

Modelling the effect of the tidal cycle on the high phytoplankton biomass area of Cape Trafalgar (SW Iberian Peninsula)

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ABSTRACT

Physical–biological interactions in the ocean are known to be crucial for understanding ecosystem processes. This is particularly relevant in the highly dynamic coastal regions, where the biogeochemical processes associated with higher-frequency perturbations such as tidal waves play a key role in primary production. In this study, we examine the influence of the tide-topography interaction on the high productivity area of Cape Trafalgar (NW limit of the Strait of Gibraltar, Iberian Peninsula) using a high-resolution ocean circulation model coupled to an ecosystem model. The obtained results highlight the relevance of the tidal cycle explaining the high phytoplankton biomass that characterises this region through an active and periodic forcing, resulting in a pulsating upwelling system. Our model shows that the interaction of the westward zonal component of the tidal current (uvel) with the submarine ridge (i.e., Barbate High) that characterises this region, which is perpendicular to the coast, results in the pumping of deep, cold, salty, and nutrient-rich waters to the well-illuminated subsurface waters, fuelling phytoplankton growth. At the same time, the interaction of the westward tidal current with the ridge leads to the development of a cyclonic eddy, which enables the redistribution of the upwelled waters over and to the east of Barbate High. The fortnightly tidal period has been identified as the most influential because (an effective) tidal-pumping process only takes place when the westward uvel is $\sim 0.42 \text{ m s}^{-1}$, a condition attained between ~ 3 days before and after the moment of maximum tidal flow during spring tides. Simultaneously, the energy and the associated horizontal and vertical mixing of the cyclonic gyre also vary with the tidal cycle, being stronger during spring tides. Both tidally driven processes, i.e., the cyclical upslope advection of deep nutrient-rich water and the influence of the cyclonic gyre, are the main mechanisms that lead to the development of a persistent phytoplankton-rich tongue over Barbate High. Consequently, Cape Trafalgar acts as a source of nutrient- and phytoplankton-rich waters to the surrounding waters.

1. Introduction

Covering 6% of the world's surface, coastal waters are a vital part of the environment and provide between 22 and 43% of global ecosystem services (Costanza et al., 2014). The highly dynamic coastal regions and the associated biogeochemical processes play a key role in phytoplankton growth, being the primary production between 3 to 5 times higher than in oceanic regions (Simpson and Sharples, 2012).

In many coastal regions, tides are an important physical forcing factor with a major impact on primary production (e.g., Pingree and Griffiths, 1978; Daly and Smith, 1993; Otto et al., 1990). The tidal cycle is responsible for a series of periodic fluctuations at different time scales. The release on internal tidal waves causes a horizontal displacement of water masses and subsequent maxima in tidal current speed, at the same time that the associated current shear results in internal turbulent

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mixing (e.g., Holloway et al., 2001; Rippeth and Inall, 2002; Dale et al., 2003). Thus, locally, tide-topography interaction can have a significant impact on biological production by fertilisation of illuminated surface waters and the resuspension of phytoplankton cells and other particles, concomitant to the consequent vertical mixing (Pingree et al., 1981; New and Pingree, 1990; Hu et al., 2008; Sharples et al., 2007; Palmer et al., 2013).

This process is especially relevant in the Strait of Gibraltar (south of the Iberian Peninsula), a shallow and narrow channel that separates the Atlantic Ocean and the Mediterranean Sea (Fig. 1). The Strait is characterised by (i) the semi-diurnal tidal cycle with a periodicity of 12 h and 25 min, which results in a horizontal displacement of water that generates an increase in the current speed and the associated vertical mixing every 6 h and 12 min; and (ii) the spring-neap tidal cycle with a periodicity of ~15 days, related to the apogee-perigee cycle (29.50-day periodicity) (García-Lafuente et al., 1990). The interaction of the tidal current with the Strait's sharp topography generates undulatory processes (e.g., internal waves) that modify the nutrient budget by recirculation from deeper to shallower layers (e.g., Macías et al., 2007b; Ramírez-Romero et al., 2012; Sánchez-Garrido et al., 2015). However, due to the hydrodynamics of this region being characterised by the superficial inflow of Atlantic water towards the Mediterranean Sea, and by the short-residence time of the mixed waters within the channel (Macías et al., 2007a; Bruno et al., 2013), the biological impact of this tide-topography interaction results in a 40% increase of productivity, not in the Strait itself, but in the adjacent Alborán Sea (western-most basin of the Mediterranean Sea) (Sánchez-Garrido et al., 2015). Thus, although the Strait is characterised by low productivity due to the surface circulation pattern, this is not the case in the coastal Cape Trafalgar region located on its north-western limit (Fig. 1b).

Cape Trafalgar has been highlighted in several studies as a hotspot of high chlorophyll concentration (e.g., Prieto et al., 1999; Echevarría et al., 2002; García et al., 2002; Navarro and Ruiz, 2006), with the along-shore tide-topography interaction being suggested as the main responsible process (Vargas-Yáñez et al., 2002; Sala et al., 2018). A distinctive feature of Cape Trafalgar is a submarine ridge, known as Barbate High (Fig. 1b), that extends offshore perpendicular to the coast. Vargas-Yáñez et al. (2002), using a 2D-model, studied the capacity of the tide-topography interaction to reproduce a situation similar to the pool off Trafalgar. Although their model was very simple, it captured the vertical excursions of heat and nutrients produced by the tide-topography interaction, cooling, and fertilising areas around the topographic accident. Later, Sala et al. (2018) examined the chlorophyll-*a* (chl-*a*) dynamics in this region using 10 years of ocean colour images (MERIS sensor, 300 m full resolution). The authors found that Cape Trafalgar is characterised by a quasi-permanent high chl-*a* concentration, with an average value of 1.44 mg m⁻³ throughout the whole studied period. A Wavelet Power Spectrum analysis showed that the temporal variability of chl-*a* concentration in Trafalgar did not show any clear pattern of variability. However, when the covariation between the zonal component of the current speed (uvel) and the chl-*a* concentration was analysed with a Wavelet Coherence analysis, it was observed a strong covariation at the fortnightly tidal cycle. In comparison, the adjacent platform of the Gulf of Cádiz showed an average chl-*a* concentration of 0.79 mg m⁻³ with a seasonal variability (Sala et al., 2018). Furthermore, Bolado-Penagos et al. (2020) analysing the trajectory of several passive drifters released close to Cape Trafalgar, together with performing a series of hydrodynamic simulations, identified the formation of a cyclonic eddy on the eastern flank of Trafalgar. This cyclonic eddy increases the residence time of the upwelled nutrient-rich waters in the region, favouring the high chl-*a* concentration that characterises this biomass hotspot. Nevertheless, although the study by Sala et al. (2018) identified a strong covariation of chl-*a* concentration with the fortnightly tidal cycle, the study was unable to analyse in detail the influence of the tidal cycle on chl-*a* dynamics, due to an inadequate sampling resolution. Furthermore,

satellite ocean colour imagery only shows the most superficial layer, making it impossible to analyse properly the influence of the tidal cycle on the phytoplankton community for the whole water column.

The correct description of physical processes and their coupling with phytoplankton growth is a complex issue in regions highly influenced by tidal forcing. On the one hand, as tidal currents become more intense, they will lead to a more active pumping of deep saltier water to shallow areas, with a nutrient enrichment of well-illuminated (i.e., euphotic) layers, and subsequent growth of primary producers with a given temporal delay of the order of days. On the other hand, the intensification of the tidal current leads to a more active horizontal mixing and dispersal, thus decreasing nutrient concentration or phytoplankton accumulation. These are two opposing mechanisms affecting the phytoplankton community that are driven by the same single process: tidal forcing. Thus, using a modelling approach is probably the best way to understand the dynamics of Cape Trafalgar, which is of great importance as a source of phytoplankton biomass in the surrounding waters. This study aims at performing accurate simulations of the study area to evaluate how the upwelling process mediated by the tide-topography interaction can affect the phytoplankton biomass in the area of Cape Trafalgar. Furthermore, our approach will allow us to study how the seasonal variability that governs the adjacent Gulf of Cádiz may influence our region of interest. For these two purposes, we have implemented a high-resolution circulation model that computes the main tidal constituents, coupled to an ecosystem model.

2. Methodology

2.1. Circulation model

The physical model used in this study is the Massachusetts Institute of Technology general circulation model (MITgcm; Marshall et al., 1997a,b). The MITgcm solves the Boussinesq form of the Navier-Stokes equations for an incompressible fluid on a staggered C-grid, with level vertical coordinates, and partial cell representation of the bottom topography. A KPP (K-Profile Parametrisation) scheme parametrises the sub-grid-scale vertical diffusion by turbulence (Large et al., 1994). Temperature, salinity, and biochemical tracers are advected with a third-order direct-space-time scheme with a flux limiter to prevent negative values in the solutions (Hundsdorfer and Trompert, 1994). This numerical scheme is stable and sufficiently diffusive without explicit horizontal diffusion. The model configuration is similar to the one applied by Sánchez-Garrido et al. (2015), which uses a curvilinear grid that is squeezed into the Strait of Gibraltar and the Alborán Sea enhancing the resolution of both areas (Fig. 1a). With a horizontal high resolution of ~0.50 km within the Strait, small-scale topographic features and processes are well captured, such as internal hydraulic jumps with the associated mixing and propagating internal waves. In the vertical dimension, there are 46 depth levels with increasing cell size from the surface to the bottom.

Initial and boundary conditions were derived from the Iberia-Biscay-Ireland (IBI) Regional Seas reanalysis product from Copernicus Marine Environment Monitoring Service (CMEMS, <http://marine.copernicus.eu/>; Sotillo et al., 2015). Daily temperature, salinity and horizontal velocity fields were computed for the model lateral forcing. At the sea surface, the model is driven by wind stress, heat fluxes, freshwater flux from precipitation, and downward short-wave and long-wave radiation. All these forcings were obtained from ERA-Interim and provided by the European Centre for Medium-range Weather Forecasts (ECMWF, <https://www.ecmwf.int/>; Dee et al., 2011). The atmospheric data has a time resolution of 6 h and a spatial resolution of 0.125 degrees. The forcing fields for open boundary and atmospheric conditions correspond to the year 2010. We decided not to use a climatological year (i.e., the average of several selected years) because averaging smooths out daily fields and would result in an underestimation of submesoscale processes (Oguz et al., 2017). Latent and sensible

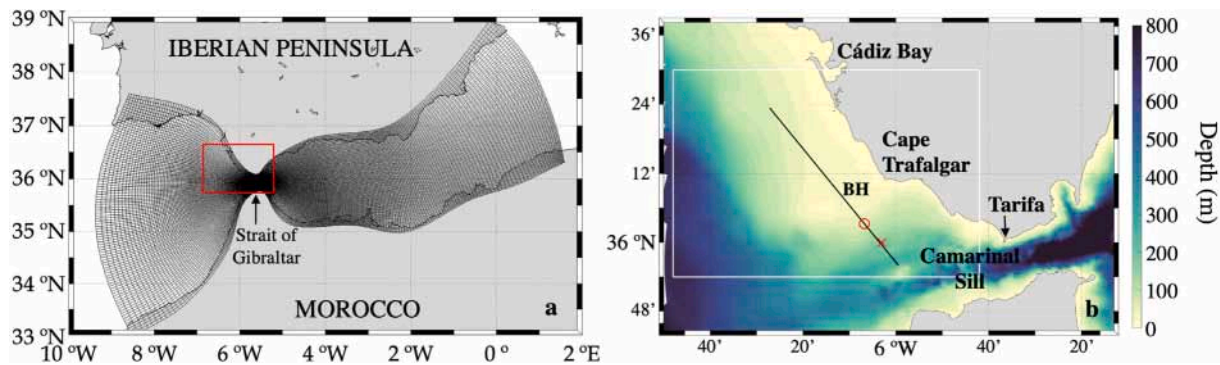


Fig. 1. (a) Model domain and computational grid. The red rectangle encloses (b) a zoom of the region of interest. The spatial and temporal evolution of different variables (i.e., salinity, nitrate concentration and total phytoplankton biomass) are analysed at the Trafalgar region (enclosed by the white box), along the black line parallel to the coast, and at the point represented by the red circle. The red cross indicates the mooring position. BH: Barbate High. More details are in Section 3.3.

heat fluxes are interactively calculated by the model using standard bulk formulas. Finally, tidal forcing is incorporated by prescribing time-dependent barotropic velocities across the open boundaries in association with the main eight tidal constituents in the region: the diurnal constituents K1, O1, P1 and Q1, with periods of 23.93 h, 25.82 h, 24.07 h, and 26.87 h, respectively; and the semi-diurnal constituents M2, S2, N2, and K2, with periods of 12.42 h, 12.00 h, 12.66 h, and 11.97 h, respectively (García-Lafuente et al., 1990). This forcing yields a spring-neap tide cycle every 29.50 days, with the tidal range fluctuating from spring tide (maximum) to neap tide (minimum) and back to the spring tide. Moreover, the tidal range and the timing change daily, shifting by about 50 min per day (Ippen, 1966). Zero-flux and non-slip conditions are applied to all solid boundaries (i.e., bathymetry) together with a nonlinear bottom drag at the seafloor. The integration time step is 15 s. To obtain a stable (repeated seasonal cycle) solution of the circulation model, the simulation was run for 3 years. Two years of spin-up were required to stabilise the numerical model and therefore they were excluded from the analysis.

The model solution was evaluated against different data sources. Modelled Sea Surface Temperature (SST) was compared to satellite-retrieved SST observations (reprocessed Pathfinder V5.3; CMEMS). The model tends to overestimate slightly the annual mean but otherwise, it captures well the spatial-temporal distribution, showing colder waters in coastal regions (due to upwelling) and warmer waters in open ocean regions for both the Gulf of Cádiz and the Alborán Sea (see Fig. A.1). We also compared temperature-salinity (TS) diagrams using modelled outputs versus *in situ* observations from the Medatlas II dataset (<http://www.ifremer.fr/medar/>). The main water masses are well reproduced by our regional model as inferred from the TS diagrams (Fig. A.2). Further details can be found in the supplementary material.

