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Effects of a Chronic Oil Spill on the Planktonic System in San Jorge Gulf, Argentina

A ONE-VERTICAL-DIMENSION MODELING APPROACH

By Philippe Klotz, Irene R. Schloss, and Dany Dumont
ABSTRACT. Known for its high biological productivity, San Jorge Gulf (SJG) in Argentinean Patagonia is also an area of oil exploitation. To understand the dynamics of the SJG plankton ecosystem under several scenarios of potential hydrocarbon (HC) contamination, we present an 11-compartment biogeochemical model coupled to a turbulence model. In the coupled model, we parameterize the main physical and biological processes related to HC contamination, such as biodegradation, growth, and mortality of phyto-, zoo-, and bacterioplankton. Planktonic responses to several levels of HC contamination are studied for two physically contrasting SJG environments, a tidally well-mixed water column and a stratified water column. Results show increasing phyto- and bacterioplankton biomass with increasing HC concentration, which in turn produces more detritus. Zooplankton communities seem to respond differently depending on HC concentration, with major indirect changes occurring in the different size classes. Effects of HC contamination on biological compartments are stronger in the stratified than in the well-mixed environment.

INTRODUCTION
Since the beginning of the twentieth century, along the Argentinian coast, particularly within San Jorge Gulf (SJG), oil production has been the main economic driver, accounting for nearly half of total oil production in the country (Ministerio de Energía y Minería, 2016). Offshore exploration and extraction of this resource pose risks to the golf’s marine ecosystem. Thus, it is essential to understand the physical and biological dynamics of this coastal system. The SJG is exposed to strong sustained westerly winds throughout the year as well as strong semi-diurnal tidal currents, and is characterized by a counterclockwise circulation (Glorioso and Simpson, 1994; Glorioso and Flather, 1995; Palma et al., 2004). In the shallow (≤80 m) southeastern area of the SJG, the bottom friction of strong tidal currents results in significant turbulent mixing (Glorioso and Flather, 1995) that creates a permanent, mainly tidally induced, frontal zone (Glorioso and Flather, 1997; Palma et al., 2004; Flores-Melo et al., 2018, in this issue) This is confirmed by sharp horizontal gradients evident in satellite observations of temperature and chlorophyll-a (Chla) concentrations in the region (Acha et al., 2004; Rivas et al., 2006, Glembocki et al., 2015). Areas of high biological productivity are often associated with high horizontal density gradients, where turbulence plays a role not only in the availability of nutrients in the euphotic zone but also in the distribution of phytoplankton biomass (Flores-Melo et al., 2018, in this issue).

It is difficult to study and quantify the effects of hydrocarbons (HC) on planktonic assemblages in situ. Micro- and mesocosm experiments provide ways to target the direct effects of contamination on plankton by controlling experimental conditions such as nutrient concentration, temperature, and light intensity, which affect the growth, mortality, and distribution of organisms. However, because coupled biological-physical processes are nonlinear, the influence of physical factors cannot be accounted for in these kinds of studies.

Dissolved HC, especially polycyclic aromatic hydrocarbons (PAHs), contribute up to 60% of the total oil composition (Fingas, 2016) and have the most negative effect on marine organisms. The hydrophobic nature of these compounds and their bioavailability once they are dissolved in the marine environment favor their adsorption by marine organisms. As a result, mortality increases through simple toxicity or by reduction of the organism’s physiological activity (Anderson et al., 1974).

In contrast, specific strains of bacterioplankton contribute to one of the most persistent disaggregation processes found during an HC contamination event: biodegradation (Beazley et al., 2012). Since the early twentieth century, study of the bacteria genome has made it possible to isolate more than 70 genera capable of degrading HC (Hassanshahian and Cappello, 2013). In the marine environment, bacterioplankton exhibit significant changes in community structure and biomass following oil spills (Hazen et al., 2010; Dubinsky et al., 2013). These marine bacteria, known as obligate HC-degrading bacteria (OHCB; Yakimov et al., 2007), are ubiquitous at very low concentrations in the water column and are characterized by high substrate specificity.

In order to assimilate HC through fast growth rates, bacterioplankton depend on the concentration of dissolved oxygen and nutrients in the water column (Valentine et al., 2012; King et al., 2015). For example, phosphorus- and nitrogen-based compounds seem to act as regulators of biodegradation processes (Atlas and Bartha, 1972; Horowitz and Atlas, 1977; Atlas, 1981), thus limiting or promoting HC incorporation into pelagic carbon biomass (Shiller and Joung, 2012). However, the scarcity of information concerning the limiting nutrient forms and their related consumption makes it difficult to estimate precise biodegradation rates associated with HC spreading in seawater.

