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Spatial patterns and scale-dependent relationships between macrozooplankton and fish in the Bay of Biscay: an acoustic study

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ABSTRACT: Macrozooplankton plays a key role in pelagic ecosystems as a link between lower trophic levels and fish. However, although its ecological role is usually considered in polar ecosystems, it is rarely considered in temperate ones. To obtain comprehensive information on the macrozooplankton distribution in the Bay of Biscay we adapted a bi-frequency acoustic method developed for the Humboldt Current system. This method can be used to extract continuous and simultaneous high-resolution information on the spatiotemporal patterns of biomass distributions of macrozooplankton and pelagic fish throughout the diel cycle. The 2 distributions were mapped using geostatistical techniques. We applied kriging with external drifts, which accounts for both diel and across-shore changes in macrozooplankton biomass. We then used a cross-variogram to determine the scale-dependent relationships between macrozooplankton and fish. The results show how macrozooplankton and fish are distributed according to the different ecological domains (coast, shelf, shelf-break and offshore) along the Spanish and French coasts. Specific macrozooplankton hotspots were observed, but macrozooplankton was generally more abundant offshore than inshore, whereas fish showed the opposite trend. This pattern was confirmed by the aggregation sizes, which increased towards oceanic waters for macrozooplankton and decreased for fish. Finally, the correlation between fish and macrozooplankton was positive on a small scale (<30 nautical miles) and negative on a large scale (>30 nautical miles).

KEY WORDS: Macrozooplankton \cdot Pelagic fish \cdot Acoustics \cdot Spatial pattern \cdot Scale-dependent interaction \cdot Bay of Biscay

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INTRODUCTION

Zooplankton plays a crucial role in structuring higher and lower trophic levels, and influences the population dynamics of exploited species (Carlotti & Poggiale 2010). Although the ecological compartment that zooplankton represents is essential for a mechanistic understanding of the ecosystem dynamics, in general the data on zooplankton are scarce and of low quality (Mitra & Davis 2010), mainly because zooplankton abundance and distribution are generally assessed based on net sampling (Omori & Hamner 1982, Wiebe et al. 1985, Ohman & Smith 1995). Although net sampling provides valuable information on zooplankton species composition, abundance and time series (Ohman & Smith 1995, Verheye & Richardson 1998), net data are discrete in space and time, and are usually biased for macrozooplankton (>20 mm in size) (Mackas & Beaugrand 2010). Furthermore, macrozooplankton can react to both visual and mechanical disturbances and avoid net sampling, particularly vertical tows (Fleminger & Clutter 1965, Brinton 1967, Debby et al. 2004, Lawson et al. 2008). Therefore, despite the importance of macrozooplankton in biogeochemical cycles and the energy flow of marine pelagic ecosystems (Robinson et al. 2010), there are not enough data, particularly high-resolution data, to fully understand its ecological role (Siegel 2000). High-resolution zooplankton data are necessary for matching physical and biological processes and determining, for instance, mesoscale oceanographic structures (Gil et al. 2002).

As acoustics are not invasive, they can be used to sample organisms that otherwise would be missed or underestimated by net sampling; for example, acoustics are routinely used to estimate krill biomass in the Antarctic (e.g. Watkins & Brierley 2002, Lawson et al. 2008). This method allows high-resolution qualitative and quantitative data of various communities of an ecosystem to be collected simultaneously, from zooplankton to large predators (Bertrand et al. 2003, Koslow 2009).

In the Bay of Biscay, most of the study effort has been focused on fish, whereas little attention has been paid to zooplankton, particularly micro- and macrozooplankton (Beaudouin 1971, 1975, 1979, Quevedo & Anadón 2000). Actually, although large macrozooplankton aggregations are regularly observed with acoustics (Lezama-Ochoa et al. 2010), a broad picture of the macrozooplankton distribution has not yet been determined.

The objective of this study was to use a bifrequency acoustic method developed in the Humboldt Current system by Ballón et al. (2011) to gain a better understanding of the macrozooplankton community distribution and its role in the Bay of Biscay ecosystem. This method can be used to obtain continuous and simultaneous high-resolution information on macrozooplankton and pelagic fish biomass distributions.

We first described the horizontal distribution patterns of macrozooplankton and fish biomass according to different spatial scales and the diel period. Then, we used the simultaneous data on macrozooplankton and fish to determine the scale-dependent relationships between them.

MATERIALS AND METHODS

Acoustic data acquisition

Acoustic data were recorded with Simrad EY60 split-beam scientific echosounders at 38 and 120 kHz (Kongsberg Simrad AS) during a scientific routine survey performed in spring (April–May) 2009 within the BIOMAN program (AZTI project) in the Bay of Biscay. BIOMAN surveys estimate the anchovy *Engraulis encrasicolus* spawning biomass using the daily egg production method. They provide a large number of acoustic measurements, mesozooplankton (0.2–20 mm in size) samples, physicochemical measurements and pelagic trawl hauls (Motos et al. 1996).

The sampling area covered the Bay of Biscay (the Cantabrian Sea and off the French coast), with the westbound survey limit at 5°W and the northbound survey limit at 47° N (Fig. 1). Sampling was performed during both the day and night. The survey design was a combination of systematic and adaptive schemes. The systematic scheme was based on cross-shelf transect lines from the coast (bottom depth ~20 m) to beyond the shelf-break. Transects were parallel, regularly spaced and perpendicular to the coast with an inter-transect distance of 15 nautical miles (n mile). Standard transects extended 6 to 10 n mile off the shelf-break if no anchovy eggs were found further from the shelf-break. Otherwise, transects were prolonged as long as eggs were detected and then stopped when no eggs had been found in 6 n mile. This adaptive scheme was adopted to ensure that the anchovy spawning area was covered completely.