Finally, thermistors and Acoustic Doppler Current Profiler (ADCP) data collected in the proximity of Cape Trafalgar from September 3rd to October 10th, 2015 (MEGAN Project; Bolado-Penagos et al., 2020), were compared with model outputs to evaluate the tidal variability (Fig. 2). The mooring was bottom-mounted at 6.05°W–36.00°N (red cross in Fig. 1b) at a bottom depth of 200 m. At this emplacement was deployed a thermistor chain of 43 thermistors disposed 3-m apart from each other, from 10 to 136 m depth and programmed with a sample interval of 1 min. Additionally, it was also deployed a 75-kHz Teledyne RD Instrument (RDI) ADCP, with a sampling interval of 3 min and a 4-m bin size covering between 40 and 140 m of the water column. Because the model outputs represent a climatological year, to compare the observed and modelled data it was identified the closest tidal moment to the observations. Model data showed a temperature overestimation of 1.21 ± 0.64 °C (Fig. 2a). Meanwhile, the current velocity data, *uvel* and *vvel* (mean meridional component of the current speed), range between very similar ranges. However, modelled *uvel* seems to slightly overestimate the observations (Fig. 2b), while *vvel*

seems to slightly underestimate them (Fig. 2c). Both tidal components, semi-diurnal and fortnightly, are well captured by the model allowing us to pursue the aim of this study.

2.2. Ecosystem model

We used a simplified configuration of the Darwin model (version 1) that simulates the planktonic ecosystem (Follows et al., 2007; Dutkiewicz et al., 2009; Vallina et al., 2014a,b), being natively coupled to the MITgcm physical ocean model described above. Four phytoplankton functional groups are defined based on nutrient requirements and uptake strategies, with a transition from gleaners to opportunists: *Prochlorococcus*, *Synechococcus*, flagellates and diatoms (see Figure 2a in Vallina et al., 2017). All groups are limited by phosphate (PO₄), nitrate (NO₃), ammonium (NH₄) and iron, while diatoms are also limited by silicate (SiO₄). The model defines one generic zooplankton herbivore that feeds upon all phytoplankton species (killing-the-winner behaviour), and one generic zooplankton carnivore feeding on the zooplankton herbivore acting as a closure trophic level for the food web. For a detailed description of the ecological model, the reader is referred to Vallina et al. (2014a,b). The model parameters are given in Table B.1.

Initial conditions for all the ecological tracers were based on a global model output from Vallina et al. (2014a). Phytoplankton initial conditions are the same for the four functional groups, applying the same biomass distribution. The same approach is used for zooplankton initial conditions. Boundary conditions were only prescribed for nutrients (PO₄, NO₃, NH₄, SiO₄) and phytoplankton biomass. Daily fields for the prescribed boundary conditions were derived from the IBI Biogeochemical Analysis and Forecast product from CMEMS (IBI-BIO). Phytoplankton boundary conditions were computed from the total biomass of the IBI-BIO model, using the same boundary condition for the four phytoplankton groups. These oceanic boundary conditions are only open for the prescribed tracers, so that those state variables with prescribed values at the boundary can leave the model domain through them. For those tracers for which there are not enough data at the boundary conditions, homogeneous Neumann boundary conditions (zero-flux) are applied (Sánchez-Garrido et al., 2015). Finally, at the sea surface, the model is forced with daily photosynthetically active radiation (obtained from ERA-Interim) and daily iron dust deposition (computed from modelled monthly data from Mahowald et al., 2003).

Due to the high computational cost of the hydrodynamic model (integration time-step of 15 s), the coupled biophysical model MITgcm + Darwin was run in offline mode (integration time-step of 240 s) using as physical forcing (inputs) the previously saved simulation (outputs) with the MITgcm hydrodynamic model on an hourly temporal resolution of temperature, salinity, vertical mixing, and velocity fields. Such

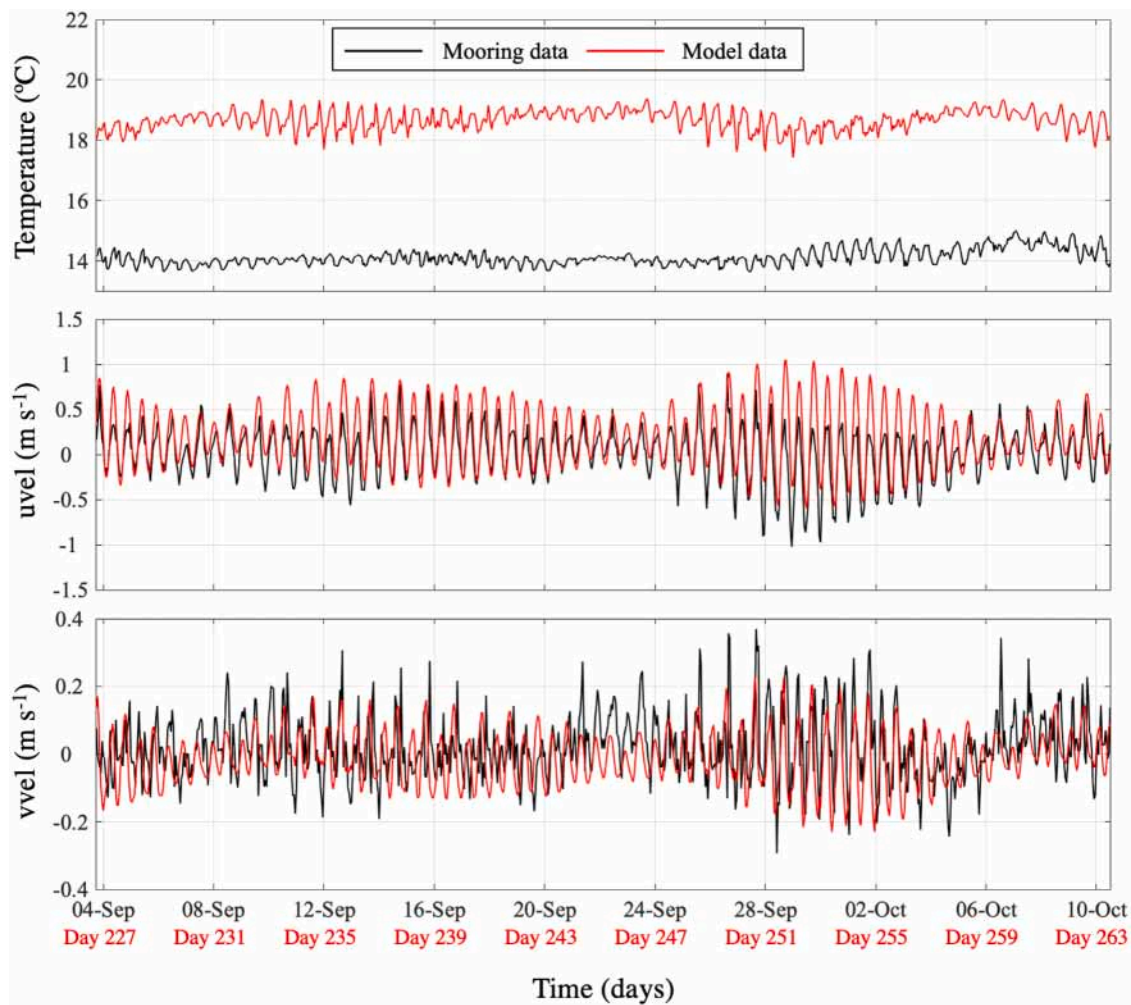


Fig. 2. Temporal evolution of (a) mean temperature ($^{\circ}\text{C}$), (b) mean zonal component of the current speed (u_{vel} ; m s^{-1}), and (c) mean meridional component of the current speed (v_{vel} ; m s^{-1}) for observed (black lines) and modelled (red lines) data. All time series were computed for the whole water column at 6.05°W – 36.06°N (red cross in Fig. 1b). The x-axis labels show in black the sampled days of the mooring and the corresponding days of the model in red.

a high temporal resolution (hourly) for the environmental variables was necessary to capture the tidally driven processes. The coupled biophysical model was run for 5 years, with 4 years of spin-up required to reach (quasi) steady-state seasonal cycles that were excluded from the analysis. Despite the relative complexity of our plankton model configuration and considering the aims of the study, we will only focus our analyses on the total phytoplankton biomass.

The seasonal variability of the ecosystem model was evaluated with satellite chl-*a* (mg m^{-3}) data, a proxy for the surface phytoplankton biomass. Ocean colour images for the study area were downloaded from the CMEMS (from 1998 to 2016). To enable this comparison, the model total phytoplankton biomass (mmol P m^{-3}) was first converted to carbon by applying the canonical Redfield ratio C:P = 106:1 (mol mol^{-1}), and then converted to chl-*a* concentration by applying a variable C:Chl ratio calculated as a function of light levels following Lefèvre et al. (2002) (see supplementary material for more details). Fig. 3 shows the comparison between the seasonal climatology derived from remote sensing images and the model-derived seasonal average of the simulated year for surface chl-*a*. The modelled chl-*a* distribution reflects well the observed oligotrophic nature of the Western Alborán Gyre. Furthermore, the model also captures the meridional gradient of chl-*a* with higher values along the northern coast of the study area, as well as the decrease west-to-east of chl-*a* concentration in the Alborán Sea, being the north-western region the richest area (e.g., Sarhan et al., 2000; Macías et al., 2008). The underestimation of chl-*a* concentration

by our simulations over the northern coastal area of the Gulf of Cádiz may be attributed to the fact that the model does not prescribe nutrient inputs from river discharges (e.g., Guadalquivir River; Huertas et al., 2005; Navarro et al., 2006).

2.3. Wavelet analysis

Wavelet analysis is a powerful tool that allows a local time-scale decomposition of the signal in both time and frequency domains simultaneously (Lau and Weng, 1995; Torrence and Compo, 1998; Klvana et al., 2004; Cazelles et al., 2008). This tool is especially relevant for the analysis of non-stationary systems (see Cazelles et al. (2008) for a review). Here, wavelet analysis is used to disentangle the temporal patterns of several state variables of the MITgcm + Darwin model, both in time and frequency domains. In particular, we use it to identify the dominant periodicities of salinity, temperature, NO_3 and phytoplankton biomass. We used the MatLab toolbox developed by Cazelles et al. (2008), applying the Morlet wavelet that provides a good balance between time and frequency localisation (Grinsted et al., 2004). In a time–frequency plane, the wavelet power spectrum represents the relative importance of frequencies at each time step. Values closer to the edge of the time series, delimited by the cone of influence, were rejected to avoid false periodic events (Torrence and Compo, 1998). For each period, the average wavelet power spectrum was calculated, identifying the averaged variance contained in all wavelet coefficients

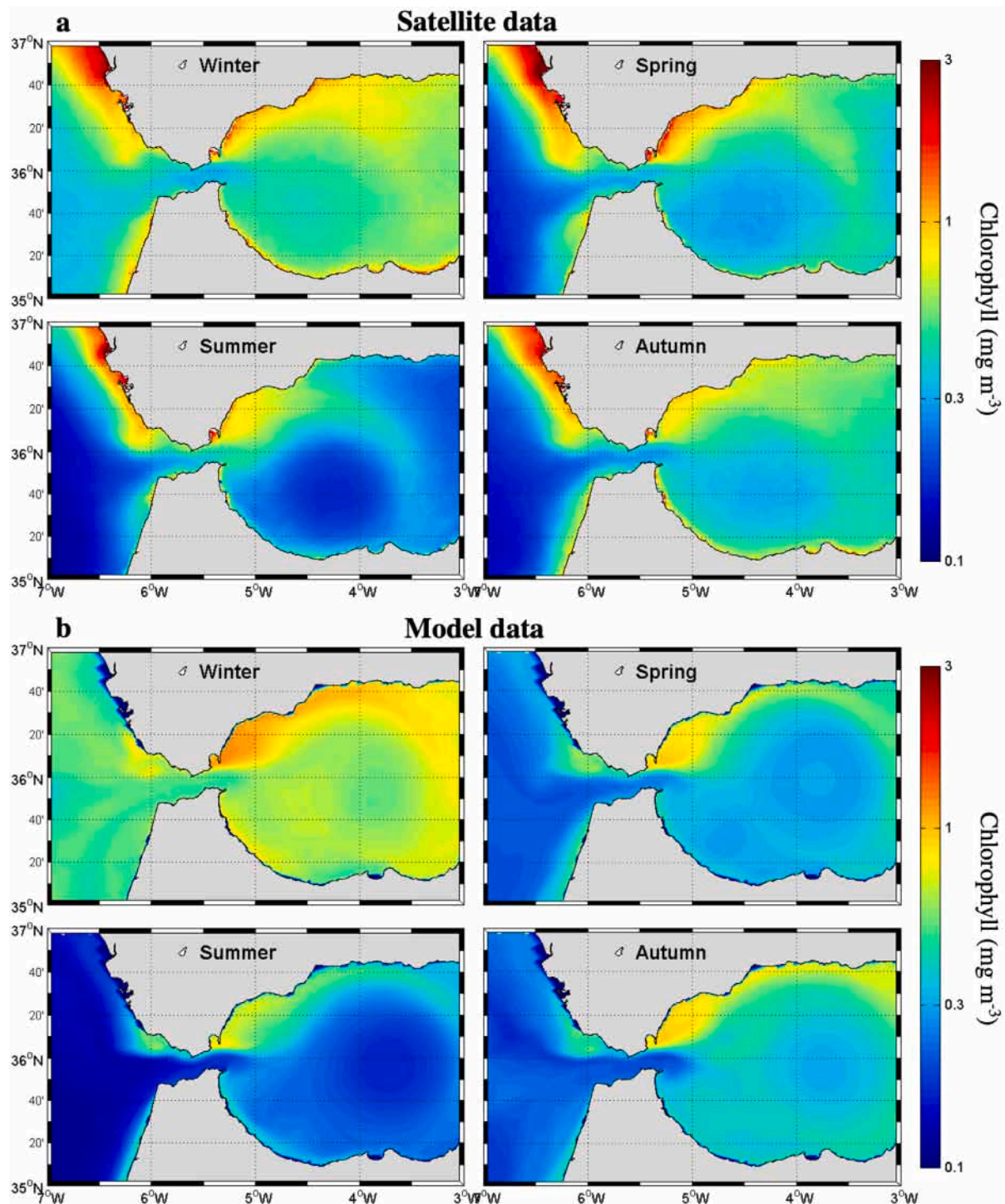


Fig. 3. (a) Seasonal climatology of surface chlorophyll (mg m^{-3}) derived from remote sensing images corresponding to the period 1998–2016. (b) Model-derived seasonal climatology of surface chlorophyll (mg m^{-3}) for the 5th year of simulation.

of the same frequency. Furthermore, a 5% significance level was determined for WPS through a bootstrapping scheme using a hidden Markov model, to assess whether the wavelet-based quantities were due to just random processes (Cazelles and Stone, 2003).

