Succession of phytoplankton assemblages in relation to estuarine hydrodynamics in the Ría de Vigo: A box model approach*

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SUMMARY: Two phytoplankton succession events (September-October 1993 and March 1994) in the Ría de Vigo were studied in relation to their hydrodynamic context. By means of a circulation box model we calculated horizontal and vertical residual fluxes as well as budgets of phytoplankton associated with them. This allowed us to assess the importance of the two factors involved in the selection processes: one hydrodynamic (dependent upon physical dispersion) and one biogeochemical (involving all other processes such as growth, mortality and grazing). The first period was characterised by a downwelling event which produced a reversal of the estuarine circulation pattern (horizontal velocity = 4 km d⁻¹ and downward velocities up to 9 m d⁻¹). This provided suitable conditions for the blooming of Gymnodinium catenatumfavoured by its capacity to swim, while the diatom community almost disappeared. The second period corresponded to an upwelling → relaxation sequence. During upwelling (horizontal velocity = 6 km d⁻¹; upward velocities up to 12 m d⁻¹), Thalassiosira spp. was dominant. It is suggested that the less energetic conditions associated with the relaxation event (horizontal velocity=1 km d⁻¹; upward velocities < 1 m d⁻¹) favoured the selection of Chaetoceros spp. due to its lower sinking rate.

Key words: estuarine circulation, succession, box model, Gymnodinium catenatum, Chaetoceros spp., Thalassiosira spp., Ría de Vigo.

INTRODUCTION

The hydrodynamic regime has a direct effect on the distribution of populations of phytoplankton since it implies a biomass advection (the so-called ‘physical dispersion’ in Painchaud et al., 1996). In addition to the hydrodynamic factor, the distribution is also determined by many other processes, which we will hereafter call “biogeochemical”. These processes include growth, mortality and grazing, passive sedimentation and swimming, which are highly dependent on the hydrodynamic conditions. The Ría de Vigo (NW coast of Spain, Fig. 1) constitutes a good example of this connection since its high biological productivity is remarkably affected by estuarine hydrodynamics (Prego, 1993; Nogueira et al., 1997; Tilstone et al., 1999). According to Dyer (1973) and Beer (1983), the Ría de Vigo can be considered as a partially mixed positive estuary, which means that typical residual transport of water along the estuary is two-layered (Prego et al., 1990). There is an inward more saline flow in the bottom layer and a seaward flow in the upper one, with the development of a zero velocity depth between them (see Fig. 2).
winds which are especially common during spring and summer lead to the upwelling of ENACW (Eastern North Atlantic Central Water), which enhances the residual circulation in two layers as it lifts cold and nutrient-rich water (Prego and Fraga, 1992) towards the Ría. Southerly winds, on the other hand, generate downwelling, forcing surface low-nutrient waters towards the head of the estuary where they pile and eventually sink, which can bring about the reversal of residual circulation (Rosón et al., 1997). Further descriptions of the hydrodynamic regime are provided in Torres López et al. (2001) and Souto et al. (2001). In general terms, a shift from diatom to dinoflagellate populations associated with the transition from upwelling (turbulence and nutrient abundance) to downwelling conditions (stratification and depletion of nutrients) can be observed (Margalef et al., 1955; Margalef, 1958; Figueiras and Pazos, 1991; Figueiras and Ríos, 1993).

For all this apparent connection, there is a lack of studies linking estuarine hydrodynamics and phytoplankton dynamics, partly due to the complexity of the system, and in general descriptive viewpoints have been maintained. To overcome this difficulty, box model approaches such as the one we propose here are available. These types of models have been applied to the study of pollutants and nutrient cycles (Prego, 1992; Álvarez-Salgado et al., 1996; Simpson and Rippeth, 1998; Rosón et al., 1999), yet recently they have also proven to be a very useful tool for gaining insight into bacterial (Painchaud et al., 1996) and phytoplankton dynamics (Tilstone et al., 2000).

