

# Intense upper ocean mixing due to large aggregations of spawning fish

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1 **Small-scale turbulent mixing plays a pivotal role in shaping the circulation and a broad range**  
 2 **of physical and biogeochemical ocean processes. Despite advances in understanding geophys-**  
 3 **ical processes responsible for this mixing, the nature and importance of biomixing – turbulent**  
 4 **mixing caused by marine biota – remains controversial. A major source of uncertainty per-**  
 5 **tains to the efficiency of biomixing – the fraction of the turbulent energy produced through**  
 6 **swimming that is spent in mixing the ocean vertically –, which the few *in situ* observations**  
 7 **available suggest to be much lower than that of geophysical turbulence. Here, we shed light**  
 8 **on this problem by analysing 14 days of continuous measurements of centimetre-scale tur-**  
 9 **bulence in a coastal upwelling area. We show that turbulent dissipation is elevated 10 to**  
 10 **100-fold (reaching  $10^{-6}$ - $10^{-5}$  W kg<sup>-1</sup>) every night during due to the swimming activity of**  
 11 **large aggregations of anchovies that gather regularly over the spawning season. Turbulent**  
 12 **mixing is invigorated concurrently to dissipation, and occurs with an efficiency comparable**

13 **to that of geophysical turbulence. Our results demonstrate that biologically-driven turbu-**  
14 **lence can be a highly effective mixing agent, and call for a re-examination of its impacts on**  
15 **productive upper-ocean regions.**

16 Turbulence is a fundamental component of the ocean's energy budget, for it mediates the  
17 transfer of kinetic energy from large (1-1000 km) to small (0.1-1 cm) scales, where such energy is  
18 dissipated as heat by molecular viscosity<sup>1</sup>. However, dissipation is just one of two possible fates  
19 of turbulent kinetic energy (TKE). As turbulent motions stir the water column, microscale phys-  
20 ical and chemical gradients are generated and ultimately eroded by molecular diffusion, thereby  
21 resulting in mixing. When acting on a stable density profile, such as the oceanic pycnocline, tur-  
22 bulent mixing drives an upward transport of mass, and hence transforms a fraction of the TKE  
23 into potential energy<sup>2</sup>. This fraction, referred to as mixing efficiency, is relatively uncertain and  
24 challenging to quantify *in situ*. Observations and idealized simulations indicate that the mixing  
25 efficiency often approaches a value of  $\sim 0.16$  (refs.3, 4), as is characteristic of shear instabilities  
26 (a major source of ocean turbulence<sup>5,6</sup>), yet there is mounting evidence that the mixing efficiency  
27 may vary extensively<sup>7</sup>.

28 The debate surrounding mixing efficiency is particularly relevant for appraising the signifi-  
29 cance of biomixing<sup>8</sup>. Although winds and tides undeniably constitute the major sources of energy  
30 for ocean mixing on a global scale<sup>9</sup>, it has been suggested<sup>10,11</sup> that swimming organisms (from  
31 zooplankton to fish and marine mammals) may also contribute a substantial energy input<sup>12</sup>, at least  
32 on regional scales<sup>13</sup>. The relevance of this suggestion was initially endorsed by dynamical<sup>13</sup> and  
33 metabolic<sup>14</sup> considerations, laboratory experiments<sup>15</sup>, and early observations of elevated TKE dis-  
34 sipation ( $\sim 10^{-5}$  W kg<sup>-1</sup>) in fish aggregations<sup>16</sup> and migrating krill swarms<sup>17</sup>. However, subsequent  
35 studies found biophysical turbulence extremely challenging to capture in lakes and oceans, indicat-  
36 ing that this phenomenon might be rarer than originally thought<sup>18-22</sup>. Further, what little evidence  
37 exists of mixing produced by biophysical turbulence suggests that the mixing efficiency of such

turbulence is very low ( $<0.01$ ) relative to that of geophysical, shear-driven turbulence. This evidence is based on the concurrent measurement of the rates of dissipation of TKE ( $\varepsilon$ , a measure of the intensity of turbulence) and of thermal variance ( $\chi$ , a measure of the intensity of mixing)—respectively quantified from observations of centimetre-scale velocity and temperature gradients—in the presence of swimming organisms<sup>16,20,22–24</sup>. Only two of these investigations reported high  $\varepsilon$  levels within fish aggregations<sup>23,24</sup> but, in both cases, these were associated with low values of  $\chi$ , i.e. weak mixing. Thus, the present balance of evidence points toward dismissing the old-proposed view of a significant large-scale influence of biomixing.

Here, we contest and redress this balance by demonstrating the occurrence of recurrent, intense and efficient biomixing in an embayment affected by wind-driven coastal upwelling pulses (Ría de Pontevedra, NW Iberia, Extended Data Figure 1)<sup>25,26</sup>. This demonstration rests on the analysis of a two-week data set of highly (temporally and vertically) resolved observations of hydrographic properties, turbulent dissipation and mixing rates, and acoustic backscatter (an indicator of fish and plankton density). The data were acquired in the summer of 2018 in three sampling periods (I01, 1–5 July; I02, 6–8 July; and I03, 9–13 July; see Methods for details of data set), and captured intense biophysical turbulence in every segment of nocturnal measurements. This allowed for an unprecedentedly detailed characterisation of biophysical turbulence, its mixing efficiency, and its biological underpinning.

## Hydrographic and turbulence environments

The Ría's hydrographic setting varied notably during the measurement campaign, evolving from a downwelling to an upwelling circulation over the fortnight of observations (Figure 1a,b). In the first sampling period (I01), southerly, downwelling-favorable winds were dominant (Figure 1a) and thermal stratification was relatively weak (Figure 1b) as a result of the import into the embay-

ment of surface shelf waters with uniform temperature of  $\sim 17^\circ\text{C}$ . After 4 July, calm conditions prevailed, and the circulation pattern reversed (Extended Data Figure 2). During I02, relatively cold waters ( $T \approx 13^\circ\text{C}$ ) upwelled into the Ría's deeper layers, giving rise to a stratified interface at  $\sim 20$  m. The inflow of warmer and fresher waters produced an additional near-surface ( $< 10$  m) stratified layer (Figure 1b). At the outset of I03, strong northerly winds led to an intensification of cold-water upwelling, which brought about a single highly-stratified layer.

The evolution of the Ría's turbulence environment bared little imprint from that of the hydrography, hinting at a non-physical origin of the turbulence. Thus, turbulent dissipation was recurrently enhanced over the entire water column every day after sunset and for a period of 5-6 hours (Figure 1c). This enhancement (termed night-time dissipation hereafter) was most striking in the embayment's interior layers (10 – 25 m), away from the direct influence of wind-induced and bottom boundary turbulence. Within this depth interval, night-time  $\varepsilon$  was elevated by 1-3 orders of magnitude above background daytime values of  $10^{-9} - 10^{-8} \text{ W kg}^{-1}$ , reaching  $10^{-7} - 10^{-5} \text{ W kg}^{-1}$  (Figure 1c, Extended Data Table 1). Background  $\varepsilon$  levels were higher during I01 (mean  $1.22 \times 10^{-7} \text{ W kg}^{-1}$ ) compared to I02 ( $1.71 \times 10^{-8} \text{ W kg}^{-1}$ ) and I03 ( $2.17 \times 10^{-8} \text{ W kg}^{-1}$ ). Night-time dissipation rates also decreased between I01 (mean  $1.62 \times 10^{-5} \text{ W kg}^{-1}$ ) and I02-I03 ( $1.96 \times 10^{-6}$  and  $0.82 \times 10^{-6} \text{ W kg}^{-1}$ , respectively).

## Sources of turbulence

To assess the energy sources of the turbulence in the Ría, we first examine the extent to which geophysical factors may explain the measured dissipation patterns. Geophysical turbulence in density-stratified waters commonly occurs when the destabilizing effect of vertical gradients of horizontal velocity (shear,  $sh^2$ , see Methods) overcomes the stabilizing effect of the vertical density gradient (stratification,  $N^2$ ). Shear instability and turbulence are predicted to develop for low,

subcritical values of the gradient Richardson number,  $Ri_g = N^2/sh^2 < 1/4$  (ref. 27). In our observational record, episodes of high near-surface  $\varepsilon$  (Figure 1c) were associated with intensified winds in particular days (Figure 1a), suggesting a physical driving of those turbulent patches. However, the occurrence of unstable conditions below 10 m did not exhibit a day-night cycle, as would be expected if shear instabilities generated the recurrent events of night-time dissipation. Instead, subcritical values of  $Ri_g$  within the water column became progressively rarer over the course of the experiment, as stratification increased (Figure 1e). Unstable conditions were relatively frequent and widespread during downwelling (I01), but retreated to the upper and bottom boundary layers during upwelling (I02 and I03). This disassociation between the observed turbulence and shear instabilities is succinctly illustrated by the weak correlation between  $\varepsilon$  and  $Ri_g$  (Spearman  $r = -0.14$ ,  $p < 0.01$ , Extended Data Figure 3), which endorses the notion that the night-time dissipation was sustained by a non-physical energy source.

A window into the nature of such source is provided by the distribution of volume backscattering strength ( $S_v$ ), a metric of the occurrence of fish, recorded with a vessel-mounted echosounder.  $S_v$  was systematically enhanced at night (Figure 1f), in remarkable concurrence with elevated turbulent dissipation. The intensity of backscatter was highly correlated with  $\varepsilon$  for all the sampled backscattering frequencies (18-200 kHz,  $r = 0.56 - 0.67$ ,  $p < 0.01$ , Extended Data Figure 3), suggesting that the night-time dissipation events were driven by fish aggregations. Consistent with this interpretation, high concentrations of European anchovy (*Engraulis encrasicolus*) eggs were detected in plankton net hauls that were performed every morning of the experiment and in the night of 8 July (Figure 2). Local spawning of the eggs is indicated by both sets of hauls. In most of the morning hauls, the majority of eggs presented an F2 development stage (indicative of a time elapsed since spawning of 4–14 hours), whereas the night-time haul (01:54 pm GMT) was dominated by freshly spawned eggs at stage F1 (corresponding to a time since spawning of  $< 4$  hours). Note, though, that lack of fish sampling gear on board prevented us from obtaining direct evidence

of the presence of fish.

