Point-by-point response to reviewer and editor comments below in blue.

Response to the editor:

In the revised manuscript we now calculate (with CO2SYS) and discuss two metrics for assessing the models’ response to alkalinity addition, i.e., OAE compared to GLODAP values: 1) the initial reduction in pCO$_2$ and the 2) the change in ocean DIC after assumed complete equilibration, µCO$_2$. The initial increase in pH and the pH increase after equilibration are calculated and discussed correspondingly.

As suggested by Reviewers 1 and 2 we have added the metric uptake efficiency, $\eta$CO$_2$, to our analysis. But unfortunately, in the first response to the reviewers the new Figure 7 showed an incorrectly calculated uptake efficiency of ~0.63. This was rightly pointed out by the editor Dr. Sulpris. We have since corrected the computation of $\eta$CO$_2$ by using TA and pCO$_2$ in CO2SYS to compute the ΔDIC directly (instead of using the computed Revelle factor to infer ΔDIC). Now, the values hover around the expected value of 0.83 (see new Figure 7):

Figure 7 NEW: Carbonate system parameters were computed for all CMIP6 ESMs, the MMM (grey line) and the GLODAP data (black line) with the CO2SYS toolbox. The results are sorted by Revelle Factor in ascending order for all panels. Shown are the TA (a), the Revelle factor (b), the TA-DIC ratio (c), initial pCO$_2$ (d), the difference in pCO$_2$ after a 100 µmol kg$^{-1}$ addition of TA (e), the uptake efficiency $\eta$CO$_2$ (f), the initial pH (g), the difference in pH for constant DIC (h), and the difference in pH for constant pCO$_2$ (i). Light blue colors indicate the unperturbed mean state in the ESMs and GLODAP, dark blue colors the initial state after OAE and green colors the state after OAE and subsequent air-sea equilibration.
We introduce the metric in the methods section:

First, we evaluate the CO2SYS output fields Revelle Factor, pH, and pCO$_2$ (partial pressure of CO$_2$ in seawater) based on the CMIP6 ESMs against the values for the GLODAP data. In a second step, we assess the initial changes in surface pCO$_2$ and pH after an addition of 100 µmol kg$^{-1}$ TA (corresponds to 102.6 mmol m$^{-3}$ TA) while keeping DIC constant. In a third step, we evaluate the CO$_2$ uptake efficiency ($\eta$CO$_2$) (Renforth and Henderson, 2017, Tyka et al., 2022) and the pH difference at constant pCO$_2$ which simulates completed air-sea CO$_2$ equilibration. Note, that this calculation has an ocean-centric perspective as it assumes constant atmospheric CO$_2$, which contradicts the motivation for ocean alkalinity enhancement to reduce atmospheric CO$_2$, and thus will only be valid for small-scale applications. The uptake efficiency metric has been previously applied in ocean model simulations with constant and non-interactive atmospheric CO$_2$ (Tyka et al., 2022; He and Tyka, 2023). We here follow this approach in our idealized calculations while acknowledging that atmospheric CO$_2$ would drop in emission-driven simulations (magnitude dependent on amount of alkalinity added; Ferrer Gonzalez et al., 2018; Lenton et al., 2018; Köhler, 2020), as in the real world, through feedbacks with the atmosphere and the land biosphere (Oschlies, 2008). The assumption of constant atmospheric CO$_2$ (and thus constant surface ocean pCO$_2$) was shown to overestimate oceanic CO$_2$ uptake by 2% on annual timescale, but by 25% on decadal timescale and further increasing on longer timescales (Oschlies, 2008).

The uptake efficiency, $\eta$CO$_2$, is the ratio of moles of CO$_2$ absorbed to moles of added alkalinity and can also be expressed as the ratio of the partial pressure sensitivities of pCO$_2$ with respect to TA and DIC (Tyka et al., 2022; Tyka et al., 2022):

$$
\eta$CO$_2$ = \Delta$DIC/\Delta$TA  #(5)
$$

For the uptake efficiency at constant pCO$_2$, the $\Delta$DIC was also computed using CO2SYS, here with TA + 100 µmol kg$^{-1}$ and the initial pCO$_2$ as input parameters.

And in the results section:

We conclude that almost all ESMs might overestimate the initial additional pCO$_2$ difference uptake in simulated OAE experiments. On the other hand, the CO$_2$ uptake efficiency computed with constant pCO$_2$ (equilibrated DIC) only differs by -0.1% to 1.1% (mean: 0.4%) from the GLODAP value, and the ESMs may thus represent equilibrium CO$_2$ uptake rather robustly.

And in the conclusions:

The CO$_2$ uptake efficiency factor, $\eta$CO$_2$, relates changes in surface DIC to alkalinity input. We computed this metric here with constant pCO$_2$ after alkalinity addition which suggests complete equilibration and neglects any reduction in atmospheric CO$_2$ due to OAE. Studies suggest that the time scale and efficiency of the equilibration can differ immensely depending on the ocean region. He and Tyka (2023) found that after one year $\eta$CO$_2$ varied between 0.2 and 0.85 and that after 10 years most locations showed an uptake fraction of 0.65–0.80. Jones et al. quantified the mean global air-sea equilibration timescale for CO2 at 4.4 months (range 0.5 to 24 months regionally). Bach et al. (2023) suggest a pragmatic time scale of 10 years for a 95% DIC equilibration after OAE measures. It is within this range of suggested equilibration time scales that the differences in simulated pCO$_2$ change between ESMs are important.

