

Nearshore Macroalgae Cultivation for Carbon Sequestration by Biomass Harvesting: Evaluating Potential and Impacts with An Earth System Model

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Key Points:

- Offshore macroalgae cultivation for CDR has a global potential of gigatonnes scale.
- Partition of marine net primary production shifts from phytoplankton to macroalgae due to shading and nutrient robbing.
- Open ocean net primary production reduces the oxygen deficit zones.

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Abstract

This study introduces an ocean-based carbon dioxide removal (CDR) approach: Nearshore Macroalgae Aquaculture for Carbon Sequestration (N-MACS). By cultivating macroalgae in nearshore ocean surface areas, N-MACS aims to sequester CO₂ with subsequent carbon storage. Utilizing an Earth System Model with intermediate complexity (EMIC), we explore the CDR potential of N-MACS alongside its impacts on the global carbon cycle, marine biogeochemistry and marine ecosystems. Our investigations unveil that coastal N-MACS could potentially sequester 0.7 to 1.1 GtC yr⁻¹. However, it also significantly suppresses marine phytoplankton net primary productivity because of nutrient removal and canopy shading, counteracting approximately 30% of the N-MACS CDR capacity. This suppression of surface NPP, in turn, reduces carbon export out of the euphotic zone to the ocean interior, leading to elevated dissolved oxygen levels and diminished denitrification in present-day oxygen minimum zones. Effects due to harvesting-induced phosphorus removal continue for centuries even beyond the cessation of N-MACS.

Plain Language Summary

Our study explores the Nearshore Macroalgae Aquaculture for Carbon Sequestration (N-MACS) as a potential marine carbon dioxide removal strategy. This approach uses ocean-based seaweed farming to capture carbon dioxide—the main greenhouse gas causing global warming—and permanently stores it post harvesting through biomass processing and carbon storage. Our simulations indicate that N-MACS has the potential to remove substantial quantities of carbon dioxide every year. Nonetheless, harvesting will also remove oceanic nutrients and decrease open ocean primary production. At the same time, N-MACS can relieve the oxygen scarcity and mitigate surface ocean acidification. Those impacts on the oceanic ecosystem and marine biogeochemistry could potentially persist for centuries, upon the cessation of N-MACS.

1 Introduction

The IPCC's Sixth Assessment Report (IPCC (2022)) stipulates global net-zero CO₂ emissions by the early 2050s to restrict global warming to 1.5°C, recognizing Carbon Dioxide Removal (CDR) as essential to counterbalance residual emissions. Ocean-based CDR approaches are gaining traction due to the ocean's inherent carbon sequestration capacity (IPCC, 2022; Keller et al., 2021; GESAMP, 2019). As the Earth's largest dynamic carbon reservoir (Falkowski et al., 2000; Sarmiento & Gruber, 2013), the ocean's expanse and natural carbon absorption capacity, combined with measures like ocean fertilization, ocean alkalinity enhancement, can substantially augment carbon sequestration efforts (Buesseler et al., 2004; Bach et al., 2019).

Macroalgae offer an avenue for ocean-based CDR due to their notable net primary production rates and high carbon-to-nutrient ratios, facilitating effective carbon sequestration (N'Yeurt et al., 2012; Fernand et al., 2017; Gao et al., 2022). The global potential carbon export by macroalgae has been estimated as 1.4 GtC per year (Krause-Jensen & Duarte, 2016; Ortega et al., 2019; Barrón & Duarte, 2015). Cultivation technologies for macroalgae are well-established (e.g., Buck and Buchholz (2004); Goecke et al. (2020); Zhang et al. (2016)), with a global harvest reaching 34.7 million tonnes wet weight (WW) in 2019 (FAO, 2018; Cai et al., 2021). Macroalgae cultivation for ocean-CDR has been considered recently (Wu et al., 2023; Fernand et al., 2017). Based on geographic location, macroalgae-based CDR can be categorized into two categories: open-ocean cultivation with deep-ocean carbon storage (Wu et al., 2023; Bach et al., 2021), and nearshore cultivation for harvesting, followed by subsequent carbon storage achieved outside of the ocean such as biochar and Bioenergy with Carbon Capture and Storage (BECCS, Roberts

et al. (2015); Bird et al. (2011); Fernand et al. (2017); Gattuso et al. (2021); Capron et al. (2020); Borchers et al. (2022); Chen et al. (2015)).

Prior to the large-scale implementation of ocean-based CDR strategies, comprehensive evaluations are essential to understand their potential and impacts on the marine environment (IPCC, 2022; Gattuso et al., 2021). Particularly, numerical simulations with Earth system models are pivotal as they, in contrast to field experiments pose, have no direct environmental impact (Oschlies et al., 2010; Keller et al., 2014; Keller, Lenton, Scott, et al., 2018; Siegel et al., 2021). Several modelling studies have examined macroalgae-based CDR strategies, revealing CDR capacities ranging from Mega (10^6) to Giga (10^9) tonnes depending on location and species. These studies, referenced as Wu et al. (2023); Bach et al. (2019) for open-ocean and Arzeno-Soltero et al. (2023); Berger et al. (2023) for nearshore areas, also underscore the constraints posed by marine physical and biogeochemical feedbacks on CDR capacity and efficiency. Furthermore, they highlight the potentially significant impacts on the global carbon cycle, marine biogeochemistry, and ecosystems through the alteration of ocean nutrient distributions and primary production patterns.

Here we evaluate ‘Nearshore Macroalgae Aquaculture for Carbon Sequestration’ (hereinafter N-MACS), operating under the assumption that the harvested carbon content will be sequestered from atmosphere and hence achieving CDR. The evaluation employs an Earth System Model of intermediate complexity, encompassing an explicit macroalgae component, to rigorously assess implications and carbon sequestration efficacy of N-MACS from 2020 to 3000, with N-MACS deployment from 2020 to 2100. Our objectives are to: a) examine the idealised large-scale CDR potential of N-MACS, and b) evaluate its effects on the global carbon cycle and marine biogeochemistry, including termination effects and millennial long-term effects.

2 Methods

We employ the University of Victoria Earth System Climate Model version 2.9 (UVic; Keller et al. (2012); Weaver et al. (2001)), an intermediate complexity Earth system model coupling a three-dimensional ocean circulation model (Pacanowski, 1996) including a dynamic thermodynamic sea ice module (Bitz & Lipscomb, 1999), a terrestrial model (Meissner et al., 2003; Weaver et al., 2001) and a one-layer atmospheric energy-moisture model (Fanning & Weaver, 1996). The horizontal resolution is 3.6° longitude \times 1.8° latitude, and the ocean component has 19 vertical layers with thicknesses ranging from 50 m near the surface to 500 m in the deep ocean. The ocean biogeochemistry module includes nutrients (nitrogen and phosphate), one general phytoplankton type, and one diazotrophic phytoplankton (i.e., nitrogen fixers), one general macroalgae (see below section), one type of zooplankton, dissolved inorganic carbon, oxygen, and total alkalinity (Keller et al., 2012; Eby et al., 2013).

