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Diatoms as a paleoproductivity proxy in the NW Iberian coastal upwelling system (NE Atlantic)

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Abstract. The objective of the current work is to improve our understanding of how water column diatom's abundance and assemblage composition is seasonally transferred from the photic zone to seafloor sediments. To address this, we used a dataset derived from water column, sediment trap and surface sediment samples recovered in the NW Iberian coastal upwelling system.

Diatom fluxes ($2.2 (\pm 5.6) 10^6$ valves $m^{-2} d^{-1}$) represented the majority of the siliceous microorganisms sinking out from the photic zone during all studied years and showed seasonal variability. Contrasting results between water column and sediment trap diatom abundances were found during downwelling periods, as shown by the unexpectedly high diatom export signals when diatom-derived primary production achieved their minimum levels. They were principally related to surface sediment remobilization and intense Minho and Douro river discharge that constitute an additional source of particulate matter to the inner continental shelf. In fact, contributions of allochthonous particles to the sinking material were confirmed by the significant increase of both benthic and freshwater diatoms in the sediment trap assemblage.

In contrast, we found that most of the living diatom species blooming during highly productive upwelling periods were dissolved during sinking, and only those resistant to dissolution and the *Chaetoceros* and *Leptocylindrus* spp. resting spores were susceptible to being exported and buried. Fur-

thermore, *Chaetoceros* spp. dominate during spring–early summer, when persistent northerly winds lead to the upwelling of nutrient-rich waters on the shelf, while *Leptocylindrus* spp. appear associated with late-summer upwelling relaxation, characterized by water column stratification and nutrient depletion. These findings evidence that the contributions of these diatom genera to the sediment's total marine diatom assemblage should allow for the reconstruction of different past upwelling regimes.

1 Introduction

Diatoms are the most important primary producers in the ocean and play a key role in biogeochemical cycles through transferring organic carbon and biogenic silica from the surface layer to the seafloor sediments (Sancetta, 1989; Romero and Armand, 2010; Tréguer and De La Rocha, 2013). The preservation of their siliceous valves in marine sediment records has promoted their use as paleoproductivity indicators. However, reconstruction of primary production still suffers from diverse uncertainties, indicating that more studies are needed to accurately discern how particular environmental conditions regulate the diatom response and how diatom's ecological traits transfer primary production signals from the water column to the sediments through exported and buried

particles. In this regard, the analysis of data provided by sediment traps has contributed significantly to improving our knowledge of this topic, because the deployment of traps still is the best approach for monitoring downward diatom fluxes as a response to oceanographic and biological processes occurring in surface waters on a long-term basis.

Coastal upwelling systems are sites with major diatom-derived primary production where seasonality is often a noticeable feature (Walsh, 1991; Falkowski et al., 1998; Capone and Hutchins, 2013). Consequently, many studies focused on how primary production signal is exported through the water column in these highly productive coastal regions and were conducted through the analysis of downward diatom flux time series (Sancetta, 1995; Lange et al., 1998; Romero et al., 2002; Abrantes et al., 2002; Venrick et al., 2003; Onodera et al., 2005). In the NW Iberian margin, despite being the most productive upwelling region in Europe (Figueiras and Pazos, 1991; Nogueira and Figueiras, 2005; Espinoza-González et al., 2012), the use of diatoms as a productivity tracer to date was based on a direct comparison of the hydrographic conditions with surface sediment (Abrantes, 1988; Bao et al., 1997; Abrantes and Moita, 1999). From these studies it was concluded that *Chaetoceros* resting spores could be used as a good tracer of upwelling patterns, in particular the position of the upwelling front. Nevertheless, none of these studies provided information on the processes regulating seasonal diatom production and export from the photic zone to the seafloor sediments. In this context, the aim of this work is to go further in this topic by presenting the first analysis of the diatom community that combines water column, sediment trap and surface sediment samples recorded in this margin. Our results will provide relevant information regarding the use of fossil diatom assemblages as a primary production paleotracer in the highly productive NW Iberian coastal upwelling system.

2 Regional setting

Our study site (RAIA station) is located in the NW Iberian continental shelf off Cape Silleiro (42°05' N, 8°56' W; 75 m water depth; Fig. 1). During spring–summer, the NW Iberian coast is characterized by prevailing northerly winds that favor upwelling of cold and nutrient-rich subsurface Eastern North Atlantic Central Water (ENACW), resulting in high primary production (Fraga, 1981; Fiuza, 1984; Tenore et al., 1995; Figueiras et al., 2002). In contrast, southwesterly winds favor coastal downwelling during autumn–winter. Hydrographically, from October to January the region is generally affected by the northward advection of warm, saline and nutrient-poor waters by the Iberian Poleward Current (IPC) (Haynes and Barton, 1990; Castro et al., 1997; Relvas et al., 2007). Later on, usually between February and March, a decrease of temperature associated with winter cooling leads to a well-homogenized mixed layer of cold and

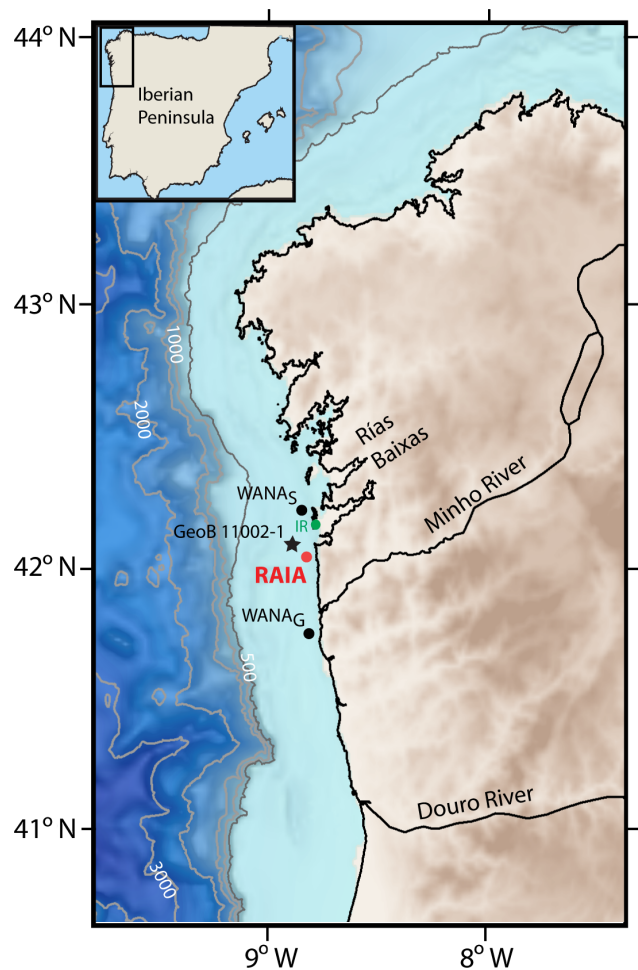


Figure 1. Map of the NW Iberian Peninsula continental margin showing the position of the mooring line (RAIA) site. WANA hindcast reanalysis points 3027034 (WANA_S, off Cape Silleiro) and 1044067 (WANA_G, off A Guarda) from which wave data were obtained, location of the irradiance Cíes station (IR) and position of the core-top sediment sample GeoB11002-1 are also shown.

nutrient-rich waters (Álvarez-Salgado et al., 2003; Castro et al., 2006). In addition, during downwelling seasons, the occurrence of southwesterly winds can generate moderate to extreme storms with wave heights >6 m, which have been simulated to produce high sediment remobilization (Vitorino et al., 2002; Jouanneau et al., 2002; Oberle et al., 2014). During these highly energetic periods, this region is also strongly influenced by the discharge of the Minho and Douro rivers (annual averages of 550 and 310 m³ s⁻¹, respectively), which is an important source of terrestrial sediment to the inner shelf. This mainly occurs during the winter months, when river inflow can reach 3850 m³ s⁻¹ for the Douro River and 1800 m³ s⁻¹ for the Minho River (Dias et al., 2002; Otero et al., 2010).

3 Material and methods

3.1 External forcing

Irradiance data were obtained from Cíes Islands meteorological station (IR; 42° 13' N, 8° 54' W; 25 m height) (Fig. 1) and are available through the Meteogalicia website (www.meteogalicia.es).

