Fine-scale vertical structure of sound-scattering layers over an east border upwelling system and its relationship to pelagic habitat characteristics

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Abstract. Understanding the relationship between sound-scattering layers (SSLs) and pelagic habitat characteristics is a substantial step to apprehend ecosystem dynamics. SSLs are detected on echo sounders representing aggregated marine pelagic organisms. In this study, SSL characteristics of zooplankton and micronekton were identified during an upwelling event in two contrasting areas of the Senegalese continental shelf. Here a cold upwelling-influenced inshore area was sharply separated by a strong thermal boundary from a deeper, warmer, stratified offshore area. Mean SSL thickness and SSL vertical depth increased with the shelf depth. The thickest and deepest SSLs were observed in the offshore part of the shelf. Hence, zooplankton and micronekton seem to occur more frequently in stratified water conditions rather than in fresh upwelled water. Diel vertical and horizontal migrations of SSLs were observed in the study area. Diel period and physicochemical water characteristics influenced SSL depth and SSL thickness. Although chlorophyll-α concentration insignificantly affected SSL characteristics, the peak of chlorophyll a was always located above or in the middle of the SSLs, regularly matching with the peak of SSL biomass. Such observations indicate trophic relationships, suggesting SSLs to be mainly composed of phytoplanktivorous zooplankton and micronekton. Despite local hypoxia, below 30 m depth, distribution patterns of SSLs indicate no vertical migration boundary. The results increase the understanding of the spatial organization of mid-trophic species and migration patterns of zooplankton and micronekton, and they will also improve dispersal models for organisms in upwelling regions.

1 Introduction

Aggregations of marine pelagic organisms in ocean water can be observed acoustically as sound-scattering layers (SSLs) (Evans and Hopkins, 1981; Cascão et al., 2017). The SSLs represent a concentrated layer of marine organisms such as zooplankton and micronekton that occur at specific depths (Benoit-Bird and Au, 2004). Nevertheless, the SSL is not a biological classification, and animals making up SSLs include various species, with correspondingly different biological, physiological, and ecological needs. The SSLs are dynamic, active, and have a particular behavior as a function of their community structure, causing changes in their vertical distribution, size, and shape over time and space (Gómez-Gutiérrez et al., 1999). Zooplanktonic and micronektonic components are fundamental to ecosystem functioning, par-
Zooplanktonic and micronektonic species provide the main trophic link between primary producers and higher trophic levels. A large amount of energy passes through zooplankton and micronekton (Steele et al., 2007). Knowledge of the vertical structure of SSLs allows us to understand their role in ecosystems, information that can be used to monitor major environmental change and variability. Most zooplankton and micronektonic taxa undergo diel vertical migration (DVM), meaning that they reside in deep waters during the day and migrate toward the surface at night to feed (Bianchi et al., 2013; Lehodey et al., 2015). DVM behaviors are influenced by environmental cues (e.g., light, nutrients, and temperature) and predator–prey interactions (Clark and Levy, 1988; Lampert, 1989). Thus, DVMs represent an essential biological process in the ocean, one that also regulates the biological carbon pump (Hidaka et al., 2001). Zooplankton and micronekton are also known to undergo diel horizontal migration (DHM), moving them to within 1 km of the shoreline each night into shallower waters (Benoit-Bird et al., 2001). DHM, like DVM, which often occur concurrently, helps organisms to find food and avoid predators (White, 1998).

The distribution of SSLs is influenced by a variety of environmental factors (Aoki and Inagaki, 1992; Baussant et al., 1992; Dekshenieks et al., 2001; Marchal et al., 1993). Changes in the structure and density of SSLs is associated with frontal zones (Aoki and Inagaki, 1992; Baussant et al., 1992; Boersch-Supan et al., 2017; Coyle and Cooney, 1993). Oceanic fronts are relatively narrow zones of enhanced horizontal gradients of physical, chemical, and biological properties (temperature, salinity, nutrients, plankton communities, etc.) that separate broader areas of different vertical structure (stratification) (Belkin et al., 2009). Upwelling fronts occur in many well-studied systems, including the upwelling off southern Senegal, south of Cap-Vert peninsula known as the “Petite Côte” (14.6°–13.5° N, 16.9°–17.6° W). Senegalese coasts are characterized by a seasonal upwelling (in winter and late spring), mainly driven by wind variability, topography, and density stratification (Estrade et al., 2008). During the upwelling season, northerly trade winds induce a strong upwelling core south of Dakar (Ndoye et al., 2014; Roy, 1998). The upwelling core is located over the shelf, and SST (sea surface temperature) is lowest on the coastal side of the shelf break, increasing in both offshore and coastal directions. Local bottom relief combined with the wind-induced upwelling establishes a typical upwelling that appears as a cold-water tongue. This cold-water tongue separates the nutrient-poor warm offshore cell with a cold nutrient-rich coastal cell functioning as a retention zone (Roy, 1998; Tiedemann and Brehmer, 2017). The Petite Côte in the Senegalese coastal shelf is a nursery area for fish and is the main area in which juveniles of numerous species, particularly small pelagic species, concentrate (Diankha et al., 2018; Thiaw et al., 2017). This area is also known to be rich in zooplankton and micronekton. Many zooplankton groups that are encountered over the Senegalese coastal shelf: Copepods, amphipods, annelids, appendicularians, chaetognaths, cirripeds, cladocerans, Decapoda, echinoderms, euphausiids, gastropods, jellyfish, Myidacea, ostracods, pelagic foraminifera, Protozoa, pteropods, and Spumellaria. Copepod is the most dominant group with a total abundance ranging from 50% to 90% (Anonymous, 2013; Ndour et al., 2018; Touré, 1971). Previous studies (Ndour et al., 2018; Tiedemann and Brehmer, 2017) on ichthyoplankton showed that Sparidae (∼50%) was predominant, followed by fewer Engraulidae (∼8%) and Soleidae (∼7%), while smaller proportions of Clupeidae and Carangidae (∼4% each) as well as Myctophidae and Sciaenidae (∼2% each) were found. Physical variability in the Senegalese coastal shelf (Capet et al., 2016; Ndoye et al., 2017) can impact marine pelagic organisms at the individual and community level (Urmy and Horne, 2016). Such an impact can be direct via advection or indirect via phytoplankton production fertilized by upwelled nutrients. Indeed, changes in physicochemical water properties and biological activities induced by upwelling plays a structuring role on the distribution of SSLs. SSL position is often reported below the thermocline, suggesting that temperature controls the SSL’s vertical distribution (Aoki and Inagaki, 1992; Baussant et al., 1992; Boersch-Supan et al., 2017; Marchal et al., 1993). Bottom depth has been identified as an additional factor structuring the vertical distribution of SSLs (Gausset and Turrel, 2001). For example, the thickness and depth of an SSL on continental shelves tend to increase with an increase in water depth (Torgersen et al., 1997), similar to patterns observed in the deep sea (Berge et al., 2014; Boersch-Supan et al., 2017). In deep-sea areas and over shelves, the maximum density of SSLs are often correlated with maximum chlorophyll-a concentrations (Berge et al., 2014; Dekshenieks et al., 2001; Holiday et al., 2010). Dissolved oxygen concentrations (above 1 mL L⁻¹, i.e., 44.661 mmol m⁻³) can also predict the lower boundary of SSL density, e.g., in eastern boundary upwelling systems (EBUSs), like the Peruvian coastal upwelling system (Bertrand et al., 2010) and the California coastal upwelling system (Netburn and Koslow, 2015).