Upon spinning up the model under pre-industrial conditions, we employed CMIP5 forcing data for the historical period (Eby et al., 2013). From 2005 to 2100, we aligned the inputs of CO₂ emissions, land-use changes, volcanic radiative forcing, and sulfate aerosols with the RCP4.5 scenario. For the period post-2300, CO₂ emissions are projected to decline linearly, reaching zero by 3000, with other forcings maintained at constant levels. RCP4.5 is a moderate emissions trajectory with a radiative forcing of 4.5 W/m^2 by 2100 (Thomson et al., 2011; Meinshausen et al., 2011).

N-MACS is an extension of the Macroalgae Open-ocean Mariculture and Sinking (MOS) framework developed by (Wu et al., 2023), featuring an idealized generic model of the Phaeophyceae (brown algae) *Sacharina* integrated with UVic. Macroalgae growth is controlled by multiple limiting factors (erosion, nutrient availability, light, and temperature) with a fixed C:N:P stoichiometric molar ratio of 400:20:1. Initial seed biomass

is deployed in each surface ocean grid box with adequate nutrients to be converted into seed biomass. The initial plantlet biomass in each N-MACS grid cell is equivalent to 0.02 mmol N m⁻³, sourced directly from the grid box's inorganic N, P, and C pools without extra nutrient or carbon input. A constant maximum biomass yield of 3,300 tDW km⁻² is set, focusing on large-scale impacts rather than optimizing farming strategies. Once biomass in a grid cell reaches this limit, macroalgae growth halts until end-of-season harvesting. In temperate zones, seeding starts on May 1st and harvesting occurs on October 31st in the northern hemisphere, while in the southern hemisphere, seeding begins on November 1 with harvesting on April 30, aligning with macroalgae growth phases. The model annually selects grid boxes with ample nutrients for reseeded, implying no further reseeding post-harvest in nutrient-depleted regions (detailed in Section 3.1, Wu et al. (2023)). Additionally, surface layer macroalgae create canopy shading effects on phytoplankton communities. Potential grazers like amphipods and gastropods (Jacobucci et al., 2008; Chikaraishi et al., 2007) are modeled within the UVic's zooplankton compartment (Keller et al., 2012). Further macroalgae model specifics, including parameters, functions, and cultivation strategies, are delineated in Wu et al. (2023, Sect. 2).

2.1 Experimental design

Our study contains a control run (Ctrl_RCP4.5) and two N-MACS simulations: the standard N-MACS simulation with all growth constraints, and a sensitivity simulation (No_Temp) with temperature constraint removed to examine the uncertainty in temperature-dependant growth rate in the modeled macroalgae. In both N-MACS simulations, macroalgae farms are limited to ocean surface zones directly along coasts between 60°S and 60°N, with grid boxes 200 to 400 km wide, aligning with Exclusive Economic Zones (EEZs) extending to 200 nautical miles from sovereign state coasts (Froehlich et al., 2019; Feng et al., 2017). It's presumed that all macroalgae production is promptly harvested post cultivation for biochar conversion or BECCS feedstock on land, indicating permanent carbon sequestration from the biomass with no nutrient return to the ocean. Meanwhile, natural macroalgae habitats are globally distributed along coastlines with species exhibiting varied temperature sensitivities (Duarte et al., 2022). The No_Temp simulation investigates the theoretical maximum coastal macroalgae biomass production with species optimally adapted to local temperatures. N-MACS CDR capacity is defined as the total carbon in harvested biomass, while its CDR efficacy is defined by the changes in combined oceanic and macroalgae carbon reservoir relative to the harvested macroalgal biomass carbon content. Our focus is on the cultivation process outcomes, excluding possible carbon leakages in post-harvest CDR applications like biochar or BECCS (Chen et al., 2015; Fernand et al., 2017; Bird et al., 2011).

3 Results & Discussions

3.1 Macroalgae model validation

The employed macroalgae model was validated against literature data and used in idealized open-ocean cultivation simulations by Wu et al. (2023). Given the notable nutrient availability differences between nearshore regions and open oceans, we compare the productivity of simulated nearshore macroalgae with relevant observational and modeling data.

Fig.1 illustrates the N-MACS distribution and its mean annual biomass yield from 2020 to 2100. Simulations indicate a total N-MACS footprint of about 24 million km², with 14 to 15 million km² yielding significant productivity (over 100 tonnes DW km⁻²yr⁻¹; Tab.1). These values are lower than other model-based estimates ranging from 48 to 100 million km² (Froehlich et al., 2019; Lehahn et al., 2016; Berger et al., 2023), hence presenting a more conservative N-MACS productivity. The reduced macroalgae farming areas in our model result from several factors: suboptimal UVic simulation of nutrient con-

centrations in nearshore regions without land run-off (Eby et al., 2009; Keller et al., 2012; Tivig et al., 2021), unique parameters for chosen brown algae species in our dynamic growth model (Froehlich et al., 2019), consistent nutrient feedback consideration unlike earlier assessments (Froehlich et al., 2019; Lehahn et al., 2016), and the assumption that farms are located within EEZs (Lehahn et al., 2016). Despite these differences, the N-MACS distribution pattern aligns with those in Lehahn et al. (2016, Fig. 3. A), Berger et al. (2023, Figure 4), Duarte et al. (2022, greenish pattern of Figure 1(a)), and Froehlich et al. (2019, Figure 1). While the total N-MACS area remains steady over time, regions of significant productivity (significant N-MACS areas) expand during the initial deployment decade (Fig.S11), resulting from dynamic nutrient cycling. Here, N-MACS suppresses phytoplankton due to canopy shading (Fig.S3), creating a nutrient surplus within its habitat that fertilizes N-MACS (see Sect.3.3).

In productive N-MACS regions, simulated macroalgae productivity averages 165 tonnes DW km⁻² yr⁻¹, rising to 223 tonnes DW km⁻² yr⁻¹ in No_Temp (Tab.1). Farmed seaweed productivity, including the modeled *Saccharina* species, varies significantly depending on species, cultivation techniques, and environmental conditions. Reported *Saccharina* yields in Europe range from 4 to 450 tonnes DW km⁻² yr⁻¹ (Peteiro et al., 2014; Buck & Buchholz, 2004), while in northeast Asia, yields can reach 2,400-3,000 tonnes DW km⁻² yr⁻¹ (Yokoyama et al., 2007; Zhang et al., 2011).