Of particular interest is the fact that wavelet analysis allows the estimation of the coupling between driver and response variables, as well as the level of synchrony between them (Keitt and Fischer, 2006; Ménard et al., 2007; Buttay et al., 2017). The wavelet coherence

analysis (WCo) was applied to analyse the synchrony between the zonal component of the current velocity (uvel; driver variable) and several model state variables: salinity, NO_3 concentration and phytoplankton biomass (response variables). Once identified the time scale at which the two non-stationary time series were locally linearly correlated, their phases, for this time scale, were extracted and compared (Cazelles et al., 2008). The phase difference between signals indicates whether or not they are in synchrony; and if they are not, the time lag between them.

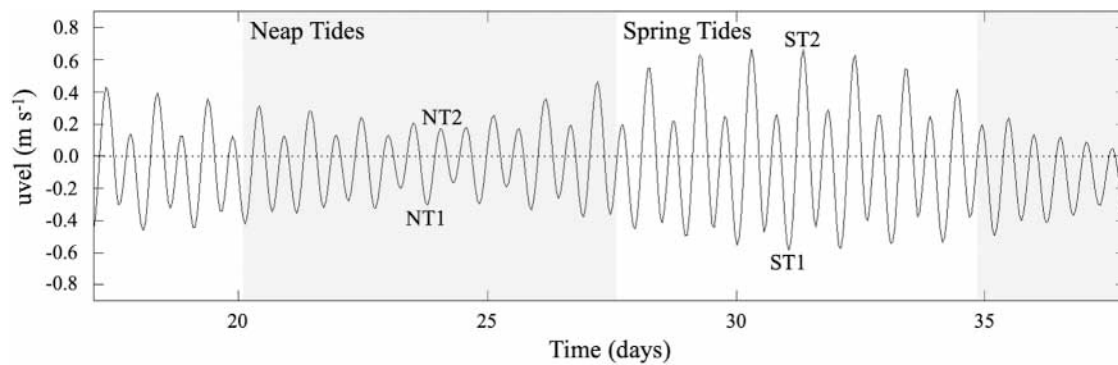


Fig. 4. Temporal evolution of the mean zonal component of the current velocity (u_{vel} ; $m\ s^{-1}$), computed for the whole water column at $6.11^{\circ}W$ and $36.06^{\circ}N$ (red circle in Fig. 1b), between days 17 and 38 of the simulated year. The black dashed line represents the zero reference, which delimits positive (negative) values that indicate flow driven eastward (westward), towards the Alborán Sea (Gulf of Cádiz). Points NT1 and NT2 indicate the examined neap tidal moments. Points ST1 and ST2 indicate the examined spring tidal moments. Grey background identifies neap tidal periods.

3. Results

3.1. Spatial and temporal variability

For the simulated year, we evaluated the dynamics of salinity, temperature, NO_3 concentration, and total phytoplankton biomass along a (i) vertical two-dimensional section perpendicular to the submarine ridge (Barbate High) and parallel to the coast (black line in Fig. 1b), and (ii) at the surface waters of the Trafalgar region (see the white box in Fig. 1b). Due to the curvilinear grid of the model, we had to perform a data interpolation to properly show the vertical section. Thus, the values shown in the surface map (left-hand panels in Figs. 5, 6 and 7) do not always match exactly those shown along the vertical section (right-hand panels in Figs. 5, 6 and 7), but the general trend can be well appreciated.

The dynamics of the spatial distribution observed for the several tracers in the Trafalgar region show a temporal variability closely related to the tidal cycle. Therefore, in this particular subsection, we place the focus on the fortnightly tidal cycle that runs approximately from day 20 at 02:00 h to day 34 at 20:00 h. We can observe that the strongest tide-topography interaction in Barbate High takes place when the tidal current flows westward. This interaction is closely related to the intensity of the tidal current, being stronger at higher u_{vel} values. Therefore, for the sake of simplicity, only the time of maximum tidal current towards the Gulf of Cádiz during neap and spring tides will be described (i.e., NT1 and ST1; Fig. 4). The time of maximum tidal current towards the Alborán Sea during neap and spring tides (NT2 and ST2, respectively) are shown in Figs. D.1 to D.3. We will use salinity as a good upwelling indicator for this region.

Fig. 5 shows the vertical (left-hand panels) and the surface (right-hand panels) distribution of salinity in the Cape Trafalgar region. Both during neap and spring tides, along the vertical section parallel to the coast, we can observe a non-symmetric distribution of salinity determined by the presence of the submarine ridge (i.e., Barbate High), which has a steeper slope on its east side ($\sim 44\%$) than on its west side ($\sim 15\%$). However, during the neap tide period, this west-to-east asymmetry is less pronounced (Fig. 5a). On the east side of the ridge, between 6.13 and $6.06^{\circ}W$, we observe the highest salinity values (left-hand panels in Fig. 5). This band of high salinity waters shows higher values at deeper layers than at the surface. During the spring tide, when the u_{vel} towards the Gulf of Cádiz is maximum ($-0.58\ m\ s^{-1}$), the upwelling of saltier waters is most intense and reaches the most superficial layers of the water column (~ 36.50 – 36.80 ; Fig. 5c). Meanwhile, during the neap tide ($-0.31\ m\ s^{-1}$), the upwelled water is less saline (coming from a shallower depth) and does not reach the most superficial layers (~ 36.40 – 36.50 ; Fig. 5a). The surface distribution of salinity displays a tongue characterised by moderately high values

relative to the surrounding waters, over and to the east of Barbate High (right-hand panels in Fig. 5). The origin of this tongue is the region characterised by the band of high salinity waters observed in the vertical section (around the black circle) and its distribution is affected by the cyclonic eddy that develops due to the interaction of the westward tidal current with the eastern edge of Barbate High (see the black arrows in the right-hand panels in Fig. 5). This eddy is more energetic (stronger horizontal mixing) during spring tides, leading to a less salty tongue (~ 36.45 ; Fig. 5d) than during neap tides when it leads to a saltier tongue (weak horizontal mixing) (~ 36.40 ; Fig. 5b).

During both neap and spring tides, NO_3 concentration along the vertical section parallel to the coast also displays a non-symmetric distribution (see left-hand panels in Fig. 6). Maximum values, between ~ 1.50 and $3.00\ mmol\ m^{-3}$, are found at the eastern side of the submarine ridge, mainly between 6.13 and $6.06^{\circ}W$, concurrent with the band of high salinity values (see left-hand panels in Fig. 5). Like salinity, this band of NO_3 -rich water presents the highest concentration values in the deeper layers and during spring tides, when the intensity of the westward tidal current is stronger. However, in contrast to salinity distribution, there is not an east-to-west gradual decrease of NO_3 concentration since the lowest values are registered over the shallowest area of Barbate High ($< 1.00\ mmol\ m^{-3}$). The superficial distribution of NO_3 , at both tidal moments, shows a spot of relatively high NO_3 concentration ($> 1.50\ mmol\ m^{-3}$) around the location of the band of NO_3 -rich water observed in the vertical section (i.e., black circle on the right-hand panels in Fig. 6). As with salinity, NO_3 surface distribution is affected by the cyclonic eddy. Thus, during neap tides, when the current speed in the cyclonic gyre is low resulting in a weaker horizontal dispersion, NO_3 concentration around the upwelling point is higher. Meanwhile, during spring tides, when the current speed in the gyre is higher, increasing the horizontal dispersion, NO_3 concentration around this point is lower. However, the presence of a NO_3 -rich tongue is not as clearly defined as the salinity tongue over Barbate High. Moreover, along the coast from the Cádiz Bay to Barbate High, following the bathymetric lines, we observe a fringe with minimum NO_3 concentration values ($< 1.00\ mmol\ m^{-3}$; Fig. 6b, d).

Total phytoplankton biomass distribution along the vertical section (see left-hand panels in Fig. 7), shows a pattern quite different from the other state variables. During both neap and spring tides, maximum phytoplankton biomass values appear over the shallowest area of the ridge ($> 0.70\ mmol\ N\ m^{-3}$). During neap tides, high biomass values are also observed in the most superficial layers of the water column to the east side of the ridge (Fig. 7a). The vertical distribution of total phytoplankton biomass along the band between 6.13 and $6.06^{\circ}W$, shows an opposite distribution to salinity and NO_3 concentration, showing low phytoplankton biomass in the recently uplifted waters (Fig. 6a, c and a, c). Thus, low phytoplankton biomass is observed at the deeper layers where NO_3 concentration is maximum, progressively increasing

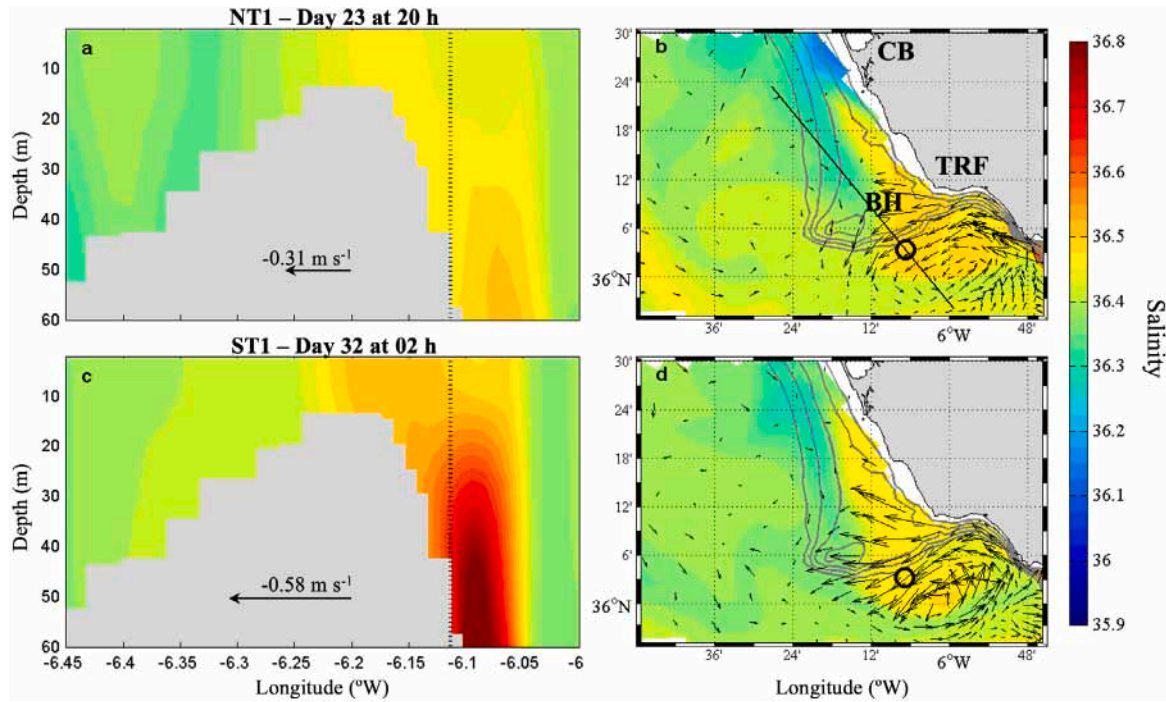


Fig. 5. Salinity distribution in the Cape Trafalgar region during the maximum westward flow of the neap tide period (NT1: a, b) and the spring tide period (ST1: c, d) (see Fig. 3). Left-hand panels show the vertical distribution along a section perpendicular to the submarine ridge (black line in panel b). Vertical dashed lines correspond to the position of open circles in the right-hand panels. Right-hand panels show the surface distribution in the Trafalgar region. Black arrows represent the magnitude and direction of the mean zonal component of current velocity (uvel; m s^{-1}), computed for the whole water column at 6.11°W and 36.06°N at each time-step. Grey solid lines represent the bathymetry (10, 20, 30, 40 and 50 m depth). BH: Barbate High. CB: Cádiz Bay. TRF: Cape Trafalgar.

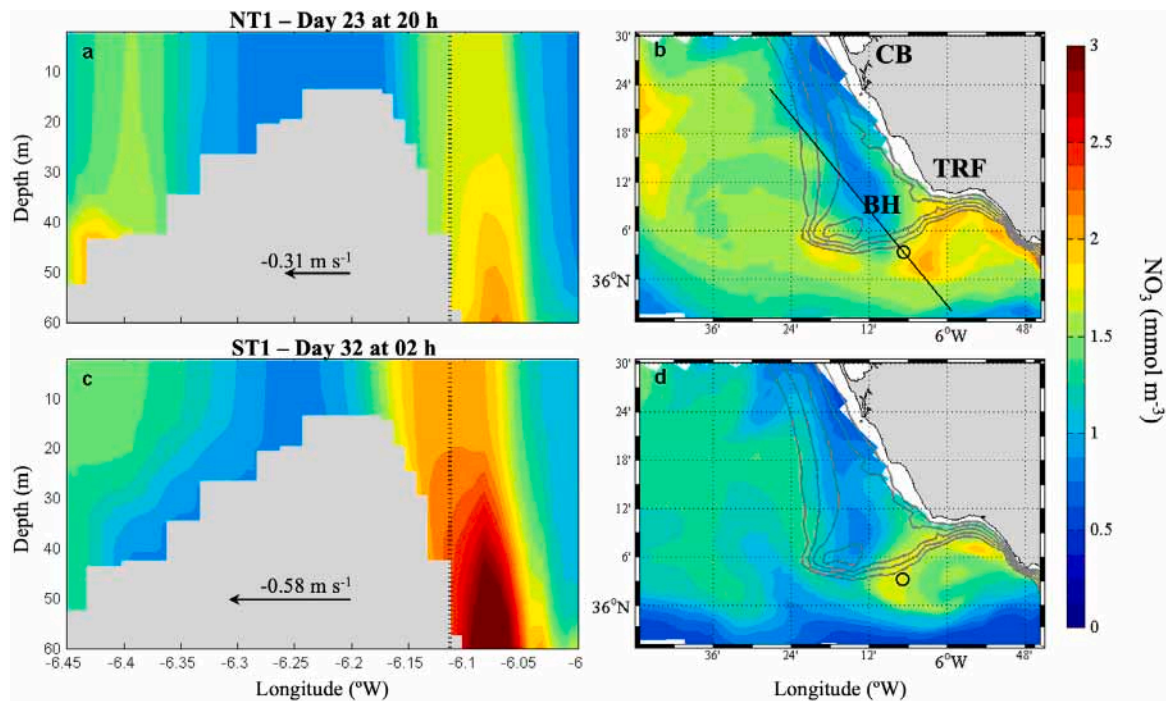


Fig. 6. Nitrate concentration (NO_3 , mmol m^{-3}) distribution in the Cape Trafalgar region during the maximum westward flow of the neap tide period (NT1: a, b) and the spring tide period (ST1: c, d) (see Fig. 3). Left-hand panels show the vertical distribution along a section perpendicular to the submarine ridge (black line in panel b). Vertical dashed lines correspond to the position of open circles in right-hand panels. Right-hand panels show the surface distribution in the Trafalgar region. Grey solid lines represent the bathymetry (10, 20, 30, 40 and 50 m depth). BH: Barbate High. CB: Cádiz Bay. TRF: Cape Trafalgar.

