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1	THE SUPPLY OF NUTRIENTS DUE TO VERTICAL TURBULENT MIXING:
2	A STUDY AT THE PORCUPINE ABYSSAL PLAIN STUDY SITE (49°N
3	16°30'W) IN THE NORTHEAST ATLANTIC
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15 Abstract

16 As part of a multidisciplinary cruise to the Porcupine Abyssal Plain (PAP) study site (49°00'N 16°30'W), in June and July of 2006, observations were made of the vertical 17 18 nitrate flux due to turbulent mixing. Daily profiles of nitrate and turbulent mixing, at 19 the central PAP site, give a mean nitrate flux into the euphotic zone of 0.09 (95% confidence intervals: 0.05-0.16) mmol N m⁻² d⁻¹. This is a factor of fifty lower than 20 21 the mean observed rate of nitrate uptake within the euphotic zone (5.1+/-1.3 mmol N $m^{-2} d^{-1}$). By using our direct observations to 'validate' a previously published 22 23 parameterisation for turbulent mixing we further quantify the variability in the vertical 24 turbulent flux across a roughly 100 km x 100 km region centred on the PAP site, using hydrographic data. The flux is uniformly low (0.08 +/- 0.26 mmol N m⁻² d⁻¹, the 25 large standard deviation being due to a strongly non-Gaussian distribution) and is 26 27 consistent with direct measurements at the central site. It is demonstrated that on an 28 annual basis convective mixing supplies at least forty-fold more nitrate to the euphotic 29 zone than turbulent mixing at this location. Other processes, such as those related with 30 mesoscale phenomena, may also contribute significantly.

31 1. Introduction

32 It may be thought surprising to claim that phytoplankton, upon whom so much life in 33 the sea depends, live on the margins of the open ocean. Yet they are typically confined to the upper 100m of a water column which extends to 4km or more. This 34 35 edge existence arises from the rapid absorption by water of the sunlight phytoplankton 36 need to photosynthesise. This would not be a problem for survival if they did not also 37 need nutrients to grow. However, the majority of nutrients used by phytoplankton 38 arise from the regeneration of decaying organic material and gravity ensures that this 39 process of nutrient recycling and accumulation occurs at depth. For phytoplankton to 40 grow it is therefore necessary to bring deep waters laden with nutrients to the surface 41 - a role fulfilled by the ubiquitous advection and mixing of water. Without this 42 physical 'supply line', phytoplankton in the open ocean would be much less abundant 43 and the dominant species very different. The small oligotrophy specialist 44 phytoplankton would dominate and even nitrogen fixers would find it difficult to 45 thrive due to the high N:P of atmospheric deposition. 46 The role of the physical circulation in controlling phytoplankton abundance

47 and productivity has been recognised for some time and a multitude of mechanisms 48 have been identified. Winter convective mixing (Williams et al., 2000), mesoscale 49 upwelling (Pollard & Regier, 1992; Allen et al., 2005) and small-scale turbulent 50 mixing (Lewis et al., 1986; Carr et al., 1995; Law et al., 2001; Law et al., 2003) are 51 three that have received perhaps the most attention. To understand the controls on 52 phytoplankton growth in a region, it is necessary to quantify the contributing flux 53 associated with each pathway. These contributions all vary in time and space. Winter 54 mixing stirs large quantities of nutrients to the surface but does so only for a relatively 55 short period each year. Mesoscale processes work throughout the year, but, though

56 they may drive large fluxes, they are intermittent in space and time. Though 57 turbulence at scales from centimetres to metres is also intermittent in space and time, 58 these scales are so much smaller than those involved in mesoscale processes that such 59 fluctuations in the circulation can be viewed as a constant background effect. For this reason, their cumulative effect is often modeled by analogy to molecular diffusion: 60 61 there too, intermittent displacements of varying size nevertheless result in dispersion at larger scales. Accordingly, mixing due to these small-scale processes is often 62 63 referred to as turbulent diffusivity, even though it is unrelated to (and much larger in 64 magnitude than) molecular diffusivity.

The turbulent diffusivity is typically 10⁻²m²s⁻¹ or more in the mixed layer but several orders of magnitude smaller deeper down (see for example Ledwell et al., 1998; Polzin et al., 1997). This reflects the major contribution of atmospheric cooling and wind-driven mixing to surface mixing with deeper turbulent motion being driven by processes such as breaking internal waves and interactions with topography.

We focus here on the turbulent flux of nitrate. We do this despite having
equivalent measurements for phosphate and silicate concentrations. The reason for
this choice is that we have simultaneous measurements for nitrate uptake. Therefore,
for nitrate it is possible to put the turbulent flux in context with the observed rate of
the nutrient's uptake by phytoplankton.

There are relatively few direct measurements of the nitrate flux due to turbulent mixing in the open ocean. In the Southern Ocean, Law et al. (2003) found the turbulent flux to account for just 8% of the nitrate required for observed carbon fixation rates. Naveira-Garabato et al. (2002) also found it to be a minor flux in the Antarctic Polar Front. In the northern North Atlantic, the flux accounted for just 16% of the observed drawdown of nitrate during a summer cruise (Law et al., 2001). In the

81 equatorial Pacific, Carr et al. (1995) found that vertical turbulent mixing could 82 account for roughly a third of the nitrate drawdown between 0 and 2°S but was a 83 negligible contributor further away from the equator. As the turbulent nitrate flux is 84 always present, it may be thought that it would be most significant in oligotrophic 85 regions, particularly during the stratified summer, where open ocean phytoplankton 86 are often nutrient limited. In the subtropical Atlantic Lewis et al. (1986) did indeed 87 find the turbulent supply to match the rate of nitrate uptake. However, the 88 measurements of nitrate uptake were substantially smaller than estimates arising from 89 tracer methods (e.g. Jenkins and Doney, 2003). There is, furthermore, little knowledge 90 concerning how vertical turbulent mixing varies at the mesoscale (Naveira-Garabato 91 et al., 2002). This is despite the long-standing paradigm that such mixing may be 92 strongly influenced by vertical gradients in horizontal currents, or shear (see for 93 example Turner, 1973). As mesoscale features such as eddies and fronts display 94 strong heterogeneity in shear, it is germane to ask if rates of turbulent mixing also 95 vary on these scales, with significantly higher mixing in regions of high strain. It 96 might be anticipated that turbulent mixing may vary considerably over a region large 97 enough to encompass varying strengths of mesoscale activity. 98 We present measurements of the vertical nitrate flux due to small-scale 99 turbulent mixing at the Porcupine Abyssal Plain (PAP) study site at 49°00'N16°30'W 100 in the Northeast Atlantic. The data were obtained as part of the D306 cruise from 23 101 June to 8 July 2006 on board RRS Discovery. It comprised a daily suite of 102 measurements at the PAP site augmented by a high resolution physical survey

spanning the last 4 days of the cruise and sampling a region roughly 100 km x 100 km

104 (Figure 1). The latter was intended to delineate the mesoscale physical structure and

105 variability of the region.

