

Ocean carbon sink assessment via temperature and salinity data assimilation into a global ocean biogeochemistry model

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Abstract. Global ocean biogeochemistry models are frequently used to derive a comprehensive estimate of the global ocean carbon uptake. These models are designed to represent the most important processes of the ocean carbon cycle, but the idealized process representation and uncertainties in the initialization of model variables lead to errors in their predictions. Here, observations of ocean physics (temperature and salinity) are assimilated into the ocean biogeochemistry model FESOM-REcoM over the period 2010-2020 to study the effect on the air-sea CO₂ flux and other biogeochemical variables. The assimilation nearly halves the model-observation differences in sea surface temperature and salinity, with modest effects on the modeled ecosystem and CO₂ fluxes. The main effects on the air-sea CO₂ flux occur on small scales in highly dynamic regions, which pose challenges to ocean models. The largest imprint of assimilation is in the Southern Ocean during winter. South of 50°S, winter CO₂ outgassing is reduced and thus the regional CO₂ uptake increases by 0.18 Pg C yr⁻¹ through the assimilation. Other particularly strong regional effects on the air-sea CO₂ flux are located in the area of the North Atlantic Current. Yet, the effect on the global ocean carbon uptake is a comparatively small increase by 0.05 Pg C yr⁻¹ induced by the assimilation, yielding a global mean uptake of 2.78 Pg C yr⁻¹ for the period 2010-2020.

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1 Introduction

The ocean plays a pivotal role in regulating the global carbon budget and thereby mitigating the impacts of anthropogenic carbon dioxide (CO₂) emissions on the Earth's climate. Since the 1960s, the ocean has absorbed consistently around 25% of anthropogenic CO₂ emissions annually (Friedlingstein et al., 2023) and has cumulatively taken up 26–34% of fossil and land-use change CO₂ emissions since the onset of the industrial revolution (Crisp et al., 2022). However, quantification of air-sea CO₂ flux still remains challenging. Air-sea CO₂ flux is usually inferred from the gradient of partial pressure (pCO₂) or fugacity (fCO₂) of CO₂ across the air-sea interface (Wanninkhof, 2014). Yet, during 2010-2020, which constitutes the best-sampled decade in terms of surface ocean pCO₂ observations so far, observations covered merely 3% of the monthly global ocean (as calculated from the 1°x1°-gridded SOCAT product; Bakker et al., 2016). While the North Atlantic and North Pacific are comparably well observed, data remain scarce in vast regions, such as the Indian Ocean, South Pacific and areas south

of 30°S during austral winter, where less than 1% of SOCAT grid cells have been sampled. Although these observations are
25 thought to be representative of a larger area (Jones et al., 2012; Hauck et al., 2020), challenges in deriving a comprehensive
global estimate of the global ocean CO₂ uptake arise due to substantial spatial and temporal pCO₂ variations and potential
biases induced by the irregular sampling pattern (Denvil-Sommer et al., 2021; Gloege et al., 2021; Hauck et al., 2023b).
Particularly in the Southern Ocean, the uncertainty is considerable (Gerber et al., 2009; Gloege et al., 2021), where estimates
of the mean flux range from -0.37 to -1.25 PgC yr⁻¹ for the period 2010-2018 (data provided by Hauck et al., 2023b).

30 In the Global Carbon Budget, estimates of the ocean carbon sink were initially derived from hindcast simulations of global
ocean biogeochemistry models (GOBMs) (Le Quéré et al., 2009; Wanninkhof et al., 2013; Hauck et al., 2020). More recently,
air-sea CO₂ flux estimates were added based on regression and machine learning techniques, interpolating pCO₂ observations
to achieve global coverage through advanced statistical methods (referred to as pCO₂ products; Rödenbeck et al., 2015).
Furthermore, atmospheric transport models that ingest atmospheric CO₂ measurements were employed to estimate the ocean
35 carbon uptake (referred to as atmospheric inversions; Peylin et al., 2013). Although the different estimation methods have
provided valuable and robust insights into large-scale patterns of oceanic carbon uptake (Gruber et al., 2009), discrepancies
have emerged. Assessments based on pCO₂-products tend to yield larger estimates of the ocean carbon sink, with stronger
trends towards more uptake, compared to estimates based on models (Friedlingstein et al., 2023; Terhaar et al., 2022). The
larger estimates are supported by ocean interior observations (Müller et al., 2023), atmospheric oxygen data and atmospheric
40 inversions (Friedlingstein et al., 2023). For the years 2010-2020, pCO₂ products included in the Global Carbon Project suggest
a mean oceanic sink of 3.0 ± 0.4 PgC yr⁻¹, while the mean of Global Carbon Project GOBMs is 2.5 ± 0.4 PgC yr⁻¹ (data
provided by Friedlingstein et al., 2023). Trends over the same time period are 0.7 PgC yr⁻¹ dec⁻¹ and 0.3 PgC yr⁻¹ dec⁻¹,
respectively.

Machine learning estimates perform well when trained with sufficient data (Gloege et al., 2021). However, their performance
45 is less reliable in data-sparse areas. Particularly in the Southern Ocean, many pCO₂-products show diverging results from
one another and are likely biased towards more ocean uptake (Hauck et al., 2023b). However even in parts of the North
Pacific, which is undersampled in the 2010s, some pCO₂ products show spurious decadal trends (Mayot et al., 2024). Models
provide process-driven estimates of the CO₂ flux across the entire global ocean, drawing from the theory of ocean dynamics,
biological and chemical processes (Hauck et al., 2020; Fennel et al., 2022). Despite the growing confidence in our mechanistic
50 understanding of the ocean carbon cycle (Crisp et al., 2022), models are also subject to uncertainty. This uncertainty stems
from uncertainties in model parametrizations, model spin-up and initial conditions, unresolved sub-gridscale processes and
uncertainties in the atmospheric forcing (Hauck et al., 2020; Terhaar et al., 2024).

Data assimilation (DA) can be employed to address the emerging discrepancies between pCO₂-products and models (Carroll
et al., 2020). Several studies assimilating ocean surface pCO₂ have focused on specific regions (e.g., a baseline state of air-
55 sea CO₂ fluxes in the Southern Ocean; Verdy and Mazloff, 2017), few years (e.g., optimized biogeochemical initial fields
for the period 2009-2011 in Brix et al., 2015) or the climatological mean state (e.g., corrections of large-scale pCO₂ model
biases in While et al., 2012). These studies capture well the assimilated pCO₂ observations, while obeying physical laws
and biogeochemical (BGC) equations. Data assimilation also provides a better understanding of various components of the

ocean carbon cycle, such as the transport of anthropogenic CO₂ in the ocean (e.g., a reconstruction of anthropogenic carbon storage since 1770 in Gerber et al., 2009), regional and interannual variability of the air-sea CO₂ flux (e.g., global reanalysis in Ford and Barciela, 2017; Carroll et al., 2020; Valsala and Maksyutov, 2010), the biological carbon pump (e.g., carbon export at a nutrient-rich and nutrient-poor site and estimation of BGC parameters related to air-sea CO₂ fluxes in Sursham, 2018; Hemmings et al., 2008, respectively) and specific ecosystems (e.g., the North West European Shelf ecosystem in Ciavatta et al., 2016, 2018). So far, however, there is no data assimilation product that provides a long-term, annually updated estimate of global ocean CO₂ uptake.

While previous studies indicate that the available BGC observations, when assimilated in isolation, are too sparse to constrain the modeled carbon cycle (Verdy and Mazloff, 2017; Spring et al., 2021), the assimilation of physical variables is expected to have a significant indirect effect on the modeled air-sea CO₂ fluxes (Bernardello et al., 2024). This is because the uptake of atmospheric CO₂ depends ultimately on the modeled physical carbon transport between the surface, the mixed layer and the deep ocean in the form of dissolved inorganic carbon (DIC) through mixing, upwelling and subduction (Doney et al., 2004). According to current knowledge, ocean physics is the dominant driver of interannual variability of the global air-sea CO₂ flux and also responsible for stagnation and acceleration of the CO₂ uptake on decadal scales (Doney et al., 2009; Keppler and Landschützer, 2019; Mayot et al., 2023; Liao et al., 2020; DeVries et al., 2017). Related to the strong control that physics exert on the interannual variability of air-sea CO₂ fluxes, it was shown in one idealized study that assimilating ocean physics at the initial state of a model simulation has a stronger and more positive impact on the modeled carbon cycle on interannual time-scales than assimilating the BGC initial state (Fransner et al., 2020). However, the relative importance of uncertainties in physical and biogeochemical fields generally remains an open research question (e.g. Séférian et al., 2014; Li et al., 2016; Lebehot et al., 2019). Therefore, we here use ensemble-based data assimilation of ocean physics into a global ocean biogeochemistry model aiming to improve the modeled air-sea CO₂ flux for the years 2010-2020. For this, we continuously assimilate temperature and salinity observations from remote-sensing at the surface and from in-situ profile measurements for eleven years and update the modelled temperature, salinity, horizontal velocities and sea surface height, using an ensemble Kalman filter variant (Nerger et al., 2012).

Several difficulties are associated with physics DA into GOBMs. A common issue is erroneous equatorial upwelling leading to unrealistically high biological productivity in the tropics (Park et al., 2018; Gasparin et al., 2021; Raghukumar et al., 2015). Furthermore, any coupled ecosystem model is adapted to its associated physical model with its strengths and weaknesses through carefully selected parameter values and a spin-up to near-equilibrium. Accordingly, the modeled carbon cycle may react very sensitive to deviations from the physical state that is typical for this model (Kriest et al., 2020; Spring et al., 2021). Potentially, this leads to biases in the carbon cycle through physics DA. The question therefore arises to what extent an ecosystem model coupled to a data-assimilated physical model also represents a more realistic biogeochemistry, and which mechanisms drive the response of the CO₂ flux in physics DA approaches. One possible driver is the physical transport of DIC and alkalinity because velocities and diffusivity are changed by the DA, affecting in particular the upwelling of carbon-rich waters and subduction, which is important to capture the ocean storage of anthropogenic carbon (Davila et al., 2022). Furthermore, physics DA may change pCO₂ directly through its temperature-dependence, an effect emphasized by Verdy and

Mazloff (2017). Additionally, the modelled biological pump might be altered, for example through the temperature-dependency
95 of phytoplankton growth or through effects of stratification on nutrient availability.

In this study, we describe the response of the model’s air-sea CO₂ flux to physics DA and identify the underlying mechanistic
drivers. To this end, we differentiate between the thermally, DIC- and alkalinity induced components and changes in lateral
and vertical transport through mixing and advection. We focus, firstly, on the global air-sea CO₂ flux. Secondly, we investigate
the Southern Ocean given the relevant impact of DA in Southern Ocean winter in our study. Thirdly, we present regions in the
100 North Atlantic given observational coverage and relevant local processes there.

2 Methods

2.1 Model FESOM-REcoM

The oceanic model component, FESOM2.1, computes the advection, diffusion, and mixing of passive biogeochemical trac-
ers. The model is based on hydrostatic primitive equations under the Boussinesq approximation and utilizes a finite-volume
105 discretization approach with surface triangles projected vertically to form prisms. Salinity (S), temperature (T), and biogeo-
chemical tracers are located at the vertices of triangles (nodes), while the horizontal velocities are centered at the triangles
(elements). The model allows for a variable mesh resolution (see Section 2.2) and incorporates parametrizations for diffusion
and eddy-stirring along isoneutral surfaces, for which parametrized mixing is scaled by mesh resolution (Danilov et al., 2017).
Vertical mixing is parametrized through the KPP scheme and the mixing depth is specified through a ‘boundary layer’ (the
110 layer of active mixing, which may have a vertical structure because the mixing of all properties across the layer is not in-
stantaneous, as opposed to the mixed layer which is defined by already well-mixed properties, Large et al., 1994), with an
additional vertical mixing scheme used in the Southern Ocean (Monin–Obukhov parametrization, Timmermann and Beck-
mann, 2004). The surface salinity (SSS) is restored towards the World Ocean Atlas climatology through a fictional surface flux
with $v_{SSS} = 50\text{m}/300\text{days}$ according to equation 1 and as in Gürses et al. (2023):

$$115 \quad (SSS_{\text{clim}} - SSS_{\text{model}}) * v_{SSS} * (h_{\text{surf}})^{-1} \quad (1)$$

with surface-layer width h_{surf} . A detailed description of FESOM2.1 and a model assessment are provided by Danilov et al.
(2017) and Scholz et al. (2019, 2022).

The ocean biogeochemistry component, the Regulated Ecosystem Model version 3 (REcoM3), describes processes in the
ocean carbon cycle and represents oceanic carbon in the form of dissolved inorganic carbon, dissolved organic carbon, plankton
120 and detritus (Gürses et al., 2023). REcoM3 contains 28 BGC tracers (listed in Appendix Table A1). There are two phytoplank-
ton groups: diatoms and small phytoplankton with implicit representation of calcifiers; two zooplankton groups: mixed and
polar macro zooplankton (Karakuş et al., 2021); and two classes of detritus. REcoM3 includes variable intracellular stoi-
chiometry with ratios of C:N:Chl:CaCO₃ for the small phytoplankton and C:N:Chl:Si for diatoms, which is propagated to
zooplankton and detritus (Schartau et al., 2007; Hohn, 2008). The publicly available Routines To Model The Ocean Carbonate
125 System (mocsy2.0, Orr and Epitalon, 2015) are used to compute pCO₂ and air-sea CO₂ flux, employing the gas-exchange

parameterization of Wanninkhof (2014). Alkalinity is restored by a fictional surface flux of 10 m yr^{-1} (as in Hauck et al., 2013; Schourup-Kristensen et al., 2014; Gürses et al., 2023). The current model version FESOM2.1-REcoM3 was assessed by Gürses et al. (2023) and previous versions were evaluated and applied in global and regional studies of the ocean carbon cycle and planktonic ecosystems (Hauck et al., 2013; Schourup-Kristensen et al., 2014; Hauck et al., 2020; Karakuş et al., 2021).

130 2.2 Simulation set-up

The model setup for both simulations closely follows Gürses et al. (2023). The mesh resolution is nominally 1 degree, ranging between 120 km and 20 km with enhanced resolution in the equatorial belt and north of 50°N (126858 surface nodes). It has 47 vertical layers with thickness ranging from 5 m at the surface to 250 m in the deep ocean, as described by Scholz et al. (2019, CORE mesh). The model time step is 45 minutes. For atmospheric forcing, JRA55-do v.1.5.0 is used, a reanalysis product
135 tailored for driving ocean-sea-ice models (Tsujino et al., 2018). The atmospheric CO_2 mixing ratio ($x\text{CO}_2$) values were taken from the Global Carbon Budget (Joos and Spahni, 2008; Ballantyne et al., 2012; Friedlingstein et al., 2023). We use model restart fields from Gürses et al. (2023) where the model was spun-up by repeating the year-1961 JRA forcing for 189 years with preindustrial atmospheric CO_2 conditions, followed by a period from 1800 to 1957 with increasing atmospheric CO_2 . Subsequently, simulations were continued with historical JRA forcing from 1958 to 2009. During the assimilation window
140 (2010-2020), we conduct two ensemble simulations to study the impact of data assimilation (DA): one without DA (referred to as FREE) and another identical setup applying DA (referred to as ASML). For each simulation, the ensemble mean for the following variables is written as output: temperature, salinity, velocity, boundary-layer depth, surface pCO_2 , DIC, alkalinity, nutrients, chlorophyll, net primary production and biological export through sinking of detritus at 190 m. For the year 2020, additional output is available for individual ensemble members, mixed-layer depth, physical sources or sinks of DIC and
145 alkalinity through horizontal and vertical advection and diffusion, and biological net sources or sinks of DIC and alkalinity through combined processes: For DIC, the net biological term is the sum of photosynthesis, respiration, remineralization of dissolved organic carbon, and formation and dissolution of calcite (Gürses et al., 2023, equation A6). For alkalinity, the net biological term is the sum of nitrogen assimilation and remineralization, and formation and dissolution of calcite (Gürses et al., 2023, equation A7).

150 2.3 Data Assimilation

2.3.1 Assimilated observations

The assimilated observations are sea surface temperature (SST), sea surface salinity and profiles of temperature and salinity. The assimilated SST observations are from the Operational Sea Surface Temperature and Ice Analysis (OSTIA) data set (CMEMS Marine Data Store; Good et al., 2020; Donlon et al., 2012; Stark et al., 2007). OSTIA provides daily gap-free
155 maps of SST at a horizontal resolution of $0.05^\circ \times 0.05^\circ$, compiled from in-situ and satellite data from infrared and microwave radiometers. The OSTIA observations were averaged to the FESOM2.1 model grid because their spatial resolution is higher than the nominal resolution of the model grid. An observation error standard deviation of 0.8°C is prescribed for the DA

following Nerger et al. (2020). Observations are excluded in the DA process if the difference between the model and observation exceeds three times the observation error standard deviation, thus 2.4°C , and at grid points with sea ice in the model, as in
160 Tang et al. (2020) and Mu et al. (2022). This exclusion keeps the model stable despite large differences between model and observations at these sites, in particular as water temperature and salinity develop differently under sea ice than under the influence of the atmosphere (Tang et al., 2020). Instead, a ‘gentler’ correction is made by assimilating neighboring points. After the initial phase, about 7% of SST observations are excluded because of the 2.4°C -threshold. Nevertheless, the data assimilation still has a strong effect in areas where these large model-observation discrepancies are typically found (North
165 Atlantic, Japan and Southern Ocean).

The assimilated SSS data is taken from the European Space Agency (ESA) Sea Surface Salinity Climate Change Initiative (CCI) v03.21 data set (Boutin et al., 2021). ESA-CCI contains daily data at a spatial resolution of 50 km, albeit not capturing temporal variability below weekly. The ESA-CCI observations are averaged to the FESOM2.1 model grid. We prescribe a constant observation error standard deviation of 0.5 psu following Nerger et al. (2024). Like for the SST data, SSS observations
170 are excluded at locations where sea ice is present in the model.

The assimilated temperature and salinity profiles are taken from the EN.4.2.2 data set (Good et al., 2013). The EN4 dataset contains quality-controlled profiles from various in-situ ocean profiling instruments. To assimilate the profiles, the observations are assigned to the respective model layers (depth range) in the vertical. In the horizontal, the model values are computed as the average of the grid points of the triangle enclosing the observation. The observation error standard deviation is set to 0.8°C
175 for temperature and to 0.5 psu for salinity, as in Tang et al. (2020).

2.3.2 Assimilation method and implementation

For the assimilation, we use the Localized Error Subspace Transform Kalman Filter (LESTKF, Nerger et al., 2012). The LESTKF sequentially updates the model forecast, incorporating observations when and where available. The model state and error covariance are represented by an ensemble simulation. Thereby, the assimilation of temperature and salinity affects the
180 state of the physical model in its whole, including the horizontal velocities and sea-surface height. A review of the LESTKF and other filters frequently used in geophysics can be found in Vetra-Carvalho et al. (2018). The assimilation is implemented using the Parallel Data Assimilation Framework (PDAF2.1), a software environment for data assimilation. PDAF is an open source project and provides fully implemented DA algorithms (Nerger et al., 2020, pdaf.awi.de). The current implementation builds on the works of Mu et al. (2022) who used DA of ocean temperature and salinity for sea-ice forecasts with FESOM2.0
185 coupled to an atmospheric model, and Tang et al. (2020) who studied the dynamic impact of oceanic DA into FESOM1.4 onto a coupled atmospheric component.

With localization of the LESTKF, the observation error is increased for an increasing horizontal distance between an observation and a model grid point, which weighs down the influence of a more distant observation. This avoids that the model is influenced by observations at distant locations through spurious ensemble estimated correlations. We use a localization radius
190 of 200 km and choose a 5th-order polynomial weighting function that mimics a Gaussian function (Gaspari and Cohn, 1999). We apply daily analysis steps at 0 UTC model time, assimilating all available observations for the day. The DA process only

directly updates the physical model variables temperature, salinity, horizontal velocities and sea surface height. After each assimilation step, corrections are applied to the analysis state to ensure the consistency of model physics: Salinity is set to a minimum value of zero and temperature to a minimum value of -2°C , if the value is otherwise below. The increment of sea surface height (SSH) is limited to two standard deviations of the ensemble. While in the simulation the correction was necessary for about 10% of SSH updates and 0.01‰ of temperature values at each step, the correction of salinity was never required. The analysis step is followed by an ensemble forecast of 1 day.

The ensemble size is 40, a compromise to balance computational resources while ensuring a sufficiently large ensemble with enough variability even in the deep ocean. The ensemble is generated through an initial perturbation of sea surface height, horizontal and vertical velocities, temperature, salinity and sea-ice concentration based on the implementation of Tang et al. (2020). This initial ensemble perturbation is generated by second-order exact sampling (Pham, 2001) from a model trajectory of FESOM2.1. With this method, the leading Empirical Orthogonal Functions (EOFs) of a model trajectory are used to generate an ensemble perturbation that contains the leading patterns of model variability. A time-scale must be chosen for the variability that is represented by the ensemble. Here, we chose variability on a weekly time-scale (Tang et al., 2020).

To maintain ensemble spread, we apply a perturbed atmospheric forcing with an autoregressive perturbation ($\text{perturb}_{e,n}$) at every model time step (n) to each ensemble member (e), with:

$$\text{perturb}_{e,n+1} = (1 - \text{arc}) * \text{perturb}_{e,n} + \text{arc} * s * \text{rand}_e \quad (2)$$

where rand is a stochastic element, again generated by second-order exact sampling from a 72-days-long trajectory of atmospheric forcing fields that captures patterns of day-to-day atmospheric variability. The autoregression coefficient (arc) can be used to tune how quickly the perturbation changes and is set to the inverse number of model steps per day. s is a scaling factor for each perturbed atmospheric forcing field. For specific humidity, downwelling longwave radiation and air temperature $s = 10$ is used. The perturbation of winds is set to the smaller value $s = 2$ because the air-sea CO_2 flux in the model is particularly sensitive to perturbations of the wind fields. Due to the functioning of the Kalman filter (which updates the model error covariance in each analysis step to reflect the new reduced uncertainty), the ensemble spread decays at each analysis step. As the method relies on a sufficiently large ensemble spread, an inflation of the ensemble covariance is applied (Pham et al., 1998). Thereby, the ensemble covariance matrix is amplified by a factor of $1/\rho$ before entering the updating step. This so-called forgetting factor downweights that past observations have reduced the model uncertainty (see e.g. Nerger et al., 2005). The forgetting factor is tuned to maintain model uncertainty, where $\rho = 1$ means no inflation and smaller values mean larger inflation. Here, we use a time-varying forgetting factor between $\rho = 0.95$ and $\rho = 1$. The strongest inflation ($\rho = 0.95$) is applied during the first two weeks of the DA process. During the following 75 days ρ is increased to 0.99. From month 17 onwards, the forgetting factor is set to either 0.99 or 1.0 depending on the ensemble standard deviation of temperature. The ensemble standard deviation of the local instantaneous air-sea CO_2 fluxes that results from the perturbation of physical fields is larger than that of the global CO_2 flux, with a mean standard deviation of $0.32 \text{ mmol m}^{-2} \text{ day}^{-1}$ for monthly means of local fluxes compared to a standard deviation of $0.0068 \text{ mmol m}^{-2} \text{ day}^{-1}$ ($0.01 \text{ Pg C yr}^{-1}$) for the annual global flux in FREE in the year 2020. The largest ensemble standard deviation is generated in the Southern Ocean, the North Atlantic and the North Pacific

(map in Fig. A1a), which corresponds to regions of high uncertainty in existing CO₂ flux estimates (Pérez et al., 2024; Hauck et al., 2023a; Mayot et al., 2024). However, the modelled standard deviation should not be understood as the true uncertainty of the model, but as a value dependent on tuning (Evensen, 2003).

2.4 Data analysis

230 We present CO₂ flux estimates for the period 2010-2020, that are compared to the 'Regional Carbon Cycle Assessment and Processes 2' (RECCAP2) global air-sea CO₂ flux estimates (DeVries et al., 2023). The RECCAP2 pCO₂ products account for oceanic outgassing of river carbon into the atmosphere. To make them comparable with our estimate stemming from a model without river carbon input, we apply a river flux adjustment (Friedlingstein et al., 2023; Regnier et al., 2022) to the RECCAP2 pCO₂ products. Thus, we quantify the anthropogenic perturbation of the ocean carbon sink (as S_{OCEAN} in the Global Carbon
235 Budget Friedlingstein et al., 2023; Hauck et al., 2020), and not the contemporary net air-sea CO₂ flux with outgassing of river carbon (as in the original RECCAP2 pCO₂ products).

To study the effect of DA on the CO₂ flux, we define regions where the effect is pronounced and where different mechanisms are active, based on the biomes defined by Fay and McKinley (2014). These are, going polewards from the subtropics in each hemisphere, the Subtropical Seasonally Stratified Biome (STSS), the Subpolar Seasonally Stratified Biome (SPSS) and the Sea-
240 Ice Biome (ICE). In the Southern Ocean (S_O), within the STSS_{S_O}, we differentiate between the area where the assimilation leads to a more positive air-sea CO₂ flux (positive: out of the ocean), referred to as STSS_{S_O}+ and the area where the assimilation leads to a more negative air-sea flux, the STSS_{S_O}- (Fig. 5a and b). In the North Atlantic (N_A), we consider four coherent regions within the STSS_{N_A} and SPSS_{N_A}, defined by the time-mean difference of the air-sea CO₂ fluxes in ASML and FREE (ΔF_{CO_2}). The Central STSS_{N_A}- and Western STSS_{N_A}+ are located in the central North Atlantic STSS_{N_A} biome and are confined by
245 $\Delta F_{\text{CO}_2} < -1 \text{ mmol C day}^{-1} \text{ m}^{-2}$ and $\Delta F_{\text{CO}_2} > 1 \text{ mmol C day}^{-1} \text{ m}^{-2}$, respectively (see Fig. 7b). The Newfoundland Basin+ and Coastal SPSS_{N_A}- are part of the SPSS_{N_A}. The former is located east of Newfoundland and south of Greenland, and is confined by $\Delta F_{\text{CO}_2} > 3 \text{ mmol C day}^{-1} \text{ m}^{-2}$; and the latter is located off the North American coast and confined by $\Delta F_{\text{CO}_2} < -1 \text{ mmol C day}^{-1} \text{ m}^{-2}$. The Central STSS_{N_A}- and Western STSS_{N_A}+ lie on the warm side of the North Atlantic Current (NAC), and the Newfoundland Basin+ and Coastal SPSS_{N_A}- lie on the cold side of the NAC, which is evident from the
250 modeled surface velocity field (Fig. A2a).

Within these regions, we identify the time of the year when the DA affects air-sea CO₂ flux and calculate the difference of ASML and FREE for physical and biogeochemical fields. In order to assess the dynamic DA effects on surface pCO₂, it is useful to distinguish between different variables that constitute the change in pCO₂. Oceanic pCO₂ varies mainly with temperature, DIC and alkalinity. Thus, we decompose changes in pCO₂ into their contributions from changes in SST, surface DIC and
255 surface alkalinity (Alk). For that, we apply the following approximations of Sarmiento and Gruber (2006) and Takahashi et al. (1993):

$$\Delta \text{pCO}_{2,\text{DIC}} = \frac{\text{pCO}_2}{\text{DIC}} * \gamma_{\text{DIC}} * \Delta \text{DIC} \quad (3)$$

$$\Delta p\text{CO}_{2,\text{Alk}} = \frac{p\text{CO}_2}{\text{Alk}} * \gamma_{\text{Alk}} * \Delta \text{Alk} \quad (4)$$

260

$$\Delta p\text{CO}_{2,\text{SST}} = p\text{CO}_2 * \exp(0.0423 * \Delta \text{SST}) \quad (5)$$

Here, differences between ASML and FREE are denoted by Δ ; else, the average of ASML and FREE is used for the computation. The sensitivities γ_{DIC} and γ_{Alk} describe how $p\text{CO}_2$ varies with changes in one variable while keeping all other variables constant. For the sensitivities, we use an approximation derived from seawater carbonate chemistry following Sarmiento and

265 Gruber (2006):

$$\gamma_{\text{DIC}} = \frac{3 * \text{Alk} * \text{DIC} - 2 * \text{DIC}^2}{(2 * \text{DIC} - \text{Alk})(\text{Alk} - \text{DIC})} \quad (6)$$

$$\gamma_{\text{Alk}} = \frac{-\text{Alk}^2}{(2 * \text{DIC} - \text{Alk})(\text{Alk} - \text{DIC})} \quad (7)$$

Based on the range of valid values for γ_{DIC} and γ_{Alk} according to the explicit formulation by Egleston et al. (2010), values
270 are excluded above 18 and below -19, respectively. This affects parts of the Southern Ocean SPSS_{SO} and ICE_{SO} biome (see white areas in Fig. 6b and c). Finally, the effect on the air-sea CO_2 flux relates directly to the $p\text{CO}_2$ -difference at each grid point, as detailed in Orr et al. (2017, equations 6-15):

$$\Delta F_{\text{CO}_2} = \alpha \cdot k_w \cdot \Delta p\text{CO}_2 \quad (8)$$

where α is the solubility of CO_2 in seawater and k_w is the gas-transfer velocity.

