

Bioaccumulation modelling and sensitivity analysis for discovering key players in contaminated food webs: the case study of PCBs in the Adriatic Sea

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Abstract

Modelling bioaccumulation processes at the food web level is the main step to analyse the effects of pollutants at the global ecosystem level. A crucial question is understanding which species play a key role in the trophic transfer of contaminants to disclose the contribution of feeding linkages and the importance of trophic dependencies in bioaccumulation dynamics. In this work we present a computational framework to model the bioaccumulation of organic chemicals in aquatic food webs, and to discover key species in polluted ecosystems. As a result, we reconstruct the first PCBs bioaccumulation model of the Adriatic food web, estimated after an extensive review of published concentration data. We define a novel index aimed to identify the key species in contaminated networks, *Sensitivity Centrality*, and based on sensitivity analysis. The index is computed from a dynamic ODE model parametrised from the estimated PCBs bioaccumulation model and compared with a set of established trophic indices of centrality. Results evidence the occurrence of PCBs biomagnification in the Adriatic food web, and highlight the dependence of bioaccumulation on trophic dynamics and external factors like fishing activity. We demonstrate the effectiveness of the introduced *Sensitivity Centrality* in identifying the set of species with the highest impact on the total contaminant flows and on the efficiency of contaminant transport within the food web.

Keywords: bioaccumulation modelling, ecological network analysis, Sensitivity Analysis, toxic keystone, Adriatic Sea, Polychlorinated Biphenyls, Linear Inverse Modelling

1. Introduction

A food web represents a comprehensive model to interpret the pattern of trophic connectedness in an ecosystem, where the biomass and energy flows are bound to a complex mixture of organic compounds. In the aquatic environment, persistent organic chemicals accumulate in the lipid tissue of organisms from dietary uptake and from exposure through water (Van der Oost et al., 2003). Bioaccumulation phenomena occur when the concentration of a toxicant in marine biota is higher than in the surrounding environmental media (Mackay and Fraser, 2000). A variety of biological and chemical factors concerning both marine organisms and chemical compounds, combined with species feeding behaviour, can differently influence the patterns of contamination (Russell et al., 1999). Feeding relationships not only expose species to contamination processes but also become a critical medium of toxicant transfer, leading to biomagnification phenomena as the result of dietary uptake (Kelly et al., 2007, Lohmann et al., 2007).

Therefore, predator-prey interactions become central to characterize contamination patterns scaling from individuals to the ecosystem level, and to predict bioaccumulation effects on ecological networks (Rohr et al., 2006). Food web members exhibit

different ecological responses to accumulated concentrations of chemicals in function of their abundance, trophic position, feeding connections, and role in maintaining ecosystem functions (Ruus et al., 2002, Walters et al., 2008). As a consequence, the contribution of trophic connections in the contamination pathways cannot be considered equal for all species. Despite the trophic role of keystone species is well-established in ecology, the crucial question of which species play a central role in the trophic transfer of contaminants remains poorly understood.

In this work we present a computational framework for bioaccumulation modelling and for analysing the trophic role of species in contaminated food webs. We focus on food-web bioaccumulation models, a class of ecological networks that represent and quantify the contaminant transfer between species by following the underlying feeding links. The bioaccumulation network is estimated from biomass and contaminant concentration data by using Linear Inverse Modelling (LIM) (Vézina and Platt, 1988, van Oevelen et al., 2010), a technique that supports incomplete and uncertain input data. In order to complete our framework, we employ ecological network analysis tools to identify the *toxic keystones*. Specifically, we apply indices of ecological centrality, typically used for trophic conservation purposes (Jordán, 2009), to provide indicators of species' importance in the bioaccumulation context. To this aim, we also introduce *Sensitivity Centrality (SC)*, a novel in-

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dex based on the sensitivity analysis of dynamic models, and suitable to express information on the temporal patterns of contamination. In our case, SC is computed from a multi-species Lotka-Volterra ODE model derived from the estimated bioaccumulation network.

We apply the framework to the case study of polychlorinated biphenyls (PCBs) bioaccumulation in the Adriatic food web. In the last decades this region has become of great interest from an ecotoxicological viewpoint, since it has been subject to dramatic changes in marine resources driven by anthropogenic and environmental perturbations. PCBs are industrial chemicals liable to contamination problems in the aquatic environments, and among other regions, they have been detected both in abiotic compartments and living organisms in the Adriatic sea (Picer, 2000). This class of persistent organic pollutants consists of 209 different congeners and are chemically characterised by high environmental persistence, being practically insoluble in water. Moreover for their lipophilic properties, PCBs readily dissolve in fats and lipids of aquatic organisms leading to bioaccumulation phenomena in species.

In our study, input biomass data is compiled from a complete and validated trophic model of the Adriatic food web (Coll et al., 2007) and we collect PCBs concentration data after an extensive review of specific experimental literature. As a result, we obtain the *first food-web bioaccumulation model of PCBs in the Adriatic ecosystem*, where missing concentration data is estimated by combining LIM with stochastic sampling techniques. Finally, we analyse toxic keystones and evaluate the newly introduced Sensitivity Centrality index on the obtained bioaccumulation network.

LIM-based methods have been extensively applied to the reconstruction of food webs from empirical data. *Ecopath* (Christensen and Walters, 2004) is one of the most established and used LIM tools, and includes the *Ecotracer* routine for bioaccumulation analysis. In this work we choose the R package *LIM* (van Oevelen et al., 2010) because it supports custom equations and multiple flow currencies (both biomass and PCBs), a crucial feature of our framework. Other LIM approaches for bioaccumulation modelling include *Toxlim* (Laender et al., 2009), a R-package implementing the OMEGA model (Hendriks et al., 2001).

A LIM model of the Venice lagoon food web is developed by Brigolin et al. (2011) to study the temporal evolution of ecosystem productivity and fishing. The estimation is based on randomly perturbing input data to avoid the bias of constraints on the obtained solutions. Our LIM trophic model is compiled taking input data from the work by Coll et al. (2007), where the Northern and Central Adriatic food web is reconstructed using *Ecopath*. In the estimation of the bioaccumulation network, we consider a different approach in order to account for uncertainty, based on the Markov Chain Monte Carlo (MCMC) sampling of the solution space.

Models for assessing and predicting PCBs bioaccumulation have been proposed for two main Adriatic areas: the Venice lagoon (Losso and Ghirardini, 2010, Dalla Valle et al., 2005, 2007) and the Po river delta (Spillman et al., 2007, 2008). These works investigate the contaminant fate and distribution in spe-

cific habitats, and the analysis is limited to the lower trophic levels of the food web.

In the trophic context, keystone species are typically studied by applying ecological network indices to food web models. Indicators based on the Mixed Trophic Impact analysis (Ulanowicz and Puccia, 1990) give a quantitative characterization of keystone species (Power et al., 1996, Ulanowicz, 2004, Libralato et al., 2006). Topological centrality indices provide instead qualitative descriptors of trophic importance (Jordán, 2009, Estrada, 2007, Bauer et al., 2010). In our framework, these indices are applied to identify key players in a contaminated food web and to this aim, we propose a novel formulation of keystone species based on sensitivity analysis.

Ciavatta et al. (2009) employ Monte Carlo-based global sensitivity analysis to study the relevance of chemical and ecological parameters in the computed concentration values of two species in the Venice lagoon food web. Similar applications of sensitivity analysis in bioaccumulation modelling are found in Carrer et al. (2000), Lamon et al. (2012), De Laender et al. (2010). In our work, we consider local sensitivity for the analysis of toxic keystones, by computing it from an ODE model parametrised with the bioaccumulation network outputs.

Within the proposed framework, our aim is twofold:

1. Reconstruct the first PCBs bioaccumulation model for the Adriatic food web, by providing a review of experimental concentration data and estimating missing concentrations.
2. Investigate the role of keystone species in the contaminant transfer through food webs by defining and evaluating a new index of species centrality.

2. Materials and Methods

Figure 1 illustrates the modelling and analysis steps implemented in our framework for the study of PCBs bioaccumulation in the Adriatic food web. First, we estimate the trophic and the bioaccumulation network using biomass and PCBs concentrations data, respectively. Second, we derive a dynamic ODE-based model from the obtained static bioaccumulation network. Last, we introduce a new index, Sensitivity Centrality, and evaluate its effectiveness through the comparison with a set of established centrality indices with respect to global network indices.

2.1. Adriatic Sea ecosystem

This study focuses on the Adriatic Sea, a distinct sub-region of the Mediterranean Sea, considered a critical water body due to the presence of intensive fishing effort (Coll et al., 2009) and river inputs flowing into the basin (Degobbis et al., 2000), which threatens the marine biological diversity (Coll et al., 2010, Penna et al., 2004). Within the Adriatic region, the freshwater discharge of the Po river exerts a strong effect on the chemical and physical dynamics of the coastal ecosystem (Marini et al., 2002). The river crosses a wide industrial and agricultural area and largely contributes to the nutrient and chemical loads flushed into the sea (Calamari et al., 2003), with a mean rate of discharge of $1500 \text{ m}^3 \cdot \text{s}^{-1}$ (Campanelli et al.,

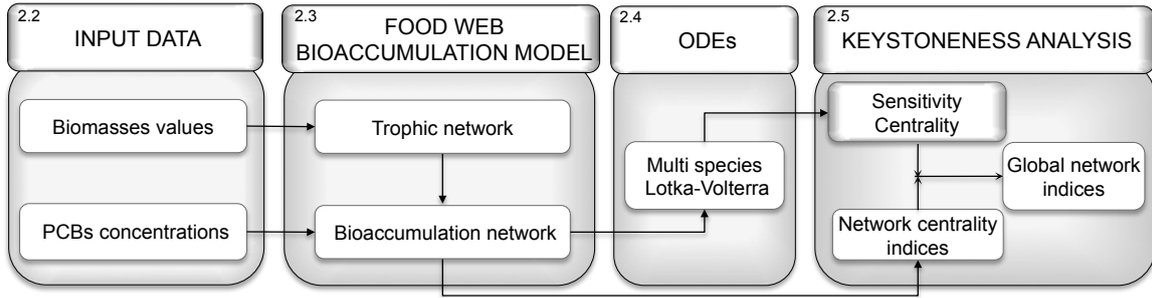


Figure 1: Bioaccumulation analysis framework. Workflow of performed analyses and corresponding section numbers

2011). The southeaster part of the Adriatic Basin is equally affected by riverine inputs, representing besides the unique canal to the open Ionian Sea (Marini et al., 2010). As a result, in the last decades a variety of organic chemicals have been detected in this region, with significant concentrations both in species and environmental compartments (Marini et al., 2012, Bellucci et al., 2002, Horvat et al., 1999, Kannan et al., 2002).