By means of a 2-D box model we will first make a quantitative description of residual circulation in the Ría de Vigo during two periods corresponding to very different contexts. The first period corresponded to a typical autumn-downwelling situation characterised by a bloom of the red-tide dinoflagellate, Gymnodinium catenatum, which developed as the diatom community virtually disappeared. The second period corresponds to the relaxation of the upwelling conditions, with an interesting shift between two genera of diatoms, Thalassiosira spp. and Chaetoceros spp. Furthermore, we will estimate the advection of these groups of phytoplankton and compare it with the net budget of their standing stock. The box model results will allow us first to separate and then to assess both the hydrodynamic and the biogeochemical effect on the evolution of phytoplankton communities. This constitutes the most innovating contribution of this work.

MATERIAL AND METHODS

The Ría de Vigo was sampled 6 times in approximately 2 weeks corresponding to 2 different periods (September 27 to October 8, 1993 and March 6 to March 24, 1994). During these periods, 4 stations were sampled (Fig. 1) aboard the “Lampadena” vessel at the same tidal cycle. A CTD-fluorescence sampler was used to obtain S-T profiles and also to determine the depths where up to 5 seawater samples were to be collected, using 5-litre Niskin bottles fitted with reversing thermometers. Aliquots were taken from the Niskin bottles and frozen for the analysis of nutrients in the laboratory using a Technicon AAII SFA auto analyser. The method of reduction to nitrites in a Cd-Cu column was used to determine nitrate concentration (Mouriño and Fraga, 1985). Phytoplankton samples were preserved in Lugol’s iodine, sedimented in 50 ml composite sedimentation chambers and eventually counted to the species level.

The Ekman cross-shore (Q_x) and along-shore (Q_y) transport component was calculated according to Wooster et al. (1976).

\[ Q_{x,y} = \frac{\rho_a C_d |V|}{\rho_w} \]  

(1)

where \( \rho_a \) is the air density (1.22 kg m\(^{-3}\)), \( C_d \) is the empirical drag coefficient (1.35 \( \times \) 10\(^{-3}\), according to Hidy, 1972), \( |V| \) is the vector corresponding to the estimated wind speed on the sea surface, with magnitude \(|V| f (9.946 \times 10^{-5} \text{s}^{-1})\) is the Coriolis parameter, and \( \rho_w \) the density of sea water (\( \rho_w \sim 1025 \text{ kg m}^{-3} \)).

Fig. 1. – Study area (Ría de Vigo, NW of the Iberian Peninsula), sampling stations and boxes. Box dimensions are given in the table.
The cross-shore transport \( (Q_x) \) is associated with winds that have a north-south component and is related to upwelling and downwelling processes which can eventually cause the enhancement or reversal of the residual circulation, as mentioned in the introduction. The alongshore component \( (Q_y) \) depends on winds with a predominant east-west component; these winds, when strong enough, can directly pull surface water into or out of the Ría and likewise alter the residual circulation pattern.

The mean integrated Brunt-Väisälä frequency \( (N^2) \), an estimate of the stability of the water column (Millard et al., 1990), was calculated according to Doval et al. (1997).

\[
N^2 = \frac{g}{z} \ln \left( \frac{\rho_s}{\rho_b} \right)
\]  

(2)

where \( g \) is gravity, \( z \) is water depth, \( \rho_s \) is surface density and \( \rho_b \) is bottom density.

**Box model of residual circulation and phytoplankton**

**Residual circulation**

Fluxes were derived from the salinity and temperature profiles using an improved 2-dimensional kinematic non-steady state mass/heat weighted box model similar to the one developed and extensively described by Rosón et al. (1997) for a similar estuarine system. The calculations are based on the mass and energy conservation principle applied to 2 conservative properties, salinity and temperature, inside closed elements or boxes defined inside the Ría on the hypothesis that changes of these properties within such elements are exclusively due to exchanges across their boundaries, and assuming that the volume of the boxes is constant over a tidal cycle (Officer, 1980).