106 We provide direct estimates of the turbulent nitrate flux into the euphotic zone 107 in this area. This is achieved using observations from turbulence profiles carried out 108 immediately subsequent to CTD casts from which water was collected for nutrient 109 analysis. These flux estimates are compared with concurrent, and co-located, 110 observations of the rate of nitrate uptake. Preliminary results on the mesoscale 111 variability of the turbulent flux of nitrate are also presented. We test the applicability 112 of published parameterisations of turbulent diffusivity (which use current shear and 113 buoyancy frequency to make predictions) and use the most accurate for our region to 114 estimate the mesoscale variability of the turbulent nitrate flux using physical 115 hydrographic data from the mesoscale survey. We are, to the best of our knowledge, 116 the first to use direct measurements to 'validate' a parameterisation for estimating the 117 turbulent diffusivity before applying it to determine mesoscale spatial variability in the turbulent flux of nutrients for the open ocean. 118

The structure of the paper is as follows. Following this introduction, in Section 2, we describe the various methods, including those used to measure turbulent mixing, nitrate concentrations and uptake and mesoscale variability in physical structure. In Section 3 we present our results, both for a comparison of turbulent nitrate supply to nitrate uptake at the central PAP site and for the mesoscale variability of the flux. A discussion of our results is found in Section 4 prior to our conclusions in Section 5.

126 **2. Methods**

127 The central measurement for this study is the vertical turbulent flux of nitrate. It has 128 already been stated that the conventional model for this physical process is by direct 129 analogy with molecular diffusion. For this reason, the changes in distribution of an

130 inert tracer *C* undergoing turbulent mixing are modeled, in the standard Fickian

131 manner, as

132
$$\frac{\partial C}{\partial t} = \frac{\partial}{\partial z} \left(\kappa(z) \frac{\partial C}{\partial z} \right)$$
(1)

where *t* is time, *z* is depth and κ is the turbulent, or effective, diffusivity. We are interested in the supply of nitrate, *N*, to the surface. Since the turbulent flux is zero at the surface, by integrating over depth we obtain the rate at which nitrate is entering that portion of ocean through its lower boundary,

137
$$F(d) = \kappa(z) \frac{\partial N}{\partial z}\Big|_{z=d}$$
 (2)

i.e. the rate of supply of nitrate via turbulent mixing to the water above depth *d* is the product of the turbulent diffusivity and the nitrate gradient at depth *d*. Therefore, to estimate F(d) we simply require vertical profiles for both nitrate and κ .

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142 2.1 Turbulent diffusivity measurements

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The microstructure profiler used to measure turbulent diffusivities (MSS90L, serial
number 10) was produced by Sea and Sun Technology GmbH in co-operation with
ISW Wassermesstechnik. The latter participated in the cruise and carried out the
measurements.

The profiler is equipped with two velocity microstructure shear sensors as well as standard high precision conductivity, temperature, depth (CTD) sensors. The sampling rate for all sensors is 1024 samples per second, with 16 bit resolution. Although it is attached to the ship by a cable (providing power and data transmission), the profiler is allowed to sink in freefall by maintaining sufficient slack cable in the water at all times. This is to minimise contamination of the signal by vibrations of the

154 profiler caused by tension in the cable. All sensors are mounted at the measuring head 155 of the profiler with the microstructure sensors placed at the tip of a slim shaft, about 156 150mm in front of the CTD sensors. This minimises contamination of any signal by 157 turbulence created by the profiler itself sinking through the water. A vibration control 158 sensor and a two component tilt sensor also provide data to remove noise 159 contamination from the signal. The general behaviour of the MSS profiler is described 160 in detail by Prandke et al. (2000). The calibration of the CTD sensors was carried out 161 by Sea & Sun Technology GmbH using standard calibration equipment and 162 procedures for CTD probes. The vibration control sensor, the tilt sensors and the shear 163 sensors were calibrated by ISW Wassermesstechnik. 164 The turbulent energy dissipation rate in 1 m depth bins has been estimated following $\varepsilon = 7.5 \cdot v \overline{(\partial u / \partial z)^2}$ where v is the kinematic viscosity and $\partial u / \partial z$ is the small-scale 165

166 current shear. The processing of the shear data has been carried out as described by 167 Prandke (2005). The turbulent diffusivity is then calculated from the dissipation rate 168 using $\kappa = \gamma \cdot \varepsilon / N^2$ (Osborn, 1980) where N is the buoyancy frequency and γ is the 169 mixing efficiency. A constant mixing efficiency of 0.2 was used.

170 Each deployment of the profiler comprised a number of profiles. This is 171 necessary because the mixing processes involved are intermittent such that 172 consecutive profiles often show considerably different, yet genuine, structure. It is 173 therefore advisable to combine several profiles for each deployment to calculate a 174 mean profile of turbulent diffusivities. Analysis of the diffusivities at the same depth 175 for different profiles of the same deployment revealed a strongly non-Gaussian 176 distribution. The observations fitted a log-normal distribution for nearly all depths and deployments when tested using a Kolmogorov-Smirnov test on the log-transformed 177 178 data. Given the relatively small number of profiles per deployment (between 5 and

179 10) we therefore followed the Baker and Gibson (1987) method for estimating the mean diffusivity. More specifically the mean is estimated as $M = \exp(m + v^2/2)$ where m 180 and v^2 are the mean and variance of the log-transformed data respectively and the 181 95% confidence intervals are given by M*exp(+/-1.96*n) where $\eta = \sqrt{\left[v^2/n + v^4/2(n-1)\right]}$ 182 183 and n is the number of data points. For this averaging process data were further 184 averaged in 4 dbar bins. Estimates formed in this way are consistently less noisy than those obtained by naively using the arithmetic mean. This is by simple virtue of the 185 186 latter method being more strongly influenced by outlying large values. For all but a few points very near to the surface the Thorpe length is less than 4m. Therefore, the 187 188 size of the overturns comprising the turbulence is smaller than the scale of vertical 189 averaging.

190

191 2.2 Nitrate measurements

192

193 Throughout the paper, nitrate represents the sum of nitrate and nitrite. Samples for 194 analysis were drawn directly into 25 ml plastic coulter counter vials from Niskin 195 bottles that had been lowered on the CTD frame. The vials were stored in the dark at 196 4°C until analysis, which commenced within 24 hours of sampling. Nitrate was 197 determined in unfiltered water samples with a Skalar Sanplus segmented flow autoanalyser and standard colorimetric techniques described by Kirkwood (1995) and 198 199 Sanders et al. (2007). Overall, the precision of the data is estimated to be better than ± 0.12 umol l⁻¹ 200

- 201 (0.6% of the top standard). Consistency of the data was ensured by the analysis of
- 202 commercial nutrient standards (Ocean Scientific International, Petersfield, Hants,

203 UK). Concentrations of nutrients determined in the commercially available nutrient204 standards were within 4% of their designated values.