2.2. Input data

The ecological classification of Adriatic species and input data used in the estimation of the trophic model are obtained from the information collected in (Coll et al., 2007), a computational study aimed at evaluating fishing impacts on the Adriatic ecosystem during the 1990s. For each functional group we consider the following data (see Table A.5 in the Appendix): biomass B , production rate P , consumption rate Q , and fishing, which consists of landing (Lan) and discard (Dis) fractions. Biomass flows are expressed in $t \cdot km^{-2} \cdot yr^{-1}$ wet weight organic matter and biomass in $t \cdot km^{-1}$. Diet composition is illustrated in Fig. A.7 (a) of the supplementary materials.

In our analysis, input parameters of the bioaccumulation model have been compiled after an extensive literature review of published PCBs concentrations data, collected over the period 1994-2002 on species throughout the Adriatic region. All data sources report both the single value and the sum of all PCBs congeners detected in the edible part and muscle tissue of sampled species, expressed in $ng \cdot g^{-1}$ and $pg \cdot g^{-1}$ wet weight. In this work we consider the sum of PCBs congeners, reported in Table 1. For groups with no concentration data available, we take published PCBs values for comparable species by following the taxonomic classification in WoRMS database¹. A detailed list of all data sources is available in Table A.6 of the supplement. Units of the bioaccumulation model are expressed in $ng \cdot g^{-1}$ wet weight for concentrations, and $ng \cdot g^{-1} \cdot t \cdot km^{-2} \cdot y^{-1}$ for contaminant flows.

2.3. Food web bioaccumulation model

We define a mechanistic model of PCBs bioaccumulation, where contaminant pathways are coupled to trophic interactions. Flow rates quantify the intensity at which the medium

(i.e. biomass or contaminant) is transferred from the source (the prey) to the target (the predator). Flows are estimated under mass-balance conditions, i.e. the total inflows of a group must equal the total outflows, and are computed from input values of concentration and biomass. External unbalanced compartments are also included, to model potentially unlimited exogenous imports and exports.

The conceptual model of the bioaccumulation network is illustrated in Figure 2. Biomass and contaminant flows from a prey i to a predator j are denoted with $b_{i \rightarrow j}$ and $c_{i \rightarrow j}$, respectively. The flows included are: Consumption ($b_{j \rightarrow i}$) and contaminant uptake ($c_{j \rightarrow i}$) from a prey j . Predation ($b_{i \rightarrow k}$) and contaminant loss ($c_{i \rightarrow k}$) due to consumption by a predator k . External inputs from Import ($b_{Import \rightarrow i}$ and $c_{Import \rightarrow i}$), used to model generic inflows like immigration. Contaminant uptake from water ($c_{Water \rightarrow i}$), which clearly does not involve a corresponding biomass transfer. External outputs to Export ($b_{i \rightarrow Export}$ and $c_{i \rightarrow Export}$), used to model generic outflows like emigration. Respiration flows ($b_{i \rightarrow Respiration}$ and $c_{i \rightarrow Respiration}$) and flows to natural detritus ($b_{i \rightarrow Detritus}$ and $c_{i \rightarrow Detritus}$), constituting together the unassimilated portion of the biomass ingested by i . Outflows due to fishing activity, that can exit the food web to the landings ($b_{i \rightarrow Landing}$ and $c_{i \rightarrow Landing}$), or enter back the biomass cycle through the discard group ($b_{i \rightarrow Discard}$ and $c_{i \rightarrow Discard}$).

On the other hand, detritus and planktonic groups are assumed to be in instant equilibrium with the water phase. Following Hendriks et al. (2001), Laender et al. (2009), their concentration C_i is determined only by the concentration in water, C_{water} (measured in $\mu g/L$), and computed as:

$$C_i = C_{water} \cdot OC_i \cdot K_{OC}, \quad K_{OC} = k_{OC/OW} \cdot K_{OW} \quad (2.1)$$

where K_{OC} is the organic carbon-water partition ratio; K_{OW} is the octanol-water partition coefficient, which measures the ratio of contaminant solubility and its value differs among PCB congeners. Since in the model the sum of congeners is considered, we set $K_{OW} = 10^6$, given that the $\text{Log}K_{OW}$ of PCBs varies between 5 and 7 (Walters et al., 2011). The other parameters are $OC_i = 0.028$ and $k_{OC/OW} = 0.41$ (Laender et al., 2009).

2.3.1. Estimation with Linear Inverse Modelling

The trophic and bioaccumulation networks are estimated from biomass and concentration data through *Linear Inverse*

¹<http://www.marinespecies.org/>

Table 1: Input PCBs concentration data ($ng \cdot g^{-1}$) by functional group, with corresponding species and references. In order to account for multiple data source, we derive concentration ranges instead of single values.

| Id.Group | Σ PCB (min-max) | Species | Sources |
|------------------------------------|------------------------|---|--|
| 1.Phytoplankton | | | |
| 2.Micro and mesozooplankton | | | |
| 3.Macrozooplankton | | | |
| 4.Jellyfish | | | |
| 5.Suprabenthos | | | |
| 6.Polychaetes | | | |
| 7.Commercial benthic invertebrates | 1.24 - 20.29 | <i>M. galloprovincialis</i> | (Perugini et al., 2004) |
| 8.Benthic Invertebrates | | <i>M. galloprovincialis</i> , <i>C. gallina</i> <i>C. gallina</i> , <i>A. tuberculata</i> , <i>E. siliqua</i> , <i>M. galloprovincialis</i> | (Bayarri et al., 2001) (Marcotrigiano and Storelli, 2003) |
| 9.Shrimps | 0.346 - 11.61 | <i>P. longirostris</i> , <i>A. antennatus</i> <i>A. antennatus</i> , <i>P. longirostris</i> , <i>P. martia</i> | (Marcotrigiano and Storelli, 2003) (Storelli et al., 2003) |
| 10.Norway lobster | 0.2 - 10.63 | <i>N. norvegicus</i> | (Bayarri et al., 2001, Perugini et al., 2004, Storelli et al., 2003) |
| 11.Mantis shrimp | 2.64 - 11.61 | <i>S. mantis</i> | (Marcotrigiano and Storelli, 2003) |
| 12.Crabs | | | |
| 13.Benthic cephalopods | 0.3 1- 6.7 | <i>T. sagittatus</i> , <i>S. officinalis</i> <i>O. salutii</i> | (Perugini et al., 2004) (Marcotrigiano and Storelli, 2003) |
| 14.Squids | 9.53-37.7 | <i>L. vulgaris</i> <i>I. coindetii</i> | (Bayarri et al., 2001) (Marcotrigiano and Storelli, 2003) |
| 15.Hake 1 | 3.183 - 31.93 | <i>M. merluccius</i> | (Storelli et al., 2003, Marcotrigiano and Storelli, 2003) |
| 16.Hake 2 | | " | " |
| 17.Other gadiformes | | | |
| 18.Red mullets | | | |
| 19.Conger eel | 22.424 - 104 | <i>C. conger</i> | (Storelli et al., 2003, 2007) |
| 20.Anglerfish | 0.2 - | <i>L. boudegassa</i> | (Storelli et al., 2003) |
| 21.Flatfish | | | |
| 22.Turbot and brill | | | |
| 23.Demersal sharks | 2 - 42 | <i>C. granulosus</i> , <i>S. blainvillei</i> | (Storelli and Marcotrigiano, 2001) |
| 24.Demersal skates | 0.45 - | <i>R. miraletus</i> , <i>R. clavata</i> , <i>R. oxyrinchus</i> | (Storelli et al., 2003) |
| 25.Demersal fish 1 | 6.687 | <i>S. flexuosa</i> , <i>H. dactyloptereus</i> | (Storelli et al., 2003) |
| 26.Demersal fish 2 | " | " | " |
| 27.Bentopelagic fish | " | " | " |
| 28.European Anchovy | 1.22 - 62.7 | <i>E. encrasicolus</i> | (Perugini et al., 2004, Bayarri et al., 2001) |
| 29.European Pilchard | 5.327 - 26.25 | <i>S. pilchardus</i> | (Perugini et al., 2004, Storelli et al., 2003) |
| 30.Small Pelagic Fish | 4.54 - 31.9 | <i>S. aurita</i> <i>S. maris</i> , <i>A. rochei</i> | (Marcotrigiano and Storelli, 2003) (Storelli et al., 2003) |
| 31.Horse Mackarel | 6.761 | <i>T. trachurus</i> | (Storelli et al., 2003) |
| 32.Mackarel | 0.95 - 80.6 | <i>S. scombrus</i> | (Perugini et al., 2004, Bayarri et al., 2001) |
| 33.Atlantic bonito | | | |
| 34.Large Pelagic Fish | | | |
| 35.Dolphins | | | |
| 36.Loggerhead turtle | 0.63 - 23.49 | <i>C. caretta</i> | (Storelli et al., 2007, Corsolini et al., 2000) |
| 37.Sea birds | | | |
| 38.Discard | | | |
| 39.Detritus | | | |

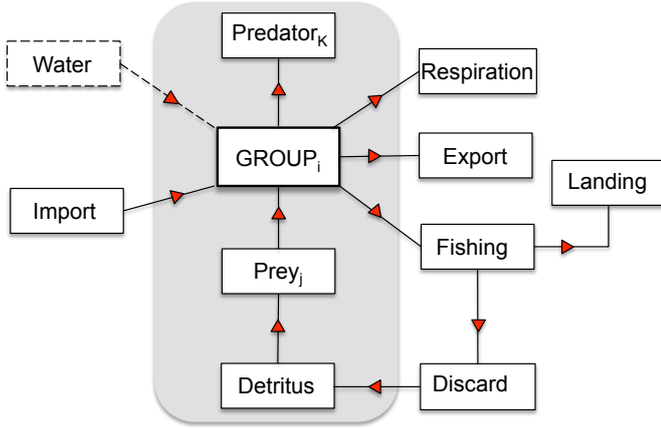


Figure 2: Conceptual bioaccumulation model. Black lines represent feeding linkages and red triangles specify the direction of trophic and contaminant flows. Mass-balanced groups are enclosed in the gray box (i , j , k refer to generic groups). External compartments (not mass-balanced) are depicted outside. The contaminant uptake from the water compartment (dashed) occurs without trophic interactions.