The upwelling index (UI), an estimate of the volume of upwelled water per kilometer of coast, was calculated according to Bakun's (1973) method:

$$UI = -((\rho_a C_D |V|)/(f \rho_{sw})) V_y, \quad (1)$$

where ρ_a is the density of the air (1.22 kg m^{-3}) at 15 °C, C_D is an empirical dimensionless drag coefficient (1.4×10^{-3}), f is the Coriolis parameter ($9.76 \times 10^{-5} \text{ s}^{-1}$) at 42° N, ρ_{sw} is the seawater density (1025 kg m^{-3}) and $|V|$ and V_y are the average daily module and northerly component of the geostrophic winds (m s^{-1}) centered at 42° N, 10° W, respectively. Positive values show the predominance of northerly winds that induces upwelling on the shelf and negative values indicate the presence of downwelling. Minho and Douro river discharge was obtained from https://github.com/PabloOtero/uptodate_rivers (Otero et al., 2010). Significant wave height data were based on WANA hindcast reanalysis of 3027034 (WANA_S, off Silleiro: 42° 15' N, 9° W) and 1044067 (WANA_G, off A Guarda: 41° 45' N, 9° W) (Fig. 1) points and supplied by Puertos del Estado (www.puertos.es).

3.2 Water column

RAIA station (75 m water depth) was visited monthly by R/V *Mytilus* from March 2009 to June 2012 except during the period December 2009–June 2010. Characterization of the water column was conducted by (i) CTD-SBE25 profiling and (ii) collection of discrete water column samples at 5 m water depth using a rosette sampler (10 L PVC Niskin bottles). Aliquots of these samples were used for inorganic nutrients, chlorophyll *a* (Chl *a*) and diatoms counting/species identification analysis.

Water column stability (0–35 m) was analyzed by using Brunt–Väisälä frequency parameter, $N^2 = [g/z] \ln(\rho_z/\rho_0)$ where g is the local acceleration of gravity, z is the water depth and ρ_z and ρ_0 are the bottom and surface density, respectively.

Inorganic nutrient concentrations were determined by segmented flow analysis with Alpkem autoanalyzers (Hansen and Grasshoff, 1983). The analytical errors were $\pm 0.05 \mu\text{mol kg}^{-1}$ for nitrate and silicate and $\pm 0.01 \mu\text{mol kg}^{-1}$ for phosphate. Final Chl *a* concentrations were determined by pigment extract fluorescence using a Turner Designs fluorometer calibrated with pure Chl *a* (see details in Zúñiga et al., 2016).

For diatom counting and identification a volume of 100 mL sample was used. The samples were preserved with

Lugol's iodine until microscopic observation. Depending on the water column, Chl *a* concentration volumes between 10 and 50 mL were deposited in composite sedimentation chambers for observation through an inverted microscope. The microorganisms were counted and identified to the species level, whenever possible, using the Utermöhl sedimentation method (Utermöhl, 1931, 1958). Centric diatom cells whose diameter did not allow for species identification were grouped as small centric diatoms. Only diatom species that appeared in more than one sample with a percentage higher than 2 % of the total abundance were considered for further analysis.

3.3 Sediment trap

At RAIA station an automated cylindrical–conical Technicap PPS 4/3 sediment trap (height/diameter ratio of 1.7 and a collecting area of 0.05 m^2) was deployed at 35 m water depth from March 2009 to June 2012. Sampling intervals ranged from 4 to 12 days. Unfortunately some data were lost due to technical problems and bad weather conditions (Zúñiga et al., 2016). Examination of CTD pressure data mounted 2 m below the trap showed that the mooring line tilting was less than 5° during 70 % of the time it was deployed. Only in exceptional hydrodynamic events that led to velocities higher than 25 cm s^{-1} was the mooring tilted 15–20°. Therefore, we assume the sediment trap was not affected by hydrodynamic biases. Sampling strategy and sample processing details are explained at length in Zúñiga et al. (2016).

Total mass flux was gravimetrically determined. Biogenic silica content was analyzed following Mortlock and Froelich (1989). The samples were treated with 2 M Na_2CO_3 for 5 h at 85 °C to extract the silica and then measure as dissolved silica by colorimetric reaction. Biogenic silica was converted from Si concentration after multiplying it by a factor of 2.4.

Sample preparation for diatom abundance and assemblage assessment was adapted from Abrantes et al. (2005). Depending on the recovered material one-fifth or two-fifth splits of the original samples were used, after HgCl_2 rinse by repeated settling in distilled water. Subsequently, organic matter and carbonates were removed by the addition of H_2O_2 (30 %) and HCl (10 %), respectively. Permanent slides were prepared using the evaporation-tray method of Battarbee (1973) and Norland optical adhesive (NOA61) as the mounting medium. Diatoms counting and species identification was performed at $1000\times$ ($10\times$ eyepieces and $100\times$ objectives), using a Nikon Eclipse E100 microscope equipped with differential interference contrast (DIC). One hundred randomly selected fields of view were counted in three replicate slides (Abrantes et al., 2005). The diatom flux was calculated as follows:

$$F = ((N)(A/a)(V)(S)(X))/D, \quad (2)$$

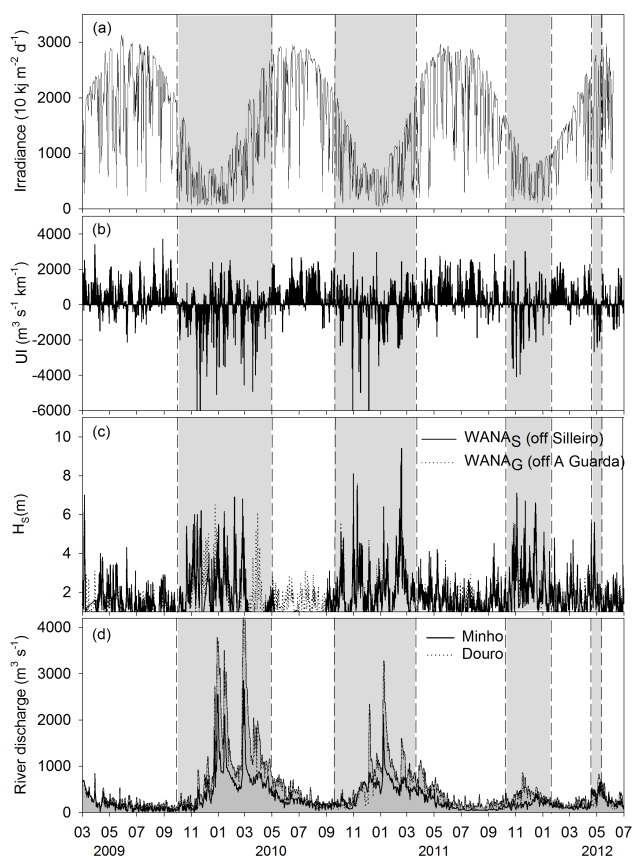


Figure 2. Temporal series of (a) total irradiance at Cíes Islands station (IR), (b) upwelling index (UI), (c) significant wave heights (H_s) obtained from the points off Silleiro and off A Guarda WANA and (d) Minho and Douro river discharge. Upwelling and downwelling periods are highlighted with white and grey bars, respectively, based on upwelling index and biogeochemical data presented in Zúñiga et al. (2016).

where the flux F is expressed as number of valves $\text{m}^{-2} \text{d}^{-1}$, N is the number of valves counted in 100 randomly selected fields of view, (a) represents the counted fraction of the total tray area (A), V is the dilution volume, S is the split fraction, X is the conversion factor from the collecting area to 1 m^2 , and D is the sampling interval in days for each sample.

Relative abundance of diatom taxa was determined following the counting procedures from Schrader and Gersonde (1978) and Abrantes (1988). For each sample, ca. 300 individuals were identified to the lowest taxonomic possible level, and raw counts were converted to percentage abundance. In samples containing low diatom abundances, the number of individuals identified was 100–200 (Fatela and Taborda, 2002). For this study only diatom species that appeared in more than one sample with a percentage higher than 2% of the total abundance were considered for further analysis.

