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Influence of Rossby waves on primary production from a coupled physical-biogeochemical model in the North Atlantic Ocean

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Abstract

How do Rossby waves influence primary production in the North Atlantic Ocean? Rossby waves have a clear signature on surface chlorophyll concentrations which can be explained by a combination of vertical and horizontal mechanisms (reviewed in Killworth et al., 2004). In this study, we aim to investigate the role of the different physical processes to explain the surface chlorophyll signatures and the consequences on primary production using a 3-D coupled physical/biogeochemical model for the year 1998.

The analysis at 20 given latitudes, mainly located in the subtropical gyre, where Rossby waves are strongly correlated with a surface chlorophyll signature, shows that vertical and horizontal processes are involved in the surface chlorophyll anomalies. Furthermore, the ecosystem response is, as expected, stronger when vertical input of dissolved inorganic nitrogen is observed.

The surface chlorophyll anomalies, induced by these physical mechanisms, have an impact on primary production. We then estimate that Rossby waves induce, locally in space and time, increases (generally associated with the wave crest) and decreases (generally associated with the wave trough) in primary production (∼±20% of the estimated primary production). This symmetrical situation suggests a net weak effect of Rossby waves on primary production.

1 Introduction

The detection of westward propagating signals in surface chlorophyll concentrations related to Rossby waves (e.g. Machu et al., 1999; Cipollini et al., 2001; Uz et al., 2001; Kawamiya and Oschlies, 2001) prompted the question of the underlying physical/biogeochemical interactions. Based on remotely sensed data and/or coupled physical/biogeochemical modelling, several studies investigated the coupled physical/biogeochemical mechanisms which might be involved (e.g. Charria et al., 2003; Killworth et al., 2004; Charria et al., 2006a). Three main processes were suggested:
(1) the upwelling mechanism associated with nutrient injection (Cipollini et al., 2001; Uz et al., 2001; Siegel, 2001), (2) the uplifting of a deep chlorophyll maximum towards the surface (Cipollini et al., 2001; Kawamiya and Oschlies, 2001; Charria et al., 2003), and (3) the meridional advection of chlorophyll by geostrophic currents associated with baroclinic Rossby waves (Killworth et al., 2004). These three processes are described using theoretical models and compared to the remotely sensed observations in Killworth et al. (2004). These authors pointed out the importance of the third process at the global scale. More recently, Charria et al. (2006a) showed a significant contribution of the vertical process of nutrient injection in the North Atlantic, north of 28° N, using remotely sensed data and theoretical models from Killworth et al. (2004). However, in the two latter studies, several assumptions (for example: a constant relaxation time for biology and a constant \( \frac{\text{Chl}}{N} \) ratio) were made. We propose here to investigate the relative contribution of these mechanisms using a more realistic 3-D coupled physical/biogeochemical modelling approach.

Furthermore, the influence of Rossby waves on biogeochemical processes has to be estimated for a better understanding of the carbon cycle through the biological pump mechanism. Kawamiya and Oschlies (2001) have found a ∼30% increase in primary production with the passage of a Rossby wave in the Indian ocean around 10° S using a coupled physical/biogeochemical model. Near the Hawaiian Ocean Time Series (HOT) station ALOHA in the Pacific Ocean, a combination of in situ high-frequency measurements (shipboard and moored sampling) and remotely sensed data (altimetry) during a long period (1997–1999) allowed Sakamoto et al. (2004) to estimate primary production enhancement up to 25% on average due to the Rossby wave’s passage. Indeed, Rossby waves can significantly contribute to supply nutrients to oligotrophic surface waters fuelling primary production. According to several authors (e.g. Oschlies and Garçon, 1998; Oschlies, 2002a,b), eddies induce increase in nutrient input into the euphotic layer. Vertical velocities associated with Rossby waves can induce a similar effect, named as the ”Rossby rototiller” by Siegel (2001), with the crucial difference that while eddies upwell water only while forming or intensifying, Rossby waves would
upwell nutrients all along their propagation path through an ocean basin. This effect is then potentially comparable to basin-scale processes as induction process in the subpolar gyre (Williams et al., 2006) or Dissolved Organic Nitrogen and Phosphorus advection in the subtropical gyre (Roussenov et al., 2006).

The present work aims at investigating Rossby wave influence on primary production using a 3-D coupled physical/biogeochemical realistic model in the North Atlantic Ocean. After a brief description of the coupled physical/biogeochemical model in Sect. 2, as well as the remotely sensed data used in the validation of simulated wave features in the model (Sect. 3), we will show the surface chlorophyll and physical signatures of Rossby waves using model results. In Sect. 4, the features of these modelled propagating signals are identified and compared to those estimated with remotely sensed data. In Sect. 5, the procedure to extract the studied regions is given followed by a results description in Sect. 6. The influence of Rossby waves on modelled primary production and the relative contribution of the underlying physical/biogeochemical processes in the oligotrophic gyre are finally discussed in the last section.