Response to Reviewer 1:

Overall, I found this to be an interesting, well-written and illuminating paper that I think will help spur improvements in model development. The separation of the TA biases into preformed TA, remineralization TA and CaCO$_3$ TA is also very useful and points to concretely implementable improvements, especially in the treatment of CaCO$_3$.

We thank the reviewer for their thoughtful and thorough review and helpful suggestions. We addressed the reviewer comments below in blue.
I recommend publication with some minor revisions (see below).

My two major comments are:

Line 254 & Figure 7

As the authors point out the biases in Revelle Factor are of great importance to mCDR. An additional metric of this that would be straightforward to add using CO2SYS and of great value to folks investigating the feasibility and cost of ocean alkalinity enhancement is the uptake efficiency factor (In our work we like to call this $\eta_{CO_2} = \frac{\partial DIC}{\partial Alk}$ at constant $pCO_2$, see https://doi.org/10.1039/D1EE01532J and https://bg.copernicus.org/articles/20/27/2023/). The metric simply indicates the number of moles of CO$_2$ taken up per mol of Alkalinity added after full equilibration (for an infinitesimal increase in Alk) and is generally ~0.8 though it is quite dependent on location (see for example He et al., 2023, https://bg.copernicus.org/articles/20/27/2023/).

I think this number is very practical because it directly represents an efficiency loss going from some alkaline substance to actual CO$_2$ drawdown and thus enters any cost estimates. Thus, I would be very curious to see how model biases affect this metric, even if just expressed as a global average or surface average.

Thank you for the suggestion. We calculated uptake efficiency $\eta_{CO_2}$ with CO2SYS and added the results to the new Figure 7 (see above)

The uptake efficiency ranges between ~0.825 and 0.835 for the models and GLODAP.

In response to Reviewer 2, we added TA in the same order to Figure 7. Models with a higher Revelle factor also have a higher uptake efficiency, $\eta_{CO_2}$, (R=0.98, Figure 7f) but overall, the CO$_2$ uptake efficiency computed with constant $pCO_2$ only differs by -0.1% to 1.1% (mean: 0.4%) from the GLODAP value, which means that the ESMs represent equilibrium CO$_2$ uptake rather robustly.

L317ff and Figure 6, panel (d). There is clearly a large amount of difference in TA* between models and also in some models these biases are clearly depth-dependent while in others they are less so. This is one of the major insights of this paper. Not being familiar with the details of each of the models tested, I am very curious about whether there is any pattern or correlation between the sophistication of the CaCO3-cycle-model in each GCM and the amount and type of bias observed?

E.g. The blue-ish models mostly overestimate TA at depth - do they have something in common in the way they treat the CaCO$_3$ dissolution? Do any of these models treat the natural occurrence and distribution of CaCO$_3$ sediments explicitly (see work by Sulpis et al and others for maps of this)? or do they only account for precipitation and redissolution? If not, then perhaps there is a spatial correlation between TA biases and occurence of CaCO$_3$ sediments? I’d love to see more discussion of this phenomenon - it’s very interesting! The discussion on L294-330 is in very general terms rather than looking at algorithm differences between the specific models that could explain the differences.

The paragraph describing the TA* bias has been reworked and amended:

The bias in TA from calcium carbonate dissolution in the interior ocean (Figure 6d) can be in absolute terms comparable to or even larger than the bias in preformed TA. The MRI model and the GFDL models have a small negative bias in TA* in the order of -10-20 mml m$^{-3}$, relatively constant with depth. The MPI and NorESM models have a slight positive TA* bias in about the same order of magnitude, also relatively constant with depth, while the UKESM, the CNRM-ESM and the IPSL-ESM exhibit TA* biases that increase with depth. The CNRM model has the largest TA* bias with about 100 mmol m$^{-3}$ at 4,000 m depth. CNRM-ESM2-1 and IPSL-CM6A-LR, have in common that they contain the same ocean model (NEMO) and the same biogeochemical model (PISCESv2). Dissolution in PISCESv2 is treated explicitly and is dependent on the calcite saturation state and the sinking speed for PIC is depth-dependent, while for other models the sinking speed is constant. In two of the models (of Figure 6d), MRI-ESM2-0 and UKESM1-0-LL, CaCO$_3$ is dissolved without a sediment, while the other models do have explicit sediment treatments where CaCO$_3$ is buried or dissolved, either
dependent on the calcite saturation state or a set rate (Planchat et al. 2023). A direct link of the treatment of CaCO$_3$ at the bottom to the bias at depths is not obvious in this case.

Figure Style comments:

As I was parsing the figures I felt some improvements in the plots could make better use of the space, aid visual parsing and generally make the paper even easier to follow. Please take these as suggestions, perhaps try them out and see if you like them.

Figure 1: Maybe expressing the MMM as a (say, dashed) line rather than an additional row would be more intuitive and allow visual comparison of each model vs the MMM?

Figure 1: does the thickness of the GLODAP line have meaning (e.g. a standard error) or is it incidental? If a standard error for the GLODAP measurement is known or can be computed it would be neat to use the thickness (using a semi-transparent color) this way (unless the certainty is so high that it reduces to a thin line of course). I think this is important as a large GLODAP uncertainty could change or weaken the conclusions.