Although N-MACS farms were initially established in all ocean grid boxes adjacent to land between 60°S and 60°N in year 2020, sustainable biomass harvests are mainly found in four regions with high nutrient availability: the Eastern Boundary Upwelling Systems in the nearshore Pacific regions of South America and the Atlantic coasts of Africa (Chavez & Messié, 2009; Fréon et al., 2009), the northeast Pacific and the Southern Ocean (Tab.S1). This is consistent with the findings of Berger et al. (2023), Arzeno-Soltero et al. (2023), and Duarte et al. (2021).

In the sensitivity study (No_Temp), where temperature no longer affects macroalgae growth, the N-MACS distribution mirrors the base case, albeit with increased biomass productivity in mid to high latitudinal coastal regions (Tab.1, Fig.S2). By employing local macroalgae species better adapted to specific temperature ranges, optimization of macroalgae cultivation and enhancement of the CDR potential of nearshore macroalgae-based strategies may be achievable.

Table 1. Summary table of N-MACS simulations. Significant N-MACS area is area with ≥ 100 tonnes DW per km^2 per year. The changes are N-MACS variations relative to Ctrl_RCP4.5.

	Unit	N-MACS	No_Temp
Total yield	Gt DW	188.96	293.40
N-MACS total area	10^6 km^2	24.34	23.65
Significant N-MACS area		14.29	15.97
Total carbon fixation in N-MACS	GtC	56.7	88.0
Annual carbon fixation (avg. 2020 to 2100)	GtC yr^{-1}	0.7	1.1
Annual unit area carbon fixation	tC $\text{km}^{-2} \text{ yr}^{-1}$	29.1	46.5
Change of global climate system in 2100 (3000 in parentheses)			
Surface averaged temperature (SAT)	$^{\circ}\text{C}$	-0.07 (-0.08)	-0.12 (-0.13)
Atmospheric CO_2 concentration	ppm	-14.2 (-12.0)	-22.6 (-18.3)
Change of global carbon reservoirs in 2100 (3000 in parentheses)			
Atmosphere		-30.1 (-25.5)	-47.9 (-38.9)
Ocean (including carbon fixation by N-MACS)	GtC	35.9 (31.4)	57.1 (48.8)
Land		-5.8 (-5.9)	-9.2 (-9.9)
Change of integrated marine biogeochemical parameters in 2100 (3000 in parentheses)			
POM export at 2km depth	GtC yr^{-1}	-4.151 (0.37)	-7.245 (0.58)
PO_4 (full depth)	Tmol	-11.64 (-11.91)	-18.10 (-18.49)
NO_3 (full depth)	Tmol	7.68 (15.78)	-62.51 (-6.01)
Phytoplankton NPP	GtC yr^{-1}	-0.36 (-0.52)	-0.50 (-0.82)

* DW: dry weight; POM: particle organic matter; tC: tonnes of carbon (10^3 Kg);
GtC: Giga (10^9) tonnes of carbon; Tmol: Tera moles (10^{12} moles).

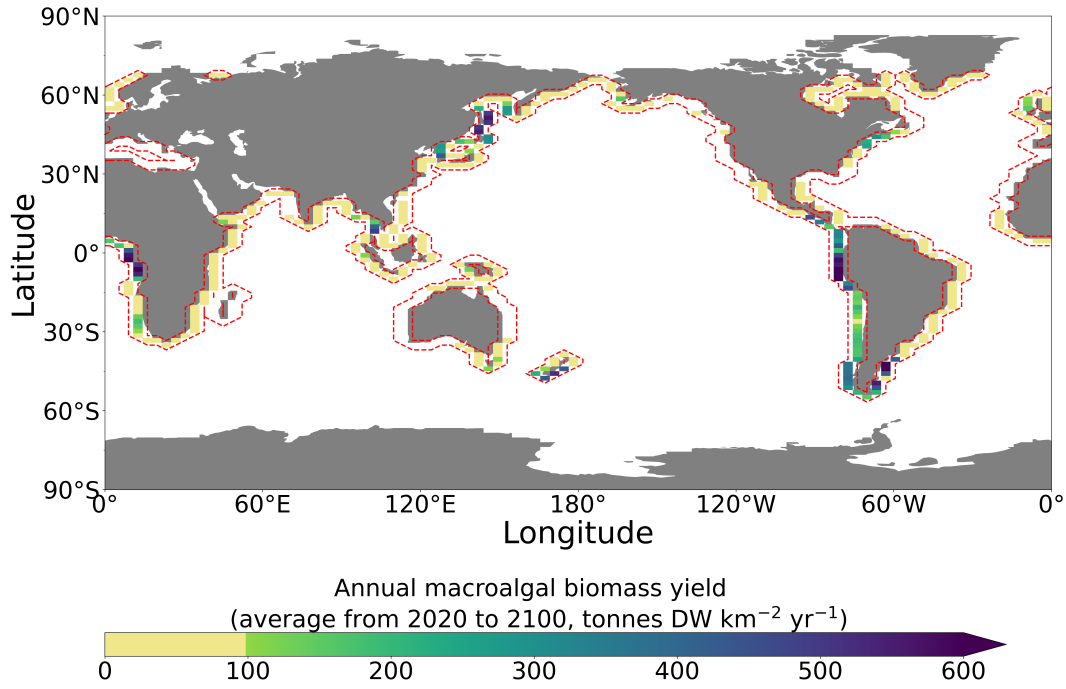


Figure 1. Annual macroalgae biomass yield (averaged from year 2020 to year 2100). Dashed red lines outline the initial seeding locations in year 2020. Regions with high macroalgae productivity include: Coasts of North Western Pacific (near northern China, Japan and Korean Peninsula), South Eastern Pacific (coasts of South America), South Eastern Atlantic (mid-south Africa coast), coast of New Zealand, and South Eastern of Australia. Yellowish areas indicate relatively lower yield (≤ 100 tonnes DW per km^2 per year).