The echosounder was calibrated in accordance with Foote et al. (1987). The water column was sampled down to depths of 300 and 500 m for the 120 and 38 kHz channels, respectively. As there was electric noise in the echograms (surveys were performed onboard commercial vessels), only the first 150 m were considered. The acoustic data were selected, classified and analyzed with Echoview (SonarData) and MATLAB (MathWorks) software.

Bi-frequency classification method

Acoustic echoes were discriminated with a bifrequency acoustic method developed by Ballón et al. (2011). This method uses the 38 and 120 kHz frequencies to extract continuous and simultaneous high-resolution information on the spatiotemporal patterns of pelagic fish and 'fluid-like' organisms



Fig. 1. (a) Bathymetry map (m) of the study area. River mouths and specific topographic features are indicated, as well as shelf areas (coasts). From light grey to dark grey, polygons represent the coast, shelf, shelf-break and offshore areas. (b) Location of the ecological zones identified in the BIOMAN campaigns

(Stanton et al. 1996), mainly euphausiids and large copepods (Fig. 2).

Except for a few modifications, the method was applied directly in the Bay of Biscay.

Pre-processing: removing noise and resampling

First, the ping number and position between echograms were synchronized using the matching ping number algorithm from Echoview. The noise from the surface (5 m depth from the transducer) and any bottom echoes were excluded. The echograms were cleaned by defining and eliminating regions containing parasite noise (unwanted signals present in the medium but independent of the echosounder transmission; Urick 1986) or a 'school tail' (diffuse ragged tail below the more solid mark of the school). The noise associated with the acoustic absorption was eliminated (Korneliussen 2000, Fernandes et al. 2006) using the following noise function:

$$20\log(R) + 2\alpha R + O \tag{1}$$

where *R* is the range (m), α is the frequency absorption coefficient (dB m⁻¹), and *O* is the offset value (dB), i.e. the assumed initial noise in the first meter.

Acoustic scattering is stochastic, and thus it is necessary to average acoustic measurements to reduce natural variations in the data (Korneliussen et al. 2008). Following the recommendations of Korneliussen et al. (2008), the bi-frequency echograms were resampled in common elementary cells with a length of 1 ping and a height of 0.80 m (from 4 raw cells 0.2 m in height). Ballón et al. (2011) used the oxycline as a natural barrier to define the vertical extension of the epipelagic community. In the Bay of Biscay, there is no physical parameter that limits the epipelagic habitat as oxygen does in the Humboldt Current system; thus, we considered the entire observed water column (150 m or the bottom depth).

Enhancing the contrast between fish and zooplankton backscatters

The difference between the mean volume backscattering strength of 2 frequencies (Δ MVBS) is commonly used to determine the characteristics of biological backscatterings (see Murase et al. 2009). Unfortunately, the discriminatory power of scatters is limited when only Δ MVBS₁₂₀₋₃₈ is applied. For instance, pneumatophores may have an increasing or decreasing trend between the frequencies of 38 and 120 kHz depending on their size, and thus can have a positive or negative Δ MVBS. The same observation can be made if the 38 to 120 kHz range for a scatter falls within the Rayleigh/geometric transition zone (Stanton et al. 1998), where backscatter efficiency oscillates.

However, a recent methodology (A. Lebourges-Dhaussy & P. G. Fernandes unpubl. data), based on the sum of the mean volume backscattering strength (+MVBS) of 2 or more frequencies, enhances the contrast between pelagic swimbladder-bearing fish and other organisms, making them easier to discriminate.



Fig. 2. Main steps of the bi-frequency classification method developed by Ballón (2010) and Ballón et al. (2011). (a) First, discriminate scatters into 'fish', 'fluid-like' and 'others' based on their scattering models. $S_V(x)$: volume scattering strength, where x is the broadcast frequency (kHz). Reproduced from Korneliussen & Ona (2003) with permission. (b) Second, refine the discrimination of these groups by using expanded echograms as a Boolean mask. (c) Finally, recompose the final echograms of each scatter following some conditional statements

Discriminating scatters

With the exception of mackerel *Scomber scombrus*, most of the pelagic fish present in the Bay of Biscay, in particular anchovy, sardine *Sardinops sagax* and horse mackerel *Trachurus trachurus*, have swimbladders. Therefore, any reference to 'fish' in this study pertains to swimbladder-bearing fish. Swimbladder-bearing fish have slightly higher backscatter at 38 than 120 kHz (Kloser et al. 2002), but there are a few cases of positive Δ MVBS₁₂₀₋₃₈ (up to ~+3 dB) in the fish data. We thus refined the fish data from the fish echograms by applying a second Boolean mask in order to keep only the targets for which

The method by Ballón et al. (2011) combines both Δ MVBS and +MVBS to discriminate and quantify the abundance of crustacean macrozooplankton. Therefore, based on observations (expert scrutinizing of the echograms) and exploratory analysis (distribution of volume scattering strength [S_v] frequencies), a threshold value of –138 dB for the sum echogram (+MVBS₁₂₀₊₃₈) was chosen and used as a Boolean mask (true for values above the threshold) to extract fish data (above –138 dB) from other scatters (below –138 dB) and create 'fish' and 'no fish' (still not free from weak fish scatters) echograms at each frequency (Fig. 2a). From this point onwards, all processes were applied to both frequencies equally.