3.4 Surface sediments

To evaluate the sedimentary record, we used one core-top sediment (0–1 cm) from box-core GeoB 11002-1 ($42^\circ 10' \text{ N}$, $8^\circ 58' \text{ W}$; 111 m) recovered near the RAIA position (Fig. 1). The sample was collected in August 2006, using a giant box corer during the GALIOMAR expedition (P342) on board of the R/V *Poseidon*. Sample cleaning and slides preparation was carried out following the methodology of Abrantes et al. (2005). Counting and identification procedures were the same as for sediment traps samples.

3.5 Statistical data analysis

Relationships between environmental variables and sediment trap diatom species relative abundances were evaluated with Pearson correlation coefficients and presented in Table 1 and 2, respectively.

In addition, the relationship between the relative abundance of the main groups of diatoms (freshwater (FW) diatoms, benthic diatoms, *Chaetoceros* resting spores, *Leptocylindrus* resting spores and *Paralia sulcata*) and the environmental variables were analyzed using the ordination technique canonical correspondence analysis (CCA) (vegan package, R-project; ter Braak, 1986; Oksanen et al., 2015). The water column environmental data were at first monthly interpolated and later averaged for the time interval recovered by each trap sample. Resulting data were subsequently integrated to 35 m where the sediment trap was moored. The multicollinearity of environmental variables was previously tested by Pearson correlations (Dormann et al., 2013) and checked after modeling using variance inflation factors (VIFs) applied to the CCA. Nine environmental variables were thus initially included in the ordination: irradiance, temperature, Brunt–Väisälä frequency parameter (N^2), Chl a , NO_3 , $\text{Si}(\text{OH})_4$, UI, Minho River flow, A Guarda wave height (H_s). Significant environmental variables were identified via a stepwise procedure, using permutation tests (999 permutations). After the selection of the significant variables, the model was tested a second time through a Monte Carlo global permutation test (999 permutations) to assess the significance of ordination axes.

The results of CCA were presented as ordination biplot diagram containing the explanatory variables plotted as arrows along with points for samples (dates) and species (main groups of diatoms). Using these diagrams, we were able to identify the relationships between species and between samples, as well as relationships of samples and species to environmental variables. Surface sediment sample location on the ordination plot was predicted using the CCA model produced with the sediment trap samples.

Table 1. Environmental variables matrix Pearson correlations. Irrad: irradiance; Temp: temperature; Sal: salinity; N^2 : Brunt–Väisälä frequency parameter; Chl a : chlorophyll a ; SPM: suspended particulate matter; POC: particulate organic carbon; UI: upwelling index; Minho: Minho River discharge; Waves: Significant wave height off A Guarda (WANA_G) station.

	Irrad	Temp	Sal	N^2	Chl a	SPM	POC	NO ₃	PO ₄	Si(OH) ₄	Oxygen	UI	Minho	Waves
Irrad	1.00	-0.394	0.487	-0.0827	0.528	-0.555	0.376	0.0167	0.0764	-0.642	0.148	0.487	-0.236	-0.357
Temp		1.00	-0.263	0.616	-0.196	0.332	0.184	-0.284	-0.195	0.351	-0.346	-0.456	0.0961	0.0557
Sal			1.00	-0.0509	0.228	-0.645	0.348	0.0839	0.263	-0.841	-0.137	0.518	-0.689	-0.27
N^2				1.00	-0.284	0.249	0.124	-0.266	0.0559	0.0825	-0.38	-0.266	-0.101	-0.0122
Chl a					1.00	-0.476	0.355	0.0608	-0.0003	-0.321	0.124	0.205	0.0433	-0.272
SPM						1.00	-0.104	0.153	0.0953	0.79	-0.223	-0.409	0.334	0.257
POC							1.00	0.24	0.341	-0.201	-0.39	0.0841	-0.246	-0.242
NO ₃								1.00	0.871	0.287	-0.677	0.182	-0.106	-0.252
PO ₄									1.00	0.137	-0.823	0.233	-0.274	-0.185
Si(OH) ₄										1.00	-0.297	-0.466	0.523	0.240
Oxygen											1.00	0.0592	0.213	0.204
UI												1.00	-0.0394	-0.116
Minho													1.00	0.139
Waves														1.00

Table 2. Pearson correlation matrix for the sediment trap diatom species relative abundances (%). FW: freshwater diatoms; ChaetoRS: *Chaetoceros* spp. resting spores; Cos.mar: *Coscinodiscus marginatus*; Cos.rad: *Coscinodiscus radiatus*; LeptoRS: *Leptocylindrus* spp. resting spores; Nav: *Navicula*; Nitzs.mar: *Nitzschia marina*; Par.sulc: *Paralia sulcata*; Pse.Pun: *Pseudo-nitzschia pungens*; Thal.ecc: *Thalassiosira eccentrica*; Thal.nitzs: *Thalassionema nitzschioides*.

	FW	Benthic	ChaetoRS	Cos.mar	Cos.rad	LeptoRS	Nav.spp	Nitzs.mar	Par.sulc	Pse.pun	Thal.ecc	Thal.nitzs
FW	1.00	0.482	-0.0085	-0.312	0.161	0.226	-0.126	0.176	0.293	0.0318	0.382	-0.0879
Benthic		1.00	-0.38	0.267	0.51	-0.377	0.2	0.0623	0.671	0.159	0.428	0.0777
ChaetoRS			1.00	-0.315	-0.388	-0.0163	-0.147	-0.389	-0.493	-0.318	-0.337	-0.214
Cos.mar				1.00	0.416	-0.137	0.517	-0.138	0.201	0.15	0.182	0.253
Cos.rad					1.00	0.283	0.267	0.0427	0.487	0.0773	0.261	0.202
LeptoRS						1.00	-0.224	-0.107	-0.495	-0.185	-0.0972	-0.173
Nav.spp							1.00	-0.0424	0.196	0.18	0.074	0.174
Nitzs.mar								1.00	0.173	0.547	0.338	0.079
Par.sulc									1.00	0.113	0.288	0.0748
Pse.pun										1.00	0.412	0.561
Thal.ecc											1.00	0.451
Thal.nitzs.												1.00

4 Results

4.1 Environmental conditions and water column characteristics

From October to April–May, the NW Iberian margin was characterized by the prevalence of low irradiance levels and southwesterly winds as shown by the negative UI values (Fig. 2a and b). During these periods the region was strongly affected by strong SW storms promoting wave heights higher than 4 m and intense Minho and Douro river discharge (Fig. 2c and d). As explained in detail in Zúñiga et al. (2016), hydrographically we can distinguish in a first phase the presence of the IPC (October–January), characterized by anomalously warm water (15–17 °C) with relatively low nutrient concentrations and Chl a (< 4 mg m⁻³) (Fig. 3). Later on, we differentiate the mixing period (from February to April–May), with temperatures of around 14 °C (due to winter cooling) and higher nutrient levels associated with in-

tense river discharge (Figs. 2d, 3a and b, Table 1). During downwelling periods, diatom abundances ranged from 5 to 135 cel mL⁻¹ with small centric diatoms accounting for the largest shares (52 ± 25 %) (Figs. 3c, 4b and Table 3). Only sporadically were *Navicula* spp. and *Paralia sulcata* highly abundant (Fig. 4). In contrast, from April–May to October, the margin was characterized by high irradiance levels and the upwelling of cold (< 14 °C) and nutrient-rich ENACW on the continental shelf that led to the development of Chl a maxima (Figs. 2a, b and 3). During these highly productive upwelling periods, diatom abundances were high (up to 7629 cel mL⁻¹) (Fig. 3c). The predominant diatoms in the water column alternated between *Chaetoceros* spp. at the onset of the upwelling season and *Leptocylindrus* spp. during the relaxation of the upwelling event when the water column became stratified (Fig. 4e and f). Other species frequently associated with upwelling favorable conditions (e.g., *Asterionellopsis glacialis*, *Detonula pumila* or *Guinardia deli-*