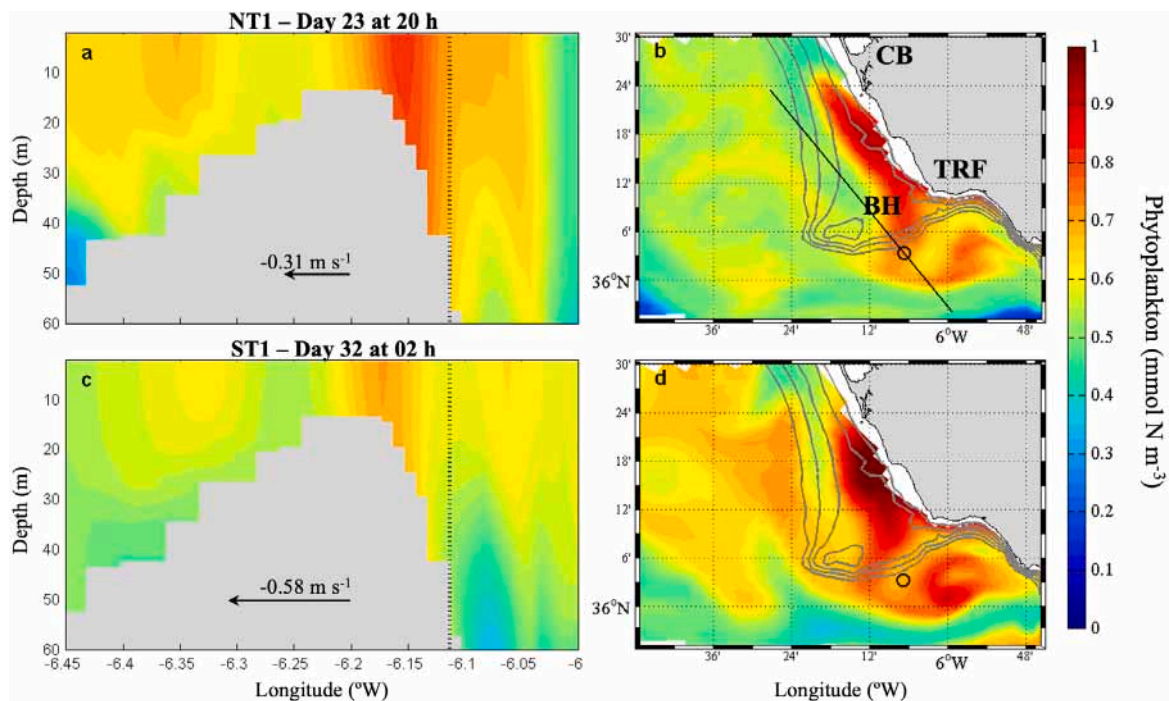


Fig. 7. Total phytoplankton biomass (mmol N m^{-3}) distribution at Cape Trafalgar region during the maximum westward flow of the neap tide period (NT1: a, b) and the spring tide period (ST1: c, d) (see Fig. 3). Left-hand panels show the vertical distribution along a section perpendicular to the submarine ridge (black line in panel b). Vertical dashed lines correspond to the position of open circles in right-hand panels. Right-hand panels show the surface distribution in the Trafalgar region. Grey solid lines represent the bathymetry (10, 20, 30, 40 and 50 m depth). BH: Barbate High. CB: Cádiz Bay. TRF: Cape Trafalgar.

to superficial layers at the same time that NO_3^- concentration decreases (see left-hand panels in Figs. 6 and 7). The surface distribution of total phytoplankton, at both tidal stages, shows a tongue over Barbate High characterised by high biomass ($>0.80 \text{ mmol N m}^{-3}$; see right-hand panels in Fig. 7) displaced to the east side of the ridge due to the influence of the cyclonic eddy (Fig. 7b, d). This tongue is richer in phytoplankton biomass during spring tides (Fig. 7d) when the input of NO_3^- -rich water is maximum (Fig. 6c). Moreover, phytoplankton biomass through this tongue seems slightly higher close to the coastline extending from Cádiz Bay to Cape Trafalgar. Also, in the rest of the domain (mainly over the continental slope), phytoplankton biomass is higher during spring tides than during neap tides. This pattern is the opposite of the one observed for the NO_3^- distribution, whose concentration over the continental shelf is slightly higher during neap tides than during spring tides (Fig. 6b, d).

The sequential analysis of the vertical and superficial distribution of these three variables shows how the band of saltier, NO_3^- -rich and with low phytoplankton biomass water, as well as the cyclonic gyre and the tongue over Barbate High (and the surrounding waters), oscillate continuously from east-to-west due to the cyclical change of direction of the zonal component of the tidal current. Thus, the ‘water masses’ in the Trafalgar region are displaced to the east when uvel flows towards the Alborán Sea (positive values), and to the west when uvel flows towards the Gulf of Cádiz (negative values). Furthermore, these east-to-west-to-east oscillations are more or less noticeable depending on the intensity of uvel. To better observe this oscillating movement that is a consequence of the tidal cycle, and all its associated processes, the reader is referred to the supplementary video that shows the computer simulation of these variables for the first 3 months of the simulation (<http://t.ly/bVQQ>).

3.2. Hovmöller diagram

In the previous subsection, we described how an entry of saltier, nutrient-rich and low phytoplankton biomass water is observed between ~ 6.13 and 6.06°W , with a time-dependent variability closely

related to the tidal cycle. To examine this process in more detail, in Fig. 8 we show the Hovmöller diagrams (depth-variability as a function of time) of several state variables and the Richardson number (R_i) together with the temporal dynamics of uvel to outline the tidal cycle. We do so for a particular location at 6.11°W – 36.06°N (see the red circle in Fig. 1b) and for the whole simulated year.

The time-dependent distribution of salinity shows a fortnightly periodicity. Higher values of salinity are observed during spring tides (Fig. 8b). During the most intense spring tide periods (every ~ 30 -days) the water is slightly saltier than during the less intense periods, although in both cases the upwelled water always reaches the most superficial layers of the water column. Whereas during the neap tide periods (i.e., weak turbulent mixing) the lowest values of salinity are observed. It should be noted that for the entire period, the salinity of the water column continuously oscillates between relatively high and relatively low values, in good agreement with the tidal current oscillations both in direction and intensity (Fig. 8a). From July to November, the upwelling events seem to be less intense than during the rest of the year, with shallower (intermediate) and less salty upwelled water than that observed in other spring tide periods, probably North Atlantic Central Water (NACW), reaching the most superficial layers. Furthermore, from October to December we observe the lowest salinity values for the whole simulation year.

The temperature at the selected location point also shows a pattern of variability associated with the fortnightly tidal cycle, although it is less appreciable than for salinity (Fig. 8c). During spring tide periods, we observe the lowest water-column temperatures and highest salinity values (Fig. 8b). The temperature is colder during the most intense spring tides. Conversely, during neap tides, the observed temperature is $\sim 2^\circ\text{C}$ warmer. As with salinity, the water temperature oscillates between relatively high and relatively low values depending on the tidal cycle (Fig. 8a). During the winter months (January to April), as expected, the coldest temperatures of the year are observed. While from July to November, in the first 30 m of the water column are observed the highest temperatures (~ 22 – 24°C) of the whole year.

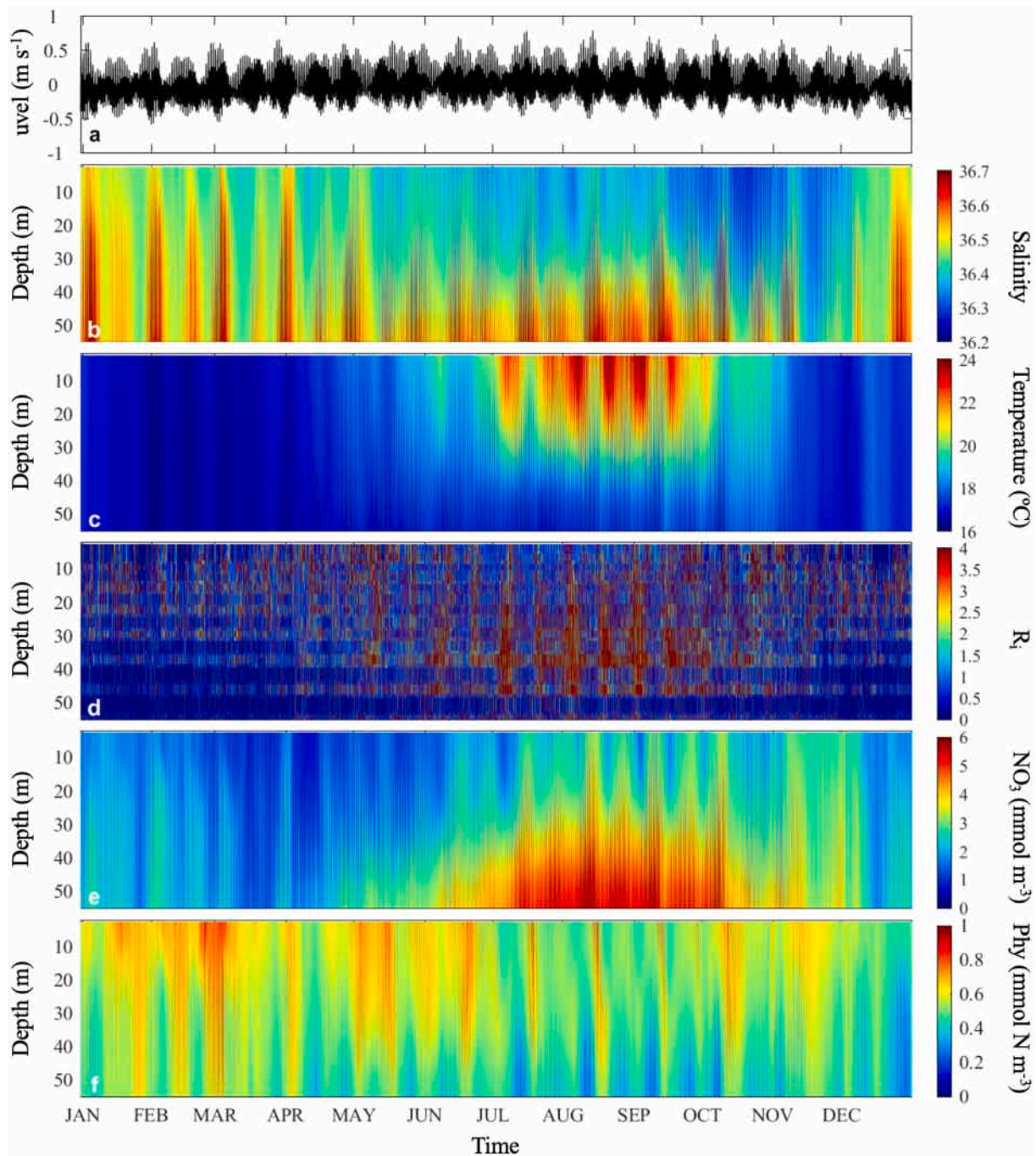


Fig. 8. (a) Temporal evolution of the mean zonal component of current velocity (u_{vel} ; m s^{-1}), computed for the whole water column at 6.11°W and 36.06°N (red circle in Fig. 1b), along the whole-year simulation. Hovmöller diagrams, computed at the same point (red circle in Fig. 1b), illustrating spatial (i.e., depth) and temporal variability of (b) salinity, (c) temperature ($^{\circ}\text{C}$), (d) the Richardson number (R_i , adimensional), (e) nitrate concentration (NO_3 , mmol m^{-3}) and (f) total phytoplankton biomass (Phy, mmol N m^{-3}).

The seasonal variability of stratification and tides modulate the intensity of diapycnal mixing over the water column, as suggested by the Richardson number, $R_i = N^2/U_z^2$ (N : Brunt-Väisälä frequency; U_z : flow shear). R_i reaches the greatest values from June to October associated to a strong summer thermocline that warrants flow stability ($R_i \gg 1/4$). During the rest of the year, the flow is also rather stable ($R_i > 1/4$), but with some small-scale potentially unstable patches of the fluid where vertical mixing is enhanced ($R_i < 1/4$). At fortnightly tidal scale, R_i shows smaller values (enhanced mixing) during spring tides. The fortnightly signal is especially noticeable during the summer months.

NO_3 concentration also displays a fortnightly periodicity (Fig. 8e). Only during the most intense spring tides, NO_3 -rich water seems to

reach the surface layers. During neap tides, NO_3 concentration throughout the water column is relatively low ($< 1.00 \text{ mmol m}^{-3}$). As with salinity and temperature, the NO_3 concentration of the whole water column continuously alternates between relatively high and relatively low values, in agreement with the tidal current oscillations (direction and intensity; Fig. 8a). Throughout the whole year, the lower NO_3 concentration values are observed from March to May (Fig. 8e). While the maximum NO_3 concentrations are observed from July to October, coinciding with the period of highest stratification of the water column (Fig. 8d). As a result, the nutrient-rich upwelled water is unable to reach the most superficial layers of the water column. The upwelling of less salty and nutrient-rich waters (compared with other upwelling

events) supports the fact that during fall there is an upwelling of NACW in this region.

Fig. 8f shows the dynamics of total phytoplankton biomass, with higher values at surface layers that tend to decrease with depth. Nevertheless, the phytoplankton biomass is quite high at all depths because the vertical domain analysed here is relatively shallow (50 m) and therefore well illuminated. As previously observed (Figs. 5 and 6), phytoplankton biomass distribution behaves oppositely to NO_3 concentration. Thus, during the neap tide periods, when NO_3 concentration throughout the water column is relatively low, phytoplankton biomass shows high values with a pretty homogeneous distribution. For phytoplankton biomass is slightly more difficult to appreciate a fortnightly periodicity than for salinity, temperature or even NO_3 concentration. Although, as with the other variables, a continuous fluctuation between high and low phytoplankton biomass is still observed. Phytoplankton biomass is lowest from July to October when stratification is highest (Fig. 8d), preventing them to access to the nutrient-rich upwelled water (Fig. 8e).