Of all the microorganism classes, phytoplankton display the highest variability in their response to HC contamination. From an ecotoxicological perspective, some phytoplankton groups appear to be more affected than others at equivalent concentrations. This is the case for diatoms, whose growth may sharply decrease (Hsiao, 1976; Nayar et al., 2005) because their siliceous walls significantly adsorb and accumulate PAHs (Siron et al., 1996; Sargian et al., 2007). However, other results show phytoplankton growth is stimulated at low concentrations of the dissolved HC fraction (Gordon and Prouse, 1973; Dunstan et al., 1975; Hsiao et al., 1978; González et al., 2013). The mechanisms associated with an increase in photosynthetic activity are poorly understood and lead to several hypotheses. For example, Baker (1971) and Cabioch et al. (1981) suggest
that extra nutrient uptake by phytoplankton is a result of the degradation of dead organisms by HC toxicity.

HC appears to affect all major zooplankton groups, but effects differ depending on species and HC concentrations. Almeda et al. (2013) observed 96% copepod mortality after 16 hours of exposure to PAH concentrations of 100 µL L⁻¹. They also calculated a mean lethal concentration (LC₅₀) of 31.4 µL L⁻¹.

Modeling allows simulation of pelagic system biogeochemical dynamics in realistic physical settings as well as simulation of the effects of contamination on plankton. The General Ocean Turbulence Model (GOTM) developed by Burchard et al. (2006) is a one-dimensional model of turbulence in the vertical dimension that can be coupled to a biogeochemical model of the nitrate-phytoplankton-zooplankton type (i.e., following Fasham et al., 1990). The coupled model can simulate the annual evolution of the mixing depth and include the processes related to the microbial loop. The effect of contaminants on the biological compartments can be further parameterized, allowing for a complete physical and biological interpretation of the effects of pollutants on plankton. Previous studies have simulated the behavior of HC in the marine environment (Reed et al., 1999), but only a few models have measured their impacts on the biogeochemical dynamics of the planktonic system (Gin et al., 2001; Valentine et al., 2012; González et al., 2013).

In this study, we combine data from the scientific literature and from the 2014 PROMESsee mission in the SJG (Flores-Melo et al., 2018, and Latorre et al., 2018, both in this issue) to simulate an offshore oil spill event. Using a one-dimensional biogeochemical-physical modeling approach, we evaluate the response of plankton to chronic HC contamination in a turbulent environment. Two mixing regimes typical of the southeastern frontal region of the SJG are compared: a seasonally stratified zone and a shallower, tidally well-mixed zone.

**MODEL DESCRIPTION**

**Physical Model and Configuration**

We used the GOTM coupled to a biogeochemical model to solve the advection-diffusion problem in the vertical dimension, assuming that state variables are horizontally homogeneous and that both mean and turbulent transport are driven by wind, buoyancy fluxes, and shear production. Horizontal advection is not taken into account. A detailed configuration of the two hydrodynamically contrasting experiments (stratified and well-mixed) can be found in the online supplementary materials.

**Biogeochemical Model and Configuration**

The biogeochemical model is composed of 11 compartments (see Figure 1), with a nitrate-phytoplankton-zooplankton-detritus (NPZD)-based structure connected to a microbial loop that reproduces nitrogen recycling from labile dissolved organic nitrogen (LDON) to ammonium (NH₄⁺). Because nitrogen is the limiting element in most marine ecosystems (Howarth, 1988), understanding its dynamics is an essential prerequisite to describing the carbon cycle (Fasham et al., 1990). Each biogeochemical variable is then expressed in nitrogen concentrations (mmol N m⁻³).

The model's biological super-compartments combine two compartments each of bacteria, phytoplankton, and zooplankton. In the bacterial super-compartment, BAC1 represents the bacterioplankton involved in the natural microbial loop that are only responsible for nitrogen recycling of detritus from LDON to ammonium. BAC2 represents the hydrocarbon-degrading or OHCB" strains that are

![Figure 1](https://example.com/figure1.png)

**FIGURE 1.** The general structure of the biogeochemical model is composed of 11 compartments: dissolved hydrocarbons, nitrates, ammonium (NH₄⁺), diatoms (DIA), flagellates (FLA), microzooplankton (MCZ), mesozooplankton (MSZ), bacteria community (BAC1), obligate hydrocarbon degrading bacteria (BAC2 [OHCB]), labile dissolved organic nitrogen (LDON), and detritus. Dashed lines illustrate the direct effects of hydrocarbons on biological compartments. Each arrow represents a flux and each compartment a concentration in mmol N m⁻³. Three super-compartments combine the two bacterial compartments (brown), the two phytoplankton compartments (green), and the two zooplankton components (orange).
found in high concentrations in oil-contaminated waters (see later section on Biodegradation).

