Responses of summer phytoplankton biomass to changes in top-down forcing:

Insights from comparative modelling

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Running head: Phytoplankton responses to top-down forcing

1
Abstract

The present study describes the responses of summer phytoplankton biomass to changes in top-down forcing (expressed as zooplankton mortality) in three ecosystems (the North Sea, the Baltic Sea and the Nordic Seas) across different 3D ecosystem models. In each of the model set-ups, we applied the same changes in the magnitude of mortality (±20%) of the highest trophic zooplankton level (Z1). Model results showed overall dampened responses of phytoplankton relative to Z1 biomass. Phytoplankton responses varied depending on the food web structure and trophic coupling represented in the models. Hence, a priori model assumptions were found to influence cascades and pathways in model estimates and, thus, become highly relevant when examining ecosystem pressures such as fishing and climate change. Especially, the different roles and parameterizations of additional zooplankton groups grazed by Z1, and their importance for the outcome, emphasized the need for better calibration data. Spatial variability was high within each model indicating that physics (hydrodynamics and temperature) and nutrient dynamics also play vital roles for ecosystem responses to top-down effects. In conclusion, the model comparison indicated that changes in top-down forcing in combination with the modelled food-web structure affect summer phytoplankton biomass and, thereby, indirectly influence water quality of the systems.
1. Introduction

Overfishing, pollution or destruction of habitats combined with climate change impose pressures on marine food webs and it is challenging to predict how changes in the strength of these human-induced pressures will impact on the trophodynamic structure and function of ecosystems (Polis et al. 2000, Shurin et al. 2002, Heath et al. 2014). Top-down forcing is defined as the regulation of lower food-web components by an upper-level predator (Pace et al. 1999). Trophic cascades occur when pressures change the biomass of one trophic level and thereby the strength of the top-down forcing across more than one trophic link in a food web (Cury et al. 2003, Huse et al. 2012). The strong decline in populations of large top-predators observed in coastal and oceanic waters and corresponding changes in top-down forcing may have severe consequences for ecosystem function (Myers & Worm 2003, Scheffer et al. 2005). Examples of top-down controlled systems are the Black Sea, the Eastern Scotian shelf off Canada and the Baltic Sea, where overfishing of the top predators directly affected the whole food web from planktivorous fish to primary producers and resulted in higher summer phytoplankton biomass (Frank et al. 2005, Casini et al. 2008, Möllmann et al. 2008, Llope et al. 2011).

High phytoplankton biomass is normally a sign of eutrophication caused by nutrient enrichment (bottom-up control) and summer phytoplankton biomass is used as an indicator of water quality in the HELCOM Baltic Sea Action Plan (HELCOM 2013). Top-down forced trophic cascades may thereby contribute to the eutrophication status and work against the goal to achieve a good ecological status in coastal and open waters according to the EU Water Framework Directive (2000/60/EC) and the EU Marine Strategy Framework Directive (2008/56/EC). On the other hand, it has been suggested that changes in fishing pressure on selected species could in turn decrease summer phytoplankton biomass and improve water clarity, as seen in lakes and some coastal ecosystems (Carpenter et al. 1985, Hansson et al. 1998, Lindegren et al. 2010, Petersen et al. 2017). Hence, knowledge on trophodynamics is
important when formulating guidelines to sustainably manage fisheries as well as meet other management goals such as to maintain high water quality (Frank et al. 2007).

Responses to changes in top-down forcing often emerge as ‘skipped-level-transmission’, i.e. different directions of change between adjacent trophic levels (Casini et al. 2008, Heath et al. 2014). The strength of the response is often dampened by each trophic level due to various compensatory mechanisms that regulate internal food-web dynamics and structure (McCann et al. 1998a, Pace et al. 1999, Shurin et al. 2002, Andersen & Pedersen 2010). The compensatory mechanisms include replacement of the affected species, regulation through density-dependent grazing pressure and loss of energy due to respiration, cannibalism and other types of mortality (Mccann et al. 1998b, Andersen & Pedersen 2010). Trophic cascades are transitory and dynamic phenomenon and, hence, exhibit variation in their strength and duration both within and between systems, the latter due to ecosystem-specific differences in food web dynamics and structure (Cury et al. 2003).

Food web models have become an important tool in examining how reductions in specific predators or prey impact on other ecosystem components (Travers et al. 2007, Daewel et al. 2014, Peck et al. 2018). A general framework and theoretical description of the different types of food web responses that can be expected in relation to changes in trophodynamic controls was provided by Cury et al. (2003). Further, theoretical modelling has produced simple rules for how perturbations at upper trophic levels can affect the strength of potential trophic cascades within specific ecosystems (Mccann et al. 1998b, Leibold et al. 2004, Wollrab et al. 2012, Heath et al. 2014). Although these models take into account trophic complexity, they do not account for any local spatio-temporal variability of the ecosystem, which may affect predator-prey interactions and, hence, trophic cascades (Frank et al. 2007, Schulz et al. 2007, Travers & Shin 2010). To fully and more realistically resolve the emergence of trophic responses in the plankton community, spatially- and temporally-explicit models are required which include both
hydrodynamics and biogeochemical processes.