 $\Delta MVBS_{120-38} < +3 \text{ dB}$ (Fig. 2a). Although this constraint (~+3 dB) may allow us to include mackerel in this group (Fernandes et al. 2006), we assume that in general any reference to fish in this study pertains to swimbladder-bearing fish.

Zooplanktonic organisms with weakly scattering material and acoustic properties similar to the medium are usually called 'fluid-like' zooplankton (Stanton et al. 1996). The fluid-like group includes euphausiids, copepods, salps, siphonophores (without gas inclusion) and other large crustacean zooplankton (e.g. squilla larvae, munidae and other decapod larvae). This group was extracted from the 'no fish' echograms by applying a Boolean mask to select the targets with a positive $\Delta MVBS_{120-38}$ (Fig. 2a). Targets with a negative $\Delta MVBS_{120-38}$ were classified as 'others' ('blue noise' in Ballón et al. 2011). This last group included all targets other than fluid-like zooplankton and swimbladder-bearing fish (mainly fish larvae and gelatinous and gas-bearing siphonophores).

Refining the discrimination

A preliminary examination of the fluid-like echograms revealed that some targets located inside or at the edge of fish schools were inaccurately selected as fluid-like organisms. In order to extract the remaining fish backscatters from the fluid-like echograms and include them in the final fish echograms, we employed the following 4-step procedure (Ballón et al. 2011) (Fig. 2b):

(1) A copy of the fish echograms was expanded by dilating the value of a cell (containing fish) to its immediate 8 'neighbour' cells (3×3 convolution matrix from Echoview);

(2) The expanded fish echograms were used to create a Boolean mask, with false values where fish were present;

(3) This mask was applied to the non-expanded echograms to create new no-fish echograms;

(4) The Boolean mask was applied again to select the targets with positive $\Delta MVBS_{120-38}$ to create new fluid-like echograms.

Although the new fluid-like echograms were almost free from fish echoes, some of the fluid-like targets were also eliminated. Therefore, an additional step was necessary to recompose the fluid-like scatter and obtain our final fluid-like echogram:

(5) Gaps in the final fluid-like echogram were filled by combining the new fluid-like echogram and its expanded version according to a logical bitmap ('select' operator from Echoview). This bitmap has true values for the cells where the original fluid-like echogram has a correspondence value in the new fluid-like echogram and false values otherwise. If there was a true value in the bitmap, then the corresponding data value from the new fluid-like echogram was used, otherwise the corresponding data from its expanded version were used (Fig. 2c).

Similarly, the same procedure was used to include the fish echoes extracted from the fluid-like echograms in the final fish echograms.

Thresholds were then applied on the final echograms of fluid-like organisms (38 kHz and 120 kHz), fish (38 kHz) and others (38 kHz) to eliminate the potential remaining high noise echoes. The upper threshold values were -21 dB for fish, -56 dB for others, and -53 dB for fluid-like organisms (120 kHz), whereas the lower threshold values were -100 dB for fish and others (Fig. 3 shows an example of the bi-frequency algorithm).

Finally, as acoustic ping-rates varied from the coast to offshore, the final echograms were averaged over 1 n mile elementary sampling distance units in order to obtain regularly spaced sample data.

Variables of interest

Acoustic biomass

The fluid-like group mainly includes euphausiids, copepods, salps and siphonophores (without gas inclusion). In the Bay of Biscay, salps are not common on the shelf but can appear on the slope and offshore (Huskin et al. 2003). Likewise, siphonophores without gas inclusion have a very low biomass (Maycas et al. 1999). Therefore, our fluid-like field was probably mainly composed of euphausiids, but also large copepods, because small fluid-like zooplankton (e.g. small copepods) cannot be properly detected and quantified at the 38 kHz frequency (Mitson et al. 1996).

The lack of *in situ* net data validation did not allow us to set the method according to a specific group (e.g. euphausiids) and model the data based on its acoustic properties; therefore, this study focused on the whole macrozooplankton community, using an acoustic biomass index (S_v in dB re. 1 m⁻¹ and s_v in m⁻¹).

The fish group corresponded to all the small pelagic fish, in particular anchovy, sardine and horse mackerel. Fish acoustic backscattering energy (S_v) was converted into an acoustic nautical area



Fig. 3. Example of an application of the bi-frequency acoustic algorithm in order to discriminate the scatter types of 'fish', 'fluid-like' (macrozooplankton) and 'others'

scattering coefficient (NASC; $m^2 n mile^{-2}$), a biomass index determined according to MacLennan et al. (2002) as:

NASC =
$$4\pi (1852)^2 s_v T$$
 (2)

where $s_v = 10^{(S_v/10)}$ and *T* is the integrated vertical distance of the elementary cells of 0.80 m.

'Acoustic size' macrozooplankton

According to Mitson et al. (1996) and Murase et al. (2009), biological backscattering size can be characterized using Δ MVBS. Therefore, the Δ MVBS₁₂₀₋₃₈ values from a discriminated group (e.g. the fluid-like group) can be directly related to size classes, assuming that large differences relate to small sizes, and small differences to large sizes.

Defining diel periods

Diel vertical migration is a common behaviour for zooplankton and nekton. Its effects can be perceived at almost all spatial scales (see Strommel diagram in Haury et al. 1978). The diel vertical migration of macrozooplankton affects the estimations of biomass, density and size, as some species may migrate beyond the range of the acoustic sample (150 m for our study). Thus, day and night data were processed independently, and data from the twilight periods ± 15 min were eliminated in order to focus on consistent light periods.