205 The consistency in the vertical profile of nitrate throughout the cruise was 206 remarkable (Fig 2a). This is despite strong evidence for advection of spatial 207 variability through the site (Painter et al., 2008b). It was therefore possible to 208 construct a statistical model to provide estimates of nitrate concentration for the 209 mesoscale survey where nitrate was only sampled at a subset of points. Predictions of nitrate, N_{pred}, from the model (fitted by minimising the least squares difference 210 211 between predictions and observations using the simplex algorithm – see for example 212 Press et al., 1992)

213
$$N_{pred} = A * \frac{P^B}{C + P^B},$$

where *P* is pressure in dbar, $A = 14.5135 \text{ mmol N m}^{-3}$, B = 1.1131 and C = 143.1686(dbar)^{*B*}, are correlated with observations with R²=0.91 (Fig 2b)¹. It is, therefore, a good first approximation to use the statistical model's vertical profile of nitrate to calculate the turbulent nitrate flux at different locations within the mesoscale survey area.

219

220 2.3 Nitrate uptake measurements

Sample water recovered from 6 light depths (97, 55, 33, 14, 4.5, 1%) was decanted directly into duplicated new 2 l acid-washed Nalgene polycarbonate incubation bottles for each light depth. One of these bottles was darkened with tin-foil and black tape to serve as a control for dark nitrate uptake. All bottles containing exactly 2 l sample water were inoculated with 100-200 μ l stock solution of K¹⁵NO₃⁻ (1

¹ It is worth noting in passing that an ISUS nitrate sensor was deployed on many of the CTD casts from which water was drawn for nitrate samples. However, perhaps by virtue of the very stable vertical structure, the ISUS data were much worse predictors of nitrate concentration than the above simple pressure-based model. For that reason the latter is used here. Subsequent work may have improved the accuracy and precision of ISUS (Sakamoto et al., 2008).

 μ mol / 100 µl), the volume of ¹⁵N spike being adjusted to ~10% of the ambient NO₃⁻ concentration.

After spiking, the incubation bottles were transferred to Perspex incubation tubes covered with neutral density filters (Lee: Misty Blue [061] and Neutral Density Grey [210 ND]) that re-constructed water column light attenuation (97, 55, 33, 14, 4.5 and 1% incoming irradiance) and removed red light. The incubators were cooled by a constant flow of surface seawater.

At the end of the ~ 10 hr incubation period, all ¹⁵N incubations were filtered 233 234 onto 25 mm ashed Whatman GF/F filters that were then stored at -20 °C for later 235 analysis by stable isotope mass spectrometry on the NOC's Stable Isotope Ratio Mass 236 Spectrometry facility (NOC's-SIRMS) using a Eurovector elemental analyser coupled 237 to a GV Isoprime mass spectrometer. The calibration standard was tyrosine, traceable 238 to International Atomic Energy Agency standards. Nitrate uptake was calculated 239 according to Dugdale and Goering (1967) and Dugdale and Wilkerson (1986) using 240 particulate N concentrations measured at the end of the incubation to account for 241 unlabelled N.

A standard astronomical formula was used to calculate the duration of daylight hours for the year day and position. The uptakes in light and dark bottles were then combined in a ratio equal to that of daylight hours to night-time hours to give the daily average uptake.

246

247 2.4 Euphotic depths

248

249 Irradiance was measured using a 4π downwelling Photosynthetically Available

250 Radiation (PAR) sensor attached to the CTD frame. Euphotic depth was calculated as

251 1% of surface irradiance.

252

253 2.5 Current velocity data

255 The raw east and north components of current velocity down to approximately 300m 256 were measured using a ship-mounted 150 kHz RDI Acoustic Doppler Current Profiler 257 (ADCP) and logged using RD Instruments data acquisition software (DAS version 258 2.48 with profiler firmware 17.20). The instrument was configured to sample over 120 259 second intervals with 96 bins of 4 m thickness, pulse length 4 m and a blank beyond 260 transmit of 4m. Spot gyro heading data were fed into the transducer deck unit where 261 they were incorporated into the individual ping profiles to correct the velocities to 262 earth co-ordinates before being reduced to 2 minute ensembles. Subsequent 263 processing steps merged the ADCP data with corrected heading information for the 264 ship's navigational GPS data-stream to obtain speed and direction. The ship's velocity 265 is also calculated from spot positions taken from the master navigation file and taken 266 from the ADCP velocities, resulting in an absolute water velocity in terms of east-267 west and north-south velocities, (see Burkill (2006)). Calibration of the 150 kHz 268 ADCP was achieved using bottom tracking data collected after departure from 269 Falmouth while crossing the continental shelf. The calibration involves the application 270 of two corrections; a misalignment angle and an amplitude factor. The misalignment 271 angle (ϕ) corrects for the rotational position of the ADCP on the ships hull relative to 272 the ships axis. The amplitude factor (A) corrects for the fore-aft tilt of the instrument 273 relative to the horizontal plane.

274

275 2.4 MVP data

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A fine scale survey of the mesoscale physics and key variables of the upper ocean was
conducted using a towed Conductivity, Temperature, Depth (CTD) device known as
the Moving Vessel Profiler (MVP). The instrument used was a BOT (Brookes Ocean

280 Technology) MVP 300 with an AML micro CTD instrument (S/N 7027). During the 281 survey, the MVP was towed behind the ship at a speed of 11-11.5 knots, undulating 282 up and down in the water column and completing a full surface-to-300m depth return 283 profile every 12-13 minutes in which time it travelled 4km. The MVP was towed along a pre-defined survey grid, Figure 1. Application of a temperature lag of 284 285 $\tau = 0.12$ s was found necessary to account for the delayed response of the temperature 286 sensor. More details can be found in the cruise report (Burkill, 2006). The MVP CTD 287 was calibrated by comparison to data from the surface thermosalinograph (TSG) data, 288 which had in turn been calibrated against data from the frame CTD casts. All MVP 289 temperature and salinity data from between 4 and 5m were extracted and merged with 290 the corrected TSG data. The difference between the MVP and corrected TSG in terms 291 of temperature and salinity was displayed in scatter plots. It was found necessary to 292 apply an offset of -0.033 to salinity but none to temperature.

293

294 **3. Results**

295

296 3.1 Turbulent diffusivities

297

A representative profile for the turbulent diffusivity, κ , is shown in Fig 3a. Despite the averaging of profiles for each deployment there is still considerable variability in mean profiles for κ between deployments. Nevertheless, there are consistent patterns. In the surface layer κ is generally of order $10^{-3} - 10^{-2} \text{ m}^2 \text{ s}^{-1}$, but occasionally much larger for the shallowest observations. At greater depth κ is of order $10^{-5} - 10^{-4} \text{ m}^2 \text{ s}^{-1}$. For the daily profiles at the central PAP site the mean depth of the euphotic zone is $50.8 \pm 5.7(\text{sd})$ dbar.