Ecosystem models of lower trophic levels depict nutrient cycling and dynamics of plankton functional types (PFTs) including primary producers and grazers with different life strategies and sizes (Blackford et al. 2004, Daewel et al. 2014). In these models, the zooplankton community ranges from unicellular fast-growing microorganisms to multicellular meso- and macrozooplankton (e.g. copepods, krill) with longer generation times, but the community is often reduced to one or a few zoo-PFTs (Maar et al. 2011, Butenschön et al. 2016). Mesozooplankton (carnivorous or omnivorous) represents, in most cases, the highest trophic level and mortality on this group represents a closure term for nutrient and carbon fluxes. A background mortality (encompassing natural mortality, predation, cannibalism, diseases, etc.) is often applied as a linear, quadratic or saturation function assuming that e.g. higher densities of zooplankton will lead to more or less strong habitat limitation effects, might attract potential predators or will increase the likelihood of infections (Edwards & Yool 2000, Fulton et al. 2003). Although different 3D ecosystem models may seem very similar, there can be important differences in their assumptions (e.g. food web structure, physiological rates, prey preferences, mortality terms) and underlying hydrodynamics, which may lead to different responses of the PFTs to changes in forcing (Fulton et al. 2003, Skogen & Moll 2005, Mitra & Davis 2010, Sailley et al. 2013). Most previous ecosystem model inter-comparisons have focused on changes in environmental drivers such as nutrient loads and climate change on water quality (e.g. Lenhart et al. 2010, Meier et al. 2012a, Skogen et al. 2014). A few studies have focused on the sensitivity of lower trophic levels to different formulations of predator-prey interactions, and the strength and complexity of zooplankton grazing dynamics (Anderson et al. 2013, Hashioka et al. 2013, Sailley et al. 2013, Le Quéré et al. 2016). These studies showed that food web dynamics, especially the predator-prey interactions, are very sensitive to the model formulations and gave different results of phytoplankton biomass within the same area. However, to our knowledge,
there has not been a comprehensive model inter-comparison study of lower trophic level responses to the same change of mortality at the highest zooplankton trophic level.

In a first step towards using ecosystem models to describe potential trophic cascades at the base of the food web induced by changes in top-down forcing, the present study applied the same zooplankton mortality scenarios across seven, previously validated 3D ecosystem models. The ecosystem models represented four types of food webs based on their trophic structure and interactions and covered three areas. The aim of the study was to predict the response of summer phytoplankton biomass to changes in top-down forcing i) among models within the same area (the North Sea or the Baltic Sea) and ii) across areas using the same model (i.e. the North Sea versus the Baltic Sea and the North Sea versus the Nordic Seas). The variability was expected to be high within areas due to differences in model formulation and across areas due to differences in ecosystem dynamics.

2. Methods and Material

2.1. Approach

We compared simulation results across seven different 3D models covering three different domains in the NE Atlantic Ocean (Figure 1) yielding 10 model set-ups in total (Table 1). The models considered in this study have been thoroughly described and validated in the list of papers given in Table 1. The data sources and time periods used for model validations are shown in Table 2. Hence, only the directly relevant features concerning the model food-web structure are outlined here (Figure 2). The models were set-up for a “Baseline” (corresponding to the published set-ups) and two “top-down” scenarios representing changes in the background mortality (model closure term) of the highest trophic level named ‘Z1’. The background mortality term encompasses a range of processes and was described either as: linear = $c \cdot Z1$, quadratic = $c \cdot Z1^2$, and saturation = $c \cdot Z1 \cdot Z1 / (Z1 + k)$ functions, where $c$ is the closure term.
constant and $k$ is the mortality half-saturation constant (Table 1). The closure term constant ($c$) was changed by +/- 20% in the P20 and M20 scenarios, respectively, which is within the natural variability of zooplankton mortality (Ji et al. 2013, Maar et al. 2014). In some models, other mortality terms such as cannibalism or death due to anoxia were explicitly described, but remained at their baseline values in the scenarios. The modelled period in Baseline and top-down scenarios covered a period of 3 years from 2003-2005 for most models except for HBM-ERGOM and POLCOMS-ERSEM using the years 2002-2004. The spin-up time varied from 2 to 10 years depending on the model.

2.2. Study areas

The three different domains covered by the models were: A) the North Sea (Delft3D-GEM, NORWECOM-NS, MIRO&CO, ECOSMO-NS, HBM-ERGOM-NS, POLCOMS-ERSEM), B) the Baltic Sea (MOM-ERGOM, ECOSMO-BS, HBM-ERGOM-BS) and C) the Nordic Seas (NORWECOM-NO) (Figure 1). Hence, the North Sea was covered by six models, the Baltic Sea by three models and the Nordic Seas by one model. The domains of the North Sea models were not identical, but outputs were estimated for the same area corresponding to the model domain of DELFT3D-GEM (-3.2-9.0°E, 49.3 -56.7°N) except for the MIRO&CO model, which only covered the southern part of the North Sea (extracted for 49.3-52.5°N). The model results for the Baltic Sea were extracted from the same area (11.0-30.3°E, 54.0-65.9°N). The Nordic Seas model domain covered the Greenland Sea, the Iceland Sea, the Norwegian Sea and the Barents Sea (-30.0-60.0°E, 61.3-80.0°N).

2.3. Model descriptions

For simplicity and ease of comparison among model responses to changes in top-down forcing, we only considered phyto- and zoo-PFTs and ignored grazing on bacteria and detritus in the analysis (Figure 2). The zoo-PFTs and phyto-PFTs were labelled as Z1-3 and P1-4, respectively, depending on the number of PFTs in each model. The Z1 group corresponds to the highest trophic level, and constitutes the group for
which the +/- 20% change in background mortality was applied in the “top-down” scenarios. Nutrient uptake by the phyto-PFTs was also simplified to competition for one resource (e.g. nitrogen or phosphorous) in the schematic representations in Figure 2. In all models, the diet of each zoo-PFT was determined dynamically by the relative abundances of its prey, but weighted with statically defined food preference factors (Table 3). In most models, there were no differences in the quality (C:N:P-ratio) of prey for zooplankton growth; in the Delft3D-GEM and POLCOMS-ERSEM, however, prey stoichiometry could vary with time, space and/or species, which could influence zooplankton growth. Complexity of the models was defined by i) the number of trophic links \((L)\) per number of PFTs \((L/S)\) and ii) connectance \((C=L/S^2)\) i.e. the ratio of trophic links over the number of possible links (Dunne et al. 2002). Among the models, four types of food webs were depicted based on the trophic structure and interactions (Figure 2, Table 1):

**Food web type 1**: One zoo-PFT (Z1) grazes on several phyto-PFTs with low \(L/S\) and low connectance in DELFT3D-GEM and MOM-ERGOM (Figure 2A).