Modelling (LIM) (van Oevelen et al., 2010), used to quantify the unknown flow rates by solving a system of linear constraints.

We first reconstruct the trophic network by solving the constraints in Table 2. Then, the computed biomass flows are used as input parameters for the estimation of the PCBs bioaccumulation network, according to the linear inverse model in Table 3.

The trophic LIM was formulated as an approximate problem, i.e. including a set of equations $A \cdot x \simeq b$ that are not solved exactly, but in a least square sense. $\|A \cdot x - b\|^2$ is minimized and solutions are accepted up to a fixed tolerance value ϵ , by imposing the constraint $\|A \cdot x - b\| \leq \epsilon$ ($\|\cdot\|$ is the Euclidean norm). In the model, only diet and fishing constraints are solved approximately, with $\epsilon = 10^{-8}$. The corresponding exact problem resulted in an empty solution space, due to the many constraints generated from the available input data.

On the other hand, the exact bioaccumulation LIM yields a non-empty solution space, and thus multiple admissible values. This is explained by the partial availability of PCBs concentration data. In this case, we derive a statistically well-founded solution through Markov Chain Monte Carlo (MCMC) sampling of the solution space and by taking the mean of the sampled values. Note indeed that the mean of valid solutions to a linear system is in turn a valid solution to the system (van Oevelen et al., 2010).

2.4. Derivation of ODE Model

From the network estimation results we define a differential equation-based bioaccumulation model to analyse the dynamics of contaminant diffusion. In this work, the model is mainly used to compute the Sensitivity Centrality index, presented in Section 2.5.

We first derive a multi-species Lotka-Volterra ODE system that describes the temporal changes in species biomass. Then, the bioaccumulation model is obtained by extending the Lotka-Volterra system with parameters from the estimated PCBs

bioaccumulation network. The resulting differential equations describe the change rate of bioconcentrations, for all groups i :

$$\dot{C}_i(t) = w_i \cdot C_{water} + g_i \cdot C_i(t) + \sum_j \left(\frac{b_{j \rightarrow i} \cdot C_j(t) - b_{i \rightarrow j} \cdot C_i(t)}{B_i} \right) \quad (2.2)$$

where $C_i(t)$ is the concentration in i at time t ; C_{water} is assumed constant; w_i is the uptake rate from water by group i ; B_i is the biomass of i ; and g_i is the growth rate of i , computed as the sum of the exogenous inflows and outflow of biomass (diet flows excluded), over the estimated biomass:

$$g_i = \frac{\sum_j b_{j \rightarrow i} - \sum_j b_{i \rightarrow j}}{B_i}$$

where j ranges among the external groups. Details on the derivation of the bioaccumulation ODEs from the trophic Lotka-Volterra model are provided in Appendix A.1.

2.5. Sensitivity-based Keystoneness Analysis

Indices of species centrality in food webs are typically employed in trophic analysis for conservation purposes. Here, we apply these notions to the study of *toxic keystones*, i.e. species with a prominent role in the trophic transfer of a pollutant through the food web. We introduce a novel index based on the sensitivity analysis of ODE models, *Sensitivity Centrality (SC)*, in order to express dynamic aspects of bioaccumulation phenomena in the analysis of keystone species. In our study, we consider the ODE model in Section 2.4. SC is able to provide an intuitive characterization of centrality in a contaminated ecosystem: a keystone species is such that a perturbation of its concentration has a high impact on the total concentration of the food web.

Sensitivity Analysis (SA) (Brun et al., 2001, Soetaert and Petzoldt, 2010) computes the variation of model outputs with respect to variations of model inputs in order to evaluate how sensitive or robust the system is to perturbations. Specifically, SC is based on *local sensitivity*, where infinitesimal changes of model parameters are considered.

The Sensitivity Centrality of a group i at time t , $SC_i(t)$ quantifies the variation of the total contaminant concentration in the food web at time t , $C_{tot}(t)$, at infinitesimal variations of the initial concentration in group i , $C_i(0)$:

$$SC_i(t) = \left| \frac{\partial C_{tot}(t)}{\partial C_i(0)} \cdot \frac{C_i(0)}{C_{tot}(t)} \right|. \quad (2.3)$$

We take the absolute value of the sensitivity in order to give the same importance to both large positive and negative effects. The term $\frac{C_i(0)}{C_{tot}(t)}$ serves as a weighting factor.

2.5.1. Network centrality indices

In order to evaluate the introduced Sensitivity Centrality, we compare its performance with respect to other well-established indices of keystoneity, that are computed from the static bioaccumulation network. The considered indices are:

Keystoneity (KS) (Libralato et al., 2006), a quantitative descriptor of species' importance, weighted by their contribution

Table 2: Linear Inverse Model for the estimation of biomass flows of a functional group i . Approximate equations are solved with a least square approach.

Mass balances: $\sum_j b_{j \rightarrow i} - \sum_j b_{i \rightarrow j} = 0$

The difference between inflows and outflows within group i is zero; j ranges among groups and externals.

Ingestion¹: $I_i = \sum_j b_{j \rightarrow i} = B_i \cdot Q_i$

The total ingestion I_i , i.e. the sum of all the consumption flows, equals the product of biomass B_i and consumption rate Q_i . j is a functional group. Ingestion constraints are not included for detritus and phytoplankton.

Unassimilated Food: $b_{i \rightarrow \text{Respiration}} + b_{i \rightarrow \text{Detritus}} = I_i \cdot (1 - g_i)$

Respiration flows and flows to detritus are a fraction of the total ingestion and accounts for the proportion of food that is not converted into biomass. $g_i = \frac{P_i}{Q_i}$ is the gross food conversion efficiency (Christensen and Walters, 2004).

Respiration-assimilation: $b_{i \rightarrow \text{Respiration}} \leq I_i \cdot g_i$

The ratio respiration/assimilation has to be lower than one to have realistic estimates (Coll et al., 2007).

Diet¹: $b_{i \rightarrow j} \approx I_j \cdot D_{ij}$.

The biomass flow from prey i to predator j is the fraction of the total ingestion of j coming from i . D is the diet composition matrix, reported in Fig. A.7 (a) of the supplementary materials.

Fishing: $b_{i \rightarrow \text{Landing}} \approx Lan_i$ and $b_{i \rightarrow \text{Discard}} \approx Dis_i$.

Input data on fishing activity (Lan_i and Dis_i) is used to constrain the exports to landings and discards.

Non-negativity of flows: $b \geq 0$

¹For species with uncertain input biomass and diet data, appropriate inequalities are set.

Table 3: Linear Inverse Model for the estimation of the PCBs concentration of a functional group i . Concentrations in detritus and planktonic groups are computed differently (see Eq. 2.1)

Mass balances: $\sum_j c_{j \rightarrow i} - \sum_j c_{i \rightarrow j} = 0$

Concentrations are estimated under mass-balance assumption. j ranges among functional groups and externals.

Concentration data: $C_i \bowtie k$

k is an input PCBs value that constrains the contaminant concentration C_i , with $\bowtie \in \{=, \leq, \geq\}$. Note that the same group can have associated arbitrarily many such constraints.

Uptake/losses from food: $c_{j \rightarrow i} = b_{j \rightarrow i} \cdot C_j$

The trophic transfer of contaminant from group j to i is the product of the corresponding biomass flow $b_{j \rightarrow i}$ and the concentration in j . It describes the PCBs uptake of predator i from prey j , and the PCBs removal from prey j due to consumption by i . The equation also characterizes the PCBs losses from j to the externals, if $i \in \{\text{Export, Respiration, Landing}\}$.

Uptake from generic imports: $c_{\text{Import} \rightarrow i} = b_{\text{Import} \rightarrow i} \cdot C_i$

The PCBs concentration in the biomass imported into group i is assumed to be the same as in i .

Uptake from water: $c_{\text{Water} \rightarrow i} = w_i \cdot C_{\text{Water}}$

w_i is the uptake rate of i from water-borne contaminant and C_{Water} is the contaminant concentration in water¹. w_i is assumed null for compartments in rapid equilibrium with the water phase.

Non-negativity of concentrations: $C_i \geq 0$

¹ If C_{water} is unknown, the constraint becomes non linear and w_i cannot be directly estimated. In that case, $c_{\text{Water} \rightarrow i}$ is estimated as a single variable.

in terms of contaminant concentration:

$$KS_i = \log(\epsilon_i \cdot (1 - p_i)), \quad p_i = \frac{C_i}{\sum_j C_j}, \quad \epsilon_i = \sqrt{\sum_{j \neq i} M_{ij}^2} \quad (2.4)$$

where p_i is the contribution of group i in the total concentration of the food web; and ϵ_i is the so-called *overall effect* of i , i.e. the sum of the indirect trophic impacts of i on all the other species. M_{ij} represents the contaminant variation in a group j at infinitesimal increases of the contaminant in i , and is calculated through the *Mixed Trophic Impact analysis* (Ulanowicz and Puccia, 1990).

Flow Betweenness Centrality (FBC) (Freeman et al., 1991), a qualitative index expressing the topological importance of a group in maintaining the flows within the food web:

$$FBC_i = \sum_{j \neq k, j \neq i, k \neq i} (max_G c_{j \rightarrow k} - max_{G \setminus i} c_{j \rightarrow k}) \quad (2.5)$$

where $max_G c_{j \rightarrow k}$ is the maximum contaminant flow between groups j and k in the food web G , and $max_{G \setminus i} c_{j \rightarrow k}$ is the maximum flow when group i is excluded from G .