3.3. Wavelet power spectrum

We performed a wavelet power spectrum analysis of the time series shown in the Hovmöller diagram to extract information about the temporal variability of salinity, temperature, NO_3 concentration and total phytoplankton biomass. For each variable, the WPS was computed using the hourly average for the whole water column.

The WPS for the salinity signal (time series) displays two strong peaks at ~ 15 -day and ~ 30 -day periodic components (Fig. 9a). The time-averaged WPS shows that these two periodicities in the signal are statistically significant, contributing on average $\sim 21\%$ and $\sim 31\%$ of the total variance, respectively (Fig. 9b). However, the ~ 15 -day WPS peak is statistically significant only during winter months, and the ~ 30 -day peak from January to mid-May and from October to December (Fig. 9a). During the whole year, a second pair of peaks are observed at the ~ 12 -h and ~ 1 -day periods (Fig. 9a). For the ~ 12 -h period, high values of WPS are observed during the most intense spring tides (i.e., approximately every 30 days), and weaker values during less intense spring tides. These discontinuous peaks of periodicities or frequencies in the WPS, disregarding the one observed (approximately) between days 15 and 23, are statistically significant. The peak in the ~ 1 -day periodicity shown by the WPS is almost coincident in time with the 12-h period, although it is not always statistically significant. However, when looking at the time-averaged WPS, these two peaks (at ~ 12 -h and ~ 1 -day) are small and therefore do not contribute significantly to the variance of the signal (Fig. 9b).

The WPS for temperature shows two relatively strong peaks at the ~ 15 -day and 30-day periodic components during the summer months, between June and October (Fig. 9c). Both peaks, locally, are not statistically significant (i.e., time and period coordinates), although the time-averaged value of the WPS does highlight these periodicities as being statistically significant (Fig. 9d). In particular, the ~ 15 -day periodicity of the temperature signal, although smaller compared to the one obtained for salinity, explains $\sim 26\%$ of the total variance. From May to November, weaker but statistically significant peaks are observed for the 12-h and 1-day periods (Fig. 9c). Like salinity, the 12-h periodicity peaks of temperature are slightly stronger during the events of most intense spring tides and weaker during less intense spring tides. The 1-day periodicity peak, which is weaker than the 12-h peak, becomes relevant only during events of the most intense spring tides. Both time-averaged WPS peaks, 12-h and 1-day, show low values not statistically significant.

The WPS for NO_3 concentration shows some strong but time-discontinuous peaks at the ~ 15 -day and ~ 30 -day periodic components of the signal, but locally they are not statistically significant (Fig. 9e). However, the time-averaged WPS peaks for these signal periods are statistically significant. The observed peak at ~ 15 -day explains $\sim 12\%$

of the total variance (Fig. 9f). A weak time-discontinuous peak that is statistically significant can be observed at the ~ 12 -h signal period (Fig. 9e). This ~ 12 -h peak shows slightly stronger values during the more intense spring tides and weaker during the less intense spring tides. At the ~ 1 -day periodic component, several peaks in the time domain are observed but only half of them are statistically significant. According to the time-averaged WPS, neither the ~ 12 -h nor the ~ 15 -days periodic components are statistically significant (Fig. 9f).

The WPS for total phytoplankton biomass shows a strong peak at ~ 15 -day periodic component, but its statistical significance is discontinuous in the time domain (from February to March, from mid-March to May, and from July to October; see Fig. 9g). On average, this periodicity explains $\sim 21\%$ of the total variance (Fig. 9h). There are other strong peaks in the WPS between the ~ 30 -day and 60-day signal periods. As with previous state variables, weaker peaks are observed at ~ 12 -h and ~ 1 -day periodicities, both discontinuous in time (Fig. 9g). At the beginning and the end of the year, the 12-h peak seems more uniform over time, without showing a discontinuous pattern as clear as for the other variables. The ~ 1 -day periodicity peak is slightly stronger and statistically significant at both the beginning and end of the simulated year. Between April and November, few of these peaks in the WPS at the ~ 12 -h or ~ 1 -day signal periods are statistically significant. Furthermore, at the time-averaged WPS, both peaks were very weak and not statistically significant (Fig. 9h). It should be noted that at several moments in the year, we can observe a peak in the WPS for the ~ 6 -h signal period which is statistically significant, coinciding with the semi-diurnal tidal cycle.

3.4. Wavelet coherence analysis

Finally, we performed a wavelet coherence analysis to study the covariance between uvel as the driving variable and salinity, temperature, NO_3 concentration, and phytoplankton biomass as the response variables. The consistency of the coherence between the driver and the different response variables can be identified from the average wavelet cross-spectrum (see left-hand panels in Fig. 10). The most consistent signal period for each pair of variables (i.e., the periodicity with the highest cross-spectrum power value) was identified, and subsequently, the corresponding oscillations were extracted for the whole period to compare their amplitudes and phases (see right-hand panels in Fig. 10).

The strongest phase coherence between the uvel and salinity signals is found at the ~ 15 -day period, with a cross-spectrum power value that explains $\sim 37\%$ of the total variance (Fig. 10a). For the entire simulation, there is an average delay of 1.05 days between the phase of the two variables, with the driver variable (uvel) peaking before the response variable (salinity) (Fig. 10b). The maximum delay between uvel and salinity signals is observed at the beginning and the end of the year, with a delay of 2.36 and 2.84 days, respectively, between driver and response. While the minimum delay is observed from June to August.

Between uvel and temperature, the strongest peak in the WCO analysis is observed at the ~ 12 -h period (Fig. 10c), although the area under this peak is not statistically significant. Therefore, as we did with salinity, we focus our analysis on the ~ 15 -day signal period. This peak has a power value that explains the $\sim 32\%$ of the total variance. During most of the simulated year, the response variable (temperature) appears before the driver variable (uvel), with an average delay of 1.97 days (Fig. 10d). However, between May and July and between September and October, the driver variable appears before the response variable, with a delay of up to 3.14 days.

The coherence between uvel and NO_3 concentration also shows the strongest and statistically significant peak at the ~ 15 -day period (Fig. 10e). The ~ 15 -day cross-spectrum power value explains $\sim 23\%$ of the total variance. Again, driver and response variables show a lack of synchrony (Fig. 10f), with the response variable appearing before or after the driver. At the beginning and the end of the year, the driver

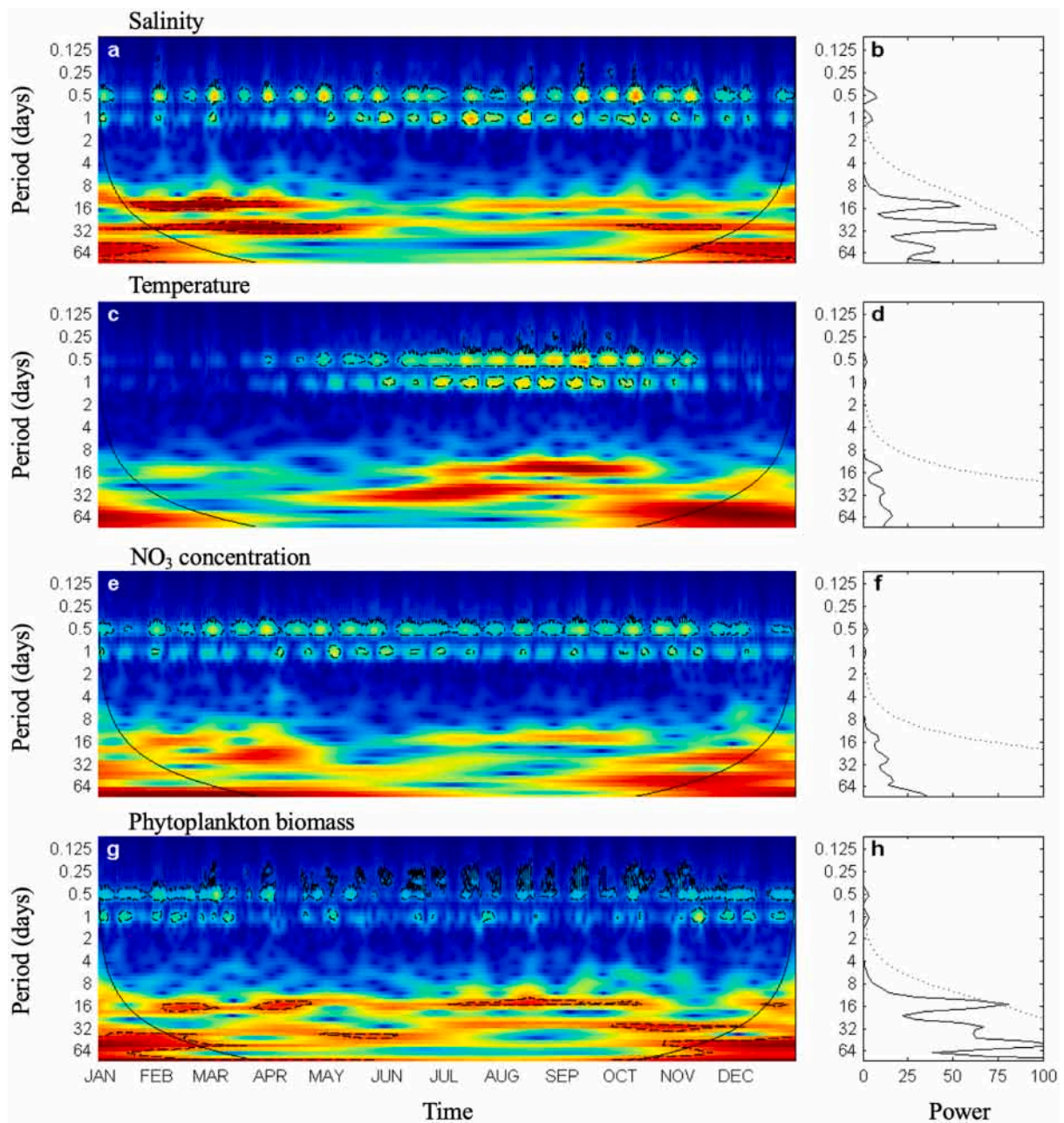


Fig. 9. Wavelet analysis computed for the 1-year simulation. Left-hand panels show the wavelet power spectrum for (a) salinity, (c) temperature ($^{\circ}\text{C}$), (e) nitrate concentration (NO_3 , mmol m^{-3}) and (g) total phytoplankton biomass (mmol N m^{-3}). The colour code varies from dark blue (low values) to dark red (high values). Black lines indicate the cone of influence. Right-hand panels show the average wavelet power spectrum for (b) salinity, (d) temperature, (f) NO_3 concentration and (h) total phytoplankton biomass. Dotted lines, at all panels, show the $\alpha = 5\%$ significance level computed based on 1000 Markov bootstrapped series.

(uvel) appears before the response variable (NO_3 concentration), with a mean delay of 2.27 days. From May to October, the response variable appears before the driver, with a mean delay of ~ 0.50 days.

Likewise, total phytoplankton biomass displays the strongest coherence with uvel at the ~ 15 -day signal period, with a cross-spectrum power value that explains $\sim 48\%$ of the total variance (Fig. 10g). The delay between both variables follows a pattern almost opposite to uvel and NO_3 concentration (Fig. 10f). The response variable (phytoplankton biomass) appears before the driver (uvel) at the beginning and the end of the simulated year, with a mean delay of 0.71 days. While from April to October, the driver variable appears before the response variable, with a mean delay of 1.38 days (see Fig. 10h).

To verify that these results are not a consequence of our approach, both the WPS and the WCo analyses were repeated using two different

time series: (i) a time series that only considers the temporal dynamics of salinity, NO_3 concentration and phytoplankton biomass at 20 m depth at 6.11°W – 36.06°N (red circle in Fig. 1b); and (ii) a time series obtained from the average values, of each variable, for the whole water column and considering the entire upwelling band region, i.e., between 6.13 and 6.06°W . In both cases, the obtained results were similar to those described here (data not shown).

4. Discussion

In the Cape Trafalgar region, the tide-topography interaction of the westward tidal current (towards the Gulf of Cádiz) with the eastern steep edge of the submarine ridge, results in a tidal pumping process that brings up deep, saltier, colder, nutrient-rich and phytoplankton-poor water onto Barbate High. This process is similar to the interaction