Figure 1, Line 264 As you note Alk and DIC are highly correlated, and they are compensating variables in the carbonate system, with respect to pH, pCO$_2$ etc.

An alternative for the two panels in Figure 1 would be to plot both together as a scatter graph with DIC on the x-axis and Alk on the y axis (or vice versa). This way the exact same information is displayed but the extent of the correlation is visually immediately apparent as well. The scatter points could be labelled directly on the graph with a floating text for example. Error bars on each pint could indicate the variance of these values over the surface average.

Thank you for the suggestion. In response, we replaced Figure 1 with a scatter plot of TA versus DIC, as we agree that this might be more intuitive. This figure also contains an error estimate for the global mean GLODAP data. While the MMM is almost within the estimated GLODAP range, we see that the difference for individual models can be quite large. We also see that TA and DIC biases are highly correlated, $R = 0.99$. 

![Graph showing TA vs DIC with error bars and model labels](image-url)
Figure 1 NEW: Global mean surface total alkalinity (TA) [mmol m\(^{-3}\)] of the 14 CMIP6 models, the mult.model-mean (MMM), and GLODAP including its error estimate versus dissolved inorganic carbon (DIC) [mmol m\(^{-3}\)].

Figure 2&5: "Absolut error" → "Absolute error"

This has been corrected.

Figure 2&5: If the whitespace between globes could be reduced at all, that would make everything bigger and easier to parse (It's already tricky without looking at the PDF on a large screen).

Space between the subplots has been reduced as much as possible.

Figure 3: Ah, I see here there is a GLODAP error estimate. Great! Could this be added to Figure 1 also?

Yes, this has been added to Figure 1. Thank you for the suggestion.

Figure 3&4: Style considerations: For a colour-blind person (like myself) it is nigh impossible to know which line is which, among similar shades/hues. I would recommend blending color with different dash/dot patterns to help with this. Perhaps the error (currently dashed lines) can be given simply by a transparent shaded area?

We added linestyles to the profiles and indicated the GLODAP error estimate with shading.

Also, a lot of features of this graph occur in the upper 0.5 km and are visually cramped in a very small area. For the same reason that most models have non-uniform vertical z slices, perhaps it would be possible to plot the vertical axis on a log scale or a mixed log-linear scale. Or split the graph into two linear regions, one for 0-500 m and one 500-4000 m?

We would like to keep this figure as is since we were interested in how well the profiles overall match.

Figure 4(f) I’d adjust the x axis to not clip the values at shallow depth. Ah - I see all the X axes are coordinated. Hmmm. Not sure how to solve this. How low do the TA values go in the Arctic Ocean (last panel)? Perhaps one could plot all these as Deltas from GLODAP the same way that Figure 6? That might help with the dynamic range of the x axis (which IMHO does not necessarily have to be the same for each subpanel). I thought Figure 6 was very nice.
We adjusted the y-axis for the Arctic in Figure 4f and applied the linestyles here as in Figure 3 and added a line for MMM.

Figure 7: Again, using the thickness of the GLODAP vertical line to indicate variance would be neat.

The estimated error for the GLODAP values based on the TA and DIC GLODAP errors (see also new Figure 1) are very small for the computed parameters. We think the figure without the error would be clearer and kept the figure without error bars.

Figure 8: I found this figure rather difficult to parse. Because the three different variables have such different dynamic range on the %-scale, especially the second one (TA-DIC ratio) is virtually impossible to read off. Is this figure really necessary? I feel like most of the information content is already contained in Figure 7.

We will drop Figure 8 and instead refer to the percentage values listed in Supplement Table S2.

L87: Some other recent studies that would be worth including that also aim to be more realistic than the earlier large-scale uniform OAE simulations, i.e. near-coastal or ship-track-based releases or regional assessments.

https://doi.org/10.1002/2017EF000659
Model-Based Assessment of the CO2 Sequestration Potential of Coastal Ocean Alkalinization, Feng, Koeve, Keller, Orschlies 2017

https://doi.org/10.1029/2022EF002816
Simulated Impact of Ocean Alkalinity Enhancement on Atmospheric CO2 Removal in the Bering Sea,
Weng et al., 2022
https://doi.org/10.5194/bg-20-27-2023
Limits and CO2 equilibration of near-coast alkalinity enhancement, He and Tyka, 2023
https://doi.org/10.5194/egusphere-egu23-9305
Atmospheric CO2 removal by alkalinity enhancement in the North Sea, Liu et al. 2023

L87 has been appended to include most of the above suggested references and now reads:

Now, more and more projects are underway or in planning that seek to apply more realistic scenarios for OAE e.g., in regional OAE applications (Butenschöhn et al. (2021), Wang et al. (2023)) or coastal applications (Feng et al. (2017), He and Tyka (2023)), which is why a model evaluation is even more important.

L149: Since the carbonate system isn’t linear wrt TA and DIC, does it make sense to first area-average the TA and DIC values and “then” put them through the CO2SYS calculation? It seems to me it would be more accurate to compute Revelle, pH, pCO2 etc for each surface location and or time and “then” do the area-weighted average of each metric. Perhaps over the range of values encountered the system is linear enough and this doesn’t make much of a difference, but I’m not sure.