As Figure 1 shows, the Ría de Vigo was divided into 3 boxes limited by a series of walls that were perpendicular to its axis and contained sampling stations. On the basis of a 2-layered circulation pattern these boxes were segmented into 2 levels (upper and bottom) separated by a zero-velocity depth \( (Z_c) \). The terms in the Volume-S-T budget for each box are presented in Figure 2. Rainfall data \( (L) \) were obtained from the Meteorological Observatory at Vigo Airport \( (42^\circ 14' \text{ N}, 8^\circ 18' \text{ W}) \) and corrected considering its altitude above sea level. Evaporation \( (E) \) was obtained with an empirical relationship depending on wind velocity and vapour pressure (Rosón et al., 1997). Continental flow \( (Q_r) \) was calculated according to Ríos et al. (1992) using rainfall in the drainage basin. Heat exchange \( (H) \) was solved as a sum of terms (irradiation, atmospheric and oceanic back radiation, conduction and reflection) whose evaluation is thoroughly explained in Rosón et al. (1997). For salinity and temperature mean values, vertical CTD profiles were used and averaged using the geometric characteristics of each box.

For the sake of brevity no thorough report of the mathematical model will be made in this paper and instead we will emphasise the application of the box model to the study of phytoplankton. A more detailed description of the calculation of residual fluxes can be found in Rosón et al. (1997) and in Pardo et al. (2001).

**Phytoplankton dynamics**

Unlike conservative properties, when calculating budgets of phytoplankton we must take into account not only the fluxes crossing the boxes and transporting phytoplanktonic populations (hydrodynamic processes), but also the biogeochemical processes. As mentioned in the introduction, these biogeochemical processes account for the inputs and outputs that occur within the boxes (e.g. growth, respiration, mortality, grazing and sedimentation) and explain the non-conservative behaviour (Smith and Hollibaugh, 1997). Hence, we can express the net temporal change of the standing stock of a certain
group of phytoplankton in a box as the sum of a hydrodynamic component, $\delta N_h$, and a biogeochemical one, $\delta N_{bg}$:

$$\frac{dN_B}{dt} V_B = \delta N_h + \delta N_{bg}$$  \hspace{1cm} (3)

where $V_B$ is the volume of the box, and $dN_B/dt$ is the total temporal change of phytoplankton abundance in a single box for a given day $t_n$, which can be expressed as an average increment between the previous sampling day, $t_{n-1}$, the present sampling day, $t_n$ and the following sampling day, $t_{n+1}$:

$$\frac{dN_B}{dt} = \frac{1}{2} \left( \frac{N_{Bn} - N_{Bn-1}}{t_n - t_{n-1}} + \frac{N_{Bn+1} - N_{Bn}}{t_{n+1} - t_n} \right)$$  \hspace{1cm} (4)

Once we have obtained residual fluxes we can easily calculate the hydrodynamic component rate $\delta N_h$ as shown in (5):

$$\delta N_h = \sum Q_n \cdot N_Q$$  \hspace{1cm} (5)

where $N_Q$ is the phytoplankton abundance carried by each flow $Q_n$.

Finally, after calculating the hydrodynamic component ($\delta N_h$) by (5) and the net temporal change $dN_B/dt$ by (4), the rate of change due to biogeochemical processes ($\delta N_{bg}$) can be simply derived by subtracting both.

$$\delta N_{bg} = \frac{dN_B}{dt} - \delta N_h$$  \hspace{1cm} (6)

Since our intention is to give an insight into the overall selection processes that occur within the system, results to be presented in this paper will refer to the Ría as a whole. Moreover, so as to be able to compare the results obtained for different populations, we will relate both the hydrodynamic and biogeochemical rates found to the total group biomass. Thus, hereafter, $\delta N_h$ and $\delta N_{bg}$ values will express a daily change of biomass in relation to the total group biomass present in the Ría. A 0.5 value means that 50% of the biomass present in the Ría is gained (if positive) or lost (if negative) in one day by the processes mentioned above.

RESULTS

In order to analyse the box-model results it is necessary to consider the meteorological, hydrographic and biological contexts of the samplings, so, a brief draft of these contexts will be included. More
detailed descriptions are available in previous works (Fermín et al., 1996; Figueiras et al., 1996; Tilstone et al., 1999; Tilstone et al., 2000).

