Considerations for hypothetical carbon dioxide removal via alkalinity addition in the Amazon River watershed

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Abstract. The Amazon River plume plays a critical role in shaping the carbonate chemistry over a vast area in the western tropical North Atlantic. We <u>conduct a explore a thought experimentsensitivity analysis</u> of <u>hypothetical</u> ocean alkalinity enhancement (OAE) via <u>hypothetical</u> quicklime addition in the Amazon River watershed, examining the response of carbonate chemistry and air-sea carbon dioxide flux to the alkalinity addition. Through a series of sensitivity tests, we show that the detectability of the OAE-induced alkalinity increment depends on the perturbation strength (or size of the alkalinity addition, ΔTA) and the number of samples: there is a 90% chance to meet a minimum detectability requirement with $\Delta TA > 15 \,\mu$ mol kg⁻¹ and sample size > 40, given background variability of 15–30 μ mol kg⁻¹. OAE-induced *p*CO₂ reduction at the Amazon plume surface would range between 0–25 μ atm when $\Delta TA = 20 \,\mu$ mol kg⁻¹, decreasing with increasing salinity (S). Adding 20 μ mol kg⁻¹ of alkalinity at the river mouth could elevate the total carbon uptake in the Amazon River plume (15 < S < 35) by at least 0.07–0.1 MtCO₂ month⁻¹, and a-A major

- 15 portion of the uptake would occur in the saltiest region (S > 32) due to its large size, comprising approximately 80% of the S > 15 plume area. However, the lowest salinity region (S < 15) has a greater drop in surface ocean partial pressure of CO₂ (pCO₂^{sw}) due to its low buffer capacity, potentially allowing for observational detectability of pCO₂^{sw} reduction in this region. Reduced outgassing in this part of the plume, while more uncertain, may also be important for total additional CO₂ uptake. Such thought experimentssensitivity tests are useful in designing minimalistic field trials and setting achievable goals for monitoring, reporting,
- 20 and verification purposes.

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1 Introduction

To meet the Paris Agreement goal of limiting global temperature change to well below 2°C (UNFCCC, 2015), reducing greenhouse gas emissions is urgently needed, but insufficient on its own. Modeling results from the IPCC (2022) estimated a total remaining carbon budget of less than 500 GtCO₂ for a >50% probability of staying below 1.5°C warming by 2100. Staying within this small remaining budget is a formidable challenge, as it requires current emissions rates, which totaled ~36 GtCO₂ yr⁻¹ in 2021 (Friedlingstein et al., 2021), to rapidly approach zero. Accordingly, even very optimistic emission reduction scenarios assume carbon dioxide removal (CDR) will be needed to remove 10–20 GtCO₂ yr⁻¹ from the atmosphere by the end of the century (NASEM, 2019; 2021). With the ocean covering ~70% of the Earth's surface and providing the largest sink for anthropogenic CO₂ emissions to date, there is growing interest in CDR solutions in the marine environment.

30 Several ocean-based CDR approaches have been suggested over the past decades to reduce CO₂ in the atmosphere (NASEM, 2021). Enhanced Weathering (EW) and Ocean Alkalinity Enhancement (OAE) are related techniques with the primary goal to accelerate the carbonate weathering process that would naturally remove atmospheric CO₂ at a very slow rate (10,000 to 100,000 years; Gonzalez and Ilyina, 2016). EW techniques involve pulverization of carbonate-rich rocks and their application on land, while OAE applies those materials to increase alkalinity at the ocean surface (Kheshgi, 1995; Bach et al., 2019Renforth and Henderson, 2017).

35 When carbonate materials are spread near the edge of a coastal watershed, the weathering process may happen on land and/or in the ocean, and the distinction between the two techniques blurs.

EW and OAE can have other co-benefits besides reducing CO_2 and mitigating global warming. Several early EW experiments via regional-scale liming focused on mitigating acidification in inland lakes and ambient streams. These studies found that the surface water pH could be elevated to desired levels and persist for years without causing deleterious effects to the environment (e.g.,

- 40 Wright, 1985; Porcella, 1989; Driscoll et al., 1996). EW deployments in agricultural settings have also been proven effective in reducing soil acidity, preventing soil erosion, and enhancing crop yields (Caires et al., 2006; Köhler et al., 2010; Kantzas et al., 2022). Similarly, OAE can increase the pH of seawater, alleviating ocean acidification, which is a major stressor for the marine ecosystem (e.g., Doney et al., 2009). Both EW and OAE approaches are still in early phases of conceptualization and most OAE studies so far have focused on numerical simulations or efforts within laboratories (Köhler et al., 2010; Gonzalez and Ilyina, 2016;
- 45 Moras et al., 2022; Wang et al., 2022). Understanding the feasibility, effectiveness, and ecological risks of OAE is required before any large-scale efforts should be implemented (NASEM, 2021). Though harmful ecosystem effects associated with highly elevated alkalinity cannot be ruled out (Bach et al., 2019), this risk must be weighed against a counterfactual in which carbon dioxide remains in the atmosphere.

Large river-dominated tropical oceans are potential test ground for OAE, for several reasons. First, mixing and subduction of surface waters into the ocean interior is minimized in tropical oceans relative to higher latitudes (Gonzalez and Ilyina, 2016; Lenton et al., 2018). Second, large rivers form surface plumes that extend thousands of kilometers offshore (Lentz and Limeburner, 1995; Coles et al., 2013). In combination, these two factors mean that added alkalinity would have a long time to absorb CO₂ at the surface ocean and impact a vast area along the plume path. During this time, the atmospheric CO₂ is continuously sequestered by the ocean until a new air-sea CO₂ equilibrium is reached. The plumes also have a heightened potential for observational tracking using surface salinity, which can be estimated from satellite observations. Finally, the carbonate-poor river waters (relative to ocean water) may help suppress secondary chemical precipitation of the added alkalinity, a risk that can reduce the efficiency of

- OAE (Bach et al., 2019; Hartmann et al., 2022). Overall, global deployment of alkalinity in riversthe transport of alkalinity in rivers to the ocean from EW deployments is seen as <u>a</u> CDR technique with the potential to scale to the gigaton level (Zhang et al., 2022).
- 60 Therefore, we examine the Amazon River-ocean continuum for its potential as a site of OAE. As the world's largest river by volume, the Amazon River represents ~20% of the global riverine discharge into the oceans (Salati and Vose, 1984). Its massive outflow (an average of ~0.2 Sverdrup; Figure S1) creates a thin surface layer of low-salinity (Figure 1) and low-carbonate plume, extending up to 1.5 × 10⁶ km² at the ocean's surface (Molleri et al., 2010). As a result, the Amazon River plume has profound influence on the carbonate dynamics and atmospheric CO₂ sequestration throughout the western tropical North Atlantic Ocean (Ternon et al., 2000; Cooley et al., 2007; Lefèvre et al., 2010; Ibánez et al., 2015; Mu et al., 2021; Olivier et al., 2022; Monteiro et al., 2020; Cooley et al., 2007; Lefèvre et al., 2010; Ibánez et al., 2015; Mu et al., 2021; Olivier et al., 2022; Monteiro et al., 2020; Cooley et al., 2007; Lefèvre et al., 2010; Ibánez et al., 2015; Mu et al., 2021; Olivier et al., 2022; Monteiro et al., 2021; Olivier et al., 2022; Monteiro et al., 2020; Cooley et al., 2007; Lefèvre et al., 2010; Ibánez et al., 2015; Mu et al., 2021; Olivier et al., 2022; Monteiro et al., 2020; Cooley et al., 2007; Lefèvre et al., 2010; Ibánez et al., 2015; Mu et al., 2021; Olivier et al., 2022; Monteiro et al., 2021; Olivier et al., 2020; Cooley et al., 2007; Lefèvre et al., 2010; Ibánez et al., 2015; Mu et al., 2021; Olivier et al., 2022; Monteiro et al., 2020; Cooley et al., 2007; Lefèvre et al., 2010; Ibánez et al., 2015; Mu et al., 2021; Olivier et al., 2020; Cooley et al., 2020; Cooley et al., 2007; Lefèvre et al., 2010; Ibánez et al., 2015; Mu et al., 2021; Olivier et al., 2022; Monteiro et al., 2021; Olivier et al., 2020; Cooley et al.,