Two phytoplankton groups, diatoms and flagellates, constitute the main size classes of organisms found in the SIG (Latorre et al., 2018, in this issue). In order to understand the indirect effects of HC on this super-compartment, both groups were parameterized with the same growth rates, initial concentrations, and settling velocities. Differences between diatoms (DIA) and flagellates (FLA) were only prescribed for mortality, zooplankton grazing preferences, natural sinking velocities, and transformation into detritus. An exudation factor reproduces the transfer of nitrogen to LDON. Given the high variability of HC effects on phytoplankton, no direct effect was parameterized and only the indirect effects were characterized.

Micro- (MCZ) and mesozooplankton (MSZ) represent two zooplankton classes in the model. Microzooplankton graze on diatoms, flagellates, detritus, and bacteria, and mesozooplankton graze on diatoms, flagellates, detritus, and microzooplankton. Zooplankton loss by exudation, natural mortality, and mortality by HC toxicity are the three sink terms leading to fluxes toward the ammonium, LDON, and detritus compartments (Table 1, Equations 1.3 and 1.4). Parameterization of nitrogen dynamics, which concern nitrate and ammonium, is described in supplementary materials.

**Parameterization of HC Effects**

To simulate the presence of HC in the GOTM, we considered a new variable: the portion of HC dissolved in seawater that results from weathering processes. As the literature suggests, the dissolved portion, also called the “water accommodated fraction,” is considered to be the most toxic HC state for marine biota (Anderson et al., 1974). Only the zooplankton super-compartment is parameterized by direct effects related to HC toxicity (see later section on Effects of HC on Zooplankton Mortality). Therefore, effects on phytoplankton dynamics after HC contamination are only indirect effects.

**Biodegradation**

Biodegradation of oil spill compounds involves a large number of specific organisms and biochemical processes. Thus, quantifying the transfer of carbon to the planktonic food web remains challenging. After several model runs, we chose the processes to integrate into the GOTM structure. Because the compounds in the extracted oil are composed of ≤1% nitrogen (Musser and Kilpatrick, 1998; Fingas, 2016; Petroleum HPV Testing Group,

**TABLE 1.** Biogeochemical equations and parameters involved in the biodegradation process (Equations 1.1 and 1.2) and the mortality of micro- and mesozooplankton by hydrocarbon toxicity (Equations 1.3, 1.4, 1.5, and 1.6). Biogeochemical fluxes from the output (o) toward the input (i) compartment follow the nomenclature: C_i,o.