September-October 1993

Meteorology, hydrography, nutrients and phytoplankton

The Ekman transport components, wind speed and the continental runoff from 24 September to 8 October are shown in Figure 3. Decreasing negative values of $Q_y$ were minimal on 5 October (-3800 m$^3$ s$^{-1}$ km$^{-1}$) and correspond to westerly winds blowing directly over the Ría. The associated alongshore transport first caused the slow-down of the positive residual circulation pattern and finally favoured its reversal and the introduction of relatively warmer, nutrient-depleted coastal water into the Ría. This was recorded in $T$, $S$ and $NO_3^-$ values (Table 1) on 1 October. The Ría was less stratified and this was reflected by $N^2$ values that were extremely low on 1 October. $Q_y$ values were generally lower (<2000 m$^3$ s$^{-1}$ km$^{-1}$) and acquired some relevance towards the end of the period, favouring the reestablishment of positive residual circulation after 4 October (we have represented $-Q_y$ so that positive values imply upwelling). This reestablishment was also supported by the simultaneous increase in continental runoff, which brought about the cooling, and above all the freshening up, of the upper inner box, along with an

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<th>$S$</th>
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<th>whole Ria per Box</th>
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**FIG. 4.** – Time course of (a) temperature ($T$, °C), (b) salinity ($S$) and (c) nitrate ($NO_3^-$, µmol kg$^{-1}$) profiles at Stn. 3. Arrows indicate sampling dates. Downwelling period (September-October 1993).
increase in NO$_3$ levels. This in turn justifies the higher N$_2$ values in Box 1, which eventually affected the whole Ría. We can observe the same processes in the T, S and NO$_3$ profiles over time for a representative station (Fig. 4). The downwelling event is recorded as a downward displacement of the isopycnals and isotherms from 29 September to 1 October, along with a decrease in nitrate levels. The entrance of coastal water leaves a homogeneous water column, which recovers its stratification at the end of the period, mainly due to the arrival of fresh water.

The last two columns in Table 1 show the selection of Gymnodinium catenatum, whose population grew sharply (by a factor of 20 in 6 days) mainly in the outermost boxes reaching a maximum on October 4. The cell abundance illustrates the dimension of the bloom: G. catenatum outnumbered the totality of the diatom community. Diatoms, on the contrary, were abundant on 29 September (mean value for the Ría = 1694 cells ml$^{-1}$, i.e. volume-averaged for the 3 boxes) but as the downwelling situation progressed, their abundance decreased by 95% (66 cells ml$^{-1}$ on 4 October). The diatom population recovered on 6 October but only in the last box. By contrast the G. catenatum population followed a decreasing pattern on 6 October: its number descended and it became more homogeneously distributed.

**Box model results**

**Residual circulation**

A general description of the flows corresponding to a positive residual circulation pattern is depicted in Figure 2. The corresponding residual surface-averaged velocities were calculated simply by dividing those flows by correspondent boundary wall surfaces and are presented in Figure 5. Arrows follow the direction of the movement and negative values indicate reversal of the residual circulation, generally associated with downwelling.

The horizontal circulation on 29 September 1993 (mean $Q_n$ absolute value for the Ría = 2.0 km d$^{-1}$, i.e. averaged for the upper and bottom walls at the 4 stations) showed the effect of westerly winds, which forced coastal water towards the Ría, blocking the positive estuarine circulation at its mouth. This can be seen in low and negative velocity values found for Box 3. In the inner Ría, however, a positive circulation pattern prevailed (bottom inward and upper outward flows). Both the vertical convective ($Q_z$) and mixing ($M$) velocities were moderate when compared with those of the following days. As westerly winds intensified on 1 October, the enhanced transport of water caused a strong reversal of the estuarine circulation, as can be seen in the external box, and horizontal velocities increased (mean value of 4.0 km d$^{-1}$). Strong negative vertical velocities (mean $Q_z$ value for the Ría = -5.7 m d$^{-1}$, i.e. averaged for the 3 boxes) indicated a net downward transport of water from upper to bottom layers. These nega-
tive velocities were concurrent with a strong deepening of the zero-velocity depth, as would be expected for a downwelling situation. Residual velocities on 4 October indicated the slight moderation of the downwelling observed as a decrease in horizontal (mean absolute value = 0.8 km d\(^{-1}\)) and vertical (mean value = -2.5 m d\(^{-1}\)) convective velocities. Coinciding with the higher runoff, stratification developed in the interior of the estuary (Box 1): the zero velocity level rose and mixing was inhibited (mean \(M\) value for the Ría = 0.7 m d\(^{-1}\)). The reestablishment of a positive circulation pattern took place on 6 October, as is reflected by almost all-positive velocities.