al., 2022).

In this study of hypothetical alkalinity addition at the Amazon River mouth, we investigate the expected changes in air-sea CO_2 flux at the offshore Amazon plume waters and examine what the perturbation size and sampling density would be needed to measure the alkalinity change and verify a resultant anomalous CO_2 flux. We make use of the knowledge from previous field

70 studies of the carbonate chemistry in the Amazon River plume, to address two main goals: 1) Analyze the potential for detectability of OAE-induced alkalinity change relative to measurement precision, background variability, and sample size; and 2) Estimate changes in ocean pCO_2 and air-sea CO_2 flux in the Amazon River plume due to hypothetical alkalinity addition at river mouth. Our effort aims to outline how one might consider the measurement, reporting and verification (MRV) needed in the context of

known background variability. We argue that this kind of "thought experiment" sensitivity analysis is a first step that could lead to more realistic numerical simulations if the system does not fail basic tests of feasibility.

2 Method

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2.1 Study Site and mixing model

We use a river-ocean conservative mixing model (Cooley and Yager, 2006) informed by direct observations collected as parts of the ANACONDAS and ROCA projects (Mu et al., 2021; Mu et al., in revision) to establish expectations of how alkalinity perturbations would influence the carbonate system in the Amazon plume (the gray and green pathways on the methods schematic shown in Figure 2). The mixing model assumes no sources or sinks of the carbonate species modeled, and the deviations from this assumption are discussed in Section 2.2. In this model, dissolved inorganic carbon (DIC) and total alkalinity (TA) are treated as conservative tracers and used to describe pCO_2 variations as a function of salinity and temperature (Figure 1). TA and DIC can be respectively expressed as:

85 $TA = [HCO_3^-] + 2[CO_3^{2-}] + minor constituents (1)$

 $TA = [Na^+] + 2[Mg^{2+}] + 2[Ca^{2+}] + \dots - [Cl^-] - 2[SO_4^{2-}] - [Br^-] - \dots$ (1)

 $DIC = [HCO_3^-] + [CO_3^{--}] + [CO_2^-]$ (2)

We adopt the explicitly conservative form of TA here (Wolf-Gladrow et al., 2007) so that TA changes due to the addition of our hypothetical calcium-based alkalinity feedstock can be fully tracked by the increase in calcium ion concentration. To establish the
baseline condition of the carbonate system in the Amazon River-ocean continuum, we adopted the river endmembers from Mu et al. (in revision; also, Table 1) and ocean endmembers (S = 36, TA = 2,369 µmol kg⁻¹, DIC = 2,025 µmol kg⁻¹) from Mu et al. (2021). The conservative mixing model can be further expressed as:

 $SSS_{mix} = S_r \times f_r + SSS_o \times f_o \qquad (3)$ $f_r + f_o = 1 \qquad (4)$

95 SSS_{mix} is the salinity for a plume sample; S_r is the salinity of the river endmember ($S_r = 0$); and SSS_o is the surface salinity of the ocean endmember. In Equation (4), f_r and f_o are the proportions of river and ocean in the sample, both of which can be solved when SSS_{mix} is known. The theoretical TA and DIC in the plume, TA_{mix} and DIC_{mix} at any given SSS_{mix} are then calculated from the known endmember properties and proportions via:

$$TA_{mix} = TA_r \times f_r + TA_o \times f_o \qquad (5)$$

100 $\text{DIC}_{\text{mix}} = \text{DIC}_{\text{r}} \times f_{r} + \text{DIC}_{\text{o}} \times f_{o}$ (6)

Eventually, pCO_2^{mix} and pH_{mix} at a given salinity and observed temperature are calculated with CO2SYS (Lewis et al., 1998) from inputs of TA_{mix} and DIC_{mix} . In this calculation we use the carbonic acid dissociation constants K1 and K2 from Mehrback (1973), refit by Dickson and Millero (1987) Millero (2010) due to their eligibility across a wide salinity range suitable for this study.

 We confined our study periods to two specific months, September 2011 and July 2012, during which measurements were made in
 both the river and throughout the Amazon River plume. <u>During the river plume/ocean expeditions, surface seawater pCO₂, temperature, salinity, and chlorophyll fluorescence were continuously measured shipboard across the Amazon-influenced regions; An oceanic station was specifically sampled for TA and DIC during each cruise as the ocean carbonate endmember for each
</u> <u>corresponding month (Mu et al., 2021).</u> Details in quantifying the river endmembers are described in Ward et al. (2015) and Mu et al. (in revision). Briefly, the Amazon River mouth region receives freshwater runoff from three channels: the north and south

- 110 channels near Macapá, and the channel of the Tocantins River near Belém (Figure S2). Sampling was conducted at the lower reaches of the three channels near the mouth (or "gateways") in each month. Six TA and DIC samples were collected at three cross-channel locations of each gateway (left bank, center, and right bank) from both surface and 50% depth (5–10 m)six river samples were collected at each gateway near the Amazon River mouth in each month and analyzed for TA and DIC. For a concise demonstration on the detectability of the alkalinity addition in the river, we chose to target use one single outer gateway (North
- 115 Macapá) as the perturbation site and consider how many samples would be needed to observe the TA perturbation relative to that gateway's background variability. However, to quantify the potential CDR induced by a TA perturbation just above detectability, we assumed the entire river mouth would be perturbed equally as in North Macapá. Because the outflow through both North and South Macapá gateways comprise > 80% of total discharge (Ward et al., 2015; Mu et al., in revision) and their chemical properties are almost identical seasonally (Mu et al., in revision), North Macapá is representative of the mouth's chemical composition. In an
- 120 actual OAE field trial, perturbing one gateway will perturb the mouth region by an amount proportional to the percentage discharge there (relative to the combined discharge), and the natural variability in the unperturbed gateways will affect the background variability at the mouth. There are key unknowns in seasonal TA contributions from other gateways to the mouth which prevents us from combining all the gateways. Therefore, the , North Macapá, to represent the Amazon River mouth in this study, while using the mean and standard deviation of the six TA values at North Macapá, respectively, were then used as the river TA endmember and their its natural background variability, respectively (Table 1). The monthly CO₂ fluxes in the plume for both
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chosen months are displayed in Table 2.