2 A coupled physical/biogeochemical model

The numerical experiments described here are performed using a 3-D coupled physical/biogeochemical model. The ocean physics is solved by the OPA 8.1 model (Madec and Imbard, 1996) in a North Atlantic configuration (MNATL – 20° S–70° N and 98.5° W–20° E). This model, initially used in the frame of the CLIPPER project and used by the operational oceanography MERCATOR project, has a 1/3° spatial resolution and 43 vertical levels with a vertical grid spacing increasing from 12 m at the surface to 200 m at the bottom. Simulations are forced by daily European Centre for Medium-Range Weather Forecasts (ECMWF) atmospheric forcing.

This physical model is coupled with a simple nitrogen-based NPZDDON (dissolved inorganic Nitrogen, Phytoplankton, Zooplankton, Detritus and Dissolved Organic Nitrogen) ecosystem model (Huret et al., 2005). The different fluxes between state vari-
ables, represented on Fig. 1 are the following: (1) Nutrients assimilation by phytoplankton during the photosynthesis, (2) Zooplankton grazing, (3) Zooplankton excretion, (4) Zooplankton mortality, (5) Non-assimilated part of zooplankton grazing (fecal pellets production), (6) Phytoplankton exudation, (7) Particles hydrolysis by bacteria, (8) Phytoplankton mortality, (9) Dissolved organic nitrogen remineralization and (10) Organic zooplankton excretion.

The tracer concentrations follow an advective-diffusive equation plus a source-minus-sink (SMS) term of exchange between different tracers. The SMS terms for each of the biological tracer in nitrogen units (mmolN m\(^{-3}\)) are:

\[
\text{SMS}(N) = \rho \text{DON} + (1 - f_2)\gamma Z - J(z, t, N)P \tag{1}
\]

\[
\text{SMS}(P) = (1 - \epsilon)J(z, t, N)P - \mu_P P - G(P)Z \tag{2}
\]

\[
\text{SMS}(Z) = f_1 G(P)Z - \gamma Z - \mu_Z Z^2 \tag{3}
\]

\[
\text{SMS}(D) = (1 - f_1)G(P)Z + \mu_P P + \mu_Z Z^2 - \mu_D D - w_S \frac{\partial D}{\partial z} \tag{4}
\]

\[
\text{SMS(DON)} = \epsilon J(z, t, N)P + f_2 \gamma Z + \mu_D D - \rho \text{DON} \tag{5}
\]

N represents dissolved inorganic nitrogen, P phytoplankton, Z zooplankton, D particulate organic matter and DON the dissolved organic nitrogen. Parameters are described in Table 1. Parameter values are deduced from Oschlies and Garçon (1999) and Huret et al. (2005), as well as from a preliminary sensitivity analysis (Charria, 2005).

The \(J(z, t, N)\) represents the phytoplankton growth function of light and nutrient limi-
Following the minimum Liebig’s Law (Liebig, 1845), the minimum between light and nutrient limitation is used:

\[ J(z, t, N) = \min(J(z, t), J_{\text{max}}L_{\text{NO}_3}) \]  

(6)

with \( J(z, t) \) the light-limited growth rate, \( L_{\text{NO}_3} \) the nutrient limitation term and \( J_{\text{max}} = ab^cT \) the light-saturated growth rate and \( T \) the temperature (°C).

The nutrient limitation follows the Michaelis – Menten formulation:

\[ L_{\text{NO}_3} = \frac{N}{N + K_N} \]  

(7)

where \( K_N \) stands for the half-saturation constant for nutrient uptake.

The light limited growth rate is based on the analytical method from Evans and Parslow (1985). One of the advantages of this method is that a diurnal cycle from daily solar fluxes is simulated analytically. In fact, the light limited growth rate \( J(z, t) \) is averaged over 24 h and over a vertical layer as:

\[ \bar{J}(z, t) = \frac{1}{\tau_{24h}} \int_0^{24h} \frac{1}{z_k - z_{k-1}} \int_{z_{k+1}}^{z_k} J(z, t) \, dz \, dt \]  

(8)

where

\[ J(z, t) = \frac{J_{\text{max}} \alpha I(z, t)}{(J_{\text{max}}^2 + \alpha^2 I(z, t)^2)^{1/2}} \]  

(9)

where \( I(z, t) \) is the local light intensity and \( \alpha \) the initial slope of the photosynthesis-light (P-I) curve. Using these Eqs. (8 and 9), the light is maximum at noon at surface (it decreases exponentially with depth). Evans and Parslow (1985) then compute analytically the limitation term integrated over the mixed layer depth and during the day using the following equation:

\[ \bar{J}(z, t) = \frac{2J_{\text{max}}}{kM} \left[ F(\beta e^{kM}, \tau) - F(\beta, \tau) - F(\beta e^{kM}, 0) + F(\beta, 0) \right] \]  

(10)
where \( M \) is the mixed layer depth, \( k = 0.04 \text{ m}^{-1} \) the light attenuation coefficient, \( \tau \) the time at noon and

\[
F(y, t) = (y^2 + t^2)^{1/2} - t \ln t + (y^2 + t^2)^{1/2} / y
\]

(11)

G(P) represents the zooplankton grazing, it is written following a Holling type III function:

\[
G(P) = \frac{gpP^2}{g + pP^2}
\]

(12)

with \( g \), the maximum grazing rate for the high \( P \) values and \( p \) the \( G(P) \) slope for the weak \( P \) values. The advection scheme is MUSCL (Monotonic Upstream centred Scheme for Conservation Laws – Lévy et al., 2001). This scheme is monotonic, positive with an implicit diffusion and a weak dispersion.