At any rate, the picture that emerges from the net samples is one of a nocturnal aggregation of anchovies for spawning being responsible for driving our observed episodes of night-time dissipation. This view is endorsed by the acoustic frequency response (FR) within the nocturnal turbulent patches, which was elevated at 18 kHz compared to higher frequencies (Extended Data Figure 4), as previously described for anchovy aggregations<sup>28</sup>. It is also advocated by previous reports of such aggregations inside the Rías of NW Iberia<sup>29,30</sup>, which reveal spawning to occur between 19:00 and 6:00 GMT (peaking at midnight)<sup>31</sup>, with a seasonal maximum in July-August<sup>32</sup>. Thus, in the following, we will consider these episodes of intense nocturnal dissipation as being triggered by biophysical turbulence, and daytime periods of weaker dissipation as being dominated by geophysical turbulence.

### Efficiency of turbulent mixing

As intense as the night-time biophysical turbulence may have been, did it effect commensurately substantial mixing? To address this question, we next characterize the mixing intensity by examining the record of temperature microstructure. This shows that the nocturnal biophysical turbulence events were associated with elevated values of the small-scale temperature-gradient variance (Extended Data Figure 5), the rate of thermal variance dissipation ( $\chi$ , Extended Data Figure 6a) and the rate of turbulent mixing of heat (quantified by the diffusivity  $K_T$ , Figure 1d). This mixing enhancement was most evident during I03 and I02, which exhibited a >10-fold increase in temperature-gradient variance above daytime levels over a broad wavenumber range (Extended Data Figure 5), as well as an amplification of  $K_T$  by two orders of magnitude (relative to daytime values of  $K_T < 10^{-6} \text{ m}^2 \text{ s}^{-1}$ , Figure 1d). The mixing impact of biophysical turbulence was muted during I01, for which daytime turbulence was more energetic than for I02-I03 (Figure 1d).

The mixing efficiency, defined here as the fraction of TKE converted to potential energy, is assessed for our entire data set by computing the flux Richardson number,  $R_f = K_T N^2 / (\varepsilon + K_T N^2)$ , where  $K_T N^2$  and  $\varepsilon$  are respectively evaluated from microstructure measurements of temperature gradient and shear.  $R_f$  varied by up to three orders of magnitude during our observational period, including episodes of both geophysical and biophysical turbulence (Figure 3a). Mixing efficiencies close to the canonical value of  $R_f = 0.16$  for geophysical, shear-driven turbulence were observed in the uppermost 10 m (mean  $R_f = 0.153$  [0.150 – 0.156] [95% confidence intervals]), where turbulence is energized directly by wind. Below the surface (10–25 m), the frequency distribution of  $R_f$  was centered at lower values (mode  $R_f \lesssim 0.1$ ), with broadly similar distributions for daytime geophysical turbulence and night-time biophysical turbulence (Figure 3a). The  $R_f$  distribution for geophysical turbulence was slightly less negatively skewed, as values larger than the mode ( $R_f \approx 0.1$ ) were more frequent than in the  $R_f$  distribution for biophysical turbulence. The average mixing efficiency during night-time biophysical turbulence events ( $R_f = 0.067$  [0.064 – 0.069]) was smaller than, but not significantly different to, the average value of  $R_f$  for background geophysical turbulence in the same depth interval ( $R_f = 0.088$  [0.086 – 0.090]). Our data thus demonstrates that, contrary to the common view at present<sup>24</sup>, biophysical turbulence can be a comparably efficient mixing agent to geophysical turbulence.

## Discussion

Our observations reveal a consistent occurrence of elevated night-time levels of biophysical turbulence (reaching rates of dissipation as high as  $10^{-6} - 10^{-5} \text{ W kg}^{-1}$ , or two orders of magnitude above daytime values) during a two-week stretch, providing compelling evidence that fish can generate intense turbulence over prolonged periods. This contrasts with the results of several past investigations, which found biophysical turbulence challenging to detect in the field<sup>19,20,22</sup>. There are, however, several threads of evidence to propose that the representativeness of our results tran-

scends the specific spatio-temporal context of our measurements. First, our observed dissipation rates are similar to those documented previously in migrating zooplankton and krill layers<sup>17,33</sup> and fish aggregations<sup>23,24,34</sup>. They also conform to predictions by empirical models of biophysical turbulence which, on the basis of the size and characteristics of the swimming organisms<sup>13,20</sup>, predict a rate of TKE production by the observed aggregations of  $\sim 10^{-6} \text{ W kg}^{-1}$  (see Methods). Finally, the aggregating fish behaviour that was responsible for the intense biophysical turbulence in our data is a recurrent feature in coastal upwelling areas in NW Iberia<sup>29</sup> and elsewhere<sup>35</sup>. This is illustrated by measurements of acoustic backscatter acquired by a moored acoustic current profiler in our study area between 26 June and 19 July, which revealed an enhancement of backscatter at almost every night in that extended period (Extended Data Figure 7).

An important result of the present work pertains to the observation of intense biophysical turbulence with a mixing efficiency comparable to that of geophysical turbulence. This finding challenges expectations from several previous studies that pointed to a reduction in mixing efficiency, and in the rate of mixing itself, in association with biophysical turbulence<sup>23,24</sup>. A theoretical explanation for such reduction was provided by Visser<sup>8</sup>, who concluded that biophysical turbulence must necessarily be inefficient because the most abundant swimming organisms (zooplankton and fish) are small, and so produce small turbulent eddies (0.01 – 0.1 m). As these overturns would not be sufficiently large to interact with buoyancy forces, they would not induce mixing, but would be rapidly destroyed by viscosity instead.

To unravel the discrepancy between Visser’s argument and our results, we examine the relationship between the mixing efficiency (again, quantified by  $R_f$ ) and a set of key turbulent parameters<sup>36</sup>. This entails projecting our data onto a space defined by the turbulent Reynolds ( $Re_T$ ) and Froude ( $Fr_T$ ) numbers<sup>37</sup> (Figure 3b). We approximate these two numbers by ratios of length scales that describe the competition between inertial, buoyancy and viscous forces in the fluid,



180 following Ivey and Imberger<sup>37</sup>. While this simplified approach has some formal limitations<sup>38</sup>,  
 181 these do not affect our analysis' findings (see discussion in Methods). The turbulent Reynolds  
 182 number, computed here as  $Re_T = (L_T/L_K)^{4/3}$ , represents the ratio of the mean size of the energy-  
 183 containing eddies (denoted by the Thorpe scale,  $L_T$ ; see Methods) to the viscosity-dominated Kol-  
 184 mogorov scale ( $L_K$ ), and measures the competition between inertia and viscosity. The turbulent  
 185 Froude number, here defined as  $Fr_T = (L_O/L_T)^{2/3}$ , quantifies the size of the observed turbulent  
 186 eddies relative to the buoyancy or Ozmidov scale ( $L_O = (\varepsilon/N^3)^{1/2}$ ), at which turbulent eddies are  
 187 strongly influenced by buoyancy<sup>39</sup>. Thus, efficient mixing is expected when  $L_T \approx L_O$  ( $Fr_T \approx 1$ ),  
 188 such that eddies interact with buoyancy forces and transport mass across the mean density gradi-  
 189 ent, and when  $Re_T$  is sufficiently large ( $Re_T \gtrsim 100$ ), such that overturning motions are not readily  
 190 damped by viscosity before they induce mixing<sup>40</sup>. According to Visser<sup>8</sup>, biophysical turbulence is  
 191 characterized by  $Fr_T \gg 1$  (i.e.  $L_T \ll L_O$ ), such that  $R_f \ll 0.16$ .

192 Bin-averaging our observational estimates of  $R_f$  in  $Re_T$  -  $Fr_T$  space reveals that the mix-  
 193 ing efficiency was maximal ( $R_f \geq 0.16$ ) when  $Re_T > 100 - 1000$  and  $Fr_T \approx 1$  (Figure 3b), as  
 194 expected<sup>37,39</sup>. These energetic conditions were only found in the surface layer ( $< 10$  m) directly  
 195 energized by the wind. During daytime, geophysical turbulence in the interior (10–25 m) was char-  
 196 acterized by  $Re_T < 100 - 1000$  and  $Fr_T \approx 0.5$ , which signal conditions under which turbulence is  
 197 susceptible to suppression by buoyancy ( $L_T > L_O$ ) and viscosity<sup>40</sup>, such that the mixing efficiency  
 198 might be curbed ( $R_f \lesssim 0.1$ ). Night-time biophysical turbulence was more energetic ( $Re_T \approx 1000$ ),  
 199 and contained overturns smaller than the Ozmidov scale ( $L_T < L_O$ ,  $Fr_T > 1 - 2$ ). For the I01 sam-  
 200 pling period, when relatively weak stratification ( $N^2 \approx 2 \times 10^{-4} \text{ s}^{-2}$ , Extended Data Table 1) and  
 201 elevated  $\varepsilon$  resulted in enhanced  $L_O \approx 1$  m and  $Fr_T = 2.63$  (Extended Data Table 1), fish-induced  
 202 eddies ( $L_T \approx 10 - 20$  cm, in line with the typical adult anchovy size in our study area<sup>41</sup> of  $\sim 12$  cm)  
 203 may have been too small to act on the background stratification, and the locus of the data in  $Re_T$  -  
 204  $Fr_T$  space suggests a reduction of  $R_f$ . In contrast, for I02 and I03,  $L_O$  was smaller ( $\sim 20$  cm) owing

to a decrease in  $\varepsilon$  and a strengthening of stratification ( $N^2 = 5 - 10 \times 10^{-4} \text{ s}^{-2}$ ), and approached the size of the fish-driven turbulent overturns ( $Fr_T \gtrsim 1$ ). Under these circumstances, the locus of the data in  $Re_T - Fr_T$  space indicates that  $R_f \approx 0.1$ , i.e. in line with the mixing efficiency of geophysical turbulence. In summary, Visser’s argument on the presumed inefficiency of biophysical turbulence does not hold in our observations because, contrary to the argument’s assumptions,  $L_O$  and  $L_T$  are comparable in a large portion of our data set (Figure 4).