Exploratory analysis

For each diel period (day/night), the distribution patterns of macrozooplankton and fish were represented using box plots according to area (Spanish or French), ecological domain (coast: 18 to 100 m depth; shelf: 100 m depth to shelf-break [~200 m depth]; slope: shelf-break (~200 m depth) to 1000 m depth; and offshore: >1000 m depth) and distance to the shelf-break. Likewise, the patterns of the acoustic size of macrozooplankton were represented using box plots according to ecological domain. Differences between distribution patterns were tested using nested ANOVA with a further post hoc Tukey's test.

The first test shows whether there is a diel effect in each area, and the second test determines whether macrozooplankton and fish biomass significantly differ between domains. However, these statistical tests assumed that the data were distributed normally, which was not the case. Therefore, macrozooplankton and fish biomass were normalized by applying a square-root transformation for macrozooplankton and a logarithmic transformation for fish. The normality of the transformed data was checked using the graphical method of the normal Q-Q plot.

Geostatistical analysis

Geostatistics are recognized to be particularly suitable for describing spatial distributions of marine populations. In particular, they provide mono- and multivariate tools for capturing and modelling the spatial structure of a population or highlighting structural relations between 2 populations. Distribution maps can then be produced by interpolation using proper kriging techniques (Simard et al. 1992, Petitgas 1993, Rivoirard et al. 2000).

Structural analysis

Classic and robust variograms (log back-transformed, Guiblin et al. 1995, Rivoirard et al. 2000) were used to describe the spatial dependence of macrozooplankton and fish, respectively. However, several factors had to be considered: (1) the presence of 2 continental shelves (Spanish and French) with distinct orientations and widths, which meant that the data had to be processed separately because the structure could differ from one area to another; (2) the possible difference between day and night structures due to the diel vertical migration and the aggregation/disaggregation phenomenon; and (3) the shelfbreak as a physical barrier that could induce different spatial structures on both sides of the break.

Therefore, following the scheme introduced in the exploratory analysis, the structural effects of the domain (Spanish and French shelves, offshore and inshore areas, and the entire Bay of Biscay) and the anisotropy or external drifts (Chilès & Delfiner 1999, Rivoirard et al. 2000) were analyzed for macrozooplankton and fish distributions. In these analyses, 2 external drifts were considered: the diel effect (Rivoirard & Wieland 2001, Wieland & Rivoirard 2001) and the vertical acoustic sampling range (150 m or the bottom depth if shallower). Indeed, the organisms were more or less likely to be detected by the echosounder according to the time of day due to vertical migration and according to the volume sampled in the water column. Finally, a cross-variogram was used to study the spatial interaction between fish and macrozooplankton (Journel & Huijbregts 1978, Cressie 1993, Goovaerts 1997).

Kriging

Ordinary kriging was used to map the fish biomass distribution based on a variogram model inferred

using a log back-transformed formula (e.g. Rivoirard et al. 2000). Indeed, no significant relationships with any external drift, which would have improved the modelling, were found for the fish biomass.

In contrast, macrozooplankton data were modelled by using ordinary kriging with a day/night indicator and the vertical range as external drifts. The variogram used in kriging with external drift should represent the structured variability of the residuals of the variable, excluding the drift. Thus, raw data were changed for the residuals obtained from a multivariate linear model ($s_v = \beta_1$ vertical range + β_2 day/night indicator + β_3 + ϵ_i ; where β_i = regression coefficients, and ε = error term) when the experimental variogram was calculated (Rivoirard & Wieland 2001, Wieland & Rivoirard 2001). This allowed us to produce kriging maps for day and night, when the macrozooplankton is expected to have a stable vertical distribution (i.e. vertical migration has finished). The 2 kriged maps use data from both the day and night, but with a correction applied to the night data for the day map and vice versa.

RESULTS

Exploratory analysis

Diel variability in coverage and biomass

The sampling coverage was different in the day and night (Fig. 4). Night data covered a smaller area than day data because of the longer day periods in summer. Macrozooplankton biomass exhibited significant differences according to diel period (Table 1), with higher macrozooplankton biomass at night (Fig. 5a), whereas fish did not show any significant differences (Table 1, Fig. 5b).

Area effect

Macrozooplankton biomass did not vary significantly according to the area (Spain or France) in either the night or day (Table 1). In each area, the diel period had a highly significant effect on macrozooplankton biomass (Table 1) with a higher biomass at night in both areas (Fig. 5c). Fish biomass showed significant differences according to the area (Table 1), with higher biomass on the French shelf, especially during the day period (Fig. 5d). The diel period did not have a significant effect on fish biomass in either area (Table 1). Highly significant differences between



Fig. 4. Day-night distribution of acoustic (a) macrozooplankton (s_v ; m^{-1}) and (b) fish biomass (NASC; m^2 n mile⁻²)

Table 1. Nested ANOVA of acoustic macrozooplankton and fish biomass according to diel period, shelf areas (Spain or France)
and ecological domains (coast, shelf, slope and offshore). Asterisks indicate significant difference: *p < 0.05; **p < 0.01; ***p <
0.001: NS: not significant

	— Die	l period ——	A	Area ———	Domain		
	df	F	df	F	df	F	
Macrozooplankton							
Diel period	1, 1095	112.39***					
Diel period × Area	1, 1093	41.2686***	2, 1093	2.2843^{NS}			
Area × Diel period	2, 1093	56.7686***	1, 1093	1.1784^{NS}			
Diel period × Area × Domain	1, 1081	142.6820***	2, 1081	1.4556^{NS}	12, 1081	25.5165***	
Fish							
Diel period	1, 1095	2.7692^{NS}					
Diel period × Area	1, 1093	3.0029^{NS}	2, 1093	47.3473***			
Area × Diel period	2, 1093	19.381***	1, 1093	58.935***			
Diel period × Area × Domain	1, 1081	12.030***	2, 1081	57.334***	12, 1081	60.498***	

fish biomass and area were observed for both the day and night (Table 1, Fig. 5d).