The interannual simulation has been initiated from Reynaud et al. (1998) climatology with a one year spin-up of the physical model in 1995. The coupled experiment started in 1996.

The initial dissolved inorganic nitrogen field is from the Conkright et al. (2002) climatology. The other biogeochemical variables need a shorter period to reach a balanced state (in order of one year). Consequently, the initial Phytoplankton, Zooplankton, and Dissolved Organic Nitrogen are respectively 0.14 mmolN m\(^{-3}\), 0.014 mmolN m\(^{-3}\) and 3 mmolN m\(^{-3}\) at surface and are decreasing exponentially with depth. Detritus are initialized to \( 10^{-4} \) mmolN m\(^{-3}\) on the whole domain. An approximately stable seasonal cycle was reached after 2 years, end of 1997. The fields used in this study represent the third year of coupled integration (i.e. 1998). Statistical analyses and comparisons with remotely sensed and in situ data showed that the model reproduces well the seasonal cycle of the ecosystem as well as the primary production in the North Atlantic biogeochemical provinces (Charria et al., 2006b). However a few biases were highlighted in the simulated fields. The Gulf Stream position is too far North as compared
to observations. This is a well known bias in this kind of general circulation model (Barnier et al., 2006). Concerning the biogeochemical fields, the northern boundary of the oligotrophic gyre is located too far South. Indeed, the northern boundary is located around 28° N in the observations and around 23° N in the simulated fields. The elevated chlorophyll concentrations in the oligotrophic gyre are linked to high inorganic nitrogen contents due to a fast remineralization loop above the nitracline associated with regenerated production.

3 Data

3.1 Chlorophyll-a concentrations

Chlorophyll-a concentrations (in mgChl m\(^{-3}\)) were obtained from the ocean colour sensor SeaWiFS (Sea-viewing Wide Field-of-view Sensor) products (level 3 binned data, monthly, version 4 – O’Reilly et al., 1998) generated by the NASA Goddard Space Flight Center (GSFC) Distributed Active Archive Center (DAAC) (McClain et al., 1998). The products are on a regular grid of 9 by 9 km for the year 1998 (similar to the simulated year). The predicted error on the single 1-km SeaWiFS estimates of chlorophyll-a concentration is 35% (McClain et al., 1998); the accuracy of 9 km gridded data is comparable or better. As our study focuses on the anomalies of chlorophyll-a concentrations, a monthly zonal average from raw data at each latitude and for each month is removed, so also removing part of the seasonal cycle. To apply the spectral analysis and filtering described below, gaps in the data, mainly due to the presence of clouds, are filled with a linear interpolation.

3.2 Sea Level Anomalies (SLA)

Sea Level Anomaly data are provided by CLS Space Oceanography Division as part of the Environment and Climate EC AGORA and DUACS projects. These SLA were
obtained from the combined processing of the Topex/Poseidon (T/P) and ERS-1/2 data. The two data sets were combined using an improved space/time objective analysis method taking into account long-wavelength-errors (noise correlated on large scales) with a 1–2 cm mean error (Le Traon et al., 1998). SLA are relative to a seven year average (1993–1999) and were mapped every 7 days for the year 1998 with a spatial resolution of 1/3°. To have the same temporal resolution as for the surface chlorophyll-a concentrations, SLA data were averaged with a monthly time step.

4 Rossby waves in the coupled model

Rossby wave features in the coupled experiment results (Sea Level Anomalies – SLA – and surface chlorophyll concentrations) are first described and compared to those deduced from remotely sensed data (Charria et al., 2006a).

The SLA are analysed to detect the westward propagating signals. SLA are estimated from the temperature and salinity fields and from the barotropic stream function computed with the pressure compensation relation (Mellor and Wang, 1996).

We will consider four main parameters (wavelength, amplitude, phase speed and phase relationship between SLA and surface chlorophyll concentrations) previously used in analysis of Rossby waves features.

The wavelengths of westward propagating signals in SLA are estimated using a 1-D continuous wavelet analysis applied for each model latitude between 10°N and 40°N. Maxima in the local wavelet power spectra are associated with wavelengths mainly located between 400 and 1000 km with a few occasions over 1000 km. They are representative of the first baroclinic mode of Rossby waves according to the linear and extended theories (Killworth and Blundell, 2003). These values are similar to those estimated from remotely sensed data with Fourier-based spectral analyses (Cipollini et al., 1997; Hill et al., 2000; Killworth et al., 2004), least-square based analysis combined with Radon transform (Polito and Cornillon, 1997) or wavelet method (Charria, 2005; Charria et al., 2006a) (Table 2).
The modelled SLA are filtered reconstructing the signal for the wavelengths located between 400 and 1200 km after a 1-D continuous spatial wavelet analysis (Torrence and Compo, 1998). To estimate the Rossby wave amplitude, the westward propagating signals are extracted (Fig. 2). The simulated waves present amplitudes slightly weaker (reaching 5 cm) than those observed (7–10 cm) in remotely sensed data (Killworth et al., 2004; Charria et al., 2006a) over the North Atlantic Ocean (Table 2). The lower amplitudes in the simulations are due to the model spatial resolution. Higher resolution would be necessary to simulate the whole Rossby waves energy.