In Delft3D-GEM (the North Sea), zooplankton (Z1) feeds on diatoms (P1), photoautotrophic flagellates (P2), the colony-forming *Phaeocystis* (P3) and dinoflagellates (P4) with different prey preferences (Table 3). The background mortality is a saturation function of the Z1 biomass.

In MOM-ERGOM (the Baltic Sea), zooplankton (Z1) feeds on diatoms (P1) and photoautotrophic flagellates (P2) with equal prey preferences and on diazotrophic cyanobacteria (P3) with lower preference (Table 3). The background mortality is a quadratic function of the Z1 biomass and a loss term due to hypoxia is expressed as increased respiration.

**Food web type 2**: Two zoo-PFTs and several phyto-PFTs interact with low \(L/S\) and low connectance as applied in the models NORWECOM (-NO, -NS) and MIRO&CO (Figure 2B).
In NORWECOM (the North Sea and the Nordic Seas), omnivorous mesozooplankton (Z1) feeds on diatoms (P1) and microzooplankton (Z2), while microzooplankton feeds on photoautotrophic flagellates (P2), all with equal prey preferences (Table 3). The background mortality is a saturation function of the Z1 biomass.

In MIRO&CO (the southern North Sea), omnivorous mesozooplankton (Z1) feeds on diatoms (P1) and microzooplankton (Z2), while microzooplankton feeds on photoautotrophic nanoflagellates (P2), all with equal prey preferences (Table 3). *Phaeocystis* colonies (P3) are not grazed, but can be disrupted into single edible cells (P2). The background mortality is a quadratic function of the Z1 biomass.

**Food web type 3**: Two zoo-PFTs and three phyto-PFTs interacts with intermediate L/S and high connectance in the models ECOSMO and HBM-ERGOM (Figure 2C).

In ECOSMO (the North Sea and Baltic Sea), omnivorous zooplankton feeds on diatoms (P1), photoautotrophic flagellates (P2), diazotrophic cyanobacteria (P3) and on herbivorous zooplankton (Z2), while Z2 feeds on all phyto-PFTs with different prey preferences (Table 3). The background mortality is a linear function of the Z1 biomass.

In HBM-ERGOM (the North Sea and the Baltic Sea), omnivorous mesozooplankton feeds on diatoms (P1), photoautotrophic flagellates (P2), diazotrophic cyanobacteria (P3) and microzooplankton (Z2), while Z2 grazes on the three phyto-PFTs. Prey preferences are lower for diazotrophic cyanobacteria and Z2 than for the other PFTs (Table 3). Background mortality is described as a saturation function of the Z1 biomass and mortality due to cannibalism and hypoxia was explicitly described for both zoo-PFTs.

**Food web type 4**: Three zoo-PFTs and four phyto-PFTS interact with high L/S and intermediate connectance in POLCOMS-ERSEM (Figure 2D).
POLCOMS-ERSEM (the North Sea) includes omnivores mesozooplankton (Z1), microzooplankton (Z2) and heterotrophic flagellates (Z3), diatoms (P1), microphytoplankton (P2), nanophytoplankton (P3) and picophytoplankton (P4) with complex predator-prey interactions and different prey preferences (Table 3, Figure 2D). The background mortality is a linear function of the Z1 biomass and cannibalism is explicitly resolved as part of the predation for all three zoo-FTPs.

2.5. Model output

Average plankton biomass was calculated from model outputs for the summer period (June to September) in the upper 50 m of the water column. The response of each PFT was estimated as the relative change (ΔC) in biomass between the mortality scenarios (SCE) and the Baseline (B) with respect to the Baseline value and normalized by the change in forcing (F=0.20) (Petersen et al. 2017):

$$\Delta C_{PFT} = \frac{(SCE-B)}{B \times F}$$  \hspace{1cm} (1)

An absolute value <1 indicates that the response is smaller than the impact, whereas an absolute value >1 means that the impact is amplified by the system. For the zooplankton group Z1, the relative change of biomass will inform on the direct effect of the change of the forcing (mortality of Z1). For other groups, these values will inform about the indirect effect of the forcing (fx change in grazing pressure from Z1) and the dampening role of the food web structure complexity. Total summer phytoplankton biomass (PHY) was estimated as the sum of all phyto-PFTs (P1+P2+P3+P4). In addition, for an inter-model and inter-region comparison, we aggregated the individual model values of $\Delta C_{PHY}$ for i) the same area over different models and ii) the same model over different areas. From that, we calculated the median $\Delta C_{PHY}$ (due to skewness of data) as a measure for the typical food web response and the data range (=maximum-minimum) as a measure of the variability of $\Delta C_{PHY}$ between individual models or regions.
In order to evaluate the major responses to changes in top-down forcing (Figure 2), we estimated the ratios of $\Delta Cs$ (from eq. 1) between each predator and its prey for all trophic links:

$$L_{p,Z} = \frac{\Delta C_{prey}}{\Delta C_{predator}}$$

(2)

We identified those direct links where the ratio was negative (i.e. opposite responses of predator and prey), which indicates a top-down forcing via skipped-level transmission (Casini et al. 2008, Heath et al. 2014). These links and their strengths are shown in Figure 2 for each model. The food web response was termed as ‘direct grazing’ if only 1 trophic link was involved, as a ‘trophic cascade’ if subsequent trophic links (e.g. Z1-Z2-P2) all responded with negative $L_{p,Z}$-ratios or as ‘combined’ if it exhibited both direct grazing and trophic cascades. Hence, food web type 1 models with one zoo-PFT (1 trophic link) can only show a direct grazing response, which in most cases would lead to a negative $L_{p,Z}$-ratio. However, it is possible that changes in grazing pressure on different phytoplankton groups competing for a limiting nutrient will favor certain phytoplankton groups and result in a positive $L_{p,Z}$-ratio for some (but not all) phyto-PFTs (Wollrab et al. 2012).

