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Implementation of an end-to-end model of the Gulf of Lions ecosystem (NW Mediterranean Sea). I. Parameterization, calibration and evaluation

Bănaru Daniela*1, Frédéric Diaz*1, Philippe Verley2, Rose Campbell3, Jonathan Navarro1, Christophe Yohia1, Ricardo Oliveros-Ramos4, Capucine Mellon-Duval5, Yunne-Jai Shin6

1Aix Marseille Université, Université de Toulon, CNRS, IRD, MIO UM 110, Mediterranean Institute of Oceanography, Marseille, France.
2IRD, UMR 123 AMAP, TA40 PS2, Boulevard de la Lironde, 34398 Montpellier Cedex 5, France.
3Ecole d’Ingénieurs en Génie des Systèmes Industriels (EIGSI). 26, rue François de Vaux de Foletier, 17041 La Rochelle cedex 1.
4Instituto del Mar del Perú (IMARPE), Gamarra y General Valle s/n Chucuito, Callao, Peru.
5UMR MARBEC (IFREMER, IRD, UM, CNRS), 34203 Sète cedex, France.
6IRD, UMR 248 MARBEC, Université de Montpellier, Bat. 24 – CC 093 Place Eugène Bataillon 34095 Montpellier Cedex 5, France.

*Corresponding Authors: Tel: +33 (0)486 090 625 (D. Bănaru) : +33 (0)486 090 564 (F. Diaz)
Email addresses: daniela.Bănaru@univ-amu.fr (D. Bănaru), frederic.diaz@univ-amu.fr (F. Diaz),

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ABSTRACT

An end-to-end model named OSMOSE-GoL has been built for the Gulf of Lions, the main French Mediterranean fishing area. This spatialized dynamic model links the coupled hydrodynamic and biogeochemical model Eco3M-S/SYMPHONIE (LTL - low trophic level model) to OSMOSE (HTL - high trophic level model). It includes 15 compartments of living organisms, five from the LTL model (i.e. nanophytoplankton, microphytoplankton, nanozooplankton, microzooplankton and mesozooplankton) and ten from the HTL model (northern krill, southern shortfin squid, European pilchard, European anchovy, European sprat, Atlantic horse mackerel, Atlantic mackerel, blue whiting, European hake and Atlantic bluefin tuna). With the exception of northern krill and European sprat, all HTL species are commercially exploited and undergo fisheries mortality pressure. The modeled species represent more than 70% of annual catches in this area. This paper presents the parameterization, calibration and evaluation of this model with satellite data for phytoplankton and with biomass, landings, diet and trophic level data for HTL groups. For most species, the diets in output of OSMOSE-GoL are similar to field and literature data in terms of dominant prey groups and species. However, some differences were observed. Various reasons may explain the mismatch between the modelled diet and field data. Benthic prey sometimes observed in the stomach content of the HTL predators were not modelled in OSMOSE-GoL. Field studies were carried out at specific periods and locations, while our data concern the period 2001-2004 and the entire modelled domain. Inter- and intra-annual variations in spatial distribution and density of prey may also explain these differences. The model estimates trophic level values similar to those cited in the literature for all the HTL compartments. These values are also close to the trophic levels estimated by a previous Ecopath model for the same area and period. Even though some improvements are still possible, this model may already be of use to explore fishery or Marine Protected Areas scenarios for socio-ecosystem management issues.
1. Introduction

End-to-end (E2E) models are particularly appropriate to disentangle the intricacy of interactions occurring between physical forcing and low and high trophic level communities in the context of a quantitative approach dedicated to Ecosystem-Based Management (EBM) (e.g. Travers et al., 2007; Rose, 2012). They use multiple field data sets and are able to assess and simulate the dynamics of the main descriptors of the ecosystem rather than evaluating single resources and single threats (Shin et al., 2010; Christensen and Walters, 2011; Collie et al., 2016). Similar approaches, driven by modeling and information are used not just in ecology, but in many other areas to improve, and in some cases save our lifes (Helbing et al., 2015).

However, the implementation of end-to-end modelling remains challenging, mainly due to the major differences between the sub-models of hydrodynamics and Low Trophic Level (LTL) organisms on the one hand, and that of High Trophic Level (HTL) organisms on the other hand (see review by Rose et al., 2010). The challenges are numerous and concern both concepts (e.g. representation of the zooplankton key level, differential scaling of processes, behavioral movement of HTL organisms, etc.) and technical issues (e.g. different programming languages and time-steps).

One of the main challenges concerns the nature of link between sub-models. Travers et al. (2009) considered two possible types of links between LTL and HTL models. In the one-way forcing mode, LTL groups’ biomasses serve as prey fields to HTL groups, without any feedback on the LTL compartments. In the two-ways coupling mode, the biomass of the LTL groups serve as prey field for HTL groups and an explicit rate of HTL-induced predation is specifically applied as feedback on each of the LTL groups.

Over the last decade, E2E modelling studies applied to regions or to the whole basin of the Mediterranean Sea have flourished in the context of EBM (Coll and Libralato, 2012). This research trend has been mainly driven by the more and more numerous observations of increasing threats and impacts on the Mediterranean marine ecosystems due to the exponential development of anthropogenic activities (e.g. Lötze et al., 2011; Coll et al., 2012). Most of these modelling studies
have been based on the Ecopath with Ecosim (EwE) model (see review of Coll and Libralato, 2012), and only a few have used alternative models, spatial and multispecies models such as the OSMOSE size-based model (Halouani et al., 2016), or age-structured models applied to single species (e.g. Santojanni et al., 2005).

Furthermore, while implementation of the EBM of the Gulf of Lions (GoL) is particularly crucial owing to its major contribution to Mediterranean fisheries catches (Sacco, 2008; Demaneche et al., 2009), this shelf area has been poorly investigated to date (Coll and Libralato, 2012). Only the recent study by Bănaru et al. (2013) dealt with the fishing impact on the trophic structure of the marine ecosystem using an EwE approach. However, this study is not spatialized and is based on some crude assumptions, concerning for example the plankton compartment (prey for the planktivorous fish species), for which the biomass level is determined from the literature and satellite imagery. A major bias of this type of modelling approach is that it does not account for the close coupling between the physical and biological (sensu largo) processes. Yet these interactions occur at multiple spatial and temporal scales in the NW Mediterranean Sea, and it has now been well-demonstrated that they have a significant impact on the dynamics and the spatial distribution of marine organisms from plankton to top predators (e.g. Fromentin et al., 2003; Niewiadomska et al., 2008; Cotté et al., 2011; Campbell et al., 2013). It is therefore necessary to represent the Gulf of Lions ecosystem more realistically, in particular the environmental forcing, the spatial dynamics of living organisms and their interactions, to enable a finer analysis of the functioning of the ecosystems and in fine to plan its optimal management for the next decades.

The E2E approach developed in this study is based on a fully dynamic coupling (i.e. two-ways coupling) of two pre-existing sub-models representing the dynamics of LTL organisms driven by hydrodynamics and climate processes on the one hand, and the dynamics of HTL organisms impacted by fishing activities on the other hand. The first model component is the Eco3M-S/Symphonie model that has been successfully used in the North-Western Mediterranean Sea to advance our understanding of the influence of hydrodynamics and atmospheric drivers on the
distribution of plankton at different spatial and temporal scales (e.g. Auger et al., 2011, 2014; Campbell et al., 2013; Carlotti et al., 2014). The second component model is the individual- and size-based model OSMOSE (Shin and Cury, 2004). This HTL model has been applied worldwide in order achieve better understanding of the functioning of diverse marine ecosystems (e.g. Travers et al., 2006; Marzloff et al., 2009; Fu et al., 2013; Grüss et al., 2015, 2016; Houxi et al., 2016).

The aim of the present paper is to document an E2E model developed for the GoL (the OSMOSE-GoL model) and based on dynamic feedback (two-ways coupling) between two pre-existing LTL and HTL sub-models. Numerous data sets available in this marine region have been used to calibrate and quantitatively evaluate both the LTL and the HTL modules of the OSMOSE-GoL model. An application of this model that consists in an analysis of the impacts of the predation pressure exerted by HTL planktivorous species on the spatial distributions, the structure of the LTL community and food webs controls is presented in a companion paper (Diaz et al., 2019).