305 Note that in the density profile shown there is no clear seasonal pycnocline 306 (Figure 3f). This was a feature of several profiles. However, strong agreement was 307 found between density-inferred mixed layer depth and the depth to which mixing was 308 enhanced near the surface over the cruise as a whole (not shown). Fluxes into the 309 mixed layer are not discussed here because there was evidence that substantial 310 production was taking place below the mixed layer (Painter et al., 2008a). Instead we 311 focus on how the turbulent nitrate supply contributes to nitrate uptake throughout the 312 euphotic zone.

It should be noted that the large variability in estimated diffusivities, even at the same depth (note the logarithmic scale), does not indicate that the instrument or technique here is insufficiently precise. Rather it reflects the strongly intermittent nature of turbulence at centimetre to metre scales – something long recognised (e.g. Gregg, 1987; Gibson, 1991; Frisch, 1995).

318

319 *3.2 Nitrate supply at fixed stations relative to nitrate uptake*

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321 Before presenting the estimates of turbulent nitrate flux into the euphotic zone, it is 322 worth making a few comments regarding the other component of the calculation, the 323 vertical profile of nitrate. Figure 3b shows the nitrate vertical profile obtained 324 immediately preceding the turbulence profile in Figure 3a. The gradual increase in 325 nitrate concentration with depth is a standard feature. For all but two of the profiles 326 obtained immediately preceding a turbulence profiler deployment, the surface nitrate concentration is of order 1 mmol N m⁻³. For the two anomalous profiles, 327 concentrations are still greater than 0.3 mmol N m⁻³. Therefore it is unlikely that 328 nitrate was limiting phytoplankton growth during the cruise period. 329

330 A consequence of the sparse vertical spacing of bottle samples inevitably 331 obtained from CTD casts is that the calculated gradients in the nitrate profile are 332 rather variable (Figure 3c). Furthermore, the small number of points means that 333 smoothing by running averages or interpolating by anything other than a linear 334 method are questionable: smoothing would be certain to move the profile away from 335 the few data points we have for each cast: a higher order interpolation would be very 336 difficult to constrain with any confidence. Although either method would give a 337 smoother nitrate profile, and hence a more regular gradient, it would do so at a cost 338 that is difficult to justify. We also choose to use individual nitrate profiles for the 339 central station rather than the statistical model discussed earlier as they were taken 340 immediately prior to turbulence profiling. It is therefore necessary to accept some 341 variability in gradients. It should be noted that unlike turbulent diffusivity we cannot 342 form error bars accounting for the variability of nitrate profiles as we only have one 343 CTD cast on which nitrate samples were taken for each turbulence station. Therefore 344 the errors in nitrate profiles, gradients and flux are likely to be larger than those 345 indicated in Figures 3, 4 and 5 where only the variability due to turbulent diffusivity is quantified. Fig 3c shows the nitrate gradient for the representative profile. For all casts 346 the gradients are typically 0.1 mmol N m⁻⁴. Fig 4 shows the mean and standard 347 348 deviation for the nitrate gradients calculated using the nitrate profiles obtained on 349 CTD frame casts preceding a turbulence deployment. For the euphotic depth horizon 350 of interest (51m), the mean and standard deviation for the gradient are 0.13 mmol N m⁻⁴ and 0.06 mmol N m⁻⁴ respectively. 351

The nitrate uptake at each depth, once more for the same representative profile, is shown in Figure 3d. There is a general decrease with depth, with a small

maxima (seen in some but not all other casts) around 15m where the turbulent mixingdrops sharply (Fig 3a).

Fig 3e shows the turbulent nitrate flux at each depth calculated using equation 2 for the same representative profile. Superimposed on this is the total nitrate uptake integrated to that depth. Note the logarithmic scale. The euphotic depth for this profile is also marked. It is apparent that for this station the turbulent supply of nitrate to the euphotic zone is almost two orders of magnitude smaller than the contemporaneous total uptake of nitrate within it.

Figure 5 demonstrates the consistency in the relationship between uptake and turbulent flux at the central PAP site for the 11 days of the study. The mean uptake rate in the euphotic zone is 5.1+/-1.3 mmol N m⁻²d⁻¹, whilst the mean turbulent flux is 0.09 mmol N m⁻²d⁻¹, with 95% confidence intervals of 0.05 mmol N m⁻²d⁻¹ and 0.16 mmol N m⁻²d⁻¹.

367

368 3.3 Estimates of turbulent diffusivity from vertical shear

369

The relationship of small scale turbulent mixing to the hydrographic properties of stratification and vertical shear has received much attention (e.g. Pacanowski & Philander, 1981; Peters et al., 1988; Turner, 1975; Yu & Schopf, 1997). In particular people have sought to relate the strength of mixing to the Richardson number, $Ri = N^2(z)/S^2(z)$,

which represents the competing influences of stratification, as represented by thebuoyancy frequency

377
$$N(z) = \sqrt{\frac{g}{\rho} \frac{\partial \rho}{\partial z}} (3)$$

378 (where g is the acceleration due to gravity, z is depth and ρ is density), and vertical 379 current shear,

$$380 \qquad S(z) = \frac{\partial |\underline{u}|}{\partial z},$$

381 (where *u* is the current velocity) in controlling the likely onset of turbulence. We test 382 the ability of three different parameterisations to estimate the turbulent diffusivity for 383 our survey. Velocity profiles from the ADCP are used to estimate S, extracting data 384 simultaneous with the turbulent profiler deployment. We begin by using hydrographic 385 data from the CTD sensors on the turbulence probe to calculate N. Using this estimate 386 for N provides the most direct and hence fairest comparison to our direct measurements of κ . More specifically, N is actually calculated as N² from the square 387 388 of equation (3) using 4 dbar binned data for ρ in order to be consistent with the 4 dbar 389 averaging used for ADCP data and for the MVP data later.

Because the calculation of both *N* and *S* require vertical derivatives they, like the nitrate gradient discussed earlier, are sensitive to relatively small distortions in the vertical profile. As data for these are much higher in frequency (every 4 dbar) than for the nitrate profiles, we can justifiably smooth these. A running average of 7 adjacent bins (28 dbar) gives the best match between direct and indirect observations of turbulent diffusivity.

Figure 6 shows the comparison of direct observations to predictions from the
(a) Pacanowski and Philander (1981), (b) Yu and Schopf (1997) and (c) Peters, Gregg

and Toole (1988) parameterisations. The Pacanowski and Philander (1981)

399 parameterisation (hereafter PP) is the most accurate. Fig 6d demonstrates that between

400 50m and 200m the PP estimate is generally within a factor of 2 of the direct estimate.