Laboratory experiments suggest that the condition,  $L_O \sim L_T$ , upon which efficient biomixing is contingent, is promoted by aggregations of swimmers, which can produce large, aggregation-scale turbulent eddies<sup>15,42</sup>. Conversely, our observations suggest an alternative route toward efficient biomixing via an increase in stratification, which reduces buoyancy length scales (i.e.  $L_O$ ) to the point that they become comparable to turbulent eddy sizes, even if these are significantly smaller than the aggregation scale. Our work thus shows that, besides biological factors –the agitated behaviour of the anchovies while spawning may have also played a role<sup>42</sup>–, the mixing efficiency of biophysical turbulence is controlled by the background stratification facilitating the injection of fish-induced TKE at the required scales. This conclusion implies that, while biomixing might be inefficient within the main open-ocean pycnocline (where  $L_O \approx 1 \text{ m}$ ; refs.<sup>8,11</sup>), it is likely to be considerably more effective in environments with stronger stratification, e.g., in seasonal pycnoclines or coastal seas (Figure 4). Such proposition appears particularly plausible in coastal regions, where riverine freshwater sources, solar heating and upwelling often give rise to highly stratified conditions<sup>43</sup>, and spawning aggregations of small pelagic fish (e.g., sardine, herring or anchovy) are especially abundant<sup>35,41,44,45</sup>. Hence, our results not only substantiate the prediction that fish aggregations can generate intense turbulent dissipation levels, comparable to storms<sup>13</sup>, but also show that elevated stratification fosters the occurrence of efficient biomixing. This highlights the potential of biophysical turbulence to drive enhanced vertical exchanges in upper-ocean areas with a rich biota, in many of which (e.g., in strongly stratified coastal waters)

vertical turbulent transfers contribute to the supply of nutrients for plankton growth<sup>46</sup> or the ventilation of de-oxygenated waters<sup>47</sup>. Therefore, biomixing could contribute promote phytoplankton growth and to reduce low-oxygen stress for higher trophic levels. The role of biomixing in shaping the physical and biogeochemical properties of productive upper-ocean regions should thus be reassessed.

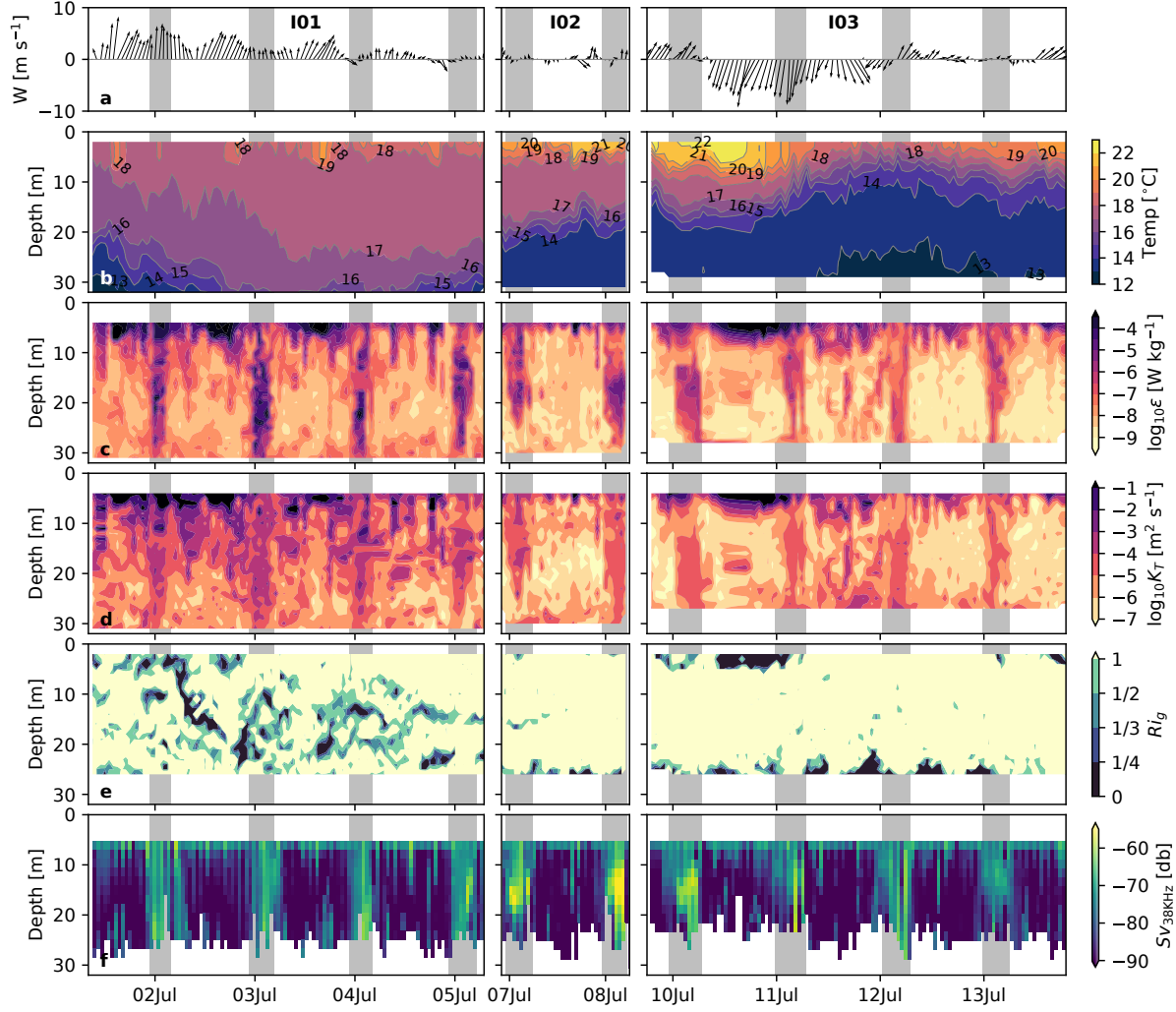
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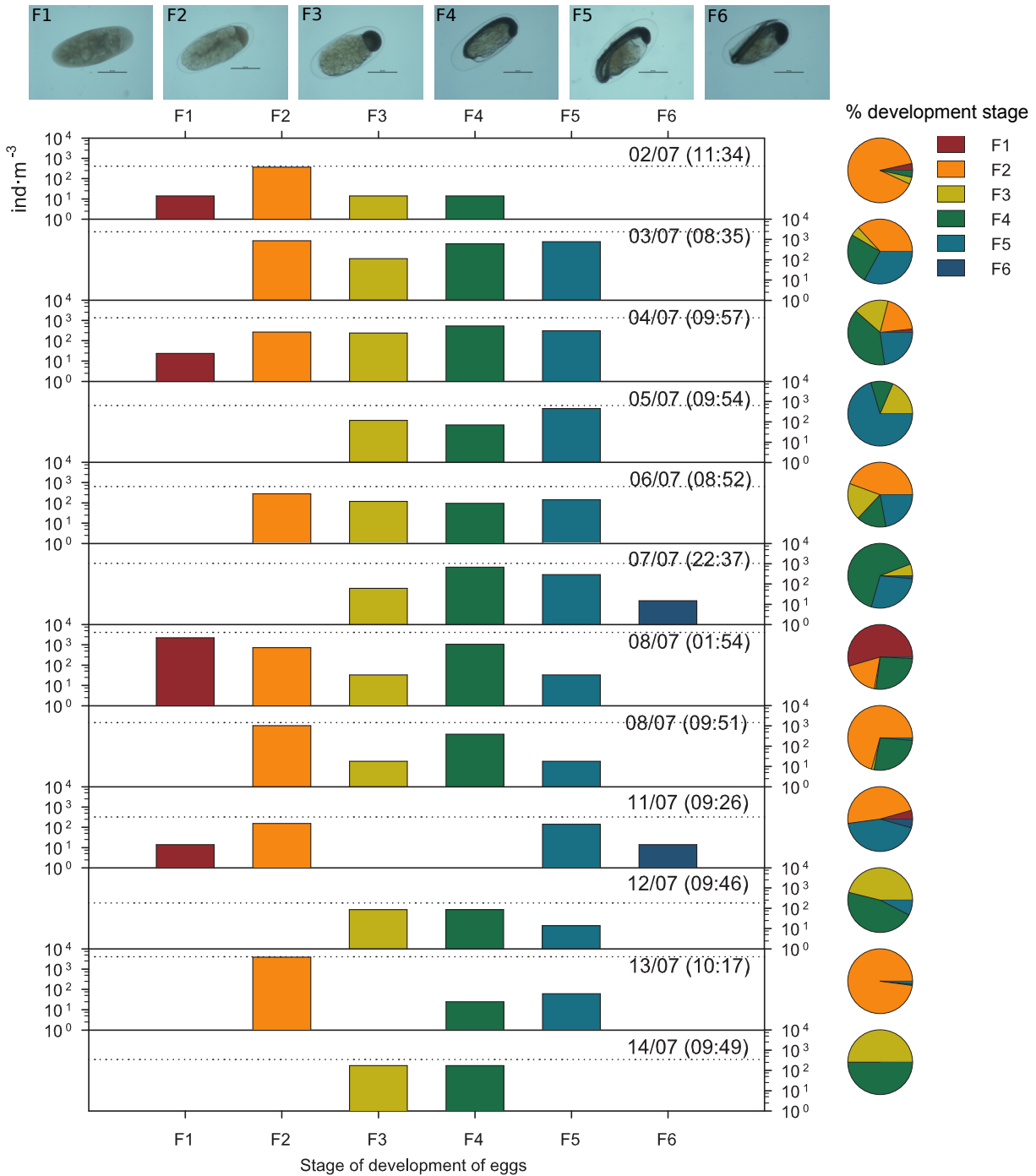
**Author contributions.** BFC and BMC conceived the study. BMC led the cruise. BFC, EN, MG, EB, AC and BMC participated in the data collection. BFC, MP, EN and MG analysed the data. All the authors contributed to the scientific discussions and interpretation of the results. BFC, BMC and ACNG wrote the

255 manuscript with contributions from all the coauthors.