This study was meant to introduce the issue of alkalinity and DIC biases in ESMs, their implications for assessing model OAE experiments in an idealized approach and to suggest some potential areas for model improvements. The suggested approach would certainly be a worthwhile follow-up study, also to look at regional differences with regard to the here described biases.
Responses to reviewer 2:

We thank Alban Planchat for his thoughtful and thorough review and helpful suggestions. We addressed the reviewer comments below in blue.

Review of Hinrichs et al. (2023)

Overall comment:
In overall, this work is interesting and innovative in its approach to bias assessment in OAE modelling studies using ESMs. Indeed, the study pushes a CMIP6 bias analysis (essentially focused on alkalinity, but also on DIC) towards a concrete, but idealized, CDR experiment. The approach used to decompose the alkalinity bias is broadly consistent with Planchat et al. (2023), thus reinforcing the message to the modelling community around the representation of alkalinity and the carbonate pump in ESMS. The writing of the manuscript could nevertheless be partly improved, in particular some parts could be restructured to gain clarity. Many technical corrections should be made.

Specific comments

A. Main comments

1. In the decomposition of the alkalinity bias, it is repeatedly mentioned that “preformed alkalinity” refers to physical biases for the ocean, but it also contains biases of both organic matter and CaCO3 production (e.g., l. 17 ; l. 121-125 ; l. 218 ; l. 291 ; l. 294)

We amended L 17 with:

The decomposition of the global mean alkalinity biases into contributions from physical processes (preformed alkalinity) which includes i) the physical redistribution of biased alkalinity originating from the soft tissue and carbonates pumps, ii) remineralization, and iii) carbonate formation and dissolution showed that the bias stemming from the physical redistribution of alkalinity is dominant.

We amended the method part L121-125 on preformed alkalinity with:

Preformed Alkalinity also refers to the physical redistribution of alkalinity biases stemming originally from soft tissue and carbonate pumps and the upwelling of water masses with biased alkalinity.

L221 was amended with the sentence:
The first part, preformed alkalinity, includes the advection and upwelling of already biased water masses.

2. The direct link made with the CO2 uptake in the OAE idealized experiment is confusing for me (e.g., 274, l. 327): a relative pCO2 difference should not directly be a relative CO2 uptake difference, or should it?

According to me, we have:

\[ \frac{d\text{DIC}}{d\text{DIC}} = 1/Re \cdot \frac{dp\text{CO2}}{p\text{CO2}} \]

An ESM with a greater relative variation of pCO2 resulting from alkalinity addition, also has a greater Revelle factor. So, the product of the right member of the above equation should at least partly compensate. To access a first approximation of the CO2 uptake associated with alkalinity addition, the relative variation of DIC should be expressed in function of the relative variation of alkalinity. I think that with what is currently offered, it is possible to assess the bias resulting from OAE in ESMS concerning acidification, but I’m confused by the direct link made with the CO2 uptake.

Thank you for bringing up this point. We have now added the parameter uptake efficiency (\(\eta\text{CO2}\)) to our back of the envelope calculation. As you said, the uptake efficiency depends on the ratio of pCO2 change over pCO2 state with respect to total DIC over alkalinity change (or addition) and the Revelle factor:

\[ \eta\text{CO2} = \frac{d\text{DIC}}{d\text{Alk}} \text{ at constant pCO2} \]

As you said, the inherent model states of pCO2, TA and DIC lead to slightly different efficiencies for a constant alkalinity addition. On your suggestion, we added the TA plot into Figure 7 as well (see above). And discuss the correlation between the parameters in the results section:
The global mean Revelle Factor from the CO2SYS computation for the GLODAP dataset is with 10.19 the third lowest in our compilation and thus almost all ESMs have a higher Revelle Factor than the GLODAP data, ranging from 10.18 to 10.54 (Figure 7b). The Revelle factor is anti-correlated to the average TA-DIC-ratio ($R=0.99$, Figure 7c). Also, the order of surface pH ($R=0.96$, Figure 7g) and $pCO_2$ ($R=0.97$, Figure 7d) values corresponds largely to each model’s rank in Revelle Factor (and thus also with TA-DIC-ratio). Models with a higher Revelle factor than GLODAP have a lower buffer capacity, which leads to already higher $pCO_2$ values (290 to 314 µatm) and lower pH (8.12 to 8.17) than in GLODAP ($pCO_2$: 292 µatm, pH: 8.16). Those models also show a greater initial reduction in surface ocean $pCO_2$ for the hypothetical addition of 100 µmol kg$^{-1}$ of TA ($R=0.99$, Figure 7e) than GLODAP (-92 µatm), ranging from a 91 µatm to a 104 µatm decrease in $pCO_2$. Models with a higher Revelle factor also have a higher uptake efficiency, $\eta_{CO_2}$ ($R=0.98$, Figure 7f). The initial change in pH after alkalinity addition (Figure 7h) is about an order of magnitude larger than the change in pH after complete air-sea equilibration at constant atmospheric $CO_2$ (Figure 7i). The respective changes in pH (unequilibrated / equilibrated at constant atm $CO_2$) have a higher correlation to TA ($R=0.92$, $R=0.99$) than to the Revelle factor ($R=0.83$, $R=0.63$).

**B. Minor comments**

1) The second part of the abstract could be improved and made more readable. In particular, I do not understand: “We find that the degree of compensation of DIC and alkalinity biases at the surface is more important for the marine $CO_2$ uptake capacity than the alkalinity biases themselves.”