3.2 CDR capacity and impacts on carbon cycle

The CDR capacity of the N-MACS approach can be quantified as the carbon contained (and securely stored) within the harvested macroalgae biomass. From 2020 to 2100, the N-MACS simulation demonstrates a total sequestration of 56.7 GtC (equivalent to 207.9 GtCO₂). In the No_Temp simulation, this capacity increases to 88 GtC due to elevated macroalgal productivity. The atmospheric CO₂ sequestration in N-MACS/No_Temp scenarios translates to a reduction in global-mean surface air temperature (SAT) by 0.07°C/0.12°C (Tab.1, Fig.S1). While this reduction in SAT alone does not enable the RCP 4.5 emission scenario to align with the Paris Agreement, the annual carbon removal (equivalent to 2.60/4.03 Gt CO₂eq) is, for example, on par with the 2022 annual CO₂ emissions from the global building sector (2.94 Gt CO₂, IEA (2023)).

The simulated global average unit-area CDR capacity is 29.1 to 46.5 tC km⁻² within N-MACS occupied regions (106.8 to 170.7 tCO₂ km⁻², Tab.1). Conversely, the global dynamic seaweed growth model of Arzeno-Soltero et al. (2023) suggested that macroalgae farming, particularly in the equatorial Pacific, could yield about 1 GtC for 1 million km² of EEZ waters, translating to 1,000 tC km⁻² yr⁻¹. These differences stem from model differences and experiment setups. Their model, incorporating four types of macroalgae species with high carbon content and yield, operates independently from dynamic nutrient changes, which we find often limits N-MACS growth, and runs for one year. Our estimation is also lower than the globally averaged per-unit-area CDR capacity of 57 tC km⁻² yr⁻¹ in Wu et al. (2023), where the identical macroalgae model of N-MACS is applied to open-ocean regions. This difference primarily arises from the diverse distribu-

tion of macroalgae farms across varying nutrient fields, as depicted by Wu et al. (2023) for open-ocean regions, contrasted with the current N-MACS in nearshore areas. The discrepancy is exacerbated by the coarse grid resolution in UVic, likely underestimating coastal productivity (Keller et al., 2012; Tivig et al., 2021). Nevertheless, the annually averaged carbon sequestration of N-MACS is estimated at 0.7 to 1.1 GtC yr⁻¹ (2.6 to 4.0 GtCO₂ yr⁻¹), surpassing the 0.37 GtC yr⁻¹ reported by Berger et al. (2023), something again attributable to the different dynamic macroalgae growth and Earth system modeling approaches.

The net increase in the oceanic carbon reservoir, consisting of water-column carbon content and the harvested macroalgae in the N-MACS (No_Temp) simulations, is 35.9 (57.1) GtC in 2100 (Tab.1), equivalent to the N-MACS induced air-sea carbon flux in the model (Fig.S6, Fig.S7). However, the increase in the oceanic plus macroalgae carbon reservoir is approximately two-thirds of the harvested macroalgae carbon, corresponding to 63.3% (64.9%) of the net carbon removed by harvesting the macroalgae. The disparity between the increase in the ocean plus macroalgae carbon pool and the carbon harvested in the form of macroalgal biomass is largely caused by backfluxes from the ocean into the atmosphere due to diminished atmospheric pCO₂ (Oschlies, 2009) and partially by the reduced phytoplankton net primary production (PNPP) from canopy shading and nutrient competition effects introduced by N-MACS (see Sect.3.3). This efficiency is somewhat higher than the CDR efficiency of 58% in Berger et al. (2023), who employed a dynamic macroalgae growth model in conjunction with a high-resolution ocean biogeochemical model with prescribed atmospheric CO₂, i.e. without back-fluxes from the ocean into the atmosphere due to diminished atmospheric pCO₂, for 5-year simulations.

Meanwhile, the increase in the oceanic plus macroalgae carbon reservoir induced by N-MACS until 2100 leads to a corresponding decline in the terrestrial carbon reservoir of 5.8 to 9.2 GtC (see Tab. 1) via an atmospheric carbon climate feedback. This response illustrates the Earth system’s endeavor to maintain equilibrium, with carbon cycling between terrestrial and oceanic reservoirs, primarily mediated by atmospheric interactions. This finding aligns with other studies, suggesting that ocean-based CDR could potentially weaken terrestrial carbon sinks, especially through the reduction of the CO₂ fertilization effect on terrestrial photosynthesis (Keller, Lenton, Littleton, et al., 2018).

During the implementation phase, an enhancement of approximately 29% (37%) in the air-to-sea downward carbon flux was observed within the macroalgae-occupied areas in N-MACS (No_Temp)(Fig.S5), aligning with the 52% enhancement reported by Berger et al. (2023). The lesser degree of carbon flux enhancement observed in our simulation within the macroalgae-occupied areas is attributed to 1) the canopy shading effect on phytoplankton in our model, reducing PNPP and subsequent carbon flux into the ocean (Fig.2d & Fig. S3); and 2) the dynamic atmospheric pCO₂ in our model compared to prescribed fixed pCO₂ in Berger et al. (2023), as well as different biogeochemical properties of macroalgae and phytoplankton in the two models. Our results further highlight the potential challenges inherent in the measurement, reporting, and verification processes when assessing carbon flux enhancements. Additionally, a slight decrease in DIC in mid and deep waters is evident in Fig.S4a, stemming from reduced water column remineralization due to the diminished downward particulate organic carbon (POC) export (see Sect.3.3).

3.3 Impacts on global marine biogeochemistry

In our simulations, the 80-year implementation of N-MACS has significantly impacted global marine biogeochemistry. This includes ocean surface nutrient distributions, surface ocean alkalinity, and dissolved oxygen concentrations at mid-depth (Fig. 2). Additionally, simulated net primary production and the distributions of ordinary phytoplankton and diazotrophs are also affected by N-MACS deployment. Notably, some of

these impacts persist until the year 3000, despite the cessation of N-MACS in 2100 (see below).

The N-MACS macroalgae model delineates two primary impacts of macroalgae on phytoplankton: nutrient competition and canopy shading (Wu et al., 2023, Sect.2.2.3). Harvesting macroalgae not only sequesters carbon but also extracts nutrients within the harvested biomass, leading to an immediate drop in global PNPP post N-MACS initiation in 2020, with a gradual reduction during N-MACS deployment till 2100 (Fig.3e). This PNPP decline predominantly occurs along coast-adjacent N-MACS areas (Fig.2d). Additionally, certain open-ocean regions beyond coastal farms exhibit a PNPP increase, notably in the Indian Ocean, eastern Atlantic near Africa, and eastern equatorial Pacific. This is attributed to nutrient leakage from N-MACS areas (see Fig.2d; further details in the subsequent paragraph). N-MACS implementation suppresses oceanic nitrogen fixers, diazotrophs, due to canopy shading and phosphate competition by macroalgae (Fig.S9). Although certain regions exhibit heightened diazotroph biomass due to increased phosphate levels (Fig.S10a&c), the overall nitrogen fixation relative to DNPP diminishes during N-MACS deployment (Fig.3h). Zooplankton, assumed capable of grazing on macroalgae (Wu et al., 2023), primarily feed on phytoplankton due to a lower macroalgae grazing preference, hence their biomass trends closely with those of phytoplankton (not shown).