2.2 Deviations from the conservative mixing model

The assumption that TA and DIC act as conservative tracers is not true in the presence of photosynthesis, respiration, <u>bio</u>calcification, <u>abiotic dissolution/precipitation of calcium carbonate</u>, and/or air-sea gas exchange. In fact, plume pCO_2 values measured from an underway system on the research vessels surveying for the ANACONDAS program were systematically lower than predicted from the mixing model (Mu et al., 2021; Figure S3). This offset was assumed to be due to net community production, an assumption supported by the strong correlation between shipboard chlorophyll and the difference between the pCO_2^{mix} calculated from conservative mixing and the measured pCO_2 (Mu et al., 2021).

A linear regression between measured SSS (15 < SSS < 35) and measured pCO_2 showed a strong linear relationship (r = 0.79 for 135 September 2011 and r = 0.97 for July 2012). Therefore, we use SSS to calculate an empirical estimate of pCO_2 ($pCO_2^{empirical}$) at every SSS level in each month (orange pathway in the schematic shown as Figure 2). SSS fields in the Amazon plume were derived from remotely sensed diffuse attenuation coefficient at 490 nm (K_d490, Figure 1). The measured-SSS vs $pCO_2^{empirical}$ regression is used to map $pCO_2^{empirical}$ at every satellite-derived SSS value in the plume (as in Figure 3a) and to calculate the empirical air-sea CO_2 flux, according to the following equation:

140 $Flux_{empirical} = k\alpha (pCO_2^{empirical} - pCO_2^{atm})$ (7)

Where k is the gas exchange coefficient calculated from gridded reanalysis wind speed and the parameterization of Sweeney et al. (2007), α is the solubility of CO₂ in seawater <u>(a function of water temperature and salinity; calculated from equations in Weiss</u>, <u>1974</u>), and *p*CO₂^{atm} is the monthly averaged *p*CO₂ in the atmosphere measured at a nearby NOAA monitoring station. Details in calculating the empirical CO₂ flux can be found in Mu et al. (2021).

145 2.3 Experimental Design

The core of our thought experiment this sensitivity analysis is to manipulate the concentrations of TA at the river mouth by the hypothetical addition of pulverized quicklime (CaO)<u>a hypothetical alkalinity feedstock. There are several feedstocks that could be</u> considered for this purpose, each with a distinct set of considerations (Hartmann et al., 2013; Renforth and Henderson, 2017). Magnesium or silicate-rich rocks such as olivine and other basalts (Hartmann et al., 2013), carbonate minerals (e.g., Rau, 2011;

- 150 Renforth et al., 2022), mineral derivatives found in some industrial waste products, or even electrochemical production of alkalinity (Tyka et al., 2022; He and Tyka, 2023) are all candidates (NASEM, 2021). The stoichiometry of the reaction, dissolution rate, risk of secondary precipitation, and contamination with metals are all specific to the particular alkalinity source chosen. Quicklime (CaO) has been identified as being promising, due to its high solubility in sea water and rapid dissolution rates (Kheshgi, 1995; Bach et al., 2022). Quicklime is made by heating limestone (primarily CaCO₃), an abundant rock at the Earth's surface. The heating
- 155 of each molecule of CaCO₃ releases one molecule of quicklime and one molecule of CO₂ (Kheshgi, 1995). Therefore, carbon capture and storage of the released CO₂ would be necessary for OAE to be effective from the application of quicklime (Paquay and Zeebe, 2013; Renforth et al., 2013; NASEM, 2021; Moras et al., 2022). Hereafter, we assume that the alkalinity feedstock for the hypothetical OAE deployment is quicklime, which simplifies the discussion of stoichiometry, precludes contamination from metals and other industrial contaminants, justifies our assumption of rapid dissolution, and informs our discussion of potential
- 160 <u>unintended consequences.</u>,

assuming <u>Assuming</u> the additional CaO converts 100% to TA according to stoichiometry. T, the dissolution of CaO consumes CO_2 and releases HCO_3^- through:

 $(CaO + H_2O \rightleftharpoons Ca(OH)_2) + 2 CO_2 \rightleftharpoons Ca^{2+} + 2 HCO_3^-$ (8)

- For every mole of additional CaO, TA is increased by 2 moles (Equation 1) while DIC remains constant (i.e., consuming 2 moles
 of CO₂ while producing 2 moles of bicarbonate in Equation 8) before any significant air-sea equilibration occurs. This assumption of a constant DIC during perturbation should hold over the time scale of the river-ocean mixing based on the following evidence:

 Because-CaO dissolution happens on hour-scales (Moras et al., 2022); 2) The time elapsed between the river outflow at the mouth and its mixing with ocean to relatively high salinity (SSS > 15) is as short as two weeks (c.f., Figure 5 from Coles et al., 2013), while the air-sea CO₂ equilibration usually takes several weeks to months in the western tropical North Atlantic (Jones et al., 2014), the assumption of a constant DIC perturbation should hold over the time scale of dissolution. We assume in this study that there is no secondary precipitation of CaCO₃, and that the additional CaO stays at the surface layer until fully dissolved. Another underlying assumption is that CaO is homogeneously mixed across the river mouth gateway, eliminating potential
 - <u>uncertainty / variability due to patchiness of the TA addition.</u> We will address key unknowns associated with these assumptions in the \underline{D} -discussion.
- 175 We explore TA perturbations over the range 1–100 μ mol kg⁻¹ (i.e., perturbation between 0.07 and 7 times the natural variability for September 2011 and 0.04 and 4 times the natural variability for July 2012). The mixing model is recalculated with each of the perturbed river endmembers to find the perturbed TA_{mix} and DIC_{mix} across the plume. Then we use CO2SYS to calculate the theoretical $pCO_2^{mix}OAE$ at each salinity stamp with the perturbed TA_{mix}, DIC_{mix} and observed temperature as inputs to compare to the unperturbed baseline. The difference of the TA-enhanced seawater pCO_2 and the baseline pCO_2 :

$$180 \quad \Delta_{OAE} p CO_2^{mix} = p CO_2^{mix_OAE} - p CO_2^{mix_base}$$
(9)

In each perturbation scenario, the added alkalinity lowers the $pCO_2^{mix}OAE$ below the unperturbed mixing curve by the amount of $\Delta_{OAE}pCO_2^{mix}$. To finally arrive at the surface ocean pCO_2 needed to calculate the air-sea fluxes according to Equation 7, we add the $\Delta_{OAE}pCO_2^{mix}$ to $pCO_2^{empirical}$:

$$pCO_2^{OAE} = \Delta_{OAE} pCO_2^{mix} + pCO_2^{empirical}$$
(10)

185 This approach (schematized in Figure 2) implicitly assumes that the biological processes that lower the measured $pCO_2^{empirical}$ below the theoretical pCO_2^{mix} would be unchanged from the deployment of additional alkalinity, an assumption that would demand careful testing in the field.