We adapted the paragraph, which now reads:

The relative biases of alkalinity versus DIC at the surface affect the Revelle factor and therefore the initial $pCO_2$ reduction after alkalinity addition. The global mean surface alkalinity bias relative to GLODAPv2 in the different models ranges from ≈-85 mmol m$^{-3}$ (-3.6%) to +50 mmol m$^{-3}$ (+2.1%) (mean: -25 mmol m$^{-3}$ or -1.1%). For DIC the relative bias ranges from -55 mmol m$^{-3}$ (-2.6%) to 53 mmol m$^{-3}$ (+2.5%) (mean: -13 mmol m$^{-3}$ or -0.6%). All but two of the CMIP6 models evaluated here overestimate the Revelle factor at the surface by up to 3.4% and thus overestimate the initial $pCO_2$ reduction after alkalinity addition by up to 13%. The uptake efficiency, $\eta_{CO_2}$, then takes into account that a higher Revelle factor and a higher initial $pCO_2$ reduction after alkalinity addition and equilibration mostly compensate, so that resulting DIC differences in the models are small (-0.1% to 1.1%). The overestimation of the initial $pCO_2$ reduction has to be taken into account when reporting on efficiencies of ocean alkalinity enhancement experiments using CMIP6 models especially as long as the $CO_2$ equilibrium is not reached.

2) (l. 25-26); and l. 27-29 are too heavy, Broke this up into two sentences.

perhaps misplaced, and with a unit error (mmol/kg instead of mmol/m$^3$ as l. 209-211).

Thanks for pointing this out, it should be mmol/m$^3$. This has been corrected.

3) For the sake of clarity and consistency throughout the manuscript, I would suggest that alkalinity and DIC should always be shared in mmol/m$^3$ rather than mixing mmol/m$^3$ and umol/kg.

Changed to mmol/m$^3$ in Figure 1 and text where applicable.

It would simply be necessary to clarify how the conversions are done for GLODAPv2 and for using CO2SYS in the methodology (see l. 28-29, l. 149-152, Fig. 1 etc.).

For the model evaluation the GLODAP TA and DIC data is converted from units of µmol kg$^{-1}$ to mmol m$^{-3}$ using the potential density computed from GLODAP salinity and temperature data. We added this to the methods section.

4) Table 1: It would be great also to share the ensemble member for each ESM, rather than writing “the first available ensemble member” l. 109, which is slightly confusing.

The ensemble member has been added to Table 1.
5) The marine biogeochemical model of CNRM-ESM2-1 (not CNRM-ESM-2-1) is PISCESv2-gas. The one for IPSL-CM6A-LR is PISCESv2. The one for NorESM2-LM/MM is iHAMOCC.

This has been corrected.

Why do you precise the grid only for a few ocean models? If you wish to precise it, add it in the “ocean horizontal resolution” column and just precise maybe whether it is a tripolar grid or not.

Added information about tripolar grids to grid column.

6) Why do you precise the grid only for a few ocean models? If you wish to precise it, add it in the “ocean horizontal resolution” column and just precise maybe whether it is a tripolar grid or not.

Added information about tripolar grids to grid column.

I. 109-119: These lines would benefit from restructuring.

The TA* method: are there limitations to this decomposition of the biases?

Koeve et al. (2014) discuss the limitations as follows: although the computation of TA* according to equations (2) to (4) reproduces tracer-based simulated TA* robustly in most of the global ocean, higher uncertainties occur in the Atlantic and in the 500m to a 1,000m layer in the Pacific and Indian ocean.

In our study we focus on the TA* results for the global mean ocean.

Another caveat that was mentioned by Koeve et al. is related to the issue that AOU overestimates true oxygen utilization by 20–25%. Hence TAR computed from AOU is probably also overestimated by this percentage. TAR is small and, in our study, we focus on the implication of TA* biases in ESMs and potential remedies for these biases.

This short discussion of the above-mentioned caveats of the Koeve method was added to the discussion part.

What are the advantages of it rather than the one suggested in Sarmiento and Gruber (2006)? Not using DIC?

In some ways the methods are similar. Both are partly based on Feely et al. (2002). E.g., the TA' computed in Koeve et al. (2014) is similar to δC_{sol}. Koeve then aims to further separate the influence of ocean physics from the impact of carbonate dissolution. Although, as the reviewer correctly points out the physical distribution of TA includes the distribution of TA biases from the biological and the carbonate pump as well. Another difference is that preformed alkalinity is computed for each grid point and therefore is resolved spatially (but here shown as global profile only).

We added a short discussion on these methods in the revised manuscript.

7) Eq. (3) and (4): why do you use phosphate for PO in Eq. (3) and then nitrate in Eq. (4) for TA* rather phosphate (or nitrate) in both cases?

We were following the equations proposed by Koeve et al. (2014) where using PO (NO) is proposed for computing TA^0 interchangeably and no3 for computing TA* in the manuscript.

Here we recomputed TA^0 and the TA* using NO not PO, below are both ways to compare.

TA^0 computed with PO:
TA^0 computed with NO:
The difference in TA* using PO versus NO is really small, since the bias profiles in PO4 and NO3 are very similar for most models and the regression coefficient for the PO (NO) part is very small. Most 'weight' in the multilinear regression is on the salinity.