In this study, we use acoustic tools (Simmonds and MacLennan, 2005a) to examine the fine-scale vertical structure of SSLs (i.e., their depth in the water column, thickness, and density) (Bertrand et al., 2013; Perrot et al., 2018). We use fine spatiotemporal resolution of acoustic data to investigate how the pelagic environment influences SSLs in the EBUS off Senegal during an upwelling event. Our objective was to model variations in SSLs structure relative to physicochemical characteristics of water masses and their locations on the shelf.
2 Materials and methods

2.1 SSLs acoustics sensing and environmental data

We performed a hydroacoustic survey along the “Petite Côte”, south of Cap-Vert peninsula off Senegal (14.6–13.5° N, 16.9–17.6° W). The survey was conducted with the research vessel *Antea* of the French National Research Institute of Sustainable Development (IRD France) during the upwelling season from 6 to 18 March 2013. The Petite Côte is a nursery area for fish and is the main area in which juveniles of numerous species (particularly small pelagic species) concentrate (Diankha et al., 2018; Thiaw et al., 2017). Strong upwelling occurs during spring, which contributes to high primary productivity, thus providing an ideal nursery area for commercially important fish species (Tiedemann and Brehmer, 2017).

We collected hydroacoustic data along three transects (T1 (North), T2 (intermediary), and T3 (south)) in 18 nautical miles (nmi) perpendicular to the coast (Fig. 1). Hydroacoustic data were continuously recorded (day and night) using a Simrad EK60 echo sounder (38, 70, 120, and 200 kHz), set at 20 log R time-varied gain function (where R is the range in meters) and using a pulse length of 1.0 ms. In this study, we used the acoustic monofrequency approach (using 38 kHz, one of the most current frequencies used in fisheries surveys) to study the spatiotemporal SSL structure. The 38 kHz frequency offers the advantages of depth penetration, covering the whole vertical range of SSLs. The multifrequency echogram was used to identify the main scatterers of SSLs and to justify the SSLs extraction threshold (see below). Transducers were calibrated following the procedures recommended in Foote et al. (1987). Considering the aft draught of the vessel, the acoustic near field, and the presence of acoustic parasites (including air bubbles) in the upper part of the water column, we have applied an offset of 10 m (acoustic data above 10 m have been deleted). Echoes along the three transects were integrated at a spatial resolution of 0.1 nmi × 1 m depth. We estimated the SSL acoustic density by calculating the nautical area scattering coefficient (NASC or $s_A$), which represents the relative biomass of acoustic targets. We assumed that the composition of the scattering layers and the resulting scattering properties of organisms in the SSLs are homogeneous within each layer we identified (sensu MacLennan et al., 2002). We analyzed integrated echoes using the in-house tool “Matecho” (Perrot et al., 2018). Matecho is an integrative processing software that allows us to manually correct echograms (e.g., by correcting bottom depths, removing empty pings, removing echogram interferences, and reducing background noise). For echo-integration accuracy, Matecho computes a quality factor (QC) (Fig. S1 in the Supplement) for each echo-integration cell, which is the number of integrated samples divided by the total number of samples in one echo-integration cell. After each echogram correction, we extracted the SSLs that were below the mean acoustic volume backscattering strength ($S_v$ in dB) threshold of −75 dB (i.e., values below −75 dB were excluded from the analysis). Cascão et al. (2017) and Saunders et al. (2013) excluded marine pelagic organisms that backscattered at −70 dB, a threshold based on the aggregative behavior of marine pelagic organisms. The SSL extraction method is based on a threshold of −75 dB and a MATLAB algorithm used in Matecho named “contourf.m” (https://ch.mathworks.com/help/matlab/ref/contourf.html, last access: 12 January 2019), which appear relevant to extract the main SSL at 38 kHz (Fig. S2). This process performs a segmentation of the echo integration from the given threshold on echo levels to extract (by calculation of isolines according to the selected $S_v$ threshold) the attached echo groups that formed the SSLs and their associated contours. Based on this contour, a set of descriptors are estimated, e.g., up and down depths of SSL and thickness. In our study, the backscattering was due to zooplankton and micronekton, as well as small pelagic fish. The inshore area is known to be rich in copepod and fish larva (Ndour et al., 2018; Tiedemann and Brehmer, 2017); however, a low sample number was collected in the coastal inshore water due to safety reasons, i.e., the research vessel investigated areas of >20 m bottom depth.