Ecological domain effect

Macrozooplankton biomass varied significantly with the ecological domain during the day and night, except in the Spanish area during the day (Tables 1 & 2). Its biomass increased regardless of the diel period across the continental shelf and towards the offshore region (Fig. 5e,g). This illustrates the importance of the shelf-break as a cut-off point with higher biomasses offshore, in particular during the night. Macrozooplankton biomass increased offshore with the distance to the shelf-break, particularly in the French area (Fig. 6a,c). In the French shelf area, in both the day and night, the macrozooplankton biomass showed a relatively smooth distribution pattern across the ecological domains (Fig. 5g) with no significant differences from the shelf onwards (Table 2). In the Spanish shelf area, the macrozooplankton distribution was less regular. There was no pattern at all during the day, and a pronounced coastoffshore gradient (Fig. 5e) with significant differences between inshore and offshore regions at night (Table 2).

Fish biomass also varied significantly with the ecological domains for both diel periods and areas, but with larger differences for areas (Tables 1 & 2). The effect that the shelf-break had on the distribution of organisms was independent of the diel period (Fig. 6b,d). Fish biomass showed the opposite pattern to the one observed for macrozooplankton, that is, it increased towards the coast during both the night and day (Fig. 5f,h). The fish biomass distribution pat-



Fig. 5. Day and night box plots of macrozooplankton and fish biomass according to areas and ecological domains. Day (white boxes) and night (grey boxes) distributions are represented according to (a,b) the total area, (c,d) the coastal areas, and (e–h) the ecological domains. Ecological domains are coast (C), shelf (Sh), slope (Sl) and offshore (O). Dot: mean; bar: median; box: 25th-75th percentiles; whiskers: 5th-95th percentiles. Macrozooplankton acoustic biomass values are in s_v (m⁻¹). Corresponding values in S_v (dB re. 1 m⁻¹) are indicated for information purposes only on the right-hand y-axis

tern, unlike the macrozooplankton pattern, was constant throughout the diel period and areas. Significant differences between ecological domains were found in almost all cases (Table 2).

Acoustic sizes

Macrozooplankton biomass and acoustic sizes showed similar across-shore distribution trends

with smaller acoustic sizes (higher $\Delta MVBS_{120-38}$) towards the coast (Fig. 7), coinciding with lower biomass (Fig. 5c).

Geostatistical analysis

Structural analysis

The Bay of Biscay has 2 distinct shelves (Spanish and French) that are more or less orthogonal to each other. Therefore, in a preliminary investigation (not shown), we studied the anisotropy and treated data from the 2 shelves separately. For each shelf, 2 directions (north-south and eastwest, corresponding grossly to the along and across contour effects) were considered when the experimental variograms were calculated. No anisotropy in macrozooplankton and fish data was found at the short-distance (i.e. small-scale) lag for either shelf. Likewise, experimental variograms along and across transects (still in the north-south and east-west directions) did not show any anisotropy at the short-distance lag when the 2 shelves were compared. Only a drift for the long-distance lag was found for the macrozooplankton over the French shelf in the east-west direction, which corresponds to the increase in biomass from the coast to offshore. For these reasons we chose to model the Spanish and French data together.

Macrozooplankton showed a significant positive correlation with vertical range (related to east–west drift over the French shelf) and diel period ($R^2 = 0.22$, depth p < 2 × 10⁻¹⁶ and time p <

 2×10^{-16} ; Fig. 8). Thus, both variables were used as external drifts to improve the kriging of the macrozooplankton. The variogram model was inferred based on the residuals and assuming isotropy for every lag. Using the vertical range as external drift meant that an isotropic model could be used for the entire Bay while accounting for the previously observed long-distance lag anisotropy.

The variogram models were fitted to a maximum distance of 150 n mile, including both day and

Table 2. Post hoc Tukey's multicomparison test of acoustic macrozooplankton and fish biomass according to diel period, shelf areas and ecological domains. Asterisks indicate significant difference: *p < 0.05; **p < 0.01; ***p < 0.001; NS: not significant

	re 75)		r							
France	Offsho $(n = 7)$		* * *	SN	NS		* *	* * *	NS	
	Slope $(n = 76)$		* * *	NS			* * *	*		
	Shelf 1 = 29)		:				•			
	ast 1 109) (r									
ight —	Co (n =									
Z	ffshore 1 = 22)		**	*	NS		•	•	NS	
	ope O = 18) (r		*	•			*	•		
, iero	lf Sl I5 (n: 15 (n:									
	Shel (n = 1		NS				NS			
	Coast (n = 6)									
	shore = 33)		:	ZZ	NS			:	* * *	
	e Off 35) (N		•	Ч	Д		·	·	•	
	Slop (n = 16)		* * *	NS			* *	* * *		
Ц.	Shelf = 263)		:				\mathbf{NS}			
	ıst 234) (n									
Day –	Co (n = 1									
	fshore 1 = 22)		\mathbf{NS}	NS	NS		÷	:	NS	
	pe Of : 18) (r		IS	IS			•			
, iero	f Slc 6) (n =		Z	Z			*	*		
	Shel (n = 1	_	NS				NS			
	Coast (n = 9)	ankton								
		lqoozo.	t		e lore		t		0	lore
		Macr	Coas	Shelf	Slopé Offsh	Fish	Coas	Shelf	Slope	Offsl