2.5.2. Global indices

Global network indices are used to derive descriptors of the structure and properties of the whole ecosystem (Kones et al., 2009). They are applied in this case to the study of contaminated food webs in order to analyse bioaccumulation phenomena at the ecosystem level. By definition, a species plays a key role in its ecosystem if its extinction (or perturbation) has a large impact on global ecosystem functions.

Following Estrada (2007), Allesina and Pascual (2009), we employ global indices to compare the introduced keystone indices. In this analysis, we simulate sequences of extinctions performed in decreasing order of importance, as ranked by each index. Then, a keystone index is considered good when the associated extinction sequence produces a substantial disruption in the global ecosystem properties of interest. The global indices analysed are:

Total system throughflow (TST). It describes the total ecosystem activity as the sum of outflows (exports included) of all groups:

$$TST = \sum_i \sum_j c_{i \rightarrow j}$$

Average path length (APL). It computes the average number of groups that each inflow passes through and thus, it measures the efficiency of contaminant transport pathways:

$$APL = \frac{TST}{\sum_i \sum_{j \in Ext} c_{i \rightarrow j}}$$

where $Ext = \{\text{Respiration, Export, Landing}\}$ is the set of external export compartments.

Link density (LD). It expresses the average number of links per

species, providing a structural descriptor of the food web:

$$LD = \frac{\sum_i \sum_j (c_{i \rightarrow j} > 0)}{n}$$

where n is the number of groups in the food network.

3. Results and Discussion

3.1. Trophic and Bioaccumulation Network

Results of the estimated Adriatic trophic and bioaccumulation networks are reported in Table 4 and graphically represented in Figure 3.

The analysis of the biomass network shows that groups occupying lower trophic positions are the most predominant in terms of biomass and flows. As visible in Fig. 3 (a), biomass values and trophic levels are negatively correlated. The only exception is the *Discard* group, which accounts for the discarded catches that enter back the food web. *Discard* is considered in this model as a detritus, and thus has a trophic level (TL) of 1. However, it clearly possesses a much lower biomass than the natural detritus (group *Detritus*) and primary producers (group *Phytoplankton*). We report that our quantitative estimations agree with the original work by Coll et al. (2007), which confirms the consistency of our trophic model.

PCBs Bioaccumulation Analysis. On the contrary, PCBs bioaccumulation values tend to increase at higher trophic levels. Thus, a phenomenon of *biomagnification* is detected, as visible in the network plot (Fig. 3 (b)). Estimated PCBs values in top predators ($TL \geq 4$) are: *Large Pelagic Fish* (70.491), *Demersal skates* (54.833), *Turbot and brill* (54.746), *Dolphins* (54.048), *Anglerfish* (53.808), and *Atlantic bonito* (52.704). Note that, *Large Pelagic Fish* (Tuna and Swordfish) shows by far the highest value, due to the concentration in groups composing its diet (mainly *European Anchovy* and *Squids*).

Moreover, the quality of input data (reported in Table 1) also influences estimated values. Specifically, the concentration in some top predators (*Squids*, *Hake 2* and *Demersal Sharks*) is limited since their valid PCBs values are constrained with low upper bounds. On the contrary, high or infinite upper bounds yield high concentration values also for groups at $TL=3$, like in *Demersal fish 2*, *Flatfish*, *Bentopelagic fish* and *Mackarel*. This is related to the fact that we use a stochastic search algorithm to solve the corresponding LIM problem, also implying that less constrained variables have higher variability (see standard deviation in Table 4).

Fishing effects on bioaccumulation. Fishing activity represents quantitatively a biomass loss and therefore a contaminant removal from the food web. In our model, relatively low PCBs values are detected for overexploited groups, i.e. where fishing rates exceed biomass. For instance, *Crabs*, *Other gadiformes* and *Red mullets* show concentrations ($< 1 \text{ ng} \cdot \text{g}^{-1}$) substantially lower than those in species belonging to the same TL, but not affected by fishing pressure. A similar tendency is observed for *Conger eel* whose resulting concentration is close to the lower

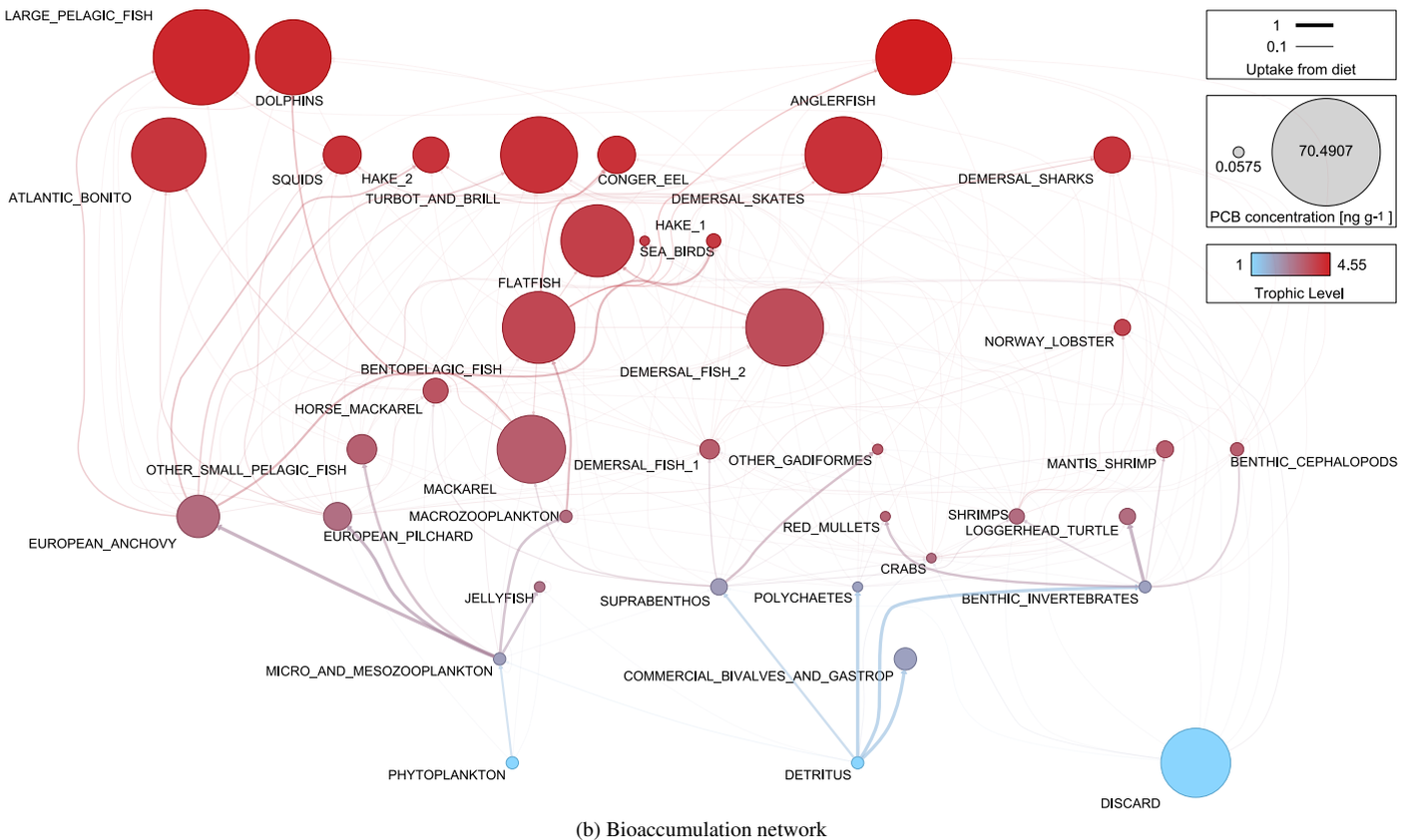
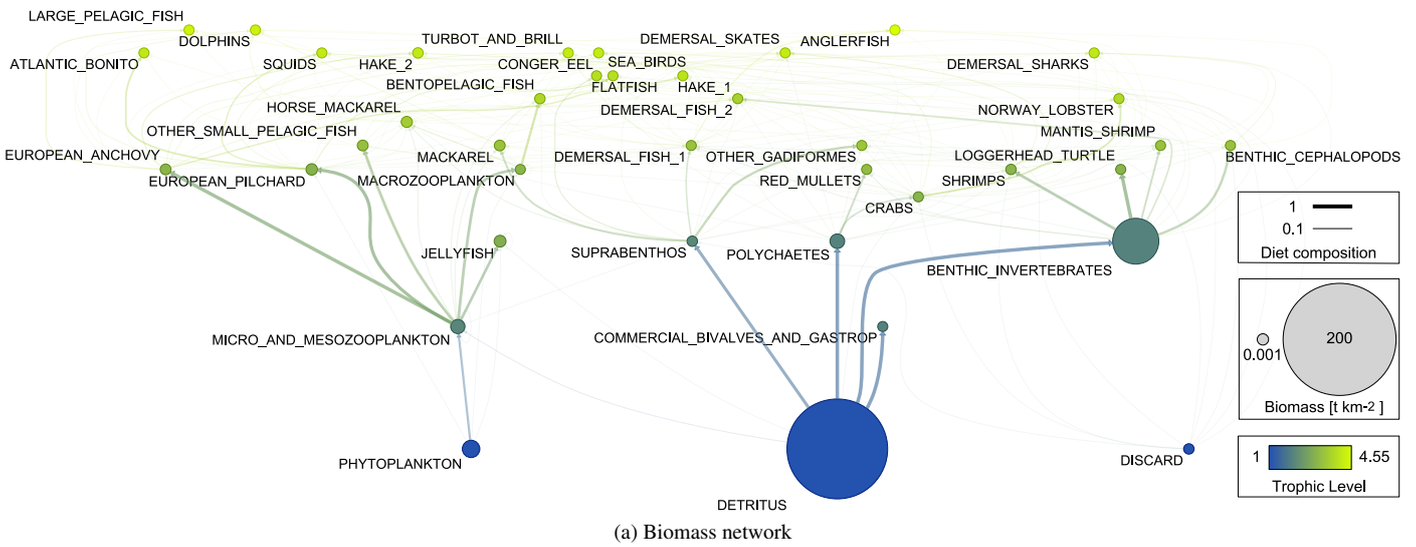


Figure 3: Estimated Adriatic trophic (a) and PCBs bioaccumulation (b) networks. Nodes represent functional groups with size proportional to the biomass content (a) or PCBs concentration (b). Edges represent feeding connections with thickness proportional to the biomass (a) or contaminant (b) flow from the prey to the predator. Flows to detritus and to fishing discards are not shown.