We collected hydrographic data using a calibrated Sea-Bird SBE 19plus conductivity, temperature, and depth (CTD) probe. The CTD specifications for temperature were...
±5.10^{-3} °C accuracy and 1.10^{-4} °C precision; for conductivity, they were ±5.10^{-3} S m^{-1} accuracy and 5.10^{-3} S m^{-1} precision; for pressure, they were ±0.1 % of full-scale range accuracy and 2.10^{-3} % precision of full-scale range precision. The CTD was equipped with sensors for fluorescence (±2.10^{-3} µg L^{-1} accuracy, and ±2.10^{-4} µg L^{-1} precision) (a measure of chlorophyll-a concentration, a proxy for phytoplankton biomass), and dissolved oxygen sensor (DO, mmol m^{-3}, Sea-Bird SBE 43, 2 % saturation for accuracy and 0.2 % saturation for precision). The CTD has been calibrated before the survey. During the survey, data delivered by the SBE 43 for DO have been corrected by Winkler titrations. From 6 to 8 March 2013, we conducted CTD casts along three transects at 36 stations. At each station, sensors measured water temperature (°C), depth (m), fluorescence (µg L^{-1}), water density (here sigma-theta, kg m^{-3}), and DO. Global High Resolution Sea Surface Temperature (GHRSSST) data were extracted from daily outputs by the Regional Ocean Modeling System group at NASA’s Jet Propulsion Laboratory (JPL OurOcean Project, 2010). Daily SST data (GHRSSST Level 4 GISST Global Foundation Sea Surface Temperature Analysis) were averaged for the 3 d of surveying using SeaDAS software version 7.2 (https://seadas.gsfc.nasa.gov/, last access: 10 November 2018) and interpolated on maps using R software (R Core Team, 2016). Cubic spline interpolations of gridded data were used within the R package Akima (Akima et al., 2016).

2.2 Data analysis

After extracting SSLs with Matecho, we developed an ad hoc MATLAB extension of Matecho named “Layer” (S1 in the Supplement). We obtained SSL thickness, minimum and maximum SSL depths (D_{min} and D_{max}, respectively), and an echo-integrated echogram from Matecho output files to provide it to another MATLAB program “ComparEchoProfil” (S2). ComparEchoProfil allows the user to fit in time and depth echo-integrated echograms to the associated CTD vertical profiles. We used the equation below to calculate thickness:

\[
\text{Thickness} = D_{max} - D_{min}
\]  

(1)

Mean nautical area backscattering coefficient (s_{A}, NASC) and mean acoustic volume backscattering strength (S_{V} in dB) profiles were based on the average of three ESUs (small-scale elementary sampling units): the ESU nearest to the CTD position (ESU_{ctd}) as well as previous and following in correspondence with CTD depths (d_{n})

\[
\bar{s}_A(d_{n}) = \sum_{i=ESU_{ctd}+1}^{i=ESU_{ctd}-1} s_A(i, d_{n}) / 3,
\]  

(2)

\[
S_V(d_{n}) = 10 \times \log_{10} \left( \sum_{i=ESU_{ctd}+1}^{i=ESU_{ctd}-1} 10^{S_A(i, d_{n}) / 10} / 3 \right),
\]  

(3)

The ComparEchoProfil displayed the profile for S_{V} in dB over an ESU of 0.1 nmi around each CTD station. The program also allowed us to display acoustic profiles for physicochemical parameters (temperature, CHL (chlorophyll), density, and DO) associated with S_{V} profiles (Fig. 2). The output included meta-information (station ID, station date, station time, latitude and longitude, diel phase (day, night), and bottom depth), all of which we associated with SSL descriptors (SSL thickness, maximum SSL depth, S_{A}, and S_{V}) based on classic fish school descriptors (Brehmer et al., 2007, 2019) and physicochemical parameters associated with each SSL.

We applied hierarchical cluster analyses (HCAs) to discriminate between water masses of inshore and offshore stations over the continental shelf based on CTD data collected at 10 m depth. HCA was based on Euclidean distance and Ward’s aggregation method (Ward, 1963). We used principal component analysis (PCA) (Chessel et al., 2013) on the same dataset to determine similarities between CTD stations relative to environmental parameters. Physicochemical parameters were standardized a priori because they were measured with different metrics.

Inshore–offshore variability of morphometric (thickness, depth) and acoustic characteristics (s_{A}) of the SSLs are investigated in the discriminated groups considering bottom depth and diel period. Diel transition periods are removed from analyses to avoid SSL density change biases due to diel vertical migrations. Transition periods are defined using sun altitude, i.e., around sunset and sunrise corresponding to a sun altitude between ±18° (Lehodey et al., 2015). Morphometric and acoustic characteristics of the SSLs are also compared between the inshore area versus offshore area and between day and night using Student’s t test whose application conditions have been verified (normal distribution and variance equality).

Echogram vs. profile coupling figures (Fig. 2) resulting from the ComparEchoProfil were analyzed to determine the relation between environmental parameters and SSLs. ANCOVA tests (analysis of covariance) (Wilcox, 2017) were implemented for SSL characteristics (thickness, depth, and density) in each discriminated area (inshore and offshore). These models were set to predict each descriptor, i.e., thickness, depth, and s_{A} as a function of temperature, density, DO, CHL, local depth, and diel period. The ANCOVA models were developed on averaged data over station. The selection of the best models was performed using stepwise procedures. Stepwise selection was based on minimizing the Akaike information criterion (AIC) (Akaike, 1974). The relative importance of each variable in total deviance explained was determined from the “relaimpo” R package (Tonbrandel and LeBreton, 2011). Validity assumptions of the models were then assessed by checking for normality of distributed errors and homogeneity of residuals (Figs. S3 to S5). For the ANCOVA, SSL density (s_{A}) was log_{10} transformed for normality assumption. For all statistical tests, the significance threshold used was 0.05.
Figure 2. Echograms and associated vertical acoustic profiles as well as physicochemical parameters (CTD data) for two example stations: (a) station 19 in the inshore area and (b) station 12 in the offshore area. For both (a) and (b), top panels are echogram data collected along the transect, i.e., 1000 ESU (elementary sampling unit) of 0.1 nmi, whereas the bottom panels depict acoustic and environmental data (depicted by the vertical red line in top panels). Environmental data for the sound-scattering layer (SSL) were collected at the stations at the locations depicted by dotted vertical lines. Data represent mean conditions for the station collected within an area of 0.1 nmi around the station: acoustic volume backscattering strength ($S_v$) SSL, temperature profile SSL, CHL profile SSL, oxygen profile SSL, and density profile SSL. The horizontal dashed lines in all profiles represent the SSL thickness, i.e., the upper and lower SSL limits.