2. Methods

2.1. The E2E modelling approach

The approach developed in this study is based on the coupling of two existing sub-models. The first is the Eco3M-S/Symphonie model (Campbell et al., 2013) that represents the dynamics of Low Trophic Level (LTL) organisms driven by hydrodynamics and climate processes. The second is the individual-based model OSMOSE (Shin and Cury, 2004; Grüss et al., 2015), that simulates the dynamics of High Trophic Level (HTL) organisms. Both models have been fully described in previous works, therefore only the main characteristics are given hereafter. They are coupled in two distinct modes (Figure 1). In the one-way forcing mode, the biomass outputs of the LTL model are provided as inputs for the OSMOSE model without any feedback on the LTL biomass. In the two-ways coupling mode, there is a dynamic feedback between the two models through the predation process: the biomass outputs of the LTL model are provided as inputs for the OSMOSE model, which provides in return an additional rate of predation by the HTL planktivorous organisms.
2.1.1. Description of the LTL model

The LTL model is composed of two coupled models: the Symphonie hydrodynamic model and the Eco3M-S biogeochemical model (Campbell et al., 2013). The meteorological and hydrodynamic processes influencing the spatial and temporal distributions of nutrients and plankton were simulated by the Symphonie model (Marsaleix et al., 2008), a 3-D primitive equation, free surface model, based on hydrostatic and Boussinesq approximation. This model has already been used to successfully represent certain physical processes in the Northwestern Mediterranean Sea (e.g. Dufau-Julliand et al., 2004; Ulses et al., 2008; Kersalé et al., 2013). The Symphonie version used here has been developed by Hu et al. (2011a). The modeled zone (711 km by 303 km) extends over the NW Mediterranean Sea, including the whole of the Gulf of Lions and parts of the Ligurian and Catalan Seas (Fig. 2). The grid used a square horizontal mesh with a spatial resolution of 3 km by 3 km. Sigma coordinates were used on the vertical dimension with a maximum of 40 levels. The model was run from January 9, 2001 to December 24, 2004. All details on the initial and boundary conditions are given in the studies of Hu et al. (2011a) and Campbell et al. (2013).

The biogeochemical model Eco3M-S is embedded in the Eco3M platform (Baklouti et al., 2006a,b), and is a multi-nutrient and Plankton Functional Types (PFT) model that simulates the dynamics of several biogeochemical decoupled cycles of biogenic elements (carbon, nitrogen, phosphorus and silica) and non-redfieldian plankton groups. The Eco3M-S version has been recently used and validated in the studies of Hu et al. (2011a,b) and Campbell et al. (2013) for the biogeochemical components and the hydrodynamics features, respectively. The model structure encompasses seven compartments of living organisms. Two of the three PFT of autotrophs of the model, from the smallest to the largest, were accounted for: (1) nano-phytoplankton, NANOPHY (2–20 μm) that dominate the biomass of phytoplankton assemblages for most of the year (Marty et al., 2002; Marty and Chiavérini, 2010), with a heterogeneous taxonomic composition (e.g. autotrophic dinobionts); and (2) the micro-phytoplankton community, MICROPHY (20-200 μm), largely dominated by...
phytoplankton silicifiers (mainly diatoms) that can for certain periods contribute to a significant part of primary production and biomass during spring bloom in the NW Mediterranean Sea (Marty et al., 2002; Marty and Chiavérini, 2010). Three of the four PFTs of heterotrophs of the model, from the smallest to the largest, were considered: (1) nano-zooplankton, NANOZOO (5-20 μm, mainly bacterivorous dinobionts and small ciliates) that consume the smallest phytoplankton groups (<2 μm) and bacteria; (2) micro-zooplankton, MICROZOO (20-200 μm, mainly most of ciliates groups and large dinobionts), having characteristics (growth, ingestion rates) close to NANOZOO but with a wider prey spectrum, especially with potential consumption of micro-phytoplankton; and (3) meso-zooplankton, MESOZOO (>200 μm, mainly copepod groups but also including amphipods) grazing on the largest categories of plankton (>20 μm, micro-phytoplankton and micro-zooplankton) and producing fast-sinking fecal pellets. All the formulations of the biogeochemical processes, as well as the whole set of parameters, have been extensively described in Campbell et al. (2013). Constant mortality rates were applied to some of phytoplankton and zooplankton groups (Table 1), representing either senescence or viral attacks or predation.

2.1.2. Description of the HTL model

The OSMOSE (Object-oriented Simulator of Marine ecOSystEms, Shin and Cury, 2001; Shin and Cury, 2004) model (version Osmose 3.2, www.osmose-model.org/downloads) is a two-dimensional spatially explicit, individual-based model (IBM), written in Java (www.osmose-model.org), and based on the main assumption of opportunistic and size-based predation. OSMOSE is a multispecies model representing the whole life cycle of several interacting species, from eggs and larvae to juveniles and adults. At the first time step following the production of eggs, the total number of eggs of each population is split into super-individuals called “schools”, spatially distributed according to the input distribution maps. At each time step, OSMOSE simulates the biological and ecological processes at the super-individual level: growth, predation and forage, reproduction, natural and starvation mortalities as
well as fishing mortality (Figure 1). The different sources of mortality of schools (fishing,
predation, starvation and diverse mortality) occur in a random order. Two types of movements are
considered in the OSMOSE-GoL model: (1) ontogenetic and seasonal migrations, taken into
account through the use of distribution maps; and (2) small-scale random diffusion, when the
distribution maps of schools (depending on fish age, stage, size, and season, year) does not change
from one time step to the next.

2.1.3. Technical details on the two-ways coupling mode

The two-ways coupling between the OSMOSE-GoL and Eco3M-S/Symphonie models was
performed through the predation process. Outputs of plankton groups provided by the LTL model
serve as prey fields for the HTL organisms, which return an additional predation mortality in the
plankton groups. To circumvent the different spatial dimensions (3D vs. 2D) and units (mmolN (or
C) m$^{-3}$ vs. tons wet weight) of the two models, plankton concentrations were vertically integrated
and converted into biomass using conversion factors (Table 1). Only a small portion of plankton
biomass is available to fish and macroinvertebrates due to various processes affecting their vertical
distribution (turbulence, migrations, etc.). Availability of plankton to HTL species is not easy to
assess in the field, and literature on this point is non-existent. Therefore, the availability parameters
$a_p$ ($p$ for a given plankton group $p$) were estimated (Table 2) via the calibration of the model (see
hereafter).

The two-ways coupling mode meant that the HTL model returns a specific mortality rate for each
plankton group over space and time, and these rates were computed from the amount of prey
ingested, as described in Travers et al. (2009). In each cell (x,y) of OSMOSE-GoL and for each
plankton group ($p$), the HTL-induced mortality rate at time $t+\Delta t$ was computed as the total biomass
of plankton eaten ($BE_{\Delta t}$) during the time step $\Delta t$ (15 days) over the mean total plankton biomass
($B_{\Delta t}$) at time $t$ multiplied by $\Delta t$ (Eq. (1)). As the maximum biomass of plankton $p$ eaten by HTL
organisms at time $t+\Delta t$ is the available biomass $a_p \cdot B(x, y, t, p)$ at time $t$, the HTL-induced
mortality rate can thus vary between 0 and $\frac{a_p}{\Delta t}$. Because this variable mortality was added to the natural mortality ($m_p$) already considered in the Eco3M-S model (Table 1), the latter rate was reduced to ($m_p - \frac{a_p}{2 \cdot \Delta t}$), with ($\frac{a_p}{2 \cdot \Delta t}$) being the median of the variable mortality due to HTL species. It was not set to zero in order to account for other sources of mortality such as predation by non-modelled organisms, senescence and starvation mortality. Outside the common domain between the Eco3M-S/Symphonie and OSMOSE-GoL models (Figure 2), the plankton mortality rate was set to $m_p$. According to the equation (1), the plankton total mortality rate thus ranges between $m_p - \frac{a_p}{2 \cdot \Delta t}$ and $m_p + \frac{a_p}{2 \cdot \Delta t}$. This rate can be either lower or higher than the initial mortality rate $m_p$ depending on the predation pressure induced by HTL organisms.

2.1.4. Design of the numerical experiment

A first spin-up period of 35 years was launched in the one-way forcing mode to reach equilibrium of the HTL model outputs. This step was achieved using the numerical fields of plankton biomass in 2001. Following this period of spin-up, the model was then run in the one-way forcing mode (years 36 and 39) using the plankton biomass in 2001, and then in the two-ways coupling mode for four years (40 to 43). For the last four years (40 to 43), the HTL model received the numerical fields of LTL biomass for the years 2001 to 2004. These years was chosen because the LTL model has been previously validated over this period (Campbell et al., 2013). The coupling simulation started on January 10, 2001 (00h00) and it ended on December 20, 2004 (00h00). Years 36 to 39 (one-way forcing) as well as the period of two-ways coupling mode (years 40 to 43) were
considered for analysis hereafter. Furthermore, a set of 50 replicated simulations was run to account for the stochasticity of the OSMOSE model.

2.2. Parameterization of the HTL model

2.2.1. Modelled domain and selected HTL groups

In this study, the OSMOSE-GoL model grid consists of 19 by 15 cells with a resolution of 12 km by 12 km covering the GoL area north of a line running from 42°04’-3°18’ to 43°05’-5°37’ (Figure 2). A set of 25 schools released per time step per species (600 per year per species) was chosen for this model as a compromise between the stochasticity of the model and numerical limitations in memory and calculation speed.