Finally, the trophic cascade (TC) -ratio was used to estimate the cascade strength of total phytoplankton biomass (PHY) in relation to changes in the highest zooplankton trophic level (Z1):

$$TC = \frac{\Delta C_{PHY}}{\Delta C_{Z1}}$$

(3)

based on the estimated $\Delta C_{PHY}$ and $\Delta C_{Z1}$ from eq. 1. When the TC-ratio is negative, Z1 and PHY change in the opposite direction (positive or negative) and vice versa. If the TC-ratio is close to one (absolute numbers), the cascade strength is strong with little or no dampening. If the TC-ratio is close to zero (i.e. low change of $\Delta C_{PHY}$), the cascade is dampened quickly, while a TC-ratio larger than one indicates that the cascade is amplified.
3. Results

3.1. Overall responses to top-down forcing

The impact of changes in top-down forcing on summer phytoplankton biomass and different PFTs was analyzed using a high diversity of ecosystem models with different representations of the structure and function of lower trophic levels. First, the highest trophic level, i.e. the zoo-PFT Z1 directly impacted by the change in forcing, showed expected responses in the opposite direction than the change in the background mortality across all models (Figure 3A). The global median $\Delta C_{Z1}$ value was -0.65 in P20 and 0.73 in M20 for all models and regions (Table 4). The absolute value of $\Delta C_{Z1}$ was less than 1 in most model results (14 out of 20) indicating an attenuation of the impact strength despite the direct impact on Z1. Amplification of the Z1-response was found in the type 3 models, i.e. ECOSMO-NS-BS (P20 and M20) and HBM-ERGOM-NS-BS (M20).

Second, the indirect effects of changes in top-down forcing were analyzed for trophic levels lower than Z1. Conversely to the direct responses, changes in total summer phytoplankton biomass ($\Delta C_{PHY}$) varied between the models for each scenario, both in direction (positive/negative) and magnitude (Figure 3B). The $\Delta C_{PHY}$ showed generally an opposite response direction to that of Z1 except for NORWECOM-NS-NO and ECOSMO-NS (P20). The median of $\Delta C_{PHY}$ was 0.10 (range: -0.36 to 0.64) in P20 and -0.11 (range: -0.35 to 0.34) in M20 across all models (Table 4). When comparing zooplankton and phytoplankton relative changes in biomass, the TC-ratio (eq. 3) was negative in most cases (i.e. opposite responses) except for the NORWECOM models (Figure 3C). ECOSMO-NS (P20) also gave a negative TC-ratio during the averaging procedure (months and grid cells) despite the negative signs of both $\Delta C_{Z1}$ and $\Delta C_{PHY}$ on total average (Table 4) probably due to non-linear responses in the food web. The median TC-ratio was -0.14 (range: -0.67 to 0.48) in P20 and -0.18 (range: -0.58 to 0.32) in M20 (Table 4) and overall
-0.17 in both scenarios. Hence, there was a general dampening of the signal from Z1 to phytoplankton biomass.

3.2. Inter-model comparison: The North Sea

3.2.1. Responses between models and food web types

The North Sea was covered by six models (DELFT3D, NORWECOM-NS, MIRO&CO, ECOSMO-NS, HBM-ERGOM-NS and POLCOMS-ERSEM) representing four different types of food webs with median $\Delta C_{PHY}$ of 0.06 and -0.15 in P20 and M20, respectively (Table 5). The inter-model variability of $\Delta C_{PHY}$ (range=1.00) was high in comparison with the other areas and the overall model variability (Table 5).

In the food web type 1 model (DELFT3D-GEM), there was only one zoo-PFT (Z1) grazing mainly on P1 (diatoms) or P4 (dinoflagellates) with negative $L_{p,z}$-ratios (Figure 2A, Table 4). The other two phyto-PFTs (P2 and P3) showed, on the other hand, a positive $L_{p,z}$-ratios, because they were influenced by the differentiated grazing pressure and competition for a limiting nutrient. The $\Delta C_{PHY}$ values had the same response direction as the North Sea median values, but were slightly higher with 0.23 and -0.25 in P20 and M20, respectively (Table 4, Figure 3A).

The food web type 2 models (NORWECOM-NS and MIRO&CO, Figure 2B) showed the strongest responses in terms of $\Delta C_{PHY}$ values and TC-ratios (Figure 3B-C). This food web type was characterized by two branches that were looped through the top consumer (Z1) and had a low connectance. Opposed to the other North Sea models, the TC-ratios in NORWECOM-NS were positive with 0.32 and 0.48 in P20 and M20, respectively (Table 4). The positive ratios could be explained by a strong trophic cascade in the Z1-Z2-P2 branch (causing the same response direction for Z1 and PHY) compared to a weaker grazing effect in the Z1-P1 branch, probably due to the high overlap of Z2 and P2 biomasses during summer (Figures 2B, 3, 4). The other type 2 model, MIRO&CO in the southern North Sea, showed on the contrary
the most negative TC-ratios among all the models with -0.58 and -0.67 in P20 and M20, respectively (Table 4). Here, the grazing response observed in the Z1-P1 branch was stronger than in the Z1-Z2-P2 branch (Figure 2B), probably because the P1 (diatom) biomass was higher than for P2 (flagellates) (Figure 4B). There was indication of a bottom-up effect (competition for nutrients) in the P2-Z2 branch, because both were affected in the same direction as Z1 (Figure 2B).