We used R software (R Core Team, 2016) for statistical analyses and to map data. We used the R package “Cluster” (Maechler et al., 2014) for HCA of CTD data, the R package “maps” (Brownrigg, 2017) to map stations, the package “ade4” (Chessel et al., 2013) to run a PCA, and the package “oce” (Kelley, 2015) to display vertical section plots of physicochemical parameters.

3 Results

3.1 Characterization of two water masses over the shelf

The HCA differentiated two groups of stations (Fig. 3a): Group 1 (G1) stations ($n = 18$) comprised four stations along transect T1, six stations along transect T2, and eight stations along transect T3. The stations of G1 were located closest to
Figure 3. Discrimination of 36 CTD stations off the Senegalese coast: (1) two groups of stations were discriminated based on temperature (temp), chlorophyll a (CHL), dissolved oxygen (oxy), and density (dens). (2) Principal component analysis of environmental parameters for all 36 stations. (a) Eigenvalue diagram; (b) factor plane; (c) correlation circle. Group 1 represents stations located in the inshore area ($n = 18$); Group 2 represents stations located in the offshore area ($n = 18$).

the coast (inshore area, from 13 to 61 m bottom depth, which encompassed the core of the upwelling (based on data for sea surface temperature) (Fig. 1). Group 2 (G2) stations ($n = 18$) comprised seven stations along transect R1, six stations along transect R2, and five stations along transect R3. These stations were located furthest from shore (offshore area), from 41 to 205 m bottom depth, which corresponds to the outer border of the upwelling zone. Considering the bathymetry, we note an overlay of the two areas discriminated between 41 and 61 m.

PCA identified the same two distinct water masses that were clustered in HCA (Fig. 3). Axis 1 of the PCA eigenvalues explained 72.8% of the inertia, whereas axis 2 explained 26.8%. On axis 1 of the PCA plot, temperature was highly correlated with density. On axis 2, temperature and DO were opposed to CHL. The distribution of these variables is related to the station groupings: G1 (inshore area) was characterized by a dense and CHL-rich water mass, whereas G2 (offshore area) was characterized by a warm and slightly oxygenated surface water mass.

Satellite measurements of SST distributions of the study area indicated the same split of stations into two groups (Fig. S6). The inshore area was characterized by low SST values (18–19°C), indicating a recently upwelled water mass, whereas an older water mass with higher SST values (20–21°C) prevailed offshore.
At transect T1, a marked frontal zone appeared isolating two water masses between the 20 and 40 m isobaths (Fig. 4a1), which separated warm surface waters from deep, cold, upwelled water masses. At transects T2 and T3, the upwelling appeared as a cold-water tongue isolating a warm water band at the coast (Fig. 4a2, a3). At T3, this cold-water tongue was expanding toward the inshore area as well as to the offshore area (Fig. 4a3). Surface water masses of the inshore area were slightly denser than water masses in the offshore area with approximately 26 and 25 kg m\(^{-3}\), respectively. For CHL, elevated concentrations were exclusively observed in the inshore area at transects T1 and T2. CHL was significantly higher in the inshore area than the offshore area with concentrations of 3.0–5.0 mg m\(^{-3}\) in the inshore area to 0.3–2.0 mg m\(^{-3}\) in the offshore area (Fig. 4c). At T3, the elevated CHL concentrations were observed in both inshore and offshore areas close to the upwelling front. CHL was higher in the upper part of the water column (0–20 m), decreasing with depth in both areas. Higher DO concentrations were observed towards both sides of the upwelling core. At T1, the upwelling front was at the most coastal part, separating the inshore area from the less oxygenated offshore area with DO concentrations of 223–312 and 178–223 mmol m\(^{-3}\), respectively. At T2 and T3, the core moved towards the offshore, separating the inshore area (DO concentrations of 178–223 mmol m\(^{-3}\)), slightly more oxygenated than the offshore area (DO concentrations of 89–178 mmol m\(^{-3}\)). DO concentration decreased from the surface to bottom in both areas.

### 3.2 Variability in vertical structure of SSLs

#### 3.2.1 Spatial variability according to water mass characteristics

Thickness and depth of the SSLs varied according to bottom depth in the inshore area and the offshore area. In the inshore area, on the northern transect T1, no SSLs were observed at coastal stations shallower than 29 m bottom depth (stations 1 and 2) (Fig. 5a). In offshore stations, starting at 41 m bottom depth, the SSLs were observed in all stations and transects (Fig. 5b), and their thickness and depth increased with bottom depth. SSL thickness and SSL depth differed significantly between the inshore area and the offshore area: the SSLs were thicker and deeper in the offshore area than in the inshore area (Fig. 6) (\(p\) value = 0.001 for both thickness and depth). An increase of SSL was observed with increasing bottom depths in the inshore area and the offshore area. The \(s_A\) comparison between the inshore area and the offshore area (Fig. 6) was not significantly different (\(p\) value = 0.833).

#### 3.2.2 Diel migration

The diel period had a significant effect on SSL thickness (\(p\) value < 0.001) and SSL depth (\(p\) value < 0.001), which
were found higher during both the night in the inshore area and the offshore area (Fig. 6). In the inshore area, during daytime, the mean depth and thickness of SSL were 19 and 11 m, respectively, while during nighttime the mean depth and thickness were 46 and 35 m, respectively. In the offshore area, SSLs were found at a mean depth and thickness of 49 and 44 m, respectively, during both the night in the inshore area and daytime, while during nighttime the mean depth and thickness of SSL were 19 and 38 m, respectively, during daytime, while during nighttime SSL depth and thickness were 86 and 75 m, respectively. Mean $s_A$ (Fig. 6) of SSLs also varied between day and night but were not significantly different ($p$ value = 0.890). In the inshore area, the mean $s_A$ was 24 m$^2$ nmi$^{-2}$ during the day and 44 m$^2$ nmi$^{-2}$ during the night. In the offshore area, the mean $s_A$ was 46 m$^2$ nmi$^{-2}$ during daytime and 25 m$^2$ nmi$^{-2}$ during nighttime. Mean $S_v$ distribution of SSLs (Fig. S7) also showed a diel variation with mean $S_v$ higher at night than during the day.