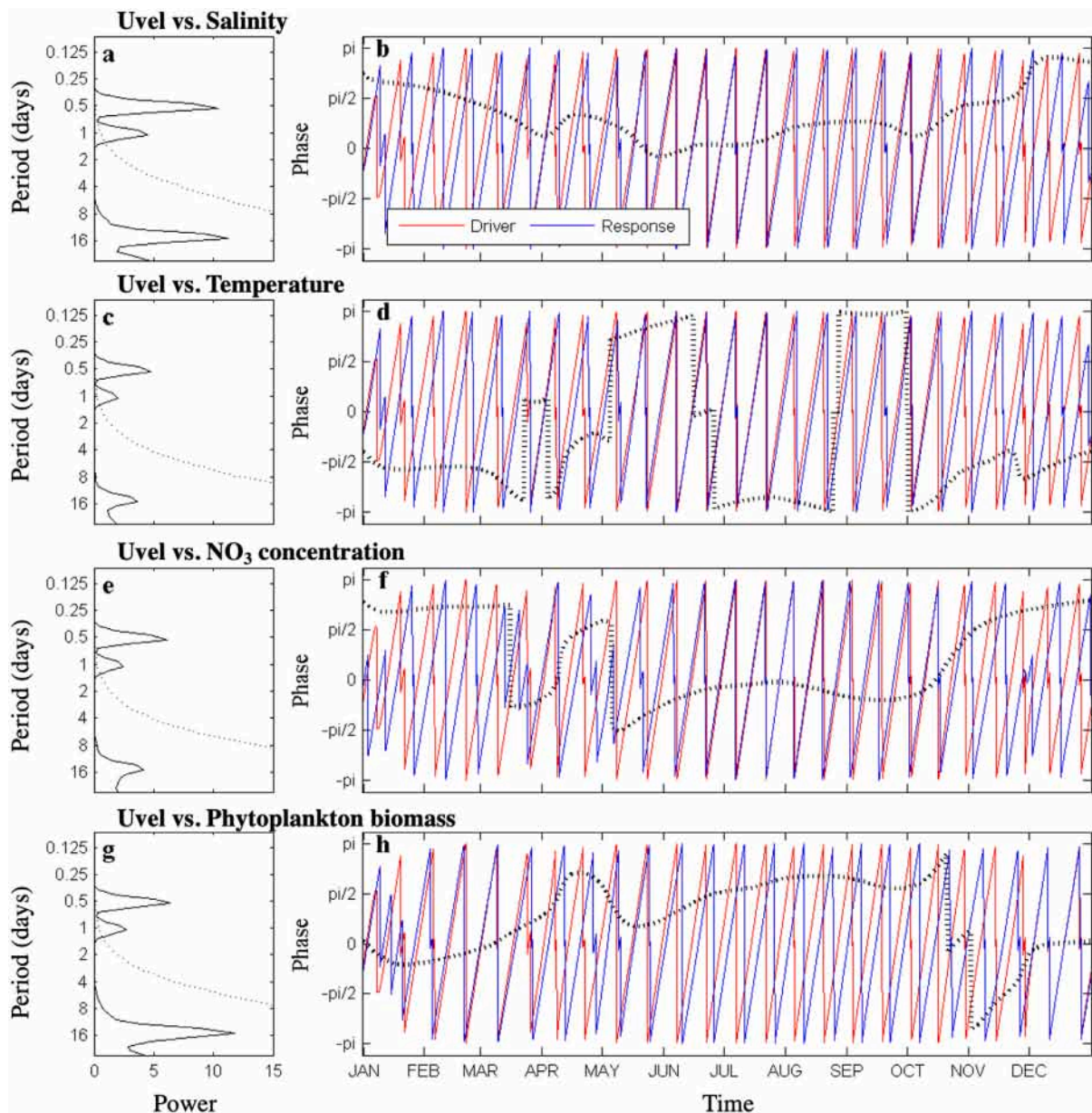


Fig. 10. Wavelet coherence analysis computed for the zonal component of current velocity (*uvel*; m s^{-1}) versus (a, b) salinity, (c, d) temperature ($^{\circ}\text{C}$), (e, f) nitrate concentration (NO_3 , mmol m^{-3}), and (g, h) total phytoplankton biomass (mmol N m^{-3}) for the 1-year simulation. Left-hand panels (a, c, e, g) show the average wavelet cross-spectrum. Dotted lines show the $\alpha = 5\%$ significance levels computed based on 1000 Markov bootstrapped series. Right-hand panels show the phases of both time series computed for the peak of highest coherence: *uvel* (red line) versus (b) salinity, (d) temperature, (f) NO_3 , and (h) total phytoplankton biomass (blue lines). Dashed lines represent the phase difference between both signals.

previously described at Georges Bank, a shallow submarine feature located on the Gulf of Maine (e.g., [Chen and Beardsley, 1998](#); [Franks and Chen, 2001](#); [Hu et al., 2008](#); [Ji et al., 2008](#)). Particularly during spring tides, denser water is advected upslope to subsurface layers leading to a spatial asymmetry between the eastern and the western side of the ridge (see left-hand panels in [Figs. 5–7](#)) ([Vargas-Yáñez et al., 2002](#)). Concurrently, the interaction of the tidal current with the eastern edge of Barbate High leads to the formation of a cyclonic eddy ([Bolado-Penagos et al., 2020](#)), which spreads the upwelled waters over and to the west of the ridge. This interaction results in the development of a water tongue with particular properties that are slightly different from the surrounding waters (right-hand panels, [Figs. 5–7](#)). The upwelling of deep, salty and nutrient-rich water is more effective during spring tides, but due to the greater horizontal mixing mediated by the cyclonic gyre the water properties are diluted. During neap tides the lower horizontal mixing leads to a saltier and enriched water tongue.

The high phytoplankton biomass observed over Barbate High (right-hand panels in [Fig. 7](#)), is favoured by the continuous and cyclical input of nutrients (e.g., [Chen and Beardsley, 1998](#); [Hu et al., 2008](#); [Ji et al., 2008](#); [Blauw et al., 2012, 2018](#)). Actually, a previous study with a similar model configuration ([Sánchez-Garrido et al., 2015](#)) shows how a simulation with tidal forcing results in a chl-*a* concentration over Barbate High ~ 3 times higher than in a simulation without tides. At the selected location point for in-depth analysis of the upwelling dynamics, we observe that NO_3 concentration and phytoplankton biomass display an anti-phase behaviour ([Fig. 8d, e](#)). The ‘water masses’ observed in this region (i.e., Trafalgar’s tongue, upwelled water) are continuously moving back and forth due to the fluctuations of the tidal current intensity and direction (see Supplementary video – <http://t.ly/bVQQ>). The pumping of NO_3 -rich waters takes place when the tide flows towards the Gulf of Cádiz while Trafalgar’s tongue moves westward. Hence, at the analysed upwelling point there is an uplifting of nutrient-rich

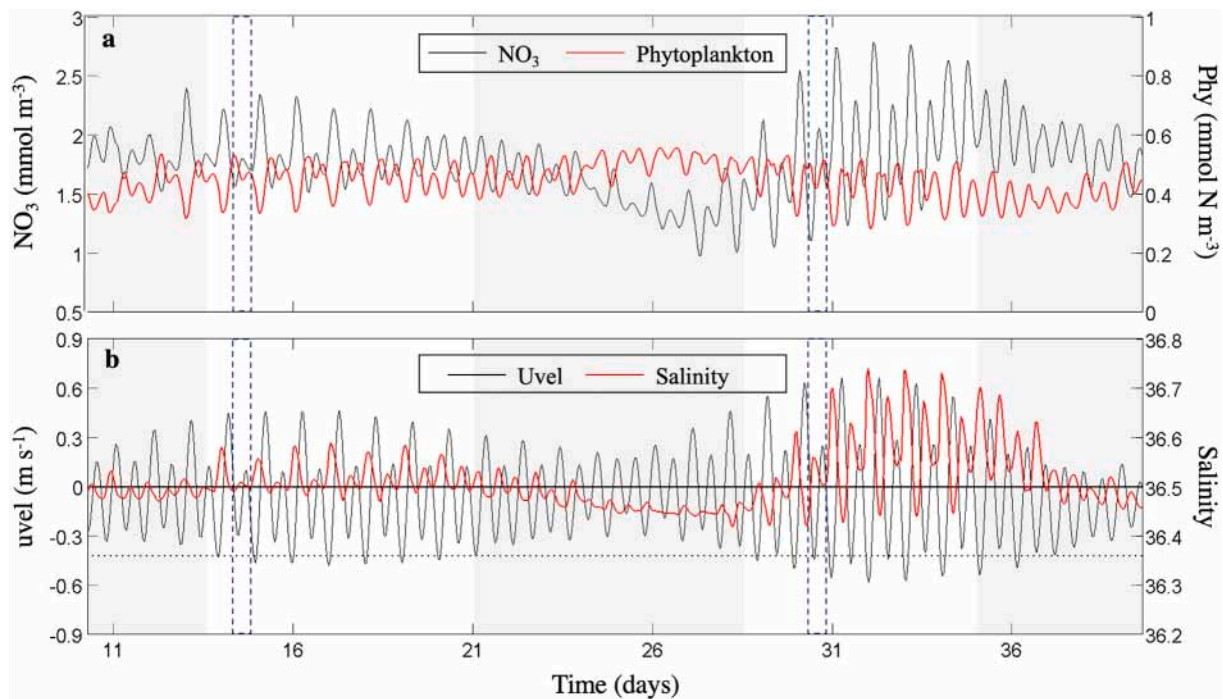


Fig. 11. (a) Temporal evolution of mean nitrate concentration (NO_3 , mmol m^{-3} ; black line) versus mean phytoplankton biomass (Phy, mmol N m^{-3} ; red line). (b) Temporal evolution of the mean zonal component of the tidal current (uvel, m s^{-1} ; black line) versus mean salinity (red line). The black horizontal line represents the zero uvel reference (delimiting positive (negative) values that indicate flow driven eastward (westward), towards the Alborán Sea (Gulf of Cádiz)). The black dashed line represents uvel equal to -0.42 m s^{-1} . Black dashed rectangles point out the first tidal-pumping event during both spring tide periods. The grey background identifies the neap tide periods. All time series were computed for the whole water column at 6.11°W – 36.06°N (red circle in Fig. 1b).

water with relatively low phytoplankton biomass and a low influence of the phytoplankton-rich Trafalgar's tongue. Nevertheless, when the tide flows towards the Alborán Sea, there is no pumping of nutrient-rich water and the phytoplankton-rich Trafalgar's tongue is advected eastwards on the analysed upwelling point, and re-distributed along the whole water column. This coincides with a maximum in phytoplankton biomass. This fact can be appreciated in Fig. 11a, where mean NO_3 concentration and phytoplankton biomass computed for the whole water column at the upwelling point (red circle in Fig. 1b) over two tidal cycles behave oppositely.

The tide-topography interaction occurs during both neap and spring tides (Figs. 5 and 6). However, during neap tides the water advection is weak, so the pumped waters (slightly more saline and moderately rich in NO_3 than the surrounding waters) do not reach the most superficial waters (Fig. 8c). The intensity of the tidal pumping of saltier, colder and NO_3 -rich waters depends on the tidal current speed, being stronger and reaching the most superficial layers when the westward current speed (towards the Gulf of Cádiz) is higher (Figs. 5, 6, 8b–d). In Fig. 8b–d, it can be observed that the tidal pumping of saltier, colder and NO_3 -rich waters takes place several days before uvel achieves its maximum speed during spring periods; when the tidal current speed is strong enough to cause a significant upwelling of deep waters. Fig. 11b shows the temporal dynamics of the average uvel (m s^{-1}) and salinity, computed for the whole water column at 6.11°W – 36.06°N (red circle in Fig. 1b) over two tidal cycles. It should be noted that the semi-diurnal tidal cycle in Trafalgar, with two high tides and two low tides per day, shows a diurnal inequality with one high-low tidal cycle more intense than the other. It also should be noted that during a semi-diurnal tidal cycle, due to the expected time delay between the maximum westward current flow (negative uvel) and the horizontal displacement of the water mass (high tide), the maximum upwelling of salty waters is observed 2–3 h after the maximum flow. Thus, each peak of maximum salinity is caused by the previous peak of maximum westward tidal current (towards the Gulf of Cádiz). Moreover, due to the diurnal inequality, the oscillations of salinity peaks are also asymmetric.

Considering Fig. 11b, it seems that the upwelling of saltier waters does not take place when the westward tidal current speed is lower than 0.42 m s^{-1} (negative uvel values, black dashed line). Black dashed rectangles mark the first effective tidal-pumping event during both spring tide periods. During the less intense spring tide, the first tidal-pumping event of deep and saltier water towards the more superficial layers occurs around day 13 at 02:30 h, raising the average salinity to 36.58. Three hours before (around day 12 at 23:36 h), the westward uvel is $\sim 0.42 \text{ m s}^{-1}$. In the previous days, characterised by lower uvel values, the upwelling of deep and saltier water does not take place. Similarly, during the most intense spring tide event, the first tidal-pumping episode takes place around day 29 at 03:48 h, as a result of uvel $> 0.42 \text{ m s}^{-1}$ (around day 29 at 01:54 h). This behaviour is observed throughout the whole year (data not shown). However, this dynamic is influenced by the summer stratification of the water column (Fig. 8d) linked with the seasonal variability of solar irradiance (and thus heat flux). In summer, although during spring tides the upwelling of salty and nutrient-rich waters continues to take place (Fig. 8b–e), the heat-flux induced stratification reduces the vertical mixing preventing the upwelled waters from reaching the most superficial layers (Fig. 8d). Hence, the phytoplankton located in the most superficial layers will not have access to the uplifted nutrients, decreasing the phytoplankton community nutrient uptake rate and biomass (Fig. 8f). This seasonal variability was previously observed in the study region using 10-years of ocean colour observations using the MEdium Resolution Imaging Spectrometer ($\sim 300 \text{ m}$ resolution Sala et al., 2018).

For all the analysed variables, salinity, temperature, NO_3 concentration and total phytoplankton biomass, the time-averaged WPS shows a peak at the fortnightly tidal cycle, explaining $\sim 21\%$, $\sim 26\%$, $\sim 12\%$ and $\sim 21\%$ of the total variance respectively (right-hand panels in Fig. 9). Although the time-averaged WPS peak observed at the ~ 15 -day signal period for temperature is smaller than for salinity, it explains a higher percentage of the total variance (Fig. 9d). However, the local WPS analysis shows statistically significant results for salinity at certain time periods of the simulated year but not for temperature (Fig. 9a and c),

confirming that temperature is not a good proxy to identify tidally driven upwelling events in this region. Salinity and temperature are intrinsic properties of the upwelled water, and no biological activity can alter them. However, upwelled nutrients may be assimilated by phytoplankton, weakening the fortnightly periodicity component of NO_3 concentration, which would explain the absence of statistically significant peaks in the WPS analysis (Fig. 9e). Furthermore, the lack of a statistically significant fortnightly periodicity component may also be related to the lack of continuity in the upwelling events, which are less intense during neap periods (Fig. 8). In fact, the WPS analysis with daily averaged time series results in stronger fortnightly signals, statistically significant over longer periods (data not shown). Supporting these findings, the WCo analysis between state variables (drivers against responses) identifies the strongest statistically significant covariation patterns at the fortnightly tidal period (see left-hand panels in Fig. 10). However, only in very few certain moments of the temporal domain, the driver variable (uvel) is in synchrony with the response variables (i.e., salinity, temperature, NO_3 concentration and total phytoplankton biomass). For most of the year, the time delay between driver and response variables varies back and forth. Although the tidal-topography interaction is the main responsible for the temporal and spatial variability in the Trafalgar region, it is not until the westward uvel reaches a threshold value of $\sim 0.42 \text{ m s}^{-1}$ that deep water is pumped to the most superficial layers. Moreover, the tidal cycle is also responsible for the horizontal displacement of water masses that modifies the properties of the water column. Finally, the seasonal variability, the Gulf of Cádiz circulation pattern, and the wind regime will also introduce changes (Vargas-Yáñez et al., 2002; Sala et al., 2018). Therefore, this asynchrony between driver and response variables is conceptually expected but difficult to anticipate before performing the simulations, due to non-linearities in the dynamics.