2.4 Sensitivity analysis evaluation for detectability of hypothetical OAE deployments

To assess the TA detectability, we evaluate perturbations in one river gateway (North Macapá). We use a simple bootstrapping
 technique (or Monte Carlo simulation) to assess the minimum TA perturbation that would be detectable with a given number of measurements, taking into consideration the background variability during a given season and reasonable measurement precision.

We randomly generate 10,000 baseline TA values under a normal distribution using the mean and standard deviation of the six measured TA values in September 2011 or July 2012. We then generate 10,000 perturbed TA values for each perturbation scenario under a normal distribution, where the mean is the addition of the baseline mean and the perturbation strength, with the same standard deviation as the baseline. The range for the tested perturbation sizes is 1–20 μmol kg⁻¹ of TA. Within the 10,000 TA pools from both the baseline and a perturbation scenario, we randomly select 1–100 values (i.e., varying sample size) from both pools, perform the Student's t-test between the selected sets, and calculate the p-value to determine if the perturbed TA set would be seen as significantly different from baseline. The t-test is repeated 12,000 times for each perturbation size and sample size, generating 12,000 p-values. The mean of those p-values are calculated for each perturbation scenario at each sample size (Figure 4). A lower p-value suggests higher likelihood that the perturbed TA values would be considered significantly different from the baseline, and therefore "detectable". We use a threshold of p < 0.1 to indicate statistical significance.

3 Results

3.1 Baseline carbon chemistry in the plume

The surface $pCO_2^{empirical}$ in the Amazon River plume derived from remotely sensed SSS generally ranged from 210–400 μ atm in 205 September 2011 (Figure 3a), with lowest values in low-salinity regions near French Guiana coast and increasing as the river mixes with ocean water (Figure 1a). Figure 3a masks out regions with surface salinity below 15 and beyond 35 psu, as these waters are out of the range of a sufficiently robust algorithm between K_d490 and SSS (Mu et al., 2021).

The distribution of plume surface pCO_2^{sw} is primarily shaped by two processes: the Amazon River-ocean mixing and biological CO_2 consumption/release in the plume waters (Mu et al., 2021). For example, zero-salinity river water near the mouth is

supersaturated with respect to atmospheric CO₂ in July 2012, where $pCO_2 > 1,000 \mu atm$ is observed (Mu et al., 2021; Mu et al., in revision) due to low carbonate buffer capacity and high microbial respiration. As the river waters mix with the ocean, the sharp increase in buffer capacity and shift towards a net autotrophic state lower the surface pCO_2 towards a minimum (below pCO_2^{atm} in July 2012; Mu et al., 2021) before rising towards the open ocean levels closer to equilibrium with the atmosphere (i.e., $pCO_2 \sim 400 \mu atm$) at higher salinity (Figure S3). The nitrogen-fixing diatom-diazotroph assemblages and other phytoplankton that are active

in the Amazon plume waters (Goes et al., 2014) further enhance the CO_2 undersaturation in mid- to mid-high salinity portions of the plume (i.e., 15 < SSS < 33) on top of the undersaturated CO_2 state caused by conservative mixing (Mu et al., 2021).

3.2 Detectability of TA perturbations

We propose that the minimum requirement for MRV in a watershed OAE experiment is that the perturbed TA in the river can be detected above background variability. To illustrate the challenge of detectability, we consider the background TA variability in

- 220 the North Macapá gateway in September 2011, with the assumption that the variability measured at that time (standard deviation 14.5 μ mol kg⁻¹) is representative of that season and gateway. It is intuitive that a large enough perturbation (commensurate with the standard deviation) is detectable with a small number of samples (Figure 4), and a small perturbation (a factor of 3 smaller than the standard deviation or smaller) cannot be detected against the background variability regardless of the number of samples. This exercise reveals the challenge of balancing the effort needed to detect the perturbation against the obstacle and risk of increasing
- the TA by a large amount.

As expected and illustrated in Figure 4, higher sample sizes and TA perturbations lead to greater detectability of the alkalinity enhancement. For practical purposes, neither sample size nor perturbation strength can be increased infinitely, so OAE experiments would likely seek a balance between the two. For example, in September 2011 (Figure 4a, c), a TA perturbation of ~7–10 μ mol kg⁻¹ at a background standard deviation of 14.5 μ mol kg⁻¹ would have been readily detectable with 40 samples. For July 2012

230 (Figure 4b, d), when background variability was higher (25.7 μ mol kg⁻¹), 40 samples would detect perturbations only if they were to exceed +20 μ mol kg⁻¹ of TA. In an actual field OAE trial, each Each gateway and season (and gateway) may have different background variability that would require measurement and characterization in advance of any perturbation to have a clear strategy for sampling.

3.3 Impact on pCO₂ and air-sea CO₂ flux

Figure 3b shows the pCO_2^{mix} decrease due to a hypothetical TA addition in the Amazon watershed, which is highest in the lowsalinity waters near the river mouth and attenuates at higher salinities due to mixing with the unperturbed ocean water. Figure 5 provides a quantification of the air-sea CO₂ flux at each salinity level for different TA perturbation strengths (20, 50 and 100 μ mol kg⁻¹). While the air-sea CO₂ flux per unit area (or flux density) is greatest at low salinities (Figure 5a, b), the large area occupied by the diluted plume (salinities greater than 2532; Figure S4) means that more than half of the total integrated air-sea CO₂ exchange - as well as the perturbation to this flux – would occur in this salty part of the plume.

Air-sea CO_2 exchange at SSS = 34-35 for July 2012 is only slightly greater than 0 (Figure 5b), but due to the large area of plume at near-oceanic salinity level, the total plume CO_2 uptake (i.e., negative flux) in lower S regions is entirely offset by the CO_2 outgassing at 34 < SSS < 35. Regardless of whether the baseline Amazon plume is a carbon sink or source, the change in air-sea CO_2 exchange due to OAE shifts toward more carbon storage by the ocean and increases linearly with the size of TA perturbation (Table 2).

4 Discussion

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4.1 Unaccounted for plume region with great CDR potential

While we explored the OAE-induced additionality of the carbon uptake (or net CDR) at different salinities in the Amazon River plume (Figure 5), it is important to note that the cumulative additional uptake in Figure 5b excludes the freshest part of the plume