Fig.3a illustrates a notable increase in surface ocean PO_4 concentrations (top 50m) following N-MACS initiation, followed by a decrease. Three primary factors underlie this PO_4 rise. Firstly, the suppression of phytoplankton by macroalgae leads to a decreased organic carbon export out of the euphotic zone. Secondly, macroalgae cannot fully utilize the *in-situ* PO_4 due to the limited growth rate and maximum macroalgae biomass (Wu et al., 2023). Lastly, the higher stoichiometric N:P ratio of 20:1 in macroalgae, compared to the Redfield ratio of 16:1 in phytoplankton, entails less PO_4 consumption per nitrogen unit for growth. This explains the increases in surface PO_4 levels in N-MACS regions shown in Fig.2c (Fig.S8c for No_Temp). Nitrate concentrations in N-MACS regions also rise due to phytoplankton inhibition and unexhausted available nitrate from macroalgae growth (Fig.2a). These disparities consequently induce lateral nutrient leakage from N-MACS areas, fertilizing the aforementioned downstream area of coastal N-MACS farms. Here, augmented PNPP consumes the displaced nutrients, driving a regional PO_4 concentration reduction (Fig.2c).

A reduction in surface PNPP within N-MACS regions triggers a decline in particulate organic matter (POM) export to ocean depths, as observed at 2000 m in Fig. 3f and Tab.1. This decline subsequently diminishes oxygen consumption via aerobic remineralization of organic carbon, thus elevating the oxygen concentration across middle and bottom waters (Fig.S4d, Fig.S12d). Notable increases in dissolved oxygen concentrations at 300m depth are apparent in the northwestern Pacific, eastern equatorial Pacific, and southern Atlantic near the South American continent (Fig.2e & Fig.3). Specifically, oxygen minimum zones (OMZs) in the North Pacific and equatorial Atlantic Ocean have shrunk compared to Ctrl_RCP4.5. The increased oxygen levels inhibit denitrification in the subsurface and the upwelling system in the eastern equatorial Pacific (Fig.2f&i, Bange et al. (2019); Ravishankara et al. (2009)), and diminished remineralization of organic carbon curtails nutrient regeneration, reducing nutrient upwelling (Fig.2g&h). This results in elevated NO_3 but reduced PO_4 compared to the Ctrl_RCP4.5 in the open ocean of the eastern equatorial Pacific (Fig.2a, c, d & f). Another factor contributing to the reduced PO_4 in the source waters of the upwelling regions is the decreased PNPP in the N-MACS areas, which lessens export and thereby reduces the PO_4 source from POM remineralization (Fig.2d, Fig.3f). Furthermore, the aforementioned decreased denitrification increases the NO_3 supply in the upwelling system to the surface, especially in oxygen-depleted regions off Peru where reduced POM remineralization leads to lesser denitrification and nitrogen loss. However, in the No_Temp simulation, amplified macroalgae

growth utilizes upwelled NO_3 before export to the open ocean, mitigating the NO_3 increase in the eastern equatorial Pacific (Fig.S8a).

Despite the reduction in mid-depth denitrification (Fig.2i), which also diminishes alkalinity production, the surface alkalinity in N-MACS increases about 1% or 10 to 20 mmol m^{-3} by 2100 (Fig.2b), due to reduced CaCO_3 generation from the PNPP reduction induced by continuous phosphate removal by N-MACS (Fig.S12, Schmittner et al. (2008, Eq.2)). Post N-MACS discontinuation in 2100, which effectively terminates canopy shading and nutrient competition effects, results in a marked resurgence in PNPP and thereby also a decreases in global surface nutrient concentrations (Fig3a, b&e). Additionally, diazotroph biomass, DNPP, and nitrogen fixation recover (Fig.S9, Fig3h). The export of PNPP and POC as well as the subsurface oxygen consumption via organic carbon remineralization also recovers (Fig3g). Additionally, the air-sea CO_2 flux reverts to baseline levels after cessation of the carbon sequestration by macroalgal harvest from the ocean (Fig.S6, S7).

By year 3000, the average surface temperature in the N-MACS/No_Temp simulations is slightly lower by -0.08/-0.13 $^{\circ}\text{C}$, respectively, compared to Ctrl_RCP4.5, maintaining the temperature reduction achieved by N-MACS in 2100 (Tab.1). After N-MACS termination in year 2100 and until year 3000, both oceanic and terrestrial carbon reservoirs shrink, with oceanic plus macroalgae carbon storage decreasing by 4.5 GtC in N-MACS and 8.3 GtC in No_Temp, and terrestrial carbon storage declining by 0.1 GtC and 0.7 GtC in N-MACS and No_Temp scenarios respectively. This leads to a 4.6 / 9.0 GtC or 2.2 / 4.3 ppm atmospheric CO_2 increase (Tab.1). Decreased global temperatures slow photosynthesis and soil respiration, in combination yielding a small reduction in the terrestrial carbon pool. The decrease in the oceanic carbon pool mainly arises from the PNPP reduction as a consequence of permanent phosphate removal during the operation of N-MACS. This enduring PO_4 removal leads to long-term alterations in marine biogeochemistry, as shown by extended simulations until year 3000 (Fig.3). Though only 0.4% of total oceanic phosphate is removed by 2100 (Fig.3c), it induces a persistent reduction in PNPP, DNPP, and nitrogen fixation (Fig.3a&h, S10b&d). This prevents PNPP and DNPP recovery to RCP4.5 levels from 2100 to 3000 (Fig. 3 e), leading to increased oxygen due to overall POC export reduction (Fig.3d&g, Fig.S12).

