the overall organic carbon sink in these regions would play a key role in the global carbon cycle in the oceans.

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