### 3.2.3 Vertical dimension of SSLs related to physicochemical profile

In both areas, SSLs were partially or completely located in areas of strong vertical gradients of temperature (thermocline), density (pycnocline), and DO (oxycline) (Fig. 2). When a strong temperature gradient was observed, usually also associated with the vertical position of the oxycline and a pycnocline, a peak of CHL was often observed and matched with the volume backscattering strength ($S_v$) peak (Fig. 2a). This observation is well illustrated in CTD stations 12, 13, 16, and 25 (Fig. S8). In the inshore area the peak of CHL concentration was always located above the SSLs (Fig. 2a), whereas in the offshore area, the peak of CHL concentration was either above the SSLs or in the middle of the SSLs (Fig. 2b). The thickest SSLs were observed in the offshore area where temperature, density, and oxygen gradients were strong.

### 3.2.4 Behavior of the SSLs relative to pelagic habitat characteristics

#### In the inshore area (G1)

In the inshore area (G1), the ANCOVA model indicated a strong effect of bottom depth and diel period on both SSL thickness and depth. For SSL thickness, the model (Tables 1, S1 in the Supplement) explained 87% of the variance ($R^2 = 0.869, p$ value = 0.001). Bottom depth explained 56% of SSL thickness, while the diel period effect accounted for 31%. The model of SSL depth (Tables 2, S2) was like those of SSL thickness; i.e., the model included bottom depth and diel period explaining 80% of the variance ($R^2 = 0.805, p$ value = 0.001). Bottom depth showed the largest effect on SSLs explaining 51% of SSL depth, while the diel period effect was estimated at 30%. For SSL acoustic density, i.e., $\log (s_A)$ (Tables 3, S3), the model explained 40% of the variance ($R^2 = 0.398, p$ value = 0.022), indicating a single effect of bottom depth on $\log (s_A)$ ($p$ value = 0.20). The bottom depth was the only variable significant in the model and explained 33% of SSL acoustic density. Temperature was insignificant in the model.

The ANCOVA models to predict SSL thickness and SSL depth can be expressed as

- SSL thickness: $\text{SSL-thickness} = -11.865 + (0.916 \times B_d) + (11.492 \times D_p)$,
- SSL depth: $\text{SSL-depth} = -4.223 + (0.954 \times B_d) + (12.864 \times D_p)$,

with $B_d$ being bottom depth in meters and $D_p$ being diel period at night.

#### In the offshore area (G2)

For offshore stations, the model showed a significant effect of diel period, temperature, water density, and DO on both thickness and depth of SSLs with similar results. Both models, SSL thickness (Tables 1, S1), and SSL depth (Tables 2,
Figure 6. Box plot (minimum, maximum, and median) of sound-scattering layers (SSLs) mean depth (m), thickness (m), and relative biomass ($s_A$ in m$^2$ nmi$^{-2}$) grouped by diel period (day and night) for (a) inshore area and (b) offshore area over the Senegalese continental shelf.

S2) included bottom depth, diel period, temperature, density, and DO explaining 85\% of variance ($R^2 = 0.855$, $p$ value = 0.001). Bottom depth and diel period accounted for 28\% and 28\%, respectively. Other significant variables were water temperature, density, and DO, which support 11\%, 10\%, and 7\%, respectively. For SSL density or log($s_A$) (Tables 3, S3), none of the predictor variables had a significant effect.

The ANCOVA models to predict SSL thickness and SSL depth can be expressed as

$$\text{SSL thickness} = 56030 + (0.21 \times B_d) + (27.35 \times D_p) + (-383.80 \times T) - (1898 \times D) - (1.76 \times O_2),$$

$$\text{SSL depth} = 56040 + (0.21 \times B_d) + (27.35 \times D_p) + (-383.80 \times T) - (1898 \times D) - (1.76 \times O_2),$$

with $B_d$ being bottom depth in meters, $D_p$ being diel period at night, $T$ being water temperature in degrees Celsius ($^\circ$C), $D$ being water density (kg m$^{-3}$), and $O_2$ being oxygen (mmol m$^{-3}$).

4 Discussion

4.1 Characterization of water masses along the Petite Côte

Upwelling is a key process for the functioning of the coastal ecosystem of Senegal and Mauritania (Capet et al., 2016; Estrade et al., 2008; Rebert, 1983). By characterizing the physicochemical parameters of the Petite Côte, we were able to discriminate two water masses: an inshore area and the offshore area, both of which could also be distinguished with SST satellite data.

Analyzing the spatial structure of SST helped to understand the upwelling dynamics along the Petite Côte. The SST pattern, measured at the time of our survey, was in line with prior studies. During the upwelling season (in winter and late spring), a tongue of cold water over the shelf isolates a coastal band of warm water from the offshore area, and there is a surface separation associated with the upwelling source over the shelf and convergence nearshore.
Table 1. Result of ANCOVA models between thickness of sound-scattering layers (SSLs) and environmental parameters (temperature, density, dissolved oxygen, chlorophyll $a$, diel period, and bottom depth) in the inshore area (G1) and the offshore area (G2). G1: multiple $R^2$ is 0.869, adjusted $R^2$ is 0.8515, and $p$ value < 0.001; G2: multiple $R^2$ is 0.8557, adjusted $R^2$ is 0.7956, and $p$ value < 0.001; significant $p$ values in bold.

| Variable          | Significance | Explained deviance (%) | Total explained variance (%) |
|-------------------|--------------|------------------------|------------------------------|
|                   | Inshore (G1) | Offshore (G2)          | Inshore (G1) | Offshore (G2) |
| Bottom depth      | 0.001        | 0.005                  | 55.86         | 28.05         |
| Diel period (night) | 0.007       | 0.008                  | 31.02         | 28.33         |
| Temperature       | 0.007        |                        | 11.29         |               |
| Density           | 0.008        |                        | 10.35         |               |
| Oxygen            | 0.007        |                        | 7.53          |               |