275 To evaluate the impact of DA on ocean physics, we compare the simulated SST and SSS to the assimilated observations (Section 2.3.1). For temperature and salinity at depth, we use the EN4-OA product (Good et al., 2013, updated to version 4.2.2). EN4-OA is an objective analysis ingesting the assimilated EN4 profile data, interpolated to global coverage on 42 depth levels. Furthermore, we compare the sea-ice concentration with remote sensing observations from OSI-SAF 2010-2020 (EUMETSAT, 2022), the mixed-layer depth in the year 2020 with the profile-observation based climatology of de Boyer Montégut et al. (2004,
280 updated version 2023) and the horizontal near-surface velocities 2010-2020 with the drifter-based climatology of Laurindo et al. (2017).

To evaluate the impact of the DA on biogeochemistry, we compare model outputs with observational datasets of surface $p\text{CO}_2$, DIC, alkalinity and surface chlorophyll. To evaluate surface $p\text{CO}_2$, we use observations from the Surface Ocean CO_2 Atlas (SOCAT Version 2023, Bakker et al., 2023, 2016), which are provided as a monthly gridded and quality-controlled
285 compilation. To assess DIC and alkalinity, we compare the modeled surface fields to the GLODAPv2.2023 bottle data (Lauvset et al., 2024b). At depth, we compare the model output to the GLODAPv2 DIC and alkalinity climatology (Lauvset et al., 2016),

which is based on observations from the period 1972-2013 and normalized to 2002. To evaluate global surface chlorophyll, we use observations from ESA-CCI, which is a multi-sensor satellite ocean-color chlorophyll-a dataset with monthly global coverage (Sathyendranath et al., 2021). In addition, for the Southern Ocean, we use the mean of three satellite products (Johnson et al., 2013) that were processed with more suitable algorithms for southern high latitudes. For each observation type (OBS), we define the improvement as:

$$\text{improvement}_{\text{OBS}} = |\text{FREE} - \text{OBS}| - |\text{ASML} - \text{OBS}| \quad (9)$$

3 Results

3.1 Effect of DA on ocean physics

Before we investigate the CO_2 flux, we first evaluate the effect of DA on the modeled physics. In particular, we compare the model output of both simulations with the assimilated observations to verify that the assimilation brings them into better agreement with the observations. The assimilation improves the agreement with the assimilated SST observations. On a global average, the SST in FREE is 0.14°C colder than the observations, which is estimated to lead to a solubility-driven global air-sea flux difference of -0.06PgCyr^{-1} (Eqs. (5) and (8)). FREE exhibits an extensive cold bias of SST in the tropics and subtropics in all ocean basins and a warm bias in the Southern Ocean south of 40°S , visible in Fig. 1a; mean state of SST in Fig. A3a). Additionally, FREE shows regional SST biases in particular near strong currents or in eddy-rich regions, such as the NAC, Kuroshio, and the Southern Subtropical Front. The assimilation reduces the SST south of 40°S and in the North Pacific, and increases the SST in the tropics and subtropics (see Fig. 1b). The effect of DA is an absolute change by 0.30°C on global average and is particularly strong in the Southern Ocean and in the North Atlantic. Through the assimilation, the model state becomes more similar to the observations globally, which is evident from the positive improvement in Fig. 1c. In total, the global mean absolute difference of SST to the observations is reduced from 0.59°C to 0.32°C . The assimilation-induced change in SST is estimated to drive a direct solubility-driven effect on the global-air sea CO_2 flux of -0.14PgCyr^{-1} (Eqs. (5) and (8)). Yet, this global attribution is subject to high uncertainty due to the non-linear dependency of pCO_2 on temperature, and because regionally large effects with opposite signs lead to uncertainty in the global mean.

The assimilation also improves the agreement with the assimilated SSS observations. Additional experiments with and without salinity restoring towards climatology show that the best agreement with the SSS-CCI observations is achieved by simultaneously using assimilation and restoring. A benefit of the additional use of restoring is the global coverage of the SSS climatology. FREE shows a global SSS bias (0.49psu , Fig. 1d). The assimilation leads to a global surface freshening (Fig. 1e). There are only a few regions where SSS in FREE is fresher than the observations and where the DA consequently increases the salinity, as for example in parts of the North Atlantic. The assimilation improves the model-observation agreement in 91% of the observed ocean area, particularly in the North Atlantic Central $\text{STSS}_{\text{NA-}}$ and in the Southern Ocean STSS_{SO} (Fig. 1f). Tests with the assimilation of temperature alone show negative side-effects of temperature assimilation on SSS in

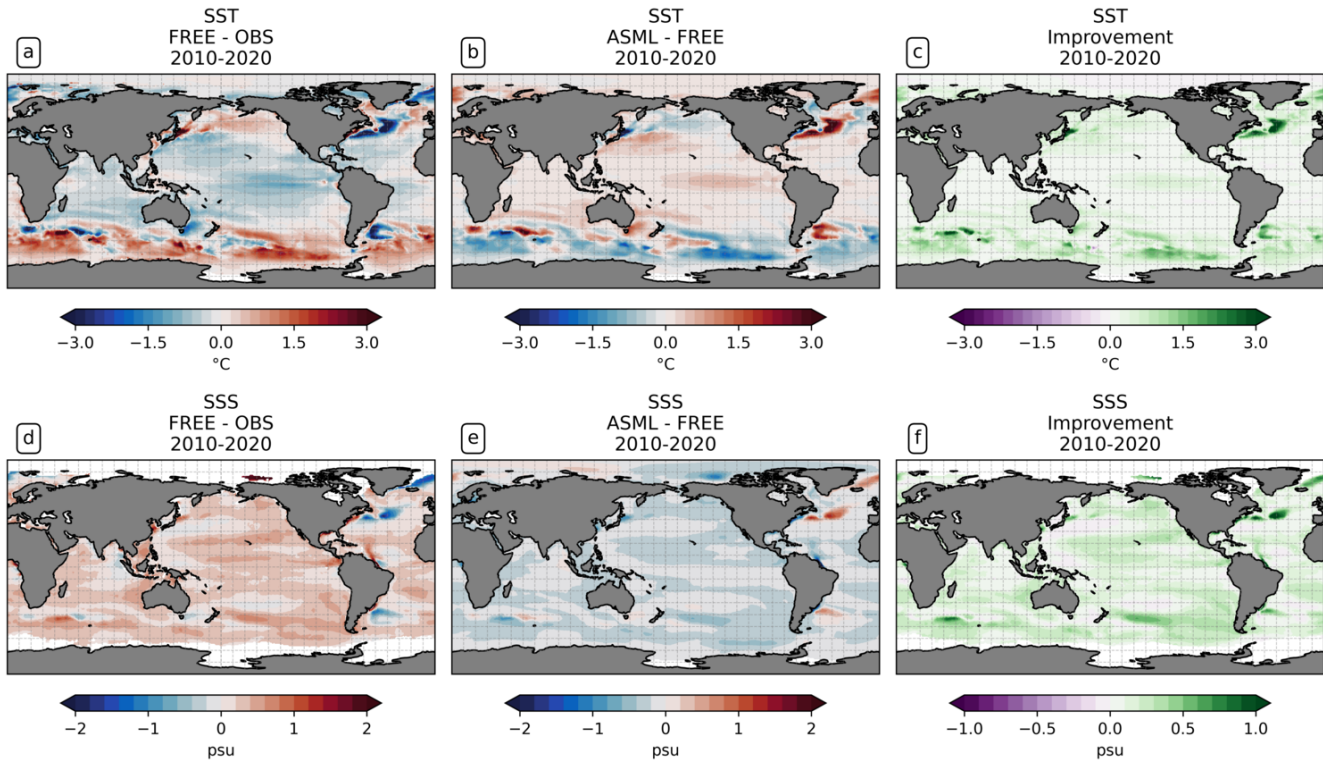


Figure 1. Effect of data assimilation on sea surface temperature (SST) and sea surface salinity (SSS). All panels show the mean over the period 2010-2020. (a) The model-observation difference in SST (FREE - OSTIA). (b) The difference ASML - FREE. (c) The improvement of monthly averaged model SST relative to OSTIA, where positive denotes that the assimilation brings the model closer to observations (Eq. (9)). (d - f) The same for SSS, computed with SSS from ESA-CCI.

some locations (not shown). In the final set-up with combined assimilation, negative effects on SSS are found in 9% of the observed area. Globally, the mean absolute difference is reduced from 0.32 to 0.17 psu relative to the SSS observations. The direct solubility-driven effect of salinity differences on the global air-sea CO₂ flux is estimated to be negligible.

The assimilation leads to a better agreement with subsurface temperature and salinity data from the EN4-OA product in the upper 1000 m. In the upper 100-200 m of the ocean, the model-observation difference in temperature follows the surface signal (compare Fig. 1a and Fig. 2a), and the difference is reduced by the assimilation (Fig. 2b and c). At intermediate depth (roughly 200-500 m), a subsurface warm bias exists in FREE in the southern hemisphere at mid-latitudes (Fig. 2; mean state in Fig. A4a). This bias affects the South Pacific, South Atlantic and southern Indian Ocean (not shown). The bias might be connected to the model's surface warm bias in the formation region of Antarctic intermediate water (Fig. 1a). Further model-observation differences exist at greater depth than 500 m, where the model's temperature is colder than the observations at almost all latitudes, but warmer than the observations north of 60°N. At most latitudes and depths, the effect of the assimilation is to reduce the model observation-differences (Fig. 2c).

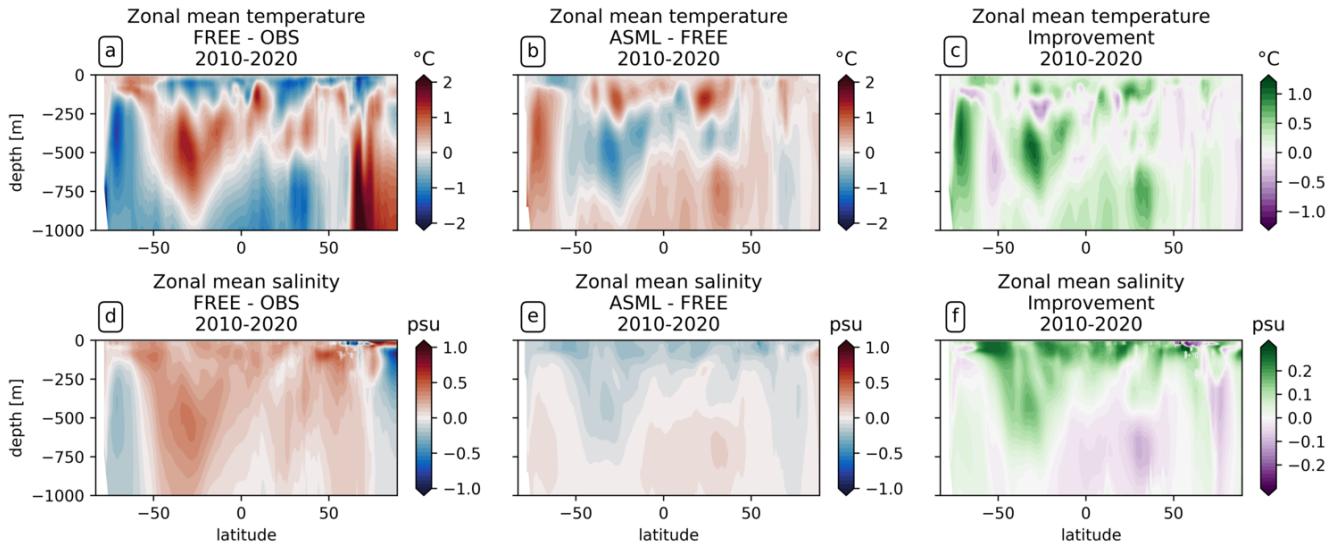


Figure 2. Effect of data assimilation on zonally averaged temperature and salinity in the upper 1000 m. All panels show the mean over the period 2010-2020. (a) The model-observation difference in temperature (FREE - EN4-OA). (b) The difference ASML - FREE. (c) The improvement of monthly averaged temperature relative to EN4-OA. (d - f) The same for salinity.

330 The model is more saline than the observations from the surface down to a depth of about 1000 m for most latitudes (Fig. 2d). This shows that the model-observation difference in this depth range follows the surface signal. The exceptions to this are at high latitudes below 200 m, where FREE is fresher than the observations. At all other latitudes, the assimilation acts towards a freshening, with the strongest effect near the surface (Fig. 2e). This improves the agreement with observations particularly near the surface (Fig. 2d). However, the improvement is smaller at depth and becomes even negative for some latitudes in greater
335 depth. This might be due to the limited amount of assimilated in-situ salinity profiles.

The effect of the assimilation on temperature and salinity is most pronounced in the upper 1000 m and, below that, mostly decreases with depth (not shown). After the second year of assimilation, the mean absolute difference between ASML and FREE stabilizes in the range $0.35 - 0.36^{\circ}\text{C}$ for SST and $0.20 - 0.25$ psu for SSS, while the effect of DA on the 3D fields keeps increasing throughout the years 2010-2020.

340 Sea ice reacts dynamically to the changed ocean physical state. In the Southern Ocean, FREE is characterized by a lower sea-ice concentration compared to OSI-SAF observations. The sea-ice extent, here defined as the area where the sea-ice concentration is more than 15%, reaches a maximum in September. The maximum extent is smaller in FREE than OSI-SAF, which is demonstrated by the 15%-line surrounding that area for FREE and OSI-SAF (Fig. 3a; mean state of sea-ice concentration in Fig. A5), and by the sea-ice concentration difference for the month September (Fig. 3b). Through DA, a higher Antarctic
345 sea-ice concentration is obtained (see Fig. 3b). This improves the agreement with OSI-SAF (Fig. 3c). During all other seasons, the assimilation leads to a higher sea-ice concentration in the Antarctic, a larger sea-ice extent and a better agreement with

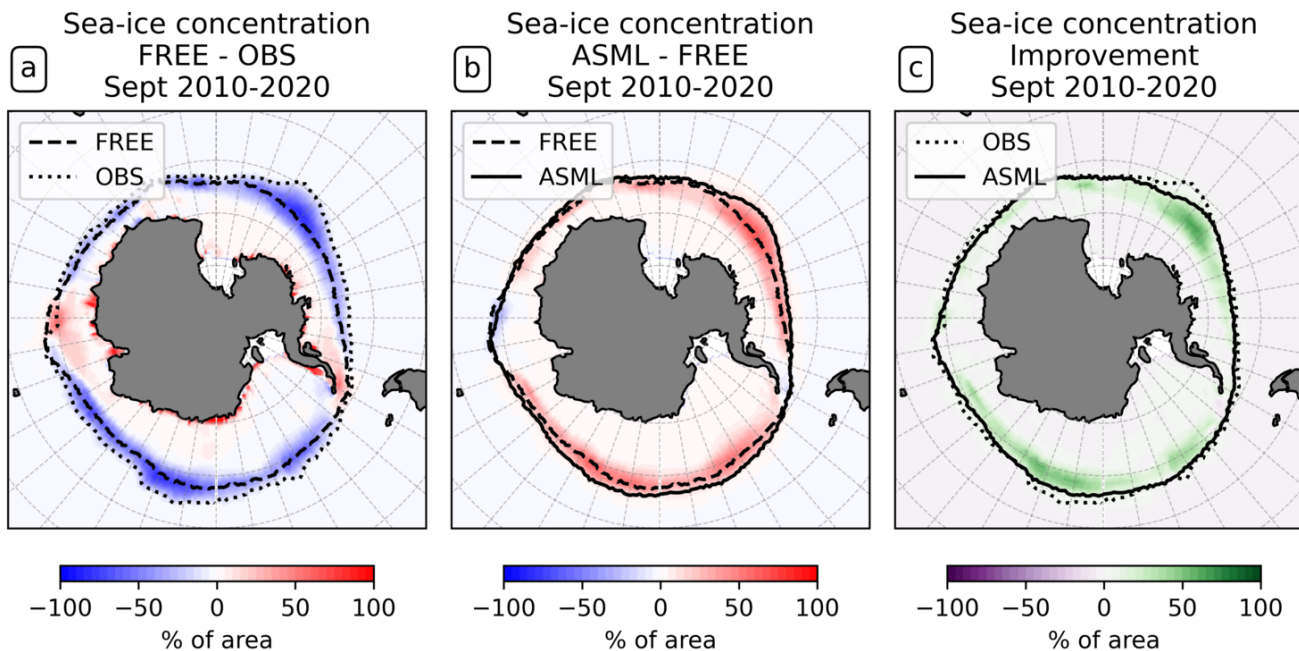


Figure 3. Effect of data assimilation on Antarctic sea-ice concentration in September. All panels show differences in the sea-ice concentration averaged for the month September over the period 2010-2020. The 15%-line for FREE, ASML and OSI-SAF observations is shown as a dashed, continuous or dotted line in panels a or b, respectively. (a) The difference between FREE and OSI-SAF observations. (b) The difference between ASML and FREE. (c) The improvement of September mean sea-ice concentration.

OSI-SAF as well (only September is shown). In the Arctic, the differences between FREE, ASML and OSI-SAF are regionally different (not shown).

The boundary-layer depth and mixed-layer depth are mostly reduced through DA. In particular, deep water formation events characterised by a mixed-layer depth of more than 1000 m or 500 m occur less frequently in ASML (not shown). This improves the agreement with the profile-observation based mixed-layer climatology of de Boyer Montégut et al. (2004), reducing the mean absolute difference to the climatology from 27 m to 19 m (comparison of mixer-layer depth in Fig. A6). In addition, the absolute difference of near-surface horizontal velocities to the drifter-observation based climatology of Laurindo et al. (2017) is reduced by about 10% through DA (comparison of surface velocities in Fig. A7). The biological productivity near the equator is stable in ASML and FREE, indicating that FESOM-REcoM does not suffer from the erroneous upwelling known from previous DA studies (Park et al., 2018). The meridional overturning, however, shows spurious structures, which may point to hidden assimilation artifacts on vertical velocities. Throughout the assimilation period, spurious, spatially limited and often deep overturning structures emerge, evolve through several months or years, and disappear in the tropical Indian, Pacific and Atlantic basin (not shown). Thereby, the surface overturning cell sometimes breaks apart where it should extend over the equator, exposing the bottom cell to the surface (Fig. A8b). Transport in the North Atlantic at 26.5°N, an indicator for the

strength of the Atlantic Meridional Overturning Circulation, is between 8-9 Sv in FREE. In ASML, during the first two years of assimilation, transport at 26.5°N decreases to below 3 Sv and, during the following years, recovers to 7-8 Sv (2016-2020). One possible cause is the effect of data assimilation on the eddy parameterisation (Gent and McWilliams, 1990). The parameterised eddy activity is relevant for the dynamics in the deep ocean, and corrupting it may have a negative impact on the large-scale oceanic circulation, as described in Sidorenko (2004, Chapter 5.5 onwards) for a previous version of the ocean model FESOM.

In summary, the ASML temperature and salinity fields from the surface to several hundred meters below, and mixed-layer depth are in good agreement with observations, and the agreement of horizontal near-surface velocities with observations is improved. This can be interpreted as an indication that the velocity field in the upper part of the ocean is also well represented. Although the spurious effects on deep ocean circulation should be further addressed in future work, we are confident that the DA provides an improved physical state in the upper ocean, which serves as an improved basis to estimate the air-sea CO₂ flux.

3.2 Effect of DA on global CO₂ flux

The ocean absorbs 2.78 Pg C_{yr}⁻¹ in ASML and 2.83 Pg C_{yr}⁻¹ in FREE during 2010-2020 (Fig. 4b), thus the assimilation decreases the global mean oceanic CO₂ uptake by 0.05 Pg C_{yr}⁻¹. The temporal evolution of the annual global CO₂ flux is similar in ASML and FREE (Fig. 4a). The first assimilation year, 2010, stands out because it is one of the very few years during which the assimilation increases the oceanic CO₂ uptake. This slightly reduces the trend in CO₂ uptake 2010-2020 from -0.40 ± 0.09 Pg C_{yr}⁻¹ dec⁻¹ in FREE to -0.38 ± 0.11 Pg C_{yr}⁻¹ dec⁻¹ in ASML (negative: into the ocean). The trend, thereby, remains within its confidence interval. Furthermore, the assimilation reduces the interannual variability of the global mean oceanic uptake slightly, demonstrated by a standard deviation of detrended annual means of 0.11 Pg C_{yr}⁻¹ in FREE and 0.08 Pg C_{yr}⁻¹ in ASML (not significantly different according to F-test). Through DA, the ensemble standard deviation of the global CO₂ flux is reduced from 1.0×10^{-2} Pg C_{yr}⁻¹ in FREE to 0.7×10^{-2} Pg C_{yr}⁻¹ in ASML in the year 2020.

The strongest time-mean air-sea CO₂ flux is found at mid and high latitudes (Fig. 4c). The large-scale pattern of the CO₂ flux is generally very similar in FREE and in ASML (FREE not shown). The largest local changes through DA, both towards stronger or weaker CO₂ fluxes, occur in the North Atlantic in the area of the NAC and in the coastal North Pacific (Fig. 4d). The most prominent large-scale effect though, is in the Southern Ocean (Fig. 4e and f). South of 50°S, the area-integrated CO₂ uptake increases by 0.18 Pg C_{yr}⁻¹ through the assimilation. In contrast, the uptake decreases by 0.07 Pg C_{yr}⁻¹ between 40-50°S. With the exception of the Southern Ocean, CO₂ uptake decreases in all world oceans by a small amount (Fig. 4d).

3.3 Effect of DA on regional CO₂ fluxes and their drivers

3.3.1 Southern Ocean

In the Southern Ocean, the ocean takes up CO₂ in the annual average (Fig. 5a), with regionally heterogeneous effects of DA (Fig. 5b). While the effect of DA on surface pCO₂ and the air-sea CO₂ flux can almost entirely be explained by the combined variation of DIC and alkalinity at most latitudes north of 40°S, the thermal effect also needs to be considered in the Southern

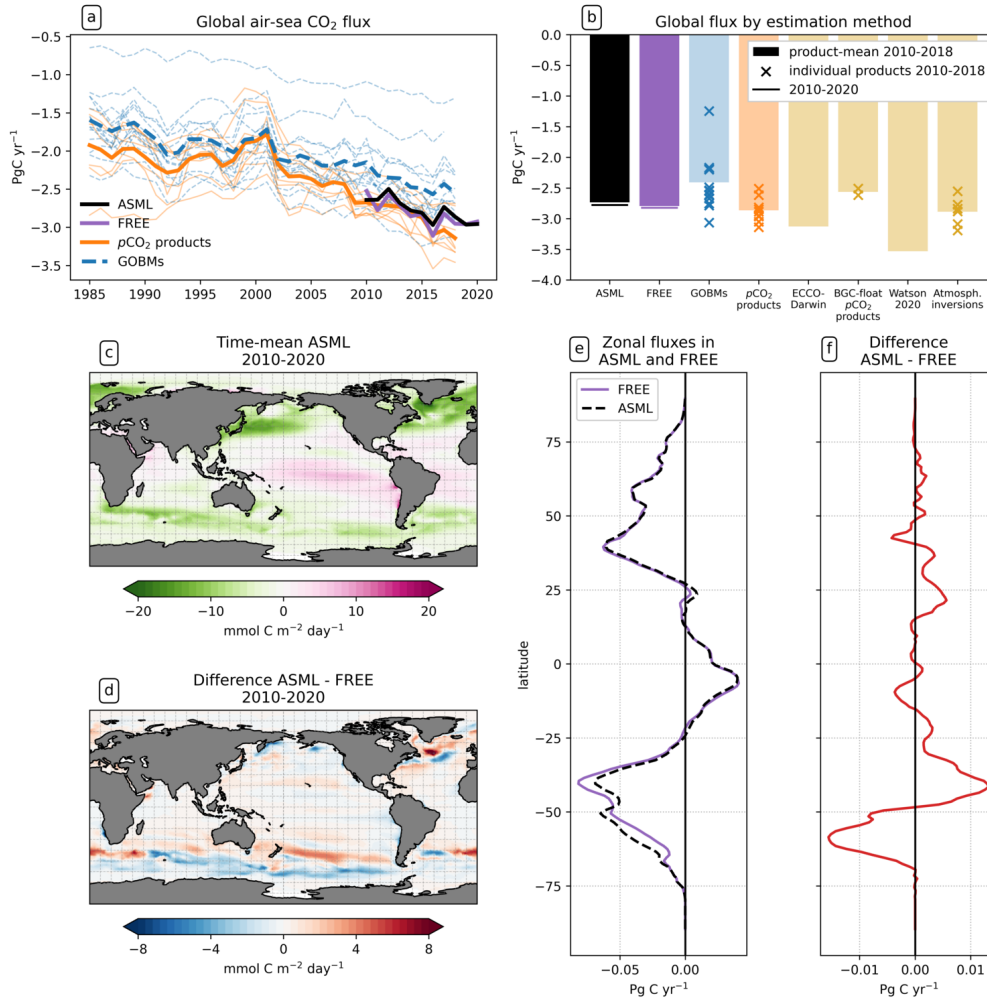


Figure 4. Effect of data assimilation on the air-sea CO₂ flux (negative: into the ocean). (a) Annual time-series of global flux in PgC yr⁻¹ in FESOM-RECoM-PDAF with ASML (black) and FREE (violet); and RECCAP2 estimates (DeVries et al., 2023) with pCO₂-products (orange) and GOBMs (blue) and their respective means (bold lines). Here, the river flux adjustment ($-0.65 \text{ PgC yr}^{-1}$) was applied to the pCO₂ products. (b) Time-mean global flux 2010-2018 in ASML (black), FREE (violet); and RECCAP estimates grouped by method (DeVries et al., 2023). Crosses represent individual estimates (e.g. individual GOBMs) and bars represent the method mean (e.g. mean of twelve GOBMs). Here, the river flux term was applied to all estimates except the models following the Global Carbon Budget methodology (Friedlingstein et al., 2023). For FESOM-RECoM-PDAF, additionally the time-mean 2010-2020 is shown (horizontal lines). (c) Spatial distribution of CO₂ flux averaged over the period 2010-2020 in ASML. (d) Spatial distribution of CO₂ flux difference between ASML and FREE averaged over the period 2010-2020 (e) Zonal averages of CO₂ flux 2010-2020 in ASML and FREE, and their difference in (f).

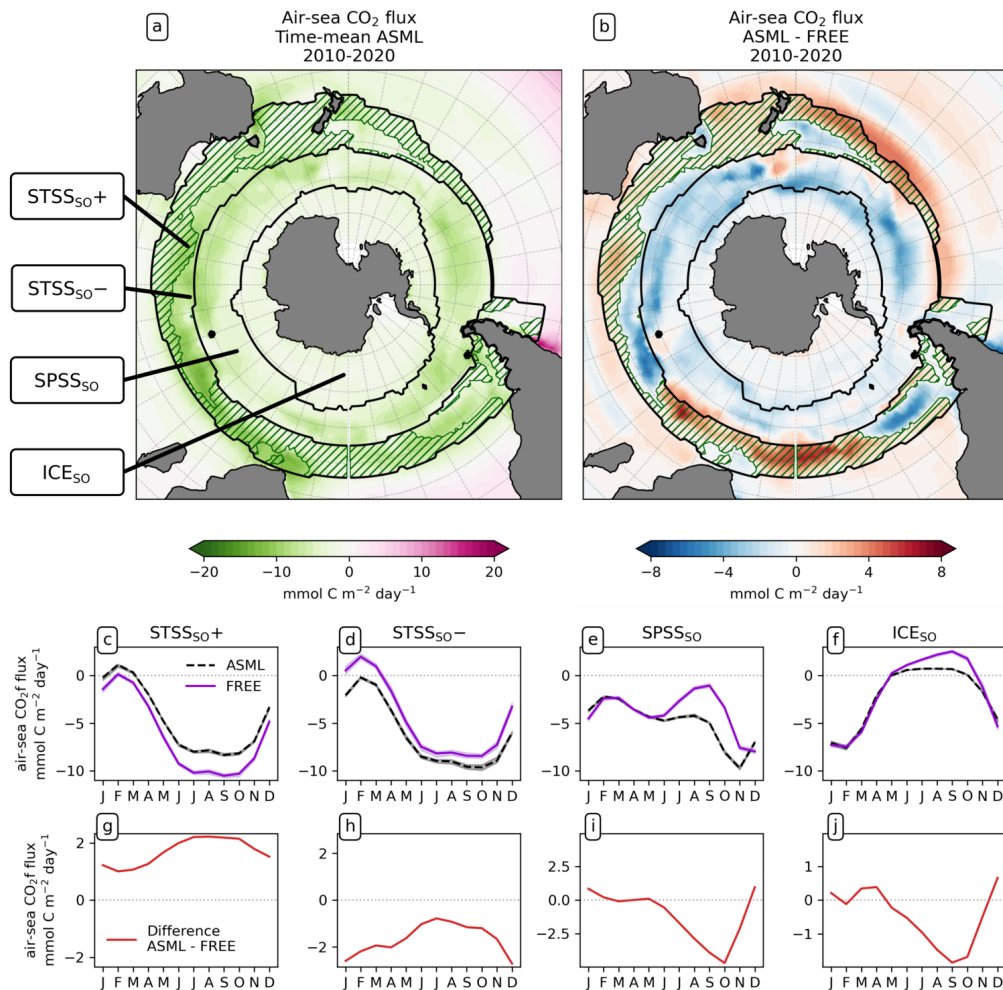


Figure 5. Effect of data assimilation on Southern Ocean CO₂ flux and its seasonality averaged over the period 2010-2020. Negative numbers indicate a flux into the ocean. Additionally, lines in a and b denote the regions, and the green hatching denotes the STSS_{S0+}. (a) Map of mean CO₂ flux in ASML. (b) Map of difference between CO₂ flux in ASML and FREE. (c - f) Seasonal cycle of air-sea CO₂ flux by region. Shading indicates the range of ensemble members in the year 2020. (g - j) Seasonal difference in air-sea CO₂ flux between ASML and FREE by region. Note the different scales.