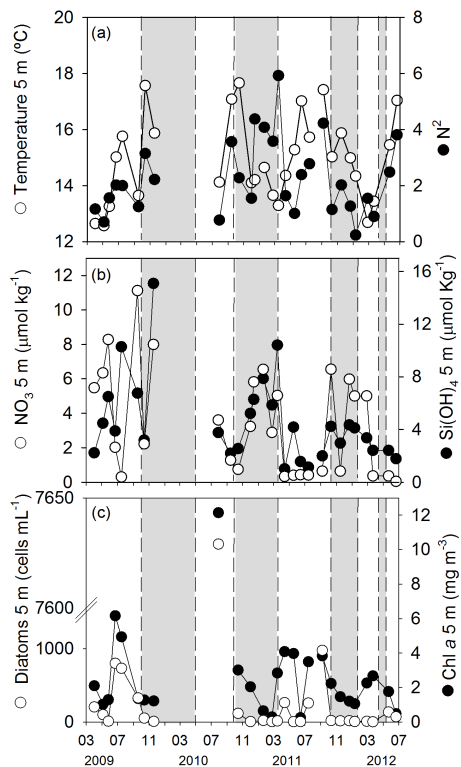


Figure 3. Temporal series of (a) temperature and water column integrated Brunt–Väisälä frequency (N^2), (b) nitrates (NO_3) and silicates (Si(OH)_4) content and (c) diatoms abundance and Chl *a* concentration, measured on samples recovered at 5 m water depth. Upwelling and downwelling periods are highlighted with white and grey bars, respectively, based on upwelling index and biogeochemical data presented in Zúñiga et al. (2016).

catula) appeared sporadically and with lower abundances (Fig. 4g, h and Table 3).

4.2 Sinking particulate material time series

The biogenic silica flux time series that ranged between 7 and $1001 \text{ mg m}^{-2} \text{ d}$, contributed from 2 to 10 % of the total material, and was closely followed by the siliceous organism fluxes calculated from microscopic counting (Fig. 5a, b and c). The contribution of diatoms to total siliceous microorganisms dominated throughout the entire period (Fig. 5c and e). Only during the 2012 upwelling season did silicoflagellates become relevant, achieving a relative abundance > 7 % (Fig. 5d). Maximum total diatom fluxes were registered under downwelling conditions (Fig. 5e). During these periods benthic and FW diatoms became relevant, contributing to the total diatom fluxes up to 24 and 17 %, respectively (Fig. 6). In contrast, during upwelling phases total diatom fluxes were relatively low, ranging around a mean seasonal value of $6 \pm 10 \times 10^5 \text{ valves m}^{-2} \text{ d}^{-1}$ (Fig. 5e and Table 3). During these periods, the diatom assemblage found in the trap samples were mainly composed of *Chaetoceros* spp. and

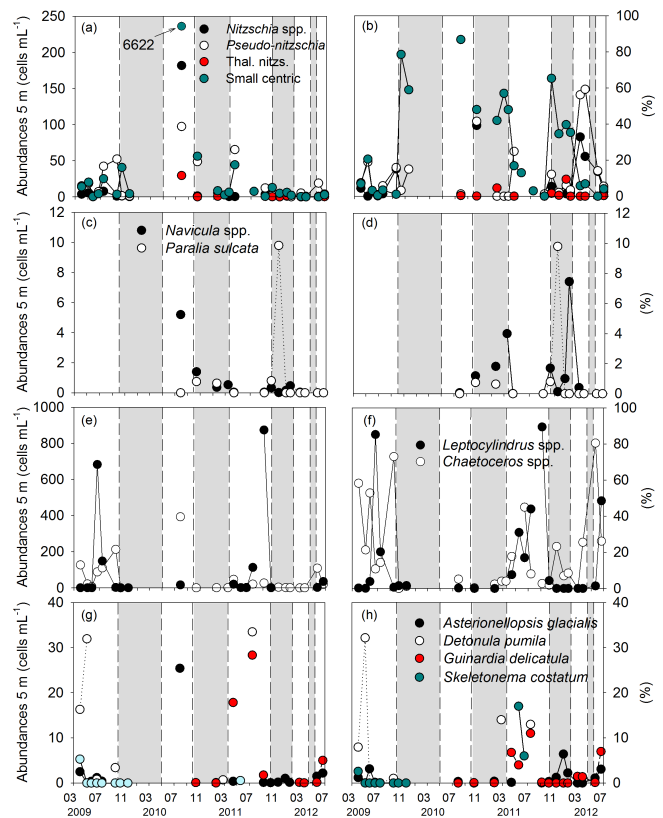


Figure 4. Time series of water column diatom abundances (a, c, e, g) and assemblages (b, d, f, h) from water column samples recovered at 5 m water depth. Water column diatom species were grouped in order to easily compare them with fossil diatom assemblage from the sediment trap samples. *Nitzschia* spp: *Nitzschia longissima*; *Pseudo-nitzschia* spp: *Pseudo-nitzschia* cf. *delicatissima* and *Pseudo-nitzschia* cf. *seriata*; Thal. nitzs: *Thalassiosira nitzschioides*; small centric includes centric diatom cells whose diameter did not allow for species identification and *Thalassiosira* spp. small; *Navicula* spp: *Navicula transitans* var. *derasa*; *Chaetoceros* (Ch.) spp: Ch. *curvisetus*, Ch. *socialis*, Ch. *didymus*, Ch. *lacinosus*, Ch. *decipiens* and small *Chaetoceros*; *Leptocylindrus* spp: *Leptocylindrus danicus*. Upwelling and downwelling periods are highlighted with white and grey bars, respectively, based on upwelling index and biogeochemical data presented in Zúñiga et al. (2016). The number and the corresponding arrow in Fig. 4a refer to small centric cells abundance in July 2010.

Leptocylindrus spp. resting spores, with mean contributions to total marine diatom fluxes of 46 and 20 %, respectively (Fig. 7 and Table 3).

4.3 Surface sediment samples

Diatom abundances in GeOB 11002-1 top sediment sample was $14 \times 10^5 \text{ valves g}^{-1}$. Marine diatom assemblage was dominated by resting spores of both *Chaetoceros* (33 %) and *Leptocylindrus* (37 %) spp., as well as *Paralia sulcata* (17 %) (Table 3). Benthic and FW diatoms had contributions < 4 %.

Table 3. Total abundance and relative contributions of the marine diatom assemblage preserved in the water column, sediment trap and core top sediment sample. SD: standard deviation. Nitzs: *Nitzschia*; Pseudo-nitzs: *Pseudo-nitzschia*; Thal. nitzs: *Thalassionema nitzschoides*; Nav: *Navicula*; Chaeto. and Lepto. spp: *Chaetoceros* and *Leptocylindrus* spp.; Aster. glac: *Asterionellopsis glacialis*; Deton. pum: *Detonula pumila*, Guin. del: *Guinardia delicatula*, Skelet. cost: *Skeletonema costatum*. Water column diatom species were grouped in order to easily compare them with fossil diatom assemblage from both the sediment trap and surface sediment samples. *Nitzschia* spp: *Nitzschia longissima*; *Pseudo-nitzschia* spp: *Pseudo-nitzschia* cf. *delicatissima* and *Pseudo nitzschia* cf. *seriata*; Thal. nitzs: *Thalassionema nitzschoides*; small centric includes centric diatom cells whose diameter did not allow for species identification and *Thalassiosira* spp. small; *Navicula* spp: *Navicula transiens* var. *derasa*; *Chaetoceros* (Chaeto.) spp: *Ch. curvisetus*, *Ch. socialis*, *Ch. didymus*, *Ch. lacinosus*, *Ch. decipiens* and small *Chaetoceros*; *Leptocylindrus* (Lepto) spp: *Leptocylindrus danicus*. Fossil diatom species from both the sediment trap and surface sediment samples were classified in three groups in order to compare them with water column diatom assemblage. *Nitzschia* spp: *Nitzschia marina*; *Pseudo-nitzschia* spp: *Pseudo-nitzschia pungens*; Thal. Nitzs: *Thalassionema nitzschoides*. Small centric includes *Coscinodiscus marginatus*, *Coscinodiscus radiatus* and *Thalassiosira eccentrica*. Chaeto. and Lepto. spp: *Chaetoceros* and *Leptocylindrus* spp. resting spores.