401 Given the large variability in direct estimates, this is perhaps surprisingly good. It

402 should be noted that a certain amount of serendipity may be involved. PP was 403 developed for a model with a horizontal grid size of 40 km x 70 km and a non-404 uniform vertical resolution of order 15 km near the surface. Here, however, shear is 405 calculated at 4 m vertical resolution (albeit after applying a moving 28 m, or 7 bin, 406 window average to smooth it first) and the horizontal resolution will typically be 407 between 10 m and 100 m (due to the mounting angle of the ADCP the horizontal scale 408 will be roughly the depth of the measurement). The difference in scales between our 409 data and PP means that a direct 1:1 relationship can not be expected. In particular, the 410 key gradients in vertical velocity and density may be significantly larger at 4 m 411 resolution than that of the Paconowski and Philander (1981) model. Given the results 412 shown in Fig.6, PP is nevertheless adopted as our parameterisation for estimating the 413 diffusive flux of nitrate across the region covered by the mesoscale survey. 414 It should be noted that the above comparison is the most direct possible. To 415 estimate turbulent fluxes during the mesoscale survey, when κ was not directly

416 measured away from the central station, it is necessary to use hydrographic data from

417 the MVP to estimate *N*. This data was averaged and smoothed in an identical manner

to that applied to the turbulence probe CTD data for consistency. Without a number of

419 direct measurements at other locations within the region it is impossible to quantify

420 any associated change in precision.

421

422 *3.4 Variability in diffusive nutrient supply across the region*

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Using MVP CTD data to calculate the buoyancy frequency and simultaneous ADCP
data to calculate the shear, we estimated the turbulent diffusivities throughout the
mesoscale survey. These estimates were combined with the previously discussed

427 statistical model for the nitrate profile to estimate the associated fluxes of nitrate into 428 the euphotic zone across the region, as shown in Fig 7. The area of each dot is 429 proportional to the flux. The largest flux (3.1 mmol N m⁻³) is very much larger than 430 the other fluxes so it is not included in the colour scale but is instead solely 431 represented by area. Note that Fig.6 indicates that our parameterisation for turbulent 432 diffusivity may often be in error by a factor of 2. Therefore the fluxes, being linear in 433 turbulent diffusivity carry the same potential error. .

434 Using the median euphotic depth of 48 dbar (estimated from all 28 CTDs casts that comprised the survey), the mean diffusive flux is 0.12 mmol N m⁻² d⁻¹. The 435 predicted nitrate concentration at this depth is 5 mmol N m⁻³ +/- 1 mmol N m⁻³. Such a 436 437 20% potential error is minor compared to those associated with the estimate of the diffusivity. The distribution of fluxes is significantly non-Gaussian with standard 438 deviation of 0.26 mmol N m^{$^{-2}$} d^{$^{-1}$}. (There is, at face-value, a single hot-spot at 49.15N 439 16.75W with flux more than 20 times greater than the mean, clearly visible in Figure 440 7. The reliability of this observation is addressed in the Discussion.) Therefore the 441 median value, 0.08 mmol N m⁻² d^{-1} , is a more representative figure. The euphotic 442 443 depth is also distributed in a non-Gaussian manner for the region. However, all 444 euphotic depths bar one (an outlier of 100 dbar) are between 40 and 70 dbar. If the calculation is repeated for these depths then the median flux is 0.10 mmol N $m^{-2} d^{-1}$ 445 with 0.10 mmol N m⁻² d⁻¹ standard deviation at 40 dbar and 0.09 mmol N m⁻² d⁻¹ with 446 0.11 mmol N m⁻² d⁻¹ standard deviation at 70 dbar. It is apparent therefore that at 447 448 these depths the estimate of nitrate flux into the euphotic zone is relatively insensitive 449 to the precise depth used for the calculation. All rates are seen to be close to those 450 estimated directly at the central PAP site in Section 3.1.

451	The near uniformly low fluxes seen in Figure 7 are consistent with the rather
452	amorphous dynamical structure of the region. Figure 7 also shows potential
453	temperature at 47m to highlight this. There are no strong frontal regions which could
454	lead to enhanced shear either by themselves or via interactions with the wind. (The
455	strong dynamical signature of an eddy in the southwest corner (Painter et al., 2008b)
456	only manifests itself below 100m). The cluster of 3 higher fluxes near 49.1°N and
457	16.7°W are actually in one of the regions with most widely spaced isotherms.

458

459

460 **4. Discussion**

461 Our estimates for the turbulent flux of nitrate into the euphotic zone for both the central PAP site and the mesoscale survey are consistent. Both give a value of 462 approximately 0.1 mmol N $m^{-2} d^{-1}$. It is worth putting this in context with previous 463 studies. Lewis et al. (1986) diagnosed a flux of 0.14 mmol N $m^{-2} d^{-1}$ in the subtropical 464 North Atlantic. Carr et al. (1995) found the flux to be between 0.1 and 1 mmol N m^{-2} 465 d⁻¹ between 0 and 2S in the equatorial Pacific but at least an order of magnitude 466 467 smaller outside this region. In the northern North Atlantic, Law et al., (2001) found a large turbulent flux of 1.8 mmol N m⁻² d⁻¹, using an SF6 tracer to infer vertical 468 turbulent mixing. Using the same technique in the Southern Ocean they estimated the 469 flux to be an order of magnitude smaller 0.17 mmol N m⁻² d⁻¹. The discrepancy was 470 471 due to a substantially smaller inferred turbulent diffusivity in the Antarctic 472 Circumpolar Current. Both of the latter two studies, however, quantified the nitrate flux into the mixed layer rather than into the euphotic zone. 473 474 Our direct estimates of the turbulent flux of nitrate into the euphotic zone are one to two orders of magnitude less than the rate of nitrate uptake within it for the 475

476 same period. This raises the obvious question of what mechanisms supply or supplied 477 the nitrate that was taken up. Over recent years it has become increasingly apparent 478 that physical processes at the mesoscale and submesoscale (Mahadevan & Archer, 479 2000; Levy et al., 2001) can induce very large vertical fluxes, associated for example 480 with the formation and interactions of eddies (McGillicuddy & Robinson, 1997; 481 Martin & Richards, 2001) and with strong frontal regions (Allen et al., 2005; Lapeyre 482 & Klein, 2006). In theory the mesoscale survey would allow the calculation of vertical 483 velocities using the Omega equation (e.g. Pollard and Regier, 1992). However, such a 484 calculation needs to be constrained by the change in distributions between multiple 485 surveys to have any degree of robustness. Consequently we have no such estimates 486 for this cruise. There is also the potential that more nutrient rich water may be 487 advected horizontally into the site. Our dataset does not allow us to quantify such a 488 flux. However, other papers in this volume (Hartman et al; Painter et al. 2008a; 489 Smythe-Wright et al.) present evidence that such nutrient rich incursions do occur. 490 Therefore, here we only compare the rate of uptake to the nitrate flux associated with 491 convective mixing the previous winter. We stress that this is a rather crude analysis 492 purely intended to determine the likely relative magnitude of fluxes. 493 The CTD profiles for the surveyed region are curious since they show no clear

signal of winter mixing, taking such a signal to be a region of homogeneous
hydrographic properties below the seasonal thermocline. This is true whether one
examines temperature, salinity, density or oxygen. We therefore turn to the Coriolis
Project (http://www.coriolis.eu.org/) for information on winter mixed layer depths.
This database stores CTD profiles from ARGOS floats, gliders, buoys, moorings and
standard ship CTD casts. We have extracted temperature data (as this is the only
parameter recorded by all available profiles) for an approximately 200kmx200km