Table 4: Results of trophic and bioaccumulation networks estimation: TL , trophic level; B biomass ($t \cdot km^{-1}$); PCBs concentration \pm standard deviation obtained with MCMC-based sampling ($\sum PCBs, ng \cdot g^{-1}$); F ($t \cdot ng \cdot y^{-1} \cdot km^{-2}$) indicates PCBs outflows due to fishing, and Dis the fraction discarded. Keystoneness indices for the bioaccumulation model (highest values in bold) are KS , Keystoneness; FBC , Flow Betweenness Centrality; and $SC(i)$, Sensitivity Centrality at year i .

| Id.Group | TL | B | $\sum PCBs$ | F | Dis | KS | FBC | $SC(1)$ | $SC(2)$ |
|------------------------------|------|--------|---------------------|--------|-------|--------------|----------------|--------------|-------------|
| 1.Phytoplankton | 1 | 16.658 | 2.247 \pm 0.955 | 0 | 0 | -1.394 | 0 | 0.001 | 0.001 |
| 2.Micro and mesozoop. | 2.05 | 9.512 | 2.247 \pm 0.955 | 0 | 0 | 0.673 | 557.983 | 0.024 | 0.024 |
| 3.Macrozoop. | 3.05 | 0.54 | 2.247 \pm 0.955 | 0 | 0 | -3.479 | 11.474 | 0.005 | 3.891 |
| 4.Jellyfish | 2.88 | 4 | 0.843 \pm 0.353 | 0 | 0 | -0.296 | 0.331 | 0.001 | 0.001 |
| 5.Suprabenthos | 2.11 | 1.01 | 5.952 \pm 3.247 | 0 | 0 | -5.62 | 192.88 | 0.006 | 0.006 |
| 6.Polychaetes | 2 | 9.984 | 0.348 \pm 0.2 | 0 | 0 | -0.849 | 5.406 | 0 | 1.358 |
| 7.Commercial benthic invert. | 2 | 0.043 | 10.502 \pm 5.369 | 0.368 | 0 | -0.717 | 0 | 0.01 | 0.01 |
| 8.Benthic Invert. | 2 | 79.763 | 2.12 \pm 0.846 | 0.696 | 0.696 | -1.475 | 38.462 | 0 | 3.899 |
| 9.Shrimps | 3.02 | 0.68 | 4.746 \pm 2.752 | 0.157 | 0.081 | -0.685 | 50.236 | 0.004 | 3.901 |
| 10.Norway lobster | 3.77 | 0.018 | 5.542 \pm 3.115 | 0.205 | 0 | -0.751 | 6.257 | 0.005 | 3.902 |
| 11.Mantis shrimp | 3.31 | 0.015 | 6.103 \pm 3.228 | 0.439 | 0 | -2.043 | 1.515 | 0.006 | 0.006 |
| 12.Crabs | 3.00 | 0.009 | 0.058 \pm 0.039 | 0.01 | 0.01 | -3.119 | 75.123 | 0 | 0.224 |
| 13.Benthic cephal. | 3.31 | 0.068 | 3.198 \pm 1.77 | 0.499 | 0.006 | -1.28 | 16.473 | 0.003 | 3.899 |
| 14.Squids | 4.14 | 0.02 | 23.289 \pm 8.172 | 0.955 | 0 | -2.519 | 34.117 | 0.022 | 0.022 |
| 15.Hake 1 | 4.00 | 0.06 | 3.852 \pm 0.617 | 0.705 | 0.27 | -2.805 | 14.277 | 0.004 | 3.9 |
| 16.Hake 2 | 4.11 | 0.56 | 21.658 \pm 5.95 | 0 | 0 | -2.676 | 2.914 | 0.017 | 0.02 |
| 17.Other gadiformes | 3.37 | 0.029 | 0.567 \pm 0.285 | 0.061 | 0.047 | -2.664 | 2.186 | 0.001 | 2.211 |
| 18.Red mullets | 3.19 | 0.025 | 0.385 \pm 0.262 | 0.043 | 0 | -2.652 | 0.917 | 0 | 1.5 |
| 19.Conger eel | 4.16 | 0.005 | 22.879 \pm 0.443 | 0.183 | 0.183 | -0.261 | 5.791 | 0.022 | 3.919 |
| 20.Anglerfish | 4.55 | 0.006 | 53.808 \pm 29.234 | 0.377 | 0 | -3.176 | 0.395 | 0.051 | 0.052 |
| 21.Flatfish | 3.89 | 0.009 | 51.436 \pm 30.137 | 2.057 | 0 | -1.643 | 16.564 | 0.049 | 3.946 |
| 22.Turbot and brill | 4.15 | 0.04 | 54.746 \pm 28.303 | 0.876 | 0 | -3.803 | 19.136 | 0.052 | 3.95 |
| 23.Demersal sharks | 4.09 | 0.018 | 21.84 \pm 11.571 | 0.175 | 0 | -3.309 | 0.224 | 0.021 | 0.021 |
| 24.Demersal skates | 4.15 | 0.003 | 54.833 \pm 29.337 | 0.11 | 0 | -1.788 | 0.073 | 0.052 | 0.053 |
| 25.Demersal fish 1 | 3.32 | 0.056 | 8.159 \pm 1.132 | 0.865 | 0.416 | -1.15 | 135.968 | 0.008 | 3.904 |
| 26.Demersal fish 2 | 3.62 | 0.24 | 55.424 \pm 28.667 | 0.942 | 0.055 | -0.258 | 96.253 | 0.052 | 3.95 |
| 27.Bentopelagic fish | 3.73 | 1.2 | 51.324 \pm 28.892 | 0.103 | 0 | -1.108 | 40.106 | 0.049 | 3.946 |
| 28.European Anchovy | 3.05 | 1.497 | 27.104 \pm 18.17 | 13.579 | 0.136 | -2.038 | 19.736 | 0.025 | 3.922 |
| 29.European Pilchard | 2.97 | 2.985 | 14.969 \pm 6.234 | 6.077 | 0.629 | -0.764 | 3.809 | 0.014 | 3.911 |
| 30.Small Pelagic Fish | 3.25 | 1.517 | 16.533 \pm 8.18 | 0.215 | 0.017 | -2.124 | 3.231 | 0.016 | 3.913 |
| 31.Horse Mackarel | 3.49 | 2.455 | 12.496 \pm 1.171 | 0.275 | 0.025 | 0.157 | 14.643 | 0.012 | 3.909 |
| 32.Mackarel | 3.32 | 1.683 | 47.96 \pm 21.422 | 1.199 | 0.384 | -0.586 | 32.264 | 0.045 | 3.943 |
| 33.Atlantic bonito | 4.09 | 0.3 | 52.704 \pm 29.504 | 0.949 | 0 | -1.075 | 0.076 | 0.05 | 0.051 |
| 34.Large Pelagic Fish | 4.34 | 0.138 | 70.491 \pm 20.461 | 1.833 | 0 | -1.804 | 0.144 | 0.058 | 0.067 |
| 35.Dolphins | 4.30 | 0.012 | 54.048 \pm 29.286 | 0.005 | 0.005 | -1.276 | 13.609 | 0.052 | 3.949 |
| 36.Loggerhead turtle | 3.01 | 0.032 | 5.478 \pm 4.412 | 0.022 | 0.022 | -0.816 | 0.423 | 0.005 | 0.005 |
| 37.Sea birds | 3.90 | 0.001 | 0.161 \pm 0.058 | 0 | 0 | -1.211 | 0.067 | 0 | 0 |
| 38.Discard | 1 | 0.733 | 49.005 \pm 0.84 | 0 | 0 | | | | |
| 39.Detritus | 1 | 200 | 2.247 \pm 0.955 | 0 | 0 | | | | |

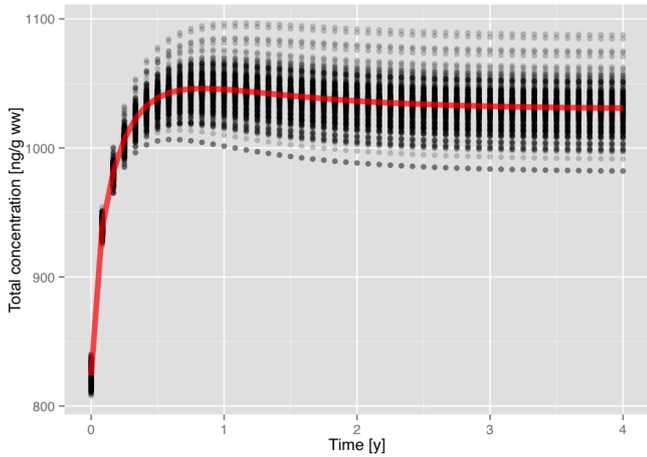


Figure 4: Temporal evolution of total PCBs concentrations, simulated through the dynamic ODE model over a period of 4 years and time step of 1 month (red line). Shaded dots indicate the results obtained after random perturbations of the initial concentration (100 uniformly distributed values for each species).

bound of the considered input PCBs values. The effects are even more evident if comparing concentrations of groups belonging to the same species but subject to different fishing pressures: *Hake 1* (< 40 cm, vulnerable to fishing, $PCB=3.852$) and *Hake 2* (> 40 cm, not vulnerable, $PCB=21.658$); or *Demersal fish 1* (overexploited, $PCB=8.159$) and *Demersal fish 2* ($PCB=55.424$).