	Total	Nitzs. spp.	Pseudo-nitzs. spp.	Thal. nitzs.	Small centric	Nav. spp.	<i>Paralia sulcata</i>	Chaeto. spp.	Lepto. spp	Aster. glac.	Deton. pum.	Guin. del.	Skelet. cost.
WATER column													
Upwelling													
cel mL ⁻¹ (±SD)	717 (1869)	17 (50)	25 (30)	10 (17)	486 (1766)	1 (2)	0 (0)	72 (108)	126 (271)	3 (8)	17 (15)	11 (12)	1 (2)
% (±SD)		3(4)	16 (24)	0.1 (0.4)	20 (31)	0.1 (0.4)	0 (0)	27 (28)	26 (33)	1 (1)	3 (9)	2 (4)	2 (9)
Downwelling													
cel mL ⁻¹ (±SD)	34 (49)	2 (2)	12 (21)	1 (1)	10 (18)	0 (0)	1 (1)	1 (1)	1 (1)	1 (1)	1 (0)	0 (0)	0 (0)
% (±SD)		12 (12)	18 (28)	2 (4)	52 (25)	2 (4)	2 (4)	7 (10)	1 (3)	2 (3)	2 (7)	0 (0)	0 (0)
TRAP													
Upwelling													
10 ⁴ valves m ⁻² d ⁻¹ (±SD)	59 (97)	0.2 (0.1)	1.1 (5.8)	0.7 (1.6)	0.5 (0.7)	0.1 (0.3)	5.3 (13)	40.3 (87.2)	23.0 (49.9)	0 (0)	0 (0)	0 (0)	0 (0)
% (±SD)		2 (1)	6 (6)	2 (4)	2 (2)	1 (1)	14 (13)	46 (25)	20 (22)	0 (0)	0 (0)	0 (0)	0 (0)
Downwelling													
10 ⁴ valves m ⁻² d ⁻¹ (±SD)	216 (462)	0.1 (0.2)	0.1 (0.2)	0.9 (2.1)	1.3 (2.1)	0.3 (0.5)	31.2 (43.6)	25.6 (40.6)	43.9 (111)	0 (0)	0 (0)	0 (0)	0 (0)
% (±SD)		1 (1)	1 (1)	2 (2)	3 (2)	1 (1)	23 (12)	28 (19)	24 (19)	0 (0)	0 (0)	0 (0)	0 (0)
SEDIMENT													
GeoB 11002-1													
10 ⁴ valves g ⁻¹	142	0	0	2	3	0	24	47	52	0	0	0	0
%		0.2	0	2	2	0	17	33	37	0	0	0	0

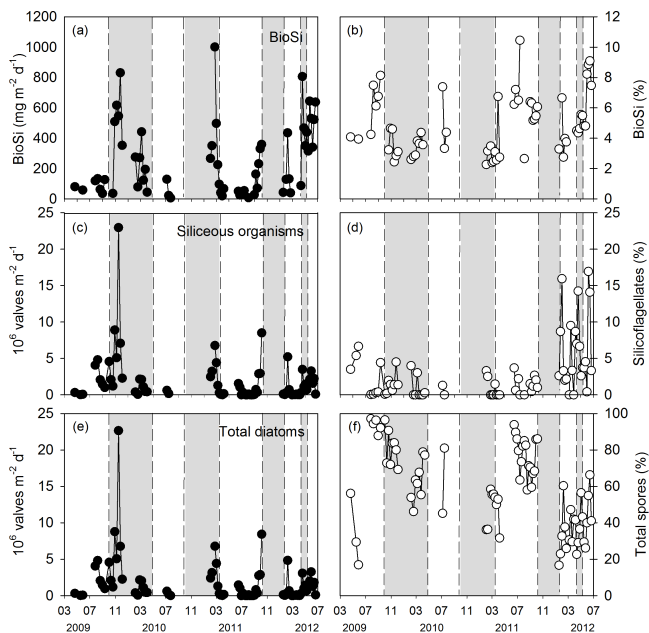


Figure 5. Time series of (a) biogenic silica (BioSi), (c) total siliceous organisms and (e) total diatom (including valves and resting spores) fluxes recorded with a PPS 4/3 sediment trap at RAIA station. Relative contribution of (b) biogenic silica to total mass flux, (d) silicoflagellates respect to total siliceous organisms and (f) resting spores to total diatoms flux is also presented. Upwelling and downwelling periods are highlighted with white and grey bars, respectively, based on upwelling index and biogeochemical data presented in Zúñiga et al. (2016).

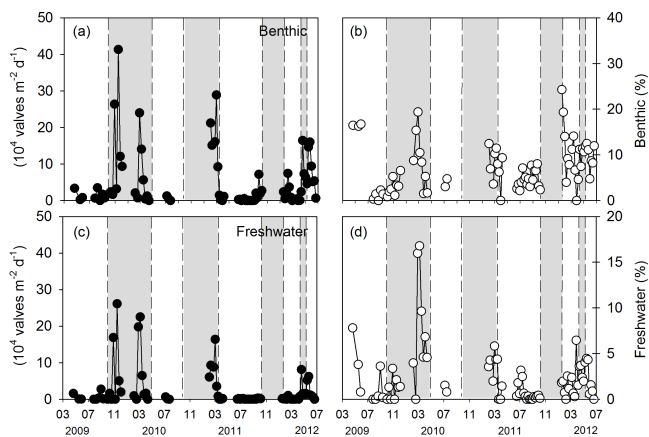


Figure 6. Time series of (a) benthic and (c) freshwater diatom fluxes (and relative contributions respect to total diatoms b, d) recorded with a PPS 4/3 sediment trap at RAIA station. Upwelling and downwelling periods are highlighted with white and grey bars, respectively, based on upwelling index and biogeochemical data presented in Zúñiga et al. (2016).

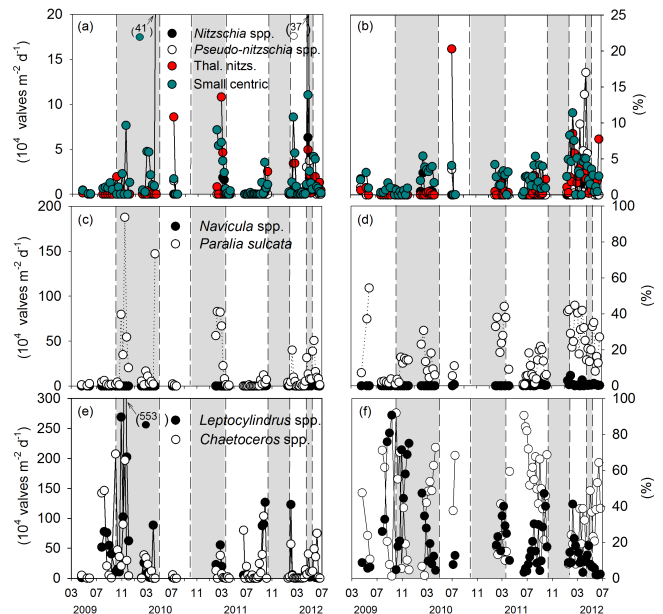


Figure 7. Time series of marine diatoms fluxes (a, c, e) and assemblages (relative contributions to total marine diatoms b, d, f) recorded with a PPS 4/3 sediment trap at RAIA station. Fossil diatom species has been classified in three groups in order to compare them with water column diatom assemblage. *Nitzschia* spp: *Nitzschia marina*; *Pseudo-nitzschia* spp: *Pseudo-nitzschia pungens*; Thal. Nitzs: *Thalassionema nitzschioides*. Small centric includes *Coscinodiscus marginatus*, *Coscinodiscus radiatus* and *Thalassiosira eccentrica*. Upwelling and downwelling periods are highlighted with white and grey bars, respectively, based on upwelling index and biogeochemical data presented in Zúñiga et al. (2016).