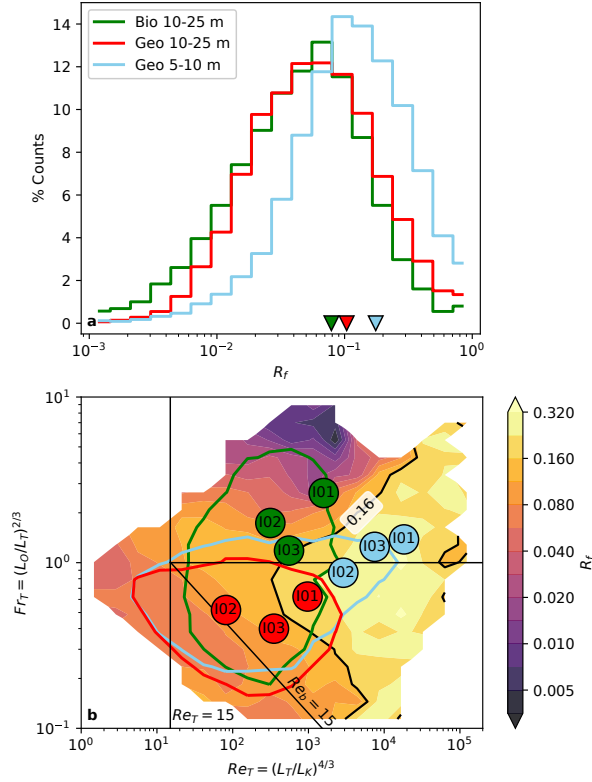
256 **Competing Interests.** The authors declare that they have no competing financial interests.



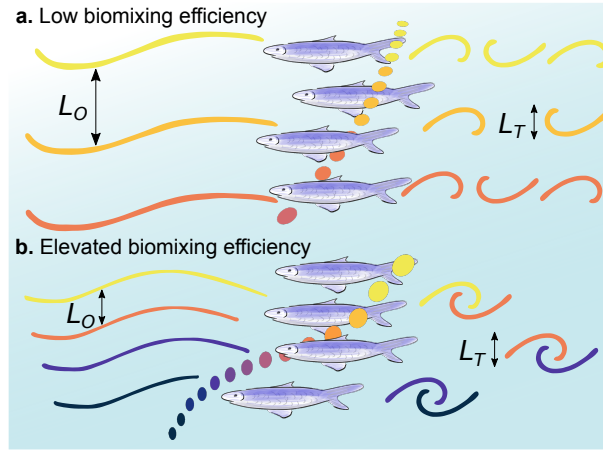
**Figure 1: Hydrography, turbulence and mixing during the REMEDIOS survey.** Time series of **a** local wind speed ( $W$ ) and direction measured at Cape Udra (Extended Data Figure 1), and hourly-mean **b** temperature (Temp.), **c** turbulent kinetic energy dissipation rate ( $\epsilon$ ), **d** turbulent heat diffusivity ( $K_T$ ), **e** gradient Richardson number ( $Ri_g$ ), and **f** volume backscattering strength at 38 kHz ( $Sv$ ), during the three sampling periods (I01, I02 and I03). Gray shading indicates night-time periods of biomixing. These periods were determined by inspection of the turbulent dissipation rate and volume backscattering strength records. The time axis is GMT time (local time = GMT + 2 hours). Note the use of logarithmic scale in panels **c**, **d**.



**Figure 2: Anchovy egg concentration.** Number of individuals per cubic metre ( $\text{ind m}^{-3}$ ) at different development stages (F1 to F6, Extended Data Table 2) for the time series of plankton net hauls carried out at the sampling location. The date and hour of sampling (GMT), and the total number of eggs (horizontal dotted line) are shown in each bar plot. The y-axis is displayed in logarithmic scale. The pie charts indicate the percentages of eggs at the development egg stages considered, relative to the total number of eggs in each sample. The horizontal black line in each egg image (in the upper axis) denotes a length of 0.5 mm. Average elapsed times since spawning corresponding to the different development stages according to ref. <sup>48</sup> are: 0 hours (F1), 4.3 (F2), 14.3 (F3), 31.5 (F4), 48.3 (F5), and 59.3 (F6).



**Figure 3: Mixing efficiency.** **a** Frequency distribution of the flux Richardson number ( $R_f$ , a measure of the mixing efficiency) in the water column interior (10-25 m depth range) in periods dominated by biophysical turbulence (green) and in periods dominated by geophysical turbulence (red), and in the near-surface layer (5-10 m) for the full time series (light blue). Mean values are indicated with triangles. **b** Bin-averaged  $R_f$  in the Reynolds number – Froude number space ( $Re_T - Fr_T$ ). The high-efficiency  $Fr_T = 1$  (horizontal line), and molecular  $Re_T = 15$  (vertical) and  $Re_b = 15$  (oblique;  $Re_b = (L_O/L_K)^{4/3}$ ) limits described by Ivey and Imberger<sup>37</sup> are shown as black lines. Mean values of  $Re_T$  and  $Fr_T$  are shown as circles for the three sampling periods (I01, I02 and I03), and for the different depth ranges. The green, red and light blue lines enclose bins in which frequency of occurrence exceeds 0.5% of the total counts for periods of biophysical turbulence (green, 10-25 m), and interior (red, 10-25 m) and near-surface (light blue, 5-10 m) geophysical turbulence.



**Figure 4: Schematic of the onset of efficient biomixing.** Schematic representation of two scenarios of biophysical turbulence in contrasting levels of stratification, underpinning low (a) and elevated (b) mixing efficiency. In the first scenario, weak stratification (illustrated here by the colored lines and dots depicting layers of different temperature) results in a large buoyancy length scale ( $L_O$ ) compared to the biologically-induced overturning scale ( $L_T$ ). In this scenario turbulent eddies are dissipated by viscosity before producing significant mixing, and biophysical mixing is inefficient. Such conditions, mimicking those described by Visser<sup>8</sup> for the main open-ocean pycnocline, are broadly captured by our observations during I01. In the second scenario, representing conditions during I02-I03,  $L_O$  shrinks as stratification increases, becoming comparable to the overturning scale. This situation allows the turbulent eddies to interact with the temperature/density profile and transport heat/mass vertically before dissipating, thus leading to an increase in the efficiency of biomixing.



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## Methods

**Sampling overview.** The REMEDIOS sampling campaign was carried out off the Galician coast (NW Iberian Peninsula) between 29 June and 18 July 2018 on board of R/V Ramón Margalef (Extended Data Figure 1). Three intensive sampling time series were performed: I01 (2018.07.02 08:00 am to 2018.07.06 08:40 am), I02 (2018.07.07 10:07 pm to 2018.07.09 05:23 am), and I03 (2018.07.10 5:50 pm to 2018.07.14 7:30 pm) at station P2-Bueu, inside the Ría de Pontevedra (42.357°N, -8.773°E, mean depth 30 m). During these sampling periods, five casts were performed every half hour with a microstructure profiler (MSS<sup>49</sup>), resulting in a total number of 1658 profiles (~50 km of microstructure data). This sequence was interrupted every 6 hours for water collection with a Rosette. Water velocity profiles were continuously recorded with a bottom-moored acoustic Doppler current profiler (ADCP), and volume backscattering strength was registered with a hull-mounted echosounder.

**Microstructure measurements.** The MSS was equipped with two shear microstructure sensors (type PNS06) and a temperature microstructure sensor (type FP07), complemented with a high-accuracy CTD and an accelerometer to assess the instrument's vibration. The various channels were sampled at 1024 Hz, and the instrument was loose-tethered and operated in free-falling mode at a nominal vertical speed of 0.6–0.7 m s<sup>-1</sup>. The dissipation rates of turbulent kinetic energy ( $\varepsilon$ ) and thermal variance ( $\chi$ ) were calculated by integrating the vertical shear and vertical temperature-gradient spectra over half-overlapping segments of 2 m. The shear microstructure signal was de-noised for instrument vibration using the accelerometer signal<sup>50</sup>. A pseudo-shear signal was also derived from the accelerometer to assess  $\varepsilon$  contamination by instrument vibration. The temperature-gradient spectra were corrected for the FP07 time-response ( $\tau = 12$  m s<sup>-1</sup>) with a double-pole function, previous to integration<sup>51</sup>. The shear spectra were integrated from a minimum wavenumber of 2 cycles per meter (cpm). The upper integration limit was estimated iteratively from an initial guess of 14 cpm until convergence to the Kolmogorov wavenumber

379  $(k_c = \frac{1}{2\pi} (\varepsilon \nu^{-3})^{1/4})$  was achieved. The upper cut-off was curtailed to a maximum value of 30 cpm,  
 380 in order to avoid the spectral region where the spatial response of the shear probe becomes lim-  
 381 iting and noise is introduced by instrument vibrations. The temperature gradient spectra were  
 382 also integrated from 2 cpm. The upper integration limit in this case was chosen as the minimum  
 383 of the following wavenumbers: the wavenumber at which the measured spectra becomes smaller  
 384 than two times an empirical noise spectrum (see Extended Data Figure 5, Supplementary Fig-  
 385 ures 1, 2); the wavenumber at which the time-response correction is larger than a factor of 100;  
 386 and a wavenumber corresponding to a frequency of 60 Hz. The missing variance at wavenumbers  
 387 beyond the integration limits was estimated by assuming that the shear and temperature spectra  
 388 respectively follow the empirical Nasmyth and Batchelor forms<sup>52</sup>. Prior to the missing-variance  
 389 correction,  $\varepsilon$  values were corrected for the probe's spatial response using the polynomial factors  
 390 given by the manufacturer<sup>53</sup>. Note that the variance correction can be substantial for high levels of  
 391 dissipation (i.e.  $\varepsilon > 10^{-6} \text{ W kg}^{-1}$ ), however, the measured spectra show excellent agreement with  
 392 the empirical universal forms for intense turbulence over the resolved wavenumber range (Supple-  
 393 mentary Figure 1), supporting the robustness of the estimation. This methodology is described in  
 394 more detail elsewhere<sup>54</sup>. During night-time biomixing periods, fish impacts on the profiler were  
 395 often apparent in the shear and pseudo-shear (accelerometer) records (Supplementary Figure 2).  
 396 These data segments were manually identified and discarded from further analysis. Impacts were  
 397 frequent during I01 (1772 of 2829 segments discarded), but rare during I02 (129 of 1866 segments  
 398 discarded) and I03 (114 of 3378 segments discarded) (Extended Data Table 1).