501 square centred on the PAP site for the period 1 February to 30 June 2006. Figure 8 502 shows the mixed layer depths for this period, calculated using a criterion of a decrease 503 in temperature of 0.05°C with respect to that at 5m depth. This is somewhat smaller 504 than has been used by others (e.g. Oka et al. (2007) use 0.2°C). However, individual 505 examination of potential temperature profiles indicated that this criterion gave the 506 most accurate diagnosis of mixed layer depth for the period studied, using the 507 criterion of matching the base of the deep winter homogenous layer. The maximum 508 mean monthly mixed layer depth is in February, extending to approximately 350 m 509 with one profile showing a 400 m mixed layer. These estimates for winter mixed layer 510 depth are consistent with climatological data from the World Ocean Atlas 511 (http://www.cdc.noaa.gov/cdc/data.nodc.woa94.html; Antonov et al., 2006; Locarnini 512 et al. 2006) which reports a mean February mixed layer depth for the same area 513 between 277 m and 472 m. We therefore take a representative value of 400m, 514 acknowledging that this might result in a small overestimate. From the deeper profiles 515 obtained during our cruise we know that the nitrate concentration at this depth is approximately 10.5 mmol N m⁻³. Using the simplest approach of multiplying nitrate 516 517 concentration at the base of the winter mixed layer by the euphotic depth, winter 518 mixing to 400m would therefore have provided a stock of approximately 504 mmol N m^{-2} within the euphotic zone (mean euphotic depth of 48dbar * 10.5 mmol N m^{-3}). 519 520 Unfortunately, oxygen profiles were not available to utilise the 'oxygen step' method of Koeve (2001). After subtracting the 127 mmol N m^{-2} still present above the 521 522 euphotic depth during the cruise, this leaves approximately 75 days supply at the mean nitrate uptake rate of 5 mmol N $m^{-2} d^{-1}$. Therefore, if nitrate uptake was roughly 523 524 the same for the 75 days preceding the start of the cruise (at the end of June) then 525 deep winter mixing would have to extend into mid-April to provide sufficient stocks

526 by itself. Except for two points (discussed below), the observations from mid-March 527 onwards reveal a mixed layer depth that is no deeper than the euphotic depth during 528 the cruise (Figure 8). Therefore, without sporadic deeper mixing between mid-March 529 and June, another mechanism must provide sufficient nitrate to the euphotic zone both 530 to match the observed uptake and to give the observed residual nitrate stock of 127 mmol N m⁻². There are two anomalously deep mixed layer depths later in the year. If 531 532 the temperature profile for the second of these (on Julian day 140) is examined more 533 closely there is evidence of a warming in the top few metres which has been missed 534 by the simple criterion used here. Hence the mixed layer is arguably much shallower 535 than the norm rather than deeper. However, the first anomalous point (on Julian day 536 130) has a homogenous profile to nearly 100m. This apparent deep mixing event 537 would deliver a substantial extra flux of nitrate to the mixed layer. We assume, once again, that nitrate uptake has been constantly 5 mmol N $m^{-2} d^{-1}$. There would, 538 therefore, remain, by Julian day 130, 229 mmol N m⁻³ of the nitrate entrained into the 539 540 euphotic zone by winter mixing. We also assume that the concentration at greater depths remains at 10.5 mmol m⁻³ on Julian day 130. Mixing to 99m, as the 541 observation suggests, would consequently introduce an extra 158 mmol N m⁻² to the 542 543 euphotic zone. This is sufficient for an extra 31 days at the uptake rate of 5 mmol N $m^{-2} d^{-1}$. Such an event would therefore allow deep winter mixing to meet the uptake 544 545 requirements given the above assumptions. However, nitrate uptake is likely to have 546 been much larger earlier in the year, during the spring (unless Fe-limited; see Moore 547 et al. (2006)). Consequently it is still most probable that another significant nitrate 548 source is required to close the budget, especially as our estimate of winter mixing using the nitrate concentration at the base of the winter mixed layer may be an over-549 estimate by as much as 50% (Koeve, 2001). The cumulative 9.5 mmol N m⁻² that 550

would result from extrapolation of our estimates for the diffusive supply over the same period (from mid-March to the end of June) is forty-fold smaller than the flux due to convective mixing even if convection stopped in mid-March. Hence, turbulent mixing will only have been a minor contribution.

555 In the absence of nitrate uptake data throughout the year, especially the spring, 556 it is impossible to put a firm estimate on the nitrate flux being delivered by pathways 557 other than winter convective mixing and small-scale turbulent mixing. Furthermore, the 127 mmol N m⁻² remaining in the euphotic zone at the end of the cruise would 558 559 only have lasted roughly 25 days at the observed uptake rate. The missing flux needs 560 to be of equivalent magnitude to the excess nitrate uptake taking place during the 561 spring period plus whatever nitrate would be utilised from the middle of August to the 562 end of the year.

563 To put the diffusive flux in a broader context, recent estimates for the rate of nitrification in the euphotic zone in the North Atlantic indicate a typical value of 0.01 564 mmol N m⁻³ d⁻¹ (D.Clark, pers.comm.; C.Fernandez I., pers.comm.). Integrating over 565 the euphotic depth gives a nitrification flux of almost 0.5 mmol N $m^{-2}d^{-1}$. This is five 566 567 times larger than the turbulent flux of nitrate reported here. It should, however, be 568 noted that there is a great deal of variability in measurements of nitrification (Yool et 569 al., 2007). Nevertheless, extant observations indicate that nitrification might be an 570 equivalent, if not larger, flux of nitrate to the euphotic zone than turbulent mixing in 571 mid / late summer for the studied area.

572 By 'validating' the most effective parameterisation for our area we have 573 estimated the variability in turbulent nitrate flux across a roughly 100 km x 100 km 574 region centred on the PAP site. There was remarkably little variability, especially 575 given the inherently variable nature of turbulent mixing. The one 'hot-spot'

576 (highlighted in Section 3.4) is instructive in the care needed for estimating turbulent 577 nutrient fluxes. The flux at this location is the largest measured in the area by some 578 margin and represents an extreme outlier. By looking at the vertical profiles for shear 579 and buoyancy frequency (not shown) it is possible to determine that it is due to small 580 deviations in shear and buoyancy frequency that may have been removed by a more 581 aggressive smoothing. This reinforces the care needed in using parameterisations of κ . 582 Inevitably subjective choices on the degree of smoothing exert a major influence on 583 the resulting estimates. For this reason, the results on variability presented here should 584 only be viewed as preliminary and individual fluxes, particularly outlying ones, 585 should be treated with care. Future fieldwork should ensure that profiles of turbulent 586 diffusivity and nitrate are obtained at a number of different locations within a region 587 as an independent check on the indirect method. In particular profiles should be 588 sought in differing physical regimes, covering a range of shear and buoyancy 589 frequency. There is little evidence of strong shear layers in the data collected by this 590 cruise, despite the presence of an eddy in the southeast corner of the surveyed region 591 (Painter et al., 2008b). Nevertheless, shear is often found to be significantly enhanced 592 in the vicinity of strong mesoscale features such as eddies and fronts (Allen & Smeed, 593 1996). The need for extra direct observations covering a range of regimes is therefore 594 even more vital in a strong dynamical region, where theory suggests the turbulent flux 595 may be more significant.