Differently from natural detritus, the *Discard* group is characterized by a significant PCBs concentration. This mainly depends on its low biomass and on its species composition that, with their concentration, contributes to a considerable total inflow of contaminant to fishing discards (see Table 4, column *Dis*). Finally, with our bioaccumulation model we can estimate the PCBs concentration in the landing fraction of the biomass exported by fishing, computed as $\sum_i c_{i \rightarrow \text{Landing}} / \sum_i b_{i \rightarrow \text{Landing}}$. The mean concentration in landings is $18.17 \text{ ng} \cdot \text{g}^{-1}$. This kind of analysis could provide an effective indicator of the chemical pollution in species of commercial interest.

3.2. ODE bioaccumulation model

We evaluate the long-term PCBs bioaccumulation dynamics in the Adriatic food web by simulating the ODE model (Eq. A.4). Figure 4 shows the results obtained under random perturbations of the initial concentration values taken from the static bioaccumulation network. We observe that the total PCBs concentration has a steep increase before reaching a plateau in the second year of the simulation. This is explained by the fact that groups in rapid equilibrium with the water phase (planktonic and detritus groups) are not mass-balanced in the bioaccumulation network. We notice that, in order to support the correctness and the applicability of the Sensitivity Centrality index (see Sect. 2.5), the qualitative dynamics of the system are relatively robust with respect to changes in the initial conditions.

3.3. Keystoneness analysis

The four networks plotted in Figure 5 show the centrality of species according to the indices of toxic keystoneess considered: *SC(1)* Sensitivity Centrality, year 1; *SC(2)* Sensitivity Centrality, year 2; *KS* Libralato's keystoneess; and *FBC* Flow Betweenness Centrality. Numerical results are reported in Table 4. Both KS and FBC identify the group *Micro and mesozooplankton* (id 2) as keystone, which is however not relevant for *SC(1)* and *SC(2)*. According to *SC(1)*, *Large pelagic fish* (34) is the most important group, but is low ranked by the other indices. This group also shows the highest estimated PCBs value. The highest *SC(2)* value is registered in both *Turbot and brill* (22) and *Demersal fish 2* (26). Remarkably, the latter occupies important positions also in other indices: 3rd in KS and *SC(1)*, 4th in FBC. As evidenced in Fig. 5 (c), FBC assigns a disproportionate centrality value (557.983) to its keystone. On the contrary, differences in KS (plot d) are less evident among groups since the index is computed with a logarithmic operation (see Sect. 2.5.1). In the evaluation of *SC(1)* (plot a), two clusters of species can be distinguished: one with higher values ranging in the interval $[0.045, 0.058]$, and the other one with $SC(1) \in [0, 0.025]$. The groups in the first cluster are (in decreasing order of importance): 34, 35, 26, 24, 22, 20, 33, 21, 27 and 32. They occupy high trophic positions ($TL \in [3.31, 4.55]$) and have initial PCBs concentrations ranging between 47.96 and $70.49 \text{ ng} \cdot \text{g}^{-1}$. *SC(2)* (plot b) performs in a similar way, identifying a larger set of important groups with values between 3.891 and 3.95, and with minor differences among species scores. Despite there are changes in species classification between *SC(1)* and *SC(2)*, groups 21, 22, 26, 27, 32 and 35 maintain top centrality positions in both rankings.

Comparison by extinction simulation. The comparison of Sensitivity Centrality with the other indices of keystoneess has been carried out by simulating the effects of successive extinctions on global network properties, performed by following the different rankings. Ideally, an effective index would identify the set of species that, when removed, produce strong perturbations at the global ecosystem level. Figure 6 illustrates the results on the total system throughflow (*TST*), i.e. the sum of contaminant flows; link density (*LD*), the average number of links per group; and the average path length (*APL*), a measure of network robustness telling the average number of groups that each inflow passes through.

In our analysis, a good index of toxic keystoneess should lead to a rapid breakdown in *TST*, since low values imply a low total rate of contaminant transfer. We observe that extinctions performed following the rankings of *SC(1)* and *SC(2)* have the highest impact, thus indicating that *SC* gives an effective measure of importance as regards the total activity of the network. In particular, they halve the value of *TST* even before the first 10 extinctions, which is instead obtained only after 17 extinctions for KS and FBC.

Similarly, the extinction of toxic keystones should yield decreased values of *LD*, since species with many trophic links have potentially a pivotal role in the contaminant transfer

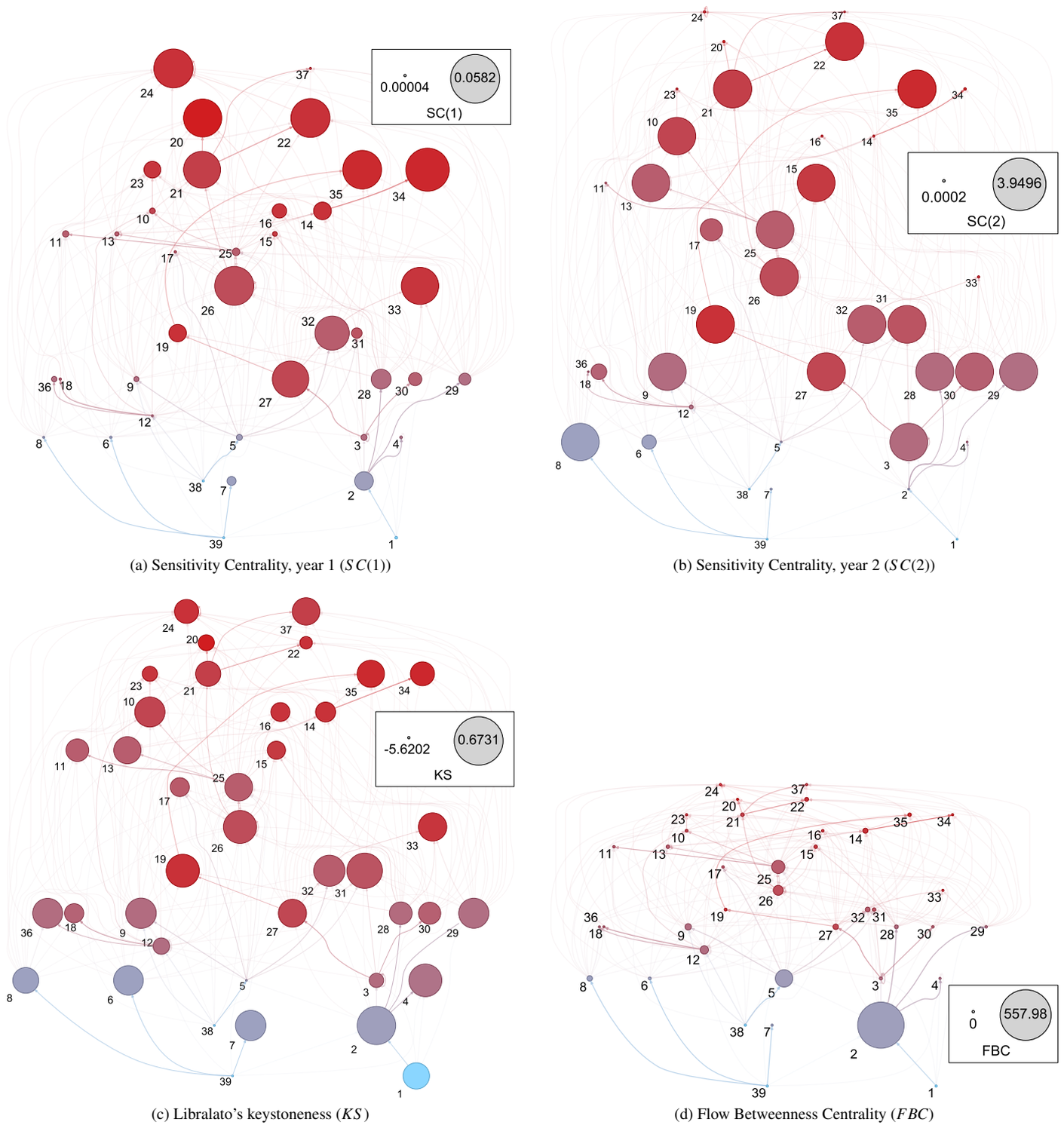


Figure 5: Ranking of species in the Adriatic bioaccumulation model according to $SC(1)$ (a), $SC(2)$ (b), KS (c) and FBC (d). In the network plots, node size is proportional to the relative importance of species given by the corresponding indices. Node colour represents the trophic level as in Fig. 3 (b).

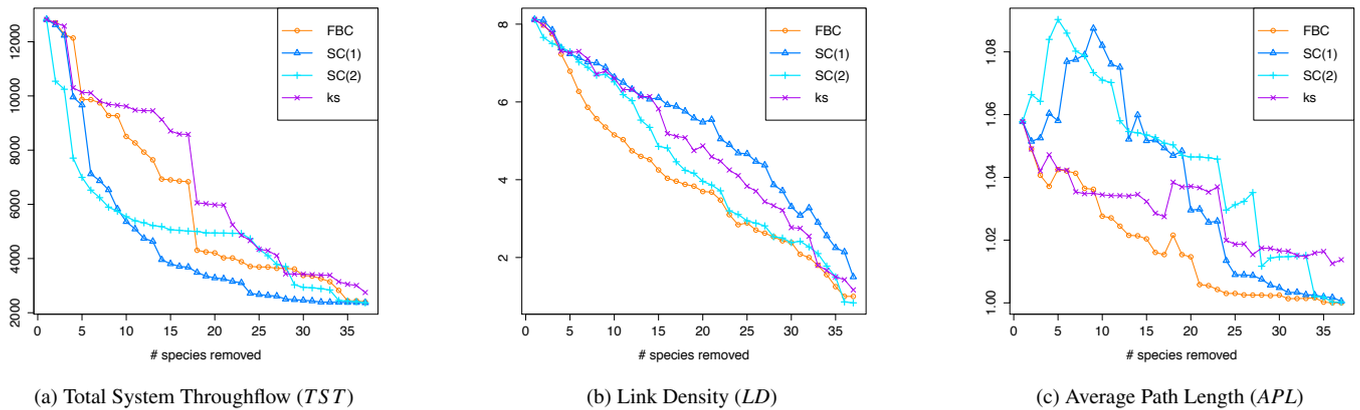


Figure 6: Simulation of the effects of extinctions on global network indices: TST (a) ($ng \cdot g^{-1} \cdot km^{-2} \cdot y^{-1}$), LD (b), APL (c). Species are removed according to the ranking of toxic keystone-ness (from the most important to the least) obtained by Flow Betweenness Centrality (FBC), orange line; Sensitivity Centrality at 1 and 2 years ($SC(1)$ and $SC(2)$), light and dark blue line, respectively; and Libralato’s keystone-ness (KS), purple line.

through the food web. In this case, FBC shows the best overall performance, even if the first 3 extinctions according to the $SC(2)$ ranking produce the strongest effect. The qualitative nature of LD explains the good performance of FBC , which is an index based on topological considerations.