Ocean (global zonal mean pCO₂-effects in Fig. A9a). In the following, we examine how the assimilation influences the air-sea CO₂ flux across individual regions in the Southern Ocean.

STSS_{S0} In the northernmost biome of the Southern Ocean, the subtropical seasonally stratified biome (STSS_{S0}), the mean oceanic CO₂ uptake is comparably high (Fig. 5a). The uptake is largest in austral winter and spring (June to November, Fig. 5c and d). The part of the STSS_{S0} characterized by a positive CO₂ flux difference between ASML and FREE (positive difference: reduced uptake through assimilation), which we call the STSS_{S0+}, roughly forms an outer northerly ring around the STSS_{S0}

biome (hatched area in Fig. 5a and b). The reduction of CO₂ uptake in the STSS_{SO+} is greatest in winter and spring from July to October (Fig. 5g).

400 The increase in pCO₂ in the STSS_{SO+} is partly driven by lowered alkalinity and partly by increased surface DIC (Fig. 6b and c). These, as well as the colder SST and fresher SSS in the STSS_{SO+} (Fig. 1b and e) are indications for a year-round stronger influence of subantarctic waters. This is evident from typical water properties in the subantarctic and subtropical Southern Ocean. In the subantarctic, surface DIC is higher, surface alkalinity is lower, temperature is colder and salinity is lower (maps of mean SST, SSS, DIC and alkalinity in Fig. A10). In the fragmented area of the STSS_{SO+}, different factors
405 contribute to regional changes of the surface DIC and alkalinity budget in ASML (sources minus sinks of DIC and alkalinity in Fig. A11). Depending on location, an increased upward transport of DIC through mixing, an increase of DIC through a reduced biological sink of DIC in spring, or a decrease of alkalinity through changes in horizontal and vertical advection dominates. The seasonality of the effect of DA on the air-sea CO₂ flux in the STSS_{SO+} (Fig. 5c and g) is determined by seasonal temperature differences between ASML and FREE (Fig. 6d and f). During summer, SST is slightly reduced (Fig. 6f), which lowers pCO₂
410 (Fig. 6a). This counteracts the effects of DIC and alkalinity on pCO₂ (Fig. 6b and c) and thus dampens the overall DA-effect on the air-sea CO₂ flux during summer.

The part of the STSS_{SO} characterized by a negative CO₂ flux difference between ASML and FREE, which we call the STSS_{SO-}, is a fragmented region and roughly consists of segments of an inner southerly ring (non-hatched area in Fig. 5a and b). Here, the increase of CO₂ uptake through DA is largest in summer and autumn (November to April, Fig. 5h). The reduction
415 of pCO₂ is driven by increased alkalinity, and partly also by lower surface DIC (Fig. 6b and c, non-hatched area). These, together with higher SST in ASML than FREE in the STSS_{SO-} regions (Fig. 1b), indicate a higher presence of subtropical waters (see characteristics of subtropical waters in Fig. A10). Where there is lower DIC in the STSS_{SO-} in ASML (Fig. 6b), this can mostly be explained by an increased biological sink of DIC, with the addition of sharply defined local changes in horizontal advection of DIC and alkalinity (Fig. A11). Additionally, seasonal temperature effects occur. During winter, SST
420 is higher in ASML than in FREE (Fig. 6e and g). This increases pCO₂ in the STSS_{SO-} (Fig. 6a), counteracting the effects of lower DIC and higher alkalinity on pCO₂ and dampening the overall DA-effect during winter.

The contrasting effects in the STSS_{SO} indicate a horizontal shift of water masses within the STSS_{SO} biome. In the center of the STSS_{SO}, the Subantarctic Front is located, which is associated with the Antarctic Circumpolar Current (ACC) and characterized by a strong gradient in SST, SSS and various other tracers (Chapman et al., 2020). Because SST and SSS are
425 directly influenced and improved by the assimilation, the position of this front is also expected to change as a result of the assimilation, leading to a horizontal relocation of waters separated by the front. With the relocation of the front, dynamic shifts in regional characteristics occur, such as the amount of DIC and alkalinity transported vertically through mixing, and biological sources and sinks of DIC and alkalinity.

SPSS_{SO} Further south, in the subpolar seasonally stratified biome (SPSS_{SO}), the ocean absorbs CO₂ all year-round
430 (Fig. 5a). The oceanic uptake is increased through the assimilation, shown by a negative difference of ASML and FREE in Fig. 5b. The largest difference between ASML and FREE is seen in spring from September to October (Fig. 5i). Due to the seasonally varying effect of DA, the seasonal cycle of the CO₂ flux in the SPSS_{SO} is altered. In ASML, the CO₂ uptake is

weakest in February, gets stronger in autumn (MAM), stagnates in winter (JJA) and resumes to grow in spring (SON), reaching peak uptake in November (Fig. 5e). In FREE, the CO₂ uptake weakens in winter, is weakest in September and gets stronger afterwards, reaching peak uptake in December.

In the SPSS_{SO}, the increased CO₂ uptake and lower surface pCO₂ during winter and spring is driven by a combination of colder temperatures and lower DIC (Fig. 6a and b), which outweighs the opposite effect of a decrease in alkalinity on pCO₂ (Fig. 6c, relative importance of thermal effect in Fig. A12a). Surface DIC is generally high due to upward transport of carbon-rich deep water (e.g. Hauck et al., 2023a). The reason for lower surface DIC in ASML is that the upward transport through mixing is reduced (Fig. A11) through a more stable stratification, which is also evident from a reduced density in the upper 300 m and an increased density below that (Fig. 6h). Thereby, the densities in the SPSS_{SO} agree better with densities calculated from EN4-OA. Boundary layer and mixed layer in winter and spring are shallower and thereby in better agreement with the observation-based climatology (Fig. A6). Vertical mixing within the boundary layer affects the vertical profiles of DIC and alkalinity, towards lower DIC in ASML above 100 m and higher DIC below (Fig. 6i). The vertical profile of DIC in ASML is closer to GLODAP DIC observations, albeit some differences to GLODAP still exist. Besides the fact that the differences in stratification and boundary-layer depth affect the vertical DIC profile, they also imply less available surface nutrients in ASML. Probably due to a combination of lower nutrient availability and colder surface temperature, ASML features lower NPP, lower chlorophyll concentrations and a lower phytoplankton biomass in the SPSS_{SO} (not shown). Thereby, the modeled biogeochemical cycle adjusts to the lower transport of nutrients to the surface by transferring less organic material to depth, ultimately acting to compensate about 60% of the difference in physical transport of DIC (Fig. A13a) and adding to the reduction in surface alkalinity (Fig. A13b). Within the SPSS_{SO} (roughly south of 50 °S), differences between FREE and ASML in terms of the temperature effect on pCO₂, vertical transport of DIC and alkalinity and biological sources and sinks are larger than at any other latitude (Fig. A13).

ICE_{SO} In the seasonally ice-covered biome (ICE_{SO}) surrounding the Antarctic continent, the time-mean CO₂ flux is smaller than in other biomes (Fig. 5a). In this region, the ocean absorbs CO₂ during summer and there is a smaller outgassing during winter (Fig. 5f), as the region is mostly ice-covered in winter (see sea-ice concentration in September in Fig. 3). In the northern part of the ICE_{SO} biome, close to the SPSS_{SO}, the effect of the assimilation is similar to the effect within the SPSS_{SO} itself (Fig. 5b). Here, the assimilation acts to increase ocean CO₂ uptake or to weaken CO₂ outgassing during winter and spring (Fig. 5i and j). Thereby, interestingly, the assimilation hinders outgassing of CO₂ from May to November in ASML in the ICE_{SO} biome (Fig. 5f; comparison of winter outgassing with other estimates in Fig. A14). The reduced outgassing and decreased pCO₂ during winter and spring is driven by similar processes as within the SPSS_{SO}. Again, lower surface DIC and colder temperatures (Fig. 6a and b) outweigh the opposite effect of a decrease in alkalinity on pCO₂ (Fig. 6c). As in the SPSS_{SO}, the reason for the decrease in pCO₂, is reduced surface DIC and increased DIC below 100 m as a result of less upward transport of DIC through mixing (Fig. A11) in a more stable stratification due to surface freshening (Fig. 1e). In addition, as the surface temperature is lower in ASML (Fig. 1b), the winter sea-ice concentration is higher (Fig. 3b), which prevents winter outgassing of CO₂. In the southern part of the ICE_{SO} biome, near the Antarctic continent, the effect of the DA on the CO₂ flux is small.

In summary, in the Southern Ocean, the main effects of the DA on the CO₂ flux are, firstly an increase of the uptake in the SPSS_{SO} caused by surface cooling and by a more stable stratification and thus less upward transport of naturally carbon-rich water through mixing, and secondly an overall lower CO₂ uptake in the STSS_{SO} as a consequence from a spatial redistribution of fluxes near the Subantarctic Front.

3.3.2 North Atlantic

In the North Atlantic, the assimilation has noticeable effects on the CO₂ flux in the area of the North Atlantic Current, where the ocean absorbs CO₂ in the annual average (Fig. 7a). During summer however, the ocean releases CO₂ while the sea surface warms (Fig. 7c-f). In the Central STSS_{NA-}, the effect of the DA is to prevent outgassing during summer (Fig. 7c and g). In the Western STSS_{NA+} and in the Newfoundland Basin+, the ocean CO₂ uptake is decreased during winter (Fig. 7d, e, h and j). The regionally different dynamics of the effects of the assimilation that drive these differences in the air-sea CO₂ flux in the North Atlantic are investigated next.

Central STSS_{NA-} In the Central STSS_{NA-}, the effect of the DA is overall towards a more negative flux of CO₂ from May to November (Fig. 7g). Thus, spring and autumn CO₂ uptake are increased and summer outgassing is prevented in ASML (Fig. 7c). The reason for decreased surface pCO₂ is higher alkalinity in ASML (Fig. 8c). In this region, the alkalinity effect, which reduces pCO₂, outweighs the opposing effects of DIC and SST on pCO₂ (Fig. 8a and b). A higher alkalinity could point to the presence of waters of subtropical origin transported northward with the NAC (Völker et al., 2002). Other fingerprints of waters transported by the NAC are a warm SST particularly in winter, a higher salinity and higher DIC than that of North Atlantic subpolar waters (maps of mean SST, SSS, DIC, alkalinity in Fig. A15; Völker et al., 2002). The assimilation causes a change in these properties, towards a higher SST, higher salinity and higher DIC in the Central STSS_{NA-}. Simultaneously, ASML represents a deeper boundary layer in this region (Fig. 8d). While changes in the North Atlantic mixed-layer depth overall result in a spatial pattern in ASML that more closely aligns with the pattern in the observation-based mixed-layer climatology, the modelled mixed layer in the simulations is still overall deeper than in the climatology, leading to less agreement in the Central STSS_{NA-} (Fig. A6). Likely facilitated by higher SST and more available nutrients through deeper mixing in winter and spring, ASML features a higher biological sink of DIC above 190 m (Fig. A16d), more biological carbon export through sinking of detritus at 190 m, more column integrated phytoplankton biomass and surface chlorophyll in spring, which is illustrated by the example of surface chlorophyll difference between ASML and FREE in Fig. 8e. In combination, the higher alkalinity associated with NAC transport and the higher biological sink of DIC result in lowered surface pCO₂ and higher oceanic uptake.

Western STSS_{NA+} In the Western STSS_{NA+}, the DA reduces the CO₂ uptake and increases pCO₂ mainly during winter, as a direct effect of increased SST (Fig. 8a). The direct thermal effect is dominant over the combined effect of DIC and alkalinity (relative importance of thermal effect in Fig. A12b). The latter have effects comparable in magnitude to SST, but mostly cancel each other out (Fig. 8b-c). The effect of DA on surface properties (SST, SSS, DIC and alkalinity) in the Western STSS_{NA+} is similar to the effect in the Central STSS_{NA-}, which indicates a higher influence of subtropical waters in both regions.

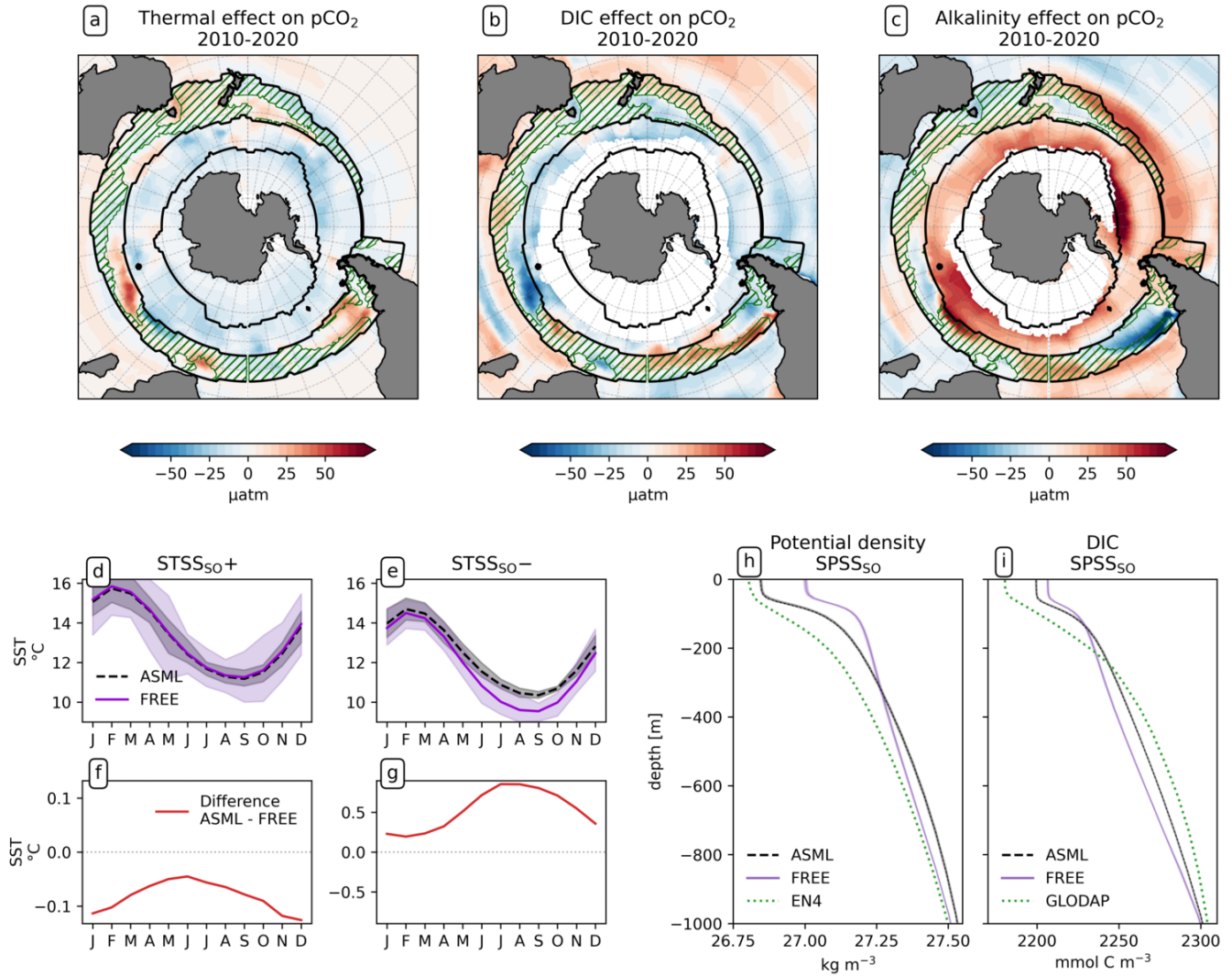


Figure 6. Drivers of the effects of data assimilation on air-sea CO_2 fluxes in the Southern Ocean. Panels a, b and c show the effects of SST, DIC and alkalinity differences between the ASML and FREE simulations on surface pCO_2 , where positive denotes an increase in pCO_2 . Hatching inside the STSS_{SO} indicates where net pCO_2 is increased through the assimilation ($\text{STSS}_{\text{SO}+}$). (d and e) Seasonal cycle of SST averaged over the regions $\text{STSS}_{\text{SO}+}$ and $\text{STSS}_{\text{SO}-}$, and (f and g) the difference between ASML and FREE for each region. (h) Potential density profiles for the SPSS_{SO} , with FREE (violet line) and ASML (dashed black line) based on daily T and S, and with EN4-OA (dotted green line) based on monthly T and S. (i) DIC profiles for the SPSS_{SO} , showing FREE (violet line), ASML (dashed black line) from 2010-2020 and climatological DIC from GLODAP. Shading in d, e, h and i indicates the range of ensemble members in the year 2020.

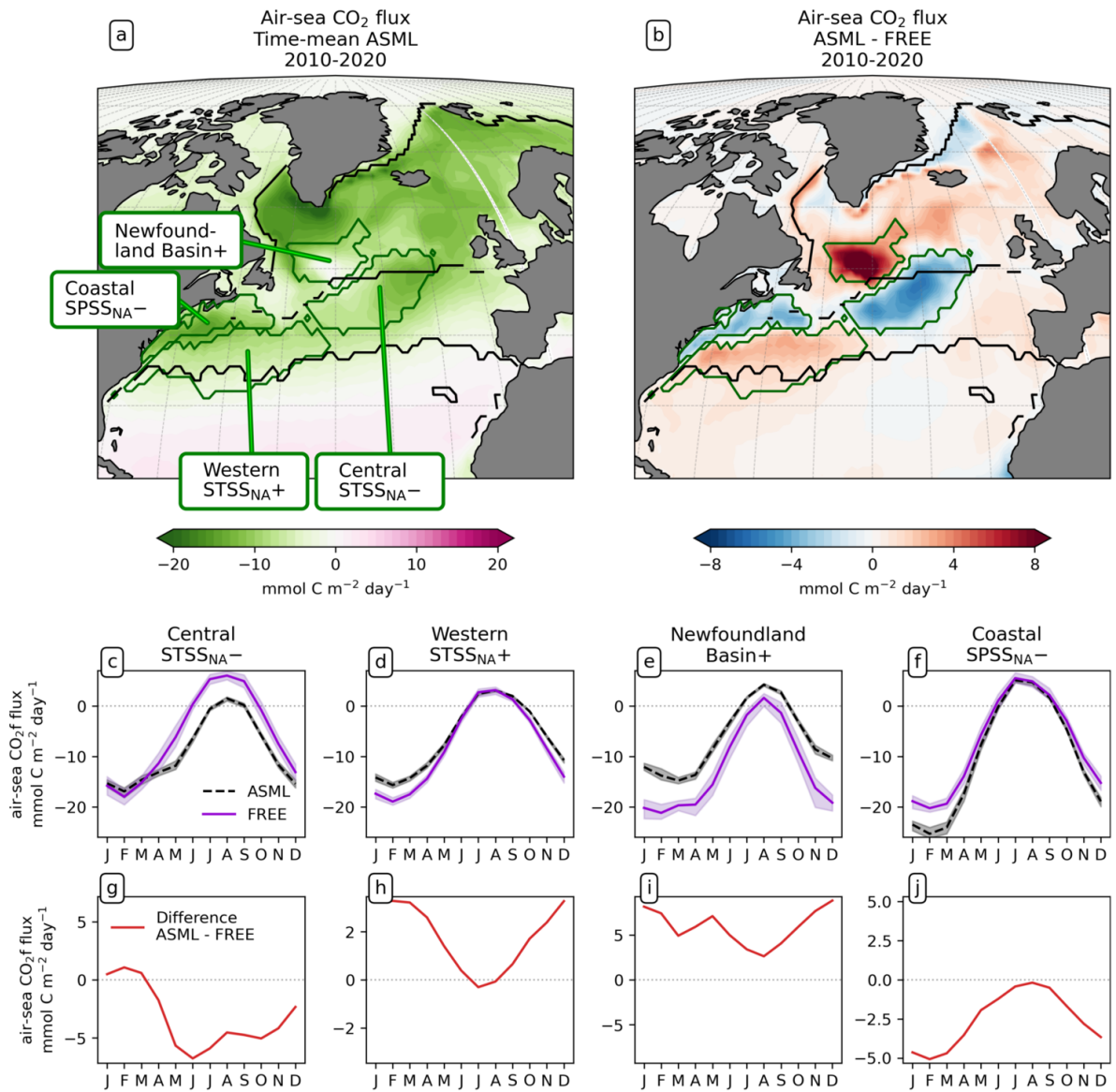


Figure 7. Effect of data assimilation on North Atlantic CO₂ flux and its seasonality averaged over the period 2010-2020. Negative numbers indicate a flux into the ocean. (a) Map of mean CO₂ flux in ASML. (b) Map of difference between CO₂ flux in ASML and FREE. (c - f) Seasonal cycle of air-sea CO₂ flux by region. Shading indicates the range of ensemble members in the year 2020. (g - j) Seasonal difference in air-sea CO₂ flux between ASML and FREE by region. Note different scales.

Newfoundland Basin+ In the Newfoundland Basin+, the dominant effect of DA is a reduction of the CO₂ uptake and an increase of pCO₂ mainly during winter, as a direct effect of increased SST (Fig. 8a). In addition, ASML also features a more stable stratification due to lower density at the surface than FREE (Fig. 8f), which mostly affects DIC at 50-400 m depth through reduced subduction of DIC (Fig. 8g). Furthermore, ASML represents less surface chlorophyll in the Newfoundland Basin+ (Fig. 8e) as a result of a redistribution of biomass from the surface to 50-400 m depth due to spring mixing (not shown). The downward mixing of biomass results in an increase of the biological sink of DIC above 50 m likely due to more primary production near the surface, and a decrease of the biological sink at 50-400 m likely due to more remineralization at this depth. However, the differences in the biological sink of DIC are compensated by mixing of DIC (profiles not shown). Overall, differences of the regional DIC profile to the observational GLODAP climatology slightly increase (Fig. 8g).

Coastal SPSS_{NA-} In the Coastal SPSS_{NA-}, pCO₂ is reduced and the ocean CO₂ uptake is increased in ASML during winter and spring (Fig. 7f and j). The reduction of pCO₂ is facilitated by colder SST (Fig. 8a). This might be due to subpolar water masses penetrating further south along the coast in ASML because the location where the current separates from the coast is further south in ASML (velocities in Fig. A2).

In summary, DA affects the CO₂ flux in the North Atlantic mainly through changes in SST, combined with changes in horizontal advection of DIC and alkalinity near the NAC. Changes in the vertical mixing of DIC and alkalinity are largely compensated by feedbacks in biogeochemical cycles. Which of these effects is dominant, however, varies from region to region.

3.4 Comparison with biogeochemical observations

3.4.1 pCO₂ (SOCAT)

To evaluate the modeled air-sea CO₂ flux based on observations, surface pCO₂ is the most informative variable, as it is closely related to the air-sea CO₂ flux. Effects of the DA on the modeled ecosystem and associated carbon fluxes, as well as thermal and dynamical effects that affect the CO₂ flux, are all included in pCO₂. The global mean of absolute monthly model-observation differences to the available SOCAT pCO₂ observations is 27.26 μatm for FREE. For ASML, the difference is slightly larger with 27.60 μatm. On global average, pCO₂ is higher than in SOCAT by 3.70 μatm in FREE and 4.59 μatm in ASML, as regions with positive and negative differences to SOCAT compensate (Fig. 9a). As an illustration of the regional changes through DA, the absolute differences in pCO₂ amount to 8.08 μatm (absolute difference ASML-FREE calculated at every grid point then averaged globally), which is ±27% of the mean absolute model-observation difference. A linear offline estimation demonstrates that this change in pCO₂ would lead to an absolute change in the air-sea CO₂ flux by 1.06 mmol C m⁻² day⁻¹ on average (Eq. (8)).

Overall, FREE and ASML show very similar regional pCO₂ differences compared to SOCAT (difference of FREE and SOCAT in Fig. 9a; difference of ASML and SOCAT not shown). In the subtropical and tropical Atlantic and the subtropical Pacific, FREE and ASML have higher pCO₂ than SOCAT, while in the equatorial Pacific, pCO₂ is lower. At high latitudes, FREE and ASML represent mostly lower pCO₂ than SOCAT.

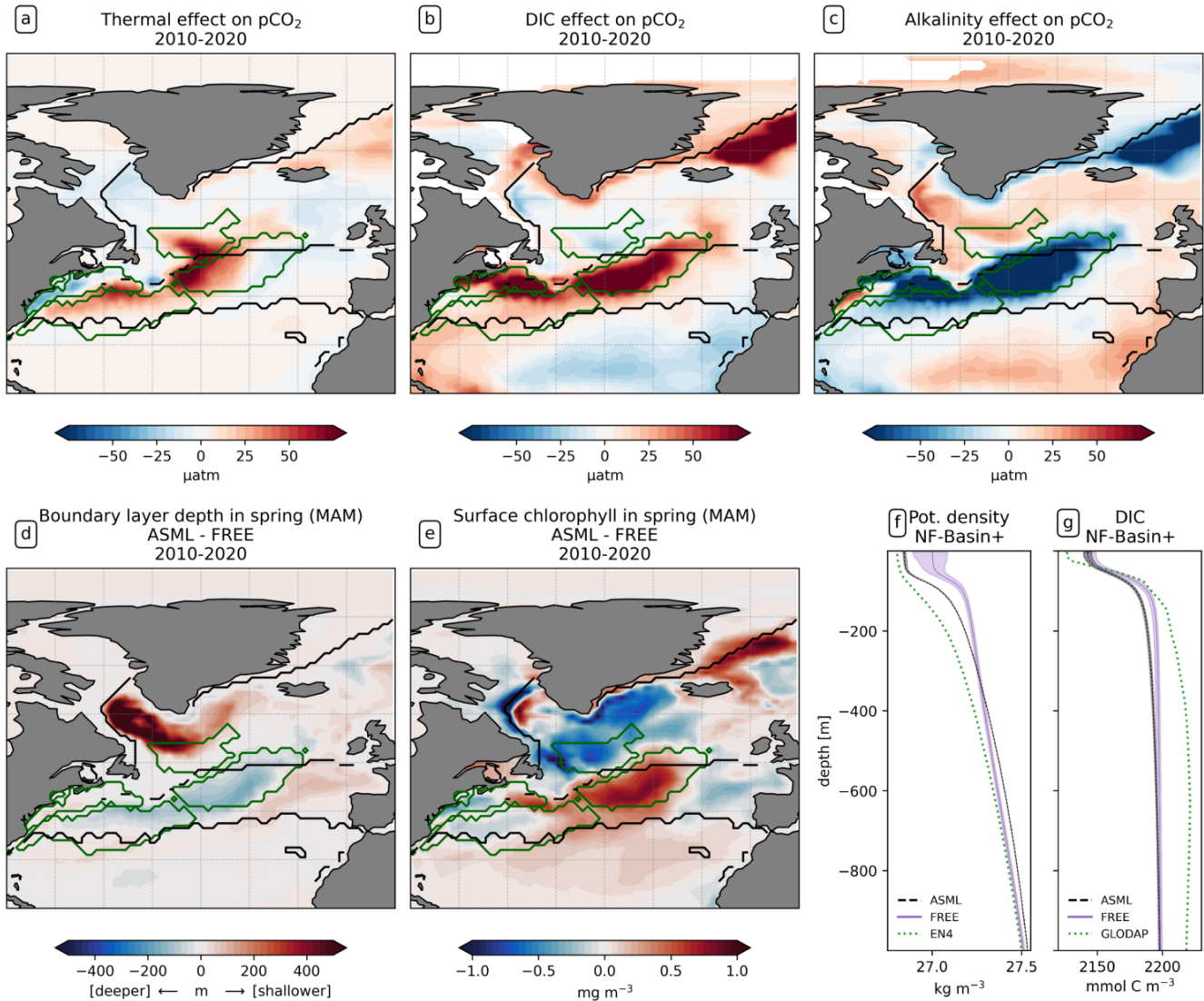


Figure 8. Drivers of the effects of data assimilation on air-sea CO₂ flux in the North Atlantic. Panels a, b and c show the effects of SST, surface DIC and alkalinity differences between ASML and FREE on surface pCO₂. (d) Difference of boundary layer depth (ASML - FREE) for spring (MAM) 2010-2020, where positive denotes a shallower boundary layer in the ASML. (e) Difference of surface chlorophyll (ASML-FREE) for spring (MAM) 2010-2020. (f) Potential density profiles for the Newfoundland Basin+ region, with FREE (violet line) and ASML (dashed black line) based on daily T and S, and with EN4-OA (dotted green line) based on monthly T and S. (g) DIC profiles for the Newfoundland Basin+ region, showing FREE (violet line), ASML (dashed black line) from 2010-2020 and climatological DIC from GLODAP. Shading in f and g indicates the range of ensemble members in the year 2020.