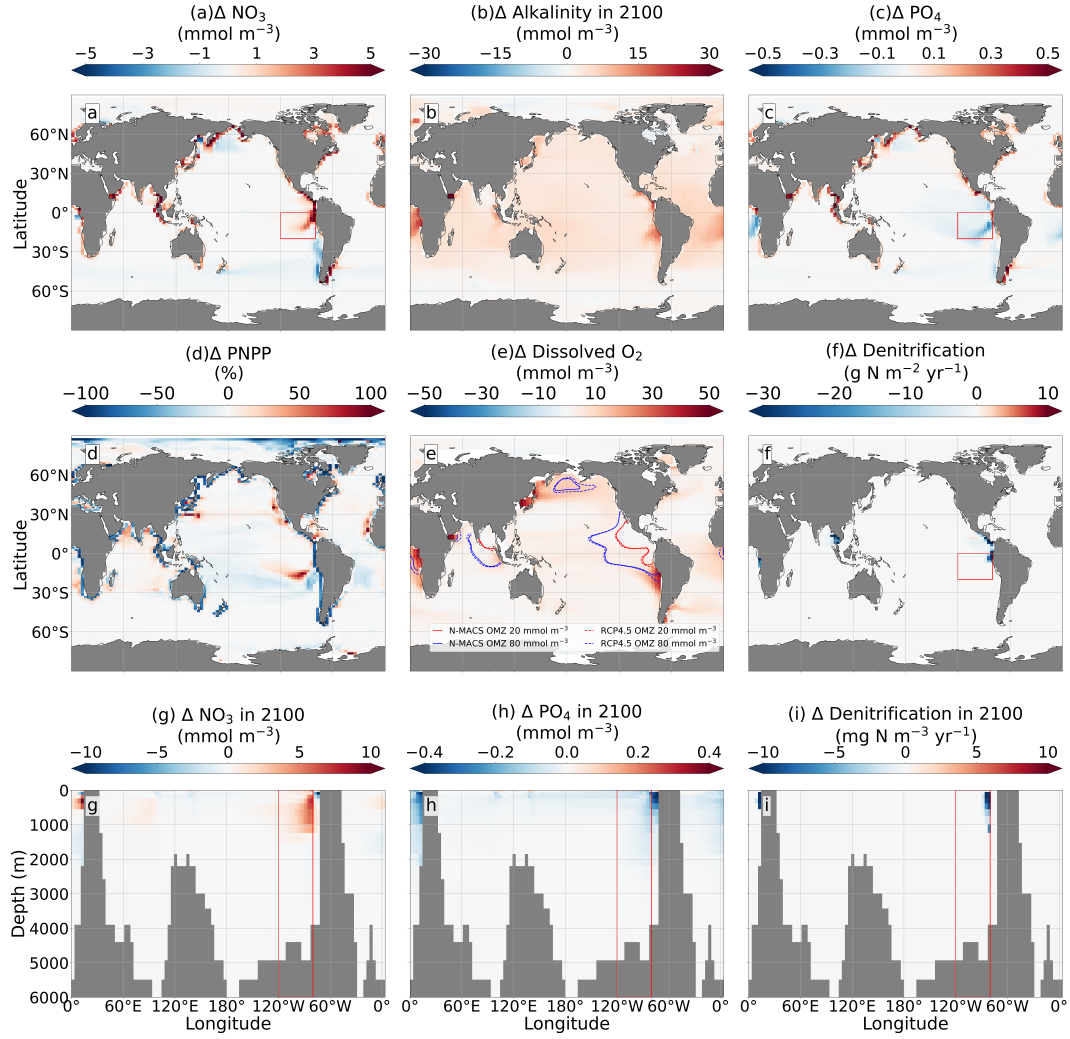


Figure 2. Differences in simulated oceanic properties in year 2100 after continuous N-MACS deployment from 2020 to 2100, with respect to Ctrl_RCP4.5 without N-MACS deployment (data averaged over this period, except for **d** and **e** representing data in 2100): **a**: Surface-layer nitrate (top 50m); **b**: Surface-layer alkalinity; **c**: Surface-layer phosphate; **d**: Phytoplankton net primary production (PNPP); **e**: Dissolved oxygen concentrations and oxygen minimum zones (OMZs) at a depth of 300m; **f**: Oceanic denitrification rates. Subfigures **g**, **h** & **i** represent latitudinally averaged data from 20°S to 0°, relative to the Ctrl_RCP4.5 scenario depicted in subfigures **a**, **c**, & **f** (highlighted by red rectangular regions between latitudes 20°S to 0° and longitudes 80°W to 120°W): **g**: Phosphate concentrations, **h**: Nitrate concentrations, **i**: Annual denitrification rates.