As simulations with OSMOSE necessitate extensive information on entire life cycles, only 10 HTL key species, being the most representative of the pelagic and demersal food web, were included in the configuration of OSMOSE GoL: northern krill (*Meganyctiphanes norvegica*, (Sars 1857)), southern shortfin squid (*Illex coindetii*, (Vérany 1837)), and eight fish species from small pelagic fish to demersal fish: European pilchard (*Sardina pilchardus*, (Walbaum 1792)), European anchovy (*Engraulis encrasicolus*, (Linnaeus 1758)), European sprat (*Sprattus sprattus*, (Linnaeus 1758)), Atlantic horse mackerel (*Trachurus trachurus*, (Linnaeus 1758)), Atlantic mackerel (*Scomber scombrus*, (Linnaeus 1758)), blue whiting (*Micromesistius poutassou*, (Risso 1827)), European hake (*Merluccius merluccius*, (Linnaeus 1758)), Atlantic bluefin tuna (*Thunnus thynnus*, (Linnaeus 1758)). These species represent more than 70% of annual catches (Demaneche *et al.*, 2009). Some of them were selected because of their importance for fisheries (European pilchard, European anchovy, Atlantic horse mackerel, Atlantic mackerel, European hake, Atlantic bluefin tuna) (SIH, 2017), others for their importance as forage species such as northern krill, European sprat and blue whiting (Bănaru *et al.*, 2013). All these species represent the most important ones in terms of structure and functioning of the food web in this area (Bănaru *et al.*, 2013).
To parameterize the model, various input information items were needed, including: (1) spatial distribution maps for different life stages and time steps (Appendix 1). These maps coded for presence vs. absence for each species. They have been obtained from geo-referenced data of the PELMED and MEDIT research surveys conducted since 1993 (IFREMER databases). Atlantic bluefin tuna is the sole highly migratory species with only a seasonal presence in the modelled area and not reproducing in the GoL (Imbert et al., 2007); (2) predation, growth and reproduction parameters (Table 4); (3) mortality parameters (Table 1); (4) fishing and reproduction seasonality (see sections 2.2.2 and 2.2.3).

2.2.2. Predation, growth and reproduction processes

In the OSMOSE model, predation is assumed to be an opportunistic process and occurs when there is both size adequacy and spatio-temporal co-occurrence between predator and prey. Within a cell of the grid, a predator can feed on a co-occurring prey if: (1) the prey is of a suitable size, that is, within a range determined by the minimum and maximum predator/prey size ratios; and (2) the vertical distribution of the prey makes it accessible to the predator, which is determined by the accessibility coefficients provided to OSMOSE-GoL. Therefore, the food web (or diet matrix) builds up as an emergent property of local trophic interactions (Travers et al., 2009).

Minimum and maximum predator/prey size ratios (Table 3) were parameterized differently for different size classes for each taxon, in order to account for ontogenetic changes in feeding behavior. These size ratios were built from local data on diet, predators and prey size (Labat and Cuzin-Roudy, 1996; Båmstedt and Karlson, 1998; Imbert et al., 2007; Le Luherne, 2012; Bourgogne, 2013; Le Bourg et al., 2015; Mellon-Duval et al., 2017).

The accessibility coefficients were set by default at 0.8 for HTL groups. For some species, these coefficients were reduced to 0.6 in relation to their exclusive benthic location during early life stages (northern krill <1 cm, southern shortfin squid <2 cm and European hake <6 cm), or to the very coastal area distribution patterns of individuals <5 cm and thus low accessibility to predation.
During the predation process, if enough prey items are present in a spatial cell, a predator feeds upon them uniformly until it reaches satiation. Predation efficiency is defined as the ingested prey biomass over the maximum biomass a predator can feed upon. For each species, the maximum ingestion rates (MIR, Table 3) have been assessed from local data (Palomares and Pauly, 1998; Bănaru et al., 2013). However, some of the considered species consume a non-negligible part of benthic prey that was not considered in the Osmose-GoL model. In order to avoid artificially increasing predation on pelagic and demersal prey, their MIR were proportionally reduced by 35% for southern shortfin squid, Atlantic horse mackerel, Atlantic mackerel and European hake (Kaci, 2012; Le Luherne, 2012; Mellon-Duval et al., 2017) and by 12.5% for blue whiting (Bourgogne, 2013). When the predator does not ingest enough food to fulfill its maintenance requirements (corresponding to a predation efficiency threshold of 0.57; Shin and Cury 2004), fish starve at a rate which decreases linearly with predation efficiency (Shin and Cury, 2004). The maximum mortality rate by starvation was fixed at 1.0 year$^{-1}$, applied during a time step in the absence of food. Predation efficiency also determines fish growth rate during a time step. When the biomass of prey eaten is higher than maintenance requirements, the growth rate of fish is positive, varying as a function of the von Bertalanffy growth rate and the predation efficiency (Shin and Cury, 2004). The growth parameters (Table 4) were computed from local studies (Farrugio et al., 1991; Campillo, 1992; SCRS, 1997; Mellon-Duval et al., 2009; GFCM-FAO, 2011a, b; PELMED and MEDITS, IFREMER campaigns). Longevity was estimated from literature data (Campillo, 1992; Labat and Cuzin-Roudy, 1996; Sánchez et al., 1998).

Predation success has also an indirect effect on the reproduction process through the biomass of spawners which, combined with relative fecundity parameters, defines the number of eggs released in the system. Any school of the key populations, whose size is greater than the sexual maturity size ($S_{mat}$), reproduces according to the spawning seasonality and to the species relative annual fecundity.
$\varphi$, number of eggs spawned per gram of mature female per year) (Shin and Cury, 2004).

Data for size at maturity ($S_{\text{mat}}$) are indicated in Table 4 (Higginbottom and Hosie, 1989; Fárrugio et al., 1991; Campillo, 1992; Labelle et al., 1997; Sánchez et al., 1998; Lleonart, 2001; Sinovčić et al., 2004; GFCM-FAO, 2011a, b; www.fishbase.org; PELMED and MEDITS, IFREMER campaigns).

Sex ratio was hypothetically fixed at 0.5.

The relative fecundity (Table 4) has been recomputed considering the number of eggs using data from Ross and Quentin (1986), Campillo (1992), Laptikhovsky and Nigmatullin (1993), Qúero and Vayne (1997), Sánchez et al. (1998), and the estimated weight of mature females based on the total length for each species.

Seasonality of reproduction was estimated using data from Farrugio et al. (1991), Campillo (1992), Labat and Cuzin-Roudy (1996), PELMED and MEDITS, IFREMER campaigns. Egg weight (0.54 mg) and size ($S$) (Table 4) were also indicated in the model (Qúero and Vayne, 1997; Le Bourg, pers. comm.) as eggs represent potential prey for the rest of the food web.

### 2.2.3. Fishing and natural mortality processes

Fishing pressure is represented through a population-specific fishing mortality rate $F$ (Table 2), affecting the number of fish per school when larger than the specified size of recruitment to fisheries. Legal size of catch ($S_{\text{rec}}$, Table 4) has been used as recruitment size in the OSMOSE-GoL model. Initial estimates of annual fishing mortality rates have been estimated for each target species, using the landings to biomass ratio obtained from stock assessments (PELMED, MEDITS, GCFM) and from the fisheries database (SIH, 2017). These mortality rates were subsequently refined through the calibration of the model (see 2.3). Fishing mortalities are assumed to be spatially homogeneous, but can vary seasonally as specified in input (Table 5). Fishing seasonality was estimated using catch data by species (SIH, 2017).

In addition to explicit predation mortality modelled in Osmose, the mortality of the first life stages (eggs and first feeding larvae, $M_0$, Table 2) is due to different natural causes (e.g. non-fertilisation of
eggs, starvation of first feeding larvae, advection, sinking) as well as predation by organisms not considered in the OSMOSE-GoL model. Additional sources of natural mortality concerning other life stages have also been accounted for (M, Table 2), including mortality due to disease, senescence and predation by organisms not represented in the OSMOSE-GoL model (e.g. birds, mammals, etc.). Since it is usually very hard to quantify these types of mortality, they have been assessed through the calibration of the model for each of the ten HTL species.

2.3. Calibration for the HTL model

The OSMOSE-GoL model has been calibrated using an optimization technique based on an evolutionary algorithm, and a maximum likelihood based objective function, so that the modeled biomass and landings of the ten HTL species remained within realistic observed ranges. The data sources used for the biomass are from the PELMED and MEDITS cruises performed every year in June and July from 2000 to 2013 and GFCM-FAO stock assessments for the GoL (Jadaud, pers. comm.; Bigot, pers. comm.; Sarrazin, pers. comm.), VPA models for hake and tuna (Jadaud, pers. comm.; Fromentin, pers. comm.) and SIH (2017) for landings.

The calibration was performed with the calibraR package (Oliveros-Ramos et al., 2015; Oliveros-Ramos and Shin, 2016; https://CRAN.R-project.org/package=calibraR) that has been specifically developed for calibrating complex ecological models, and osmose2R (http://www.osmose-model.org, https://CRAN.R-project.org/package=osmose), which includes a set of R functions for interfacing Osmose with calibraR.

The calibration step aimed at providing estimates of the following parameters of the model: (i) availability coefficients of plankton groups to HTL species, (ii) larval mortality rate for each species and (iii) fishing mortality rates for each species (Table 3). These parameters have been selected because they are highly model-dependent, i.e. their meaning depends on the model structure and assumptions, and there are no reasonable initial estimates for plankton accessibility and larval mortality (Oliveros-Ramos et al., 2015). The objective function minimized by the algorithm is built
automatically by the R package by aggregating lognormal-likelihood functions that quantify the fit between model outputs and field observations. Biomass and landings from year 2001 were given to the algorithm for every modeled species as the observed data, and the OSMOSE-GoL model was forced with the LTL biomass of the Eco3M-S/Symphonie model from the same year. In this way, a “steady-state” calibration of the OSMOSE-GoL model has been achieved for year 2001, that is the first year of the coupled simulation.