<table>
<thead>
<tr>
<th>Equation</th>
<th>Parameter</th>
<th>Definition</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>CHC_{i,o} = νb_2 min_(C_{i} + C_{i}^{min})</td>
<td>C_HC</td>
<td>Hydrocarbon concentration</td>
<td>mmol N m⁻³</td>
</tr>
<tr>
<td></td>
<td>νb_2</td>
<td>Maximal uptake rate of OHCB</td>
<td>day⁻¹</td>
</tr>
<tr>
<td></td>
<td>min_(C_{i}^{HC})</td>
<td>Minimal HC to LDON uptake ratio for “OHCB” Bacteria</td>
<td></td>
</tr>
<tr>
<td></td>
<td>K_{i}^{HC}</td>
<td>Half saturation constant of traditional bacteria uptake</td>
<td></td>
</tr>
<tr>
<td></td>
<td>K_{i}^{OHCB}</td>
<td>Half saturation constant of OHCB uptake</td>
<td></td>
</tr>
<tr>
<td></td>
<td>C_{i}^{min}</td>
<td>Minimal bacterial concentration (traditional &amp; “OHCB”)</td>
<td>mmol N m⁻³</td>
</tr>
<tr>
<td></td>
<td>C_{3}</td>
<td>Microzooplankton concentration</td>
<td>mmol N m⁻³</td>
</tr>
<tr>
<td></td>
<td>C_{4}</td>
<td>Mesozooplankton concentration</td>
<td>mmol N m⁻³</td>
</tr>
<tr>
<td></td>
<td>C_{7}</td>
<td>Detritus concentration</td>
<td>mmol N m⁻³</td>
</tr>
<tr>
<td></td>
<td>C_{11}</td>
<td>Hydrocarbon concentration</td>
<td>mmol N m⁻³</td>
</tr>
<tr>
<td></td>
<td>(1 - ρ)</td>
<td>Grazing efficiency coefficient</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>g_{21}^{max}</td>
<td>Maximal mesozooplankton ingestion rate</td>
<td>day⁻¹</td>
</tr>
<tr>
<td></td>
<td>ρ_{g}</td>
<td>Mesozooplankton grazing preference on microzooplankton</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>Fac_{2}</td>
<td>Grazing preference normalization factor for mesozooplankton</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>μ_{21}^{HC} + μ_{12}^{HC}</td>
<td>Fractional micro- &amp; mesozooplankton loss of ammonium</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>δ</td>
<td>Fractional micro- &amp; mesozooplankton loss of LDON</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>C_{i}^{min}</td>
<td>Minimal bacterial concentration (traditional &amp; “OHCB”)</td>
<td>mmol N m⁻³</td>
</tr>
<tr>
<td></td>
<td>K_{6}^{HC}</td>
<td>Half saturation constant of micro- &amp; mesozooplankton loss</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>K_{6}^{HC}</td>
<td>Half saturation constant of micro- &amp; mesozooplankton loss by HC toxicity</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>n_{HC}</td>
<td>Slope factor of micro- &amp; mesozooplankton mortality for a sigmoid curve</td>
<td>day⁻¹</td>
</tr>
<tr>
<td></td>
<td>μ_{21}^{HC}</td>
<td>Maximal microzooplankton mortality rate</td>
<td>day⁻¹</td>
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<tr>
<td></td>
<td>μ_{21}^{HC}</td>
<td>Natural microzooplankton mortality rate</td>
<td>day⁻¹</td>
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hypothesized that the LDON compart-
processes is not known. Therefore, we
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pounds. However, the form of nitro-
consumption of nitrogen-based com-
amonium uptake by bacteria (BAC1).
Equation 1.2 in
substrate with BAC1, thus affecting the
rioplankton (BAC2), but they compete for
applies a co-limitation on HC consump-
the minimal HC to LDON uptake ratio
Equation 1.1).
Table 1
Biodegradation also involves the
consumption of nitrogen-based com-
ounds. However, the form of nitro-
gen utilized by OHC bacteria in these
processes is not known. Therefore, we
hypothesized that the LDON compart-
ment (Figure 1), which contains the most
bioavailable form consumed by bacteria,
apply a co-limitation on HC consump-
HC directly stimulates OHC bacte-
rioplankton (BAC2), but they compete for
substrate with BAC1, thus affecting the
whole bacterioplankton super-compartment. Equation 1.2 in Table 1 describes
the minimal HC to LDON uptake ratio
for the OHCB compartment.

Effects of HC on Zooplankton Mortality
Results from micro- and mesocosm
experiments show different but directly
noticeable effects of HC on the survival of
zooplankton species. Finding that
copepod species seemed to be strongly
affected, Almeda et al. (2013) proposed
a sigmoid numerical model of mortality
with increasing HC concentration.
Therefore, we integrated this relation-
ship between mesozooplankton mor-
tality and HC concentration with exist-
ing natural mortality rates $\mu_{21}$ and $\mu_{22}$
(Table 1, Equations 1.5 and 1.6). Because
no specific mortality rates were available
for microzooplankton species,
we applied the same HC toxicity to
micro- and mesozooplankton (Table 1, Equations 1.3 and 1.4).

CHRONIC OIL SPILL SCENARIO
Because chronic oil spills account for
the majority of HC pollution in seawater
(Potters, 2013), we chose to simulate a
chronic oil spill that replicates an offshore
oil leak. HC is incorporated into the system
in a subsurface 5 m layer with a small
time relaxation (seven days). In order to
target the austral summer, the contamination
period was set from August 1, 2013, to
July 31, 2014. To evaluate different
responses of the planktonic system to the
contamination, we performed one con-
rol run at three levels of contamination
(0, 10, 20, and 50 mmol N m$^{-3}$). Finally,
each contamination level was run in the
two physical configurations (stratified and
well-mixed). For example, CST$_{10}$ will be
referred as the control (0 mmol N m$^{-3}$)
simulation in the STratified experiment
using the latest Complete version of the
biogeochemical model (Table S1).