8) L. 139, you explicitly mention the fact that TA* “is computed as residual after rearranging Eq. (2)”. This is questioning, since it is finally on the dissolution of CaCO3 that we have the more certainty over its affects on alkalinity (+2 eq for 1 eq of CaCO3 dissolved). On the contrary, the effect of remineralisation, for example, is more complicated to extract from ESMs because of the N-reactions whether or not they are taken into account in the biogeochemical models. It is therefore perhaps a pity not to take advantage of the ease with which the carbonate pump can be used in terms of its effect on alkalinity with this method.

Yes, as the reviewer points out, there are uncertainties related to N-reactions, i.e., of reaction of PON to ammonium to nitrate. However, on the one hand, this is a fast reaction and on the other hand most models don’t simulate ammonium explicitly. We thus think that the uncertainty that goes with neglecting the effect from N-reactions is small at present. Koeve et al. (2014) states that the “global mean ratio of the alkalinity effect stemming from organic matter remineralization to TA background is only 0.005”. Contributions from N2-fixation, denitrification, sulfate reduction, and shelf alkalinity fluxes are even smaller or of local importance only. Koeve et al. (2014) also tested the a posteriori TA* method used here against explicitly modeled tracer of TA0, TA' and TA* with good results on a global scale.
Could you explain why you have chosen these values rather than the mean surface ESM values for instance?

We chose to apply the same typical values for all models. We now did a test and applied the actual MMM surface salinity (34.5), temperature (18.4) and PO (0.5) values for the calculation and found that the differences in the results are very, very small.

<table>
<thead>
<tr>
<th>Model</th>
<th>% difference in pCO₂ after alkalinity addition using S=34, T=15, PO=2</th>
<th>% difference in pCO₂ after alkalinity addition using S=34.5, T=18.4, PO=0.5</th>
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</thead>
<tbody>
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<td>MRI-ESM2-0'</td>
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<td>-0.99</td>
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<td>IPSL-CM6A-LR'</td>
<td>7.70</td>
<td>7.57</td>
</tr>
<tr>
<td>CanESM5'</td>
<td>6.90</td>
<td>6.83</td>
</tr>
<tr>
<td>MPI-ESM1-2-HR'</td>
<td>10.63</td>
<td>10.44</td>
</tr>
<tr>
<td>MPI-ESM1-2-LR'</td>
<td>12.98</td>
<td>12.76</td>
</tr>
</tbody>
</table>

This sentence should be part of the methodology.
Sentence was moved to methods part.

The very high TAn value observed at the ocean surface for GFDL-ESM4 is linked to very low salinity values (even possibly null) in the Baltic Sea if I remember well. You might be able to neglect this closed sea for this ESM when averaging to avoid this issue. You could also use a salinity threshold maybe.
Thank you for the suggestion, this issue has been fixed by using a salinity threshold of 10, which we noted in the methods section.

I would combine both figures so that the first panel is not repeated.
We would like to keep the comparison of TA and TAn separate from the comparison of global to regional profiles.

It could be mentioned that the bias observed for CNRM, IPSL and UK is associated with the ocean model NEMO.
Most models reproduce this pattern, while the CNRM, IPSL and UK ESMs simulate a strong increase of TA below about 2,000 m depth (Figure 4b). Those three ESMs have a NEMO ocean model in common.

TA biases likely lead DIC biases, as DIC can adjust through gas-exchange of CO2 to maintain a surface chemical equilibrium with the atmospheric CO2 concentration.
The suggested sentence has been added.
Besides, this aspect could be slightly more highlighted to understand why you decompose the alkalinity biases and not the DIC biases in addition, whereas you consider both surface alkalinity and DIC in the following with CO2SYS.
We focused on TA because of its conservative nature and because we came from the background of alkalinity enhancement studies. One result then became that DIC biases can act in a compensating way.
The decomposition of the TA biases (Figure 6a) shows that in the upper 1 km most of the models’ alkalinity biases are due to their preformed TA (Figure 6b). Per definition, models with a negative surface TA bias have a negative bias in preformed TA. Below about 1,000 m depth TA stays constant with depth. TA biases from the representation of organic matter remineralization processes are in the order of 5 to 10 mmol m\(^{-3}\) and play only a negligible role in absolute terms (Figure 6c). The bias in TA from calcium carbonate dissolution in the interior ocean (Figure 6d) can be in absolute terms comparable to or even larger than the bias in preformed TA. The MRI model and the GFDL models have a small negative bias in TA* in the order of ~10-20 mmol m\(^{-3}\), relatively constant with depth. The MPI and NorESM models have a slight positive TA* bias in about the same order of magnitude, also relatively constant with depth, while the UKESM, the CNRM-ESM and the IPSL ESM exhibit TA* biases that increase with depth. The CNRM model has the largest TA* bias with about 100 mmol m\(^{-3}\) at 4,000 m depth. CNRM-ESM2-1 and IPSL-CM6A-LR, have in common that they contain the same ocean model (NEMO) and the same biogeochemical model (PISCESv2). Dissolution in PISCESv2 is treated explicitly and is dependent on the calcite saturation state and the sinking speed for PIC is depth-dependent, while for other models the sinking speed is constant. In two of the models (of Figure 6d), MRI-ESM2-0 and UKESM1-0-LL, CaCO\(_3\) is dissolved without a sediment, while the other models do have explicit sediment treatments where CaCO\(_3\) is buried or dissolved, either depend on the calcite saturation state or a set rate (Planchat et al. 2023). A direct link of the treatment of CaCO\(_3\) at the bottom to the bias at depths is not obvious in this case.