- 250 (SSS < 15). Due to high organic carbon remineralization in shallow waters (Mu et al., 2021), the quasi-linear SSS versus pCO_2 empirical relationship collapses at SSS < 15 and prevents the establishment of empirical pCO_2 in this region. Therefore, we excluded SSS < 15 in our analyses (e.g., in Figure 3, coastal areas < 50 m deep near the mouth are masked) while acknowledging that because of the strong CO₂ outgassing in this region, the baseline air-sea CO₂ flux in the entire plume (0 < SSS < 35) will be shifted towards more CO₂ release, should the SSS < 15 water be included. However, once TA is added, surface pCO_2 will decline
- 255 in the entire plume regardless of salinity, causing greater net CO_2 drawdown across the plume. In other words, the OAE-induced CO_2 uptake increase calculated for 15 < S < 35 plume water could substantially underestimate the true additionality when SSS < 15 is not considered. The area of the S < 15 water is small compared to the 15 < SSS < 35 plume (i.e., < 15% of the SSS > 15 plume area), but its lower buffer capacity also means pCO_2 is much more sensitive to TA addition, and therefore could still contribute a considerable amount to CDR.
- 260 <u>A basic scaling would suggest that 10% of the plume area with an OAE-induced surface pCO_2 decrease of 30 µatm would have as large an impact on area-integrated ocean uptake as the 15 < SSS < 35 portion of the plume with a 3 µatm decrease. Therefore, including the freshest part of the plume and calculating its reduction in outgassing could easily double our estimate of additional CO_2 uptake for the entire plume. Lacking both knowledge of the spatial distribution of salinity and a robust estimate for pCO_2 in this low salinity region prevents us from quantifying its CDR potential in a rigorously manner.</u>
- 265 <u>However, we can still conduct a back of the envelope calculation to estimate the distribution of CDR between SSS < 15 and 15 < SSS < 35 regions of the plume. With an addition of 20 μ mol kg⁻¹ TA and an averaged Amazon freshwater discharge of 0.18 Sverdrup (Figure S1), we can calculate the total transport of this TA addition of 9.3 × 10⁹ mol month⁻¹, given sufficient time for air-sea re-equilibration and the subsequent DIC increase due to OAE. If we assume the maximum DIC increase is 0.8 times the Δ TA (Wang et al., 2022), the maximum CDR resulting from TA addition would be $0.8 \times 9.3 \times 10^9$ mol month⁻¹ = 7.5 × 10⁹ mol</u>
- 270 month⁻¹, or 0.3 MtCO₂ month⁻¹. Given the estimated enhanced CO₂ flux in the S > 15 region is approximately 0.07~0.1 MtCO₂ month⁻¹ (Table 2), ~0.2 Mt of CO₂ uptake per month would be expected to take place in the low salinity region (SSS < 15), which is consistent with the estimation based on direct air-sea CO₂ flux change. This calculation highlights the potentially important role of the freshest part of the plume in the net CDR by the entire plume, which could be disproportionately large relative to its small size.

275 4.2 Exploring the minimum TA detectability

Here we propose that the minimum MRV requirement for an OAE experiment in a river be that the TA increase can be detected in the fresh river endmember, and we showed what size perturbation and sampling density would meet that minimum for the Amazon River. We found that the minimum detectable TA addition at the Amazon River mouth ranges between 10–20 μ mol kg⁻¹ with a reasonable sample size of 40 during field campaigns before and during the perturbation (Figure 4c, d). We focus on

- 280 detectability results for TA, since that is the variable directly targeted by any OAE approach. However, we present similar detectability analyses for the increase in pH (Figure S5) and decrease in pCO_2 in the river mouth (Figure S6) in the Supplementary Materials, which shows that the perturbation to these variables from a 10–20 μ mol kg⁻¹ is also readily detectable with a reasonable number of samples. With commercially available autonomous sensors for pH and pCO_2 (Kapsenberg et al., 2017; Sabine et al., 2020), these might be attractive proxies for the added TA.
- 285 One may further hope that MRV efforts for OAE deployed at the river mouth would document reductions in plume pCO_2 . We rely on the observed pCO_2 , a mixing model, and satellite-derived SSS to deduce where one might expect pCO_2 reductions to be measurable relative to background concentrations. Sensors for measuring pCO_2^{sw} aboard ships and uncrewed surface vehicles like

Saildrone and Waveglider have a target accuracy of 2 μ atm (Sabine et al., 2020). Figure 3 shows that, if Amazon River water TA were continuously increased by 20 μ mol kg⁻¹, a broad region of the plume would see pCO_2^{sw} decreases measurable with current

- 290 sensor technology, if they could sustain regular sampling over the SSS < 30 region before, during, and after the OAE deployment. However, like with the detectability of the TA in the river mouth, the *p*CO₂ perturbation must be detectable given background variability, and does not depend only on sensor accuracy. Here, a measure of that background variability is the <u>root-mean-square</u>
 - error (RMS-errorE) between the $pCO_2^{\text{empirical}}$, calculated from the linear regression of the *in situ* observations of pCO_2 and SSS, and the actual observations of pCO_2 (pCO_2^{measured} , Figure 2). For September 2011, the RMS error of this regression is ~22 μ atm for 15 < SSS < 35 (data not shownFigure S7). Thus, we would expect that a perturbation approximately this size would be
- 295 for 15 < SSS < 35 (data not shownFigure S7). Thus, we would expect that a perturbation approximately this size would be detectable with sufficient samples at the river mouth (Figure S6). Figure 3 and S7 suggests that a region of the plume near the river mouth approximately 15 < SSS < 20 could meet this criterion due to 1) lower salinity and higher perturbed TA and 2) relatively lower RMS error in *p*CO₂^{empirical} and thus low background *p*CO₂ variability, given a TA perturbation of +20 µmol kg⁻⁺ in the river. This speculation is nicely verified in a minimum OAE detectability test using *p*CO₂ in the plume waters as the proxy (Figure S8), where the reduction in *p*CO₂ at ΔTA = +10 µmol kg⁻¹ is readily detectable with 10 samples for 15 < SSS < 20 waters (Figure S8c),
 - contrasting sharply with the poor detectability at higher salinity (30 < SSS < 35; Figure S8d).

It is encouraging that SSS can be estimated from satellite measurements, allowing for field campaigns – including uncrewed vehicles piloted from operators onshore – to find and sample in the plume where an OAE-induced pCO_2 reduction may be detected. Nevertheless, Figure 5 shows that nearly half the additional ocean CO₂ uptake due to the OAE experiment could happen in the part of the plume with 30 < SSS < 35, where detecting the perturbation in TA and plume surface pCO_2 is impossible with the accuracy

of existing sensors or the laboratory analysis of bottle measurements.

Finally, we consider the amount of limestone (CaCO₃) required to produce pulverized quicklime as the alkalinity source material that sustains a set TA perturbation size (Table 2). The limestone equivalents needed to create a detectable perturbation of $+20 \,\mu$ mol kg⁻¹ of TA is 0.4 Mt per month using river discharge numbers from September 2011, which is approximately equal to 0.04% of

- 310 US industrial limestone production in the year 2021 (USGS, <u>https://pubs.usgs.gov/periodicals/mcs2021/mcs2021-stone-crushed.pdf</u>). Using river discharge for July 2012, the requirement grows to 0.7 Mt limestone per month. In terms of OAE-induced additional CO₂ uptake, every 0.1 MtCO₂ month⁻¹ represents more than 30 times greater CDR than the CO₂ captured by one ORCA Direct Air Capture plant in a full year (<u>https://en.wikipedia.org/wiki/Orca (carbon capture plant</u>)), yet amounts to less than what Brazil emitted on an average 2 hour basis in 2021 (<u>https://www.icos-cp.eu/science-and-impact/global-carbon-budget/2021</u>). It is
- 315 worth noting that concerns have been raised over the feasibility of CaO liming considering the energy need and CO₂ release associated with the production of this alkalinity source (e.g., <u>Paquay and Zeebe, 2013</u>; Renforth et al., 2013; Voosen, 2022), but limestone calcination coupled with heat recovery and carbon capture & storage technologies is found to significantly reduce energy costs and CO₂ emission during CaO production, potentially making this liming approach more sustainable (Foteinis et al., 2022).