Table 2. Result of ANCOVA models between depth of sound-scattering layers (SSLs) and environmental parameters (temperature, density, dissolved oxygen, chlorophyll $a$, diel period, and bottom depth) in the inshore area (G1) and the offshore area (G2). G1: multiple $R^2$ is 0.8056, adjusted $R^2$ is 0.7797, and $p$ value is 0.001; G2: multiple $R^2$ is 0.8557, adjusted $R^2$ is 0.7956, and $p$ value is 0.000; significant $p$ values in bold.

| Variable          | Significance | Explained deviance (%) | Total explained variance (%) |
|-------------------|--------------|------------------------|------------------------------|
|                   | Inshore (G1) | Offshore (G2)          | Inshore (G1) | Offshore (G2) |
| Bottom depth      | 0.001        | 0.005                  | 55.86         | 28.05         |
| Diel period (night) | 0.021       | 0.008                  | 31.02         | 28.33         |
| Temperature       | 0.007        |                        | 11.29         |               |
| Density           | 0.008        |                        | 10.35         |               |
| Oxygen            | 0.007        |                        | 7.53          |               |

tial difference of CHL concentration between the inshore area and the offshore area is the result of upwelled water carrying nutrients to the coast, which is separated by water mass fronts. Nutrient-rich water, supplied to the sunlit surface layer by wind-driven upwelling, stimulates the growth of phytoplankton that ultimately fuel diverse and productive marine ecosystems (Jacox et al., 2018). There is a link between the accumulation of biological material and the location of the coastal band of warm water. This coastal band between coast and the upwelling core has been regarded to function as a retention area in which nutrient particles are trapped (Demarcq and Faure, 2000; Roy, 1998). The nutrient utilization is optimized by retentive physical mechanisms in the coastal area, which enhances microbial remineralization of particulate organic matter and zooplankton excretion and then regenerates production through ammonium consumption (Auger et al., 2016). This causes an increase in primary production and results in a surplus of phytoplankton biomass in inshore areas. Low DO concentrations observed in the upwelling core separating more oxygenated water masses have been reported in previous studies (Capet et al., 2016; Teisson, 1983) over the Petite Côte. Once a water mass becomes isolated from the atmosphere, its oxygen content starts to decrease due to biological remineralization of dissolved organic matter (Emerson et al., 2008; Machu et al., 2019). These low-oxygen bottom waters are transported to the inner shelf during upwelling-favorable wind events. Moreover, temporal stability of the upwelling core is also noticeable over periods of several days to weeks, and export from the shelf to the open ocean is retarded (Capet et al., 2016). Thus, in such favorable conditions of continuous food supply, photosynthesis may foster an enrichment of DO in the inshore. This is in line with high CHL levels observed towards both sides of the upwelling core, particularly in the inshore area.

4.2 Spatial variation of the SSLs off the Petite Côte of Senegal

We measured a longitudinal gradient of the thickness of the SSLs over the continental shelf. The SSLs were concentrated in a narrow band in the inshore area, whereas the SSLs were wider in the offshore zone. The absence or weakness of SSLs in the inshore area (in contrast to the more stratified water column in the offshore area) may have been due to turbulence in the water column (Sengupta et al., 2017), coupled with a well-mixed surface layer. In the inshore area, it is
likely that turbulence and the probable low residence time of marine pelagic organisms advected from outside this area and both inhibited SSL formation. Indeed, in such upwelling systems, in addition to the retention mechanism that has been recognized by several authors (Aristegui et al., 2009; Capet et al., 2016; Mbaye et al., 2015; Roy, 1998), there is also an offshore Ekman transport mechanism (Aristegui et al., 2009; Estrade et al., 2008) that contributes to cross-shore exchanges. Otherwise, different animals can respond very differently to different physical forcing. Many authors have stressed that SSLs need stable hydrological conditions to form (Aoki and Inagaki, 1992; Baussant et al., 1992; Marchal et al., 1993). As an example, in Monterey Bay (California), Urmy and Horne (2016) observed a decline in acoustic backscatter intensity in the upper part of the water column immediately following an upwelling event. In a more recent study, Benoit-Bird et al. (2019) found that when upwelling was strong both krill and anchovies were found in small, discrete aggregations, while during upwelling relaxation and reversals forage biomass was more diffusely distributed. Therefore, we assume that the increase of SSL thickness with depth from inshore to offshore off Senegal is caused by upwelled waters that disrupt the vertical stability of the water column. Therefore, although the SSLs are first constrained by the bottom depth (i.e., room available), we assume that the increase of SSL thickness with depth from inshore to offshore off Senegal is caused by upwelled waters that disrupt the vertical stability of the water column.

4.3 Diel temporal variation of SSLs

In our study area, the diel period consistently exhibited pronounced effects on SSL thickness and depth. Deeper nighttime SSLs have a greater thickness than daytime SSLs. The diel difference in thickness and depth is due to the well-known DVM patterns performed by many marine species. DVM is a behavioral mechanism usually characterized by an ascent during nighttime for feeding and a descent to avoid predation by visual predators during daytime known as type I (Bianchi et al., 2013; Haney, 1988; Lehodey et al., 2015). Some planktonic and micronektonic organisms have been reported to exhibit reverse DVM (type II), i.e., ascending in the morning and descending in the evening or early night, which is the opposite pattern observed with vertically migrating animals (Cushing, 1951; Ohman et al., 1983). The main SSLs scatter much more strongly at 38 kHz than at 70 or 120 kHz (Fig. 2); the backscattering response is probably dominated by animals with swim bladders such as fish larvae and small fish (Simmonds and MacLennan, 2005a). Indeed, the Petite Côte is a nursery area for fish and is the main area in which juveniles of numerous species concentrate (Diakha et al., 2018; Thiaw et al., 2017). Moreover, Tiedemann and Brehmer (2017) have reported fish larvae (Sardinella aurita, Engraulis encrasicolus, Trachurus trachurus, Trachurus tereca, Microchirus ocellatus, and Hygophum macrochi) all along our study area. Previous studies have reported that DVM of plankton may increase coastal retention in the inshore area (Brochier et al., 2018; Mbaye et al., 2015; Rojas and Landeta, 2014). Diel variation was also observed for SSL acoustic density, which showed opposite patterns in the two areas, i.e., higher up in the water column during night than day in the inshore area and higher up during days than at night in the offshore area. Tiedemann and Brehmer (2017) observed that all fish larvae in the offshore area, except Trachurus trachurus, exhibited a DVM type II, and their observations are in accordance with the DVM pattern of SSL acoustic density reported in our study. Another possible explanation of this observed diel variation is the horizontal migration. DHMs are known as nocturnal horizontal migration of both plankton and consumers into shallow and inshore waters (Benoit-Bird et al., 2001; Benoit-Bird and Au, 2006). DHMs have been observed in marine copepods (Suh and Yu, 1996) which represent the main zooplankton group in the study area (Ndour et al., 2018; Rodrigues et al., 2017). It is hypothesized that these inshore–offshore migrations are a strategy for avoiding visual predators (White, 1998), and they result in increased access to food resources relative to simple vertical migration (Benoit-Bird et al., 2008). Otherwise, DVM of marine pelagic organisms may not be the only factor causing diel backscatter variations.