399 **Mixing characterization.** The stability with respect to shear-driven turbulence was characterized  
 400 with the gradient Richardson number,  $Ri_g = N^2/sh^2$ , where  $N^2 = -g/\rho(\partial_z \rho)$  is the buoyancy  
 401 frequency, with  $\rho$  as the potential density, and  $sh^2 = (\partial_z u)^2 + (\partial_z v)^2$  is the squared vertical shear,  
 402 with  $u$  and  $v$  as the zonal and meridional velocity components. The turbulent diffusivity for heat  
 403 was calculated using the Osborn-Cox<sup>55</sup> relation as  $K_T = 0.5\chi/(\partial_z T)^2$ , where  $\partial_z T$  is the back-

ground temperature gradient. Mixing efficiency was quantified with the flux Richardson number ( $R_f = K_T N^2 / (\varepsilon + K_T N^2)$ ), and the vertical size of turbulent overturns with the Thorpe length scale ( $L_T$ ).  $L_T$  was computed by comparing the measured potential density with an adiabatically resorted density profile<sup>56</sup>. Other relevant scales for turbulence are the Ozmidov ( $L_O = (\varepsilon N^{-3})^{1/2}$ ) and the Kolmogorov ( $L_K = (\nu^3 \varepsilon^{-1})^{1/4}$ ) length scales, where  $\nu$  is the kinematic viscosity,  $\sim 10^{-6} \text{ m}^2 \text{ s}^{-1}$ .

**Acoustic backscatter and fish density.** A Simrad EK80 echosounder operated the split-beam transducers of 18, 38, 70, 120 and 200 kHz at continuous wave mode with maximum pinging rate, registering 50 metres of data in the vertical. Pulse duration was set to 1 ms for all the frequencies, while the beam width was  $11^\circ$  for 18 kHz and  $7^\circ$  for higher frequencies. Transmitting power was 1600, 1600, 675, 225 and 135 W, respectively, for each frequency. The standard sphere calibration procedure was performed after the survey<sup>57</sup>. A mean volume backscattering coefficient ( $S_v$ , dB re  $1 \text{ m}^2 \text{ m}^{-3}$ , dB hereafter) was averaged (in the linear domain) in bins of 2 metres by 30 minutes, after removing the 6 metres where ringing noise (remaining vibration of the echosounder while already listening) affected the 18 kHz echogram.

**European anchovy eggs** Sampling of *Engraulis encrasicolus* eggs was performed by means of oblique hauls from the surface down to 3 m above the bottom ( $\sim 30 \text{ m}$  depth) with a double-WP2 plankton net (HydroBios;  $3.95 \text{ m}^2$  mouth area;  $200 \mu\text{m}$  mesh-size), at a descending/ascending rate of 50/30 metres per minute and a trawling speed of 2 knots. Each net carried a mechanical flowmeter (General Oceanics) to estimate the volume of sampled water (between ca. 17 and  $32 \text{ m}^3$ ). Collected samples were preserved with buffered formaldehyde (4% final concentration). Counting and definition of the development state of eggs were carried out with a stereoscopic binocular (Nikon SMZ-10). Sample aliquots of 20 mL, from a solution of 400 mL of the whole sample, were used for those purposes. To illustrate development state (Extended Data Table 2), microphotographs were taken with a stereoscopic binocular and dedicated image acquisition software (Nikon SMZ-1270 and NIS-Elements). Development state was classified in stages according



to morphological properties of the embryo<sup>58</sup>. The eggs collected during the survey were classified into 6 stages (F1 to F6), focusing on the characterization of development in the first 12 hours after spawning and considering a reference temperature of 17 °C (ref. <sup>48</sup>). Stages applied here (F1 to F6), their equivalence to the stages ('stageing') proposed in ref. <sup>58</sup> (I to XI), stage duration after spawning in hours (for 17 °C ambient temperature) and morphological characteristics of the embryo [from Table A1.3 of ref. <sup>41</sup>] are given in Extended Data Table 2.

**Currents and continuous backscatter.** A RD Instruments ADCP (300 kHz) was bottom-moored looking upwards in station P2-Bueu before the start of the cruise (2018.06.26), and was recovered after the end of the cruise (2018.07.19) from R/V Kraken. The three-dimensional current was recorded every 5 minutes as the average of 120 individual pings in 70 layers of 0.5 m, spanning the water column from 4 m above the bottom to the surface. ADCP backscatter was converted to volume backscattering strength ( $S_v$ , dB) following refs.<sup>60,61</sup>.

**TKE production by anchovy aggregations.** TKE production by the anchovy aggregations was estimated as  $P_{TKE} = 0.072/3 n U^{14/5} L^{9/5} \nu^{0.2} \approx 10^{-6} \text{ W kg}^{-1}$ , following refs.<sup>13,20</sup>, where  $\nu \approx 1 \times 10^{-6} \text{ m}^2 \text{ s}^{-1}$  is the molecular viscosity of seawater. A characteristic fish size of  $L = 12 \text{ cm}$  (corresponding to the modal size of 1-year-old anchovy adults<sup>41</sup>) and a swimming speed of  $U \approx 36 \text{ cm s}^{-1}$  (equivalent to three body lengths per second<sup>64</sup>) were used in the calculation. Fish concentration in the aggregation ( $n$ , individuals per  $\text{m}^3$ ) was calculated using the relationship between target strength for a given species (TS) and the observed volume backscattering ( $S_v = TS + 10 \log_{10}(n)$ ). Using the target strength at 38 kHz employed for the PELACUS surveys in this area<sup>41</sup> ( $TS = 20 \times \log_{10}(L) - 72.6 = -51.02$ ), the recorded mean  $S_v$  in the shoal at 38 kHz (-53.63 dB for I02) would convert into a concentration of  $n = 0.5$  individuals per  $\text{m}^3$ .

**Mixing efficiency in the  $Re_T$ - $Fr_T$  diagram** In order to synthesize the variability of the flux Richardson number ( $R_f$ , a measure of the mixing efficiency) in our cruise measurements, and

453 rationalize the relatively high efficiency diagnosed for biophysical mixing, we placed our data in  
 454 a Reynolds-Froude ( $Re_T$ - $Fr_T$ ) numbers diagram. Projection onto  $Re_T$ - $Fr_T$  space is useful to de-  
 455 scribe the balance of forces in a turbulent fluid which, in turn, underpins the efficiency of mixing.  
 456 The state-of-the-art profiling instruments enabling quantification of turbulence in natural waters,  
 457 such as the MSS employed in this study, do not directly measure  $Re_T$  and  $Fr_T$ . Thus, following  
 458 common practice in ocean turbulence works<sup>37</sup>, we approximate these quantities using turbulence  
 459 length scales that can be directly assessed from the measurements: the Ozmidov ( $L_O = (\varepsilon N^{-3})^{1/2}$ ),  
 460 Kolmogorov ( $L_K = (\nu^3 \varepsilon^{-1})^{1/4}$ ) and Thorpe ( $L_T$ , see Methods) length scales, such that:

$$Re_T = \left( \frac{L_T}{L_K} \right)^{4/3} \quad (1)$$

461 and

$$Fr_T = \left( \frac{L_O}{L_T} \right)^{2/3}. \quad (2)$$

462 Recent evidence from direct numerical simulations<sup>38</sup> (DNS) suggests that these approximate  
 463 definitions of  $Re_T$  and  $Fr_T$  are only valid in a weakly-stratified regime, defined by  $L_T < L_O$ .  
 464 According to these authors, the length scale dependency of  $Re_T$  and  $Fr_T$  differs for a highly-  
 465 stratified regime ( $L_T > L_O$ ), which would apply to a significant portion of our data. In such a  
 466 regime,

$$Re_T = \left( \frac{L_T^3}{L_O^2 L_K} \right)^{4/3} \quad (3)$$

467 and

$$Fr_T = \left( \frac{L_O}{L_T} \right)^2. \quad (4)$$

468 Here, we assess the impact that this regime shift has for our conclusions by re-calculating  $Re_T$   
 469 and  $Fr_T$  using equations 3-4. The results (Supplementary Figure 3) show that the diagram is  
 470 "stretched" relative to Figure 3b, owing to the stronger power dependencies of the scaling func-

tions in the highly-stratified regime. However, the key patterns supporting our findings remain unchanged.

Specifically, biophysical turbulence (green contour) spans a wide range of  $Fr_T$  values, with some data points intruding into an area with  $Fr_T > 1$  and very low mixing efficiency, but with many others located closer to  $Fr_T = 1$  (and even  $Fr_T < 1$ ) where mixing efficiency is higher. In contrast, interior geophysical turbulence (red contours) appears in an area with  $Fr_T < 1$  and possible turbulence suppression by buoyancy forces (around the oblique line representing  $Re_b \approx 10$ , where buoyancy forces suppress mixing). Geophysical turbulence in the wind-influenced surface layer displays data points in an energetic region of higher efficiency. In summary, although the shape of the diagram is slightly different, our key conclusions –i.e. that biomixing is occasionally suppressed because  $Fr_T > 1$  (particularly during I01), but not as stronger stratification reduces  $Fr_T$  during I02-I03; and that geophysical turbulence in the interior is partly suppressed by buoyancy, thereby making the efficiency of biophysical and geophysical mixing comparable– hold irrespectively of the approach chosen to estimate the turbulence parameters.

**Data availability.** The data that support the findings of this study are available at a Zenodo repository, doi:10.5281/zenodo.5559023.