596

597 **5.** Conclusions

598

599 We have presented results on both direct and indirect estimates of the nitrate flux due 600 to vertical turbulent mixing for the PAP site and its environs. Comparison of the direct

601 estimates to simultaneous nitrate uptake measurements indicates that the turbulent 602 flux is a small contributor to the nitrate budget at the PAP site. A rough calculation 603 suggests that winter mixing is a major contributor. A simple analysis, however, 604 indicates that other mechanisms, most likely related to mesoscale physical 605 phenomena, may be equally significant. 606 We have also 'validated' a parameterisation for indirect estimation of 607 turbulent mixing and used this to estimate the nitrate flux into the euphotic zone 608 throughout the survey region. These indirect estimates agree with direct estimates in 609 terms of magnitude. They also vary very little across the region. 610 611 Acknowledgements 612 We would like to thank Jon Sharples and Mark Moore for useful discussions 613 regarding the processing of turbulent diffusivity data, Alberto Naveira-Garabato for 614 615 advice on the application and limitations of parameterisations for turbulent mixing, 616 Robin Hankin for his statistical expertise and finally, but most definitely not least, the 617 captain and crew of RRS Discovery for providing such a superb platform for 618 oceanographic research. This manuscript contributes to NOCS Theme 2 of the Oceans 619 2025 core programme funded by the Natural Environmental Research Council, UK. 620 Adrian Martin was also part funded by an NERC Advanced Research Fellowship 621 (NER/J/S/2001/00708). 622 References 623

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773 FIGURE CAPTIONS

774

775 Figure 1.

Map showing the survey track for the cruise. The central PAP site is at 49°50'N
16°30'W.

778

779 Figure 2.

780 (a) Variation of 'nitrate' (NO3+NO2) concentration versus pressure using data from

all profiles taken during the cruise (dots). The solid line shows the predicted nitrate

concentration using the fitted model, $N_{pred} = AP^B/(C+P^B)$ where A= 14.5135 mmol N

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m^{-3}, B= 1.1131 and C= 143.1686 (dbar)<sup>B</sup>. (b) Actual versus predicted nitrate
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concentrations (dots) with 1:1 line superimposed. The correlation has $R^2=0.91$. Units

785 are mmol N m^{-3} for both plots.

786

787 **Figure 3.**

788 Various profiles versus depth for a representative station on year day 179: (a)

turbulent diffusivity with mean for station as solid line and 95% confidence limits as

dashed lines; (b) 'nitrate' (actually NO3+NO2); (c) nitrate gradient; (d) nitrate uptake;

- (e) nitrate flux at each depth (mean thin solid line and 95% confidence intervals as
- dashed) plus cumulative nitrate uptake integrated from surface (thick solid line) and
- euphotic depth (horizontal solid line) calculated as 1% of surface irradiance; (f)

794 potential density.

795

796 Figure 4.

Mean (solid) and mean +/- one standard deviation (dashed) profiles of nitrate gradient
from CTD frame casts at central PAP site.

799

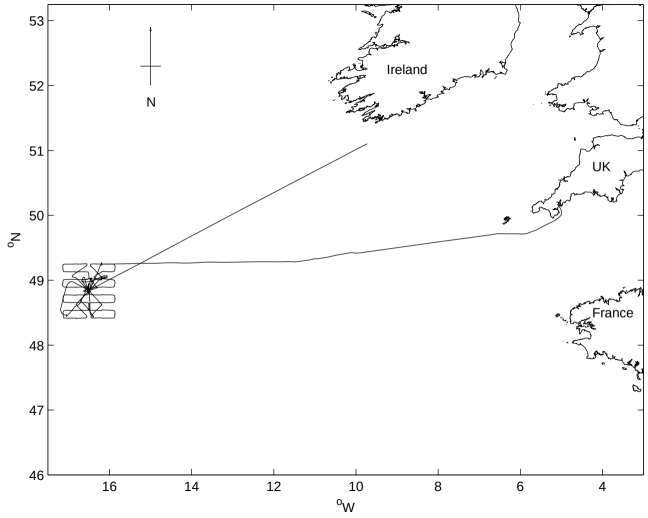
800	Figure 5.
801	Time series of flux and uptake at the central PAP site covering year days 179-187 (28
802	June to 6 July 2007). Turbulent nitrate flux at base of euphotic zone (48m) as thin
803	solid line with 95% confidence interval as dashed lines. Nitrate uptake integrated from
804	surface to base of euphotic depth (determined as 1% surface irradiance for each
805	profile independently) as thick solid line. Note the logarithmic y axis and that no
806	uptake data are available for year day 183.
807	
808	Figure 6
809	Direct observations of turbulent diffusivity versus estimates from the
810	parameterisations of (a) Pacanowski and Philander (1981), (b) Yu and Schopf (1997)
811	and (c) Peters, Gregg and Toole (1988). The solid line is 1:1 and the dashed lines
812	indicate $\frac{1}{4}$, $\frac{1}{2}$, 2 and 4 times the 1:1 relationship. (d) shows the fractional difference
813	versus depth between the direct observations and the best parameterisation (PP).
814	
815	Figure 7
816	Variability in the turbulent flux of nitrate (in mmol N $m^{-2} d^{-1}$) at the base of the mean
817	euphotic depth (48m) across the mesoscale survey region. The area of each dot (with
818	one for each CTD profile provided by the MVP) is proportional to the corresponding
819	flux. The greyscale also indicates the flux but it is chosen to fit all but the largest flux
820	$(3.1 \text{ mmol N m}^{-2} \text{ d}^{-1})$ for clarity. The dashed lines mark contours of potential

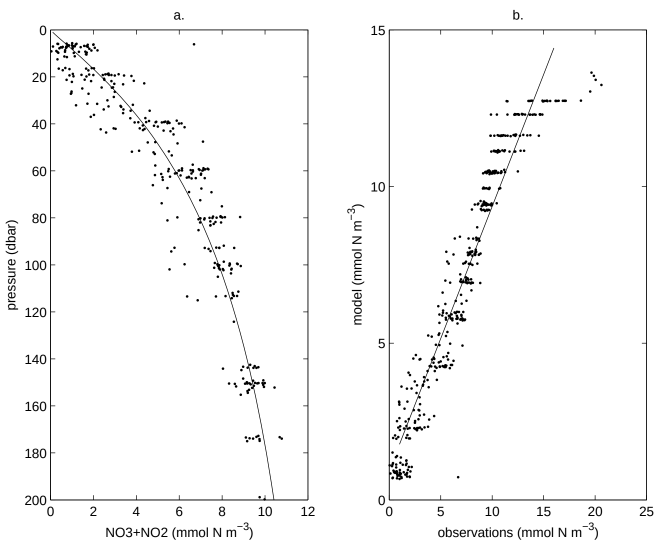
821 temperature to indicate the dynamical physical processes influencing this depth.