The food web type 3 models were characterized by having two zoo-PFTs and high connectance. ECOSMO-NS showed skewed responses of $\Delta C_{PHY}$ with negative values in both scenarios (-0.18 and -0.19 in P20 and M20, respectively) (Table 4). The Z1 PFT (omnivorous zooplankton) dominated the total summer zooplankton biomass (97%) and P2 (flagellates) the total summer phytoplankton biomass (96%) (Figure 4), which allowed different types of model responses to changes in top-down forcing. When the grazing impact by Z1 decreased (P20), this allowed Z2 (herbivorous zooplankton) to increase considerably ($\Delta C_{Z2}=5.76$) and initiated a strong trophic cascade in the Z1-Z2-P2 branch leading to negative $\Delta C_{PHY}$ values (same direction between Z1 and PHY). In the opposite situation (M20), the increased grazing pressure by Z1 reduced both Z2 ($\Delta C_{Z2}=-2.60$), P1 and P2 leading to negative $\Delta C_{PHY}$ values (opposite direction between Z1 and PHY) (Figure 3, Table 4). In the other food web type 3 model, HBM-ERGOM-NS, the resultant $\Delta C_{PHY}$s were 0.10 and -0.11 in P20 and M20, respectively, and hence less strong than for ECOSMO-NS (Table 4). The major grazer pathways (Z1 grazing on P2 and Z2) were similar to the M20 response in ECOSMO-NS (Figure 3C). The zoo-PFTs were also subject to cannibalism. The Z1 (mesozooplankton) biomass (69% of total zoo-PFTs) and P2 (flagellates) biomass (73% of total phyto-PFTs) also dominated the summer plankton biomasses, but to a lesser extent than in ECOSMO-NS (Figure 4).

In the food web type 4 model (POLCOMS-ERSEM), the predator-prey interactions were the most complex with several major grazer pathways in the scenarios compared to the other food webs (Figure
2D). There was a direct grazing response between Z1 and P2 in addition to trophic cascades in the branches Z1-Z2-P3 and Z1-Z2-Z3, both involving the Z2 group (Figure 3D). In addition, the three zoo-PFTs were subject to cannibalism. The different responses counteracted each other and overall resulted in a weak $\Delta C_{\text{PHY}}$ of 0.03 and -0.03 in P20 and M20, respectively (Table 4).

3.2.2. Spatial patterns of phytoplankton biomass changes in the North Sea

Spatial patterns of responses (magnitude and direction) in summer phytoplankton biomass ($\Delta C_{\text{PHY}}$) to changes in top-down forcing in the North Sea were highly variable between the models (Figure 5). In five of the six North Sea models (DELFT3D-GEM, NORWECOM-NS, MIRO&CO, HBM-ERGOM-NS and POLCOMS-ERSEM), 87-100% of the area showed the same response direction of $\Delta C_{\text{PHY}}$ in P20 and M20 (Table 6), but with opposite response direction for NORWECOM-NS (as explained before). ECOSMO-NS showed, on the other hand, the highest spatial heterogeneity of $\Delta C_{\text{PHY}}$ with only 67% and 76% in P20 and M20, respectively, of the areas showing the same (negative) response direction (Figure 5D). The high spatial variability in ECOSMO-NS was due to the competition between two pathways in the food web (Z1-P2 or Z1-Z2-P2), which had different outcomes in different areas. Hence, local areas could have either negative or positive values within the same model and scenario and, hence, not necessarily in the same response direction as the area-averaged result (Figure 3B-C). The strongest responses (either negative or positive) were observed in the German Bight and the English Channel (southeastern shallowest parts) in most models except for POLCOMS-ERSEM with lowest responses in this area.

3.3. Inter-model comparison: The Baltic Sea

The Baltic Sea was covered by three models (MOM-ERGOM, ECOSMO-BS and HBM-ERGOM-BS), which represented two different types of food webs (1 and 3) with $\Delta C_{\text{PHY}}$ median of 0.11 and -0.11 in P20 and M20, respectively (Table 5). Despite their differences, the three models provided similar results of
zooplankton and total phytoplankton changes in the Baltic Sea (Figure 3). The variability of $\Delta C_{PHY}$ was relatively low (range=0.33) compared to the North Sea and the overall model variability (Table 5). The overall TC-ratios were negative (from -0.09 to -0.19), i.e. the opposite response between Z1 and phytoplankton biomass (Table 4). The direct grazing response by Z1 on different phytoplankton prey dominated the models, whereas the trophic cascade through Z2 (food web type 3 models) was weaker (Figures 2A and C). Spatially, the responses were in the same direction in MOM-ERGOM and HBM-ERGOM-BS, whereas in ECOSMO-BS the response was more diverse (Table 6, Figure 6). However, in all three models, the highest changes ($\Delta C_{PHY}$) were estimated in the high productive coastal and shallow areas and in the same response direction (Figure 6).

3.4. Inter-regional model comparison

Three of the models covered two areas; NORWECOM-NS-NO, ECOSMO-NS-BS and HBM-ERGOM-NS-BS. The internal model variability expressed as the range of $\Delta C_{PHY}$ within each model (including the two areas) was highest for NORWECOM with the range= 0.70 followed by ECOSMO with the range= 0.38 and lowest for HBM-ERGOM with the range= 0.21 (Table 5). NORWECOM exhibited four to five times stronger responses of $\Delta C_{PHY}$ in the North Sea than for the Nordic Seas (Figures 3B, C). The inter-regional variability of $\Delta C_{PHY}$ in NORWECOM was lower than the inter-model variability of the North Sea models, but higher than for the Baltic Sea models (Table 5). In ECOSMO, the internal variability of $\Delta C_{PHY}$ was higher than for the Baltic Sea models, but lower than for the North Sea models (Figure 3, Table 4). In HBM-ERGOM, variability of $\Delta C_{PHY}$ was generally lower than for the Baltic Sea and North Sea models (Figure 3B, C).
4. Discussion

4.1. Top-down mediated trophic cascades

The present study provides new knowledge on the role of top-down mediated trophic cascades and its parameterization in lower trophic level ecosystem models by applying the same top-down scenarios to a wide range of 3D dynamic ecosystem models. The top-down scenarios were expressed as a change in the background mortality of the highest trophic level, Z1, which showed opposite responses to the change in forcing (Figure 3A). In most cases, there was an attenuation of the response by Z1 due to a dilution effect at the open boundaries, other mortality terms (e.g. anoxia, cannibalism) or complex food web interactions (e.g. bottom-up effects) counterbalancing the top-down effects (Wollrab et al. 2012). Amplification of the Z1-response was found in the type 3 models (i.e. ECOSMO and HBM-ERGOM) probably due to positive feedback mechanisms from the trophic cascades in the food web (Wollrab & Diehl 2015). Overall, there was a general dampening of the signal from the highest trophic level (Z1) to phytoplankton biomass (i.e. an absolute TC-ratio less than one, Figure 3C), which is a common feature of aquatic food webs due to internal trophic interactions (Shurin et al. 2002, Andersen & Pedersen 2010, Heath et al. 2014). However, phytoplankton responses varied in both magnitude and direction (positive/negative) relative to the change in Z1 depending on the food web structure and trophic coupling represented in the models (Figures 2, 3B).