4.3 Constraints on OAE deployments

- 320 So far, we have focused most of our efforts on describing the minimum detectable OAE perturbation. It is also important to consider what might constrain the upper limit of TA addition (i.e., through liming), such as ecosystem disturbances and secondary precipitation of calcium carbonate. Ideally, OAE would not put the ambient marine ecosystems at risk. The minimum detectable perturbation described above is so-close to the background TA variability that we suspect it would have, and so has a greater likelihood for <u>negligible small</u> impact on the ecosystem relative to larger deployments. Heavy metals in some source alkaline
- 325 minerals are problematic (NASEM, 2021), a consideration avoided with the use of 100% pure CaO compound as the source of TA

addition. The highest perturbation we explored was +100 μ mol kg⁻¹ of TA added to the river endmember, which would lead to initial pH increase of 2.0 at the river mouth, a pCO₂^{sw} decrease of up to 100 μ atm (data not shown) in the Amazon plume, and a total enhanced atmospheric carbon uptake of ~0.35 MtCO₂ month⁻¹ across the plume (15 < SSS < 35, with considerable additional contribution likely in the freshest plume waters) during September 2011 (Table 2 and Figure 5c). While the This level of initial pH increase at the TA addition site may be be likely concerning for local marine communities, although the effect would be quickly

Another constraint on the upper limit of TA addition through ocean liming is secondary CaCO₃ precipitation and the resulting reduction of atmospheric CO_2 uptake. Moras et al. (2022) suggested that runaway $CaCO_3$ during OAE may be avoidable if the saturation state of aragonite (Ω_A) is kept below 5 for natural seawater. Assuming such a Ω_A threshold for barely avoiding CaCO₃

dispersed by the river, and is and diminish reduced with increasing distance from the river due to mixing with the ocean.

- 335 precipitation is also applicable in fresher Amazon plume and river water, we have calculated the corresponding maximum TA perturbation allowed to keep Ω_A below this threshold according to the method in Mucci (1983), with the results shown in Table 3. Because the Ω_{A} increases with salinity, TA perturbation that likely avoids secondary CaCO₃ precipitation becomes substantially smaller close to <u>SSS</u> = 35 (i.e., ~100 μ mol kg⁻¹, compared to ~260 μ mol kg⁻¹ at S = 0). Adding 20 μ mol kg⁻¹ of TA in September 2011 would only increase Ω_A by a maximum of 0.3 (Table 3), and Ω_A stays below 5 everywhere in the plume for TA additions of
- 340 100 μ mol kg⁻¹ or smaller. An important caveat on assuming a Ω_A threshold of 5 is that the Amazon River at the mouth contains heavy loads of mineral particles and will likely facilitate the formation of heterogeneous nucleation and precipitation for CaCO₃ (Renforth and Henderson, 2017; Moras et al., 2022), thereby lowering the maximum TA addition allowed. Further study is needed to ensure Ω_A is always kept below proper precipitation threshold in all regions of the river plume.
- Adding TA to the land surrounding the Amazon watershed has could have possible advantages over spreading over across the open 345 ocean, involving co-benefits to the terrestrial ecosystems and agriculture, such as increased crop yield in agricultural watersheds (Caires et al., 2006; Hartmann et al., 2013; Beerling et al., 2020; Kelland et al., 2020), reduced soil run-off (Taylor et al., 2016), and reduction of N_2O production (Beerling et al., 2018). Introducing the lime-TA on land might also slow its dissolution in the river and allow its reaction with CO₂ to form bicarbonate in pore water, helping reduce the risks associated with large perturbations to pH and TA, like secondary precipitation or phytoplankton community perturbations. Basaltic rock with slower dissolution rates, 350 but more favorable stoichiometry from CDR than carbonates (Beerling et al., 2018), might also provide an alternative TA feedstock for land-based deployment. However, deployment on land would likely increase the difficulty to quantify the effect of TA addition for MRV purposes.

4.4 Additional challenges

One of the underlying assumptions for this analysis is that CaO reacting with CO₂ takes place in a time frame faster than air-sea 355 re-equilibration to occur, and the layer of plume water dwells at the surface for a relatively long period of time. If the TA-enhanced surface plume is not exposed to the atmosphere long enough to allow for full air-sea CO₂ equilibration (e.g., due to water mass subduction), CO₂ removal efficiency will decrease in these regions (Jones et al., 2014). A logical next step to characterize the spatial footprint of the hypothetical experiment and the total time scale for equilibration would be to simulate the time-evolving patch of the alkalinity anomaly, the corresponding pCO_2 perturbation, and the total excess DIC in an ocean general circulation 360 model with the OAE experiment relative to a control simulation.

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We argue that the TA-enhanced waters do not need to remain continuously within the surface layer to induce ocean CDR. The change in the air-sea flux can be realized whenever the perturbed waters influence the surface, even if that is far from the plume or years after the deployment. Though the implied dilution of the signal or delay in causing additionality of the CDR may not

change the net impact of the OAE experiment if integrated over a sufficiently long time period, it would make MRV of the total impact essentially impossible through observational methods alone.

Another assumption in this work was that the photosynthesis, which is known to drive pCO_2 beneath the conservative mixing curve (Mu et al., 2021), would be unchanged by the alkalinity addition. This assumption is premised on the argument that perturbations barely detectable above background variability are unlikely to strongly change the ambient ecosystem. an argument supported by a number of recent studies. For example, Regardless, ecosystem impacts would require careful monitoring in any field experiment.

Recent <u>Regardless, ecosystem impacts would require careful monitoring in any field experiment.</u> results from a mesocosm experiment with Tasmanian coastal waters show statistically significant, though relatively moderate, changes to phytoplankton community structure and function at high TA perturbation values of +495 μmol kg⁻¹, which is 25 times higher than the minimum detectable perturbation explored here (Ferderer et al., 2022). Another microcosm incubation study reveals the alkalinity enhancement inducing minor but largely negligible short-term effects on primary productivity and no measurable changes in biological CaCO₃ precipitation, even at a TA perturbation level of +2000 μmol kg⁻¹ (Subhas et al., 2022). Regardless, ecosystem impacts would require careful monitoring in any field OAE experiment.

If the alkalinity source for OAE is quicklime, managing the heat released through its exothermic reaction during dissolution in water at a rate of 64 kJ mol⁻¹ CaO is another important consideration. Taking the highest perturbation scenario explored in Table 2 that increases TA in the Amazon watershed by $100 \,\mu$ mol kg⁻¹, we estimate 310 MW total heat released during the duration of the

380 dissolution. In order for the heat released from the CaO reaction to be lower than 10% of the daily solar radiation at the equator ~400 W m⁻² (a level that would presumably be swamped by natural variability in the river's heat budget), the TA injection would have to occur over an area of river larger than about 8 km², e.g., 2 km of river width and 4 km along its length. Therefore, it is possible to minimize the environmental risk due to heat released during quicklime dissolution by spreading it over a large enough area at the river mouth.