In the Southern Ocean, the simulations represent lower $p\text{CO}_2$ than SOCAT in the SPSS_{SO} and ICE_{SO} biomes in the annual mean (Fig. 9c), which is dominated by summer differences to SOCAT (not shown) when most observations are available. Through the assimilation, $p\text{CO}_2$ is slightly increased in summer and mostly reduced in winter (not shown), leading to an overall better agreement with SOCAT (Fig. 9e). In contrast, in the STSS, FREE and ASML represent higher $p\text{CO}_2$ than SOCAT, and through the assimilation, the agreement with SOCAT decreases.

In the North Atlantic, the simulations and SOCAT show a similar large-scale pattern, namely that $p\text{CO}_2$ is higher in the subtropics (ASML: around $400\mu\text{atm}$) than in the subpolar regions (ASML: around $280\mu\text{atm}$). Yet, this latitudinal difference of $p\text{CO}_2$ is stronger in the simulations compared to SOCAT, meaning that in the subtropics, $p\text{CO}_2$ in the simulations is higher than in SOCAT (Fig. 9d), while it is lower in the subpolar regions. Furthermore, in both simulations there is a pronounced $p\text{CO}_2$ surface gradient in the NAC and North Atlantic Subpolar Gyre region, whose position is changed by the assimilation, and which appears to be further northward in SOCAT. Thereby, the assimilation overall leads to a better agreement with SOCAT, in particular through a decrease of $p\text{CO}_2$ in the Central STSS $_{\text{NA}^-}$, where the average difference is reduced from $26\mu\text{atm}$ (FREE - SOCAT) to $1\mu\text{atm}$ (ASML - SOCAT). However, in the Newfoundland Basin $^+$, the average difference is reversed from $-17\mu\text{atm}$ (FREE - SOCAT) into $13\mu\text{atm}$ (ASML - SOCAT), which is associated with a larger absolute discrepancy of ASML and SOCAT.

3.4.2 DIC and alkalinity (GLODAP)

DIC and alkalinity are two of the most important variables from which $p\text{CO}_2$ is derived (Section 3.3). Comparing them with observations provides more insights into the strengths and weaknesses of the modeled carbonate system than a comparison with $p\text{CO}_2$ observations alone. The FESOM-REcoM simulations represent higher surface DIC than GLODAP bottle observations (Lauvset et al., 2024a, gridded monthly-means) on average (Fig. 10a), with a global mean surface difference FREE-GLODAP of 6.46mmol C m^{-3} for DIC. Although fewer DIC observations are available than $p\text{CO}_2$ observations, similarities between the respective model-observations differences for DIC and $p\text{CO}_2$ can be recognized. For example, DIC in the model is lower in the tropical and subtropical Atlantic than GLODAP, and higher in the polar Atlantic. This is consistent with SOCAT $p\text{CO}_2$ observations in the same areas. The model-observation differences to GLODAP DIC and SOCAT $p\text{CO}_2$ are also consistent with each other in the north Pacific. The assimilation induces absolute changes in surface DIC of 6.33mmol C m^{-3} on global average, with regional differences in sign. These changes slightly reduce the mean absolute difference to the surface observations from $32.78\text{mmol C m}^{-3}$ to $32.15\text{mmol C m}^{-3}$, and yield a mixed picture of the improvement (Fig. 10b).

While the trend in surface DIC due to anthropogenic input makes it necessary to compare the model with contemporaneous observations at the ocean surface, a comparison with climatological data is meaningful below a depth of approximately 200 m. In fact, the modeled global distribution of DIC at depth is overall similar to that in the GLODAP climatology for both simulations (zonal mean DIC surface to 1000 m depth in Fig. A17). For example, the model results and GLODAP data sets show that DIC is lowest in the isopycnals of the subtropical gyres ($2050 - 2150\text{mmol C m}^{-3}$; Fig. A17a) and that DIC mostly increases with depth and is higher in the Pacific (2420mmol C m^{-3} at 1000 m in the North Pacific) than in the Atlantic (2320mmol C m^{-3} below 3000 m in the South Atlantic). Yet, depending on the ocean basin and depth, there can be both nega-

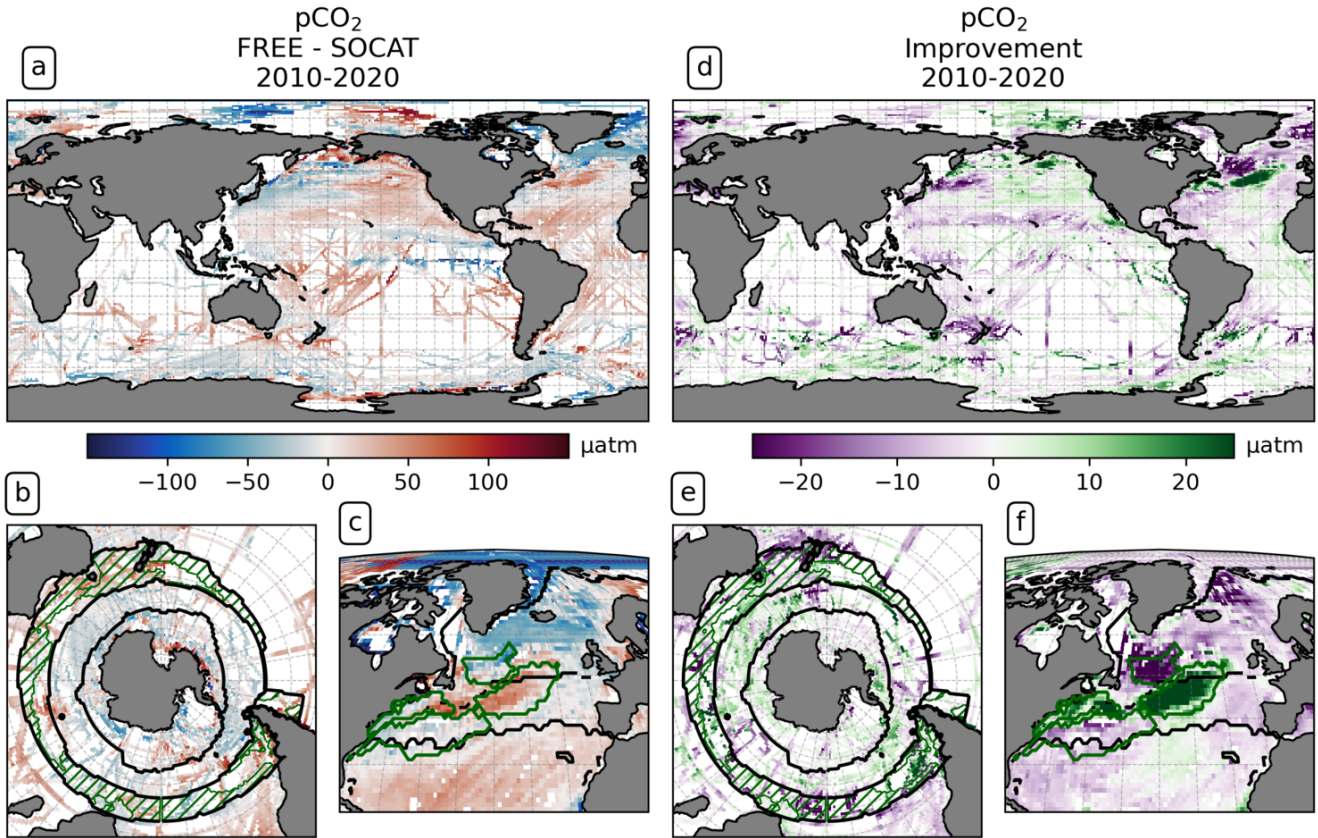


Figure 9. Partial pressure of CO₂ (pCO₂) at the surface averaged over the years 2010-2020. Panels (a-c) show the difference between FREE and SOCAT observations in (a) the global ocean, (b) Southern Ocean and (c) North Atlantic; panels (d-f) show the impact of the assimilation as 'improvement' relative to SOCAT observations computed from monthly mean pCO₂ in the same regions. Positive values (green color) denote a reduced difference to SOCAT.

570 tive and positive differences between the simulations and the GLODAP climatology, which are in the order of 20 mmol C m^{-3} (Fig. A17c). On a global average, the assimilation leads to an increase in DIC between 200-600 m depth and a reduction of DIC between the surface and 200m, with the largest effect in the upper 400 m (Fig. A17b). This leads to an improved agreement with the GLODAP climatology, with the largest global mean improvement at a depth of 400 m ($2.5 \text{ mmol C m}^{-3}$; Fig. A17d). Below 1000 m depth, the global mean absolute difference FREE-ASML of DIC and alkalinity is only $1 - 2 \text{ mmol m}^{-3}$ and is therefore substantially smaller than at the surface.

575 The comparison with GLODAP bottle alkalinity at the surface shows a similar spatial patterns as for DIC (see Fig. 10a and c). The magnitude of the bias is also comparable ($14 \text{ mmol Alk m}^{-3}$). The global mean of the absolute difference ASML-FREE of surface alkalinity is $7.72 \text{ mmol Alk m}^{-3}$. The assimilation leads to a reduction of the absolute difference of the model alkalinity to GLODAP from $34.34 \text{ mmol Alk m}^{-3}$ to $32.60 \text{ mmol Alk m}^{-3}$. Since the effects of physics assimilation on

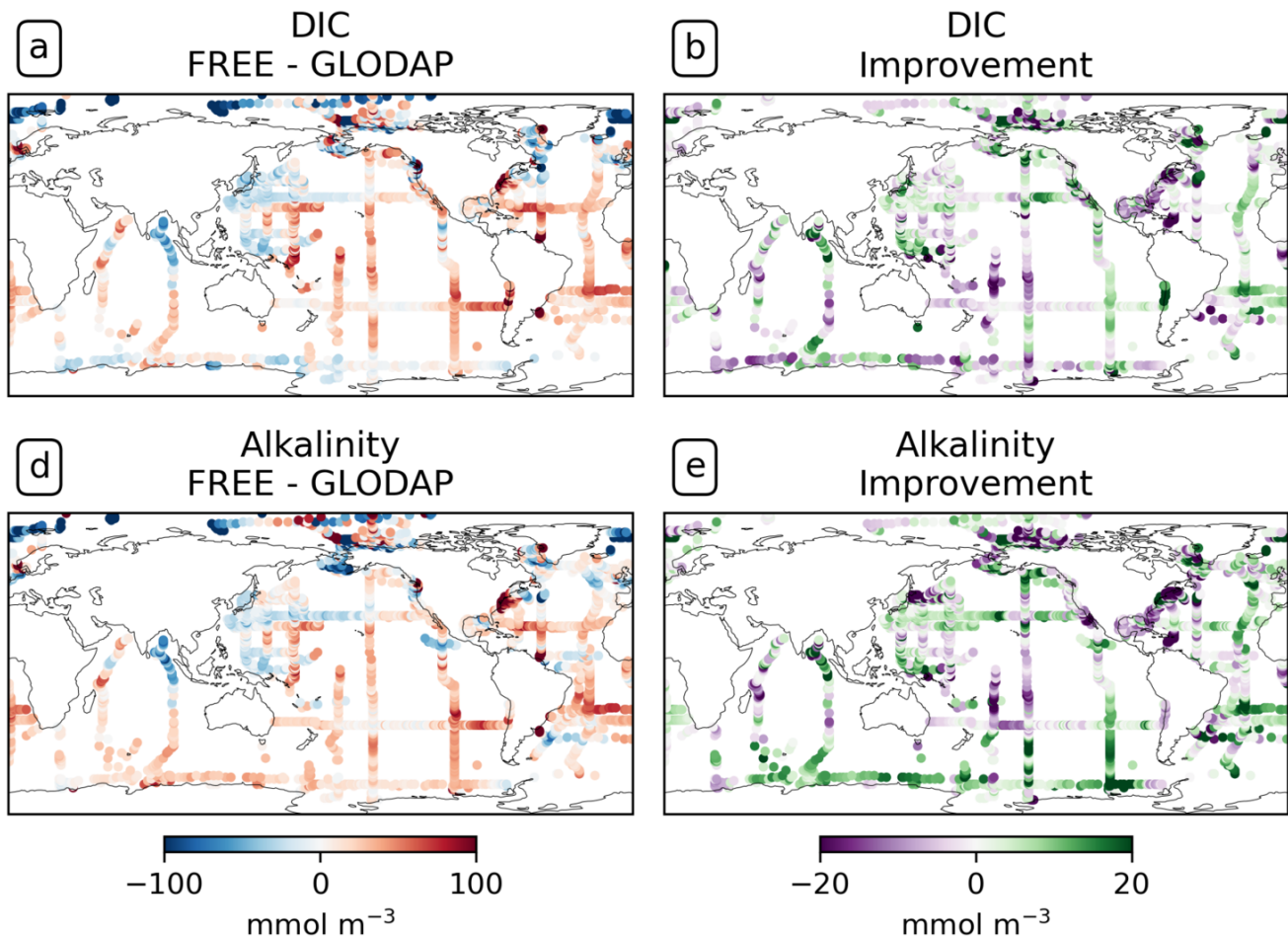


Figure 10. Comparison of the model result with surface DIC and alkalinity bottle observations from GLODAP over the years 2010 to 2020. (a) Difference of DIC between FREE and GLODAP. (b) Improvement of monthly surface DIC relative to GLODAP. (c and d) For alkalinity.

alkalinity and DIC are regionally consistent, regions of improved or deteriorated agreement with GLODAP often coincide for both variables (compare Fig. 10b and d). Because changes of DIC and alkalinity have an opposing effect on the CO_2 flux, it is likely that their correlation results in compensating effects. A linear estimate shows that the joint effect of DIC and alkalinity changes is responsible for a change in the CO_2 flux in the order of $1.22 \text{ mmol C m}^{-2} \text{ day}^{-1}$ on average, and, globally integrated, the assimilation-induced changes in DIC and alkalinity lead to an estimated net increase of the air-sea CO_2 flux in the order of $0.50 \text{ Pg C yr}^{-1}$ (Eqs. (3), (4) and (8)). However, this linear offline estimate is subject to a large uncertainty because regionally large effects with opposite sign lead to uncertainty in the global mean.

585 3.4.3 Surface chlorophyll (OC-CCI)

The representation of chlorophyll by the model is of interest as a proxy for primary production. Surface chlorophyll reflects the phytoplankton state and biomass, and therefore, effects of the DA on the biological model state can be seen in the total surface chlorophyll concentration. A comparison of the modeled surface chlorophyll with remotely-sensed chlorophyll from OC-CCI reveals that both simulations feature a higher surface chlorophyll concentration than OC-CCI (FREE-OBS in Fig. 11a and
590 c). In FREE, the difference to OC-CCI is 0.02 mg m^{-3} on global average, with low deviations in the tropics and an enhanced difference north of 30°N (0.12 mg m^{-3}) and south of 30°S (0.24 mg m^{-3}). Apart from this, both simulations capture the global distribution of chlorophyll well. The simulations show the seasonal maxima in each hemisphere around one month earlier in the year (not shown). South of 30°S , FREE is in better agreement with chlorophyll-a from Johnson et al.'s (2013) Southern Ocean specific chlorophyll product (Fig. 11b) than with OC-CCI data (Fig. 11a).

595 On global average, the assimilation slightly reduces the differences between model and OC-CCI data, from a global mean absolute difference of 0.31 mg m^{-3} to 0.29 mg m^{-3} . The assimilation changes the chlorophyll concentration by an absolute value of 0.05 mg m^{-3} on average, which is 15% of the global mean absolute difference to OC-CCI. There are regions in which assimilation leads to a reduction in chlorophyll and thus to better agreement with the satellite products, for example in the North Atlantic Subpolar Gyre and the Southern Ocean SPSS_{SO} (Fig. 11e and f). In contrast, the model reacts to the DA with
600 an increase in chlorophyll in the North Atlantic Central STSS_{NA-} and the Argentine Basin, which leads to poorer agreement.

4 Discussion

The improvement in temperature and salinity overall leads to a heterogeneous picture in biogeochemistry. While near-surface temperature and salinity fields are improved through DA almost everywhere, the global mean absolute difference of modeled surface pCO_2 to SOCAT remains similar in ASML compared to FREE, and this also applies to the model-observation differ-
605 ences for surface chlorophyll, DIC and alkalinity (Section 3.4). Where improvements in one BGC variable occur, these do not necessarily lead to consistent improvement in all BGC variables. For example, the representation of pCO_2 improves while that of chlorophyll deteriorates in the North Atlantic Central STSS_{NA-} (Fig. 11f and Fig. 9f). In the Southern Ocean SPSS_{SO}, the reduction of modeled surface chlorophyll in spring and the increase of pCO_2 in summer lead to a better agreement with pCO_2 observations, yet the available observations of DIC and alkalinity do not resolve the regional scales to evaluate the correspond-
610 ing changes in these variables (Fig. 9, Fig. 10 and Fig. 11f). The uncertainty represented by the ensemble is reduced by the DA, which has the most obvious effect on the directly assimilated fields (SST in Fig. 6d and e and density in Fig. 8f). The ensemble standard deviation of the CO_2 flux, where it is large in FREE, is constrained by the DA to globally more uniform and smaller values (Fig. 5c-f, Fig. 7c-f and Fig. A1). Only in the North Pacific, the standard deviation of CO_2 fluxes is equally high in ASML and FREE, precisely in a region that also presents a challenge for pCO_2 products (compare Fig. A1 and Mayot
615 et al., 2024, Figure 5a). In the rest of the ocean, the reduced uncertainty represented by the ensemble does not necessarily coincide with improved agreement with BGC observations. One possible reason for improvement of model-data mismatch in one variable with worsening in another may lie in inconsistencies between the observational datasets. Another reason may be

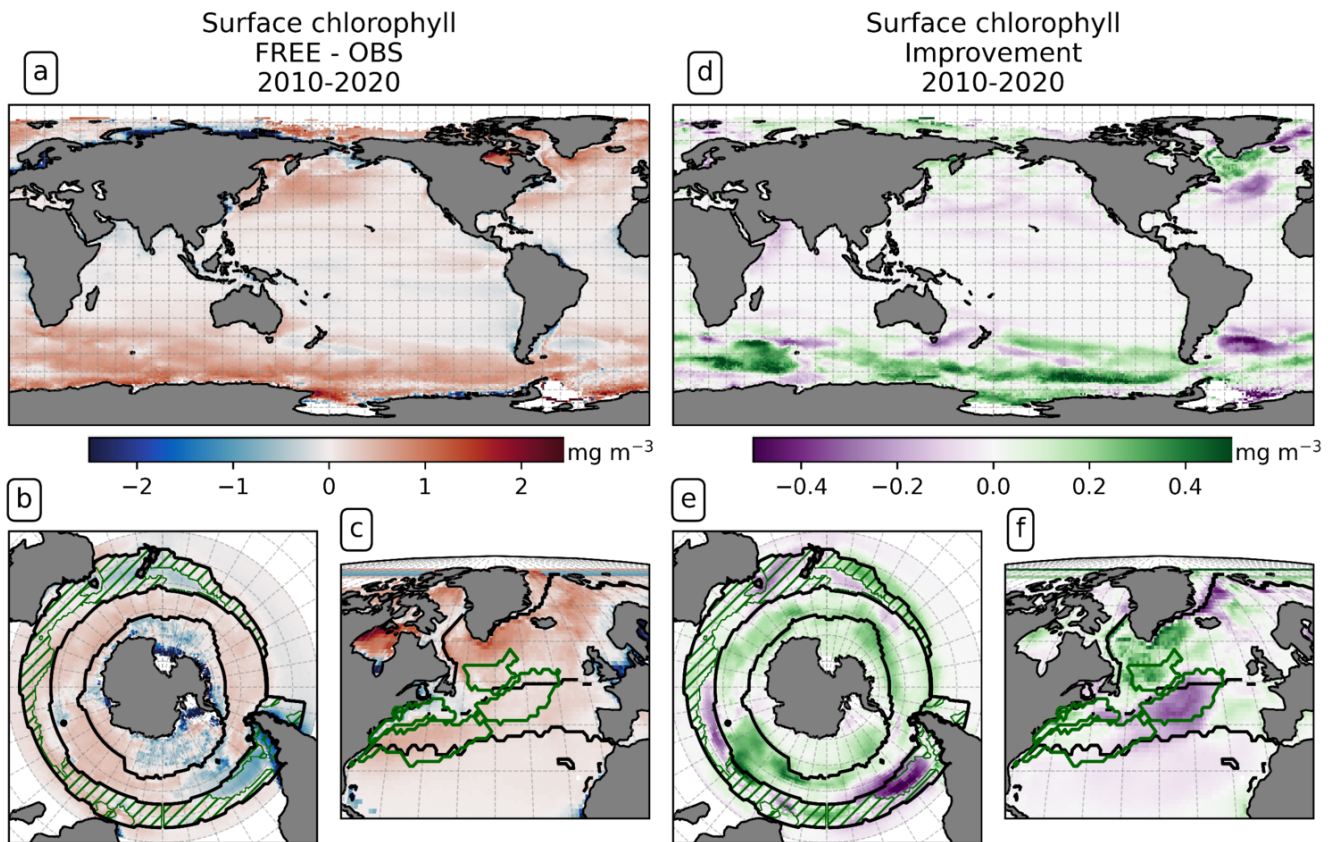


Figure 11. Surface chlorophyll averaged over the years 2010-2020: (a-c) difference between FREE and SOCAT observations in (a) the global ocean, (b) the Southern Ocean and (c) the North Atlantic; (d-f) impact of the assimilation as 'improvement' relative to the observations in the same regions. Panels (a, c, d) and (f) compare to monthly OC-CCI observations, panels (b) and (e) refer to the climatology for 1998-2019 by Johnson et al. (2013).

missing processes in the model and the use of constant BGC model parameters. Those parameters are responsible for linking changes between ecosystem variables and in reality, they vary across space and time depending on species composition in the ecosystem (Mammun et al., 2023, Chapter 3). Overly simplified links between ecosystem variables can lead to cancelling errors, which means that the state of one variable may worsen as a result of improving the other through DA (as in Ford and Barciela, 2017).

The major effects of physics DA on BGC variables seem to be related to changes of SST and are largely uniform over the full period of DA (Section 3.4). Surface chlorophyll changes follow SST changes (Figs. 1 and 11). The modeled phytoplankton growth is temperature-dependent (Gürses et al., 2023). Furthermore, indirect temperature effects on plankton dynamics due to stratification and mixing changes contribute, albeit those can have heterogeneous effects and the correlation of chlorophyll and boundary-layer depth is less clear (not shown). The changes of surface DIC and alkalinity show similar spatial patterns

with regional heterogeneity (Section 3.3), again with the major changes being coherent with the changes in SST (Fig. 1). Furthermore, the effects of the assimilation on DIC and on temperature in the upper 1000 m correlate regionally: Cooling through DA at intermediate depth (Fig. 2b) is usually accompanied by higher DIC in ASML (Fig. A17b), while warming through DA near the surface occurs together with reduced DIC in ASML. An overall more stable ocean stratification in the upper hundreds of meters explains why. On global average, the assimilation leads to lower DIC above 200 m and higher DIC between 200-600 m depth. In regions of substantial DA effects on vertical transport of DIC, as for example in the Central STSS_{NA}– or in the SPSS_{SO} (Section 3.3), the modelled biogeochemical cycles adjust dynamically to the altered vertical transport. The resulting changes in biological sources and sinks of DIC compensate for 20-70% of the changes in vertical transport of DIC (Fig. A13a). In addition to changes in stability and mixing, the assimilation affects the distribution of DIC and alkalinity through local changes in near-surface horizontal transport. As the horizontal distribution of surface DIC, alkalinity and SST is governed by latitudinal gradients and common pathways of transport (Figs. A10 and A15), all of them undergo similar changes as the SST field is modified. An exception to this is in the STSS_{SO}, where regional shifts along contrasting surface gradients of DIC, alkalinity and temperature affect the respective variables differently (Section 3.3). These shifts change the spatial pattern of air-sea CO₂ fluxes. With the exception of the Southern Ocean, zonally averaged changes in surface pCO₂ are dominated by the combined effects of surface alkalinity and DIC on pCO₂ (Fig. A9a). Because alkalinity and DIC are usually modified according to the same pattern through mechanisms acting on both, their effects on pCO₂ are anticorrelated (Fig. A9b). The direct thermal effect on pCO₂ can still be the largest locally, for example in the North Atlantic Newfoundland Basin+ (Fig. A12b). While the DA dynamically induces changes in surface pCO₂ everywhere, the strongest effects on the air-sea CO₂ flux are at high latitudes, where pCO₂ changes are amplified by high wind velocities.

The net effect of DA on the global air-sea CO₂ flux varies from year to year between -0.12 PgCyr^{-1} and 0.15 PgCyr^{-1} , which is small compared to the changes in regional CO₂ fluxes. The global net effect of lateral redistribution of alkalinity and DIC at the ocean surface is a result of compensation between regions where alkalinity and DIC are added and removed. Similarly, regional SST effects on surface pCO₂ mostly balance out globally, because DA primarily induces a correction of regional SST biases, reducing the mean absolute difference to the observations from 0.59°C to 0.32°C , rather than changing the global mean SST, which differs by only 0.02°C between FREE and ASML. DA-induced differences in vertical transport of DIC are comparably large south of 50°S , but approximately 95% of them are balanced globally by opposing changes in vertical transport further north (vertical transport of DIC in Fig. A13a). In particular, the effect of DA on subduction of DIC through vertical advection into the ocean's deeper layers (not shown), which is the rate-limiting step on oceanic uptake of anthropogenic CO₂ emissions (DeVries, 2022), appears small, which may be due to an insufficient amount of deep observations. Besides, experiments on longer time scales might be necessary to generate a visible effect of deep circulation changes on the ocean's carbon cycle (Cao et al., 2009), which could however lead to imbalances in the CO₂ flux (Lebehot et al., 2019; Kriest et al., 2020; Primeau and Deleersnijder, 2009). Another possible reason why the DA effect on the global CO₂ flux in our simulation is small, is the variable stoichiometry in REcoM. The dynamic biological functioning reduces the sensitivity of critical fields, like DIC, to physical changes (Buchanan et al., 2018). Furthermore, negative feedback effects between surface alkalinity, DIC, atmospheric pCO₂ and air-sea fluxes might reduce the overall response (Bunsen et al., 2024).