The model results emphasized that the second zooplankton group (Z2) played an important role as mediator of trophic cascades. In most cases, the Z2 exhibited opposite grazing pressures on the phyto-PFTs compared to Z1 (Figure 2, Table 4) and thereby dampened the total phytoplankton biomass responses to changes in Z1 (i.e. less negative TC-ratio). In NORWECOM, the trophic cascades in the Z1-Z2-P2 branch even overruled the direct grazing pressure by Z1 on phytoplankton biomass and caused a positive TC-ratio. The Z2-PFT covers a wide range of species with different sizes, growth rates, prey
preference, mixotrophy and feeding strategies (Hansen 1991), which makes it difficult to define them as a functional group and to parameterize the general physiological processes (Anderson et al. 2013, Sailley et al. 2013). According to our model results from NORWECOM, MIRO&CO and HBM-ERGOM, Z2 had a biomass similar to Z1 (31-77% of total), whereas in ECOSMO, Z2 comprised 3% of the zooplankton biomass (Figure 4A). However, the Z2-PFT plays different roles in different models as e.g. herbivorous zooplankton in ECOSMO, ciliates and heterotrophic dinoflagellates in HBM-ERGOM and a broader range of microzooplankton in NORWECOM. POLCOMS-ERSEM included both microzooplankton (Z2) and heterotrophic flagellates (Z3), comprising 16% and 28%, respectively, of total zooplankton biomass (Figure 4A). Consequently, a direct comparison of Z2 between models is complicated by the lack of common metrics (Sailley et al. 2013). There is, to our knowledge, no coherent data set on microzooplankton (Z2) for the North Sea and the Baltic Sea (Quéré et al. 2005, Bils et al. 2017). An accurate model parameterization of this PFT is challenging without the necessary calibration data. This issue is reflected in the relatively few attempts to validate the biomass of different zoo-PFTs based on available national monitoring-, research- and ‘other’ data sets (Table 2). For the Z1-PFT (often assumed to represent copepods), more data sets from e.g. the Continuous Plankton Recorder (Pitois & Fox 2006), national monitoring data and research data were available allowing a validation of Z1 biomass in most models (Table 2).

4.2. Effects on summer phytoplankton biomass

The change in summer phytoplankton biomass showed overall opposite responses relative to the change of Z1. Hence, a higher mortality of Z1 (scenario P20) was shown to increase the summer phytoplankton biomass and thereby affect the water quality negatively in marine waters as suggested by previous studies (Frank et al. 2005, Casini et al. 2008, Llope et al. 2011). In the North Sea, a previous
model study using size-selective predation by Atlantic herring (*Clupea harengus*), four zoo-PFTs and two phyto-PFTs also found that top-down control should be present in the system (Koslow 1983). However, analysis of field data from 50 years showed no sign of top-down forced trophic cascades affecting the phytoplankton despite intense fishing activities in the North Sea (Koslow 1983, Reid et al. 2000). These studies suggested that trophic cascades in the North Sea disappeared due to ecosystem-wide fishing on most species, that zooplankton was more food limited than predation limited or that complex food web interactions dampened the response (Koslow 1983, Reid et al. 2000, Andersen & Pedersen 2010). For the Baltic Sea, the scenarios results in P20 (decrease of Z1) are consistent with field data over a 30-year period showing that overfishing of cod population directly affected its main prey, European sprat (*Sprattus sprattus*), and indirectly caused a reduction in summer zooplankton and higher summer phytoplankton biomass (Casini et al. 2008, Möllmann et al. 2008). The responses were analyzed with respect to the summer period, but will probably be different other times of the year due the seasonal plankton community succession (*e.g.* the spring diatom bloom) and change in environmental conditions (*e.g.* light, nutrient levels).

4.3. Inter-model comparison: the North Sea

The highest inter-model variability was found for the North Sea covered by six models (Table 5). The inter-model median for the North Sea predicted that changes in top-down forcing could affect the summer phytoplankton biomass although the responses varied in magnitude and direction both spatially and between models. The high variability in model responses was mainly due to the food web type 2 models (NORWECOM-NS and MIRO&CO, Figure 2B), which showed the strongest responses in terms of $\Delta C_{phy}$ values and $TC$-ratios (Figures 3B-C). These models with two branches were very sensitive to top-down forcing, because it was possible for one of the two branches to dominate the response depending on the spatial-seasonal overlap of predator-prey distributions. A dominant trophic cascade
response in one of the branches is consistent with theoretical models for an odd-length food chain (McCann et al. 1998a, Cury et al. 2003, Wollrab et al. 2012, Heath et al. 2014). In the food web type 1 model DELFT3D-GEM, the differentiated grazing pressure and competitive ability for a shared nutrient favored some phyto-PFTs on behalf of others in agreement with theoretical models (Wollrab et al. 2012). In food web type 3 models, ECOSMO-NS and HBM-ERGOM-NS, the different model parameterizations of grazing interactions (e.g. grazing rates, prey preferences and cannibalism) resulted in a different sensitivity of the lower trophic levels to changes in top-down forcing in P20, whereas the responses in M20 were more similar. The missing trophic cascade in HBM-ERGOM-NS was due to internal attenuation of the signal from top-down forcing by the prescribed cannibalism within Z2 (not present in ECOSMO-NS) indicated by the lower $\Delta C_{Z2}$ compared to ECOSMO-NS (Table 4). It was previously demonstrated that inclusion of zooplankton interference (e.g. cannibalism) weakened trophic interactions in a theoretical food web model and better matched the patterns found in nature (Mccann et al. 1998b). Hence, the food web type 3 responses were more complex than the more rigid food web types 1 and 2 models and therefore more difficult to predict in relation to changes in top-down forcing.