RESULTS
Physical Environment Dynamics
In all simulations, wind stress shows a sea-
sonal pattern, with the strongest average
values during austral winter. However, in the
stratified experiment, which aims to
reproduce the stratified side of the tidal
front in the southeastern region of the SJG,
episodic wind events strongly influence
upper layer mixing (Figures S2a and S3a)
and result in deepening of biotic and
abiotic variables at weekly timescales. In
this same experiment, temperature shows a
marked two-layer stratification pattern
from November 2013 to April 2014
(Figure S2b). In the well-mixed exper-
iment, temperature is almost vertically
homogeneous, with maxima reaching
6.6°C and 15.8°C in September and
March, respectively (Figure S3b). No vertical stratification is evident in either
experiment (Figures S2c and S3c).

Horizontal currents are forced by external pressure gradients oscillating at
the $M_2$ tidal period. Maximum current
speeds occur in the upper 20 m of the
water column and decrease toward the
seafloor (Figure S3e) due to bottom friction (Figure S1).

Distribution of HC
In the stratified experiment, verti-
cal distribution of dissolved HC fol-
ows a summertime two-layer stratifi-
cation close to the thermocline depth
(Figure 2a–c). During winter, vertical mixing allows the deepening of low HC
concentrations, which reach the bottom
of the water column between August and
October 2013. In the well-mixed exper-
iment, almost vertically homogeneous
concentrations can be observed from the
beginning of the simulation for all levels
of contamination with only subtle differ-
ces in the upper 30 m (Figure 2d–f).

In the stratified experiment, the low-
est level of HC contamination (CST$_{10}$,
Figure 2a), ranges from 1 mmol N m$^{-3}$
to 6 mmol N m$^{-3}$, last measurable in sur-
face waters at the end of September 2013.
From the end of August to mid-October
2013, 2 mmol N m$^{-3}$ are found between
50 m and 80 m depth. By November 16,
2013, the system is clear of contamina-
tion. However, during the first week of
May 2014, low concentrations between
0 m and 40 m appear again and last until
the end of July that year. In the well-
mixed experiment (CWM$_{50}$, Figure 2c),
values range from 4 mmol N m$^{-3}$ to
5 mmol N m$^{-3}$ from late September to
mid-October 2013. Thereafter, a quick
reduction in concentrations is followed
by the disappearance of HC two weeks
earlier than in the stratified experiment.

At the highest level of contamina-
tion, contrasting results can be observed
between the two experiments. In the
stratified experiment (CST$_{50}$, Figure 2c),
bio-degradation processes never seem
thoroughly effective throughout the sim-
ulation, and the maximal HC concen-
tration at 32 mmol N m$^{-3}$ remains from
early October to early December 2013
between 0 m and 20 m depth. Conversely,
in the well-mixed experiment (CWM$_{50}$,
Figure 2f), most of HC present is

This paper is structured as follows:

1. **Introduction**
   - Background on HC pollution and its impact on the marine ecosystem.
   - Objective of the study.

2. **Materials and Methods**
   - Experimental design and setup.
   - Data collection and analysis.

3. **Results**
   - Environmental conditions during the simulation.
   - Distribution of HC in the stratified and well-mixed experiments.

4. **Discussion**
   - Comparison of results between stratified and well-mixed experiments.
   - Implications for marine ecosystem health.

5. **Conclusion**
   - Summary of findings.
   - Implications for future research.

Additional references may include:

- Almeda et al. (2013)
- Potters, 2013
- Burchard et al.
- Liao et al.

**Table 1**
Describes the minimal HC to LDON uptake ratio for the OHCB compartment.

**Table S1**
Contains the latest Complete version of the biogeochemical model for simulations.

**Figures S1-S3**
Illustrate the distribution of HC and environmental variables over time.

**Equations 1.1-1.4**
Formulate the biogeochemical model and its components.
degraded over the entire water column after five months. Moreover, maxima are significantly lower, with 24 mmol N m\(^{-3}\) from the first week of October to the first week of November over the top 22 m of the water column.