The first sentences are too repetitive compared to what was already described in the previous sections.

Removed the repetitive sentence.

I. 226-228: This sentence is out of topic as you are talking about biases in this section. It would be great also to share some values.

The sentence has been removed and some values are added, see new paragraph above.

16) Fig. 6: The addition of MMM would be meaningful on these plots.

The MMM has been added to Figure 6.
17) l. 237: I am not convinced of the veracity of the causal sequence (cf. “thus”). Could you detail it?
Biases in simulated surface TA and surface DIC have implications for the individual models’ efficiency of OAE in terms of Revelle factor and change in pCO$_2$ and thus in the marine CO$_2$ uptake capacity. This is now shown in the newly added µCO$_2$ metric in Figure 7.

18) l. 240-242: “All panels are sorted by Revelle factor in ascending order.” This is enough in the legend. Sentence was removed.

19) “The Revelle factor ... at the ocean surface.” should be part of the Introduction. Removed here.

20) Fig. 7: Could you set “MMM” in bold so that we can easily spot it? Changed the figure to also have vertical MMM bars. Please see above.

21) I also suggest to keep the x-axis increasing in the last panel, to avoid confusions. Okay, this has been changed.

22) l. 260-261: Too repetitive with what was already said for Fig. 1.

23) l. 262: It is less than 13.0 % in Fig. 8, I think: t should be about 10.7 %.
Figure 8 has now been removed following the suggestion by Reviewer 1. In the initial figure though, the model with largest deviation, MPI-ESM-LR, of ~13% was missing. Thanks for pointing this out.
All values are listed in Table S2. The maximum value is 12.98%.

24) It would be great to have also the TA bias in Fig. 7 with the ESMs ordered in the same way. Thank you for the suggestion. TA has been added to Figure 7, which helps to illustrate our findings.

25) Could you set “MMM” in bold so that we can easily spot it? Could you adapt the y-axis scale to avoid the blank space to the top? Figure 8 has been removed in response to a comment by reviewer 1.

26) Discussion and conclusions: In overall, this section could be improved in terms of organization and content. There are some repetitions and it is sometimes difficult to follow you. In particular, l. 310-323 are quite messy for instance and the points overlap each other sometimes. We sorted the suggestions for improvement into points that relate to tuning and points that relate to implementing or expanding parametrizations.

In the first point, are you pointing towards including aragonite as well as calcite?

The first point was a suggestion to tune the calcification rate. What would be the effect of reducing the calcification rate in the Southern Ocean, where deep waters are upwelled?

Biases related to primary production and the calcification rate would be distributed, relatively speaking, equally at the ocean surface and in the upwelling regions of the Southern Ocean there is very little calcification anyway.

For almost all ESMs the TA bias is the same sign in the SO as in most of the global ocean (see Figure 2) so that tuning the calcification rate would have the same directional impact. Hopefully, this answers the question, otherwise we are not quite sure what specifically the reviewer is asking.

27) l. 320: are you talking about calcite dissolution (effectively sometimes explicitly modelled) or calcite production (always implicitly modelled in that case)? The carbon cycle formulation could be expanded to also simulate aragonite, i.e. including both aragonite formation and dissolution. Indeed, it is more important to separate aragonite and calcite for dissolution than for CaCO$_3$ formation.

28) Another important point, which is not mentioned, is the burial and dissolution at the seafloor.

We added a bullet point to the list:

- The representation of CaCO$_3$ treatment at the bottom-sediment interface (dissolution, sedimentation, sediment weathering) is important for the total alkalinity budget and also for upper ocean alkalinity especially in more shallow regions where alkalinity-enriched waters (through dissolution) can recirculate to the upper ocean more quickly (Gehlen et al., 2008).

29) Fig. S2: It is not compared to the GLODAP climatology here, as opposed to Fig. 6.

Figure S2 shows the results from the TA* method for the models and GLODAP side-by-side. GLODAP is the thick black line.

30) Table S1: Not useful, I think.

We would like to keep it, just in case.

31) There might be some issues in the References. There is at least one for Planchat et al., 2023, since some authors are missing.

Corrected the reference.

Technical corrections
All technical corrections have been addressed in the revised manuscript. Thanks again for your thorough review!

The following remarks are valid for the whole manuscript, and only a few occurrences will be mentioned below:

- There is a regular lack of punctuation, mainly commas.
- Beware of citations: a number of references appear with double brackets in the text.
In intercomparison studies of ESMs, it may be more accurate and preferable to refer to ESMs throughout the study rather than models.

If you decide not to put a space before '%', do so everywhere.

I suggest to write 'TA' method' instead of 'TA Method'.

The p in pCO2 should be in italic.

When the abbreviation/acronym has been defined, it can then be used directly (especially in the legend of the figures).

Try to keep the text independent of the figures and mention the figures in parentheses instead of within the text directly.

Write "Revelle factor" instead of "Revelle Factor"

Always write "TA-to-DIC-ratio" instead of mixing the way you mention it.