Propagation velocities (or phase speeds) associated with Rossby waves are calculated from the filtered time/longitude diagrams with a 2-D Radon transform method (Challenor et al., 2001). The modelled velocity values are between 3 and 4 cm s\(^{-1}\) (Table 2). They fall in the same range as those estimated in previous studies (Table 2). However, a narrower interval is observed for the present study partly due to the time period over which the phase speeds are estimated.

Similar analyses, based on the longitude/time diagrams and using the same wavelet filter, are performed on the simulated surface chlorophyll concentration fields. Wavelengths between 400 and 1500 km are observed in agreement with those deduced from the SeaWiFS data for the same year. These wavelengths are similar to the SLA south of 35°N and are slightly higher north of this latitude. Concerning the amplitudes associated with westward propagations, they are similar in the filtered modelled concentrations and in the filtered remotely sensed concentrations estimated by the SeaWiFS sensor. Note that chlorophyll anomalies are used, not absolute values; areas of high absolute chlorophyll concentrations may not necessarily show large anomalies.

Now we will examine the relationships between SLA and surface chlorophyll concentrations. Following Killworth et al.’s (2004) approach, the coupled processes involved can be estimated from the relationships between SLA and surface chlorophyll concentrations. As shown by Charria et al. (2006a), the phase relationships between Rossby waves (signal in SLA) and their chlorophyll signature can be highly variable. We compare the phase relationships using a cross-wavelet analysis between filtered
SLA and filtered surface chlorophyll concentrations. From 23° N to 35° N, the modelled phase relationships (phase(Chlorophyll) – phase(SLA)) are mainly between $\pi/2$ and $\pi$ (Fig. 3a). By contrast, between 16° N and 23° N in the simulated fields (Fig. 3a) and between 10° N and 28° N in the remotely sensed data (Fig. 3b), values of phase relationship are between $-\pi/2$ and 0. According to the theoretical approach from Killworth et al. (2004), we can deduce that between 16° N and 23° N in the simulations (between 10° N and 28° N in the data), the meridional chlorophyll advection should be the dominant process as confirmed by Charria et al. (2006a). However, the phase relationships deduced from the modelled fields north of 23° N do not allow to detect a clear dominant process when the meridional chlorophyll gradients are positive due to the same phase relationship for vertical and horizontal processes (Killworth et al., 2004).

After the analysis of modelled Rossby waves and chlorophyll signature features and the comparison with previous studies, we can conclude that Rossby waves are well represented in the model despite a southwards shift of the boundary between two different phase regimes due to modelled biogeochemical conditions.

5 Region selection with significant Rossby wave signatures

To quantify the influence of Rossby waves on primary production, our analysis is focused on the three parts of the wave chlorophyll signature: the positive chlorophyll anomaly linked to the wave’s crest (hereafter CA$^+$), the negative anomaly linked to the wave’s trough (CA$^-$) and the lack of chlorophyll anomaly (CA0). The accurate identification of these wave patterns in particular regions needs the following pre-processing on the simulated fields.

Before extracting any wave, the previously spatially filtered fields (see section 4) at each time step and each latitude are used to detect clear wave propagations in SLA and surface chlorophyll concentrations. A cross-spectrum analysis between SLA and the decimal logarithm of surface chlorophyll concentrations (similar to the analysis applied on remotely sensed data in Killworth et al., 2004) is then performed on longitude/time
windows. Windows are 60 grid points wide (around 20 degrees in longitude). The cross-spectrum energy is integrated over Rossby wave period and wavelength (3–24 months/400–1200 km). Figure 4 shows the amplitude of the integrated cross-spectrum peak. Larger values are observed in the north-west part of the domain where Rossby waves have larger amplitudes. Then, several maxima can be extracted for different latitudes. We decided to focus on regions centred around the mid-Atlantic ridge and not influenced by the coastal effects. A few latitudes are then selected between 30°W and 60°W where the cross-spectrum amplitudes are above \(4.35 \, \text{m} \log_{10}(\text{mgChl} \, \text{m}^{-3})\). An homogeneous zonal distribution is also preferred. Based on these latitudes, the extent of the time/space region is refined over the time period where positive/negative chlorophyll anomalies can be clearly identified and followed in time. Finally, 20 time/space regions are chosen over the basin (Fig. 5). These regions are equally distributed to cover different latitudes where Rossby waves propagations are well correlated with their chlorophyll signature.