**Effects of HC on Abiotic Compartments**

In the control experiment CWM\(_{00}\), by mid-January 2014, 8 mmol N m\(^{-3}\) of nitrate are still available, while this value is reached in late October 2013 in CWM\(_{20}\). In the stratified experiment, CST\(_{00}\) and CST\(_{20}\) results show concentrations reaching 12.5 mmol N m\(^{-3}\) in the bottom layer (50 m to 80 m water depth) from mid-December 2013 to mid-May 2014. Between 0 m and 20 m, minima between 0.5 mmol N m\(^{-3}\) and 2 mmol N m\(^{-3}\) are found from mid-December 2013 to the end of March 2014 for the CST\(_{00}\) simulation, and from mid-October 2013 to the end of March 2014 for the CST\(_{20}\) simulation.

In comparison to the CST\(_{00}\) simulation, other simulation results show elevated concentrations of ammonium. CST\(_{10}\), CST\(_{20}\), and CST\(_{50}\) present, respectively, values two, three, and seven times higher than the control CST\(_{00}\) (Figure 3c). On April 13, the maximum reaches 15 mmol N m\(^{-3}\) at 30 m depth for the CST\(_{20}\) simulation (Figure 3d). These values can be explained by a sustained bacterial activity below the thermocline (Figure 4f). During the same period, vertical detritus dynamics (Figure S4d) follow a similar pattern, confirming the active microbial loop in the HC contaminated experiments. In contrast, above the thermocline, lower concentrations, ranging from 2 mmol N m\(^{-3}\) to 4 mmol N m\(^{-3}\), decrease to 0 mmol N m\(^{-3}\) in December 2013 and January 2014. In the well-mixed experiment, vertical detritus distribution follows a very homogeneous pattern. Results from simulation CWM\(_{20}\) show values seven times higher than the control in early June 2014.

LDON, as part of the biodegradation process, shows interesting variations, especially in the stratified experiment. In CST\(_{00}\), a slow depletion from 0.3 mmol N m\(^{-3}\) in September 2013 to 0.05 mmol N m\(^{-3}\) in May 2014 is evident between 0 m and 80 m depth (Figure 3a). In CST\(_{20}\), there is rapid depletion (from 0.30 mmol N m\(^{-3}\) to 0.05 mmol N m\(^{-3}\)) in the top 25 m of the water column from October 12, 2013, to the end of March 2014 (Figure 3b). This trend follows the presence of HC toward the beginning of winter 2014 (Figure 2c), diminishing until LDON is absent in the upper 60 m in June 2014. Furthermore, these concentrations do not show strong dependence on stratification.

**Effects of HC on Planktonic Compartments**

The response of the phytoplankton, zooplankton, and bacterioplankton (BAC1) compartments to HC was studied. In Figures 4 and 5, we chose to compare results from simulations at 20 mmol N m\(^{-3}\) of contamination to the control simulations.

In the stratified experiment, phytoplankton show very high concentrations at all levels of contamination. Results from the CST\(_{20}\) simulation show maxima up to 8.7 times higher than the control values between 0 m and 30 m depth (Figure 4b). Moreover, the bloom periods shift from September (1.2 mmol N m\(^{-3}\)) and December 2013 (1.5 mmol N m\(^{-3}\)) for CST\(_{00}\) to February (11.4 mmol N m\(^{-3}\)) and March 2014 (13 mmol N m\(^{-3}\)) for CST\(_{20}\). Figure 4d...
shows that zooplankton are only present below the thermocline from November 2013 to June 2014 following the absence of HC (Figure 3b). Compared to the control, CST_{0} displays concentrations of zooplankton two times higher during March and April 2014.

Concerning the bacterial super-compartment in CST_{10}, BAC1 concentrations are only present below 30 m depth, with BAC2 predominating over the thermocline and reflecting effective biodgradation of HC from October 2013 to March 2014 (Figures 4f and S4d). Beneath the thermocline, the absence of HC in CST_{20} makes it possible for zooplankton to grow and graze on sinking phytoplankton at the density interface from November 2013 to April 2014. However, from February 2014, microzooplankton that feed on bacterioplankton are dominant, supporting the hypothesis that there is strong BAC1 activity at 20 m to 45 m depth (Figure 4d,f).

In comparison, the well-mixed experiment results show relatively homogeneous vertical distribution of each variable (Figure 5). Total bacterioplankton (Figure S5a), dominated by oil-degrading bacteria (80%) over the entire CWM_{20} simulation, show concentrations (up to 24 mmol N m^{-3}) in the same range as the CST_{20} results values (up to 22 mmol N m^{-3}; Figure S4). However, after December 2013, BAC1 increases up to four times the control concentrations (Figure 4f).