APL measures the efficiency of contaminant transport pathways, that is the number of trophic links needed to transfer the contaminant between any pair of groups. Therefore after the extinction of a toxic keystone an increased value is expected. For this network property, $SC(1)$ and $SC(2)$ have the best performance suggesting that our Sensitivity Centrality is effective in identifying species that promote the overall efficiency of contaminant transfer.

3.4. Implementation

We implemented the computational framework in R software with following packages: *LIM* (van Oevelen et al., 2010), estimation of the trophic and bioaccumulation network; *NetIndices* (Kones et al., 2009), global network indices (TST , LD and APL) and trophic levels; *enaR* (Borrett and Lau, 2012), MTI analysis (required to compute KS); *sna* (Butts, 2008), FBC estimation; and *FME* (Soetaert and Petzoldt, 2010), ODE simulations and sensitivity analysis. Network plots have been generated with *Graphviz* (Ellson et al., 2002).

4. Conclusions

Recent advances in the application of food web ecology to ecotoxicological research have highlighted the influence of contaminants on the ecosystem structure and functions, and improved the ability of models to predict bioaccumulation phenomena in aquatic food webs. Ideally, an integrated pipeline of methods and information ranging from the molecular to the ecological level can effectively support environmental decision-making with a more comprehensive understanding of contamination patterns in an ecosystem. In this context, the main contribution of this study is the combination of computational and

network analysis tools to estimate bioaccumulation in contaminated food webs and to identify keystone species in contamination pathways. We reconstructed the first food web-based bioaccumulation model for the Adriatic sea, providing a state-of-the-art review of PCBs experimental concentration data in species. Further, we defined a novel index based on sensitivity analysis, Sensitivity Centrality, to analyse toxic keystone-ness. Our estimations evidenced that species at higher trophic positions exhibit higher PCBs concentrations, thus suggesting the occurrence of biomagnification phenomena in the Adriatic food web. Finally, we showed that the newly introduced index is able to identify species that have a large impact on the total amount of contaminant flows and on the PCBs transfer efficiency through trophic pathways.

We believe that the concept of sensitivity gives a simple yet mathematically well-grounded characterization of toxic keystones and more generally, that the problem of identifying the key species in a contaminated ecosystem could be better addressed by using indices that incorporate both network-level and dynamic information. These kinds of analyses could be more meaningful than just evaluating concentrations on single species, and could reveal which are the bioindicators to monitor and control in a polluted ecosystem for both conservation and remediation purposes (Taffi et al., 2014). Likewise, clarifying species centrality in the contaminant transfer could help in predicting the temporal evolution of bioaccumulation under different scenarios of natural or anthropogenic perturbations.

Acknowledgements

This work has been supported by the RITMARE Flagship Project funded by the Italian Ministry of University and Research.

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Appendix A. Supplementary Materials

Appendix A.1. Derivation of dynamic bioaccumulation ODEs

We define a dynamic bioaccumulation model on top of a multi-species Lotka-Volterra system used to describe the temporal changes in species biomass. In its general form, the Lotka-Volterra system is formulated as:

$$\dot{B}_i(t) = B_i(t) \left(g_i - \sum_j A_{ij} B_j(t) \right) \quad (\text{A.1})$$

where $B_i(t)$ is the biomass of species i at time t ; g_i is the intrinsic growth rate of i ; and A is the interaction matrix. In particular A_{ij} describes the predation effect of species j on species i . Parameters of the ODE model are derived from the estimated food web model:

- $B_i(0) = B_i$, initial biomass values are those in the static food web estimated with LIM;
- $g_i = \frac{\sum_j b_{j \rightarrow i} - \sum_j b_{i \rightarrow j}}{B_i(0)}$, with j ranging among the external groups; the growth rate of i is the sum of exogenous inflows and outflow, over the estimated biomass of i ;

- $A_{ij} = \frac{b_{i \rightarrow j} - b_{j \rightarrow i}}{B_i(0) \cdot B_j(0)}$, the interaction rate between prey i and predator j is calculated as the net flow from i to j divided by the estimated biomasses of i and j .

Additionally, we define the biomass flow rate from group i to j at time t , $b_{i \rightarrow j}(t)$, which is non-linear with respect to the biomasses of i and j , as:

$$b_{i \rightarrow j}(t) = \frac{b_{i \rightarrow j}}{B_i(0) \cdot B_j(0)} \cdot B_i(t) \cdot B_j(t)$$

in such a way that Eq. A.1 can be rewritten as:

$$\dot{B}_i(t) = g_i \cdot B_i(t) + \sum_j b_{j \rightarrow i}(t) - \sum_j b_{i \rightarrow j}(t)$$

Therefore, the dynamics of the contaminant concentration in species i , $C_i(t)$, is given by the net sum of contaminant flows, over the biomass of i :

$$\dot{C}_i(t) = w_i \cdot C_{water} + \frac{g_i \cdot B_i(t) \cdot C_i(t) + \sum_j b_{j \rightarrow i}(t) \cdot C_j(t) - \sum_j b_{i \rightarrow j}(t) \cdot C_i(t)}{B_i(t)} \quad (\text{A.2})$$

where C_{water} is the concentration in water (assumed constant) and w_i is the uptake rate from water by group i . As done for the biomass equations, the initial concentrations correspond to those estimated in the static bioaccumulation network: $C_i(0) = C_i$, for each group i .

Finally, expanding the interaction terms, Eq. A.2 is equivalent to the following:

$$\dot{C}_i(t) = w_i \cdot C_{water} + g_i \cdot C_i(t) + \sum_j \left(\frac{b_{j \rightarrow i}}{B_i(0) \cdot B_j(0)} \cdot B_j(t) \cdot C_j(t) \right) - \sum_j \left(\frac{b_{i \rightarrow j}}{B_i(0) \cdot B_j(0)} \cdot B_j(t) \cdot C_i(t) \right) \quad (\text{A.3})$$

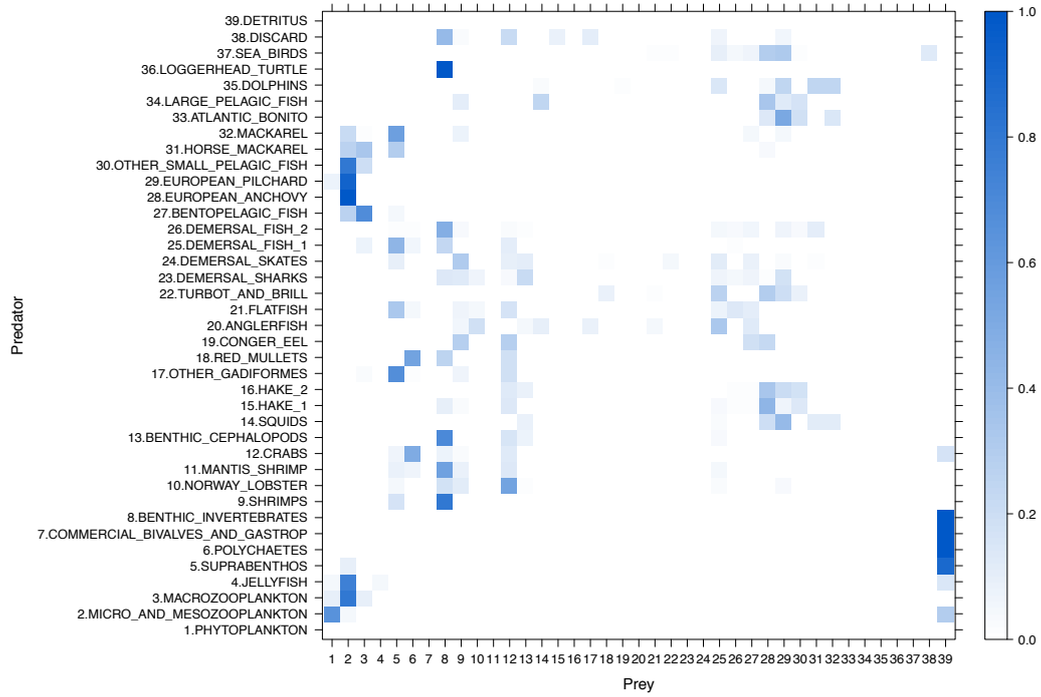
We focus on the temporal changes in concentrations independent of the biomass variations, thus assuming constant species biomass ($B_i(t) = B_i(0), \forall t$), which gives the following system of linear differential equations:

$$\dot{C}_i(t) = w_i \cdot C_{water} + g_i \cdot C_i(t) + \sum_j \left(\frac{b_{j \rightarrow i} \cdot C_j(t) - b_{i \rightarrow j} \cdot C_i(t)}{B_i(0)} \right) \quad (\text{A.4})$$

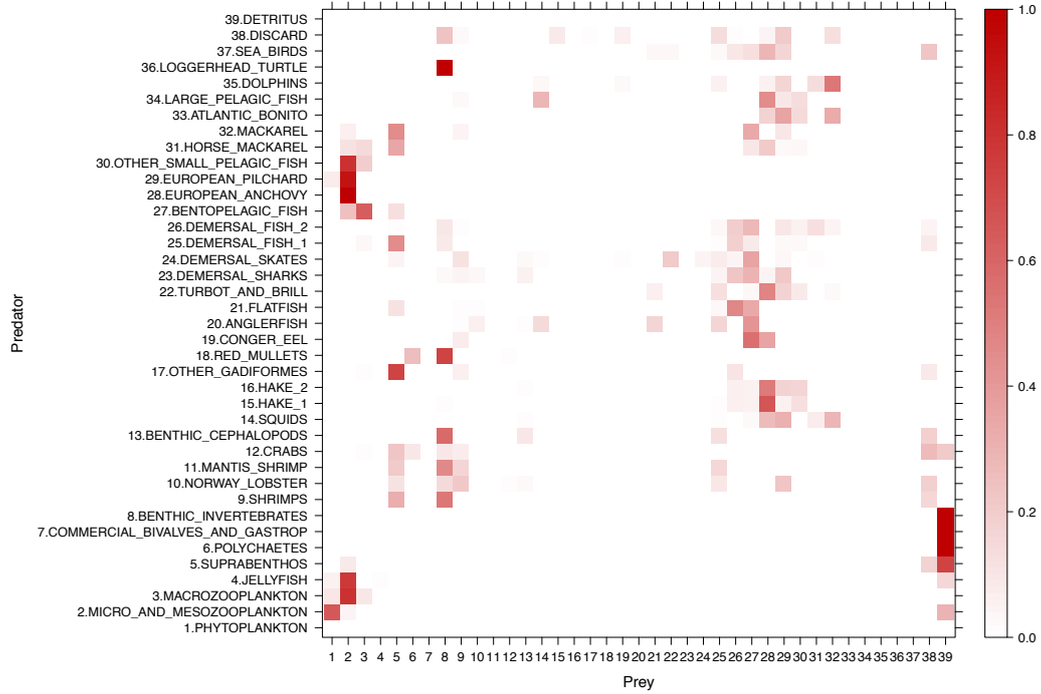
Note that this simplification does not change the quantitative dynamics of the model, because biomasses have been estimated under mass-balance conditions.