night data. They consisted of a nugget, representing unresolved small-scale variation and observation error, and a structured part. The fish model had a 3-component structure: a nugget effect of 60 000 and 2 spherical models with sills of 30 000 and 98 000 and ranges of 5 and 50 n mile (Fig. 9b). The macrozooplankton model had 3 components: a nugget effect of 0.50×10^{-17} and 2 spherical models with sills of 2.80×10^{-17} and 2 spherical models with sills of 2.80×10^{-17} and 2.75×10^{-17} and ranges of 14 and 80 n mile (Fig. 9a).

Fish and macrozooplankton biomasses showed opposite inshore–offshore trends, as macrozooplankton increased towards oceanic waters and fish decreased. This also occurred for the spatial structures of these 2 populations. Experimental variograms were calculated and models were fitted (nested structures with a nugget, a short- and a long-range spherical component) for inshore and offshore data for each population (Fig. 10).

Offshore, both short- and long-range structures for macrozooplankton showed similar variability. Therefore, both scales of the spatial structure were equally represented in this region. However, inshore, the short-range structure showed larger variability than the long-range structure; thus, the small scale of the spatial structure was more important inshore.

Offshore, fish showed the same pattern as macrozooplankton, with similar variability for both structures. Inshore, the long-range structure showed larger variability aggregate.

Macrozooplankton and fish horizontal distribution

Interpolated maps of acoustic macrozooplankton and fish biomass confirmed the previous results but provided far more details on the spatial patterns (Fig. 11). In general, fish and macrozooplankton exhibited opposite spatial patterns as zooplankton biomass increased towards oceanic waters, whereas fish biomass decreased.

Zooplankton diel vertical migration was particularly noticeable during the night, in particular in the offshore area. Therefore, 2 separate maps, one for day and one for night data, were used to represent the macrozooplankton distribution.

Some hotspots were detected on both the fish and macrozooplankton maps. The shelf-break was a key feature, both for macrozooplankton and fish. For instance, high values of macrozooplankton biomass occurred at the extreme northeast of the map in the slope/offshore regions (~46–47° N), and at the Fer à cheval and Capbreton canyon locations (Fig. 11).



Distance to the shelf-break (n mile)

Fig. 6. Day and night box plots (limits as in Fig. 5) of macrozooplankton and fish biomass according to the distance to the shelfbreak. Patterns during the day (white boxes) and night (grey boxes) are shown for (a,b) Spain and (c,d) France. Acoustic macrozooplankton biomass values are in s_v (m⁻¹). Corresponding values in S_v (dB re. 1 m⁻¹) are indicated for information purposes only on the right-hand y-axis



Fig. 7. Day (white) and night (grey) box plots (limits as in Fig. 5) of difference in mean volume backscattering strength (Δ MVBS₁₂₀₋₃₈) (as an indicator of acoustic sizes) for the macrozooplankton in (a) Spain and (b) France in relation to the 4 ecological domains: coast (C), shelf (Sh), slope (Sl) and offshore (O)

Another noticeable macrozooplankton aggregation occurred in oceanic waters in the Cantabrian Sea between $2^{\circ}30'$ W and $3^{\circ}45'$ W.

The 2 main aggregations of fish biomass occurred in the coastal areas, north of the French shelf $(46-47^{\circ} \text{ N})$ and along the Spanish coast $(2-3^{\circ} \text{ W})$. A third aggregation was located between 45 and 46° N, within the 70–200 m bottom-depth zone.

Cross-scale correlation

A cross-variogram between fish and macrozooplankton was calculated to determine the cross-scale correlation between these 2 populations (Fig. 12). Fish and macrozoo-

plankton were positively correlated at a small scale (~2 to 30 n mile) and negatively correlated at a larger scale (>30 n mile).

Fig. 8. Drift from diel period and vertical range. Scatter plot between the square root of acoustic macrozooplankton biomass (s_v) and vertical range, with day values in grey, and night values in black. A linear model has been fitted and represented: $\sqrt{s_v} \approx$ indicator of day + vertical range



DISCUSSION

A bi-frequency classification method was applied to acoustic survey data collected during spring 2009 in the Bay of Biscay (Fig. 3). Although it was applied for only 1 yr, this method produced 4 major results: (1) the first high-resolution, largescale acoustic description of the macrozooplankton biomass distribution; (2) the comparison of macrozooplankton and fish biomass distribution patterns in relation to the diel cycle; (3) the spatial modelling of these distributions using geostatistics; and (4) evidence of a correlation between pelagic fish and macrozooplankton biomass, positive at a small scale (<30 n mile) and negative at a larger scale (>30 n mile).