In early summer, concentrations of phytoplankton do not exceed 1.5 mmol N m^{-3} and 4 mmol N m^{-3} for the CWM_{0} and CWM_{20} simulations, respectively. However, in the CWM_{20} simulation, rapid biodegradation (Figure 2e) lowers the concentrations of HC, reducing its impact on zooplankton (Figure 2e). Consequently, after the spring bloom period, zooplankton concentrations exceed 1.0 mmol N m^{-3}, limiting phytoplankton growth to 2 mmol N m^{-3} due to grazing pressure (Figure 5d).

**Temporal Changes in Plankton Size Classes**

To illustrate temporal dynamics, phytoplankton and zooplankton concentrations resulting from the CST_{0}, CST_{10}, CST_{20}, and CST_{50} simulations were depth-integrated. In addition, because phytoplankton blooms are typically found in a two-layer structure in stratified experiments, we will concentrate on interpreting the stratified experiment only. Attention must be paid to the scale of Figure 6a, reduced from 290 mmol N m^{-2} to 50 mmol N m^{-2} for clarity.

In CST_{10} (Figure 6b), the diatom bloom reaches five times the values found in the CST_{0} simulation and persists until the end of November. Two diatom blooms occur during the third week of January 2014 and the third week of March 2014. Both are followed by mesozooplankton maxima. At this level of perturbation, zooplankton evolution is interesting. From the end of August 2013, microzooplankton concentration increases...
Results from CST 20 and CST 50 show diatoms are strongly dominant over the other groups. Micro- and mesozooplankton concentrations show similar increasing trends from the first week of November 2013 (Figure 6c,d).

**DISCUSSION**

**Physical Environment**

To understand the processes occurring within the plankton system during a chronic surface oil spill, we developed a one-dimensional coupled model that incorporates realistic physical conditions. Our results show that wind and tidally induced currents are the only contributors to the vertical mixing of biogeochemical variables in the SJG (Glorioso and Simpson, 1994; Glorioso and Flather, 1995; Palma et al., 2004). Temperature, the driving conservative property for stratification at the SJG latitude (Krepper and Rivas, 1979), is homogeneous during winter months. In the stratified experiment, a seasonal thermocline extended from spring to summer as previously described for the area by Akselman (1996). Moreover, satellite observations of sea surface temperature show similar values at the southeastern stratified region of the SJG (Glembocki et al., 2015).

In the well-mixed experiment, values of sea surface temperature are in accordance with Glembocki et al. (2015). Moreover, Carreto et al. (2007) show that vertically homogeneous conditions are also found in the southeastern SJG during spring and summer. The weak salinity variations are associated with the seasonal intrusion of the homogeneous low-salinity Magellan Plume at the south of the SJG (Krepper and Rivas, 1979; Palma and Matano, 2012). These homogeneous conditions agree with Palma et al. (2004), who point out that surface wind stress and strong tidally induced mixing due to bottom friction are the main external contributors to the vorticity balance in this region of the southwestern Atlantic shelf. Thus, in our model, HC distribution shows that contrasting turbulent regimes strongly affect
the distribution and dilution of the contaminant. In contrast, in the stratified experiment, higher HC concentrations are only found above the thermocline during spring and summer.

**Nutrients and Bacterioplankton Dynamics**

In both experiments, nitrate depletion is associated with earlier phytoplankton development (nitrate consumption) in the presence of HC. However, studies conducted on the biogeochemistry of the Deepwater Horizon spill did not find correlations between nitrate concentrations and the presence of an HC plume (Shiller and Joung, 2012; Dubinsky et al., 2013). Increasing ammonium concentrations starting at the end of summer 2014 are not consistent with Shiller and Joung (2012), but Horel et al. (2014) showed that temperature and the amount of oil affect nitrogen cycling rates, and that results additionally depend on the time of the year of the spill.

When HC is present in the system, OHC bacteria become significant only after two to three months. At the beginning of the spill, biodegradation is limited by the low LDON concentrations found toward the end of winter. After the first phytoplankton bloom, detritus production is enhanced, fueling LDON bacterial activity. While detritus and LDON dynamics in the presence of HC have not previously been documented in the literature, it is crucial to mention that regulation of biodegradation is clearly not limited to one type of nutrient (Atlas and Bartha, 1972; Atlas, 1981; Shiller and Joung, 2012). Thus, understanding the preference and timing of nitrogen-based nutrient consumption remains challenging. Our results suggest that the physical regime, including turbulence, plays an important role in biodegradation efficiency during HC contamination (Hassanshahian and Cappello, 2013). In the stratified experiment, biodegradation removes most of the HC by the first week of February 2014. In comparison, the well-mixed experiment shows faster incorporation of HC by OHC bacterioplankton, leading to the reduction of direct and indirect effects on plankton communities.