For each area, dissolved inorganic nitrogen and phytoplankton modelled source and sink terms, as well as associated advective and diffusive terms, are detailed. Each flux is vertically integrated over 112 m depth. Values obtained are then averaged in longitude over the positive or the negative chlorophyll anomalies. The value for the lack of chlorophyll anomaly (case CA0) is estimated from an average of values from the western boundary of the trough, between the crest and the trough and from the eastern boundary of the crest. The three integrated and averaged values (for \(\text{CA}^-\), \(\text{CA}^0\) and \(\text{CA}^+\)) are then integrated over the time period. The integrated terms (biogeochemical, advection and diffusion) are then obtained for each region over the chlorophyll signature of the wave crest (\(\text{CA}^+\)), the wave trough (\(\text{CA}^-\)) and for assumed background conditions (CA0).
6 Results

Two kinds of terms are then extracted: the biological fluxes between the ecosystem components and the physical advection/diffusion terms. To understand the origin of higher chlorophyll concentrations due to the wave’s passage, the horizontal and vertical advection terms as well as the vertical diffusion are analysed (the MUSCL advection scheme doesn’t resolve any horizontal diffusion).

6.1 Chlorophyll wave crest and primary production

First, the terms for the chlorophyll signature of the wave crest (CA+) are analysed. The influence on primary production is obtained for each region (Fig. 6) and then the underlying processes investigated (Figs. 7–9). Results are very sensitive to the latitude and the time period considered. Nevertheless, we can consider five main subgroups for different zonal bands.

In the northern part of the domain, north of 39 N, two main regions are considered (40 N–41.3 W and 39.5 N–34.3 W). A +10.5% increase in primary production is obtained for the northernmost latitude (40 N–41.3 W). This value can be associated with the meridional advection processes (dissolved inorganic nitrogen and phytoplankton). Indeed, the phytoplankton meridional advection and the inorganic nitrogen meridional advection are reaching respectively more than +140% and +78% of the CA0 condition, respectively (Fig. 7a). At the opposite, a strong decrease in primary production (−34.3%) can be noticed in the southernmost latitude (39.5 N–34.3 W) (Fig. 7b). This decreasing production is mainly explained by a strong diminution of the vertical inorganic nitrogen advection and the zonal phytoplankton advection. The CA+ is then explained by a simple meridional chlorophyll advection (Fig. 7b). These first results

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1 Regions are identified using the latitude and the easternmost position of the wave during the given period.

2 The percentage of increase (or decrease) is the difference between CA+ (or CA0) divided by the absolute value of CA0 and multiply by 100.
illustrate the complex variability in the Rossby wave influence on primary production. Furthermore, these two close latitudes are corresponding to very different time periods (winter at 40 N–41.3 W and spring at 39.5 N–34.3 W) associated with two different biogeochemical regimes (bloom in spring). In spring, the primary production associated with the CA+ will then decrease due to the limiting effect of the vertical inorganic nitrogen advection. The combination of these different factors linked to biogeochemical background and Rossby wave features can easily explain such a difference between these two situations.

Further south, 8 extracted regions between 29 and 39 N are considered together (38.9 N–42 W/37.1 N–32.7 W/34.7 N–37 W/34.7 N–40 W/32.2 N–31.3 W/31.9 N–41.7 W/29.9 N–32.7 W/29.6 N–47.3 W). In this band, primary production is slightly increasing between CA0 and CA+ (+0.2% at 38.9 N–42 W; +12.3% at 37.1 N–32.7 W; +7.2% at 34.7 N–37 W; +2.7% at 32.2 N–31.3 W; +3.1% at 31.9 N–41.7 W and +9.4% at 29.6 N–47.3 W) except for the 34.7 N–40 W and 29.9 N–32.7 W regions where the primary production is decreasing (−8.3% and −10.6% respectively). For these different cases, the primary production increase is mainly driven by the horizontal and vertical advection of phytoplankton and dissolved inorganic nitrogen. For example, in Fig. 8a (37.1 N–32.7 W case), the CA+ is associated with a strong enhancement of the meridional phytoplankton advection. By contrast, in Fig. 8b, we observe that the primary production diminution is due to decreases in phytoplankton and inorganic nitrogen vertical advection as well as horizontal phytoplankton and inorganic nitrogen advection.

Between 25 N and 26 N, two wave signatures are followed for two very close latitudes (25.8 N–38.7 W and 25.5 N–44.7 W) and with a common initial time but for a longer time period in the case of the easternmost longitude (from 26 March to 10 June at 25.5 N–44.7 W and to 13 August at 25.8 N–38.7 W). Surprisingly, the effects of the wave passage on the primary production are completely different. For the easternmost area, we observe a strong decrease in primary production associated with CA+ (−19.9%). At the other location, we found a +20.3% augmentation. The source of the increase is
associated with vertical and horizontal sources of dissolved inorganic nitrogen. Wave signatures are tracked for different time periods at 20.6°N and 18.7°N. First, at 20.6°N, primary production enhancements are observed. There are estimated at +3.4% over a slightly longer time period (26 July to 18 December) and +12.7% over a shorter time period (18 August to 18 December). The vertical diffusion of phytoplankton allows to explain this production increase. The difference between these two values is associated with the integration time period and highlight a negative effect on primary production in the CA+ between mid-July and mid-August 1998 for this latitude. At 18.7°N, no significant primary production trend is observed in the western part (+1.6%) and a −13.9% decrease is obtained in the eastern part. This last decrease is associated with a diminution of the meridional and vertical advection of phytoplankton and dissolved inorganic nitrogen during the wave passage.