**Phytoplankton dynamics**

The change in biomass associated with both the hydrographic (\(\delta N_h\)) and the biogeochemical component (\(\delta N_{bg}\)) for both the diatom and the *G. catenatum* assemblages in the whole of the Ría is shown in Figure 5 (autumn period). Notice the differences between absolute values of both components, \(\delta N_h\) always being lower than \(\delta N_{bg}\). This highlights the importance of biogeochemical processes occurring within the Ría during this downwelling period. Physical dispersion processes associated with residual flows, on the contrary, did not have such a marked effect on the variation of phytoplanktonic communities. On 29 September \(\delta N_h\) positive values evidenced a net entrance of diatoms and *G. catenatum* associated with water transport, which represented respectively 4% and 9% of their standing stock. Under downwelling conditions (1 and 4 October), \(\delta N_h\) values kept relatively low and positive. On 6 October, there was a net hydrodynamic exportation of *G. catenatum*, as shown by the negative \(\delta N_h\) value (= -0.18).

The rate of change that is not dependent upon physical dispersion, \(\delta N_{bg}\), showed much higher values. Diatom abundance decreased 21% due to biogeochemical processes on 29 September, whereas *G. catenatum* assemblage accumulated and its standing stock doubled. On 1 October, when the circulation was inverted, the differences persisted: the diatom abundance diminished sharply (68% of the total cell abundance was lost) and the *G. catenatum* population increased by 45%. Negative \(\delta N_{bg}\) values were found for both communities on 4 October, although the loss was negligible in *G. catenatum* (\(\delta\)

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**Fig. 6.** – Ekman transport components, wind speed and direction calculated at 43°N 11°W and continental runoff in the Ría de Vigo. Arrows indicate sampling dates. Upwelling-relaxation period (March 1994).
$N_{bg} = -0.07$) as compared to the diatom community ($\delta N_{bg} = -1.00$). On 6 October loss processes could still be detected as negative $\delta N_{bg}$ values for both groups involved in the study.

### March 1994

**Meteorology, hydrography, nutrients and phytoplankton**

The Ekman transport components (Fig. 6) on days previous to the sampling showed the influence of decreasing northeasterly winds, which eventually changed to south-west and caused the relaxation of a previous upwelling period. After 9 March the winds turned back to the north-east, which caused upwelling. This was detected in the Ría from 11 to 16 March and can be seen in the increase in salinity and NO$_3^-$ levels in the 3 boxes, whereas temperature values decreased (Table 2). Notice the uplifting of the T, S and NO$_3^-$ isolines for the same period (Fig. 7). From 18 March on, the influence of westerly winds was recorded as a reduction in $Q_x$ and $Q_y$ values, which again produced the relaxation of the upwelling towards the end of the period. This can be seen in the decreasing NO$_3^-$ levels on 22 March, while salinity and temperature varied slightly. Continental runoff showed very low values (decreasing from 14 to 5 m$^3$ s$^{-1}$) and had little influence on the circulation pattern.

### Table 2.

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Fig. 7. – Time course of (a) temperature (T, °C), (b) salinity (S) and (c) nitrate (NO$_3^-$, µmol kg$^{-1}$) profiles at Stn. 3. Upwelling-relaxation period (March 1994).
The stratification increased at the end of the sampling period, as we can see in the salinity and temperature profiles at Station 3 (Fig. 7).

The phytoplankton abundance values presented in Table 2 show a succession event between two genera of diatoms. On 11 March Thalassiosira spp. was more abundant than Chaetoceros spp. (values for the whole Ría), especially in Box 3. However, from 14 March Thalassiosira spp. standing stock followed a decreasing trend, whereas Chaetoceros spp. abundance increased by a factor of 3 during the same sampling period (from 347 to 1136 cells ml⁻¹).