The overall impact of the DA on the air-sea CO₂ flux on a global scale is modest (0.05 Pg C yr⁻¹) compared to the differences between other estimates (e.g., a standard deviation of 0.45 Pg C yr⁻¹ of GOBMs in DeVries et al., 2023). The global air-sea CO₂ flux estimates of FREE and ASML fall in the range of previous model estimates and in the range of previous pCO₂ products (Fig. 4a and b) for the period 2010-2018, during which comparable estimates are available (DeVries et al., 2023). We compare here to two other data assimilating BGC model approaches, namely ECCO-Darwin (global; Carroll et al., 2020) and B-SOSE, which is restricted to the Southern Ocean (Verdy and Mazloff, 2017). Both approaches use Linearized Least Squares Optimization data assimilation methods (4D-var/adjoint and Green's function, Wunsch, 1996; Menemenlis et al., 2005). However, the largest difference to our study is probably that they assimilate BGC observations in addition to physical data. Thus, as expected, the effect on pCO₂ in our study is smaller (3%) than in ECCO-Darwin and B-SOSE where a reduction in pCO₂ model-data misfit of 6% and 64% was reported, respectively (here given as quadratic misfit). The global CO₂ flux (2010-2018) is smaller in FESOM2.1-REcoM3-PDAF (-2.73 Pg C yr⁻¹ in FREE and -2.78 Pg C yr⁻¹ in ASML) than in ECCO-Darwin (-3.13 Pg C yr⁻¹). The discrepancy between the CO₂ flux estimates based on models and pCO₂-products is an area of active research and not fully resolved (Friedlingstein et al., 2023; DeVries et al., 2023). On the one hand, model biases in the Atlantic Meridional Overturning Circulation, in Southern Ocean ventilation and possibly biases in the surface ocean carbonate chemistry were suggested as reasons why models might underestimate the global mean CO₂ uptake in recent decades (Friedlingstein et al., 2023; Terhaar et al., 2024, 2022). On the other hand, the sparsity of observations is a concern for the pCO₂ products. According to one testbed simulation, the pCO₂ products reflect the global mean and the seasonal cycle relatively well, while the decadal variability may be overestimated (Gloege et al., 2021). An overestimation of the decadal trend, as suggested by Hauck et al. (2023b), could explain the high estimates of the pCO₂ products for the present-day global mean CO₂ flux. In contrast, for the North Atlantic, it was argued that pCO₂ is comparatively well constrained by observations in the last decade but not in the 1980s, which has an erroneous influence on the long-term trend (Pérez et al., 2024).

The effects of data assimilation on the CO₂ flux are most pronounced in the Southern Ocean STSS_{SO} and SPSS_{SO} in winter. Verdy and Mazloff (2017) also found the largest effects of assimilation on the CO₂ flux in this region. Although the region is of crucial importance for the global ocean carbon sink, it also has the greatest uncertainty due to the lack of ship-based winter observations (Friedlingstein et al., 2023; Hauck et al., 2020). In the last decade, the number of winter observations has increased due to the introduction of biogeochemical Argo floats (Johnson et al., 2017; Williams et al., 2017), although the float-based pCO₂ derived from pH measurements and estimated alkalinity is subject to higher uncertainty compared to direct pCO₂ measurements (Williams et al., 2017; Bakker et al., 2016). Machine learning approaches incorporating BGC Argo float observations suggest a stronger winter outgassing around Antarctica, particularly south of 50°S in the SPSS_{SO} and ICE_{SO} biomes, for 2015-2017 (Bushinsky et al., 2019; Gray et al., 2018). This results in a lower estimate of annual Southern Ocean CO₂ uptake in the float products. One suggestion in the literature is that model inadequacies in the representation of mixing and upwelling in the Southern Ocean might cause the discrepancy between float products and models (Gray et al., 2018). However, improvements in the modeled ocean physics and changes in mixing through data assimilation do not lead to closer agreement between the FESOM-REcoM estimate and the float products (comparison of FESOM-REcoM, float products and B-SOSE in Fig. A14). In contrast, ASML shows even weaker winter outgassing and stronger summer uptake south of 50°S

than FREE, which brings the FESOM-REcoM estimate further away from the float products. However, ASML is brought close to B-SOSE in terms of winter outgassing in the Antarctic polar ocean south of 60°S and winter uptake in the STSS_{SO} around 40°S . Additionally, airborne CO_2 flux estimates and direct pCO_2 measurements stemming from a sail drone have questioned the estimates of winter outgassing based on the BGC floats, either attributing the high pCO_2 values to possible biases in the floats' measuring devices or to anomalously high pCO_2 in the years 2015-2016 (Long et al., 2021; Sutton et al., 2021).

5 Conclusion

We apply data assimilation of temperature and salinity into a global ocean-biogeochemical model to improve the physical state for the years 2010-2020. The simulation is then assessed with regard to the effects on the biogeochemical variables. The experiments show that the effect of data assimilation (DA) on biogeochemical variables is mostly related to temperature changes. While the air-sea CO_2 flux and pCO_2 are directly affected by sea surface temperature, the DA also induces indirect changes to pCO_2 through dissolved inorganic carbon (DIC) and alkalinity. Globally integrated, these are more relevant for pCO_2 than the direct temperature effect. Yet, which of these factors has a dominant effect on pCO_2 varies locally. The assimilation leads to regional shifts in areas of CO_2 outgassing and uptake. Local effects on the air-sea CO_2 flux are particularly large in dynamic regions such as the North Atlantic Current and near the Subantarctic Front, whose pathways are challenging for the model to resolve without DA. The largest effect on the air-sea CO_2 flux occurs in the Southern Ocean during winter. In the simulation with assimilation, the uptake south of 50°S is increased due to shallower mixing and surface cooling, and the uptake northward of that (40 - 50°S) is weakened. In this area of the ocean, the uncertainty in current estimates of CO_2 fluxes is particularly high. Overall, the uncertainty inherent to the biogeochemical model appears to be larger than the uncertainties induced through physical biases in the free running model. Locally, the changes in surface pCO_2 , chlorophyll, alkalinity, and DIC caused by the assimilation range between about 15 and 30% of the mean absolute model-observation difference. Yet, local improvements in one variable do not necessarily come along with improvements across other observed biogeochemical variables. Therefore, globally, physics DA does not generally improve the difference between the model and observations. In total, the effect of physics DA on the global ocean carbon uptake is with $0.05 \text{ Pg C yr}^{-1}$ small compared to the spread between previous estimates of models, pCO_2 products and other DA estimates. While the assimilation of temperature and salinity improves the representation of these two and also of mixed-layer depth, sea-ice concentration and horizontal near-surface velocities, possible errors in the vertical velocities and overturning circulation are not eliminated. Further biogeochemical variables are only indirectly affected. To this end, the additional assimilation of biogeochemical observations is an obvious next step to reduce the uncertainty stemming from the ecosystem model and to improve the model-observation differences for biogeochemical variables.

Code and data availability. The code used to perform the free simulation and the data assimilation is available at [10.5281/zenodo.11495274](https://zenodo.org/record/11495274). This code archive additionally contains a notebook to produce the manuscript figures from the model output. The processed model output data underlying the figures of this manuscript are available at [10.5281/zenodo.11495081](https://zenodo.org/record/11495081).

730 **Appendix A**

Table A1. List of tracers in REcoM3

Tracers in REcoM3
Dissolved inorganic nitrogen and carbon (DIN, DIC)
Dissolved organic nitrogen and carbon (DON, DOC)
Alkalinity
Oxygen
Iron
Silicate
Intracellular concentrations of nitrogen, carbon, chlorophyll, and calcium in small phytoplankton (PhyN, PhyC, PhyChl, PhyCalc)
Intracellular concentrations of nitrogen, carbon, chlorophyll, and silicate in diatoms (DiaN, DiaC, DiaChl, DiaSi)
Intracellular concentrations of nitrogen and carbon in each of two zooplankton groups (HetN, HetC, Zoo2N, Zoo2C)
Two size classes of detritus for nitrogen, carbon, silicate, and calcium (DetN, DetC, DetSi, DetCalc; and DetZ2N, DetZ2C, DetZ2Si, DetZ2Calc)

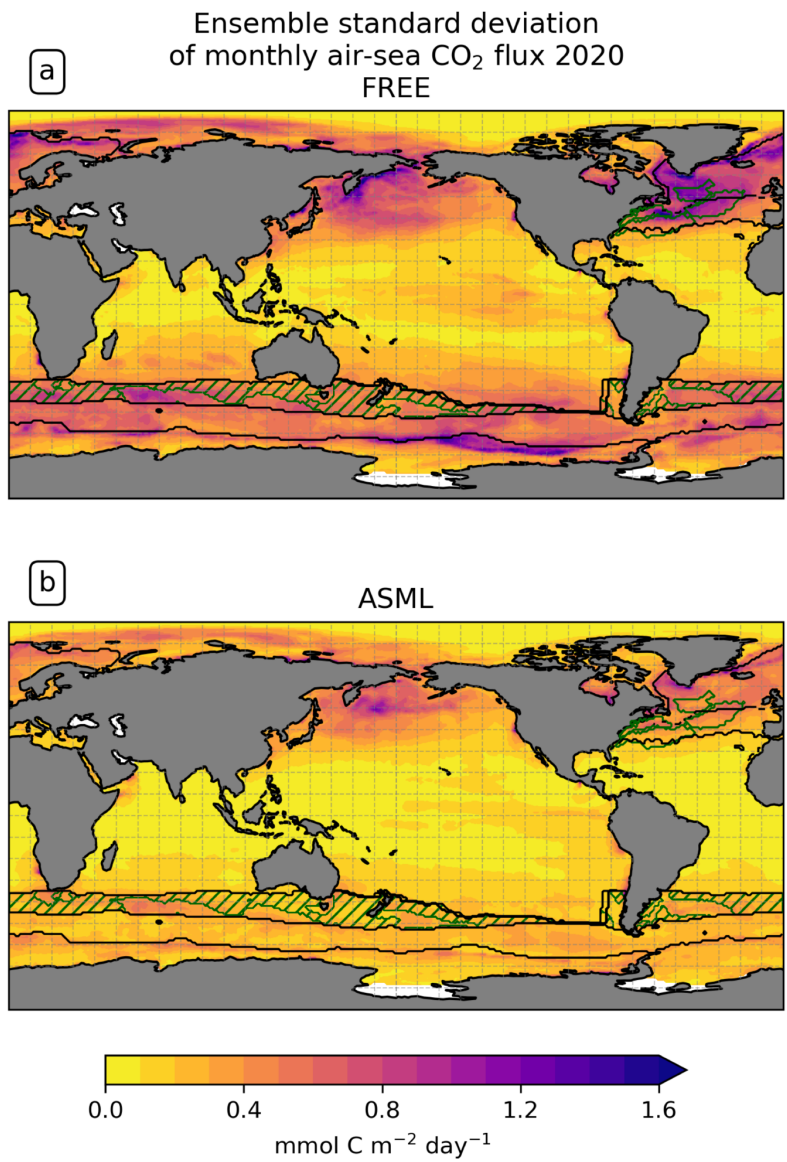


Figure A1. Ensemble standard deviation of monthly air-sea CO₂ flux in the year 2020 in (a) FREE and (b) ASML.

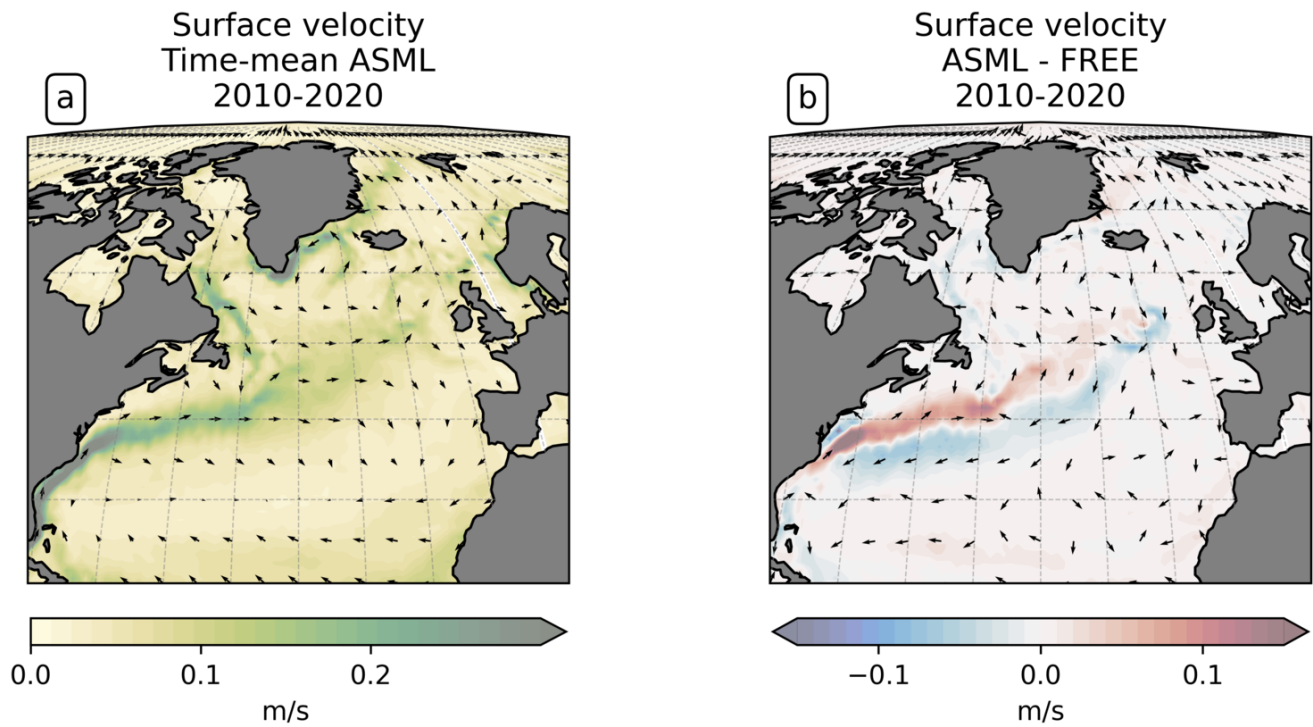


Figure A2. North Atlantic surface velocities, (a) time-mean in ASML and (b) difference ASML-FREE.

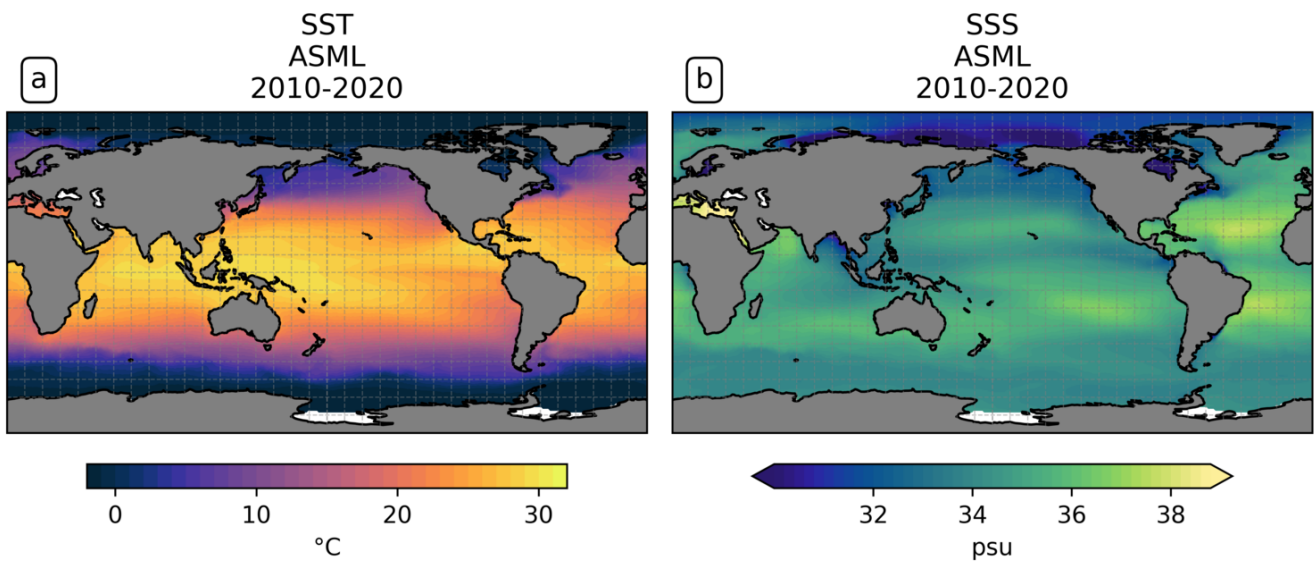


Figure A3. Time-mean sea surface (a) temperature and (b) salinity in ASML.

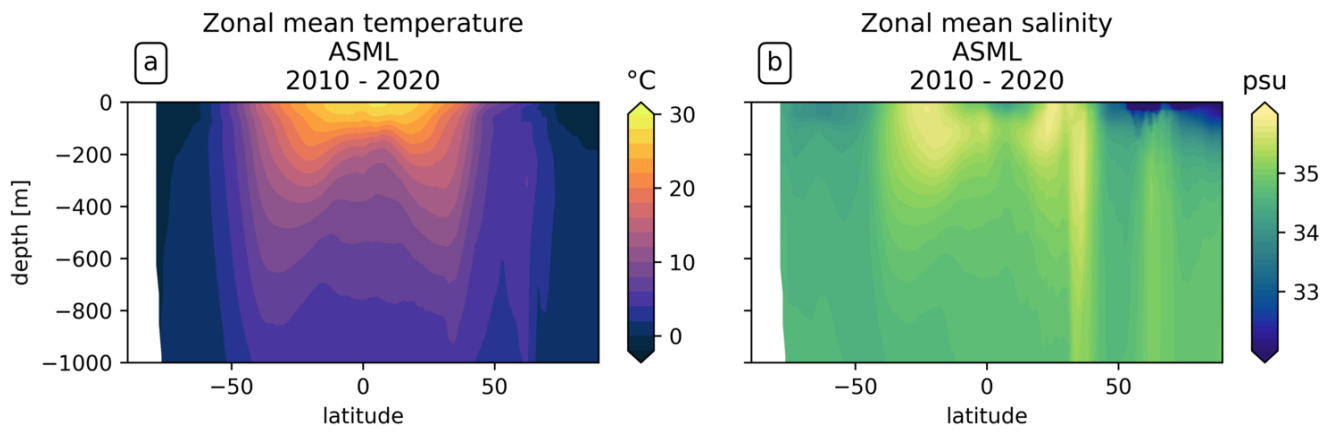


Figure A4. Zonally averaged time-mean (a) temperature and (b) salinity in ASML.

Sea-ice concentration

ASML

Sept 2010-2020

a

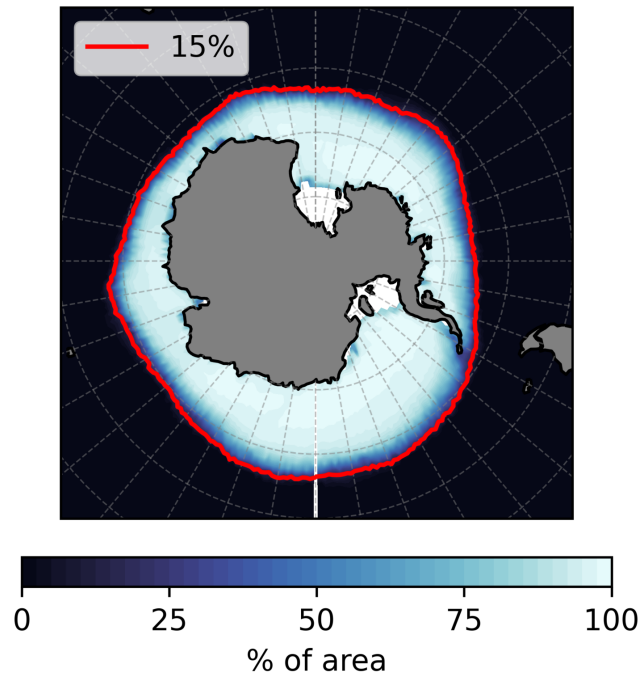


Figure A5. September mean Antarctic sea-ice concentration in ASML.

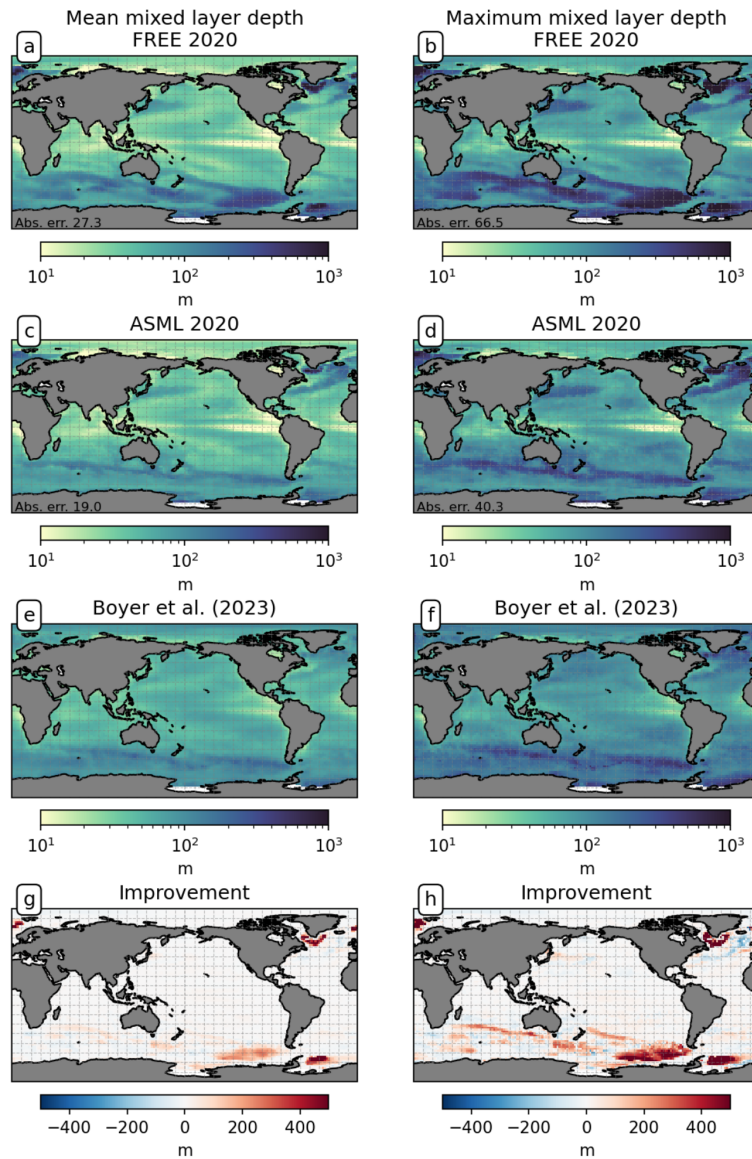


Figure A6. Mixed-layer depth (a,b) in FREE and (c,d) ASML in the year 2020, (e,f) de Boyer Montégut et al.'s (2004) profile-based climatology v2023 and (g,h) the improvement through DA relative to the climatology. On the left: time-mean mixed layer, on the right: maximum of monthly-mean mixed layer. For FREE and ASML (a,b,c,d), the mean absolute difference to the climatology is given in the bottom-left corner.

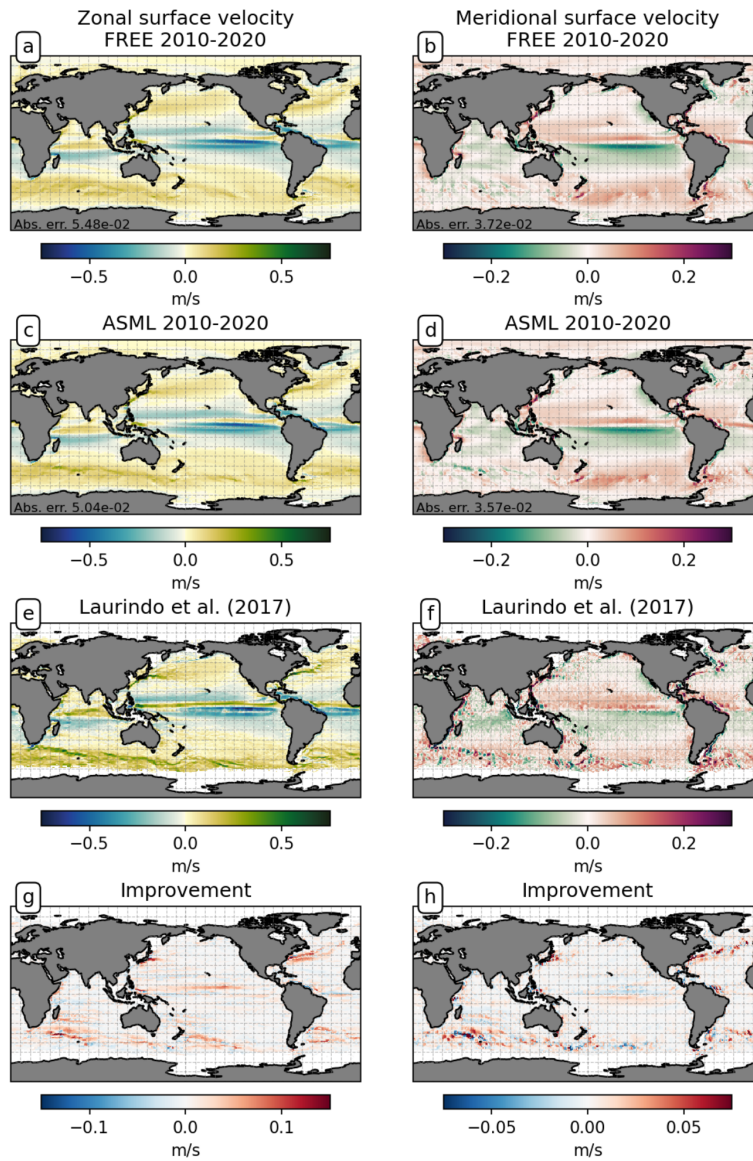


Figure A7. Near-surface velocities (a,b) in FREE and (c,d) ASML for the period 2010-2020, (e,f) Laurindo et al.'s (2017) climatology from drifter observations and (g,h) the improvement through DA relative to the climatology. On the left: zonal velocities, on the right: meridional velocities. For FREE and ASML (a,b,c,d), the mean absolute difference to the climatology is given in the bottom-left corner.

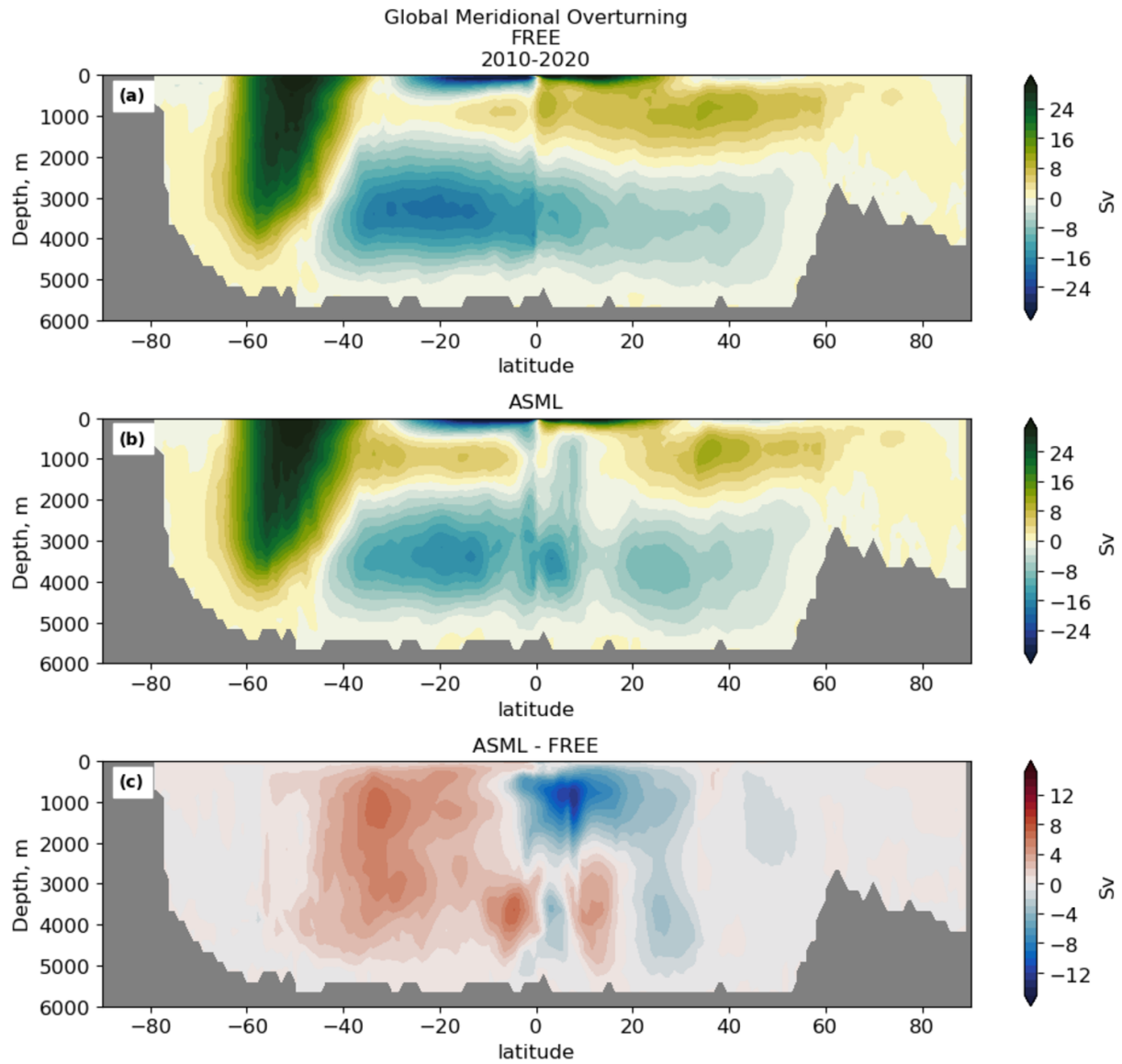


Figure A8. Global meridional overturning in (a) FREE, (b) ASML and (c) difference ASML-FREE.