Finally in the southern part of the domain (9°N–15°N), the analysed wave signatures are initially tracked east (14.8°N–34.7°W and 10°N–37.3°W) and over (14.8°N–41°W and 11.3°N–42.7°W) the mid-Atlantic ridge. The westernmost waves are observed earlier during the year than the easternmost propagations. In every case, an increase in primary production associated with CA+ is noticed: +15.8% at 14.8°N–34.7°W, +8.4% at 14.8°N–41°W, +18.2% at 11.3°N–42.7°W and +7.3% at 10°N–37.3°W. It appears that wave signatures from the eastern part of the basin observed during autumn have a weaker impact on primary production than those observed in spring and summer. This is in agreement with a more stratified ocean in summer. Indeed, when we detail the processes involved, the vertical input of dissolved inorganic nitrogen explains the production increase (Fig. 9a). By contrast, during the less productive season, the primary production increase associated with the wave signature is due to meridional phytoplankton advection and vertical phytoplankton diffusion (Fig. 9b). There is no significant inorganic nitrogen advective input.
6.2 Chlorophyll wave trough and primary production

After analyzing the processes explaining the surface chlorophyll positive anomaly and the impact on the primary production, the opposite situation during the chlorophyll negative anomaly (CA−) is explored. It generally appears in the southern part of the domain that the CA− is associated with a negative effect on primary production (Fig. 6). This first result confirms that the vertical advection of dissolved organic nitrogen is never the only process involved during the wave’s passage. In fact, this mechanism inducing a inorganic nitrogen upwelling has an effect on primary production. This effect needs then to be offset by different mechanisms as horizontal advection to explain the decrease in primary production measured as well as the negative surface chlorophyll anomaly.

Except in a few regions, the primary production trend (increase or decrease) in the CA− is then opposite to the situation in the CA+. This opposition is also observed when we consider each mechanism separately.

More surprisingly, we can remark that when the primary production is decreasing over the CA+, the situation in the CA− is opposite and we generally observe an increase in primary production associated with the CA−. This result confirms the important role of meridional advective mechanisms as suggested by Killworth et al. (2004) and Charria et al. (2006a). In fact, in the case of an horizontal advection, the chlorophyll is similar to a passive tracer and only reacts to the physical forcing without significant changes in the biogeochemical processes. These purely physical processes are theoretically symmetrical between the crest and the trough of a Rossby wave. This bipolar system reflects a symmetric effect of the Rossby waves on primary production which suggests that the net effect of Rossby waves on primary production would be very small.
7 Discussion

In the present work, the different biogeochemical-physical mechanisms explaining the surface chlorophyll signature of Rossby waves are investigated through the exploration of the effect of these waves on primary production. The use of a realistic coupled physical/biogeochemical model in the North Atlantic allows to detail separately the positive and negative surface chlorophyll anomalies associated with these waves.

The biogeochemical advective and diffusive fluxes analysed for 20 areas highlight the fact that the increase of phytoplankton biomass in the CA+ is due to a complex combination of vertical and meridional processes depending on the considered latitude. This surface chlorophyll positive anomaly is associated with various patterns of primary production depending upon season and region (latitude and longitude). Previous studies with remotely sensed data and theoretical modelling from Killworth et al. (2004) and Charria et al. (2006a) showed that the Rossby wave signature in chlorophyll concentration can be explained by the meridional chlorophyll advection mostly south of 28 N. When we compare these results to the present work (Fig. 10), the role of the horizontal advection is also noticed but several regions highlight also a strong effect of the dissolved inorganic nitrogen vertical advection. Based on the 20 present regions analysed, the different relative contributions north and south of 28 N (as described in Charria et al., 2006a) are not fully reproduced. This can be partly explained by the limited time period and the specific location of the selected regions. Furthermore, several assumptions were made to compare remotely sensed data and theoretical modelling in Killworth et al. (2004) and Charria et al. (2006a). Indeed, a simple term was used to represent the source and sinks biogeochemical terms in the theoretical model and several statistical assumptions were made to fit the model outputs with observations. However, the present results of the realistic simulations confirm the weak effect of the vertical advection of phytoplankton, apparent in the whole domain (Fig. 10).