The calibration step enabled us to obtain a set of parameters so that biomass and landings of the considered HTL species best range between the minima and maxima of the observed biomass in the GoL. This parameter set was then used in the coupled model.

2.4. Datasets for the evaluation of the E2E model

2.4.1. Evaluating phytoplankton biomass of the LTL model using remote sensing satellite data

Ocean color data from the GlobColour project (www.glocolour.info/) were used in this study to evaluate the realism of the modelled surface chlorophyll \( a \) concentrations (considered as a proxy of phytoplankton biomass). The GlobColour product takes advantage of gathering data sets derived from several ocean color sensors (ENVISAT, MODIS, MERIS and SeaWiFs). Three sensors (SeaWiFS, MERIS and MODIS) showed a temporal overlap of functioning from April 2002 to December 2010, and it was possible to perform a quantitative comparison between satellite and modeled chlorophyll \( a \) values over most (990 days) of the modelled period (1440 days).

The GlobColour product provided a weighted mean \( \langle Chl_w \rangle \) and a weighted error \( \epsilon(\langle Chl_w \rangle) \). The detailed computation of these parameters is given in the Product User Guide of GlobColour available on the aforementioned web site.

In order to accurately compare the concentrations of the satellite-derived chlorophyll \( a \) to those modeled, the numerical concentrations of total chlorophyll (sum of chlorophyll \( a \) concentrations of pico-, nano- and microphytoplankton) were averaged over the first optical layer (\textit{sensu} Bricaud et al., 2010), \( \langle Chl_{opt1} \rangle \). The details of the computation are given in the recent study of Campbell et al.
The model-derived chlorophyll $a$ values were re-interpolated point by point at the size of
each pixel (1.1 km) of the satellite image on the modeled domain. The period of assessment (990
days) was characterized by more than a third (~34%) of cloudy days without useable pixels on the
modeled area, therefore the comparison was made on the remaining days, corresponding to a total
of 236877 pixels.

A metric of model to data comparison (C parameter) was built and used hereafter. This metric
assessed the size of the discrepancies between the predicted and satellite values as follows:

$$C = \begin{cases} 0, & \text{if } Chl_{opt,1} \in [Chl_w - \epsilon(Chl_w), Chl_w + \epsilon(Chl_w)] \\ \min \left\{ \left| \frac{Chl_{opt,1}}{Chl_w} - \left( \frac{Chl_w}{Chl_w} + \epsilon(Chl_w) \right) \right|, \left| \frac{Chl_{opt,1}}{Chl_w} - \left( \frac{Chl_w}{Chl_w} - \epsilon(Chl_w) \right) \right| \right\}, & \text{otherwise} \end{cases}$$

2.4.2. Evaluating the HTL model using diet and trophic level data

Diet data for each of the modeled HTL species obtained as output of the OSMOSE-Gol
model were compared with in situ field data obtained from stomach content analyses (Le Bourg et
al., 2015; Bănaru, 2015; Bănaru, pers. comm.) or literature data (Båmstedt and Karlson, 1998; Sara
and Sara, 2007; Mellon-Duval et al., 2017) for similar individual sizes.

Estimated trophic levels (TL) from the OSMOSE-Gol model were compared with TL values from
the literature based on the stomach content (Båmstedt and Karlson, 1998; Stergiou and Karpouzi,
2002; Bănaru, 2015; Bănaru et Harmelin-Vivien, 2017) and compared with those of a local
ECOPATH model (Bănaru et al., 2013).

The aim of these comparisons was to check whether the parameterization and the processes of the
model were able to reproduce trophic interactions close to those observed in field data.

3. Results and discussion

3.1. Calibrated biomass and landings of the HTL model against data

The calibration of the model enabled us to estimate some parameters of the HTL model such
as the accessibility coefficients of the plankton and larval mortality of HTL species, and to refine the
fishing mortality estimates. The success of this calibration step lies in the ability of the model, once
run with this set of calibrated parameters, to provide results remaining within realistic ranges for the
biomass and landings of HTL species. The modeled median biomass (Figure 3) and landings (Figure
4) of the ten HTL species and their corresponding envelopes delimiting 0.25 and 0.75 percentiles are
presented for years 36 to 39 in the one-way forcing configuration. The modeled biomass and landings
for years 40 to 43, resulting from the two-ways coupling mode, are presented and discussed in the
companion paper (Diaz et al., 2019) for the detailed comparison with one-way forcing mode effects.
Most of the modeled biomass of HTL species showed seasonal patterns of change in median values
within the ranges of the field observations (Figure 3), except for that of European hake, which is
slightly overestimated. The biomass of certain species such as the European hake is known to be
underestimated, due mainly to the difficulty of catching large individuals during scientific surveys
(Bigot, pers. comm.). The median simulated biomass of northern krill (Figure 3A) showed a seasonal
dynamic with the highest values occurring in winter and the lowest at the end of the spring. The
seasonal patterns of change of southern shortfin squid (Figure 3B) showed median levels of biomass
around 2.5 x 10^4 tons, with maximum values at the end of summer and minimum values at the end of
winter. The stock of European pilchard (Figure 3C) showed cyclical variations around 2 x 10^5 tons,
with maxima reached in spring and minima at the beginning of the winter. The median biomass of
European anchovy (Figure 3D) showed cyclical seasonal variations centered on 10^6 tons, and
characterized by highest biomass from mid-winter to the end of spring, and the lowest during autumn.
The seasonal variation of the European sprat (Figure 3E) biomass was very low. The median levels
were around 1 x 10^4 tons, with a barely discernible peak at the end of summer. The seasonal dynamics
of Atlantic horse mackerel (Figure 3F) showed an increase in biomass at the end of autumn. A very
weak seasonal signal in the biomass of Atlantic mackerel was simulated (Figure 3G), with the highest
biomass extending from winter to mid–spring, and a minimum occurring from mid–summer to mid–
autumn. The biomass of blue whiting (Figure 3H) did not show any clear seasonal pattern and
oscillated around $4 \times 10^4$ tons. The temporal changes of European hake biomass (Figure 3I) also showed a very weak seasonal signal, with values around $1.4 \times 10^4$ tons. By contrast, the biomass of Atlantic bluefin tuna (Figure 3J) showed a marked seasonal cycle, with a minimum of biomass during winter ($-5 \times 10^3$ - $-5.5 \times 10^3$ tons), followed by a sharp increase to reach a first seasonal maximum at the beginning of the spring ($-5.6 \times 10^3$ - $-6.2 \times 10^3$ tons). Another biomass minimum occurred at the beginning of summer, and then the annual peak (around $7 \times 10^3$ tons) is observed in autumn. There were no seasonal biomass field data to use for comparison with model output, as scientific surveys were only organized during summer.

On the whole, for most of the HTL species, the whole or at least *pro parte* interannual dynamics of modeled median landings (and the corresponding range of percentiles) were within the ranges of observed data (Figure 4). However the median landings of southern shortfin squid (Figure 4B) and of Atlantic mackerel (Figure 4F) were much higher than those observed, by a factor of 16 ($\approx 800$ tons) and 2.5 ($\approx 500$ tons) with regard to observed landings, respectively, without any marked seasonal pattern. A possible explanation for this discrepancy could be the underestimation of the landings of southern shortfin squid and Atlantic mackerel in SIH (2017) landings databases, as for many other demersal and benthic species, because fishermen often directly sell their fish at local markets (CRPMEM PACA, 2016). Landings of northern krill and European sprat (Figures 4A and 4E) are not computed by the model as they are not landed by fishermen. The simulated temporal dynamics of planktivorous fish such as European pilchard and European anchovy showed marked and steady seasonal cycles. The median landings for European pilchard (Figure 4C) were minimum ($\approx 180$ tons) at the beginning of winter. They rapidly increased during winter and reached a relative maximum ($\approx 400$ tons) at the beginning of spring. Then, they briefly fell below 400 tons and again increased up to their absolute maximum ($\approx 750$ tons) at the beginning of summer. They sharply decreased from the middle of summer to the end of the autumn down the absolute minimum. Maximum simulated values were higher than the maximum observed values during the first half of summer. The simulated landings of European anchovy (Figure 4D) also showed a minimum ($\approx 100$ tons) at the beginning of
winter, but the catch then increased very rapidly up to a first peak of ~180 tons, and an absolute maximum (~250 tons) at the end of spring. The summer season is marked by a sharp decrease in landings, and a third peak of landings (~200-220 tons) in autumn. The whole set of seasonal modeled values remained within the range of observed landings. The seasonal changes in median landings for Atlantic horse mackerel (Figure 4F) were characterized by two maxima in spring and autumn. The spring maximum comprised between 20 and 28 tons was generally higher than the autumn maximum (<20 tons). In parallel, minima (around 10 tons or lower) were simulated during winter and summer. Only the maximum values of landings during spring and autumn fell within the range of observed landings. The temporal patterns of change in simulated landings for blue whiting (Figure 4H) showed a very marked seasonal trend. Median landings were close to zero from the beginning of summer to the end of autumn, they sharply increased to reach ~40 tons at the beginning of spring, and then drastically dropped during spring. The modeled landings were underestimated during the first half of the year. The median landings of the European hake (Figure 4I) remained around 100 tons from mid-winter to the end of summer within the range of observed landings during this period of year. They fell sharply during autumn to well below the lower limit of observed data, with a minimum of ~10 tons, before increasing again at the beginning of winter. Modeled landings of Atlantic bluefin tuna (Figure 4J) were close to zero only over a short period of the year, mainly from the end of autumn to the beginning of winter. Catches sharply increased up to 250 tons during spring, before slightly declining to just below 200 tons at the beginning of summer. Summer was marked by an abrupt increase in the catches, reaching 500 tons at the end of summer. This seasonal cycle is characterized by simulated landings that were close to field data.