i. The acoustic target strength can be strongly dependent on the aspect at which a target is insonified. Target strengths of zooplankton and micronekton can vary by several orders of magnitude between extreme tilt angles, i.e., horizontal vs. head up or head down (Benoit-Bird and Au, 2004; Yasuma et al., 2003). Target strength is not independent of depth, as migrations through the hydrostatic depth gradient can alter, e.g., swim bladder volume (Fässler et al., 2009). This can bias target strengths, in particular near the resonance frequency, leading to artificial increases of backscatter at a particular depth (Davison et al., 2015; Godø et al., 2009; Kloser et al., 2002).

ii. In the inshore area the CTD sampling was mainly achieved during the daytime, which may have biased the observed DVM type I.

iii. Otherwise, plankton such as fish larvae are able to perform a DVM type II by ascending in the upper 10 m of the water column at night, i.e., in the echo sounder offset.

4.4 Effect of environmental parameters on SSLs

4.4.1 SSLs related to physicochemical parameters in the vertical dimension

Previous studies have shown that hydrographic structures of the water column influence SSL vertical structure (Balino
and Aksnes, 1993; Berge et al., 2014; Gausset and Turrel, 2001). In our case study the results show that vertical distribution of SSLs was linked to strong vertical gradients of temperature, DO, and water density (Fig. 2). The peak of $S_N$ was sometimes very close to the strong gradient of water temperature, density, CHL, and DO (Fig. S8). The depth of SSLs has been reported to be related to the thermocline (Marchal et al., 1993; Yoon et al., 2007). In more stratified areas, SSL vertical distribution was limited by a strong thermocline and when thermocline was not well marked (low gradient) SSLs occupied the entire water column (Lee et al., 2013). Olla and Davis (1990) and Rojas and Landau (2014) suggested that the thermocline is a physical barrier that acts above or below in the vertical distribution of some fish larvae, while other studies (Gray and Kingsford, 2003; Tiedemann and Brehmer, 2017) showed no effect of a thermocline on vertical larval fish distributions. In this study, the SSL was correlated to temperature in the offshore stratified area, but it did not act as a physical barrier limiting vertical distribution. Previous studies (Bertrand et al., 2010; Bianchi et al., 2013; Netburn and Koslow, 2015) have suggested that vertical distributions of SSL organisms are limited by mid-water DO concentrations which constrain SSL depth. These authors found a relationship between SSL depths and hypoxia. However, in our study, we found correlations between SSLs (depth, thickness) and DO as expected, but the vertical distribution of SSLs was not constrained by DO. SSLs were also observed in some hypoxic stations (DO < 1.42 mL L$^{-1}$, i.e., 63.42 mmol m$^{-3}$); consequently, DO was not a limiting factor for SSLs organisms. Fish larvae respond to oxygen gradients by moving upwards or laterally (Breitburg, 2002, 1994). Vertical movement of fish larvae may also be related to the avoidance of predators, which are limited to well-oxygenated layers. The high phytoplankton concentration found in this study, particularly in the inshore area, may be interpreted as a potential food source for fish larvae, which are able to perform DVM towards the surface. The vertical position of SSLs compared to the CHL concentration peak can be explained by trophic relationships between phytoplankton, zooplankton, and micronekton. It is understood that zooplanktivorous micronekton migrates upward in the water column to forage on mesozooplankton while the mesozooplankton is migrating toward the surface to graze upon the phytoplankton. This trophic relationship may explain the link in vertical position of the SSLs with the phytoplankton peak reported in this study.

### 4.4.2 Behavior of SSLs relative to pelagic habitat characteristics

In the inshore area, where SSLs were sparsely present (or sometimes non-existent) bottom depth and diel period were the main environmental parameters influencing the vertical distribution (thickness and depth) of the SSLs. Bottom depth has been shown to regulate the vertical distribution of SSLs in the water column (Donaldson, 1967; Gausset and Turrel, 2001; Torgersen et al., 1997). In our study, all stations indicated a single SSL, while in deep water more thick and deep SSLs are often partitioned into multiple layers (Ariza et al., 2016; Balino and Aksnes, 1993; Cascão et al., 2017; Gausset and Turrel, 2001). Diel period is the second most important parameter acting on SSL thickness and depth through the DVM phenomenon. In well-mixed water masses, temperature, density, and oxygen had no effect on the SSLs. The insignificant effect of temperature, oxygen, and water density on the SSLs in the inshore area is explained by the presence of less marked and superficial clines because of the newly upwelled water. As stated above, SSLs probably need stable conditions to occur.