Validation of the bi-frequency classification method

The main difficulty in validating the acoustic results is the availability of independent biological observations. Net sampling is the most common tool for sampling marine pelagic organisms. Unfortunately, the confidence levels in net biomass estimations are either high or low depending on the type of nets and the target species (Fleminger & Clutter 1965, McGowan & Fraundorf 1966, Gorbatenko & Dolganova 2006). The problem is particularly crucial for macrozooplankton, which are capable of avoiding nets (Debby et al. 2004). This and the lack of comprehensive information on macrozooplankton in the Bay of Biscay make it difficult to determine which groups compose the observed macrozooplankton. Therefore, we will discuss our results according to the well-known broad patterns of macrozooplankton distribution (e.g. Blackburn 1979, Barange & Pillar 1992, Swartzman et al. 2005, Ballón et al. 2011).

It was less difficult, however, to differentiate the pelagic swimbladder-bearing fish because the main research institutes in this area routinely study the fish biomass distribution with acoustic methods (from a single frequency) and use ground-truth catch data for validation (Massé 1996, Massé et al. 1996, Boyra

Fig. 9. Experimental variograms and corresponding models used for the kriging for (a) macrozooplankton (inferred from the residuals) and (b) fish (log-back transformed). Symbol area is proportional to the number of pairs involved in the calculation for each distance lag. Variograms are normalized to the sample variance for comparison purposes only





Fig. 10. Inshore–offshore structural analysis of macrozooplankton and fish. (a,d) Proportional representation of biomass; inshore values (0–150 m) are in grey whereas offshore values (>150 m bottom depth) are in black. Normalized experimental variograms of the (b,e) inshore and (c,f) offshore biomass and their corresponding models. Symbol area is proportional to the number of pairs involved in the calculation for each distance lag. Fish variograms are log back-transformed

& Uriarte 2006, G. Boyra et al. unpubl.). Thus, although the distribution and ecology of pelagic fish have been studied in the Bay of Biscay for over 2 decades, little is known about macrozooplankton. expected (e.g. Ballón et al. 2011) and could be due to the fact the acoustic vertical range covers most of the water column inshore and most of the migrating communities are distributed offshore.

Diel patterns of distribution

Macrozooplankton

As expected, the macrozooplankton biomass in the upper layer (from the surface to 150 m) was significantly higher during the night than during the day. The nocturnal macrozooplankton biomass is composed of migratory and non-migratory macrozooplankton (Fernández de Puelles et al. 1996, Poulet et al. 1996, Maycas et al. 1999). Migratory organisms were not included in the biomass estimated during the day, as they were mainly distributed at approximately 400 m, which is deeper than our acoustic range of 150 m.

Based on the difference between the day and night biomass, we estimated the migrating macrozoo-plankton biomass to be 19% offshore and 10% inshore. These inshore–offshore differences were

Fish

No significant day-night effect was observed in the fish vertical distribution. This is probably because the bulk of pelagic fish biomass was located within our acoustic detection range of 150 m (Massé 1996, Massé et al. 1996). The slightly higher level of fish biomass during the day could be due to unsampled biomasses at the surface (0-5 m) or a stochastic difference between day-night sampling coverage.

Horizontal distribution

When mapping, the spatial distributions of macrozooplankton and pelagic fish may be questionable. Indeed, macrozooplankton and pelagic fish are known to form aggregations/schools during the day that disperse during the night, and also to perform





Fig. 11. Horizontal biomass distribution during spring 2009. Acoustic macrozooplankton biomass (s_{vi} ; m⁻¹) during the (a) day and (b) night, and (c) acoustic fish biomass (NASC; m² n mile⁻²) during the day–night period

diel vertical migration. However, the scale at which these phenomena occur is mostly below our sampling lag (elementary sampling distance unit < 1 n mile). Thus, the modelled spatial structures (using day and night data) are acceptable and correspond to at least that of the clusters of macrozooplankton or fish aggregations.

Macrozooplankton

The highest biomasses of macrozooplankton were found in areas adjacent to the shelf-break: Capbreton and Cap Ferret canyons, the Fer à cheval region, the Spanish slope and the northwestern French shelf. The shelf-break thus constitutes an important feature for retaining and accumulating macrozooplankton offshore and over the slope in the Bay of Biscay. Similar observations have been made in other systems (e.g. Bakun 1996, Genin 2004, Zhu et al. 2009, Ballón et al. 2011). The acoustic sizes of macrozooplankton were also higher offshore (smaller $\Delta MVBS_{120-38}$). This suggests a shift in species composition towards organisms such as euphausiids or large copepods. The significant shelf-break effect on macrozooplankton biomass and acoustic size is similar to the effect that has been observed on the composition of mesozooplankton communities, which shift from small to large mesozooplankton (Sourisseau & Carlotti 2006, Albaina & Irigoien 2007, Irigoien et al. 2009).

Fish

Fish were distributed differently than macrozooplankton. High biomasses were found in the coastal regions, mostly over the French shelf area. The densest fish aggregations were observed inshore on the northern platform (46 and 47° N), in the inner shelf area in front of the Gironde River plume (45° 10' N), in



Fig. 12. Isotropic cross-variogram illustrating the cross-scale correlation between fish and macrozooplankton biomass. Dashed lines represent the positive, null and negative sample cross-variances, from top to bottom

the outer shelf area between 70 and 150 m depth, and along the Spanish coast (2–3° W). Other, less dense aggregations were found offshore on the southern platform related to the Adour River plume and shelf edge fronts (which defined the area between the Capbreton and Cap Ferret canyons, and Fer à cheval region). Similar patterns were observed during both annual spring surveys (PELGAS and BIOMAN) performed to estimate the biomass of small pelagic species in the Bay of Biscay (ICES 2009).