### 3.2. Comparing phytoplankton surface biomass of the LTL model with ocean color data

The seasonal cycle of surface satellite-derived chlorophyll \(a\) content averaged over the modeled area from April 2002 to December 2004 (Figure 5) is typical of those usually observed at oceanic mid-latitudes (e.g. Siegel et al., 2002), with average minima during summer months and
maxima from winter to late spring. The very low summer concentrations (~10^{-1} mg m^{-3} or lower) are considered as representative of the Mediterranean Sea (Bosc et al., 2004). As expected, the seasonal pattern found during the study period was close to the “bloom” trophic regime according to the classification of D’Ortenzio and Ribera d’Alcalà (2009) for the Mediterranean Sea.

An interannual variability of the seasonal cycle could be detected, with the year 2003 being characterized by a longer summer period of low chlorophyll \( a \) than 2002 and 2004. The year 2003 did not show any clear autumnal bloom, while it was particularly marked in mid-September 2002 and September and November 2004. The inter-annual variability in the phytoplankton surface content in this area of the Mediterranean Sea is a common feature that has been already observed in some other remote sensing time-series (Bosc et al., 2004; Auger et al., 2014) and in situ data (e.g. Marty and Chiavérini, 2010; Gernez et al., 2011).

Beyond the seasonal dynamics, the time-series highlights the variability of mean values at very high frequency (i.e. day-to-day), whatever the season considered. Most of the time, large error bars also suggest a strong spatial heterogeneity of phytoplankton content in the NW Mediterranean Sea. This variability at high frequency and the strong spatial heterogeneity are mainly due to the multiple physical forcing occurring in this marine area, such as changing wind gusts that drive intricate (sub)-mesoscale hydrodynamic processes (e.g. eddies, see Hu et al., 2011a,b) and upwellings (Millot, 1999), river inputs discharging large amounts of nutrients throughout the year (e.g. Minas and Minas, 1989) and the large-scale circulation of the Northern Current sometimes intruding on the shelf (Petrenko, 2003). All the aforementioned physical processes interact together at different spatial and temporal scales, and play a key role in setting ocean dynamics, heat transport and biogeochemical budgets through intense upwelling of nutrients, subduction of plankton and horizontal stirring (see review of Lévy, 2008).

The LTL model was able to reproduce the seasonal signal and the inter-annual variability of phytoplankton surface concentrations. The model also captured the daily variability. However, the modeled concentrations were, most of the time, underestimated compared to the satellite-derived...
means concentrations, especially during late spring and summer. It is crucial to keep in mind that ocean color data in the Mediterranean Sea have to be considered with caution, because the algorithms used to derive the chlorophyll $a$ concentrations perform poorly in this regional sea (Gregg and Casey, 2004; Volpe et al., 2007 and references therein). These low performances are generally attributed to the particular composition (i.e. inorganic and organic matter) of the water column causing an overestimation of chlorophyll $a$ surface concentrations for low values especially of chlorophyll (i.e. during summer and in oligotrophic waters). This bias between in situ data and values derived from algorithms would be furthermore accentuated in the area of the Rhone freshwater influence according to the study of Ouillon and Petrenko, (2005). The C parameter, assessing error between model output and data (see section 2.4.2.), was null during ca. 23% of the comparison period. This result means that the modeled chlorophyll value remained within the range of the error bars given by the GlobColour algorithm during a quarter of the simulation period. Furthermore, figure 6 shows the temporal changes in the GlobColour errors on chlorophyll $a$ concentrations and of the C parameter. The GlobColour errors and the C parameter were the lowest ($<1.25 \times 10^{-1} \text{mg m}^{-3}$) during summer, especially in 2003 and 2004, when they were the most variable and highest from autumn to mid-spring. The C parameter was lower than the satellite sensor errors during ca. 71% of the comparison period, which is a rather satisfactory result.

3.3. Comparing modeled and observed diet of the HTL species

Figure 7 shows the modelled diet of every size class of the ten HTL species. In order to describe the diets emerging from the OSMOSE-GoL model, prey groups were classified into three categories according to their proportion (in biomass) in the predator's diet: main prey (>50%), secondary prey (from 10 to 50%), accessory prey (from 1 to 10%).

According to the model, the main prey of the Northern krill is microphytoplankton (~56%), while the secondary prey is microzooplankton (~35%). Mesozooplankton remains an accessory prey (~8%). This modelled diet, composed of phyto- and zooplankton in almost equal parts, is consistent
with the observations of Båmstedt and Karlson (1998) on krill of the Northeast Atlantic. According
to this field study, northern krill may not survive on a diet only based on phytoplankton and needs
to consume at least 40% of zooplankton.

The modelled southern shortfin squid mainly feeds on small teleosts (~77%) such as planktivorous
European pilchard (~36%) and European anchovy (~21%), but also on juveniles of Atlantic horse
mackerel (~9%) and blue whiting (~8%). Small squid (i.e. cannibalism) (~11%) are secondary prey,
while northern krill (~8%) and mesozooplankton (~4%) are accessory prey of southern shortfin
squid. This modelled diet is consistent with some in situ datasets (Bănaru, 2015) showing a diet
mainly composed by ~80% of teleosts. In the latter field study, southern shortfin squid prey on
certain other groups, such as benthic decapods, but these groups were not represented in the
OSMOSE-GoL model.

According to the model outputs, European pilchard larvae (<3 cm) mainly feed on
mesozooplankton (~81%). This trophic behaviour for larvae are close to the results of Borme et al.
(2009) in the Mediterranean Sea, showing a diet almost exclusively (~99%) composed of copepods
in the size range of mesozooplankton. Larger individuals significantly change their diet, feeding on
smaller prey, with microzooplankton (~51% of the diet of juveniles ranging between 3 and 12.5 cm,
and ~52% for adults >12.5 cm) as main prey, and microphytoplankton as secondary prey (~44% of
the diet of juveniles and ~43% for adults). The latter results are not in line with some recent dietary
field studies based on the analysis of stomach content in the Gulf of Lions. According to the study
of Le Bourg et al. (2015), juveniles of European pilchard as well as adults may prey quasi-
exhaustively on mesozooplankton (~98% and 100%, respectively). However this mismatch between
model outputs and observations has to be moderated, because the study of Bode et al. (2004)
showed that the size of the prey generally decreases with increasing body length of European
pilchard. Hence, adults may consume a larger part of phytoplankton according to the latter study. As
phytoplankton prey are actually more easily digested, they are therefore more difficult to identify in
stomach contents, which may explain the differences between the model and observations for this
fish species. Moreover, the study of Pethybridge et al. (2014) using analysis of fatty acids in adults of European pilchard confirms the assumption that its diet is mainly based on microplankton. The modelled European anchovy, including its larvae stage, has a diet similar to those of the juveniles and adults of European pilchard. All stages mainly feed on microzooplankton (~55% of the diet for larvae <3 cm, ~49% for juveniles ranging between 3 and 12.5 cm and ~44% for adults >12.5 cm). Their secondary prey is the microphytoplankton group (~39% for larvae, ~40% for juveniles and ~36% for adults). Mesozooplankton is only an accessory prey, representing about 5% of the diet for larvae and juveniles, and around 7% for adults. The simulated diet of the larvae is close to observed data. Analyzing fatty acids content, Rossi et al. (2006) showed that microzooplankton and microphytoplankton are the prey most consumed by anchovy larvae. However, the modelled diet for adult anchovy is rather different from those resulting from the analysis of stomach content (Borme et al., 2009; Le Bourg et al., 2015). In the observations, as for the European pilchard, the proportions of microzooplankton and microphytoplankton may be underestimated due to a rapid digestion of these types of prey.

The modelled diet of European sprat juveniles is similar to those of juveniles and adults of the European pilchard and of all size classes of the European anchovy. The diet of the juveniles is composed of microzooplankton (~48% of the diet) and microphytoplankton (~34%). The proportion of consumed mesozooplankton (~14%) is higher than in the diet of the juveniles of the European sardine and anchovy, but is much lower than that observed in their stomach content (~100%), according to the study of Le Bourg et al. (2015). The modelled diet of European sprat larvae (<3 cm) is mainly composed of mesozooplankton (~81%), which is consistent with previous observations (Dickmann, 2005). Mesozooplankton is furthermore the main prey for adults but in much higher proportions (~68%) than for the adults of European pilchard and anchovy (~4.5% and ~7%, respectively). Adults also consume, as secondary prey, some European pilchard larvae (~16%). In the model, the European sprat (including larvae and adults) thus appears to be more carnivorous than the European sardine and anchovy. This modelled output is in line with the dataset.
of Pethybridge et al. (2014) based on the comparative analysis of the fatty acids content of these three pelagic species. In the model, pilchard eggs and larvae are another significant prey (~16%) for the adults of European sprat, as also observed in their stomach content (Le Bourg et al., 2015).