The second part of this work addresses the local impact of Rossby waves on primary production following the time period and location. When we quantify the local enhance-
ment of primary production due to the wave passage, values between $-34.3\%$ (latitude 39.5 N–34.3 W) and $+20.3\%$ (latitude 25.5 N–44.7 W) in percentage of increase and between $-0.108 \text{ mmolN m}^{-2} \text{ d}^{-1}$ (latitude 39.5 N–34.3 W) and $+0.098 \text{ mmolN m}^{-2} \text{ d}^{-1}$ (latitude 37.1 N–32.7 W) are obtained. These values represent the difference between CA+ and CA0 primary productions (divided by the CA0 primary production for the percentages). It appears that the increases are stronger when the dissolved inorganic nitrogen vertical advection is involved (i.e. between 9 N and 15 N). Furthermore, this last process seems to be systematically involved in spring and summer when the increases of primary production are exceeding 5%. By contrast, decreases in primary production associated with the CA+ are generally due to a decrease of the vertical advection and diffusion of dissolved inorganic nitrogen. For the other situations, the primary production decreases associated with CA+ can be explained by a contribution of the meridional advection of phytoplankton. The different dominant processes explaining primary production decrease and the CA+ signature are probably the result of a strong stratification in summer which will induce a de-correlation of the processes in the mixed layer and below. Kawamiya and Oschlies (2001), with the same kind of coupled physical/biogeochemical model applied in the Indian Ocean, showed that Rossby waves can induce a $\sim30\%$ increase of primary production associated with the uplifting of the deep chlorophyll maximum during the wave passage. This value obtained with a different approach in a different basin is comparable to our estimations. In the Pacific, Sakamoto et al. (2004), based on local measurements at the ALOHA site and on an estimated relationship between primary production and Sea Surface Height anomalies estimated near the HOT station ALOHA, showed similar increases in primary production reaching 25%. They were linked to nitrate injection in the euphotic layer during two well identified passages of Rossby waves. Using the same relationship, these authors conclude a limited role of Rossby waves inducing a mean primary production enhancement lower than 5–10 % over the 1997–1999 period. These values are also comparable to our estimations over the selected transects in the North Atlantic Ocean. Then, in our study similar increases are obtained using a different approach in a
different ocean but decreases are also simulated. Consequently, our results suggest a net weak effect of Rossby waves on primary production.

These estimations need however to be considered carefully. These results using a realistic ocean general circulation model in the North Atlantic are obtained on given latitudes mainly located in low production provinces for a given time period. Furthermore, the method used to identify wave propagation in surface chlorophyll concentrations is based on several assumptions necessary to perform this first detailed analysis of the coupled physical/biogeochemical processes involved during the Rossby wave propagation. In particular, a recent study from Chelton et al. (2007) suggests that most westward propagating features observed in remotely sensed sea surface height are the signature of non linear propagating eddies. In the North Atlantic, more than 25% of the variance is explained by eddies instead of Rossby waves west of 53 W and north of 32 N from the western to eastern coasts. These observations are based on the Okubo-Weiss parameter which is a measure of the relative importance of deformation and rotation (see Fig. 3 in Chelton et al., 2007). In our studied domain, 14 out of 20 regions are located south of 32 N and east of 53 W which is a region where non linear eddies are not predominant. North of 32 N, our approach does not allow to certify that Rossby wave signature is analysed instead of eddy signature. However, Sweeney et al. (2003) and Mouriño-Carballido and McGillicuddy (2006) showed that in the case of eddies, the primary production associated with the negative chlorophyll anomaly will be similar to the background primary production. In the present work, we showed that generally, the negative chlorophyll anomaly is associated with a decrease in primary production which suggests a wave-like process acting instead of non linear eddies.

8 Conclusions

Rossby waves, a process previously observed with remotely sensed data, were modelled using a 3-D coupled physical/biogeochemical model in the North Atlantic ocean. Waves are well reproduced as well as their signature in surface chlorophyll concentra-
The analyses based on 20 regions showed the important contribution of horizontal advection and of vertical inorganic dissolved nitrogen advection. The main mechanism involved differs according to the biogeochemical conditions (due to latitude and time period). Furthermore, Rossby waves have a non negligible influence on primary production which can be quantified using coupled 3-D modelling. Indeed, positive surface chlorophyll anomalies are generally associated with an increase in primary production. This approach also allowed to explore the influence of wave trough on primary production and a negative effect was observed. The net impact of Rossby wave on primary production, based on the 20 regions analysed, seems to be weak due to the symmetrical shape of the wave effects (between crest and trough). However, as suggested by Sakamoto et al. (2004), inputs of dissolved inorganic nitrogen can induce shifts in the phytoplankton community structure and consequently increase the exported production. This approach using different particular cases needs to be extended to the whole basin in order to quantify the net basin scale effect of Rossby waves on primary production. To perform this estimation, we need to investigate what happens when the Rossby wave effect is removed. This calls for a more systematic study with coupled physical/biogeochemical models where Rossby waves can be switched on and off.

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Table 1. Parameters of the ecosystem model.