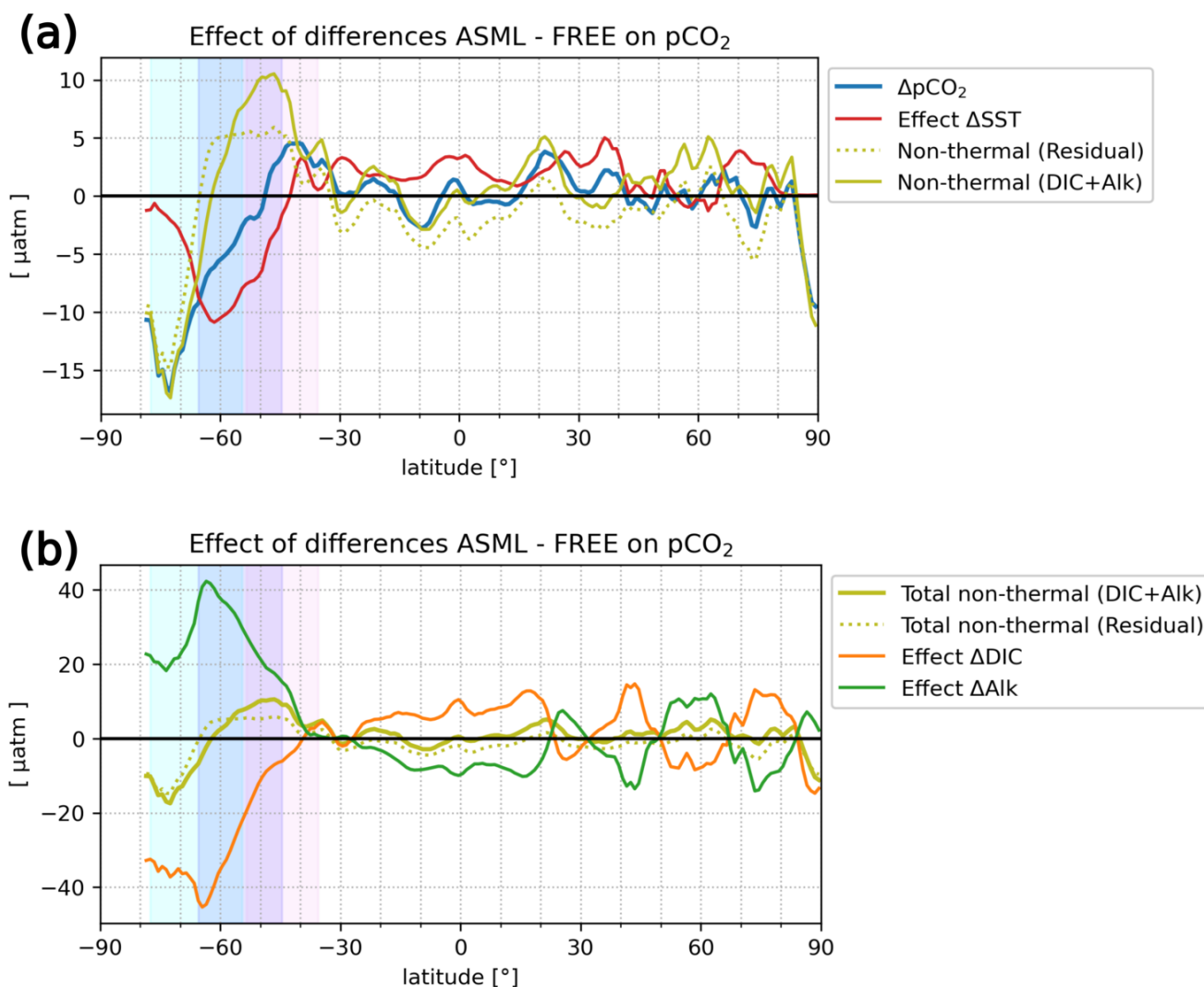


Figure A9. The net difference (ASML-FREE) of surface pCO₂ by latitude (panel a, blue line), and the offline-approximated effects causing that pCO₂ difference for the period 2010-2020: Thermal effect (panel a, red line); non-thermal effect calculated, firstly, as the residual i.e. net-minus-thermal (panels a and b, light-green dotted lines) and, secondly, as the sum of alkalinity and DIC effects (panels a and b, light-green solid lines); and effects of alkalinity and DIC individually (panel b, orange and dark-green lines). The shaded areas in the background indicate the zonal extent of defined biomes in the Southern Ocean: ICE_{S0} in light-blue, SPSS_{S0} in blue and STSS_{S0} in pink. Colors blend where the regions overlap.

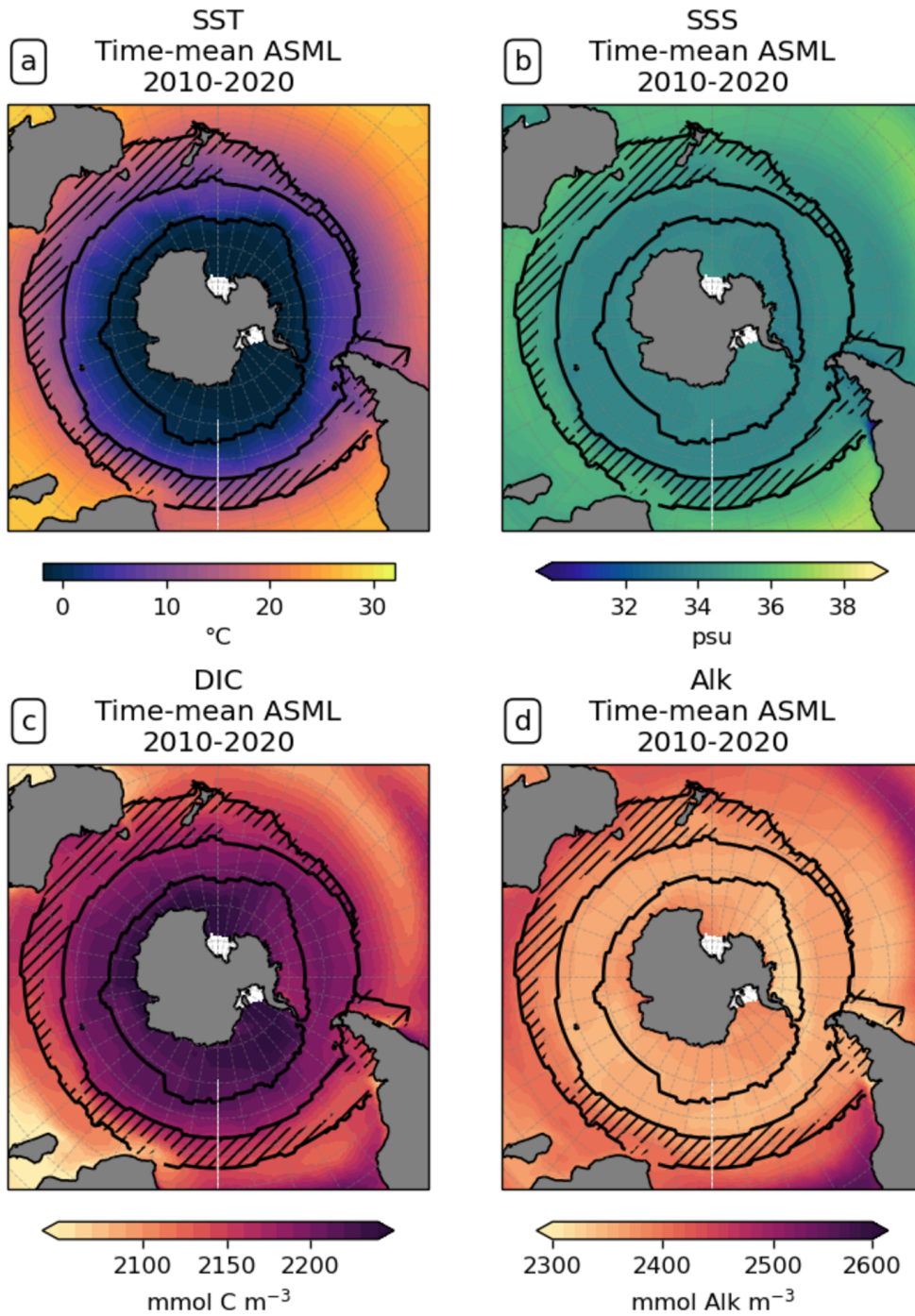


Figure A10. Southern Ocean time-mean sea surface in ASML, (a) temperature, (b) salinity, (c) DIC and (d) alkalinity.

Sources and sinks of DIC and alkalinity at 0-190m ASML - FREE

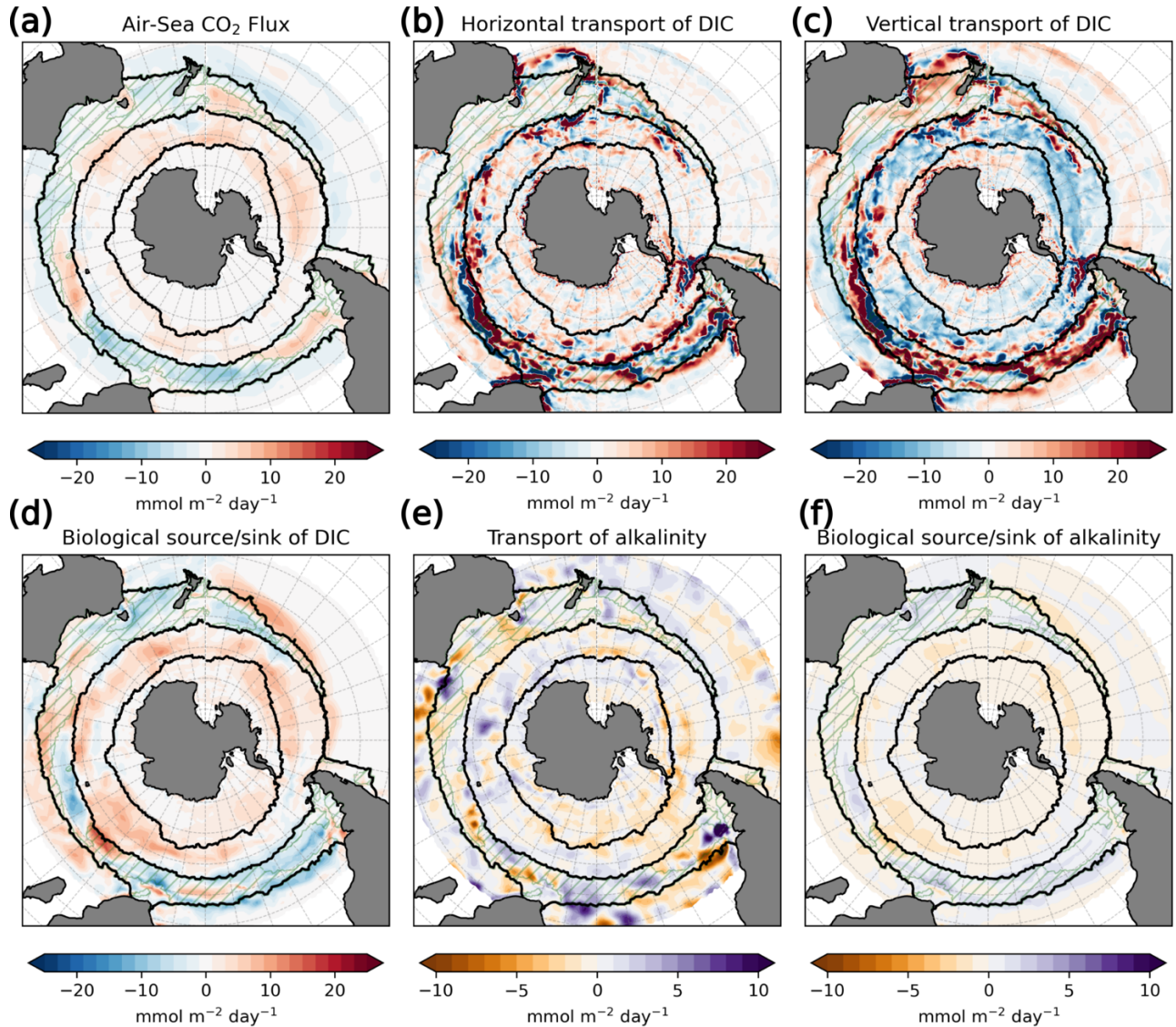


Figure A11. The difference (ASML-FREE) of source and sink terms for the ocean's DIC and alkalinity content integrated over 0-190 m in the Southern Ocean in the year 2020. Transport terms include advection and diffusion of DIC and alkalinity. Biological terms for DIC are the sum of: photosynthesis, respiration, remineralization of dissolved organic carbon, and formation and dissolution of calcite. Biological terms for alkalinity are the sum of: nitrogen assimilation and remineralization, and formation and dissolution of calcite.

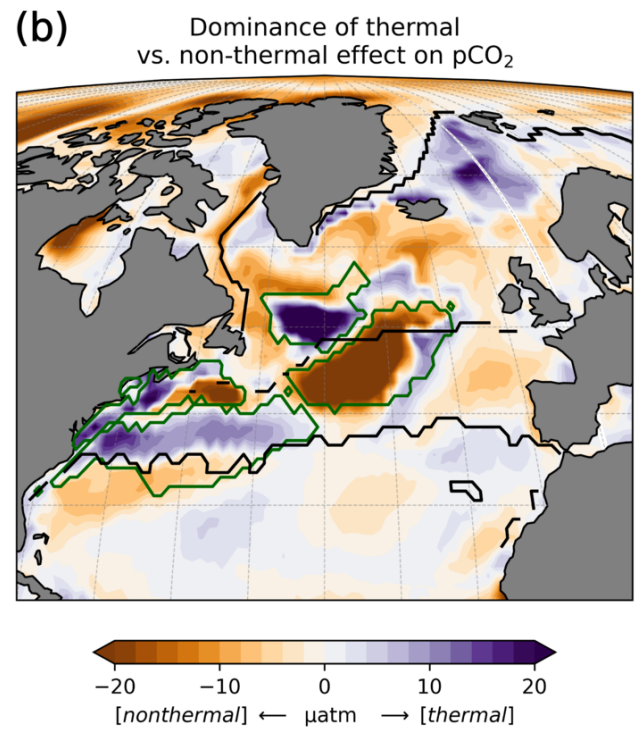
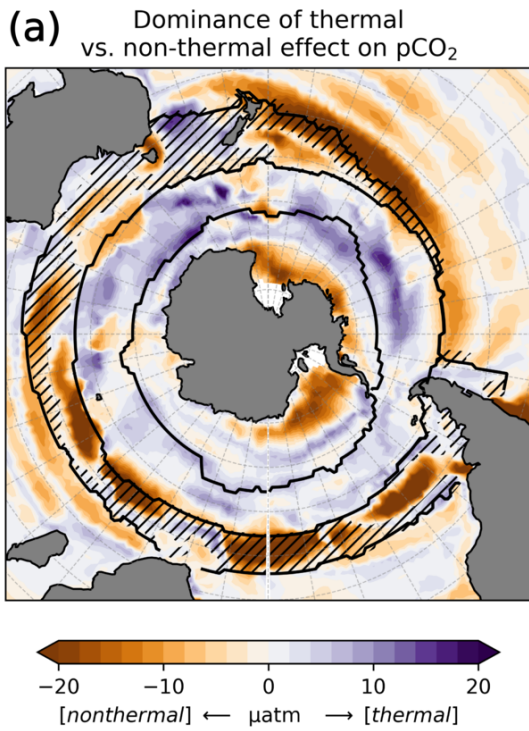


Figure A12. Linear offline estimate of the dominance of thermal versus the non-thermal effect through the assimilation on pCO₂ in the Southern Ocean and North Atlantic for the period 2010-2020.

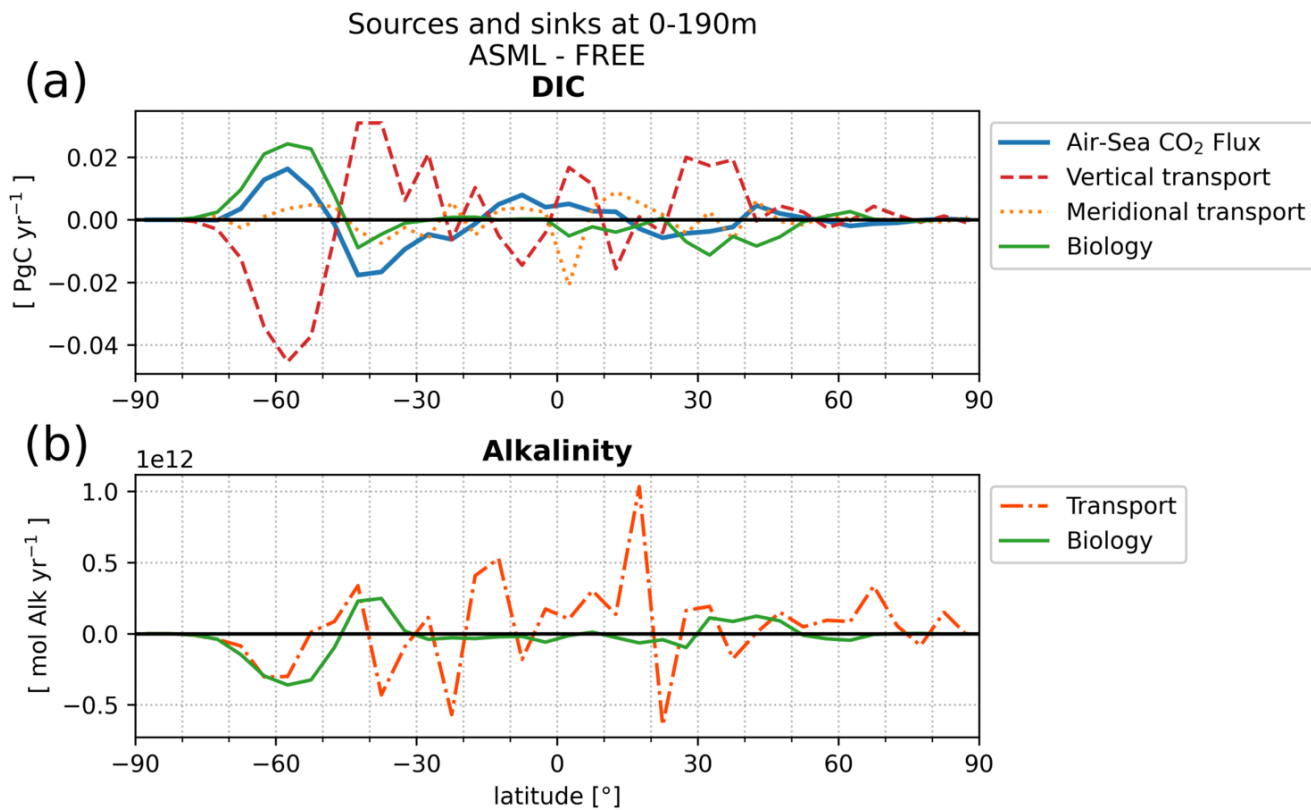


Figure A13. The difference (ASML-FREE) of source and sink terms for the ocean's (a) DIC and (b) alkalinity content integrated over 0-190 m per 1° latitude in the year 2020. Transport terms include advection and diffusion of DIC and alkalinity. Meridional transport is averaged across bins of 5° latitude. In panel b, vertical and horizontal transport are summed up for readability. Biological terms for DIC are the sum of: photosynthesis, respiration, remineralization of dissolved organic carbon, and formation and dissolution of calcite. Biological terms for alkalinity are the sum of: nitrogen assimilation and remineralization, and formation and dissolution of calcite.

JJA 2015-2018 air-sea CO₂ flux

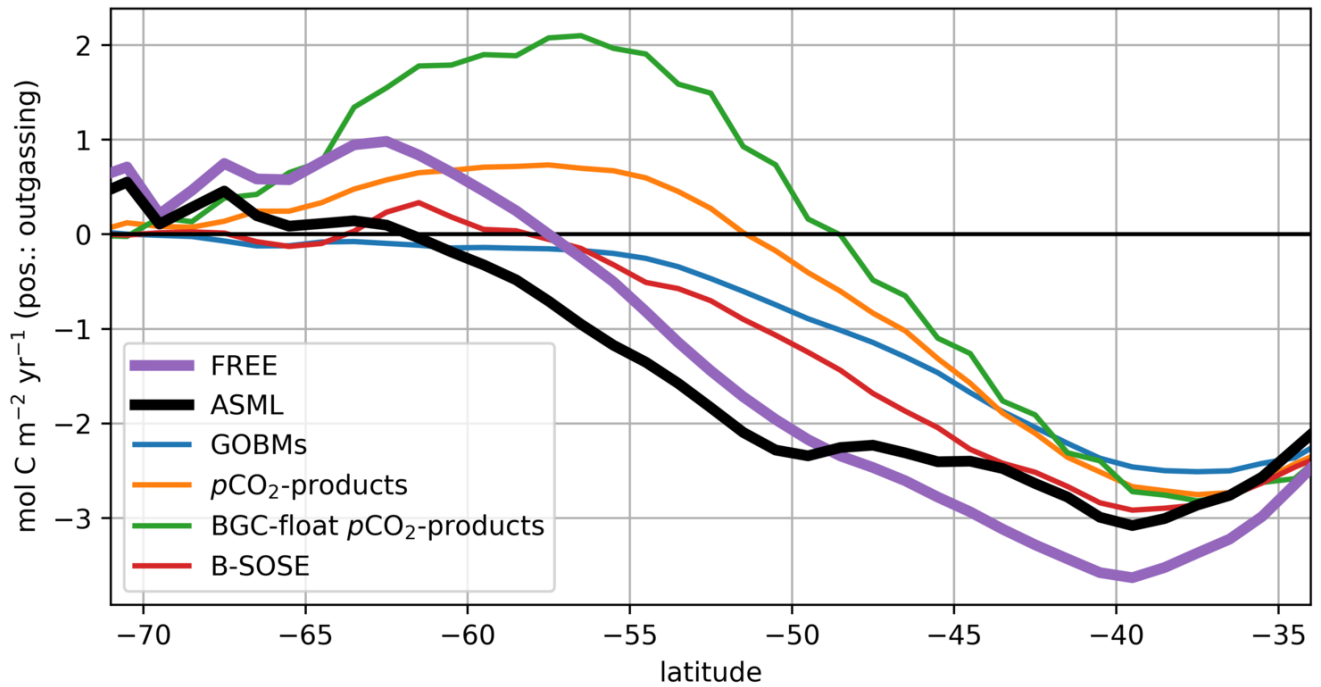


Figure A14. Zonally averaged winter (JJA) air-sea CO₂ flux in FREE, ASML and previous estimates (Hauck et al., 2023a; Verdy and Mazloff, 2017).

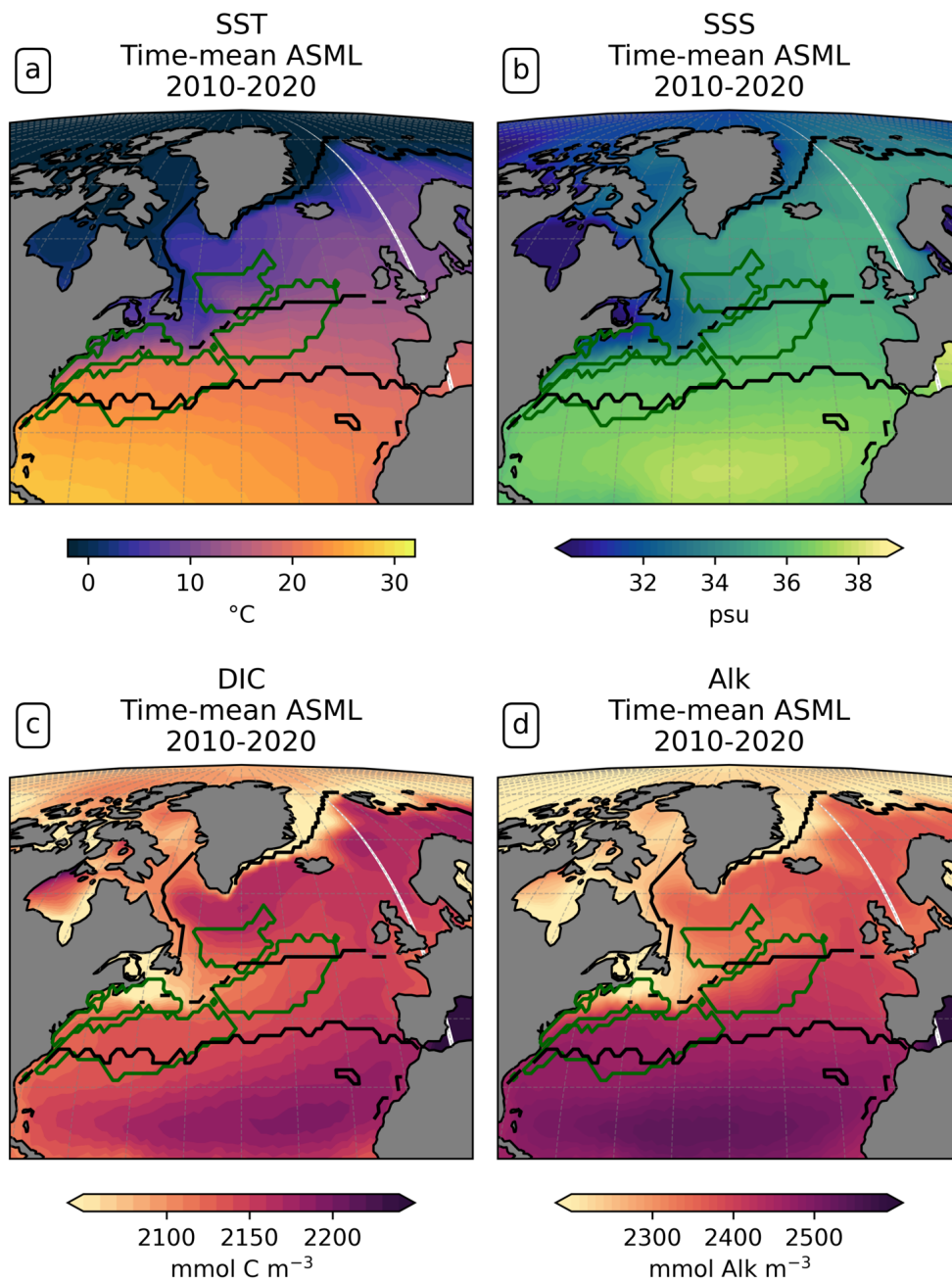


Figure A15. North Atlantic time-mean sea surface in ASML, (a) temperature, (b) salinity, (c) DIC and (d) alkalinity.