The diet of the juveniles of Atlantic horse mackerel predicted by the model is composed of a dominant proportion of mesozooplankton (~62%) and a lesser proportion of fish larvae (~25%), while the study of Le Luherne (2012) based on the analysis of stomach content showed a dominant consumption of northern krill (~71% in proportion) and a lesser proportion (~21%) of benthic decapods (not modelled). These differences may result from opportunistic predation behaviour (making the diets highly variable between the sampling stations inside the GoL domain), as well as from some differences between the spatial domain of the model (entire GoL) and the specific locations of the sampling stations used in the study of Le Luherne (2012). Adults of Atlantic horse mackerel (>16 cm) have a modelled diet which is quite different from that of juveniles. They mainly feed on southern shortfin squid (~37%) and northern krill (~28%). Some other prey such as juveniles of European sprat and hake and Atlantic mackerel supplement their diet (~17%). The cumulated percentage of consumed fish is rather high (34%). The modelled proportion of northern krill consumed is close to that resulting from the analysis of in situ stomach content (~21%; Le Luherne, 2012). By contrast, the latter empirical study did not show the presence of southern shortfin squid in the stomach of analysed individuals. Differences between model outputs and observations may again result from differences in diet when estimated from the entire modelled domain rather than from some particular locations as in the study of Le Luherne (2012). Differences in the diet of predators may also result from inter-annual variations in the spatial distribution and density of prey. Furthermore, a major bias in the modelling of the diet of Atlantic horse mackerel adults is that a dominant proportion of consumed prey (~54% of macrozooplankton in the form of pteropods and ~21% of decapods, Le Luherne, 2012) is not represented as potential prey in the model.

In the model, the juveniles of Atlantic mackerel (<20 cm) mainly consume certain planktivorous
fish, a result which is consistent with observations (Le Luherne, 2012). However, the modelled diet appears to be dominated by European pilchard (~38%) and European anchovy (~22%), while the analysis of stomach content data rather shows a high consumption of sprats (~37%) and to a lesser extent, anchovies (~16%). Accessory prey are diverse (i.e. mesozooplankton, northern krill, southern shortfin squid, Atlantic horse mackerel, blue whiting, other prey), each of them accounting for 6 to 7% of the modelled diet. The differences in the consumed fish species between the model and observations may be attributable to differences in species composition between the modelled period (2001-2004) and the field studies that were carried out in 2011-2012 (Le Luherne, 2012). Since 2008, the biomass of European pilchard has strongly declined in the Gulf of Lions, while that of sprat has increased (Van Beveren et al., 2014). Macrozooplankton and groups of benthic species that are not represented in the model account for significant proportions of the stomach content (~27%, ~19% respectively, Le Luherne, 2012). The model succeeded in predicting the functional group consumed (i.e. planktivorous fish) for the juveniles of Atlantic mackerel. Adults of Atlantic mackerel (>20 cm) are even more piscivorous than juveniles, since their modelled diet is almost 90% composed of fish. The most consumed fish are firstly European pilchard (~46%) and secondarily, European anchovy (~26%), in the model. Comparatively, the proportion of consumed teleosts observed in stomach content amounts to almost 90% in the study of Le Luherne (2012), but the teleost species could not be identified due to too advanced digestion of prey. Juveniles of blue whiting (<15 cm) have a rather diversified diet in the model. They feed on certain plankton groups (mesozooplankton at ~19%, and northern krill at ~25%), on southern shortfin squid (~10%) and on different species of fish (~46%). The most consumed fish species are European pilchard (~27%), Atlantic horse mackerel (~10%) and blue whiting (~6%). This result is different from those obtained in the study of Bourgogne (2013) who observed only decapods (not represented in the model) in the stomach content of analysed juveniles. In the model, adults of blue whiting (>15 cm) mainly feed on fish (~65%) secondarily on northern krill (~26%) and southern shortfin squid (~10%). The most preyed species of fish are European pilchard (~32%), Atlantic horse
mackerel (~11%), blue whiting (~12%) and European anchovy (~8%). The study of Bourgogne (2013) showed a proportion of northern krill (~33%) close to that provided by the model. The main difference between model and observations is in the dominant prey. The proportion of teleosts consumed amounts to ~65% in the model, while stomach content data show a lower proportion (~35%, Bourgogne, 2013). In parallel, benthic decapods (not modelled) have been found in a significant proportion (~32%) in the stomach content of analysed adults, which may explain the mismatch between the modelled diet of adults and field data.

In the model, the main prey of the European hake juveniles are teleosts (~90%). Among these teleosts, the most consumed species are firstly the European pilchard (~42%), secondly the European anchovy (~24%) and thirdly, the blue whiting (~13%). In parallel, a recent study of stomach content from the same area (Mellon-Duval et al., 2017) revealed in situ diet mainly composed of teleosts (>80% and to 100%). In more detail, the European pilchard was observed in the highest proportion (22 to 74%), but the European anchovy and the blue whiting were also detected in lesser proportions (6 to 30% and 1 to 3%, respectively). The model results also match the observations of two other recent studies, estimating at ~92% the proportion of teleosts in the diet of juveniles (Merquiol, 2016; Bănaru and Harmelin-Vivien, 2017). In the model, adults, as juveniles, feed mostly on teleosts (~93%). The teleost prey are firstly European pilchard in a proportion of ~43%, and secondly, blue whiting at ~13%. Other fish species such as European sprat, Atlantic mackerel and European hake supplement their diet up to 16%. Mellon-Duval et al. (2017) found from the analysis of stomach content a roughly similar proportion of European pilchard (~38%) and slightly higher proportion of blue whiting (~26 to ~30%).

The modelled diet of Atlantic Bluefin tuna is almost exclusively piscivorous (~93%) and composed of small teleosts. European pilchard is the main prey (~51%), while European anchovy (~23%) and blue whiting (~10%) are the secondary prey. Southern shortfin squid is only an accessory prey (~7%). This preference for planktivorous teleosts, mainly European anchovy, has already been shown from the analysis of the stomach content of individuals caught in the Gulf of Lions (Imbert
et al., 2007). Once again, the minor mismatches observed between the modelled diet and that from
the field data may result from a dataset based on sampling carried out outside the temporal window
of the simulation period. This fast-swimming pelagic tuna species has been shown to have a high
variability in its distribution at different spatial and temporal scales in the GoL (Royer et al., 2004).

3.4. Comparing TL in the E2E modelled food web with literature data

The E2E modelled food web (Figure 8) is composed of 15 compartments organised into four
trophic levels (TL). This food web length is rather common in natural ecosystems (Hastings and
Conrad, 1979). Five plankton groups represent the lowest TL ([1-2]), while ten invertebrates and
teleost groups represent the highest TL (>3). The highest flows of predation occur between the
phyto- and zooplankton groups (>10^6 tons y^-1), and sharply decrease with increasing TL, which is a
common feature observed in trophic pyramids (Odum, 1959). Primary producers such as nano- and
microphytoplankton representing the first TL are mainly consumed by nano- (TL=2.0), micro-
(TL=2.0) and mesozooplankton (TL=2.2). Among these groups, mesozooplankton has the highest
TL, as it consumes both phyto- and zooplankton. Planktivorous species such as northern krill
(TL=2.5), European pilchard (TL=2.7), European anchovy (TL=2.8) and European sprat (TL=2.9)
mainly feed on micro- and mesozooplankton. The European pilchard has a more omnivorous diet,
with a consumption of microphytoplankton of the same order of magnitude as that of
microzooplankton, or even higher than that of mesozooplankton. For these planktivorous species,
TL increases from the northern krill to the European sprat, in agreement with literature data
(Båmstedt and Karlson, 1998; Stergiou and Karpouzi, 2002; Le Bourg et al., 2015).

Among the meso-predators, the Atlantic horse mackerel has the lowest TL (TL=3.5), followed by
those of Atlantic mackerel (TL=3.9), southern shortfin squid (TL=4.0), blue whiting (TL=4.0),
Atlantic bluefin tuna (TL=4.0) and European hake (TL=4.1). These TLs are close to field data
(Stergiou and Karpouzi, 2002; Sara and Sara, 2007; Bănaru, 2015; Bănaru and Harmelin-Vivien,
2017; Mellon-Duval et al., 2017). Some larvae of these teleosts are consumed by cannibalism and
also by other species of similar, lower and higher TL.

On the whole, the modelled TLs are in agreement with literature data and ECOPATH-GoL outputs for most of the 10 HTL species (Figure 9). Some interesting points can however be highlighted. The modelled TLs of southern shortfin squid are slightly higher than those in the literature (Bănaru et Harmelin-Vivien, 2017) and from ECOPATH outputs (Bănaru et al., 2013). The predator-prey size ratios of the present model may be better adjusted in order to obtain a modelled TL for southern shortfin squid closer to field and literature data.