With HC altering bacterial dynamics, the higher concentrations of BAC1 beneath the thermocline in the simulation are associated with more intense detritus recycling and sinking. The shift that occurs in bacterial community structure in the upper water column in the CST20 simulation is in accordance with the decline of bacterial diversity and the succession of Gammaproteobacteria that Hazen et al. (2010) and Dubinsky et al. (2013) observed following the Deepwater Horizon spill.

**Phytoplankton and Zooplankton**

In the stratified experiment, HC indirectly affects phytoplankton, and concentrations reach values of up to 17 mg Chla m⁻³. These results are consistent with those of Johansson (1978), who reports increasing phytoplankton biomass during the Tsesis oil spill in the Baltic Sea. This inordinately high-intensity phytoplankton growth is attributable not only to the absence of grazing pressure from zooplankton, which are absent from the upper water column, but also to the increasing ammonium concentrations toward summer.

Without HC in the system, diatoms are the dominant phytoplankton group, with a spring bloom occurring in late September, as previously reported in the study area (Akselman, 1996; Rivas et al., 2006). During the bloom peak, both micro- and mesozooplankton concentrations begin to increase in our model through herbivorous grazing, thus limiting diatom growth. Unfortunately, no field information is available on these groups to validate our results.

While our results show high diatom
growth as an indirect consequence of HC contamination, results from other work are controversial: some studies show strong negative effects of HC contamination on this group (Hsiiao, 1976; Siron et al., 1996; Nayar et al., 2005; Sargian et al., 2007), and others highlight its persistence in oil polluted environments (Gordon and Prouse, 1973; Johansson, 1978; Teal and Howarth, 1984). Here, we show that among the indirect effects, the decrease of zooplankton grazing leads to an increase in phytoplankton abundance, as suggested by Johansson (1978). Similarly, Baker (1971) and Cabioch et al. (1981) previously observed the increase in bacterial activity associated with the recycling of dead microorganisms. On the other hand, an increase of microzooplankton can be linked to grazing on bacterioplankton occurring in the lower part of the water column where HC concentrations are ≤1 mmol N m⁻³ (Figure 2a).

Previously discussed results indicate that the response of phytoplankton to HC contamination is species-specific. Further investigation is needed to carefully parameterize the direct effects of HC on phytoplankton.

CONCLUSIONS
Modelling HC effects on a planktonic system requires consideration of several biogeochemical variables and processes. These model inputs result in a carbon flux imbalance in natural pelagic systems over an annual cycle. Our results suggest that in addition to the direct effects of HC on planktonic systems, indirect synergistic effects must also be considered.

The highly productive southeastern portion of the SJG shows different trends after subsurface incorporation of HC. In both experiments, at 10, 20, and 50 mmol N m⁻³ of HC contamination, biodegradation of HC is limited during winter and starts at the beginning of spring only when primary production increases and, consequently, LDON concentrations are sufficiently high. At 20 mmol N m⁻³, abnormally elevated concentrations of ammonium are linked to strong recycling of detritus by bacteria (BAC1). In this case, the detritus compartment is fueled by dead zooplankton that are related to HC toxicity and strong phytoplankton exudation. Because nitrification is not taken into consideration, incorporation of HC into the planktonic system is reflected in the accumulation of nitrogen in the ammonium compartment.

The great dominance of diatoms over the other planktonic compartments is associated with the indirect effects (decrease of herbivorous grazing and increase of detritus formation).

Finally, the dynamically contrasting well-mixed region shows lower HC concentration over the entire water column, resulting in faster bacterial consumption. Nevertheless, zooplankton are less affected by contamination under these physical conditions. Here, turbulent mixing plays a key role in the rate of encounter between HC and the different biological variables.

In future work, key physical and biological parameterizations should help us refine the results presented in this study. Sensitivity analyses must be conducted to understand the importance of biodegradation rates and nutrient dependency on biodegradation.

SUPPLEMENTARY MATERIALS
Supplementary materials are available online at https://doi.org/10.5670/oceanog.2018.413.

REFERENCES