385 5 Conclusions

- We propose conduct a thought experiments ensitivity analysis of alkalinity enhancement in the Amazon River watershed, evaluating the detectability of added TA, and predicting its influence on the atmospheric CO₂ uptake in the offshore plume. Adding 20 μ mol kg⁻¹ of TA in a month at the Amazon watershed could increase the CO₂ uptake by the river plume by at least 0.07–0.1 MtCO₂ month⁻¹. The full CDR might be as much as three times as high as this estimate due to reduced outgassing in the freshest part of 390 the plume where we lack a robust empirical pCO_2^{sw} -salinity relationship and therefore cannot robustly quantify the anomalous airsea exchange in this region. We found a TA perturbation of $+10-20 \ \mu \text{mol kg}^{-1}$ in the river is readily detectable with at least 40 samples, given background variability of 15–30 μ mol kg⁻¹. By adding a TA perturbation barely detectable relative to the background TA variability, the likelihood of substantial environmental disturbance at the injection site would be minimized. However, observing the corresponding pCO_2 perturbation in the plume presents a greater observational challenge. Even with the 395 highly sensitive pCO_2 sensors available today (e.g., Sabine et al., 2020), detectability of the pCO_2 anomaly is plausible only in the freshest fresher part of the plume given dilution of the signal at higher salinities and large background variability. Finally, Quantifying the total additional CO₂ uptake from this type of OAE cannot rely on observations alone; idealized conceptual models like the one presented here and more sophisticated circulation-biogeochemical ocean models will always be required to understand the total CDR because the additional CO₂ uptake is likely to occur over long time periods and at very diluted perturbation levels. 400 It is also clear that the basic scientific feasibility we aimed to assess here does not reveal the full feasibility of such interventions:
 - ocean-based CDR research and deployment require care and attention to ethical, political and governance issues.

Data availability

Data for the air–sea CO_2 flux calculations in the Amazon River plume are from Mu et al. (2021). River mouth data are reported in a manuscript currently under revision in Global Biogeochemical Cycles that will be resubmitted in May 2023. All codes are

405 available at https://drive.google.com/drive/folders/1YYjcT5ZYvyHoJaKmRl3Fd8aagMhtRwCA?usp=share_link.

Author Contribution

JBP and LM conceptualized the framework of this experiment. LM, JBP, and HW designed the methodology. LM performed the analytical calculations and numerical simulations. LM wrote the manuscript with support from JBP and HW. JBP supervised the project.

410 **Competing Interests**

The authors declare that they have no conflict of interest.

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Tables

| Sample No. | Total Alkalinity (µmol kg ⁻¹) | | | | |
|------------|---|--------------|--|--|--|
| | September 2011 | July 2012 | | | |
| 1 | 279.8 | 362.3 | | | |
| 2 | 286.8 | 322.2 | | | |
| 3 | 310.8 | 301.2 | | | |
| 4 | 296.3 | 327.7 | | | |
| 5 | 307.5 | 296.9 | | | |
| 6 | 275.5 | 295.8 | | | |
| Mean (std) | 292.8 (14.5) | 317.7 (25.7) | | | |

590 Table 1. Baseline total alkalinity (μmol kg⁻¹) determined at the N. Macapa gateway at the Amazon River mouth during September 2011 and July 2012 (Mu et al., in revision). The means represent the river's alkalinity endmember in that month.

Table 2. Summary table for air-sea CO₂ flux in the Amazon River plume (15 < SSS < 35) and the mass of CaCO₃ mineral required for each TA perturbation size (Δ TA) in each month. Positive fluxes indicate CO₂ outgassing and negative fluxes (–) indicate ocean CO₂ uptake. Note that the post-perturbation CO₂ flux and CaCO₃ demand increase linearly with increase in

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 $\Delta TA.$

| | | Sej | ptember 2011 | | July 2012 | | | |
|--------------------------------|-----|--------------------------------------|------------------------------------|---------------------|--------------------------------------|------------------------------------|---------------------|--|
| | | CO ₂ flux | | CaCO ₃ | CO ₂ flux | | CaCO ₃ | |
| | | mmol m ⁻² d ⁻¹ | MtCO ₂ mo ⁻¹ | Mt mo ⁻¹ | mmol m ⁻² d ⁻¹ | MtCO ₂ mo ⁻¹ | Mt mo ⁻¹ | |
| ΔTA umol kg ⁻¹) | 0 | -0.43 | -0.47 | 0 | 0.26 | 0.24 | 0 | |
| | 10 | -0.46 | -0.50 | 0.2 | 0.20 | 0.19 | 0.3 | |
| | 20 | -0.49 | -0.54 | 0.4 | 0.15 | 0.14 | 0.7 | |
| | 50 | -0.60 | -0.65 | 1.1 | -0.02 | -0.01 | 1.7 | |
| 3 | 100 | -0.76 | -0.83 | 2.2 | -0.27 | -0.26 | 3.5 | |

Table 3. Theoretical aragonite saturation state (Ω_A) at the surface of the Amazon River-ocean continuum, calculated from the river-ocean mixing model (see Section 2.1 and Figure 2) for different TA perturbation strengths (Δ TA) at various salinities in September 2011 using constant water temperature of 29 °C, according to the equation $\Omega_A = [Ca^{2+}][CO_3^{2-}] / K_{sp}^{A(*)}$. [Ca²⁺] is calculated by adding the additional [Ca²⁺] due to TA perturbation (i.e., Δ TA/2) to the unperturbed [Ca²⁺]; the latter term is comprised of relative contributions from both the river and ocean [Ca²⁺] endmembers forat a specific salinity. [Ca²⁺] of 0.15 mmol kg⁻¹ is estimated used as the river endmember approximating results from Drake et al. (2021), and the global ocean average [Ca²⁺] of 10.28 mmol kg⁻¹ is used as the ocean endmember. [CO₃²⁻] is derived from theoretical TA and DIC in the river-ocean endmember mixing model; the temperature- and salinity- dependent stoichiometric solubility product for aragonite, $K_{sp}^{A(*)}$, is calculated from Mucci (1983). The following equations in Mucci (1983) are involved in the determination of $K_{sp}^{A(*)}$ for each salinity: $logK_{sp}^{A(*)} - logK_{sp}^{A(0)} = (b_0 + b_1SSTK + b_2/SSTK) S^{0.5} + c_0S + d_0S^{1.5}$ and $logK_{sp}^{A(0)} = -171.945 - 0.077993SSTK + 2903.293/SSTK + 71.595logSSTK; SSTK is the sea surface temperature in Kelvin (i.e., SSTK = 302.15 K); b, c, and d are constants determined reported in Mucci (1983).$

| ΔΤΑ | $\Omega_{\rm A}$ (at various salinities) for Sep-2011 | | | | | | | |
|-----------------------|---|-----|------|------|------|------|--|--|
| µmol kg ⁻¹ | S=0 | S=5 | S=10 | S=20 | S=30 | S=35 | | |
| 0 | 0.0 | 0.1 | 0.5 | 1.8 | 3.2 | 3.8 | | |
| 20 | 0.0 | 0.1 | 0.7 | 2.1 | 3.4 | 4.0 | | |
| 50 | 0.0 | 0.3 | 1.0 | 2.4 | 3.8 | 4.4 | | |
| 100 | 0.1 | 1.0 | 1.7 | 3.0 | 4.4 | 5.0 | | |
| 200 | 2.9 | 2.6 | 3.1 | 4.3 | 5.6 | 6.2 | | |
| 260 | 5.1 | 3.6 | 4.0 | 5.1 | 6.4 | 6.9 | | |