| Symbol | Parameter | Definition |
|--------|-----------|------------|
| **Phytoplankton coefficients** | | |
| $\alpha$ | $0.025 \, \text{d}^{-1}/(\text{W m}^{-2})$ | Initial slope P-I curve |
| PAR | 0.43 | Photosynthetically active radiation coefficient |
| $k_c$ | $0.03 \, \text{m}^{-1}(\text{mmol m}^{-3})^{-1}$ | Light attenuation by phytoplankton |
| a | $0.6 \, \text{d}^{-1}$ | Maximum growth rate parameters |
| b | 1.066 | |
| c | $1 \, (^\circ \text{C})^{-1}$ | |
| $1/k_w$ | 25 m | 1/Light attenuation due to water |
| $k_N$ | 0.5 mmol m$^{-3}$ | Half-saturation constant for N uptake |
| $\mu_P$ | $0.03 \, \text{d}^{-1}$ | Specific mortality rate |
| $\epsilon$ | 0.01 | Exudation fraction of primary production |
| **Zooplankton coefficients** | | |
| $f_1$ | 0.75 | Assimilation efficiency |
| g | $2 \, \text{d}^{-1}$ | Maximum grazing rate |
| $\rho$ | $1 \, (\text{mmol m}^{-3})^{-2} \, \text{d}^{-1}$ | Prey capture rate |
| $\mu_z$ | $0.2 \, (\text{mmol m}^{-3})^{-1} \, \text{d}^{-1}$ | Mortality rate |
| $\gamma$ | $0.03 \, \text{d}^{-1}$ | Excretion rate |
| $f_2$ | 0.25 | Organic fraction of excretion |
| **Detritus coefficients** | | |
| $\mu_d$ | $0.23 \, \text{d}^{-1}$ | Hydrolysis rate |
| $w_S$ | $5 \, \text{m d}^{-1}$ | Sinking velocity |
| **DON coefficients** | | |
| $\rho$ | $0.025 \, \text{d}^{-1}$ | Remineralization rate |
Table 2. Summary of Rossby wave features compared to previous studies. Extracted wavelengths and speeds are representative of the North Atlantic basin (mainly between 10° N and 40° N) except for the Cipollini et al. (1997) study based on the North-East Atlantic at 34° N.

| Amplitude (cm) | Wavelengths λ (km) | Phase Speeds $c_\varphi$ (cm s$^{-1}$) | Time period | References |
|----------------|---------------------|------------------------------------------|-------------|------------|
| 0–5            | 400–1200            | 3–4                                      | 1998        | Present Study (MNATL) |
| 1–10           | 500–1000            | 2–8                                      | 1993–2001   | Charria et al. (2006a) |
| 0–7            | 400–900             | 2–8                                      | 1997–2002   | Killworth et al. (2004) |
| 200–2000       | 3–17                | 1991–1996                                |             | Hill et al. (2000) |
| 390–520        | 0.9–3               | 1992–1995                                |             | Cipollini et al. (1997) |
| 1.5–11.6       | 400–4600            | 1.2–31.25                                | 1992–1995   | Polito and Cornillon (1997) |
Fig. 1. NPZDDON model (Huret et al., 2005).
**Fig. 2.** Examples of longitude-time plots of simulated (top) and remotely observed (bottom) Sea Level Anomalies for the 25.15° N latitude after filtering with spatial 1-D wavelet analysis.
Fig. 3. Spatial phase relationships for the year 1998 between chlorophyll concentrations and SLA, averaged in time, obtained from a cross wavelet analysis for: the coupled model (a) and remotely sensed data (b).
Fig. 4. Cross-spectral amplitude (in m log10(mgChl m$^{-3}$)) integrated over Rossby wave period and wavelength (3–24 months/400–1200 km) for the 11 July 1998.
Fig. 5. Position (white lines and red lines) of selected regions compared to the model bathymetry (top) and time extension of these regions (bottom). Different colours are only used to distinguish regions around close latitudes and are similar in top and bottom figures.
Fig. 6. Primary production increases (in %) following the latitude over the positive (blue Δ) and the negative (red ◢) chlorophyll anomalies. Fluxes are integrated over the time period, 112 m depth and averaged in longitude. The percentage of increase is derived from the value for the lack of chlorophyll anomaly.
Fig. 7. Meridional (yad), zonal (xad) and vertical (zad) advection of phytoplankton (phy) and dissolved inorganic nitrogen (no3), vertical diffusion of phytoplankton (zdf.phy) and inorganic nitrogen (zdf.no3) and primary production (IPP) at 40 N–41.3 W from 11 January to 3 February 1998 (a) and 39.5 N–34.3 W from 3 May to 17 May 1998 (b). These fluxes are integrated over the time period, 112 m depth and averaged in longitude over the positive (blue △) and the negative (red ▽) chlorophyll anomalies. The percentage of increase is derived from the value for the lack of chlorophyll.
**Fig. 8.** Same fluxes as represented in Fig. 7 at 37.1 N–32.7 W from 28 December 1997 to 18 February 1998 (a) and 29.9 N–32.7 W from 5 April to 8 July 1998 (b).
Fig. 9. Same fluxes as represented in Fig. 7 at 11.3°N–42.7°W from 8 March to 15 April 1998 (a) and 10°N–37.3°W from 30 September to 8 December 1998 (b).
Fig. 10. Relative contribution (in %) of the three assumed processes which can explain the chlorophyll signature of Rossby waves: horizontal advection, upwelling of dissolved inorganic nitrogen in the euphotic zone and uplifting of the subsurface chlorophyll maximum. The black dots represent the results from Charria et al. (2006a) north of 28 N (a) and south of 28 N (b) between 9.5 N and 40.5 N. The superimposed red crosses represent the 20 regions analysed in this study with the 3-D model outputs. The horizontal advection is the sum of the zonal and meridional phytoplankton and inorganic nitrogen advections. The upwelling of nitrates is the sum of vertical advection and diffusion for dissolved inorganic nitrogen respectively. The uplifting of chlorophyll is the sum of vertical advection and diffusion for phytoplankton. The relative contributions are computed from the mechanisms associated with the wave crest in each region.