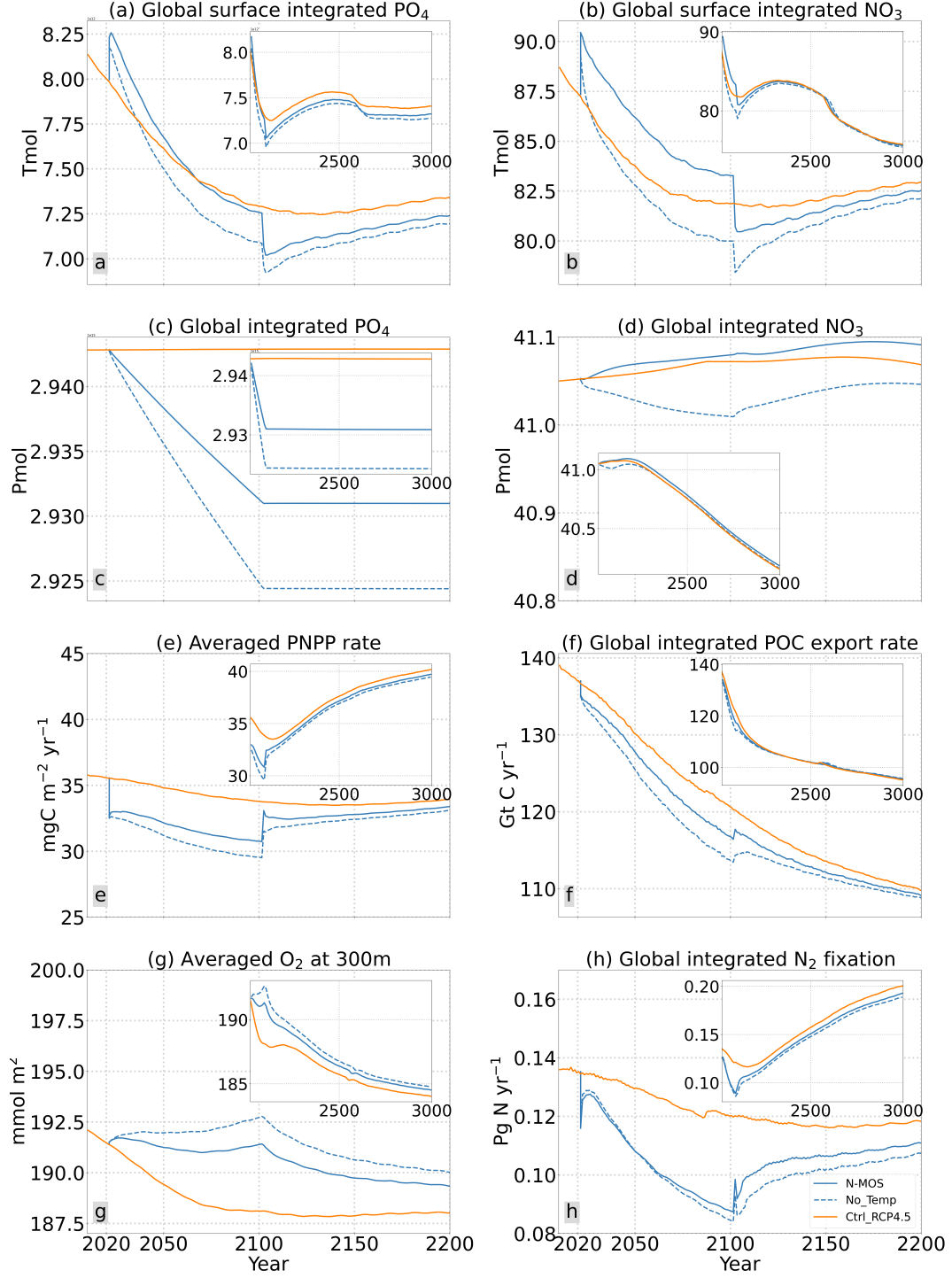


Figure 3. Temporal evolution of globally integrated nutrients, Phytoplankton Net Primary Production (PNPP), and Particulate Organic Carbon (POC) Export at 2,000m depth: Comparison of N-MACS (solid blue), No_Temp (dashed blue), and Ctrl_RCP4.5 Baseline Simulation (orange). Insets in each panel extend the timeline to the year 3000. **a & c:** Permanent removal of PO_4 from the surface, **b & d:** Surface NO_3 levels and global NO_3 trends (increase in N-MACS, decrease in No_Temp). **e:** Surface PNPP (see also Fig.2d). **f:** The export of POC at 2,000m depth. **g:** The averaged O_2 concentration at 300m depth. **h:** Globally integrated Nitrogen fixation.

4 Conclusion & Outlook

Our analysis highlights the substantial annual gigatonne-scale CO₂ sequestration potential of N-MACS, though with marine biogeochemical and global carbon cycle feedbacks reducing the additional air-sea CO₂ flux by 35% compared to carbon removal via harvesting. Large-scale N-MACS deployment considerably alters marine biogeochemistry and ecosystems, suppressing PNPP, elevating dissolved oxygen concentrations, reducing denitrification, and decreasing surface ocean alkalinity. Terminating N-MACS in 2100 triggers a transient rebound in surface PNPP and a decrease in the air-sea CO₂ flux, yet long-term effects like nutrient depletion and increased oxygen levels persist for centuries. Promising regions for macroalgae production include the upwelling systems in South America, Africa's Atlantic coasts, the Northeast Pacific, and the Southern Ocean.

Our simulations have certain limitations: Given that the UVic operates on a coarse grid resolution ($1.8^\circ \times 3.6^\circ$), it inadequately represents the physical and biogeochemical processes of the coastal ecosystem in the marine ecosystem model (Keller et al., 2012). While not significantly impacting our current global and millennial scale simulations, it may affect coastal macroalgae farming simulations when considering nutrient fluxes in coastal areas (e.g., Van Der Molen et al. (2018)). Possible improvements to our model include a consideration of a wider range of macroalgae species (Arzeno-Soltero et al., 2023; Duarte et al., 2022), explicit accounting of iron limitation (Paine et al., 2023; Anton et al., 2018), dynamic cellular stoichiometry, and current impacts on macroalgae frond erosion (Frieder et al., 2022; Broch & Slagstad, 2012). Acknowledging both remineralization-resistant particulate and dissolved organic carbon release from macroalgae and subsequent deep-water may be crucial for comprehending the CDR capacity (Pedersen et al., 2021; Ortega et al., 2019; Duarte & Krause-Jensen, 2017; Wada & Hama, 2013). Further considerations include macroalgae halocarbon emissions (Baker et al., 2001; Leedham et al., 2013; Jia et al., 2022) and alterations in ocean surface albedo and local ecosystem (Bach et al., 2021; Boyd et al., 2022). Herein it's assumed that no nutrients from the harvested biomass are returned to the ocean, which significantly impacts the simulated biogeochemistry. Thus, evaluating nutrient extraction and return strategies is imperative if N-MACS is pursued as a sustainable CDR approach.

Governance and societal facets need consideration in macroalgae-based CDR, particularly due to potential spatial competition between macroalgae cultivation and fisheries, especially along the Peruvian coast (Gattuso et al., 2021; Ricart et al., 2022; Merk et al., 2022). A Comprehensive Life Cycle Analysis (LCA) considering energy consumption biomass conversion efficiency, and financial cost is pivotal (Fernand et al., 2017; Melara et al., 2020; Capron et al., 2020; Hughes et al., 2012; Aitken et al., 2014).

5 Open Research

The data files used in this paper are available through GEOMAR at (Wu, 2024).

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Figure 1.