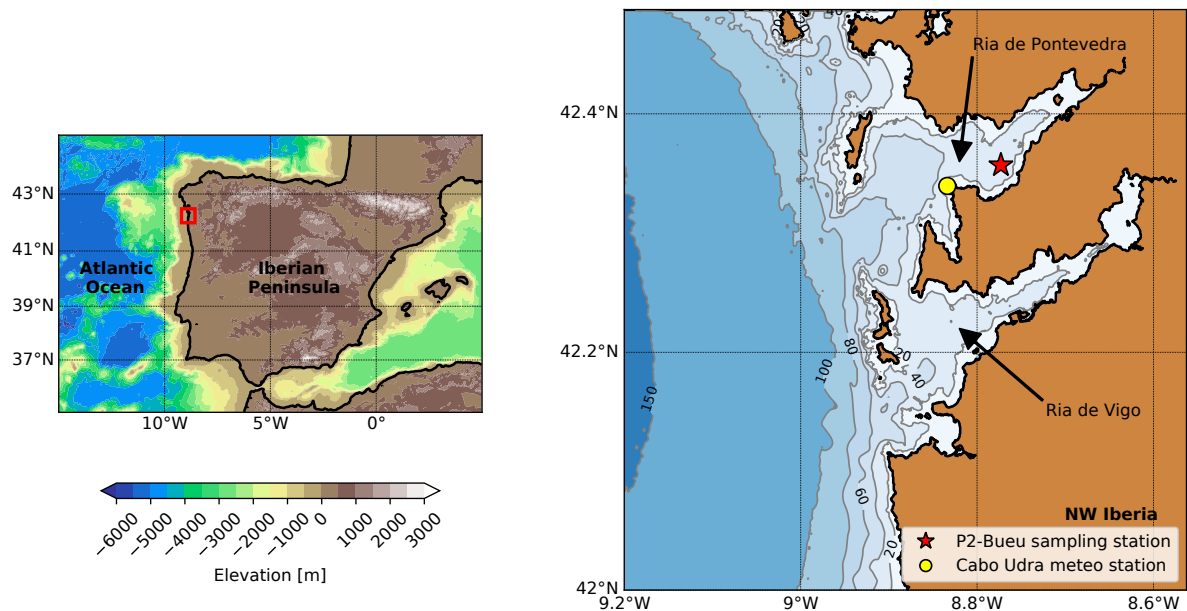
**Code availability.** The scripts used for microstructure data processing are freely available at [https://github.com/bieitofernandez/MSS\\_processing](https://github.com/bieitofernandez/MSS_processing)

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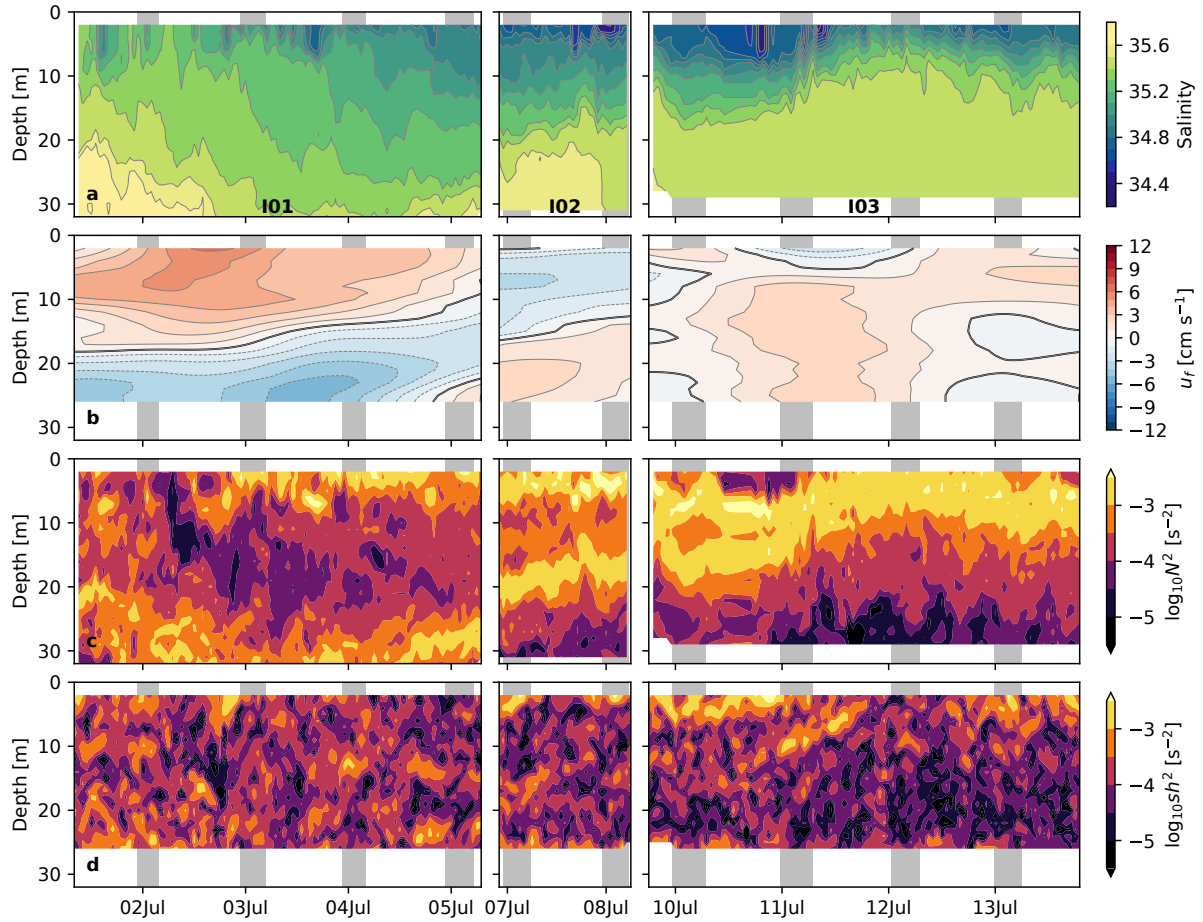
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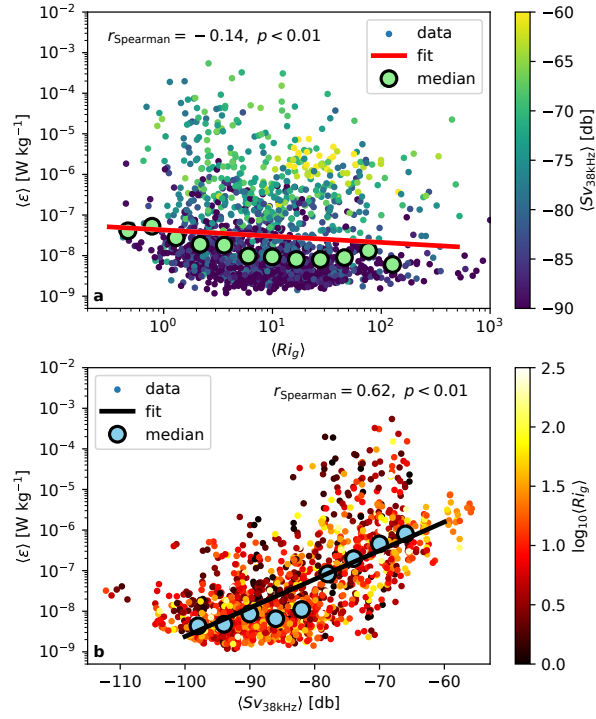
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**Extended Data Figure 1: Location of survey.** Map of the location of the REMEDIOS sampling station P2-Bueu (red star, 42.357°N, -8.773°W, mean depth 30 m) in the Ría de Pontevedra (off the Galician coast, NW Iberian Peninsula). The location of the closest Meteogalicia ([www.meteogalicia.gal](http://www.meteogalicia.gal)) meteorological station (Cape Udra, 42.340°N, -8.884°E) is also shown.

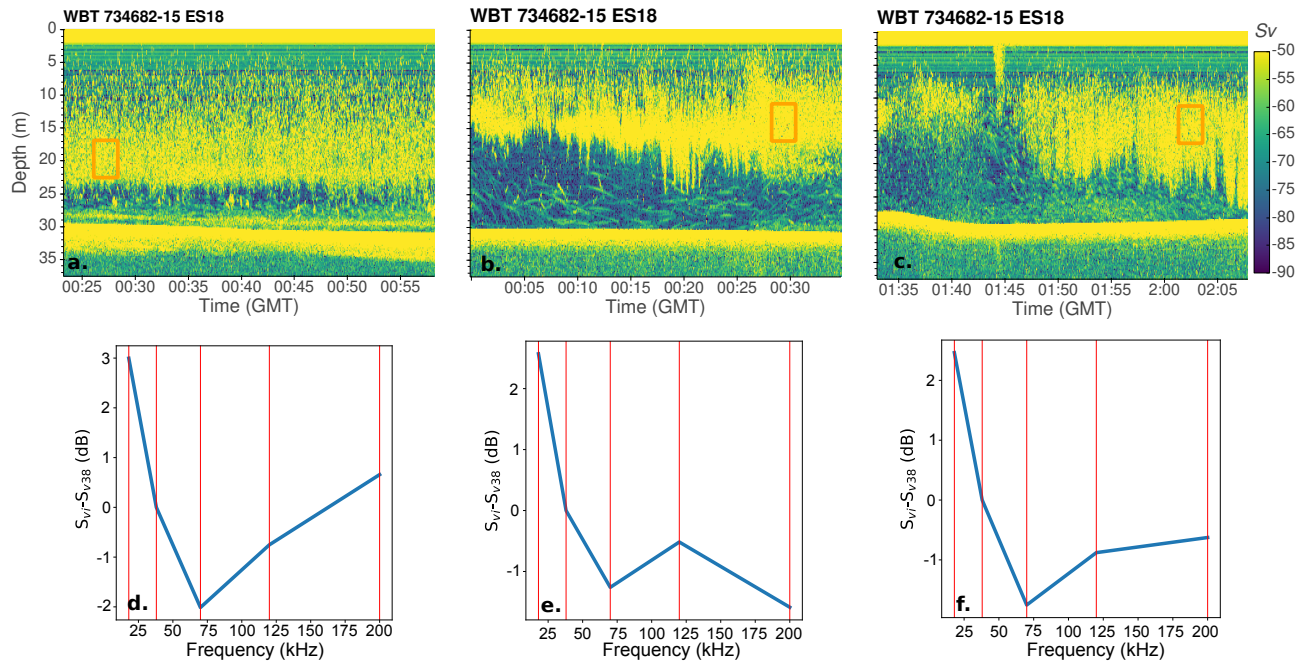


**Extended Data Figure 2: Hydrographic setting.** Hourly mean time series of **a** salinity, **b** de-tided eastward velocity ( $u$ ), **c** squared buoyancy frequency ( $N^2$ ), and **d** squared vertical shear of horizontal velocity ( $sh^2$ ) during the three sampling periods (I01, I02 and I03). Gray shading indicates night-time biomixing events. These periods were determined by inspection of the turbulent dissipation rate and volume backscattering strength records. De-tided residual velocity was calculated with a 24/25/24 h Godin filter. Positive eastward velocity imports offshore waters into the Ría, and negative westward velocity exports onshore waters out of the Ría. Note the use of logarithmic scale in panels **c** and **d**.