Sources and sinks of DIC and alkalinity at 0-190m ASML - FREE

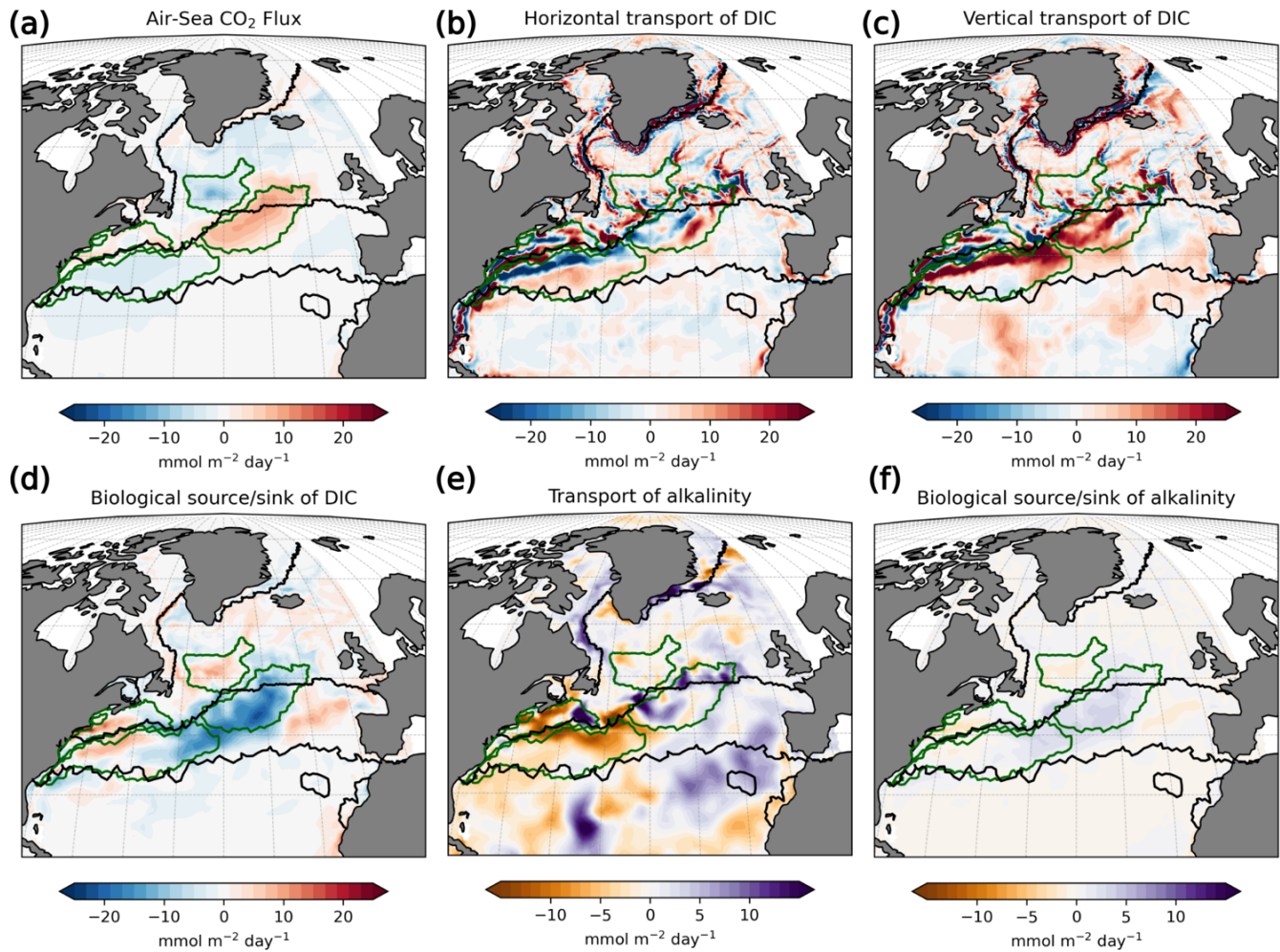


Figure A16. The difference (ASML-FREE) of source and sink terms for the ocean's DIC and alkalinity content integrated over 0-190 m in the North Atlantic in the year 2020. Transport terms include advection and diffusion of DIC and alkalinity. Biological terms for DIC are the sum of: photosynthesis, respiration, remineralization of dissolved organic carbon, and formation and dissolution of calcite. Biological terms for alkalinity are the sum of: nitrogen assimilation and remineralization, and formation and dissolution of calcite.

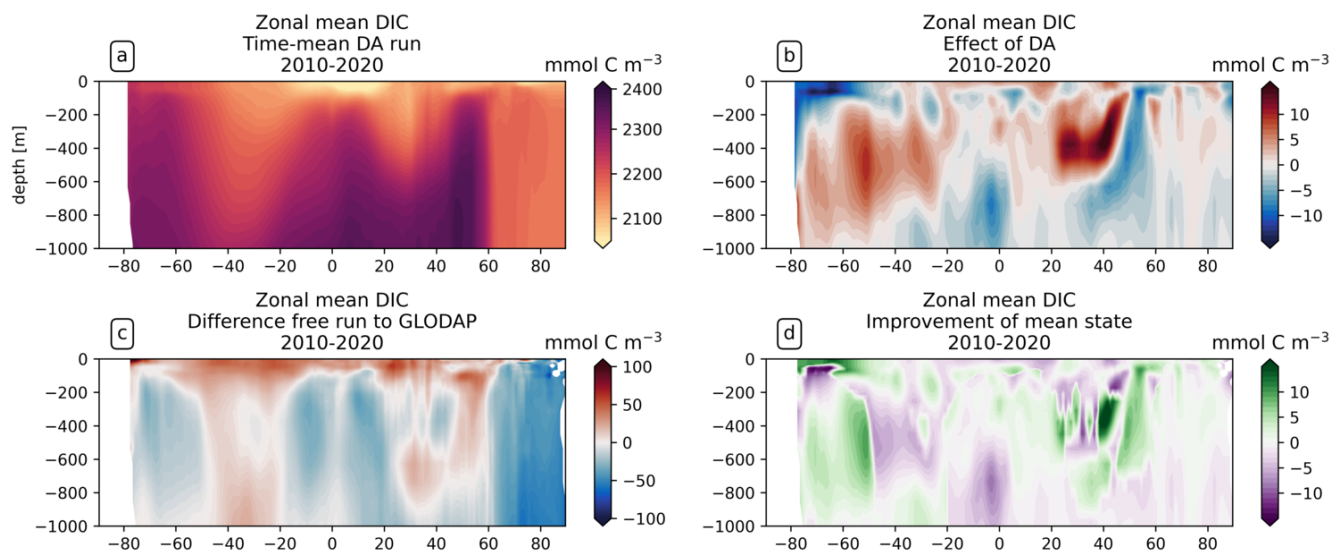


Figure A17. Zonally averaged DIC, (a) time-mean in ASML, (b) difference ASML-FREE, (c) difference FREE-OBS compared to the GLODAP climatology for DIC (Lauvset et al., 2016) and (d) improvement respective to the GLODAP climatology.

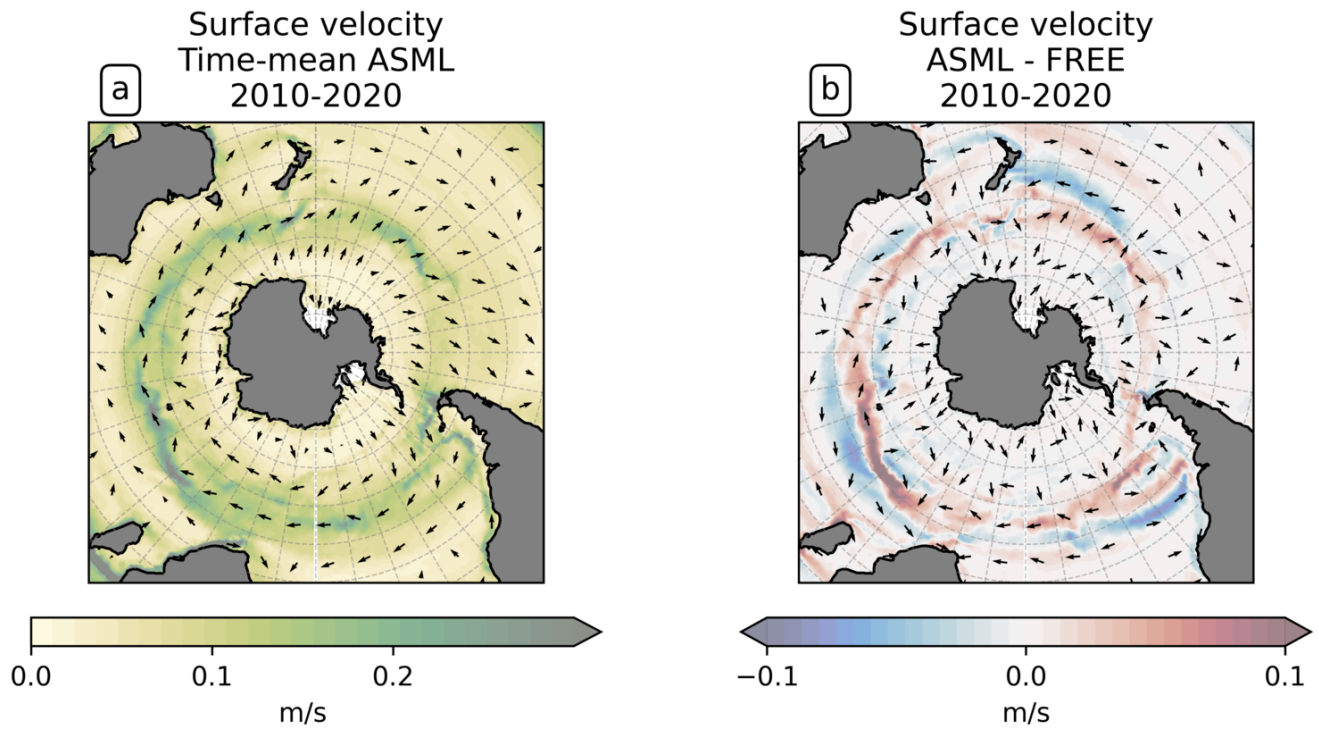


Figure A18. Surface velocities in the Southern Ocean, (a) time-mean in ASML and (b) difference ASML-FREE.

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References

- 745 Bakker, D. C. E., Pfeil, B., Landa, C. S., Metzl, N., O'Brien, K. M., Olsen, A., Smith, K., Cosca, C., Harasawa, S., Jones, S. D., Nakaoka, S., Nojiri, Y., Schuster, U., Steinhoff, T., Sweeney, C., Takahashi, T., Tilbrook, B., Wada, C., Wanninkhof, R., Alin, S. R., Balestrini, C. F., Barbero, L., Bates, N. R., Bianchi, A. A., Bonou, F., Boutin, J., Bozec, Y., Burger, E. F., Cai, W.-J., Castle, R. D., Chen, L., Chierici, M., Currie, K., Evans, W., Featherstone, C., Feely, R. A., Fransson, A., Goyet, C., Greenwood, N., Gregor, L., Hankin, S., Hardman-Mountford, N. J., Harlay, J., Hauck, J., Hoppema, M., Humphreys, M. P., Hunt, C. W., Huss, B., Ibáñez, J. S. P., Johannessen, T., Keeling, R., Kitidis, V., Körtzinger, A., Kozyr, A., Krasakopoulou, E., Kuwata, A., Landschützer, P., Lauvset, S. K., Lefèvre, N., Lo Monaco, C., Manke, A., Mathis, J. T., Merlivat, L., Millero, F. J., Monteiro, P. M. S., Munro, D. R., Murata, A., Newberger, T., Omar, A. M., Ono, T., Paterson, K., Pearce, D., Pierrot, D., Robbins, L. L., Saito, S., Salisbury, J., Schlitzer, R., Schneider, B., Schweitzer, R., Sieger, R., Skjelvan, I., Sullivan, K. F., Sutherland, S. C., Sutton, A. J., Tadokoro, K., Telszewski, M., Tuma, M., van Heuven, S. M. A. C., Vandemark, D., Ward, B., Watson, A. J., and Xu, S.: A multi-decade record of high-quality $f\text{CO}_2$ data in version 3 of the Surface Ocean CO_2 Atlas (SOCAT), Earth System Science Data, 8, 383–413, <https://doi.org/10.5194/essd-8-383-2016>, 2016.
- 750 Bakker, D. C. E., Alin, S. R., Bates, N., Becker, M., Feely, R. A., Gkritzalis, T., Jones, S. D., Kozyr, A., Lauvset, S. K., Metzl, N., Munro, D. R., Nakaoka, S.-i., Nojiri, Y., O'Brien, K. M., Olsen, A., Pierrot, D., Rehder, G., Steinhoff, T., Sutton, A. J., Sweeney, C., Tilbrook, B., Wada, C., Wanninkhof, R., Akl, J., Barbero, L., Beatty, C. M., Berghoff, C. F., Bittig, H. C., Bott, R., Burger, E. F., Cai, W.-J., Castaño Primo, R., Corredor, J. E., Cronin, M., De Carlo, E. H., DeGrandpre, M. D., Dietrich, C., Drennan, W. M., Emerson, S. R., Enochs, I. C., Enyo, K., Epherra, L., Evans, W., Fiedler, B., Fontela, M., Frangoulis, C., Gehrung, M., Giannoudi, L., Glockzin, M., Hales, B., Howden, S. D., Ibáñez, J. S. P., Kamb, L., Körtzinger, A., Lefèvre, N., Lo Monaco, C., Lutz, V. A., Macovei, V. A., Maenner Jones, S., Manalang, D., Manzello, D. P., Metzl, N., Mickett, J., Millero, F. J., Monacci, N. M., Morell, J. M., Musielewicz, S., Neill, C., Newberger, T., Newton, J., Noakes, S., Ólafsdóttir, S. R., Ono, T., Osborne, J., Padín, X. A., Paulsen, M., Perivoliotis, L., Petersen, W., Petihakis, G., Plueddemann, A. J., Rodriguez, C., Rutgersson, A., Sabine, C. L., Salisbury, J. E., Schlitzer, R., Skjelvan, I., Stamataki, N., Sullivan, K. F., Sutherland, S. C., T'Jampens, M., Tadokoro, K., Tanhua, T., Telszewski, M., Theetaert, H., Tomlinson, M., Vandemark, D., Velo, A., Voynova, Y. G., Weller, R. A., Whitehead, C., and Wimart-Rousseau, C.: Surface Ocean CO_2 Atlas Database Version 2023 (SOCATv2023) (NCEI Accession 0278913), <https://doi.org/10.25921/r7xa-bt92>, 2023.
- 760 Ballantyne, A. P., Alden, C. B., Miller, J. B., Tans, P. P., and White, J. W. C.: Increase in observed net carbon dioxide uptake by land and oceans during the past 50 years, *Nature*, 488, 70–72, <https://doi.org/10.1038/nature11299>, 2012.
- 770 Bernardello, R., Sicardi, V., Lapin, V., Ortega, P., Ruprich-Robert, Y., Tourigny, E., and Ferrer, E.: Ocean biogeochemical reconstructions to estimate historical ocean CO_2 uptake, *Earth System Dynamics*, 15, 1255–1275, <https://doi.org/10.5194/esd-15-1255-2024>, 2024.
- Boutin, J., Vergely, J.-L., Reul, N., Catany, R., Koehler, J., Martin, A., Rouffi, F., Arias, M., Chakroun, M., Corato, G., et al.: ESA Sea Surface Salinity Climate Change Initiative (Sea_Surface_Salinity_cci): Weekly and Monthly Sea Surface Salinity Product, v03.21, from 2010 to 2020, NERC EDS Centre for Environmental Data Analysis, <http://dx.doi.org/10.5285/5920a2c77e3c45339477acd31ce62c3c>, 2021.
- 775 Brix, H., Menemenlis, D., Hill, C., Dutkiewicz, S., Jahn, O., Wang, D., Bowman, K., and Zhang, H.: Using Green's Functions to initialize and adjust a global, eddy ocean biogeochemistry general circulation model, *Ocean Modelling*, 95, 1–14, <https://doi.org/10.1016/j.ocemod.2015.07.008>, 2015.
- Buchanan, P. J., Matear, R. J., Chase, Z., Phipps, S. J., and Bindoff, N. L.: Dynamic Biological Functioning Important for Simulating and Stabilizing Ocean Biogeochemistry, *Global Biogeochemical Cycles*, 32, 565–593, <https://doi.org/10.1002/2017GB005753>, 2018.

- 780 Bunsen, F., Nissen, C., and Hauck, J.: The Impact of Recent Climate Change on the Global Ocean Carbon Sink, *Geophysical Research Letters*, 51, e2023GL107 030, <https://doi.org/10.1029/2023GL107030>, 2024.
- Bushinsky, S. M., Landschützer, P., Rödenbeck, C., Gray, A. R., Baker, D., Mazloff, M. R., Resplandy, L., Johnson, K. S., and Sarmiento, J. L.: Reassessing Southern Ocean Air-Sea CO₂ Flux Estimates With the Addition of Biogeochemical Float Observations, *Global Biogeochemical Cycles*, 33, 1370–1388, <https://doi.org/10.1029/2019GB006176>, 2019.
- 785 Cao, L., Eby, M., Ridgwell, A., Caldeira, K., Archer, D., Ishida, A., Joos, F., Matsumoto, K., Mikolajewicz, U., Mouchet, A., Orr, J. C., Plattner, G.-K., Schlitzer, R., Tokos, K., Totterdell, I., Tschumi, T., Yamanaka, Y., and Yool, A.: The role of ocean transport in the uptake of anthropogenic CO₂, *Biogeosciences*, 6, 375–390, <https://doi.org/10.5194/bg-6-375-2009>, 2009.
- Carroll, D., Menemenlis, D., Adkins, J. F., Bowman, K. W., Brix, H., Dutkiewicz, S., Fenty, I., Gierach, M. M., Hill, C., Jahn, O., Landschützer, P., Lauderdale, J. M., Liu, J., Manizza, M., Naviaux, J. D., Rödenbeck, C., Schimel, D. S., Van der Stocken, T., and Zhang, H.:
- 790 The ECCO-Darwin Data-Assimilative Global Ocean Biogeochemistry Model: Estimates of Seasonal to Multidecadal Surface Ocean pCO₂ and Air-Sea CO₂ Flux, *Journal of Advances in Modeling Earth Systems*, 12, e2019MS001 888, <https://doi.org/10.1029/2019MS001888>, 2020.
- Chapman, C. C., Lea, M.-A., Meyer, A., Sallée, J.-B., and Hindell, M.: Defining Southern Ocean fronts and their influence on biological and physical processes in a changing climate, *Nature Climate Change*, 10, 209–219, <https://doi.org/10.1038/s41558-020-0705-4>, 2020.
- 795 Ciavatta, S., Kay, S., Saux-Picart, S., Butenschön, M., and Allen, J. I.: Decadal reanalysis of biogeochemical indicators and fluxes in the North West European shelf-sea ecosystem, *Journal of Geophysical Research: Oceans*, 121, 1824–1845, <https://doi.org/10.1002/2015JC011496>, 2016.
- Ciavatta, S., Brewin, R. J. W., Skákala, J., Polimene, L., de Mora, L., Artioli, Y., and Allen, J. I.: Assimilation of Ocean-Color Plankton Functional Types to Improve Marine Ecosystem Simulations, *Journal of Geophysical Research: Oceans*, 123, 834–854,
- 800 <https://doi.org/10.1002/2017JC013490>, 2018.
- CMEMS: Operational Sea Surface Temperature and Ice Analysis (OSTIA), <https://doi.org/10.48670/moi-00165>, 2023.
- Crisp, D., Dolman, H., Tanhua, T., McKinley, G. A., Hauck, J., Bastos, A., Sitch, S., Eggleston, S., and Aich, V.: How Well Do We Understand the Land-Ocean-Atmosphere Carbon Cycle?, *Reviews of Geophysics*, 60, e2021RG000 736, <https://doi.org/10.1029/2021RG000736>, 2022.
- 805 Danilov, S., Sidorenko, D., Wang, Q., and Jung, T.: The Finite-volume Sea ice–Ocean Model (FESOM2), *Geoscientific Model Development*, 10, 765–789, <https://doi.org/10.5194/gmd-10-765-2017>, 2017.
- Davila, X., Gebbie, G., Brakstad, A., Lauvset, S. K., McDonagh, E. L., Schwinger, J., and Olsen, A.: How Is the Ocean Anthropogenic Carbon Reservoir Filled?, *Global Biogeochemical Cycles*, 36, e2021GB007 055, <https://doi.org/10.1029/2021GB007055>, 2022.
- de Boyer Montégut, C., Madec, G., Fischer, A. S., Lazar, A., and Iudicone, D.: Mixed layer depth over the global
- 810 ocean: An examination of profile data and a profile-based climatology, *Journal of Geophysical Research: Oceans*, 109, <https://doi.org/https://doi.org/10.1029/2004JC002378>, 2004.
- Denvil-Sommer, A., Gehlen, M., and Vrac, M.: Observation system simulation experiments in the Atlantic Ocean for enhanced surface ocean pCO₂ reconstructions, *Ocean Science*, 17, 1011–1030, <https://doi.org/10.5194/os-17-1011-2021>, 2021.
- DeVries, T.: The Ocean Carbon Cycle, *Annual Review of Environment and Resources*, 47, 317–341, [https://doi.org/10.1146/annurev-](https://doi.org/10.1146/annurev-environ-120920-111307)
- 815 [environ-120920-111307](https://doi.org/10.1146/annurev-environ-120920-111307), 2022.
- DeVries, T., Holzer, M., and Primeau, F.: Recent increase in oceanic carbon uptake driven by weaker upper-ocean overturning, *Nature*, 542, 215–218, <https://doi.org/10.1038/nature21068>, 2017.

- DeVries, T., Yamamoto, K., Wanninkhof, R., Gruber, N., Hauck, J., Müller, J. D., Bopp, L., Carroll, D., Carter, B., Chau, T.-T.-T., Doney, S. C., Gehlen, M., Gloege, L., Gregor, L., Henson, S., Kim, J. H., Iida, Y., Ilyina, T., Landschützer, P., Le Quéré, C., Munro, D., Nissen, C., Patara, L., Pérez, F. F., Resplandy, L., Rodgers, K. B., Schwinger, J., Séférian, R., Sicardi, V., Terhaar, J., Triñanes, J., Tsujino, H., Watson, A., Yasunaka, S., and Zeng, J.: Magnitude, Trends, and Variability of the Global Ocean Carbon Sink From 1985 to 2018, *Global Biogeochemical Cycles*, 37, e2023GB007780, <https://doi.org/10.1029/2023GB007780>, 2023.
- 820 Doney, S. C., Lindsay, K., Caldeira, K., Campin, J.-M., Drange, H., Dutay, J.-C., Follows, M., Gao, Y., Gnanadesikan, A., Gruber, N., Ishida, A., Joos, F., Madec, G., Maier-Reimer, E., Marshall, J. C., Matear, R. J., Monfray, P., Mouchet, A., Najjar, R., Orr, J. C., Plattner, G.-K., Sarmiento, J., Schlitzer, R., Slater, R., Totterdell, I. J., Weirig, M.-F., Yamanaka, Y., and Yool, A.: Evaluating global ocean carbon models: The importance of realistic physics, *Global Biogeochemical Cycles*, 18, <https://doi.org/10.1029/2003GB002150>, 2004.
- 825 Doney, S. C., Lima, I., Feely, R. A., Glover, D. M., Lindsay, K., Mahowald, N., Moore, J. K., and Wanninkhof, R.: Mechanisms governing interannual variability in upper-ocean inorganic carbon system and air–sea CO₂ fluxes: Physical climate and atmospheric dust, *Deep Sea Research Part II: Topical Studies in Oceanography*, 56, 640–655, <https://doi.org/10.1016/j.dsr2.2008.12.006>, 2009.
- 830 Donlon, C., Martin, M., Stark, J., Roberts-Jones, J., Fiedler, E., and Wimmer, W.: The Operational Sea Surface Temperature and Sea Ice Analysis (OSTIA) system, *Remote Sensing of the Environment*, <https://doi.org/10.1016/j.rse.2010.10.017>, 2012.
- Eggleston, E. S., Sabine, C. L., and Morel, F. M. M.: Revelle revisited: Buffer factors that quantify the response of ocean chemistry to changes in DIC and alkalinity, *Global Biogeochemical Cycles*, 24, <https://doi.org/10.1029/2008GB003407>, 2010.
- EUMETSAT: OSI SAF Global medium resolution sea ice concentration climate data record 2002–2020 (v3.0), Ocean and Sea Ice Satellite Application Facility, https://doi.org/10.15770/EUM_SAF_OSI_0015, 2022.
- 835 Evensen, G.: The Ensemble Kalman Filter: Theoretical Formulation and Practical Implementation, *Ocean Dynamics*, 53, 343–367, <https://doi.org/10.1007/s10236-003-0036-9>, 2003.
- Fay, A. R. and McKinley, G. A.: Global open-ocean biomes: mean and temporal variability, *Earth System Science Data*, 6, 273–284, <https://doi.org/10.5194/essd-6-273-2014>, 2014.
- 840 Fennel, K., Mattern, J. P., Doney, S. C., Bopp, L., Moore, A. M., Wang, B., and Yu, L.: Ocean biogeochemical modelling, *Nature Reviews Methods Primers*, 2, 1–21, <https://doi.org/10.1038/s43586-022-00154-2>, 2022.
- Ford, D. and Barciela, R.: Global marine biogeochemical reanalyses assimilating two different sets of merged ocean colour products, *Remote Sensing of Environment*, 203, 40–54, <https://doi.org/10.1016/j.rse.2017.03.040>, *earth Observation of Essential Climate Variables*, 2017.
- Fransner, F., Counillon, F., Bethke, I., Tjiputra, J., Samuelsen, A., Nummelin, A., and Olsen, A.: Ocean Biogeochemical Predictions—Initialization and Limits of Predictability, *Frontiers in Marine Science*, 7, 508 048, <https://doi.org/10.3389/fmars.2020.00386>, 2020.
- 845 Friedlingstein, P., O’Sullivan, M., Jones, M. W., Andrew, R. M., Bakker, D. C. E., Hauck, J., Landschützer, P., Le Quéré, C., Luijkx, I. T., Peters, G. P., Peters, W., Pongratz, J., Schwingshackl, C., Sitch, S., Canadell, J. G., Ciais, P., Jackson, R. B., Alin, S. R., Anthoni, P., Barbero, L., Bates, N. R., Becker, M., Bellouin, N., Decharme, B., Bopp, L., Brasika, I. B. M., Cadule, P., Chamberlain, M. A., Chandra, N., Chau, T.-T.-T., Chevallier, F., Chini, L. P., Cronin, M., Dou, X., Enyo, K., Evans, W., Falk, S., Feely, R. A., Feng, L., Ford, D. J., Gasser, T., Ghattas, J., Gkritzalis, T., Grassi, G., Gregor, L., Gruber, N., Gürses, Ö., Harris, I., Hefner, M., Heinke, J., Houghton, R. A., Hurtt, G. C., Iida, Y., Ilyina, T., Jacobson, A. R., Jain, A., Jarníková, T., Jersild, A., Jiang, F., Jin, Z., Joos, F., Kato, E., Keeling, R. F., Kennedy, D., Klein Goldewijk, K., Knauer, J., Korsbakken, J. I., Körtzinger, A., Lan, X., Lefèvre, N., Li, H., Liu, J., Liu, Z., Ma, L., Marland, G., Mayot, N., McGuire, P. C., McKinley, G. A., Meyer, G., Morgan, E. J., Munro, D. R., Nakaoka, S.-I., Niwa, Y., O’Brien, K. M., Olsen, A., Omar, A. M., Ono, T., Paulsen, M., Pierrot, D., Pockock, K., Poulter, B., Powis, C. M., Rehder, G., Resplandy, L., Robertson, E., Rödenbeck, C., Rosan, T. M., Schwinger, J., Séférian, R., Smallman, T. L., Smith, S. M., Sospedra-Alfonso, R., Sun, Q.,
- 855

- Sutton, A. J., Sweeney, C., Takao, S., Tans, P. P., Tian, H., Tilbrook, B., Tsujino, H., Tubiello, F., van der Werf, G. R., van Ooijen, E., Wanninkhof, R., Watanabe, M., Wimart-Rousseau, C., Yang, D., Yang, X., Yuan, W., Yue, X., Zaehle, S., Zeng, J., and Zheng, B.: Global Carbon Budget 2023, *Earth System Science Data*, 15, 5301–5369, <https://doi.org/10.5194/essd-15-5301-2023>, 2023.
- 860 Gaspari, G. and Cohn, S. E.: Construction of correlation functions in two and three dimensions, *Quarterly Journal of the Royal Meteorological Society*, 125, 723–757, <https://doi.org/10.1002/qj.49712555417>, 1999.
- Gasparin, F., Cravatte, S., Greiner, E., Perruche, C., Hamon, M., Van Gennip, S., and Lellouche, J.-M.: Excessive productivity and heat content in tropical Pacific analyses: Disentangling the effects of in situ and altimetry assimilation, *Ocean Modelling*, 160, 101768, <https://doi.org/https://doi.org/10.1016/j.ocemod.2021.101768>, 2021.
- Gent, P. R. and McWilliams, J. C.: Isopycnal Mixing in Ocean Circulation Models, *Journal of Physical Oceanography*, 20, 150–155, 865 [https://doi.org/10.1175/1520-0485\(1990\)020<0150:IMIOCM>2.0.CO;2](https://doi.org/10.1175/1520-0485(1990)020<0150:IMIOCM>2.0.CO;2), 1990.
- Gerber, M., Joos, F., Vázquez-Rodríguez, M., Touratier, F., and Goyet, C.: Regional air-sea fluxes of anthropogenic carbon inferred with an Ensemble Kalman Filter, *Global Biogeochemical Cycles*, 23, <https://doi.org/10.1029/2008GB003247>, 2009.
- Gloege, L., McKinley, G. A., Landschützer, P., Fay, A. R., Frölicher, T. L., Fyfe, J. C., Ilyina, T., Jones, S., Lovenduski, N. S., Rodgers, K. B., Schlunegger, S., and Takano, Y.: Quantifying Errors in Observationally Based Estimates of Ocean Carbon Sink Variability, *Global* 870 *Biogeochemical Cycles*, 35, e2020GB006788, <https://doi.org/10.1029/2020GB006788>, 2021.
- Good, S., Fiedler, E., Mao, C., Martin, M., Maycock, A., Reid, R., Roberts-Jones, J., Searle, T., Waters, J., While, J., and Worsfold, M.: The Current Configuration of the OSTIA System for Operational Production of Foundation Sea Surface Temperature and Ice Concentration Analyses, *Remote Sens.*, 12, 720, <https://doi.org/10.3390/rs12040720>, 2020.
- Good, S. A., Martin, M. J., and Rayner, N. A.: EN4: Quality Controlled Ocean Temperature and Salinity Profiles and Monthly Objective 875 Analyses with Uncertainty Estimates, *Journal of Geophysical Research: Oceans*, <https://doi.org/10.1002/2013JC009067>, 2013.
- Gray, A. R., Johnson, K. S., Bushinsky, S. M., Riser, S. C., Russell, J. L., Talley, L. D., Wanninkhof, R., Williams, N. L., and Sarmiento, J. L.: Autonomous Biogeochemical Floats Detect Significant Carbon Dioxide Outgassing in the High-Latitude Southern Ocean, *Geophysical Research Letters*, 45, 9049–9057, <https://doi.org/10.1029/2018GL078013>, 2018.
- Gruber, N., Gloor, M., Fletcher, S. E. M., Doney, S. C., Dutkiewicz, S., Follows, M. J., Gerber, M., Jacobson, A. R., Joos, F., Lindsay, K., 880 Menemenlis, D., Mouchet, A., Müller, S. A., Sarmiento, J. L., and Takahashi, T.: Oceanic sources, sinks, and transport of atmospheric CO₂, *Global Biogeochemical Cycles*, 23, <https://doi.org/10.1029/2008GB003349>, 2009.
- Gürses, O., Oziel, L., Karakuş, O., Sidorenko, D., Völker, C., Ye, Y., Zeising, M., Butzin, M., and Hauck, J.: Ocean biogeochemistry in the coupled ocean–sea ice–biogeochemistry model FESOM2.1–REcoM3, *Geoscientific Model Development*, 16, 4883–4936, <https://doi.org/10.5194/gmd-16-4883-2023>, 2023.
- 885 Hauck, J., Völker, C., Wang, T., Hoppema, M., Losch, M., and Wolf-Gladrow, D. A.: Seasonally Different Carbon Flux Changes in the Southern Ocean in Response to the Southern Annular Mode, *Global Biogeochemical Cycles*, 27, 1236–1245, <https://doi.org/10.1002/2013GB004600>, 2013.
- Hauck, J., Zeising, M., Le Quéré, C., Gruber, N., Bakker, D. C. E., Bopp, L., Chau, T. T. T., Gürses, Ö., Ilyina, T., Landschützer, P., Lenton, A., Resplandy, L., Rödenbeck, C., Schwinger, J., and Séférian, R.: Consistency and Challenges in the Ocean Carbon Sink Estimate for the 890 Global Carbon Budget, *Frontiers in Marine Science*, 7, 571720, <https://doi.org/10.3389/fmars.2020.571720>, 2020.
- Hauck, J., Gregor, L., Nissen, C., Patara, L., Hague, M., Mongwe, P., Bushinsky, S., Doney, S. C., Gruber, N., Le Quéré, C., Manizza, M., Mazloff, M., Monteiro, P. M. S., and Terhaar, J.: The Southern Ocean Carbon Cycle 1985–2018: Mean, Seasonal Cycle, Trends, and Storage, *Global Biogeochemical Cycles*, 37, e2023GB007848, <https://doi.org/10.1029/2023GB007848>, 2023a.