**Box model results**

*Residual circulation*

Unlike the autumn situation, a positive circulation pattern was present through the whole March sampling period (Fig. 8). The relatively low horizontal and vertical fluxes on 11 March 1994 (mean velocity, 3.2 km d⁻¹ and 2.6 m d⁻¹ respectively) might be regarded as a consequence of the lack of continuity of the winds blowing on the previous days. The highest horizontal residual velocities in the Ría were observed on 14 March (mean velocity, 6.0 km d⁻¹), indicating upwelling. The uplifting of deep water brought about the intensification of the circulation, which was also apparent in the enhanced vertical upward transport (maximum velocity, 14.0 m d⁻¹ in Box 3) and mixing (up to 7.1 m d⁻¹ in Box 2). Values on 16 March were typical of an upwelling scenario too, but slightly lower. The circulation was slower than on the previous day and only mixing increased, especially in the inner box (mean value 11.1 m d⁻¹), a process presumably facilitated by the already eroded stratification. The horizontal velocities found on 22 March were substantially lower (mean velocity for the Ría, 1.2 km d⁻¹) and corresponded to an upwelling relaxation event (Qₓ and Qᵧ values close to zero). As the pumping ceased, mixing and vertical transport diminished (mean values, 1.8 and 0.9 m d⁻¹ respectively).

*Phytoplankton dynamics*

On 11 March δNₓ negative values expressed the initial exportation of Thalassiosira spp. and Chaetoceros spp. communities, which means that 10% and 15% of the respective biomass was lost through advection. The loss was particularly strong on 14 March (79% for Thalassiosira spp. and 56% for Chaetoceros spp.), when the upwelling was more intense. As the upwelling weakened on 16 March, a balance between hydrodynamic losses and gains was established (as indicated by low δNₓ values). Finally, on March 22 there was an input of Thalassiosira spp. biomass (representing 11% of the total biomass present in the Ría) while almost no change was recorded for Chaetoceros spp.

The effect of biogeochemical processes on the two diatom assemblages is expressed by δN(bg) values. Beginning on 11 March and especially on 14 March...
March, our results show the accumulation of *Thalassiosira* spp. and *Chaetoceros* spp. biomass in the Ría. On 14 March *Thalassiosira* spp. standing stock increased by 54% regardless of physical dispersion processes, whereas that of *Chaetoceros* spp. increased by 61%. The greatest differences arose in the second part of the sampling period. Thus, while *Chaetoceros* spp. accumulation continued, the biogeochemical rate of change for *Thalassiosira* spp. (negative on 16 and 22 March) indicated a strong loss process.

**DISCUSSION**

Separating the effects of hydrodynamic and biogeochemical processes on phytoplankton assemblages becomes a difficult issue in upwelling zones due to the tight coupling between these two processes. In general, hydrographic data and phytoplankton species distributions can merely provide a static description, which is then used to infer the effect of hydrodynamic processes on the evolution of the phytoplankton populations. Box models can supply more quantitative information so that a more dynamic view can be achieved. In this paper we used an improved 2-D kinematic box model conceived to describe the residual circulation and calculate flows of phytoplankton. The box model results allowed us to estimate the relative influence of the hydrodynamic factors (related to movements of water) and biogeochemical factors (comprising all other processes such as growth, mortality, passive sedimentation and grazing) on the succession events which characterised the sampling periods.