4.4 Relationships between sediment trap main diatom groups and environmental variables

CCA stepwise procedure identified five significant variables for the abundance of the main diatom groups (p value < 0.05), Minho River flow (Minho River), temperature (Temp), Chl a , NO_3 and $\text{Si}(\text{OH})_4$ (Fig. 8). The first two canonical axes explained 48.7 and 40.4 %, i.e., 89 % of the modeled inertia, and consequently only those two axes were considered. The CCA model with the five variables explained 46 % of the total inertia. The first canonical axis showed a positive gradient with Temp and Chl a , opposite to Minho River discharge. FW diatoms, benthic diatoms and *Paralia sulcata* (Parsul) were negatively positioned in the first canonical axis, thus indicating a positive relationship with the Minho River and a negative relationship with Temp and Chl a . The second canonical axis showed a negative gradient with NO_3 and $\text{Si}(\text{OH})_4$ and a negative relationship between these variables and *Chaetoceros* resting spores (ChaeRS). Conversely, *Leptocylindrus* resting spores (LepRS) were positively related with NO_3 and $\text{Si}(\text{OH})_4$. The temporal distribution of the sediment trap samples confirmed that FW diatoms, benthic diatoms and

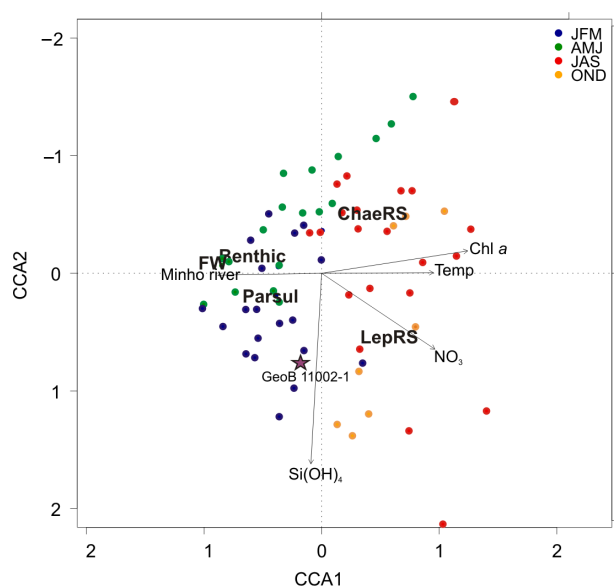


Figure 8. RDA biplot results of the canonical ordination (only significant variables shown) for main fossil sediment trap diatom groups (freshwater (FW) diatoms, benthic diatoms, *Paralia sulcata* (Parsul), *Chaetoceros* spp. spores (ChaRS) and *Lepidocylindrus* spp. spores (LepRS), and forward-selected environmental variables (chlorophyll *a* (Chl *a*), temperature (Temp), nitrates (NO_3) and silicates ($\text{Si}(\text{OH})_4$) and Minho River flow). JFM: January–February–March; AMJ: April–May–June; JAS: July–August–September; OND: October–November–December. Position of the predicted surface sediment sample GEOB 11002-1 is also shown.

Parsul occurred mainly during downwelling months, while ChaRS and LepRS were associated with upwelling periods (Fig. 8). In addition, this figure also identifies LepRS with late-summer periods. The position of the core top sample (GEOB 11001-2) on the CCA ordination reflected an integration between upwelling and downwelling seasonal processes (Fig. 8).

5 Discussion

Diatoms exported out from the photic zone ($2.2 (\pm 5.6) 10^6 \text{ valves m}^{-2} \text{ d}^{-1}$), similar to those registered in other coastal upwelling systems (Sancetta, 1995; Lange et al., 1998; Romero et al., 2002; Abrantes et al., 2002; Venrick et al., 2003; Onodera et al., 2005), showed contrasting results compared to diatom abundances on the surface waters (Figs. 3c, 5c and e). In fact, the a priori contradictory observation of maximum diatom fluxes during autumn–winter when irradiance conditions were unfavorable for phytoplankton growth and Chl *a* showed minimum levels (Figs. 2, 3c and 5e), can be explained by the input of allochthonous sources associated with the inner continental margin hydrodynamics (Zúñiga et al., 2016). Furthermore,

Zúñiga et al. (2016) have also showed how seasonal intensification of primary production promoted biogenic settling particles during spring–summer seasons, clarifying why diatom assemblages' dominant species recorded in the trap material were the same as in the water column (Figs. 4 and 7). With this in mind, our results confirm the major influence of both hydrodynamic and biogenic processes over the diatoms abundance, assemblage composition and export in this coastal upwelling system.

5.1 Sediment trap diatom assemblage as a tracer for allochthonous sources in sinking material

During highly hydrodynamic downwelling periods, higher wave heights, indicative of strong storms, co-occurred with maximum fluxes of benthic diatoms (Figs. 2c and 6a), whose natural habitat is the sediment interface. This finding, along with the fact that during these high-energy episodes lithogenic particle fluxes achieved their maximum levels (as shown in Zúñiga et al., 2016), may only be explained if strong storms resuspended surface sediments covering the Iberian continental shelf (Dias et al., 2002; Vitorino et al., 2002; Jouanneau et al., 2002; Oberle et al., 2014). Furthermore, stormy conditions were accompanied by intense Minho and Douro river discharge which impacted the water column thermohaline structure (Fig. 2c and d). The increase in FW diatoms associated with river runoff reinforce that those continental inputs are an additional source of terrestrial material to the inner continental shelf (Figs. 2d and 6c). Indeed, canonical analysis of sediment trap samples revealed a high correlation between benthic and FW diatoms, corroborating the co-occurrence of both resuspension processes and river discharge during downwelling periods (Fig. 8 and Table 2).

Additional evidence of resuspension resulted from the analysis of the marine diatom assemblage collected in the sediment trap. *Paralia sulcata* was sporadically found in the water column diatom assemblage during the 2009–2012 studied years (Fig. 4c and Table 3). This meroplanktonic and shadow species was, by contrast, common in sediments and contributed significantly to the trap diatom fluxes during downwelling phases (Fig. 7c and Table 3). All this indicates that this resistant-to-dissolution species can be easily resuspended from the sediments under highly hydrodynamic conditions (Bernárdez et al., 2010; Zúñiga et al., 2011). This is also supported by the positive relationship found between *Paralia sulcata* and benthic diatoms in the trap samples (Fig. 8 and Table 2).

Also of interest is the positive correlation between FW and benthic diatoms to *Thalassiosira eccentrica* (Table 2), a species which is known to occur in areas where nutrient input is continuous throughout the year, such as in areas influenced by river discharge (Moita, 1993; Abrantes and Moita, 1999).

5.2 Seasonal succession of diatom species during upwelling seasons: the imprint of the fossil diatom assemblage

During the studied period, the water column diatom community was modulated by the seasonality of environmental variables, with the highest abundances always recorded during upwelling periods, when irradiance and water column characteristics promote favorable conditions for diatom growth (Figs. 2 and 3). In this regard, a detailed analysis of the marine diatom assemblage as a whole revealed that most living diatom species linked to upwelling favorable conditions were either not present (e.g., *Asterionellopsis glacialis*, *Detonula pumila*, *Guinardia delicatula* and *Skeletonema costatum*) or appeared with a significantly lower contribution (e.g., *Nitzschia* spp., *Pseudo-nitzschia* spp. and small centric) in the diatom assemblages in both the sediment trap and the surface sediment samples (Table 3). This confirms that selective dissolution processes acted on thin-walled, less-silicified diatoms and, thus, only the robust and heavily silicified frustules and resting spores will be available to be exported and buried. Although this may lead to underestimation of primary production from fossil diatoms, analysis made over the sediment trap data demonstrates that the vertical sinking of the highly resistant *Chaetoceros* and *Leptocylindrus* spp. resting spores (positively positioned in CCA1) (Fig. 8) occurred in agreement with their dominance in the upper water column, and therefore both diatom genera are good sedimentary indicators of high primary production in the NW Iberian margin. This brings new relevant information to previous works carried out along this margin, which have only considered *Chaetoceros* spp. resting spores as a tracer of the coastal upwelling regime (Abrantes, 1988; Bao et al., 1997; Abrantes and Moita, 1999).