- Hauck, J., Nissen, C., Landschützer, P., Rödenbeck, C., Bushinsky, S., and Olsen, A.: Sparse observations induce large biases in estimates of the global ocean CO₂ sink: an ocean model subsampling experiment, *Philosophical Transactions of the Royal Society A*, 381, 20220063, <https://doi.org/10.1098/rsta.2022.0063>, 2023b.
- Hemmings, J. C. P., Barciela, R. M., and Bell, M. J.: Ocean Color Data Assimilation with Material Conservation for Improving Model Estimates of Air-Sea CO₂ Flux, *Journal of Marine Research*, 66, 87–126, <https://doi.org/10.1357/002224008784815739>, 2008.
- Hohn, S.: Coupling and decoupling of biogeochemical cycles in marine ecosystems, Ph.D. thesis, University of Bremen, 2008.
- Johnson, K. S., Plant, J. N., Coletti, L. J., Jannasch, H. W., Sakamoto, C. M., Riser, S. C., Swift, D. D., Williams, N. L., Boss, E., Haëntjens, N., Talley, L. D., and Sarmiento, J. L.: Biogeochemical sensor performance in the SOCCOM profiling float array, *Journal of Geophysical Research: Oceans*, 122, 6416–6436, <https://doi.org/10.1002/2017JC012838>, 2017.
- Johnson, R., Strutton, P. G., Wright, S. W., McMinn, A., and Meiners, K. M.: Three Improved Satellite Chlorophyll Algorithms for the Southern Ocean, *Journal of Geophysical Research: Oceans*, 118, 3694–3703, <https://doi.org/10.1002/jgrc.20270>, 2013.
- Jones, S. D., Le Quéré, C., and Rödenbeck, C.: Autocorrelation characteristics of surface ocean pCO₂ and air-sea CO₂ fluxes, *Global Biogeochemical Cycles*, 26, <https://doi.org/10.1029/2010GB004017>, 2012.
- Joos, F. and Spahni, R.: Rates of change in natural and anthropogenic radiative forcing over the past 20,000 years, *Proceedings of the National Academy of Sciences*, 105, 1425–1430, <https://doi.org/10.1073/pnas.0707386105>, 2008.
- Karakuş, O., Völker, C., Iversen, M., Hagen, W., Wolf-Gladrow, D., Fach, B., and Hauck, J.: Modeling the Impact of Macrozooplankton on Carbon Export Production in the Southern Ocean, *Journal of Geophysical Research: Oceans*, 126, e2021JC017315, <https://doi.org/10.1029/2021JC017315>, 2021.
- Keppler, L. and Landschützer, P.: Regional wind variability modulates the Southern Ocean carbon sink, *Scientific reports*, 9, 7384, <https://doi.org/10.1038/s41598-019-43826-y>, 2019.
- Kriest, I., Kähler, P., Koeve, W., Kvale, K., Sauerland, V., and Oschlies, A.: One size fits all? Calibrating an ocean biogeochemistry model for different circulations, *Biogeosciences*, 17, 3057–3082, <https://doi.org/10.5194/bg-17-3057-2020>, 2020.
- Large, W. G., McWilliams, J. C., and Doney, S. C.: Oceanic vertical mixing: A review and a model with a nonlocal boundary layer parameterization, *Reviews of Geophysics*, 32, 363–403, <https://doi.org/10.1029/94RG01872>, 1994.
- Laurindo, L. C., Mariano, A. J., and Lumpkin, R.: An improved near-surface velocity climatology for the global ocean from drifter observations, *Deep Sea Research Part I: Oceanographic Research Papers*, 124, 73–92, <https://doi.org/10.1016/j.dsr.2017.04.009>, 2017.
- Lauvset, S. K., Key, R. M., Olsen, A., van Heuven, S., Velo, A., Lin, X., Schirnack, C., Kozyr, A., Tanhua, T., Hoppema, M., Jutterström, S., Steinfeldt, R., Jeansson, E., Ishii, M., Perez, F. F., Suzuki, T., and Watelet, S.: A New Global Interior Ocean Mapped Climatology: The 1°x1° GLODAP Version 2, *Earth System Science Data*, 8, 325–340, <https://doi.org/10.5194/essd-8-325-2016>, 2016.
- Lauvset, S. K., Lange, N., Tanhua, T., Bittig, H. C., Olsen, A., Kozyr, A., Álvarez, M., Azetsu-Scott, K., Brown, P. J., Carter, B. R., Cotrim da Cunha, L., Hoppema, M., Humphreys, M. P., Ishii, M., Jeansson, E., Murata, A., Müller, J. D., Perez, F. F., Schirnack, C., Steinfeldt, R., Suzuki, T., Ulfsbo, A., Velo, A., Woosley, R. J., and Key, R.: The annual update GLODAPv2.2023: the global interior ocean biogeochemical data product, *Earth System Science Data Discussions*, 2024, 1–32, <https://doi.org/10.5194/essd-2023-468>, 2024a.
- Lauvset, S. K., Lange, N., Tanhua, T., Bittig, H. C., Olsen, A., Kozyr, A., Álvarez, M., Azetsu-Scott, K., Brown, P. J., Carter, B. R., Cotrim da Cunha, L., Hoppema, M., Humphreys, M. P., Ishii, M., Jeansson, E., Murata, A., Müller, J. D., Pérez, F. F., Schirnack, C., Steinfeldt, R., Suzuki, T., Ulfsbo, A., Velo, A., Woosley, R. J., and Key, R. M.: The annual update GLODAPv2.2023: the global interior ocean biogeochemical data product, *Earth System Science Data*, 16, 2047–2072, <https://doi.org/10.5194/essd-16-2047-2024>, 2024b.

- Le Quéré, C., Raupach, M. R., Canadell, J. G., Marland, G., Bopp, L., Ciais, P., Conway, T. J., Doney, S. C., Feely, R. A., Foster, P., Friedlingstein, P., Gurney, K., Houghton, R. A., House, J. I., Huntingford, C., Levy, P. E., Lomas, M. R., Majkut, J., Metzli, N., Ometto, J. P., Peters, G. P., Prentice, I. C., Randerson, J. T., Running, S. W., Sarmiento, J. L., Schuster, U., Sitch, S., Takahashi, T., Viovy, N., Van Der Werf, G. R., and Woodward, F. I.: Trends in the sources and sinks of carbon dioxide, *Nature Geoscience*, 2, 831–836, <https://doi.org/10.1038/ngeo689>, 2009.
- Lebehot, A. D., Halloran, P. R., Watson, A. J., McNeill, D., Ford, D. A., Landschützer, P., Lauvset, S. K., and Schuster, U.: Reconciling Observation and Model Trends in North Atlantic Surface CO₂, *Global Biogeochemical Cycles*, 33, 1204–1222, <https://doi.org/10.1029/2019GB006186>, 2019.
- 935 Li, H., Ilyina, T., Müller, W. A., and Sienz, F.: Decadal predictions of the North Atlantic CO₂ uptake, *Nature Communications*, 7, 1–7, <https://doi.org/10.1038/ncomms11076>, 2016.
- Liao, E., Resplandy, L., Liu, J., and Bowman, K. W.: Amplification of the ocean carbon sink during El Niños: Role of poleward Ekman transport and influence on atmospheric CO₂, *Global Biogeochemical Cycles*, 34, <https://doi.org/10.1029/2020GB006574>, 2020.
- Long, M. C., Stephens, B. B., McKain, K., Sweeney, C., Keeling, R. F., Kort, E. A., Morgan, E. J., Bent, J. D., Chandra, N., Chevallier, F., Commane, R., Daube, B. C., Krummel, P. B., Loh, Z., Luijkx, I. T., Munro, D., Patra, P., Peters, W., Ramonet, M., Rödenbeck, C., Stavert, A., Tans, P., and Wofsy, S. C.: Strong Southern Ocean carbon uptake evident in airborne observations, *Science*, 374, 1275–1280, <https://doi.org/10.1126/science.abi4355>, 2021.
- 945 Mamnun, N., Völker, C., Krumscheid, S., Vrekoussis, M., and Nerger, L.: Global Sensitivity Analysis of a One-Dimensional Ocean Biogeochemical Model, *Socio-Environmental Systems Modelling*, 5, 18 613, <https://doi.org/10.18174/sesmo.18613>, 2023.
- Mayot, N., Le Quéré, C., Rödenbeck, C., Bernardello, R., Bopp, L., Djeutchouang, L. M., Gehlen, M., Gregor, L., Gruber, N., Hauck, J., Iida, Y., Ilyina, T., Keeling, R. F., Landschützer, P., Manning, A. C., Patara, L., Resplandy, L., Schwinger, J., Séférian, R., Watson, A. J., Wright, R. M., and Zeng, J.: Climate-driven variability of the Southern Ocean CO₂ sink, *Philosophical Transactions of the Royal Society A: Mathematical, Physical and Engineering Sciences*, 381, 20220 055, <https://doi.org/10.1098/rsta.2022.0055>, 2023.
- Mayot, N., Buitenhuis, E. T., Wright, R. M., Hauck, J., Bakker, D. C. E., and Le Quéré, C.: Constraining the trend in the ocean CO₂ sink during 2000–2022, *Nature Communications*, 15, 1–11, <https://doi.org/10.1038/s41467-024-52641-7>, 2024.
- 955 Menemenlis, D., Fukumori, I., and Lee, T.: Using Green's Functions to Calibrate an Ocean General Circulation Model, *Monthly Weather Review*, 133, 1224–1240, <https://doi.org/10.1175/MWR2912.1>, 2005.
- Mu, L., Nerger, L., Streffing, J., Tang, Q., Niraula, B., Zampieri, L., Loza, S. N., and Goessling, H. F.: Sea-Ice Forecasts With an Upgraded AWI Coupled Prediction System, *Journal of Advances in Modeling Earth Systems*, 14, e2022MS003 176, <https://doi.org/10.1029/2022MS003176>, 2022.
- 960 Müller, J. D., Gruber, N., Carter, B., Feely, R., Ishii, M., Lange, N., Lauvset, S. K., Murata, A., Olsen, A., Pérez, F. F., Sabine, C., Tanhua, T., Wanninkhof, R., and Zhu, D.: Decadal Trends in the Oceanic Storage of Anthropogenic Carbon From 1994 to 2014, *AGU Advances*, 4, e2023AV000 875, <https://doi.org/10.1029/2023AV000875>, 2023.
- Nerger, L., Hiller, W., and Schröter, J.: PDAF - THE PARALLEL DATA ASSIMILATION FRAMEWORK: EXPERIENCES WITH KALMAN FILTERING, in: *Use of High Performance Computing in Meteorology*, pp. 63–83, WORLD SCIENTIFIC, Singapore, https://doi.org/10.1142/9789812701831_0006, 2005.
- 965 Nerger, L., Janjić, T., Schröter, J., and Hiller, W.: A Unification of Ensemble Square Root Kalman Filters, *Monthly Weather Review*, 140, 2335–2345, <https://doi.org/10.1175/MWR-D-11-00102.1>, 2012.

- Nerger, L., Tang, Q., and Mu, L.: Efficient ensemble data assimilation for coupled models with the Parallel Data Assimilation Framework: example of AWI-CM (AWI-CM-PDAF 1.0), *Geoscientific Model Development*, 13, 4305–4321, <https://doi.org/10.5194/gmd-13-4305-2020>, 2020.
- 970 Nerger, L., Tang, Q., and Mu, L.: The PDAF model binding for AWI-CM (AWI-CM-PDAF version 1.0 update 1), <https://doi.org/10.5281/zenodo.3822030>, 2024.
- Orr, J. C. and Epitalon, J.-M.: Improved routines to model the ocean carbonate system: mocsy 2.0, *Geoscientific Model Development*, 8, 485–499, <https://doi.org/10.5194/gmd-8-485-2015>, 2015.
- 975 Orr, J. C., Najjar, R. G., Aumont, O., Bopp, L., Bullister, J. L., Danabasoglu, G., Doney, S. C., Dunne, J. P., Dutay, J.-C., Graven, H., Griffies, S. M., John, J. G., Joos, F., Levin, I., Lindsay, K., Matear, R. J., McKinley, G. A., Mouchet, A., Oschlies, A., Romanou, A., Schlitzer, R., Tagliabue, A., Tanhua, T., and Yool, A.: Biogeochemical protocols and diagnostics for the CMIP6 Ocean Model Intercomparison Project (OMIP), *Geoscientific Model Development*, 10, 2169–2199, <https://doi.org/10.5194/gmd-10-2169-2017>, 2017.
- 980 Park, J.-Y., Stock, C. A., Yang, X., Dunne, J. P., Rosati, A., John, J., and Zhang, S.: Modeling global ocean biogeochemistry with physical data assimilation: a pragmatic solution to the equatorial instability, *Journal of Advances in modeling earth systems*, 10, 891–906, <https://doi.org/10.1002/2017MS001223>, 2018.
- Pérez, F. F., Becker, M., Goris, N., Gehlen, M., López-Mozos, M., Tjiputra, J., Olsen, A., Müller, J. D., Huertas, I. E., Chau, T. T., Cainzos, V., Velo, A., Benard, G., Hauck, J., Gruber, N., and Wanninkhof, R.: An Assessment of CO₂ Storage and Sea-Air Fluxes for the Atlantic Ocean and Mediterranean Sea Between 1985 and 2018, *Global Biogeochemical Cycles*, 38, e2023GB007862, <https://doi.org/10.1029/2023GB007862>, 2024.
- 985 Peylin, P., Law, R. M., Gurney, K. R., Chevallier, F., Jacobson, A. R., Maki, T., Niwa, Y., Patra, P. K., Peters, W., Rayner, P. J., Rödenbeck, C., van der Laan-Luijkx, I. T., and Zhang, X.: Global atmospheric carbon budget: results from an ensemble of atmospheric CO₂ inversions, *Biogeosciences*, 10, 6699–6720, <https://doi.org/10.5194/bg-10-6699-2013>, 2013.
- 990 Pham, D. T.: Stochastic Methods for Sequential Data Assimilation in Strongly Nonlinear Systems, *Monthly Weather Review*, 129, 1194 – 1207, [https://doi.org/10.1175/1520-0493\(2001\)129<1194:SMFSDA>2.0.CO;2](https://doi.org/10.1175/1520-0493(2001)129<1194:SMFSDA>2.0.CO;2), 2001.
- Pham, D. T., Verron, J., and Roubaud, M. C.: A singular evolutive extended Kalman filter for data assimilation in oceanography, *Journal of Marine Systems*, 16, 323–340, [https://doi.org/10.1016/S0924-7963\(97\)00109-7](https://doi.org/10.1016/S0924-7963(97)00109-7), 1998.
- Primeau, F. and Deleersnijder, E.: On the time to tracer equilibrium in the global ocean, *Ocean Science*, 5, 13–28, <https://doi.org/10.5194/os-5-13-2009>, 2009.
- 995 Raghukumar, K., Edwards, C. A., Goebel, N. L., Broquet, G., Veneziani, M., Moore, A. M., and Zehr, J. P.: Impact of assimilating physical oceanographic data on modeled ecosystem dynamics in the California Current System, *Progress in Oceanography*, 138, 546–558, <https://doi.org/https://doi.org/10.1016/j.pocean.2015.01.004>, combining Modeling and Observations to Better Understand Marine Ecosystem Dynamics, 2015.
- 1000 Regnier, P., Resplandy, L., Najjar, R. G., and Ciais, P.: The land-to-ocean loops of the global carbon cycle, *Nature*, 603, 401–410, <https://doi.org/10.1038/s41586-021-04339-9>, 2022.
- Rödenbeck, C., Bakker, D. C. E., Gruber, N., Iida, Y., Jacobson, A. R., Jones, S., Landschützer, P., Metzl, N., Nakaoka, S., Olsen, A., Park, G.-H., Peylin, P., Rodgers, K. B., Sasse, T. P., Schuster, U., Shutler, J. D., Valsala, V., Wanninkhof, R., and Zeng, J.: Data-based estimates of the ocean carbon sink variability – first results of the Surface Ocean pCO₂ Mapping intercomparison (SOCOM), *Biogeosciences*, 12, 7251–7278, <https://doi.org/10.5194/bg-12-7251-2015>, 2015.
- 1005

- Sarmiento, J. L. and Gruber, N.: Carbon Cycle, in: *Ocean Biogeochemical Dynamics*, chap. 8, pp. 318–358, Princeton University Press, 2006.
- Sathyendranath, S., Jackson, T., Brockmann, C., Brotas, V., Calton, B., Chuprin, A., Clements, O., Cipollini, P., Danne, O., Dingle, J., Donlon, C., Grant, M., Groom, S., Krasemann, H., Lavender, S., Mazeran, C., Mélin, F., Müller, D., Steinmetz, F., Valente, A., Zühlke, M., Feldman, G., Franz, B., Frouin, R., Werdell, J., and Platt, T.: ESA Ocean Colour Climate Change Initiative (Ocean_Colour_cci): Version 5.0 Data, NERC EDS Centre for Environmental Data Analysis, <https://doi.org/10.5285/1dbe7a109c0244aaad713e078fd3059a>, 2021.
- Schartau, M., Engel, A., Schröter, J., Thoms, S., Völker, C., and Wolf-Gladrow, D.: Modelling carbon overconsumption and the formation of extracellular particulate organic carbon, *Biogeosciences*, 4, 433–454, <https://doi.org/10.5194/bg-4-433-2007>, 2007.
- 1015 Scholz, P., Sidorenko, D., Gurses, O., Danilov, S., Koldunov, N., Wang, Q., Sein, D., Smolentseva, M., Rakowsky, N., and Jung, T.: Assessment of the Finite-volume Sea ice-Ocean Model (FESOM2.0) – Part 1: Description of selected key model elements and comparison to its predecessor version, *Geoscientific Model Development*, 12, 4875–4899, <https://doi.org/10.5194/gmd-12-4875-2019>, 2019.
- Scholz, P., Sidorenko, D., Danilov, S., Wang, Q., Koldunov, N., Sein, D., and Jung, T.: Assessment of the Finite-Volume Sea ice–Ocean Model (FESOM2.0) – Part 2: Partial bottom cells, embedded sea ice and vertical mixing library CVMix, *Geoscientific Model Development*, 15, 335–363, <https://doi.org/10.5194/gmd-15-335-2022>, 2022.
- 1020 Schourup-Kristensen, V., Sidorenko, D., Wolf-Gladrow, D. A., and Völker, C.: A Skill Assessment of the Biogeochemical Model REcom2 coupled to the Finite Element Sea Ice Ocean Model (FESOM 1.3), *Geoscientific Model Development*, 7, 2769–2802, <https://doi.org/10.5194/gmd-7-2769-2014>, 2014.
- Séférian, R., Bopp, L., Gehlen, M., Swingedouw, D., Mignot, J., Guilyardi, E., and Servonnat, J.: Multiyear predictability of tropical marine productivity, *Proceedings of the National Academy of Sciences*, 111, 11 646–11 651, <https://doi.org/10.1073/pnas.1315855111>, 2014.
- 1025 Sidorenko, D.: The North Atlantic circulation derived from inverse models, Ph.D. thesis, University of Bremen, 2004.
- Spring, A., Dunkl, I., Li, H., Brovkin, V., and Ilyina, T.: Trivial improvements in predictive skill due to direct reconstruction of the global carbon cycle, *Earth System Dynamics*, 12, 1139–1167, <https://doi.org/10.5194/esd-12-1139-2021>, 2021.
- Stark, J., Donlon, C., Martin, M., and McCulloch, M.: OSTIA: An Operational, High Resolution, Real-Time, Global Sea Surface Temperature Analysis System, in: *Oceans 07 IEEE Aberdeen, Conference Proceedings. Marine Challenges: Coastline to Deep Sea, IEEE*, 2007.
- 1030 Sursham, D.: Improving the Simulation and Understanding of Biologically Driven Carbon Pumps in Marine Ecosystems using an Ensemble-Based Data Assimilation Method, Ph.D. thesis, University of Reading, 2018.
- Sutton, A. J., Williams, N. L., and Tilbrook, B.: Constraining Southern Ocean CO₂ Flux Uncertainty Using Uncrewed Surface Vehicle Observations, *Geophysical Research Letters*, 48, e2020GL091 748, <https://doi.org/10.1029/2020GL091748>, 2021.
- 1035 Takahashi, T., Olafsson, J., Goddard, J. G., Chipman, D. W., and Sutherland, S. C.: Seasonal variation of CO₂ and nutrients in the high-latitude surface oceans: A comparative study, *Global Biogeochemical Cycles*, 7, 843–878, <https://doi.org/10.1029/93GB02263>, 1993.
- Tang, Q., Mu, L., Sidorenko, D., Goessling, H., Semmler, T., and Nerger, L.: Improving the ocean and atmosphere in a coupled ocean–atmosphere model by assimilating satellite sea-surface temperature and subsurface profile data, *Quarterly Journal of the Royal Meteorological Society*, 146, 4014–4029, <https://doi.org/10.1002/qj.3885>, 2020.
- 1040 Terhaar, J., Frölicher, T. L., and Joos, F.: Observation-constrained estimates of the global ocean carbon sink from Earth system models, *Biogeosciences*, 19, 4431–4457, <https://doi.org/10.5194/bg-19-4431-2022>, 2022.

- Terhaar, J., Goris, N., Müller, J. D., DeVries, T., Gruber, N., Hauck, J., Perez, F. F., and Séférian, R.: Assessment of Global Ocean Biogeochemistry Models for Ocean Carbon Sink Estimates in RECCAP2 and Recommendations for Future Studies, *Journal of Advances in Modeling Earth Systems*, 16, e2023MS003 840, <https://doi.org/10.1029/2023MS003840>, 2024.
- 1045 Timmermann, R. and Beckmann, A.: Parameterization of vertical mixing in the Weddell Sea, *Ocean Modelling*, 6, 83–100, [https://doi.org/10.1016/S1463-5003\(02\)00061-6](https://doi.org/10.1016/S1463-5003(02)00061-6), 2004.
- Tsujino, H., Urakawa, S., Nakano, H., Small, R. J., Kim, W. M., Yeager, S. G., Danabasoglu, G., Suzuki, T., Bamber, J. L., Bentsen, M., Böning, C. W., Bozec, A., Chassignet, E. P., Curchitser, E., Boeira Dias, F., Durack, P. J., Griffies, S. M., Harada, Y., Ilicak, M., Josey, S. A., Kobayashi, C., Kobayashi, S., Komuro, Y., Large, W. G., Le Sommer, J., Marsland, S. J., Masina, S., Scheinert, M., Tomita, H., 1050 Valdivieso, M., and Yamazaki, D.: JRA-55 based surface dataset for driving ocean–sea-ice models (JRA55-do), *Ocean Modelling*, 130, 79–139, <https://doi.org/10.1016/j.ocemod.2018.07.002>, 2018.
- Valsala, V. and Maksyutov, S.: Simulation and assimilation of global ocean pCO₂ and air–sea CO₂ fluxes using ship observations of surface ocean pCO₂ in a simplified biogeochemical offline model, *Tellus B: Chemical and Physical Meteorology*, 62, 821–840, <https://doi.org/10.1111/j.1600-0889.2010.00495.x>, 2010.
- 1055 Verdy, A. and Mazloff, M. R.: A data assimilating model for estimating Southern Ocean biogeochemistry, *Journal of Geophysical Research: Oceans*, 122, 6968–6988, <https://doi.org/10.1002/2016JC012650>, 2017.
- Vetra-Carvalho, S., van Leeuwen, P. J., Nerger, L., Barth, A., Altaf, M. U., Brasseur, P., Kirchgessner, P., and Beckers, J.-M.: State-of-the-art stochastic data assimilation methods for high-dimensional non-Gaussian problems, *Tellus A: Dynamic Meteorology and Oceanography*, 70, 1–43, <https://doi.org/10.1080/16000870.2018.1445364>, 2018.
- 1060 Völker, C., Wallace, D. W. R., and Wolf-Gladrow, D. A.: On the role of heat fluxes in the uptake of anthropogenic carbon in the North Atlantic, *Global Biogeochemical Cycles*, 16, 85–1–85–9, <https://doi.org/10.1029/2002GB001897>, 2002.
- Wanninkhof, R.: Relationship between wind speed and gas exchange over the ocean revisited, *Limnology and Oceanography: Methods*, 12, 351–362, <https://doi.org/10.4319/lom.2014.12.351>, 2014.
- Wanninkhof, R., Park, G.-H., Takahashi, T., Sweeney, C., Feely, R., Nojiri, Y., Gruber, N., Doney, S. C., McKinley, G. A., Lenton, A., 1065 Le Quéré, C., Heinze, C., Schwinger, J., Graven, H., and Khatiwala, S.: Global ocean carbon uptake: magnitude, variability and trends, *Biogeosciences*, 10, 1983–2000, <https://doi.org/10.5194/bg-10-1983-2013>, 2013.
- While, J., Totterdell, I., and Martin, M.: Assimilation of pCO₂ data into a global coupled physical-biogeochemical ocean model, *Journal of Geophysical Research: Oceans*, 117, <https://doi.org/10.1029/2010JC006815>, 2012.
- Williams, N. L., Juranek, L. W., Feely, R. A., Johnson, K. S., Sarmiento, J. L., Talley, L. D., Dickson, A. G., Gray, A. R., Wanninkhof, R., 1070 Russell, J. L., Riser, S. C., and Takeshita, Y.: Calculating surface ocean pCO₂ from biogeochemical Argo floats equipped with pH: An uncertainty analysis, *Global Biogeochemical Cycles*, 31, 591–604, <https://doi.org/10.1002/2016GB005541>, 2017.
- Wunsch, C.: *The Ocean Circulation Inverse Problem*, Cambridge University Press, Cambridge, England, UK, <https://doi.org/10.1017/CBO9780511629570>, 1996.