The modelled TLs of juveniles and adults of European pilchard are close to the minimum values estimated from the analysis of the stomach content (Stergiou and Karpouzi, 2002) or using δ^{15}N stable isotope ratios (Bănaru, 2015), while the corresponding larvae show their numerical TLs close to the observed highest values. In this case too, the predator-prey size ratios of the model may be better adjusted.

The TLs of all species (except pilchard, and to a lesser extent blue whiting) increase with increasing size of individuals. In order to better account for ontogenetic changes in TL (Chassot et al. 2008; Reed et al. 2017), it would be useful in the future to better refine field estimation of trophic levels by size.

It is interesting and encouraging to note that the TLs computed from two very different approaches of E2E modelling (i.e. OSMOSE vs. ECOPATH) are very close for most of HTL species, except for European pilchard and blue whiting. Even if the represented periods in these models partly overlap (2001-2004 for OSMOSE and 2000-2009 for ECOPATH) and the modelled area is the same, major differences in conception between these two models (opportunistic, size-based diet in OSMOSE and fixed diet in ECOPATH) may explain these small differences in trophic level results. OSMOSE is an individual size-based, spatial and dynamic model, while ECOPATH is a mass-balanced model, with no size groups, non-spatialized and offering a static snapshot of the system. Moreover, ECOPATH includes more groups and species (benthic groups, marine mammals, birds and others), which are not represented in OSMOSE. Data gaps concerning the diet of some species were
highlighted by Bânaru et al. (2013), and new recent diet data were included in the OSMOSE model parametrization.

4. Conclusion

The GoL is a highly exploited area for fisheries (Demaneche et al., 2009; Bânaru et al., 2013). Here, as in other parts of the world, fishing has reduced the biomass of top predators (Aldebert, 1997; Piroddi et al., 2017), with potential cascading effects on the flows and biomass in the food web (Pauly et al., 1998; Cury et al., 2003; Bânaru et al., 2010; Ferretti et al., 2010). Climate variations impact the GoL ecosystems through river inputs (Ludwig et al., 2009) or hydrological processes (Hermann et al., 2008). As shown in other areas, the impact of fishing combined with climate changes induces sometimes unexpected effects in the ecosystems (Travers-Trolet et al., 2014; Auber et al., 2015). The end-to-end modelling approach aims at understanding and anticipating some of these processes.

This paper presents the first spatialized dynamic coupled end-to-end ecosystem model for the GoL. The modeled groups and species represent more than 70% of annual catches in this area, and they encompass the pelagic and demersal ecosystem trophic structure organised into four trophic levels. The assessment of both LTL and HTL groups and species of the E2E OSMOSE-GoL model showed a satisfactory agreement with literature, satellite and field local data in terms of biomass, landings and diet. The model has been parametrized with the best available local data. Following the calibration stage, realistic ranges for the biomass and landings have been obtained for most species and groups. Biomass for the European hake and landings for the Atlantic mackerel are however slightly overestimated. At first sight, it might be considered that these differences are acceptable owing to uncertainties existing in biomass estimations and landings data. However, some improvements of the model are obviously still possible. Predator-prey size ratios may be refined to produce a better correspondence between model and field data. Sex-ratio established by default at 0.5 may be refined with field data when available. The distribution maps currently based on
presence-absence may be replaced with density-based maps. Fishing mortality considered uniform for the entire domain may be also spatialized.

Previous versions of the OSMOSE model have already been applied in different ecosystems to address various questions regarding: i/ the assessment of the ecosystem trophic structure (Marzloff et al., 2009; Grüss et al., 2015; Houaoui et al., 2016, Fu et al., 2017), ii/ the effect of Marine Protected Areas (Yemane et al., 2009), iii/ the combined effects of fishing and climate change (Travers et al., 2009; Fu et al., 2013; Travers-Trolet et al., 2014; Fu et al., 2018) iv/ the simulation of fishing scenarios (Shin et al., 2004; Travers et al., 2010; Smith et al., 2011; Grüss et al., 2016), and v/ the testing of indicator performance (Travers et al., 2006; Shin et al., 2018). However, the present version of E2E OSMOSE for the GoL may go further, in the analysis of the fine impact of the predation pressure exerted by HTL planktivorous species on the spatial distributions and the structure (size, trophic shortcut, etc.) of the plankton community. It may also contribute to the understanding of complex processes of simultaneous bottom-up and top-down controls in this exploited ecosystem (Diaz et al., 2019). Moreover, this type of E2E model may allow for the quantitative assessment of the combined effects of fishing and climate change scenarios on the ecosystem dynamics and for the computation of model-based indicators used to assess whether an ecosystem and its services are used sustainably, and then maintained (Coll et al., 2015). The impact of existing or future Marine Protected Areas (Gulf of Lions, Fisheries Restricted Area, etc.) and spatio-temporal management measures on the structure and the functioning of the ecosystem may also be tested in future.

This coupled E2E model may be extended to the entire Mediterranean Sea and compared with existing Ecopath with Ecosim ecosystem model configurations (Coll and Libralato, 2012; Piroddi et al., 2017).

Acknowledgements
This study was funded by the project EMIBIOS (End-to-end Modelling and Indicators for BIODiversity Scenarios, FRB contract no. APP-SCEN-2010-II) and by the EU FP7 project PERSEUS (Policy-oriented marine Environmental Research for the Southern European Seas, Theme “Oceans of Tomorrow” OCEAN.2011-3 Grant Agreement No. 287600). It benefited from and contributed to MERMEX WP2 collaborative work and to the MERMEX IPP “Interactions plancton-planctonophages” project. We wish to thank J.M. Fromentin, J.L. Bigot, C. Saraux, A. Jadaud for providing some data and expertise. The authors acknowledge T. Ballerini for her contribution to the coupling code and suggestions that improved the parametrization of the OSMOSE-GoL model, the assistance of staff maintaining the clusters DATARMOR of the IFREMER and HPC Platform of the OSU Institut PYTHEAS (Aix-Marseille University, INSU-CNRS) for providing the computing facilities, as well as for technical assistance. Thanks are also addressed to Michael Paul for English corrections.

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**LEGENDS OF TABLES**

**Table 1** Characteristics and parameters of the five LTL groups of the Eco3M-S model.

**Table 2** Parameter values of the OSMOSE-GoL model obtained from fitting the model to observed data. \( M_0 \): survival rate; \( M_p \): mortality rate due to predation from other species that are not explicitly considered in the model; \( F \): annual fishing mortality rate; \( a_p \): availability coefficients of plankton groups to HTL species.

**Table 3** Size classes of the different species implemented in the OSMOSE-GoL model, minimum and maximum predator/prey size ratios. MIR= Maximum ingestion rate.

**Table 4** Input parameters of the OSMOSE-GoL model for each of the 10 HTL species modelled. \( K \) (growth rate), \( L_\infty \) (asymptotic size) and \( t_0 \) (time at null size): the von Bertalanffy growth parameters; \( b \): the exponent of the allometric length–weight relationship; \( c \): const. of proportionality of the allometric length-weight relationship; \( s_{mat} \): size at maturity; \( \phi \): relative fecundity; \( S \): egg size; \( a_{max} \): longevity; \( s_{rec} \): legal size of recruitment for fisheries catch.

**Table 5** Seasonality of the fishing activity in the GoL for target species.
<table>
<thead>
<tr>
<th>Size range</th>
<th>Eco3M-S mortality rate, $m_p$ (d$^{-1}$)</th>
<th>Trophic level</th>
<th>Conversion factor (mg$_{ww}$ mmolN$^{-1}$)</th>
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</thead>
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<tr>
<td>NANOPHY</td>
<td>0.000$^b$</td>
<td>1.0$^c$</td>
<td>993.75$^{d,e,f}$</td>
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<tr>
<td>MICROPHY</td>
<td>0.075$^b$</td>
<td>1.0$^c$</td>
<td>993.75$^{d,e,f}$</td>
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<tr>
<td>NANOZOO</td>
<td>0.043$^b$</td>
<td>1.5$^c$</td>
<td>832.50$^{p,h}$</td>
</tr>
<tr>
<td>MICROZOO</td>
<td>0.070$^b$</td>
<td>2.0$^c$</td>
<td>832.50$^{p,h}$</td>
</tr>
<tr>
<td>MESOZOO</td>
<td>0.033$^{a,b}$</td>
<td>2.5$^c$</td>
<td>150.00$^{**}$</td>
</tr>
</tbody>
</table>

*Units: m$^3$ mmolC$^{-1}$ d$^{-1}$ (predation rate), **Units: mg$_{ww}$ mmolC$^{-1}$, $^a$Sieburth et al. (1978), $^b$Campbell et al. (2013), $^c$Arbitrarily set, $^d$Dalsgaard and Pauly (1997), $^e$Walsh (1981), $^f$Bundy (2004), $^g$Gifford and Caron (2000), $^h$Hardsted-Roméo (1982).
### Table 2

<table>
<thead>
<tr>
<th>Species</th>
<th>Mortality</th>
<th>Availability coefficients</th>
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<tr>
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<td>$M_0$</td>
<td>$M_s$</td>
</tr>
<tr>
<td></td>
<td>($y^{-1}$)</td>
<td>($y^{-1}$)</td>
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<td>5.558</td>
<td>0.365</td>
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<td>European anchovy</td>
<td>6.609</td>
<td>0.228</td>
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<tr>
<td>European sprat</td>
<td>4.549</td>
<td>0.404</td>
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<tr>
<td>Atlantic horse mackerel</td>
<td>1.271</td>
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<tr>
<td>Atlantic mackerel</td>
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<td>0.000</td>
</tr>
<tr>
<td>NANOPHY</td>
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<td>-</td>
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<tr>
<td>MICROPHY</td>
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<tr>
<td>NANOZOO</td>
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<td>MESOZOO</td>
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## Table 3