This distribution agrees with the habitat preferences of the main pelagic species (anchovy, sardine and horse mackerel) during spring. Small clupeiformes, especially young anchovies and sardines, prefer coastal habitats and river mouths (Motos et al. 1996, Planque et al. 2007), which has been related to the favourable conditions (increased biological production) produced by the high river runoff (Gironde, Loire and Adour; Fig. 1), by nutrient inputs from deeper waters (coastal upwellings) and by low vertical mixing (haline and thermal stratification, Lazure & Jegou 1998, Puillat et al. 2004). This makes the French inner shelf a more suitable area for these fish during the spawning season than the Spanish shelf.

However, horse mackerel and older anchovies (≥ 2 yr old) show more affinity for outer-shelf regions, and occasionally approach the shelf-break (Abaunza et al. 2003), which, according to Motos et al. (1996), occurs in the Fer à cheval region, along the French shelf near Capbreton canyon and in the Cantabrian Sea, west of 2°W.

Fish-macrozooplankton interactions

Aggregation scale

During recent decades, numerous studies have focused on characterizing the spatial patterns of the pelagic fish in the Bay of Biscay (e.g. Petitgas 1998, 2000, Petitgas et al. 2003, Woillez 2007). Nonetheless, little is known about the spatial patterns of zooplankton (López-Jamar et al. 1991), and nothing about those of macrozooplankton. The present study revealed that the macrozooplankton biomass was structured spatially at 2 different ranges: a short one at 14 n mile and a longer one at 80 n mile. The fish spatial structure also showed 2 different ranges: a small one at 5 n mile and a larger one at 50 n mile. Previous studies have also observed a 2-scale spatial structure: 4 and 19 n mile for pelagic fish in Petitgas (2000), and 10 and 75 n mile for anchovy in Woillez (2007). These differences illustrate the annual variability of the aggregation patterns related to changes in the specific composition of the main communities and to the oceanographic forcing (Bertrand et al. 2008).

It is interesting to note that fish and macrozooplankton aggregation scales were similar offshore (5-4 n mile for the small-medium scale and 40-35 n mile for the large scale). Bertrand et al. (2008) described similar associations between fish and zooplankton aggregation scales in the Humboldt Current system, and suggested that mesoscale (10s of km) physical forces (e.g. upwelling/downwelling, tidal fronts and river discharges) shape the distribution of zooplankton patches given that there is motivation for feeding, which indirectly influences the fish patch size. However, we would also expect to obtain the same aggregation scales between macrozooplankton and fish inshore, where most of the fish biomass occurs, but this was not the case. Young anchovies and sardines mainly forage on small-sized prey, e.g. micro- and meso-zooplankton (ca. small-size copepods) that dominate inshore zooplankton (Irigoien et al. 2009), which are not well represented in our data (zooplankton $>\sim 2$ mm). However, older anchovies (≥ 2 yr old) and horse mackerel, which are usually located at the shelf-break (Motos et al. 1996), forage on larger prey (e.g. Blaxter & Hunter 1982, Van der Lingen et al. 2009) belonging to mesozooplankton (ca. mediumsize copepods) and macrozooplankton (ca. large copepods and euphausiids). This may explain why we found the same aggregation scales between fish and macrozooplankton in the offshore region where the fish prey field is accurately observed, and not inshore where the fish prey field is not or partially observed.

Scale-dependent predator-prey relationships

The cross-variograms showed that macrozooplankton and fish biomass were positively correlated at small scales (<30 n mile) and negatively correlated at larger scales (>30 n mile). These results appear to contradict the classic theory on predator-prey spatial interactions (Rose & Leggett 1990) in which negative correlations occur at small scales and positive ones at larger scales. In our case, the negative relationship observed on a large scale illustrates that fish and macrozooplankton are distributed in different regions (inshore and offshore). The positive relationship between macrozooplankton and fish biomass at the small scale indicates that, locally, fish concentrate at macrozooplankton patches. Genin (2004) observed similar positive local relationships along topographically generated fronts, for instance seamounts, canyons and shelf-breaks. Zooplankton aggregations have a certain degree of permanence above these features, and attract planktivorous fish species (Sameoto 1976, Simmard et al. 1986, Bertrand et al. 2008). However, our data do not address the real small scale sensu Rose & Leggett (1990). The smallest scale considered was 1 n mile (cross-variogram lag), which is probably appropriate for determining that fish are attracted to macrozooplankton hotspots; however, it does not allow us to study the very small scale where fish and macrozooplankton interact, and where depletion (negative relation) can be observed.

CONCLUSIONS

Simultaneous high-resolution data on both macrozooplankton and pelagic fish were used to provide a better understanding of the spatial processes and interactions in the Bay of Biscay pelagic ecosystem. Macrozooplankton and fish showed opposite largescale distribution patterns as macrozooplankton biomass was higher offshore than inshore and the reverse was true for fish. The shelf-break is a key feature for macrozooplankton because it is both a hotspot and the limit between low and high abundance. The aggregation scales followed a similar trend because larger aggregations predominated offshore for macrozooplankton and inshore for fish. These opposed large-scale distribution patterns lead to a negative correlation between the fish and macrozooplankton distributions at scales >30 n mile. However, the correlation was positive at scales <30 n mile, which indicates that fish concentrate where macrozooplankton is more abundant. Further

analyses that include environmental factors need to be performed in order to better understand the mechanism responsible for the observed patterns. The availability of ground-truth data of suspected macrozooplankton echosigns should allow us to transform an acoustic biomass into an absolute biomass, which will be another step towards understanding trophic interactions better and, more generally, extending ecosystem management by including non-exploited organisms.

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