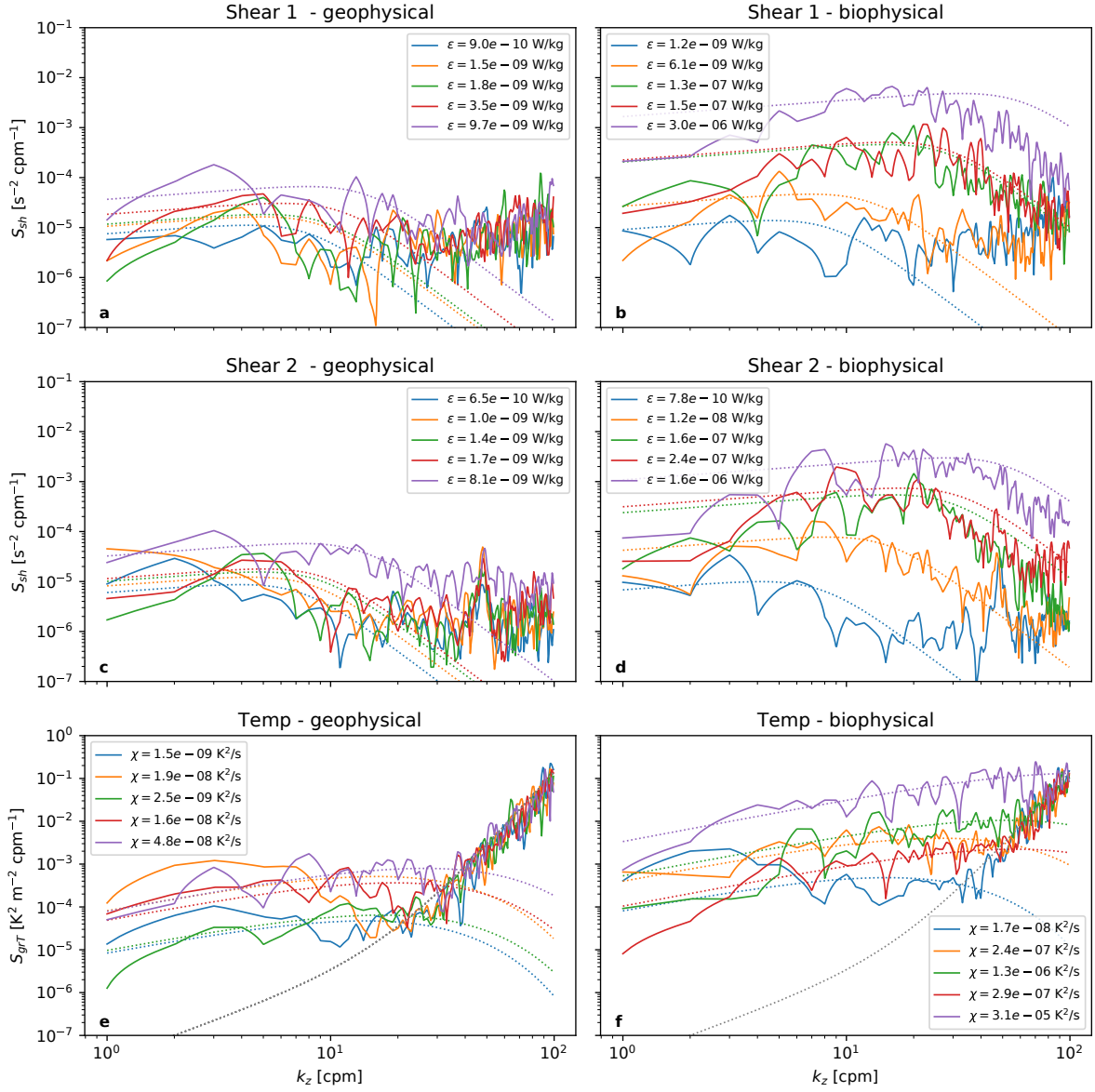


**Extended Data Figure 3: Sources of turbulence.** Depth-averaged (10–25 m)  $\varepsilon$  vs. **a** depth-averaged  $Ri_g$  and **b** 38 KHz volume backscattering strength ( $Sv$ ).  $\varepsilon$  median values in bins of  $Ri_g$  and  $Sv_{38\text{kHz}}$  are indicated as large circles. Linear fits in logarithmic scale and Spearman correlation coefficients are shown. The color scale represents  $Sv_{38\text{kHz}}$  and  $Ri_g$  in panels **a** and **b**, respectively.

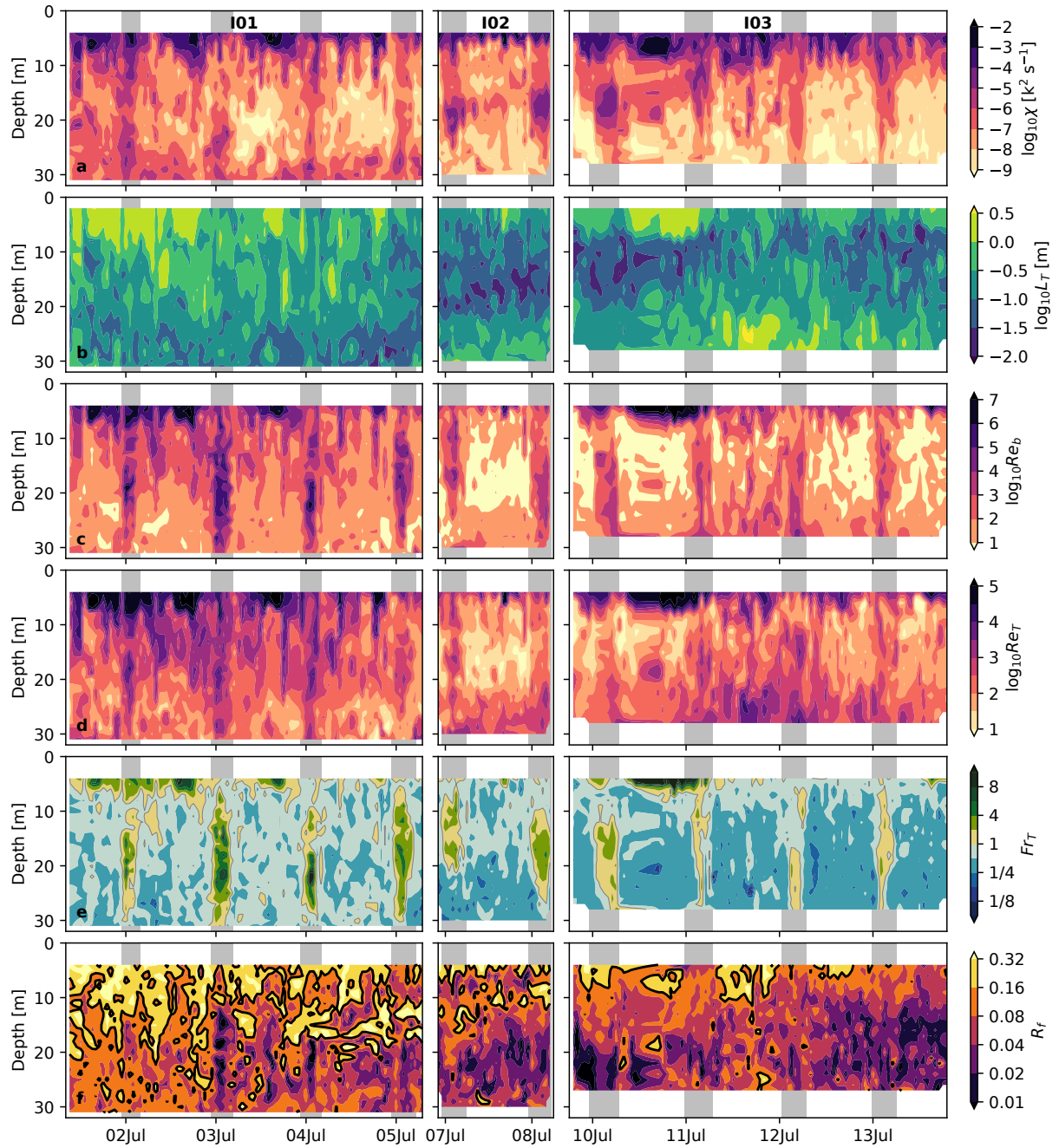




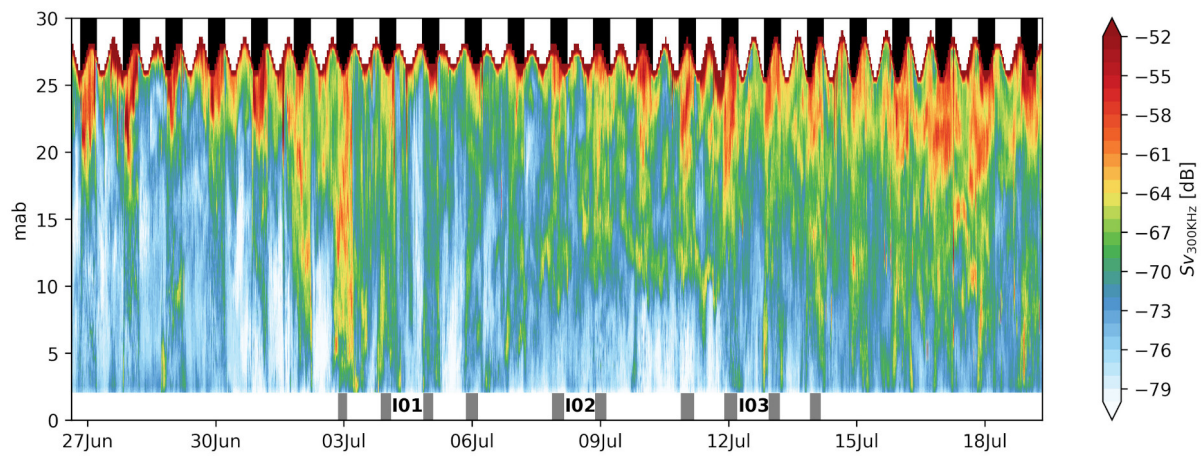
**Extended Data Figure 4: Acoustic backscatter frequency response.** Three examples of night-time echograms at 18 KHz, recorded during sampling periods I01 (a, 4 July), I02 (b, 8 July) and I03 (c, 12 July). Panels d-f show the mean frequency response ( $S_v$  at each frequency minus  $S_v$  at 38 kHz) for the region enclosed by the orange squares in panels a-c.