Furthermore, during the highly productive upwelling periods, we found how the environmentally controlled succession between *Chaetoceros* and *Leptocylindrus* spp. blooms (Figueiras and Rios, 1993; Escaravage et al., 1999; Casas et al., 1999; Nogueira and Figueiras, 2005) is clearly reflected in the export of their corresponding resting spores (Figs. 4, 7 and 8). Indeed, our sediment trap dataset reflected how the sinking fluxes of *Chaetoceros* spp. resting spores were mostly associated with the onset of the upwelling period, when irradiance conditions are favorable and persistent northerly winds lead to the upwelling of nutrient-rich subsurface ENACW waters on the shelf (Figs. 3, 4, 7 and 8). In contrast, *Leptocylindrus* spp. resting spore fluxes were significantly associated with late-summer and autumn when more frequent relaxation of winds promoted water column stratification and nutrient depletion (Figs. 3, 4, 7 and 8). In this regard, our finding linking environmentally controlled diatom blooms and resting spores' vertical export give us the opportunity to reinterpret previous studies published on this margin with regards upwelling-related paleosignals, where, despite not being discussed, a downcore alternation between

the resting spores accumulation rates of these two genera is shown (Bernárdez et al., 2008; Abrantes et al., 2011).

6 Conclusions

Even though the remobilization of bottom sediments by re-suspension processes may distort diatoms accumulation and preservation rates in the NW Iberian margin, our results prove that surface sediments from the inner continental shelf integrate both hydrodynamic and biological processes occurring in the upper water column (Fig. 8). This provides relevant information in relation to the use of fossil diatoms in marine sediment records for determining allochthonous sources and primary production paleosignals in this margin. Main observations may be summarize as follows: (i) high diatom fluxes during downwelling periods did not represent water column primary production signal but rather reflected highly energetic episodes that proved that resuspended sediment and river inputs were the main sources of diatoms to the sinking fluxes. This fact is also evidenced by the significant increase of both benthic and freshwater diatoms in the sediment trap assemblage. (ii) During highly productive upwelling periods, diatom's export signal, mainly represented by the highly resistant-to-dissolution *Chaetoceros* and *Leptocylindrus* spp. resting spores, mirrored diatom community structure in the upper water column, and thus both diatom genera may be considered as good sedimentary imprints of highly productive upwelling conditions. (iii) The seasonal succession as a response to particular environmental conditions in both the bloom and export of *Chaetoceros* and *Leptocylindrus* spp. postulates that contributions of these diatom genera to the total marine diatom assemblage in the sediment records should allow the identification of paleoceanographic conditions attributed to different patterns (onset versus relaxation) of the upwelling regime.

Appendix A

Table A1. List of diatom species found in both the RAIA sediment trap and surficial sediment sample (Geo B 11002). Species that appeared in more than one sediment trap sample with a percentage higher than 2 % of the total abundance are highlighted in bold. Ecology preferences: B is benthic; MP is meroplanktonic; P is planktonic; CO is coastal; O is open ocean; C is cosmopolitan; M is marine; MB is marine to brackish; BF is brackish to freshwater; BR is brackish; FW is freshwater.

Diatom species	Ecology	Sediment trap	Sediments
<i>Achnanthes brevipes</i> C. Agardh	B-MB	×	×
<i>Achnanthes</i> sp. (cf. FW)	B-FW	×	
<i>Actinocyclus curvatulus</i> Janisch	P-M-C	×	×
<i>Actinocyclus octonarius</i> Ehrenberg	P-M-C	×	
<i>Actinoptychus senarius</i> (Ehrenberg) Ehrenberg	MP-C	×	×
<i>Actinocyclus</i> sp.	×	×	
<i>Amphora gracilis</i> Ehrenberg	B-FW	×	
<i>Amphora marina</i> T. V. Desikachary and P. Prema	B-MB	×	
<i>Amphora</i> sp.	B	×	
<i>Anorthoneis excentrica</i> (Donkin) Grunow	B-M	×	
<i>Asteromphalus flabellatus</i> (Brébisson) Greville	P-M	×	
<i>Asteromphalus</i> sp.	P-M-O	×	
<i>Aulacoseira</i> cf. <i>granulata</i> (Ehrenberg) Simonsen	P-FW	×	×
<i>Aulacoseira</i> sp.	P-FW	×	×
<i>Azpeitia neocrenulata</i> (VanLandingham) (G. Fryxell and T. P. Watkins)	P-M	×	
<i>Azpeitia nodulifera</i> (A. Schmidt) G. A. Fryxell and P. A. Sims	P-M	×	
<i>Bacillaria paxillifera</i> (O. F. Müller) T. Marsson	B-P-M	×	
<i>Bacteriastrum hyalinum</i> Lauder	P-M	×	
<i>Caloneis</i> sp.	B-MB	×	
<i>Campylodiscus incertus</i> A. W. F. Schmidt	P-M	×	
<i>Campyloneis grevillei</i> (W. Smith) Grunow and Eulenstein	B-M	×	
<i>Campylosira cymbelliformis</i> Grunow ex Van Heurck	B-MB-CO	×	
<i>Catacombas gaillonii</i> (Bory) D. M. Williams and Round	B-M	×	
<i>Cerataulus smithii</i> Ralfs ex Pritchard	B-MB	×	
<i>Chaetoceros lorenzianus</i> Grunow	P-M-CO	×	
<i>Chaetoceros</i> sp.	P-M-CO	×	
<i>Chaetoceros</i> sp. (resting spores)	M-CO	×	×
<i>Cocconeis disculoides</i> (Hustedt) Stefano and Marino	B-M	×	
<i>Cocconeis guttata</i> Husted and Aleem	B-M	×	
<i>Cocconeis hoffmannii</i> Simonsen	B-M	×	
<i>Cocconeis neodiminuta</i> Krammer	B-FW	×	
<i>Cocconeis placentula</i> Ehrenberg	B-FW	×	×
<i>Cocconeis pseudomarginata</i> Gregory	B-M	×	
<i>Cocconeis scutellum</i> Ehrenberg	B-M	×	×
<i>Cocconeis speciosa</i> Gregory	B-M	×	
<i>Cocconeis stauroneiformis</i> (W. Smith) H. Okuno	B-M	×	
<i>Cocconeis</i> sp.	B	×	
<i>Coscinodiscus gigas</i> Ehrenberg	P-M	×	
<i>Coscinodiscus marginatus</i> Ehrenberg	P-M-OC	×	×
<i>Coscinodiscus</i> cf. <i>oculus-iridis</i> (Ehrenberg) Ehrenberg	P-M-OC	×	
<i>Coscinodiscus radiatus</i> Ehrenberg	P-M-OC	×	×
<i>Coscinodiscus</i> sp.	P-M	×	
<i>Ctenophora pulchella</i> (Ralfs ex Kützing)	P-BF	×	
<i>Cyclotella meneghiniana</i> Kützing	P-CO-BF	×	×
<i>Cyclotella plitvicensis</i> Husted	P-FW	×	
<i>Cyclotella stelligera</i> Cleve and Grunow in Van Heurck	P-FW	×	
<i>Cyclotella radiosa</i> (Grunow) Lemmermann	P-FW	×	
<i>Cyclotella</i> sp.	P-FW	×	
<i>Cyclostephanos</i> sp.	P	×	
<i>Cymbella affinis</i> Kützing	B-FW	×	×
<i>Cymbella</i> sp.	B	×	

Table A1. Continued.