**September-October 1993**

Residual fluxes at the beginning of the autumn period (29 September) showed the blockage of the circulation caused by south-western winds. Such hydrodynamic conditions hardly brought about any variation in diatom and *G. catenatum* abundance linked to water transport (\(\delta N_{\text{b}}\) close to zero). Nonetheless, the hydrodynamic context might help to explain the differences between the biogeochemical rates of change found for the two phytoplankton groups. The relatively stagnant conditions did not favour the permanence of the initially abundant diatom community, which, due to its higher aggregation and sedimentation rates (Smetacek, 1985; Passow, 1991), sank in the water column towards levels of low irradiance and eventually died. Conversely, the growth of another red tide species such as *G. catenatum* is facilitated by low turbulence and stratification (Figueiras and Rios, 1993; Fermín et al., 1996) and might explain the high positive \(\delta N_{\text{b}}\) value. The intensification of the downwelling on 1 October brought about the reversal of the circulation, and a downward displacement of water associated with a homogenisation and a nutrient depletion (Tilstone et al., 1999) of the water column, which again hindered the development of the diatoms. *G. catenatum*, conversely, has shown itself capable of counteracting these displacements due to its ability to migrate vertically (Fraga et al., 1989; Figueiras et al., 1994), which is favoured by the weakness of the physical gradient (Fermín et al., 1996). In addition, the hydrodynamic variation rate is also positive, which supports the accumulation of these dinoflagellates in the Ría and their selection. On 4 October, downwelling conditions persisted and the biogeochemical rate of change accounted for the disappearance (due to mortality and grazing) of the diatom community, which occupied a poorly irradiated and nutrient depleted medium (Tilstone et al., 1999). Red-tide events appear associated with coastal downwelling and have been related to open ocean blooms which were later introduced to the Ría by the reversal of circulation (Fraga et al., 1988; Fraga et al., 1989; Tilstone et al., 1994). Other authors, however, have pointed out that dinoflagellates were able to develop inside the Ría (Figueiras and Pazos, 1991; Figueiras et al., 1994; Fermín et al., 1996) though they have a lower growth rate than diatoms (Banse, 1982). Up to now the matter remained unsolved due to the difficulties involved in estimating growth rates in the field and assessing the effect of the circulation pattern. On that score our results showed that the development of the red-tide forming dinoflagellate, *G. catenatum* was not exclusively a consequence of water inward transport, but mainly of the *in situ* growth of the standing stock inside the Ría (positive \(\delta N_{\text{b}}\) values on 29 September and 1 October). Red tide species are commonly assumed to be relatively slow growing. However, Smayda (1997) suggests that some dinoflagellates are capable of rapid growth and have growth rates for the genera *Gymnodinium* which are compatible with the \(\delta N_{\text{b}}\) values we found. On 6 October, the reestablishment of the positive estuarine circulation and the supply of nutrients propitiated the relatively higher accumulation of diatoms while *G. catenatum* was displaced out of the Ría. Horizontal and vertical
velocities were similar to those found in previous works for periods with similar hydrographic conditions (Rosón et al., 1997; Villarino et al. 1995).

March 1994

Environmental conditions were clearly different during the March sampling period, with the predominance of an upwelling event, which intensified on 14 March and then relaxed. Upwelling lifted colder and richer waters towards the Ría. This led to the enhancement of positive residual circulation and the exportation of biomass, as can be inferred from negative values of $\delta N$. It also involved availability of nutrients, and enough turbulence to maintain phytoplankton populations in the upper layer. These suitable conditions for growth help to explain the high accumulation rates $\delta N$ found for both genera. Horizontal and vertical fluxes were reduced on March 16 and 22, indicating relaxation of upwelling simultaneously to the shift between Chaetoceros spp. and Thalassiosira spp. The new, less energetic, hydrodynamic conditions involved a reduction in the rate of biomass loss through exportation and, above all, turned out to be much less favourable for the development of Thalassiosira spp. On the other hand, Chaetoceros spp. kept accumulating through biogeochemical processes, yet at a lower rate. Although the two genera coexist in upwelling systems (Hood et al., 1991; López-Jamar et al., 1992), in this case they show very different responses, probably due to their different morphological features (Margalef, 1978). Centric diatoms such as Thalassiosira spp. form large chains by cytoplasmatic extensions which favour their aggregation and sedimentation (Riebsell, 1989; Passow, 1991; Alldredge et al. 1995), probably also aided by the high cellular abundance during the first sampling days. By contrast, the long aerolated setae characteristic of Chaetoceros spp. can facilitate its flotation by causing a spiral rotation of the diatom chain (Goldsmith, 1966), allowing them to remain in the photic zone. Former work in the area has shown that Chaetoceros spp. are selected during upwelling-downwelling relaxation events coinciding with high primary production values in the Ría (Tilstone et al. 1999).

It is worth noting that although the selection of Gymnodinium catenatum and Chaetoceros spp. appeared to happen mainly through biogeochemical processes, these processes can only be fully understood if we take into account the hydrodynamic phenomena that are related to downwelling and upwelling-relaxation. Using the box model to differentiate between hydrodynamic and biogeochemical factors leads us once again to stress the tight relationship between the two.

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