**Extended Data Figure 5: Microstructure spectra.** Randomly selected wavenumber ( $k_z$ , units: cycles per meter, cpm) spectra of vertical shear (a-d) and temperature gradient (e, f) microstructure between 10 and 25 m depth, during the third sampling period (I03). Periods dominated by geophysical turbulence are shown in the left column, and those dominated by biophysical turbulence (gray shading in Figure 1), in the right column. The corresponding universal spectra are indicated by dotted coloured lines, and the computed dissipation rates of turbulent kinetic energy ( $\epsilon$ ) and thermal variance ( $\chi$ ) are reported. Spectra recorded with the two shear sensors over the same portion of the water column are shown a, b and c, d, respectively. Empirical spectra of thermistor noise are represented by the gray dotted line e, f.



**Extended Data Figure 6: Turbulence and mixing parameters.** Time series of hourly mean **a** rate of dissipation of thermal variance ( $\chi$ ), **b** Thorpe scale ( $L_T$ ), **c** buoyancy Reynolds number ( $Re_b$ ), **d** turbulent Reynolds number ( $Re_T$ ), **e** turbulent Froude number ( $Fr_T$ ), and **f** flux Richardson number ( $R_f$ , a proxy for mixing efficiency) during the three sampling periods (I01, I02 and I03). Gray shading indicates night-time biomixing. Note the use of a logarithmic scale in all panels.



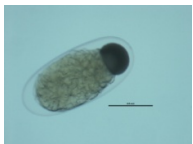





**Extended Data Figure 7: ADCP backscatter.** Time series of volume backscattering strength ( $S_v$ , dB) measured with a 300 KHz bottom-moored ADCP. Nights and biomixing events during the sampling periods (I01, I02 and I03) are indicated with black and gray shading, respectively.

**Extended Data Table 1: Mean turbulent properties.** Mean values [95% confidence intervals in brackets], and/or median values ( $^\dagger$ ) for the three sampling periods (I01, I02 and I03) and for times of biophysical and geophysical turbulence. Unless indicated, the averaging was performed over the 10–25 m depth range. The number of segments in which turbulent quantities were calculated, and were unaffected ( $n_{\text{good}}$ ) or affected ( $n_{\text{impacts}}$ ) by impacts against the instrument, are indicated. The affected data were discarded and not used for computing averages.  $N^2$  is the buoyancy frequency (a measure of stratification);  $sh^2$ , the vertical shear of the horizontal velocity;  $Ri_g = N^2/sh^2$ , the gradient Richardson number;  $\varepsilon$  the turbulent kinetic energy dissipation rate;  $\chi$ , the thermal variance dissipation rate;  $K_T$ , the turbulent heat diffusivity;  $R_f$  the flux Richardson number or mixing efficiency;  $L_T$ , the Thorpe scale;  $L_O$  the Ozmidov scale;  $Fr_T$ , the turbulent Froude number;  $Re_T$  the turbulent Reynolds number; and  $Re_b$  the buoyancy Reynolds number.

|  | I01   |  | I02  |  | I03  |  |
|--|---|--|--|--|--|--|
|  | Geophysical   | Biophysical  | Geophysical  | Biophysical  | Geophysical  | Biophysical  |
| $n_{\text{good}} (n_{\text{impacts}})$ | 8517 (0)  | 1772 (1057)  | 2573 (0)   | 1737 (129)   | 7531 (0)   | 3264 (114)   |
| $N^2$ [ $\text{s}^{-2}$ ]              | 2.15 [2.10 – 2.21] $\times 10^{-4}$                                     | 2.00 [1.93 – 2.1] $\times 10^{-4}$                           | 7.73 [7.44 – 8.07] $\times 10^{-4}$                          | 7.79 [7.37 – 8.11] $\times 10^{-4}$                          | 4.59 [4.45 – 4.72] $\times 10^{-4}$                          | 5.11 [4.92 – 5.32] $\times 10^{-4}$                          |
| $sh^2$ [ $\text{s}^{-2}$ ]             | 2.22 [2.16 – 2.28] $\times 10^{-4}$                                     | 2.47 [2.38 – 2.59] $\times 10^{-4}$                          | 1.85 [1.47 – 1.67] $\times 10^{-4}$                          | 1.68 [1.58 – 1.78] $\times 10^{-4}$                          | 1.30 [1.24 – 1.38] $\times 10^{-4}$                          | 1.44 [1.38 – 1.54] $\times 10^{-4}$                          |
| $Ri_g$                                 | 1.15 $^\dagger$   | 1.03   | 6.43   | 5.60   | 4.3  | 3.7  |
| $\varepsilon$ [ $\text{W kg}^{-1}$ ]   | 1.22 [0.83 – 2.27] $\times 10^{-7}$<br>6.52 $\times 10^{-9}$ $^\dagger$ | 1.62 [1.14 – 2.59] $\times 10^{-5}$<br>8.80 $\times 10^{-8}$ | 1.71 [1.18 – 2.86] $\times 10^{-8}$<br>3.35 $\times 10^{-9}$ | 1.96 [1.50 – 2.43] $\times 10^{-6}$<br>7.73 $\times 10^{-8}$ | 2.17 [1.83 – 2.70] $\times 10^{-8}$<br>2.18 $\times 10^{-9}$ | 0.82 [0.55 – 1.41] $\times 10^{-6}$<br>4.25 $\times 10^{-8}$ |
| $\chi$ [ $\text{K}^2 \text{s}^{-1}$ ]  | 1.19 [0.90 – 1.74] $\times 10^{-6}$<br>1.36 $\times 10^{-8}$ $^\dagger$ | 1.73 [1.30 – 2.41] $\times 10^{-6}$<br>8.64 $\times 10^{-8}$ | 4.38 [2.44 – 8.90] $\times 10^{-7}$<br>1.37 $\times 10^{-8}$ | 1.10 [0.94 – 1.42] $\times 10^{-5}$<br>3.41 $\times 10^{-7}$ | 5.40 [3.98 – 8.76] $\times 10^{-7}$<br>5.80 $\times 10^{-9}$ | 4.23 [3.69 – 5.09] $\times 10^{-6}$<br>1.50 $\times 10^{-7}$ |
| $K_T$ [ $\text{m}^2 \text{s}^{-1}$ ]   | 1.30 [0.63 – 2.76] $\times 10^{-3}$<br>4.85 $\times 10^{-6}$ $^\dagger$ | 1.30 [0.51 – 3.03] $\times 10^{-3}$<br>3.79 $\times 10^{-5}$ | 1.55 [0.63 – 3.99] $\times 10^{-5}$<br>2.61 $\times 10^{-7}$ | 1.58 [0.75 – 5.28] $\times 10^{-4}$<br>1.33 $\times 10^{-5}$ | 1.39 [1.12 – 1.92] $\times 10^{-5}$<br>3.08 $\times 10^{-7}$ | 3.52 [3.10 – 4.37] $\times 10^{-5}$<br>7.33 $\times 10^{-6}$ |
| $R_f$                                  | 0.137 [0.133 – 0.141]   | 0.092 [0.083 – 0.099]  | 0.068 [0.064 – 0.073]  | 0.081 [0.075 – 0.087]  | 0.053 [0.052 – 0.054]  | 0.052 [0.050 – 0.055]  |
| $L_T$ [m]                              | 0.342 [0.333 – 0.350]   | 0.235 [0.225 – 0.245]  | 0.099 [0.093 – 0.103]  | 0.115 [0.110 – 0.123]  | 0.283 [0.273 – 0.292]  | 0.222 [0.212 – 0.230]  |
| $L_O$ [m]                              | 0.195 [0.187 – 0.210]   | 1.35 [1.20 – 1.85]   | 0.028 [0.027 – 0.032]  | 0.23 [0.20 – 0.26]   | 0.058 [0.0558 – 0.062]                                       | 0.226 [0.214 – 0.240]  |
| $Fr_T$                                 | 0.63 [0.62 – 0.64]  | 2.63 [2.48 – 2.91]   | 0.52 [0.51 – 0.53]   | 1.76 [1.66 – 1.88]   | 0.40 [0.400 – 0.41]  | 1.19 [1.14 – 1.23]   |
| $Re_T$                                 | 1093 [1009 – 1205]  | 1716 [1482 – 2300]   | 85 [78 – 94]   | 321 [290 – 370]  | 343 [320 – 365]  | 548 [510 – 586]  |
| $Re_b$                                 | 1961 [1271 – 3462]  | 160189 [102140 – 271970]                                     | 57 [31 – 141]  | 4002 [2828 – 5677]   | 113 [95 – 149]   | 1699 [1454 – 2079]   |

**Extended Data Table 2: Anchovy development stages.** Development stages of the European anchovy (*Engraulis encrasicolus*) eggs considered in the present work (F1 to F6), equivalence with the stages proposed in ref. <sup>58</sup> (Stageing) (I to XI), elapsed time since spawning according to ref. <sup>48</sup> for a reference ambient temperature of 17°C, description of the development of the embryo, according to Table A1.3 from ref. <sup>41</sup>, and images of development stages (F1 to F6) (the bar inserted in the pictures is 0.5 mm length)

| Stage | Stageing | Elapsed time [hours] | Description  | Images  |
|-------|----------|----------------------|--|---|
| F1    | I        | 0                    | Cell division has not begun. The cytoplasm of the single cell appears as a clear hemisphere at one pole, although may be displaced to other locations. Unfertilized eggs are included in this stage. |    |
| F2    | II       | 4.3                  | Cell division starts. The blastodisc has a mulberry-like appearance. Blastula cells are very small, but it is possible to distinguish them.  |    |
| F3    | III      | 14.3                 | The eggs have appearance of tissue rather than a collection of individual cells. The segmentation cavity is visible. The blastodermal cap is $\leq 1/3$ of the yolk mass.                            |  |
| F4    | IV-VI    | 31.5                 | The blastodermal cap is $> 1/3$ of the yolk mass. There is a rapid differentiation at this stage. At the end of this stage the angle between the tail and the yolk is $\geq 90^\circ$                |  |
| F5    | VII-IX   | 48.3                 | The tip of the tail is free from the yolk. At the end of this stage, the curvature of the tail is evident and the gut is apparent along the ventral surface of the tail.                             |  |
| F6    | X-XI     | 59.3                 | The free portion of the tail is considered to extend from the body and not from the fin-fold. The tail length is $\geq 3/4$ of the yolk-sac. At the end of this stage hatching takes place.          |  |