<table>
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<th>Species</th>
<th>Size classes (cm)</th>
<th>Predator/prey size ratios</th>
<th>MIR</th>
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<td>188</td>
<td>4</td>
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<td>17</td>
<td>1</td>
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<tr>
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<td>65</td>
<td>13</td>
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<td></td>
<td>3-12.5</td>
<td>1139</td>
<td>15</td>
</tr>
<tr>
<td></td>
<td>&gt;12.5</td>
<td>1621</td>
<td>60</td>
</tr>
<tr>
<td>European anchovy</td>
<td>&lt;3</td>
<td>500</td>
<td>4</td>
</tr>
<tr>
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<td>3-8</td>
<td>500</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>&gt;8</td>
<td>806</td>
<td>7</td>
</tr>
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<td>European sprat</td>
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<td>2</td>
</tr>
<tr>
<td></td>
<td>3-11.6</td>
<td>517</td>
<td>10</td>
</tr>
<tr>
<td></td>
<td>&gt;11.6</td>
<td>517</td>
<td>10</td>
</tr>
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<td>&lt;16</td>
<td>40</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>&gt;16</td>
<td>100</td>
<td>4</td>
</tr>
<tr>
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</tr>
<tr>
<td></td>
<td>&gt;20</td>
<td>61</td>
<td>2</td>
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<td>3</td>
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<td></td>
<td>&gt;15</td>
<td>32</td>
<td>4</td>
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<td>1</td>
</tr>
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<td></td>
<td>&gt;36</td>
<td>14</td>
<td>1</td>
</tr>
<tr>
<td>Atlantic bluefin tuna</td>
<td>-</td>
<td>20</td>
<td>4</td>
</tr>
<tr>
<td>Species</td>
<td>K   (y(^{-1}))</td>
<td>L(_\infty) (cm)</td>
<td>t(_0) (y)</td>
</tr>
<tr>
<td>------------------------------</td>
<td>------------------</td>
<td>-------------------</td>
<td>-----------</td>
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<tr>
<td>Northern krill</td>
<td>1.680</td>
<td>3.462</td>
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<tr>
<td>Southern shortfin squid</td>
<td>0.930</td>
<td>17.400</td>
<td>0.087</td>
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<td>European pilchard</td>
<td>0.334</td>
<td>19.925</td>
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<td>European anchovy</td>
<td>0.609</td>
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<td>0.093</td>
<td>318.85</td>
<td>-0.97</td>
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**Table 4**
<table>
<thead>
<tr>
<th>Fishing mortality / time steps</th>
<th>Southern shortfin squid</th>
<th>European pilchard</th>
<th>European anchovy</th>
<th>Atlantic horse mackerel</th>
<th>Atlantic mackerel</th>
<th>Blue whiting</th>
<th>European hake</th>
<th>Atlantic bluefin tuna</th>
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<td>5, 6</td>
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<td>0.053</td>
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<td>0.003</td>
<td>0.005</td>
<td>0.000</td>
</tr>
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</table>
LEGENDS OF FIGURES

Fig. 1. Processes taken into account within each of the two models and processes linking the two models (dashed-line arrows). The time step of Eco3M-S/SYMPHONIE (left hand side, Campbell et al., 2013) is one hour while that of OSMOSE (right hand side, adapted from Travers-Trolet et al., 2014) is 15 days. The two-ways coupling mode is used throughout the predation process, where the biomass of the plankton groups serve as prey field for fish schools, cephalopods and krill (“Prey availability” arrow), while an explicit rate of HTL-induced predation is specifically applied as feedback on each of the aforementioned five plankton groups (“Predation mortality” arrow). In the one-way forcing mode, plankton biomasses serve as prey fields to fish schools, cephalopods and krill (“Prey availability” arrow), without any feedback on the plankton prey compartments.

Fig. 2. The Eco3M-S/Symphonie model domain in the NW Mediterranean Sea is delineated by the black thin line. The OSMOSE-GoL model domain over the Gulf of Lions is delimited by the black bold line. The map of grid points in the OSMOSE-GoL domain is given in the small panel in the right edge of the figure. The black arrow indicates the main flow of the Northern Current (NC). Rivers taken into account by the model are named. Bathymetry with isobaths 50, 100 and 1000 m is shown in the modeled area.

Fig. 3. Temporal patterns of change in the simulated biomass of the 10 HTL species during the last four years of spin-up over the whole modeled domain. The solid blue line shows the median value computed from the 50 simulation replicates. The lower and upper limits of the grey range delineate the 0.25 and 0.75 (resp.) percentiles computed from the 50 replicates. The two horizontal dotted black lines represent the range of observed biomass (see references 2.3).

Fig. 4. Temporal patterns of change in the simulated landings of the 10 HTL species during the last
four years of spin-up over the whole modeled domain. The solid blue line shows the median value computed from the 50 simulation replicates. The lower and upper limits of the grey range delineate the 0.25 and 0.75 percentiles computed from the 50 simulation replicates. The horizontal black lines represent monthly observed landings (see references 2.3).

Fig. 5. Time-series of satellite-derived (blue) and modeled (red) chlorophyll surface concentrations (mg m\(^{-3}\)) over the whole modeled domain. Thick lines represent mean values and shaded area (grey) is the range of error bars computed from the GlobColour algorithm. Missing data in the time-series are explained by cloudy days without any useable pixels.

Fig. 6. Time-series of the averaged error (mg m\(^{-3}\)) on satellite-derived chlorophyll (blue dots) and of the C parameter (metric of data-to-model distance in mg m\(^{-3}\), red dots) on the whole modeled domain.

Fig. 7. Mean diet (over the years 36 to 43) of larvae, juveniles and adults of the 10 HTL species considered in the OSMOSE-GoL model.

Fig. 8. Representation of the trophic levels and the main fluxes of matter between the compartments of the OSMOSE-GoL model (TL = trophic level, LTL = low trophic levels, HTL = high trophic levels). Size of the arrows is related to the intensity of predation fluxes. Fluxes lower than 10 tons y\(^{-1}\) are not quoted in the diagram. For sake of clarity, only fluxes that account for at least 10% of predation total flux for a given prey are shown.

Fig. 9. Trophic levels vs. total length of the ten HTL species represented in the OSMOSE-GoL model (there was a high inter-annual overlap of the trophic levels). Horizontal dashed black lines indicate the minimum and maximum values from the literature data based on the analysis of
stomach content and corresponding to a range of total length. Horizontal solid black lines indicate
the trophic levels from the ECOPATH model of the GoL (Bănaru et al., 2013).
Fig. 2.
Fig. 3.
Fig. 4. 

A. Northern krill

B. Southern shortfin squid

C. European pilchard

D. European anchovy

E. European sprat

F. Atlantic horse mackerel

G. Atlantic mackerel

H. Blue whiting

I. European hake

J. Atlantic bluefin tuna
Fig. 5.

Satellite-derived
Modeled
Mean values
Error bars

Chl (mg m⁻³)

10⁸
10⁶
10⁴
10²

Fig. 6.

Chl satellite-derived C parameter (data-to-model distance)
Micro phytoplankton
Micro zooplankton
Meso zooplankton
Northern krill
Southern shortfin squid

Prey groups:

- European pilchard < 3 cm
- European pilchard 3-12.5 cm
- European pilchard > 12.5 cm
- European anchovy < 3 cm
- European anchovy 3-12.5 cm
- European anchovy > 12.5 cm
- European sprat < 3 cm
- European sprat 3-11.6 cm
- European sprat > 11.6 cm
- Atlantic horse mackerel < 16 cm
- Atlantic horse mackerel > 16 cm
- Atlantic mackerel < 20 cm
- Atlantic mackerel > 20 cm
- Blue whiting < 15 cm
- Blue whiting > 15 cm
- European hake < 36 cm
- European hake > 36 cm
- Atlantic bluefin tuna > 36 cm

Other prey
Fig. 8.

TROPHIC LEVELS (TL)

4

- SOUTHERN SHORTFIN SQUID
- BLUE WHITING

3

- ATLANTIC MACKEREL
- ATLANTIC HORSE MACKEREL

NORTHERN KRILL

- EUROPEAN PILCHARD
- EUROPEAN ANCHOVY

- ATLANTIC MACKEREL
- BLUE WHITING

- EUROPEAN SPRAT
- ATLANTIC MACKEREL

2

- NANOZOOPLANKTON
- MICROZOOPLANKTON

1

- NANOPHYTOPLANKTON
- MICROPHYTOPLANKTON

\[ >10^8; \uparrow 10^7 - 10^8; \uparrow 10^6 - 10^7; \uparrow 10^4 - 10^5; \uparrow 10^3 - 10^4; \uparrow 10^2 - 10^3; \uparrow 10^{-2} \text{ tons y}^{-1} \]
Fig. 9.