The WCo analysis between uvel (as driver variable) against NO_3 concentration and phytoplankton biomass (as response variables) shows an opposite pattern (disregarding May) (Fig. 10 f, h). When NO_3 oscillations appear later than the driver, the phytoplankton biomass oscillations appear earlier than the driver, and vice versa. In any case, the maximum of phytoplankton biomass must be a response to the previous NO_3 tidal pumping. This is explicit in the primary production equations of the ecosystem model used to perform these simulations. In the Cape Trafalgar region, there seems to be always a phytoplankton community ready to assimilate nutrients when there is a new input. After a neap tide period with a minimum nutrient input (Figs. 6c, 8c), during which the previously pumped nutrients have been consumed or dispersed, a new nutrient input during spring tides (when westward uvel $>0.42 \text{ m s}^{-1}$) can be efficiently assimilated by the pre-existing phytoplankton community. In fact, some modelling exercises have shown how patterns of periodic inputs of nutrients into the surface (especially tidal pulses) lead to the apparition of larger values of phytoplankton biomass than scenarios associated with a continuous supply; furthermore, it can favour some taxonomic groups as diatoms and larger zooplankton (Macías et al., 2010).

Although the fortnightly period is shown to be the most influential scale of the tidal variability in our study area, as previously found by Sala et al. (2018), it should be highlighted that the WPS and WCo analyses show a peak at the ~ 12 -h period which confirms the influence of the semi-diurnal tidal cycle on the simulated state variables (see Fig. 9 and left-hand panels in Fig. 10). However, the time-averaged WPS and WCo analyses for the entire year show that this periodicity is not statistically significant, probably due to the discontinuity of the tidal-pumping events, which do not take place when uvel was $<0.42 \text{ m s}^{-1}$.

Previous studies have highlighted that the tidal pumping of nutrient-rich waters could be enhanced by other mechanisms not addressed in our study, such as river discharges (Huertas et al., 2005; Navarro et al., 2006) and wind-driven upwellings (Vargas-Yáñez et al., 2002; Sala et al., 2018). Our model configuration does not include nutrient inputs

from river discharges (e.g., Guadalquivir River). This fact facilitates the aim of this study by removing its possible influence on the time series analysis. However, the lack of nutrient river discharges results in an underestimation of primary production of the northern coastal areas in the Gulf of Cádiz, which can be observed in Fig. 3. The superficial circulation pattern of the Gulf would advect the productive water that flow from the rivers towards the Trafalgar region, increasing its primary production. Therefore, it would be interesting for future works to compare our results with a model that prescribes nutrient inputs from rivers to evaluate and quantify their relative importance. Regarding the wind forcing, since it was explicitly resolved in our model, we have extracted the zonal component of the wind at 10 m (u_{10}) for the analysed area. The comparison between both u_{10} and salinity time series does not show a clear pattern of variability (Fig. E.1), and the lack of correlation was supported by a WCo analysis (data not shown). Nevertheless, its influence may introduce ‘noise’ in our analysis but, as the seasonal variability, without undermining the tidal impact in the region. However, Bolado-Penagos et al. (2020) observed that easterly winds increase the retention capacity of Cape Trafalgar. Moreover, due to the presence of the ridge that characterises this region a stacking of waters may take place leading to a deepening of the interface. Therefore, it would be interesting to carry out a more detailed analysis with a different model setup.

Finally, it is worth mentioning that several authors have described a secondary impact of the tidal cycle in the Strait of Gibraltar that would increase the primary production in the Alborán Sea. Due to the velocity divergences developed in the surface layer when undulatory features are generated over the Camarinal Sill, a lateral advection of patches with high chl-*a* concentration was observed from coastal areas to the centre of the Strait (Macías et al., 2006; Vázquez et al., 2009; Ramírez-Romero et al., 2012; Bruno et al., 2013). Thus, Cape Trafalgar is considered to act as a source of nutrient- and biomass-rich waters that periodically will flow to the Alborán Sea. Moreover, a recent study (Bolado-Penagos et al., 2020) has described a transport process of phytoplankton along the coastal margin between Cape Trafalgar and the Alborán Sea. These transport events would imply a loss of nutrient and phytoplankton-rich waters towards Alborán that could be introducing some degree of ‘noise’ in our wavelet analysis (WPS and WCo).

5. Conclusions

The results obtained in this study confirm the importance of the tidal cycle for the productivity of the Cape Trafalgar region through an active and periodic forcing resulting in a pulsating upwelling pattern. The interaction of the westward tidal current with the eastern edge of the ridge perpendicular to the coast that characterises this region results in the upwelling of deep, saltier and nutrient-rich waters that favour the development of a tongue with high phytoplankton biomass over Barbate High.

The fortnightly period is confirmed as the most influential scale of the tidal cycle when the tidal-pumping process takes place during spring tides characterised by a westward tidal current $>0.42 \text{ m s}^{-1}$, between ~ 3 days before and after the moment of maximum westward flow. This tidal interaction implies cyclical fertilisation of the illuminated surface waters in the region, favouring the growth of the phytoplankton community. After an (effective) tidal-pumping event during spring tides, periods of minimum NO_3 concentration lead to phytoplankton nutrient-limitation at the surface and low primary production. However, these periods do not last long (~ 4 days).

The seasonal variability of solar radiation affects the Cape Trafalgar region only during summer. From July to September, the stratification of the water column increases the vertical thermal gradient in the water column, which decreases the vertical uplifting of nutrient-rich water into the first 30 m of the water column. For this reason, the upwelled nutrients driven by the tidal-topography interaction remain unconsumed below the euphotic layer, doubling their concentration by accumulation. Meanwhile, the phytoplankton biomass decreases due to a lower input of nutrients to the most superficial layers.

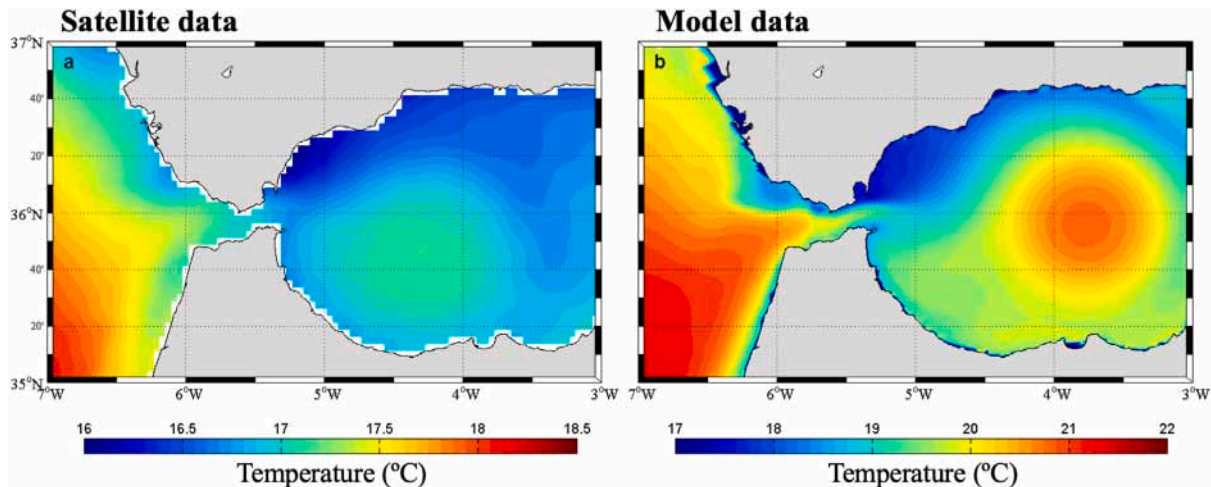


Fig. A.1. (a) Sea surface temperature (SST, °C) climatology derived from remote sensing images corresponding to the period 2001–2010. (b) Model-derived climatology of SST for the 3rd-year simulation. Please note the different scale ranges between both subplots.

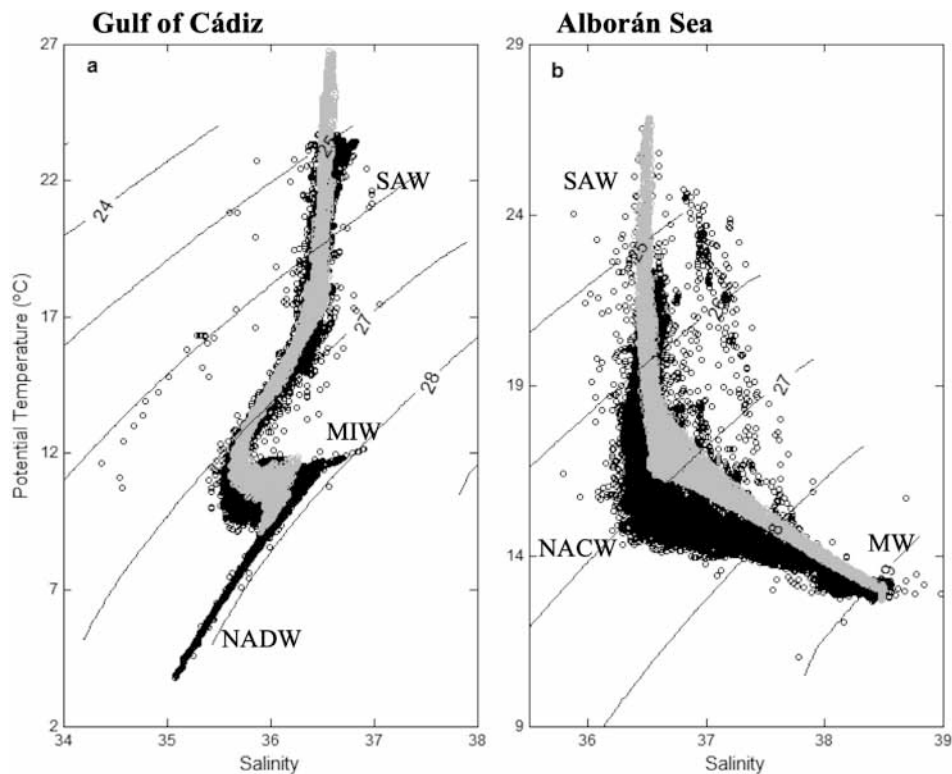


Fig. A.2. Temperature–salinity diagrams comparing the Medatlas II database (black points) with the model outputs (grey points), representing the water mass characteristics of the (a) Gulf of Cádiz region, and the (b) Alborán Sea region. MIW: Mediterranean Intermediate Water. MW: Mediterranean Water. NACW: North Atlantic Central Water. NADW: North Atlantic Deep Water. SAW: Superficial Atlantic Water.

CRedit authorship contribution statement

Iria Sala: Conceptualization, Methodology, Validation, Formal analysis, Investigation, Data curation, Visualization, Writing – original draft, Writing – review & editing. **Sergio M. Vallina:** Conceptualization, Methodology, Validation, Formal analysis, Data curation, Supervision, Writing – review & editing. **Marina Lévy:** Supervision, Writing

– review & editing. **Marina Bolado-Penagos:** Validation, Writing – review & editing. **Carlos M. García:** Conceptualization, Funding acquisition, Project administration, Writing – review & editing. **Fidel Echevarría:** Conceptualization, Funding acquisition, Project administration, Writing – review & editing. **José C. Sánchez-Garrido:** Methodology, Formal analysis, Data curation, Supervision, Writing – review & editing.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

Data will be made available on request.

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Appendix A. Validation of the circulation model

Sea Surface Temperature (SST). SST data for the study area and the period 2001–2010 were downloaded from the Copernicus Marine Environmental Monitoring Service (CMEMS; <http://marine.copernicus.eu/>). This product is based on the reprocessed Pathfinder V5.3 (PFV53) Advanced Very-High-Resolution Radiometer (AVHRR) data combined with a bias-corrected version of the CMEMS-NRT-L4 data, using an Optimal Interpolation algorithm to provide a full-time series of consistent daily gap-free maps with a resolution of ~ 4.63 km.

Fig. A.1 shows the comparison between the measured and modelled annual climatological values of SST. The climatological SST distribution is similar in both satellite data and the model, with colder coastal waters and warmer open ocean regions at both the Gulf of Cádiz and the Alborán Sea. Furthermore, it should be highlighted the presence of colder waters in the coastal region of Trafalgar, our area of interest. However, the model overestimates the annual mean satellite SST by ~ 2 °C.

Temperature–salinity diagrams. Temperature and salinity data for the study area were extracted from the Medar/Medatlas II project, whose objective is to make available a comprehensive data product of temperature, salinity and biochemical data in the Mediterranean and the Black Sea, through wide cooperation of the Mediterranean countries (<http://www.ifremer.fr/medar/>). Temperature and salinity data were collected from 1911 to 1987. Temperature–salinity diagrams were computed for the centre of both sub-basins, the Gulf of Cádiz and the Alborán Sea, from observed and modelled data (**Fig. A.2**). The main water masses present in both sub-basins were quite well reproduced by our regional model (grey points).

Table B.1

List of ecological model parameters. The values are shown in units of nitrogen. The corresponding values for phosphorus, silica and iron can be derived using a Redfield N:Si:P:Fe ratio of 16:16:1:0.001 (mol mol⁻¹). Dia.: Diatoms. DIM: dissolved inorganic matter. Din.: Dinoflagellates. DOM: dissolved organic matter. Mes.: Mesozooplankton. Mic.: Microzooplankton. n.d.: no dimensions. Phy.: Phytoplankton. POM: particulate organic matter. Pro.: *Prochlorococcus*. Syn.: *Synechococcus*. Zoo.: Zooplankton.