In the food web type 4 model (POLCOMS-ERSEM), the more complex and compensating food web interactions dampened the response to changes in top-down forcing. This is in agreement with previous studies, which showed that increased complexity in model structures stabilized the system with a higher resistance to perturbations (Mccann et al. 1998b, Vallina & Le Quéré 2011, Vallina et al. 2017).

In top-down scenarios, the skipped-level-transmission complicates the prediction of responses at various trophic levels and the choice of model parameters and food web structure becomes even more important and should be appropriate for the study area. In comparison, ecosystem models are expected to give more similar results in bottom-up scenarios (e.g. changes in nutrient inputs), because of the direct link between nutrients and phytoplankton and responses typically will be in the same direction as the forcing (Cury et al. 2003, Heath et al. 2014). In addition to the food web definition, additional model
components, such as the choice of nutrient cycles, remineralization rates and benthic-pelagic coupling would be indirectly affected by changes in the Z1 mortality and additionally lead to response variability among different ecosystem models. Hence, a priori assumptions on food web structure and parameterization will influence cascades and pathways in model estimates and, thus, become highly relevant when examining ecosystem pressures such as fishing and climate change.

4.4. Spatial patterns

Spatial variability of the phytoplankton responses was generally high in the North Sea and the Baltic Sea indicating that not only the specification of trophic links, but also temperature and nutrient dynamics play vital roles for ecosystem response to top-down effects. Strong horizontal gradients of temperatures and nutrient concentrations are typically found at the interface between coastal and offshore areas. Especially in the North Sea, bathymetric features and strong tides result in the formation of mixed, stratified and frontal regions that are characterized by very different hydrographic features (Otto et al. 1990). Higher temperatures can strengthen trophic interactions since zooplankton respiration increases faster with warming than primary production (Scheffer et al. 2001, Maar & Hansen 2011, Svensson et al. 2017). Likewise, higher nutrient concentrations may strengthen or weaken different grazer pathways and thereby interact with the top-down forced trophic cascades (Wollrab & Diehl 2015, Petersen et al. 2017). Furthermore, spatial variations in top-down responses are also caused by spatial patterns in background mortality in the field (Fulton et al. 2003, Travers et al. 2009, Maar et al. 2014), a feature that could not be investigated by our experimental set-up. These diverse spatial responses make it difficult to support a coherent management plan with respect to sustainable fishing and water quality.

4.5. Inter-regional model comparisons
Strongest responses of NORWECOM were found in the shallow, well-mixed and highly productive southern North Sea (Figure 5C), whereas the phytoplankton responses were lower and more homogenously distributed in the Nordic Seas (Figure 6D). This difference could be related to the higher nutrient inputs and higher temperatures in the North Sea that can strengthen trophic interactions compared to the more nutrient limited, colder and less productive Nordic Seas (Frank et al. 2007, Wollrab & Diehl 2015, Svensson et al. 2017). Hence, the higher variability of the North Sea models and partly the Baltic Sea models compared to the internal model variability still indicates that a priori assumptions in the food web structure and dynamics will influence trophic cascades and pathways arising from the same change in top-down forcing (Sailley et al. 2013, Heath et al. 2014). In principle, the model response of a single model can be highly variable in different regions, indicating that ecosystem functioning and hydrographic characteristics, and not only model formulations, were driving the responses to top-down forcing.

4.6. Conclusions

There was a high variability in the responses across models, especially in the North Sea, due to the different food web structures and trophic couplings represented in the models. Especially the role of the zoo-PFTs seems important for the resulting model response to top-down forcing and highlights a challenge in ecosystem modelling. The model-dependent responses to the same forcing demonstrate the uncertainty that arises from simplifying real-world ecosystems into numerically tractable model systems. This needs to be taken into account when confronting models tuned to present-day dynamics using what-if scenarios, e.g. by discussing their outcome as a possible, rather than a predicted, reaction of the ecosystem. In the present state of model development, an ensemble of model simulations seems as a reasonable approach to reduce inherent uncertainty in model estimates and provide weight of evidence (Lenhart et al. 2010, Meier et al. 2012b, Queiros et al. 2016, Yun et al. 2017). The behavior of
the relatively simple ecosystem models used here, should be compared to potentially more complex behavior and trophic responses of trait-based models, which can include many more groups (or a blending across groups in terms of size-spectrum models) to better understand how changes in top-down forcing may cascade through the plankton community. For a better representation of trophic cascades, the ecosystem models could be coupled to higher trophic level models (e.g. fish production models) or provide input to ‘End-to-End’ models considering food web interactions for all trophic levels including human pressures (Fulton 2010, Shin et al. 2010, Utne et al. 2012).
Acknowledgements

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References


Shin YJ, Travers M, Maury O (2010) Coupling low and high trophic levels models: Towards a pathways-orientated approach for end-to-end models. Prog Oceanogr 84:105-112


Table 1. Overview of the applied models and their domains, food web types (see Figure 2), number of zoo-PFTs, number of total PFTs (S) i.e. not including resources, number of trophic links (\(L\)), links per PFT (\(L/S\)), connectance (\(C=L/S^2\)), mesozooplankton background mortality function and model references. The model domains are the North Sea (NS), the Baltic Sea (BS) and the Nordic Seas (NO).