Diatom species	Ecology	Sediment trap	Sediments
<i>Delphineis minutissima</i> (Husted) Simonsen	P-M-CO	×	×
<i>Delphineis surirella</i> (Ehrenberg) G. W. Andrews	P-M-CO	×	
<i>Detonula pumila</i> (Castracane) Gran	P-M-CO	×	
<i>Dimeregramma minor</i> (Gregory) Ralfs ex Pritchard	B-M	×	
<i>Diploneis</i> cf. <i>bombus</i> (Ehrenberg) Ehrenberg	B-MB	×	×
<i>Diploneis didymus</i> (Ehrenberg) Ehrenberg	B-MB-BFW	×	
<i>Diploneis smithii</i> (Brébisson) Cleve	B-MB-BFW	×	
<i>Diploneis</i> cf. <i>stroemii</i> Husted	B-M	×	
<i>Diploneis suborbicularis</i> (W. Gregory) Cleve	B-M	×	
<i>Diploneis weissflogii</i> (A. W. F. Schmidt) Cleve	B-M	×	
<i>Diploneis</i> sp.	B	×	
<i>Dytilum</i> sp.	P-M-CO	×	
<i>Encyonema</i> sp.	B-FW	×	
<i>Epithemia</i> sp.	B-BFW	×	
<i>Eunotia</i> cf. <i>pectinalis</i> (Kützing) Rabenhorst	B-FW	×	
<i>Eunotia praerupta</i> (Grunow)	B-FW	×	
<i>Eunotia</i> sp.	B-FW	×	
<i>Fragilariforma constricta</i> (Ehrenberg) D. M. Williams and Round	B-FW	×	
<i>Fragilaria crotonensis</i> (Kitton) Cleve and Möller	B-BFW	×	
<i>Fragilaria inflata</i> (Heiden) Hustedt	B-FW	×	
<i>Fragilaria investiens</i> (W. Smith) Clever-Euler	B-M	×	
<i>Fragilaria schulzii</i> Brockmann	B-M	×	
<i>Fragilaria</i> sp.	B-C	×	
<i>Fragilariforma virescens</i> (Ralfs) D. M. Williams and Round	B-P-FW	×	
<i>Gomphonema</i> sp.	B	×	×
<i>Gomphonema</i> cf. <i>acuminatum</i> (Ehrenberg)	B-BFW	×	
<i>Gomphonema</i> cf. <i>constrictum</i> (Ehrenberg)	B-FW	×	
<i>Gomphonema parvulum</i> (Kützing) Kützing	B-FW	×	
<i>Grammatophora angulosa</i> (Ehrenberg)	B-M	×	
<i>Grammatophora marina</i> (Lyngbye) Kützing	B-M	×	×
<i>Grammatophora oceanica</i> (Ehrenberg) Cleve	B-M	×	
<i>Grammatophora</i> sp.	B-M	×	
<i>Grammatophora serpentina</i> (Ehrenberg) Hartley	B-M	×	
<i>Haslea</i> sp.	B-P-M-MB	×	
<i>Hantzschia</i> sp.	B-M-MB	×	
<i>Hemidiscus cuneiformis</i> Wallich	P-M-O	×	
<i>Hemiaulus</i> sp.	P-M-CO	×	
<i>Hyalodiscus scoticus</i> (Kützing) Grunow	P-M-CO	×	
<i>Leptocylindrus</i> sp. (resting spores)	M-CO	×	×
<i>Licmophora abbreviata</i> (C. Agardh)	B-M	×	
<i>Licmophora</i> sp.	B-M	×	
<i>Luticola mutica</i> (Kützing) D. G. Mann	B-FW	×	
<i>Martyana martyi</i> (Héribaud-Joseph) Round	B-BR-FW	×	
<i>Melosira moniliformis</i> (O. F. Müller) C. Agardh	B-M	×	
<i>Melosira varians</i> C. Agardh	P-FW	×	
<i>Melosira westii</i> W. Smith	P-M-CO	×	×
<i>Melosira</i> sp.	B-P-M-BR-FW	×	×
<i>Navicula bacillum</i> Ehrenberg	B-P-M-CO	×	
<i>Navicula</i> cf. <i>cancellata</i> Donkin	B-P-M-CO	×	
<i>Navicula cincta</i> (Ehrenberg) Ralfs	B-FW	×	
<i>Navicula mutica</i> Kützing	B-FW	×	
<i>Navicula</i> cf. <i>pennata</i> A. Schmidt	B-P-M-CO	×	
<i>Navicula</i> sp.	B-P	×	
<i>Nitzschia angularis</i> W. Smith	P-M-CO	×	
<i>Nitzschia longissima</i> (Brébisson) Ralfs	P-M	×	

Table A1. Continued.

Diatom species	Ecology	Sediment trap	Sediments
<i>Nitzschia macilenta</i> W. Gregory	M-CO	×	
<i>Nitzschia marina</i> Grunow	P-M	×	×
<i>Nitzschia umbonata</i> (Ehrenberg) H. Lange-Bertalot	M-FW	×	
<i>Nitzschia</i> sp.	P-M-O	×	×
<i>Odontella aurita</i> (Lyngbye) C. Agardh	P-M-CO	×	
<i>Odontella longicuris</i> (Greville) M. A. Hoban	P-M-CO	×	
<i>Odontella</i> sp.	P-M	×	
<i>Opephora marina</i> (W. Gregory) Petit	B-M	×	
<i>Paralia sulcata</i> (Ehrenberg) Cleve	MP-CO	×	×
<i>Petronella humerosa</i> (Brébisson ex W. Smith) Stickle and D. G. Mann	B-M	×	
<i>Pinnularia borealis</i> Ehrenberg	B-FW	×	
<i>Pinnularia</i> sp.	B	×	
<i>Pleurosigma elongatum</i> W. Smith	P-BR	×	
<i>Pleurosigma normanii</i> Ralfs in Pritchard	P-M	×	
<i>Pleurosigma</i> sp.	P	×	
<i>Pleurosira laevis</i> (Ehrenberg) Compère	P-BR	×	
<i>Porosira glacialis</i> (Grunow) Jørgensen	P-BR	×	
<i>Podosira stelliger</i> (Bailey) Mann	P-M	×	×
<i>Proboscia alata</i> (Brightwell) Sundström	P-M	×	
<i>Psammodiscus nitidus</i> (Gregory) Round and Mann	MP-M-CO	×	
<i>Pseudo-nitzschia pungens</i> (Grunow ex Cleve) G. R. Hasle	P-M-C	×	
<i>Pseudo-nitzschia seriata</i> (Cleve) H. Peragallo	P-M	×	
<i>Rhaphoneis amphicerus</i> (Ehrenberg) Ehrenberg	B-M	×	
<i>Rhabdonema arcuatum</i> (Lyngbye) Kützing	B-M	×	
<i>Rhabdonema minutum</i> Kützing	B-M	×	
<i>Rhabdonema</i> sp.	B-M	×	
<i>Rhizosolenia bergonii</i> Peragallo	P-M-O	×	
<i>Rhizosolenia hebetata</i> (Bailey) Gran	P-M-O	×	
<i>Rhizosolenia</i> sp.	P-M-O	×	
<i>Rhoicosphenia abbreviata</i> (C. Agardh) Lange-Bertalot	B-BR-FW	×	
<i>Rhoicosphenia marina</i> (Kützing) M. Schmidt	B-M-CO	×	
<i>Roperia tessellata</i> (Roper) Grunow ex Pelletan	P-M-O	×	×
<i>Staurosirella pinnata</i> (Ehrenberg) D. M. Williams and Round	MP-FW	×	
<i>Staurosirella</i> sp.		×	
<i>Stellarima stellaris</i> (Roper) G. R. Hasle and P. A. Sims	P-M	×	
<i>Stephanodiscus astrea</i> (Ehrenberg) Grunow	P-C	×	
<i>Stephanodiscus</i> sp.	P-C	×	×
<i>Stephanopyxis turris</i> (Greville) Ralfs in Pritchard	P-M	×	
<i>Surirella</i> sp.	B-M-BR	×	
<i>Synedra</i> sp.		×	
<i>Ulnaria ulna</i> (Nitzsch) Compère	B-FW	×	×
<i>Tabellaria fenestrata</i> (Lyngbye) Kützing	B-FW	×	
<i>Tabellaria flocculosa</i> (Roth) Kützing	B-FW	×	
<i>Tabellaria</i> sp.		×	
<i>Tetracyclus glans</i> (Ehrenberg) F. W. Mills	B-FW	×	
<i>Thalassiosira eccentrica</i> (Ehrenberg) Cleve	P-M-O	×	×
<i>Thalassiosira</i> cf. <i>leptopus</i> (Grunow) Hasle and G. Fryxell	P-M-O	×	
<i>Thalassiosira lineata</i> Jousé	P-M-O	×	
<i>Thalassiosira</i> sp.	P	×	×
<i>Thalassionema nitzschioides</i> (Grunow) Mereschkowsky	P-M	×	×
<i>Toxarium undulatum</i> Bailey	P-M	×	
<i>Trachyneis aspera</i> (Ehrenberg) Cleve 1894	B-M-BR	×	×
<i>Triceratium favus</i> Ehrenberg	P-M	×	
<i>Tryblionella angustata</i> W. Smith	P-M	×	
<i>Tryblionella navicularis</i> (Brébisson) Ralfs	B-BR	×	

Competing interests. The authors declare that they have no conflict of interest.

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