In the offshore area, where vertical gradients were marked, the main parameters structuring SSL thickness and depth were bottom depth and diel period but also water temperature, density, and DO. DVM behaviors are influenced by environmental cues (e.g., light, nutrients, and temperature) and predator-prey interactions (Clark and Levy, 1988; Lampert, 1989). Relative changes in light intensity are identified as the most important proximate stimuli driving DVM, in-

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Table 3. Result of ANCOVA models between sound-scattering-layer (SSL) density (log($s_N$)) and environmental parameters (temperature, density, dissolved oxygen, chlorophyll $a$, diel period, and bottom depth) in the inshore area (G1) and the offshore area (G2). G1: multiple $R^2$ is 0.398, adjusted $R^2$ is 0.3178, and $p$ value is 0.022; G2: multiple $R^2$ is 0.3448, adjusted $R^2$ is −0.01258, and $p$ value is 0.490; significant $p$ values in bold.

| Variable          | Significance | Explained deviance (%) | Total explained variance (%) |
|-------------------|--------------|------------------------|-----------------------------|
|                   | Inshore (G1) | Offshore (G2)          | Inshore (G1)                | Offshore (G2)          |
|                   |              |                        |                             |                           |
| Bottom depth      | **0.008**    | 0.357                  | 33.06                       | 7.56                      |
| Temperature       | 0.119        | 0.273                  | 6.73                        | 5.17                      |
| Diel period (night) | 0.007    | 0.546                  | 7.22                        |                           |
| Density           | 0.008        | 0.250                  | 5.56                        |                           |
| Oxygen            | 0.007        | 0.166                  | 5.19                        |                           |
including the amplitude of the migration as well as the timing of the up- and downward movement (Meester, 2009). SSL vertical distribution is also known to be a function of temperature (Bertrand et al., 2010; Hazen and Johnston, 2010; Netburn and Koslow, 2015). Overnight, depths of the SSLs are strongly correlated to the depth of thermal and density gradients (Boersch-Supan et al., 2017; Cascão et al., 2017; Marchal et al., 1993). In the offshore area, the results suggest that DO also influences SSL depth and SSL thickness. In well-oxygenated continental shelf waters, DO influences SSLs but does not limit their vertical distribution. Some previous work in French Polynesia (Bertrand et al., 2000), and in the southern California current ecosystem (Netburn and Koslow, 2015), showed that the oxygen minimum zone (OMZ) acts like a barrier to SSLs in their vertical distribution. Bianchi et al. (2013) suggest that distribution of the open-ocean OMZ may modulate the depth of migration at the large scale so that organisms within SSLs migrate to shallower waters in low-oxygen regions and to deeper waters in well-oxygenated waters. For both areas, CHL concentration was the only predictor that was not included in any of the final models. However, by coupling echogram and profile data (Fig. 2), we can argue that a relation between CHL and SSL exists even if it was not significant in the models, because CHL and SSL biomass peaks matched, i.e., always located above or in the middle of the SSLs. Moreover, a simple linear model between CHL and SSL structure (depth and thickness) was significant in the inshore area, suggesting that the effect of CHL on full models was masked by autocorrelation between predictive variables.

Fish larvae vertical distributions have been related to the distribution of their prey and predator, and it has been argued that the presence and position of the thermocline is an important feature in their vertical distribution (Haney, 1988; Röpke, 1993). Other studies have shown that the thermocline has only a limited role in the vertical distribution patterns of fish larvae (Gray, 1996; Gray and Kingsford, 2003). Indeed, in coastal areas, where the structure of the water column is less regular than in the open sea, the vertical distribution of fish larvae depends on the physics of the water column (Sánchez-Velasco et al., 2007) but also on the behavior of each species (Fortier and Harris, 1989). According to Sánchez-Velasco et al. (2007), the vertical distribution of fish larvae is closely related to the changes in the water column structure, with most fish larvae concentrated in the stratum of maximum stability. Therefore, the vertical stratification level in water column is strongly related to vertical distribution of these organisms.

Furthermore, the vertical distribution of SSLs can be influenced by mixed layer depth (MLD). The MLD is one of the primary factors affecting the vertical distribution of zooplankton. Lee et al. (2018) have shown that the weighted mean depths of SSLs exhibit a strong linear relationship with the MLD, meaning that the MLD could be a significant environmental factor controlling the habitat depth of marine pelagic organisms. A recent study (Stranne et al., 2018) has shown that the MLD can be tracked acoustically at high horizontal and vertical resolutions. The method was shown to be highly accurate when the MLD is well defined and biological scattering does not dominate the acoustic returns. However, in our study area, biological scattering dominated the acoustic records and due to the upwelling acoustic methods were not appropriate to determine MLD.

5 Conclusions

Using our echogram vs. profile coupling approach, we were able to examine fine-scale processes affecting SSL distributions. SSLs were influenced by turbulence level in the upwelling, which led to an offshore advection of SSL organisms. SSL distributions were mainly structured by bottom depth, diel period, and the level of vertical stratification in water column. SSL acoustic density variation suggested different diel migrations: a normal and reverse DVM, and/or a DHM. Such an observation should be considered in modeling exercises to better understand DVM implications in ecosystem functioning. Further investigations should integrate small-scale turbulence measurements to better describe the fine-scale spatiotemporal variability of SSLs and their relationship to the pelagic environment. Information on SSL species composition and morphological characteristics will provide an accurate description of their fine-scale relationships in the pelagic habitat.

Code availability. “Matecho” is an open-source tool available at https://svn.mpl.ird.fr/echopen/MATECHO/ (last access: 15 February 2019) (login: userecho, password: echopen). Other MATLAB codes used in this work are “Layer” and “ComparEchoProfil” and are shared in the Supplement sections S1 and S2 of this paper. “Matecho” (Perrot et al., 2018) is an Open-Source Tool available at: https://svn.mpl.ird.fr/echopen/MATECHO/ (login: userecho, password: echopen).

Sample availability. The public cannot access our data because they belong to the partners who funded the oceanographic cruise.

Supplement. The supplement related to this article is available online at: https://doi.org/10.5194/os-16-65-2020-supplement.

Author contributions. ND set the methodology, analyzed data, and redacted the paper and the review. PB was cruise leader on the ECOAO sea survey, defined the sampling design, collected the data, defined the methodology, supervised the work and the review, and took charge of the acquisition of the financial support for the project leading to this publication. MT helped with data processing and analyses, paper redaction, and the review. YP developed the “Matecho” software tool and MATLAB code and contributed to the redaction and data collection. AS, AT, and SEA contributed to the PhD
supervision of ND. AM and CM helped with statistical analyses and OS performed the early PCA on CTD data.

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