Parameter		Symbol	Value	Units
Phy. Pro.	Max. specific growth rate	μ_{max}^{pro}	0.50	d ⁻¹
Phy. Syn.	Max. specific growth rate	μ_{max}^{syn}	1.00	d ⁻¹
Phy. Fla.	Max. specific growth rate	μ_{max}^{fla}	2.00	d ⁻¹
Phy. Dia.	Max. specific growth rate	μ_{max}^{dia}	4.00	d ⁻¹
Phy. Pro.	Half-sat. for DIM uptake	ks^{pro}	0.10	μM
Phy. Syn.	Half-sat. for DIM uptake	ks^{syn}	0.30	μM
Phy. Fla.	Half-sat. for DIM uptake	ks^{fla}	0.60	μM
Phy. Dia.	Half-sat. for DIM uptake	ks^{dia}	1.20	μM
Phy.	Assim. efficiency	β_{phy}	0.80	n.d.
Phy.	Mortality specific rate	m_{phy}	0.05	d ⁻¹
Phy.	Mortality fraction to POM	ω_{phy}	0.50	n.d.
Phy.	Non-assim. fraction to POM	ψ_{phy}	0.50	n.d.
Zoo. Mic.	Max. specific ingestion rate	g_{max}^{mic}	4.00	d ⁻¹
Zoo. Mes.	Max. specific ingestion rate	g_{max}^{mes}	2.00	d ⁻¹
Zoo. Mic.	Half-sat. for ingestion	kg_{mic}	0.50	μM
Zoo. Mes.	Half-sat. for ingestion	kg_{mes}	0.30	μM
Zoo. Mic.	Preference for Small Phy.	ρ_{11}	1.00	n.d.
Zoo. Mes.	Preference for Small Phy.	ρ_{12}	1.00	n.d.
Zoo. Mic.	Preference for Large Phy.	ρ_{21}	1.00	n.d.
Zoo. Mes.	Preference for Large Phy.	ρ_{22}	1.00	n.d.
Zoo.	Assim. efficiency	β_{zoo}	0.40	n.d.
Zoo.	Mortality specific rate	m_{zoo}	0.05	d ⁻¹
Zoo.	Mortality fraction to POM	ω_{zoo}	0.50	n.d.
Zoo.	Non-assim. fraction to POM	ψ_{zoo}	0.50	n.d.
POM	Sinking rate	$ w $	0.50	m d ⁻¹
POM	Degradation rate to DOM	m^{POM}	0.10	d ⁻¹
DOM	Degradation rate to DOM	m^{DOM}	0.10	d ⁻¹

Appendix B. Ecosystem model: Ecological parameters

See **Table B.1**.

Appendix C. Validation of the ecological model

Following **Lefèvre et al. (2002)**, modelled chlorophyll was computed applying a variable C:Chl ratio calculated as a function of light levels following:

$$R_t = R_{max} - (R_{max} - R_{min}) \cdot \left(\frac{I_{max} - I_t}{I_{max} - I_{mix}} \right), \quad (C.1)$$

R represents the C:Chl ratio. R_{max} and R_{min} vary depending on phytoplankton size ($R_{max}^{sma} = 200.00$ mg C mg Chl⁻¹; $R_{min}^{sma} = 166.67$ mg C mg Chl⁻¹; $R_{max}^{lar} = 111.11$ mg C mg Chl⁻¹; $R_{min}^{lar} = 90.91$ mg C mg Chl⁻¹).

I is the photosynthetically active radiation (PAR; $\mu\text{Ein m}^{-2} \text{s}^{-1}$) at the sea surface, derived from ERA-Interim and provided by the European Center for Medium-range Weather Forecast (ECMWF; <https://www.ecmwf.int/>; **Dee et al., 2011**). I_{max} , I_{min} and I_t , which represents the daily PAR, are obtained for each point of the model grid and at noon.

Appendix D. Eastward time-steps during neap and spring tides

See **Figs. D.1–D.3**.

Appendix E. Analysis of the zonal component of the wind at 10 m

See **Fig. E.1**.

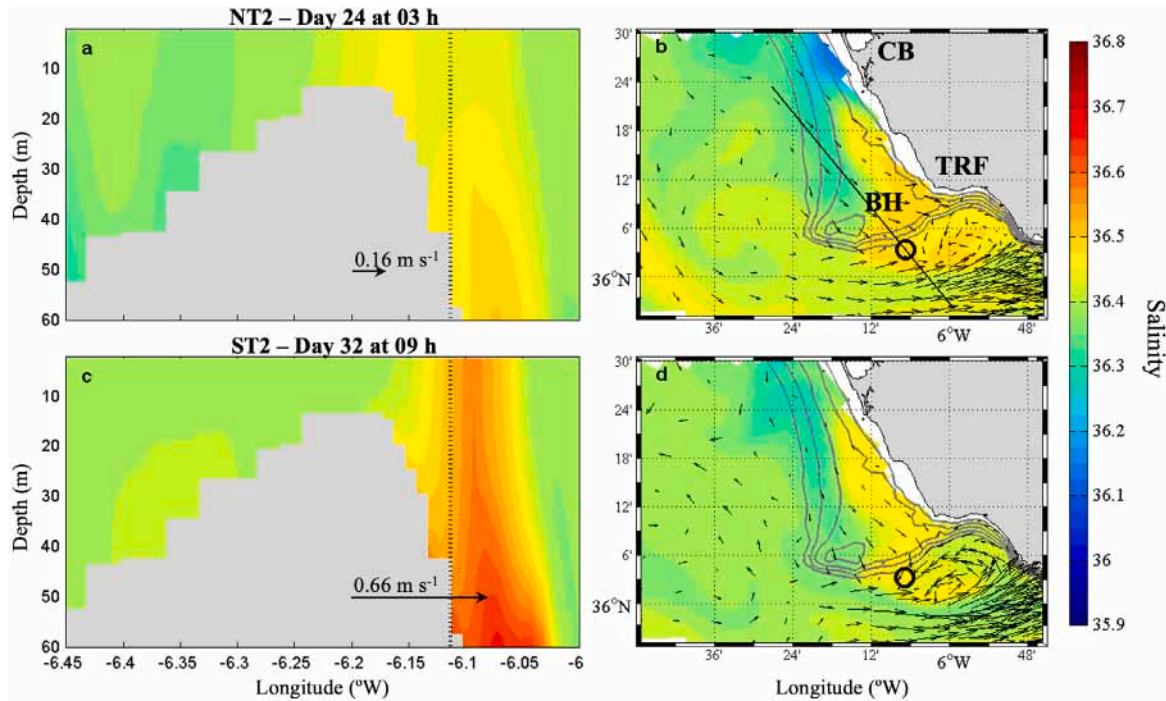


Fig. D.1. Salinity distribution at the Cape Trafalgar region during the maximum eastward flow of the neap tide period (NT2: a, b) and the spring tide period (ST2: c, d) (see Fig. 3). Left-hand panels show the vertical distribution along a section perpendicular to the submarine ridge (black line in panel b). Vertical dashed lines correspond to the position of open black circles in right-hand panels. Right-hand panels show the surface distribution. Black arrows represent the magnitude and direction of the mean zonal component of current velocity (uvel; m s^{-1}), computed for the whole water column at 6.11°W and 36.06°N at each time-step. Grey solid lines represent the bathymetry (10, 20, 30, 40 and 50 m depth). BH: Barbate High. CB: Cádiz Bay. TRF: Cape Trafalgar.

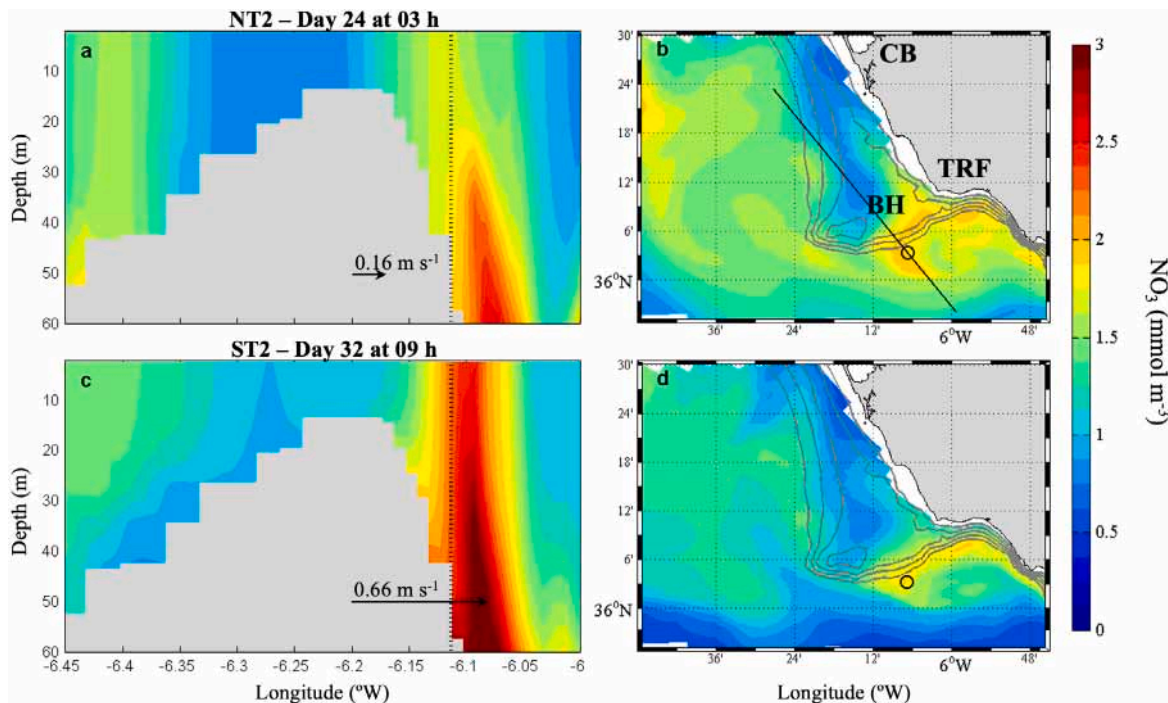


Fig. D.2. Nitrate concentration (NO_3 , mmol m^{-3}) distribution at the Cape Trafalgar region during the maximum eastward flow of the neap tide period (NT2: a, b) and the spring tide period (ST2: c, d) (see Fig. 3). Left-hand panels show the vertical distribution along a section perpendicular to the submarine ridge (black line in panel b). Vertical dashed lines correspond to the position of open black circles in right-hand panels. Right-hand panels show the surface distribution. Grey solid lines represent the bathymetry (10, 20, 30, 40 and 50 m depth). BH: Barbate High. CB: Cádiz Bay. TRF: Cape Trafalgar.

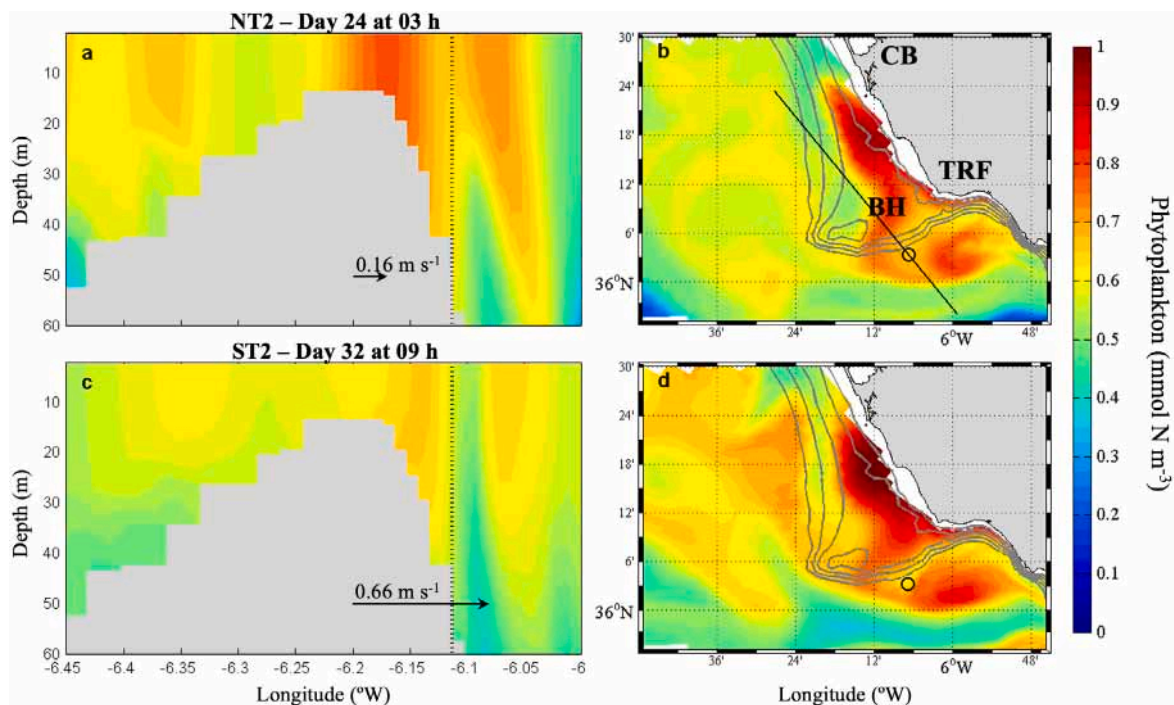


Fig. D.3. Total phytoplankton biomass (mmol N m^{-3}) distribution at the Cape Trafalgar region during the maximum eastward flow of the neap tide period (NT2: a, b) and the spring tide period (ST2: c, d) (see Fig. 3). Left-hand panels show the vertical distribution along a section perpendicular to the submarine ridge (black line in panel b). Vertical dashed lines correspond to the position of open black circles in right-hand panels. Black arrows (a, c) represent the magnitude and direction of the mean zonal component of current velocity (uvel; m s^{-1}), computed for the whole water column at 6.11°W and 36.06°N at each time-step. Right-hand panels show the surface distribution. Grey solid lines represent the bathymetry (10, 20, 30, 40 and 50 m depth). BH: Barbate High. CB: Cádiz Bay. TRF: Cape Trafalgar.

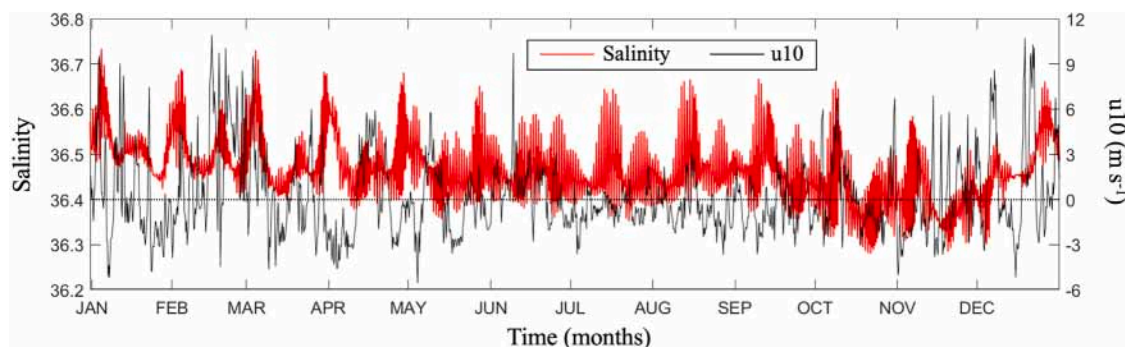


Fig. E.1. Temporal evolution of salinity versus the zonal component of the wind at 10 m (u_{10} , m s^{-1}) every 6 h at 6.11°W and 36.06°N (red circle in Fig. 1b). The black horizontal line represents the zero u_{10} reference delimiting positive (negative) values that indicate westerly (easterly) winds.

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