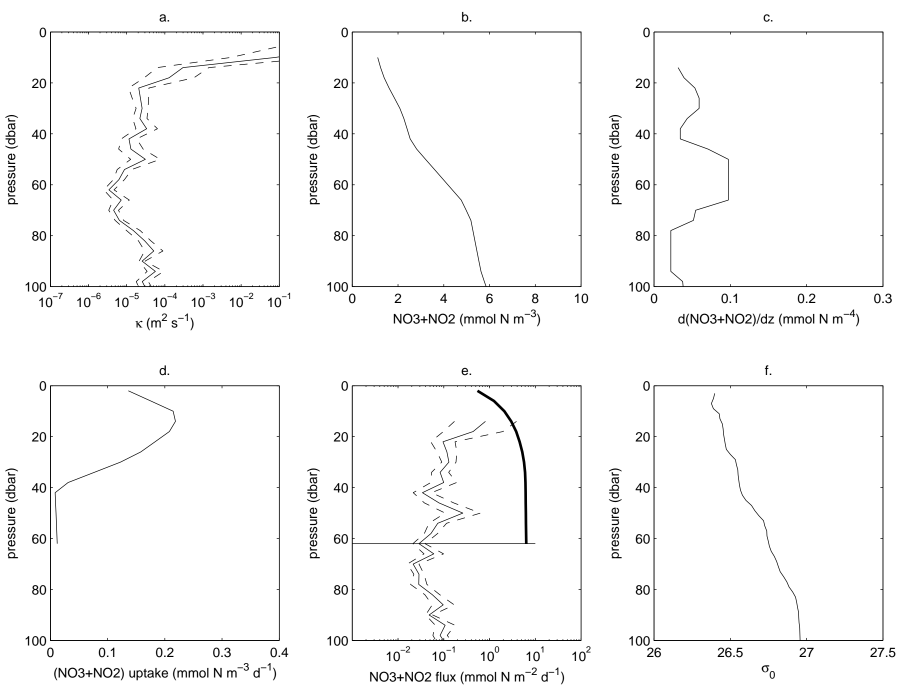
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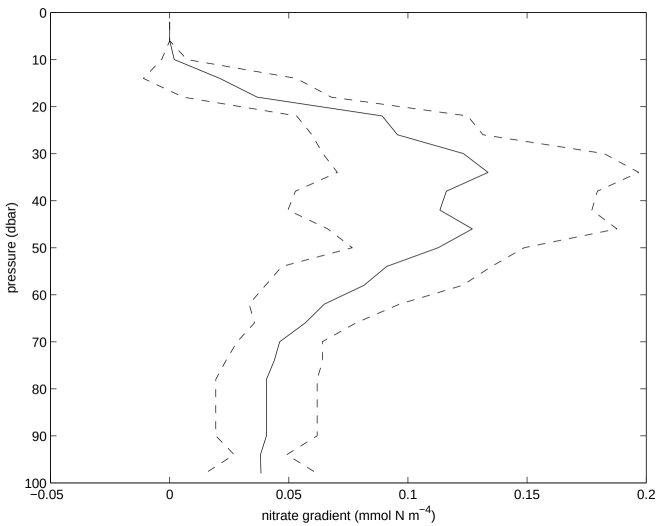
823 **Figure 8**

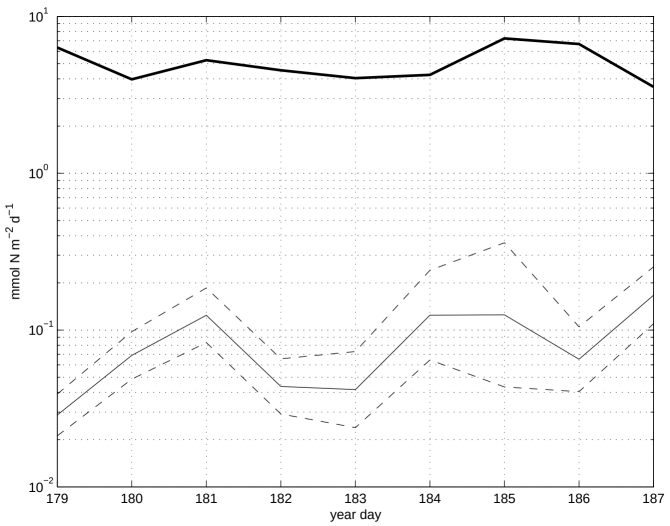
- 824 Mixed layer depth versus year day for a period in 2006. The most consistent criterion
- for estimating mixed layer depth was found to be the depth at which the temperature
- is 0.05 lower than at 5m depth. The hydrographic data is a combination of profiles
- from ARGOS floats and gliders and is taken from the Coriolis database
- 828 (http://www.coriolis.eu.org/). Also shown as a solid line is the mean euphotic depth
- 829 for the cruise discussed here.
- 830
- 831
- 832

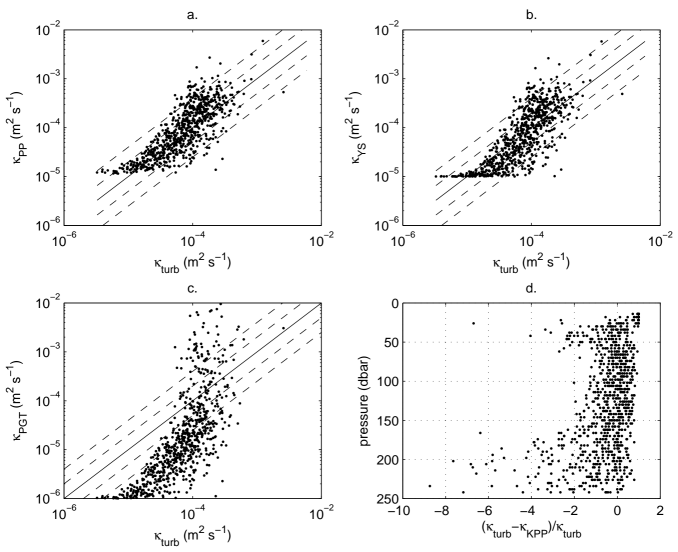












mmol N $m^{-2}d^{-1}$