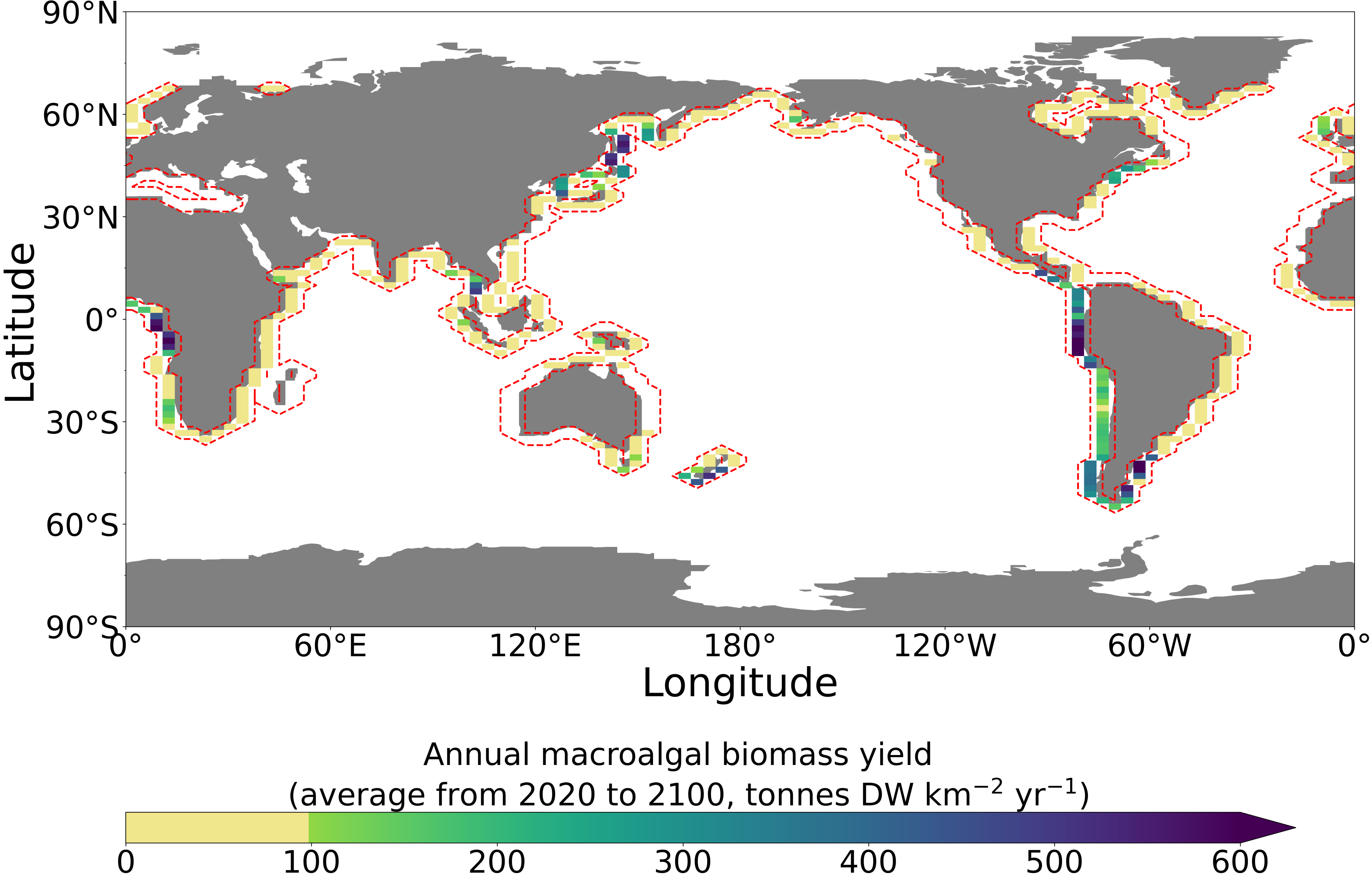


Figure 2.

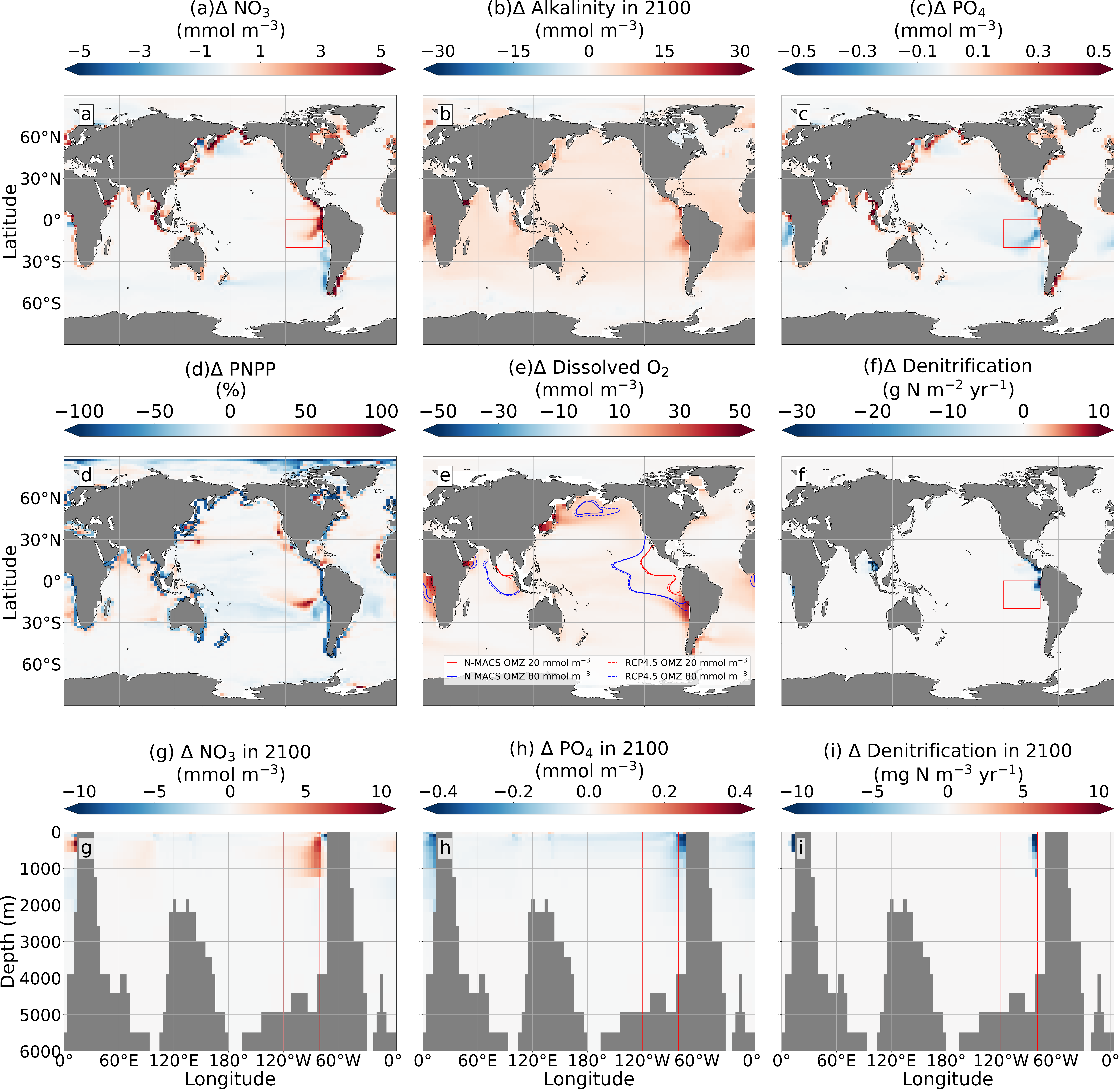


Figure 3.