Table A.5: Input data for the estimation of the trophic network from (Coll et al., 2007). B , biomass ($t \cdot km^{-2}$); P , production rate (yr^{-1}); Q , consumption rate (yr^{-1}); Lan , landed fraction of biomass exported by fishing ($t \cdot km^{-2} \cdot yr^{-1}$), and Dis the fraction discarded ($t \cdot km^{-2} \cdot yr^{-1}$).

| Id.Group | B | P | Q | Lan | Dis |
|------------------------------------|---------------|-------|-------|-------|--------|
| 1.Phytoplankton | 16.658 | 69.03 | | | |
| 2.Micro and mesozooplankton | 9.512 | 30.43 | 49.87 | | |
| 3.Macrozooplankton | 0.54 | 21.28 | 53.14 | | |
| 4.Jellyfish | 4 | 14.6 | 50.48 | | |
| 5.Suprabenthos | 1.01 | 8.4 | 54.36 | | |
| 6.Polychaetes | 9.984 | 1.9 | 11.53 | | |
| 7.Commercial benthic invertebrates | 0.043 | 1.06 | 3.13 | 0.035 | |
| 8.Benthic Invertebrates | 79.763 | 1.06 | 3.13 | | 0.328 |
| 9.Shrimps | | 3.21 | 7.2 | 0.016 | 0.017 |
| 10.Norway lobster | 0.018 | 1.25 | 4.56 | 0.037 | |
| 11.Mantis shrimp | 0.015 | 1.5 | 4.56 | 0.072 | |
| 12.Crabs | 0.009 | 2.44 | 4.73 | 0.002 | 0.177 |
| 13.Benthic cephalopods | 0.068 | 2.96 | 5.3 | 0.154 | 0.002 |
| 14.Squids | 0.02 | 3.11 | 26.47 | 0.041 | |
| 15.Hake 1 | 0.06 | 1 | 4.24 | 0.113 | 0.07 |
| 16.Hake 2 | | 0.5 | 1.85 | | |
| 17.Other gadiformes | 0.029 | 1.59 | 4.37 | 0.025 | 0.083 |
| 18.Red mullets | 0.025 | 1.9 | 8.02 | 0.112 | |
| 19.Conger eel | 0.005 | 1.92 | 6.45 | | 0.008 |
| 20.Anglerfish | 0.006 | 1.04 | 4.58 | 0.007 | |
| 21.Flatfish | 0.009 | 1.43 | 9.83 | 0.04 | |
| 22.Turbot and brill | | 1.43 | 5.34 | 0.016 | |
| 23.Demersal sharks | 0.018 | 0.63 | 4.47 | 0.008 | |
| 24.Demersal skates | 0.003 | 1.11 | 7.08 | 0.002 | |
| 25.Demersal fish 1 | 0.056 | 2.4 | 7.68 | 0.055 | 0.051 |
| 26.Demersal fish 2 | | 2.4 | 5.68 | 0.016 | 0.001 |
| 27.Bentopelagic fish | | 1.07 | 7.99 | 0.002 | |
| 28.European Anchovy | 1.019 - 6.611 | 0.87 | 11.02 | 0.496 | 0.005 |
| 29.European Pilchard | 2.985 - 7.803 | 0.75 | 9.19 | 0.364 | 0.042 |
| 30.Small Pelagic Fish | 0.413 - 1.517 | 1.1 | 11.29 | 0.012 | 0.001 |
| 31.Horse Mackarel | 0.659 - 2.455 | 0.99 | 7.57 | 0.02 | 0.002 |
| 32.Mackarel | 0.452 - 1.683 | 0.99 | 6.09 | 0.017 | 0.008 |
| 33.Atlantic bonito | 0.3 | 0.39 | 4.54 | 0.018 | |
| 34.Large Pelagic Fish | 0.138 | 0.37 | 1.99 | 0.026 | |
| 35.Dolphins | 0.012 | 0.08 | 11.01 | | 0.0001 |
| 36.Loggerhead turtle | 0.032 | 0.17 | 2.54 | | 0.004 |
| 37.Sea birds | 0.001 | 4.61 | 69.34 | | |
| 38.Discard | 0.733 | | | | |
| 39.Detritus | 200 | | | | |



(a) Diet composition matrix



(b) Contaminant uptake from diet matrix

Figure A.7: Level plots of the diet composition matrix in the trophic network (a) and of the contaminant uptake rate from diet relative to the PCB bioaccumulation network. Darker cells indicate feeding links where the contribution of the prey in the diet/PCBs concentration of the predator is higher. Diet composition is taken from Coll et al. (2007), while the uptake rate of a predator j from a prey i , U_{ij} , is the contaminant flow from i to j scaled by the sum of the inflows of j .

Table A.6: Summary of the studies considered for the parametrisation of the PCBs bioaccumulation model. We report references; periods and areas of analysis; species considered; kinds of tissues sampled; units of measurement; and PCBs congeners detected.

| Source | Period of analysis | Area | Species | Tissue | Units | PCBs congeners |
|------------------------------------|--------------------|--------------------------------|--|--------|---------|---|
| (Perugini et al., 2004) | 2002 | Center | <i>M. galloprovincialis</i> , <i>N. norvegicus</i> , <i>M. barbatus</i> , <i>S. officinalis</i> , <i>E. fying squid</i> , <i>E. encrasicolus</i> , <i>S. pilchardus</i> , <i>S.r scombrus</i> | edible | ng/g ww | 28, 52, 101, 118, 138, 153, 180 |
| (Marcofrigi and Storelli, 2003) | 1997-1998 | Adriatic, Ionian Sea | <i>M. Merluccius</i> , <i>M. pouassou</i> , <i>P. bleinnoides</i> , <i>S. smaris</i> , <i>S. pilchardus</i> , <i>E. encrasicolus</i> , <i>L. caudatus</i> , <i>H. dactylopterus</i> , <i>L. budegassa</i> , <i>T. trachurus</i> , <i>A. rochei</i> , <i>Raje spp</i> , <i>P. glauca</i> , <i>S. acanthias</i> , <i>S. blainvillei</i> , <i>S. canicula</i> , <i>G. melastomus</i> , <i>C. gallina</i> , <i>A. tuberculata</i> , <i>E. siliqua</i> , <i>M. galloprovincialis</i> , <i>O. salatii</i> , <i>I. coindet</i> , <i>S. mantis</i> , <i>P. longirostris</i> , <i>A. antennatus</i> | muscle | ng/g ww | 8, 20, 28, 35, 52, 60, 77, 101, 105, 118, 126, 138, 153, 156, 169, 180, 209 |
| (Bayarri et al., 2001) | 1997-1998 | North, Center, South | <i>L. vulgaris</i> , <i>M. galloprovincialis</i> , <i>N. norvegicus</i> , <i>M. barbatus</i> , <i>C. gallina</i> | edible | ng/g ww | 28, 52, 101, 118, 138, 153, 180, 138, 163 |
| (Storelli et al., 2003) | 2001 | South | <i>C. conger</i> , <i>H. dactylopterus</i> , <i>L. budegassa</i> , <i>M. barbatus</i> , <i>S. flexuosa</i> , <i>P. bleinnoides</i> , <i>P. erythrinus</i> , <i>R. clavata</i> , <i>R. oxyrinchus</i> , <i>R. miraletus</i> , <i>S. pilchardus</i> , <i>M. merluccius</i> , <i>S. aurita</i> , <i>S. scombrus</i> , <i>T. trachurus</i> , <i>A. antennatus</i> , <i>N. norvegicus</i> , <i>P. longirostris</i> , <i>P. maritima</i> | muscle | pg/g ww | 126, 138, 153, 156, 169, 180, 209 |
| (Storelli et al., 2007) | 1999 | South | <i>A. anguilla</i> | muscle | ng/g ww | 52, 70, 77, 101, 105, 118, 126, 138, 153, 180 |
| (Storelli and Marcotrigiano, 2001) | 1999 | South | <i>C. granulatus</i> , <i>S. blainvillei</i> | muscle | ng/g ww | 8, 20, 28, 35, 52, 60, 77, 101, 105, 118, 126, 138, 153, 156, 169, 180, 209 |
| (Storelli et al., 2007) | 1999 | Adriatic, Ionian Sea | <i>C. caretta</i> | muscle | ng/g ww | 8, 20, 28, 35, 52, 60, 77, 101, 105, 118, 126, 138, 153, 156, 169, 180, 209 |
| (Corsolini et al., 2000) | 1994 | Adriatic, Baltic, Northern Sea | <i>C. caretta</i> | muscle | ng/g ww | 153, 137, 138, 180, 170, 194, 60, 118, 105, 156, 189, 77, 126, 169 |