<table>
<thead>
<tr>
<th>Model name</th>
<th>Model domain</th>
<th>Food web type</th>
<th>Zoo-PFTs</th>
<th>S</th>
<th>L</th>
<th>L/S</th>
<th>L/S^2</th>
<th>Background mortality</th>
<th>References</th>
</tr>
</thead>
<tbody>
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<td>NS</td>
<td>1</td>
<td>1</td>
<td>5</td>
<td>4</td>
<td>0.80</td>
<td>0.16</td>
<td>saturation</td>
<td>(Los et al. 2008, Blauw et al. 2009, Los &amp; Blaas 2010)</td>
</tr>
<tr>
<td>MIRO&amp;CO</td>
<td>S. NS</td>
<td>2</td>
<td>2</td>
<td>5</td>
<td>4</td>
<td>0.80</td>
<td>0.16</td>
<td>quadratic</td>
<td>(Lancelot et al. 2005, Lacroix et al. 2007)</td>
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<td>2</td>
<td>4</td>
<td>3</td>
<td>0.75</td>
<td>0.19</td>
<td>saturation</td>
<td>(Aksnes et al. 1995, Skogen et al. 1995, Skogen et al. 2007, Skogen &amp; Mathisen 2009, Hjøllo et al. 2012)</td>
</tr>
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<td>2</td>
<td>5</td>
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<td>1.40</td>
<td>0.28</td>
<td>linear</td>
<td>(Schrum et al. 2006, Daewel &amp; Schrum 2013, Daewel et al. 2015)</td>
</tr>
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<td>NS, BS</td>
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<td>2</td>
<td>5</td>
<td>7</td>
<td>1.40</td>
<td>0.28</td>
<td>saturation</td>
<td>(Maar et al. 2011, 2014, 2016, Petersen et al. 2017)</td>
</tr>
<tr>
<td>MOM-ERGOM</td>
<td>BS</td>
<td>1</td>
<td>1</td>
<td>4</td>
<td>3</td>
<td>0.75</td>
<td>0.19</td>
<td>quadratic</td>
<td>(Neumann et al. 2002, 2015, 2017)</td>
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Table 2. Data sources used for model validation of nutrient concentrations, Chl $\alpha$ concentration, primary production, biomass of different zoo-PFT’s and Z1 biomass. I=ICES data, H=HELCOM data, N=National monitoring data, C=Continuous Plankton Recorder (CPR) data, W=World Ocean Atlas, R= research projects, RS = remote sensing data, L=literature values and O=other data (e.g. PhD project). The last column shows the validation period for the different variables. Validation references are shown in Table 1.

<table>
<thead>
<tr>
<th>Model name</th>
<th>Nutrients$^a$</th>
<th>Phyto-PFT$^b$</th>
<th>Chl $\alpha$$^c$</th>
<th>Primary production$^d$</th>
<th>Zoo-PFT$^e$</th>
<th>Z1 biomass$^f$</th>
<th>Validation period</th>
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<tr>
<td>DELFT3D-GEM</td>
<td>N, R</td>
<td>N, R</td>
<td>N, R, RS</td>
<td>-</td>
<td>Not relevant</td>
<td>L</td>
<td>a,c) 1975-2012 b) 1998 f) 2008</td>
</tr>
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</table>
Table 3. Prey preferences of the different zoo-PFTs in the food web models (see Figure 2). In some models, the zoo-PFTs are subject to cannibalism.

<table>
<thead>
<tr>
<th>Predator Type</th>
<th>P1</th>
<th>P2</th>
<th>P3</th>
<th>P4</th>
<th>Z1</th>
<th>Z2</th>
<th>Z3</th>
<th>P1</th>
<th>P2</th>
<th>P3</th>
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<th>P3</th>
<th>P4</th>
<th>Z3</th>
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<td>0.30</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
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<td>-</td>
<td>-</td>
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<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
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<td>-</td>
<td>-</td>
<td>-</td>
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<td>-</td>
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<td>-</td>
<td>0.00</td>
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<tr>
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<td>-</td>
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<td>1.00</td>
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<td>0.00</td>
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<td>-</td>
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<td>0.15</td>
<td>-</td>
<td>0.25</td>
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<td>0.00</td>
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<td>0.40</td>
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<td>0.40</td>
<td>-</td>
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Table 4. The ΔC for each PFT and TC-ratios in scenarios P20 (top) and M20 (bottom) averaged from June to September in the upper 50 m and the median for all models.

<table>
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<tr>
<th>P20 scenario</th>
<th>Domain</th>
<th>Food web type</th>
<th>ΔC_{C1}</th>
<th>ΔC_{C2}</th>
<th>ΔC_{C3}</th>
<th>ΔC_{ZOO}</th>
<th>ΔC_{P1}</th>
<th>ΔC_{P2}</th>
<th>ΔC_{P3}</th>
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<td>-</td>
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<td>-</td>
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<td>3.90</td>
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<td>-</td>
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<td>-</td>
<td>-0.36</td>
<td>0.48</td>
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<th>Domain</th>
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<th>ΔC_{C1}</th>
<th>ΔC_{C2}</th>
<th>ΔC_{C3}</th>
<th>ΔC_{ZOO}</th>
<th>ΔC_{P1}</th>
<th>ΔC_{P2}</th>
<th>ΔC_{P3}</th>
<th>ΔC_{P4}</th>
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<th>TC-ratio</th>
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Table 5. Aggregated $\Delta C_{PHY}$ results. The number ($N$) of all results (P20 and M20), the number of models, the median for P20 and M20 and the range=maximum-minimum values for both P20 and M20 for i) each model area and ii) for the models covering two areas.

<table>
<thead>
<tr>
<th>Area</th>
<th>$N$</th>
<th>Number of models</th>
<th>Median P20</th>
<th>Median M20</th>
<th>P20+M20 range</th>
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<tr>
<td>North Sea</td>
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Table 6. The % of grid cells in the model domains with negative values of $\Delta C_{PHY}$ and TC-ratio.

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<th>Model</th>
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<th>M20 $\Delta C_{PHY}$</th>
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<th>M20 TC-ratio</th>
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<td>100</td>
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