I. 4: 1 should in exponent

I. 9-15: commas are missing

I. 10: … on alkalinity …

I. 10: That is why, in the search for …

I. 11: … Ocean Alkalinity Enhancement (OAE) …

I. 12: … exists on how …

I. 13: … 14 CMIP6 Earth system models (ESMs) …

I. 17: … shows …

I. 36: … overestimate it in the …

I. 40-42: commas are missing

I. 45: … species composing the so-called …

I. 45-47: Suggestion: The oceanic uptake of anthropogenic carbon leads to an increase in aqueous CO2 and thus DIC. By changing the chemical equilibria between the carbonate species, this results in ocean acidification with a decrease in pH.

I. 48: … over acids (proton donors) …

I. 48: … role in the partitioning of the DIC pool, especially in the form …

I. 49-56: Very nice paragraph, but the first sentence could potentially be improved. Suggestion: The carbonate system is key in driving the seawater ability to resist a change in its chemistry, also called its buffering capacity. In particular, the Revelle factor is the sensitivity of pCO2 to changes in DIC. A low Revelle factor indicating a high buffering capacity and vice versa.

I. 55: … the potential CO2 uptake …

I. 68: … much research has focused on …

I. 72: … (OAE; Köhler et al., 2013 …

I. 81: … set-ups (e.g., Ilyina et al., 2013; …; Burt et al., 2021).

I. 86-87: … underway, or in planning, seeking to apply … for OAE (e.g., Butenschön et al., 2021), highlighting the importance and urgency of a robust ESM evaluation.

I. 93: They report an improvement in …

I. 109: “the first available ensemble member” is confusing and not precise.

I. 111: (CDO; Schulzweida, 2022)

Eq. (1): Precise that S is the salinity.

I. 115-119: Suggestion: … against gridded observational products: (i) TA, DIC and pH from the GLODAPv2.2016b Mapped Climatology (GLODAP in the following; Lauvset et al., 2016); (ii) oxygen and nutrients from the World Ocean Atlas 2018 dataset (WOA; Garcia H.E., 2019) and GLODAP; and (iii) salinity and temperature from the Polar science center Hydrographic Climatology (PHC3.0; Steele et al., 2001) and WOA. For the evaluation of global mean vertical profiles, the model data are interpolated onto the same 33 vertical levels used in the GLODAP climatology.

I. 125: … calcium carbonate (CaCO3) dissolution (TA*)

I. 128: potential temperature

I. 129: if NO is mentioned, then it should be defined to explicitly mention the difference with PO.

I. 131, 136: could you add references for the ratios r_1, r_2, … that you consider?

I. 131-132: “onto upper ocean TA values” and then “interior ocean” is a bit contradictory

I. 140: … to 10 of the 14 CMIP6 ESMs (… NorESM2-MM and UKESM1-0-LL)

I. 143: 2.3. Theoretical model sensitivity to OAE

I. 145: OAE. Thus, …

I. 147-148: Suggestion: this toolbox, from the combination of two of the carbonate system variables, compute the entire ocean CO2 system.

I. 149-152: Suggestion: … 1,026 kg/m3 (Fig. 1). First, we evaluate the CO2SYS output fields: Revelle
factor, pH and pCO2 (partial pressure of CO2 in seawater) based on the CMIP6 outputs. Then, we assess the changes...

1. 155 ... (MMM). Black vertical ... value. (b) Same ...
2. 156: potential temperature
3. 164: ... at the surface (Fig. 1a and 2)
4. 165: (Fig. 1a and 2)
5. 170: ... in NorESM-LM/MM and CanESM5; Fig. S1).

Fig. 2: Panel 1, title: GLODAP
Panel 2, title: GLODAP error; colorbar title: absolute error.
General colorbar title: Surface TA bias [mmol/m3]
Legend: Surface distribution of TA in GLODAP (top left), its error estimate (top center) and the CMIP6 multi-model-mean (MMM) bias (top right), as well as the respective biases of the ESMs.
Same comments for Fig. S1, where TAn should be directly mentioned.
Same comments for Fig. 5.
Try to homogenize the legends too
1. 177: ... the increase of TA with depth deeper ...
2. 179: ... TA overall. This indicates that their global inventory of TA is respectively too low and too high ...
3. 182: ... in the GFDL ESMs ...
4. 185-186 1,024; 1,028; 1,035
5. 187: ... at depth, or vice versa, ...
6. 188: ... in the upper ocean. The surface minima ...

Fig. 3: Could the black and grey lines have no transparency and be to the forefront so that we can clearly see both of them?
Legend: ... and TAn, (b) ...
1. 199: ... by the ESMs, referring to circulation biases.
2. 200: ... Ocean), between ... depth, ...
3. 208: ... and local distribution ...
4. 216: 3.2 Decomposition of the vertical alkalinity biases
5. 219: ... (CaCO3) dissolution and ...

Fig. 6: subplot titles: I suggest to simply write “TA/TA0/TAr/TA* bias”
1. 235: Impact of biases on OAE efficiency
2. 236: ... surface TA and DIC ...
3. 238: ... was conducted, using surface TA and DIC, to calculate the full carbonate system (see Methods and Fig. 1).
4. 248: ... GLODAP data, ranging from ...
5. 278-279: ... mmol/m3; i.e., YY %)
6. 294-295: In the sub-surface and the deep ocean, biases in TA are also driven by the CaCO3 dissolution, while contributions from remineralization of organic matter are negligible