Figure 1. Sea surface salinity (SSS) for the Amazon River plume (15°N–3°S, 43–60°W) during (**a**) September 2011, and (**b**) July 2012, derived from remotely sensed diffuse attenuation coefficient at 490 nm. Area likely affected by the Orinoco River plume (< 150 km off the coastline between 55–60°W) is excluded from this study. Oceanic regions in light blue on the maps indicate the satellite-derived SSS data are unavailable, mostly due to muddy nearshore waters or clouds. See Mu et al. (2021) for further details.



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Figure 2. Schematic diagram for the method used to calculate carbonate chemistry and air–sea CO₂ fluxes before and after a hypothetical alkalinity addition (OAE) in the Amazon River-ocean continuum. The baseline and OAE pathways (gray and green) use the conservative mixing model (Equations 3-6) and CO2SYS to calculate the baseline and hypothetical OAE pCO_2 . The difference between the baseline and OAE mixed models ($\Delta_{OAE}pCO_2^{mix} = pCO_2^{mix_OAE} - pCO_2^{mix_base}$) is added to the empirically-derived pCO_2 at every SSS. The resulting pCO_2^{OAE} is used to calculate the air-sea CO₂ flux across the surface of the plume for each hypothetical OAE experiment.



Figure 3. *p*CO₂ at the surface of the Amazon River plume and its predicted response to alkalinity addition based on satellitederived SSS. (a) Spatial distribution of unperturbed $pCO_2^{empirical}$ (μ atm) at the surface of the Amazon River plume (15 < SSS < 35) during September 2011, with SSS outside of the plume range removed from this map as the remotely sensed K_d490 vs SSS regression is sufficiently robust only when 15 < SSS < 35. (See Figure 2 and Section 2.2 for details on how the mapped quantity is defined, and Mu et al. (2021) for additional details). (b). Predicted changes in the plume surface pCO_2 (i.e., $\Delta_{OAE}pCO_2^{mix}$ from Equation 9) due to addition of 20 μ mol kg⁻¹ of alkalinity at the Amazon River mouth during September 2011.



Figure 4. Detectability of hypothetical TA enhancement relative to background variability. (**a**)(**b**). The theoretical shift of TA $(\mu \text{mol } \text{kg}^{-1})$ means and distributions for September 2011 and July 2012 due to the addition of 20 μ mol kg⁻¹ of TA at the Amazon River mouth. Black lines indicate the *in situ* TA measurements at the mouth on which the baseline data distributions are based. The standard deviations in the perturbation scenarios are assumed the same as those from the baseline. (**c**)(**d**). The p-value maps from t-tests performed between the baseline and TA perturbation scenarios at various sample sizes and TA perturbation strengths for September 2011 and July 2012. Areas in yellow (p < 0.1) indicate conditions where an analyst would conclude that the TA perturbation was detected relative to the baseline condition with 90% certainty.



Figure 5. The impact of TA addition on ocean uptake of CO_2 across the Amazon River plume. (Top) Air-sea CO_2 flux density (mmol m⁻² d⁻¹) in the Amazon River plume at different sea surface salinities for the baseline and multiple TA perturbation scenarios in (a) September 2011 and (b) July 2012. Standard deviations of the fluxes within each salinity band (1 psu apart) are represented by the shaded areas. Negative values indicate atmospheric CO_2 sinks. (Bottom) Cumulative ocean CO_2 uptake (MtCO₂ mo⁻¹) in the Amazon plume due to TA additions in (c) September 2011 and (d) July 2012. SSS ranging between 15 and 35 is used while SSS < 15 and > 35 is omitted, because the remotely sensed K_d490 vs SSS regression is sufficiently robust only when 15 < SSS < 35 according to the Amazon River plume definition from Mu et al. (2021).



Figure S1. Monthly discharge of the Amazon River (Sv; $1 \text{ Sv} = 10^6 \text{ m}^3 \text{ s}^{-1}$) at the Óbidos gauge station (~650 km upriver from the mouth) in 2011 (red line), 2012 (blue line), and climatology averaged over 1968–2012 (black line), in reference to the mean annual discharge 1968–2012 (dotted line).



Figure S2. Map of the Amazon River lower reach. Geochemical surveys (in particular, river TA and DIC measurements) were conducted at the outer gateways (North Macapá, South Macapá, and Belém) during the September 2011 and July 2012 *River Ocean Continuum of the Amazon* (ROCA) project. This figure is reproduced from Ward et al. (2015) and Mu et al. (in revision).



Figure S3. Surface pCO_2 in the Amazon River plume at 15 < salinity < 35 and river-ocean mixing curve before and after the TA perturbation at the mouth for (a) September 2011 and (b) July 2012. TA perturbation strength is 20 µmol kg⁻¹. The TA and DIC for the baseline mixing curves (solid red and blue lines) are calculated from the river and ocean endmember values, from which $pCO_2^{mix_base}$ at each salinity stamp is calculated using the CO2SYS program. The dashed lines show the OAE-perturbed mixing curve at $\Delta TA = 20 \ \mu mol \ kg^{-1}$. Black triangle markers show the empirical $pCO_2 (pCO_2^{empirical})$ at the plume surface computed from satellite-derived salinity and measured $pCO_2 (pCO_2^{measured}; magenta dots)$ versus salinity regression, and open cyan circles represent the OAE-perturbed plume $pCO_2 (pCO_2^{OAE})$ at $\Delta TA = 20 \ \mu mol \ kg^{-1}$. Salinity as low as 15 was observed in the July 2012 survey, while no salinity under 25 was recorded during the September 2011 cruise.



to compute the percentages. This figure is reproduced from Mu et al. (2021).



Figure S5. Detectability of OAE-induced pH variation in the North Macapá gateway relative to its background variability. (a)(b). The theoretical shift of pH (μ mol kg⁻¹) means and distributions for September 2011 and July 2012 due to the addition of 20 μ mol kg⁻¹ of TA at the Amazon River mouth. Black lines indicate baseline pH calculated by baseline TA and DIC at the river mouth. The mean and standard deviations of pH in the perturbation scenarios (dash-dot lines) are calculated from perturbed TA and DIC values. (c)(d). The p-value plots from t-tests simulated between baseline and perturbed pH induced by TA perturbation scenarios at various sample sizes and TA perturbation strengths for September 2011 and July 2012. Areas in yellow (p < 0.1) indicate conditions where an analyst would conclude that the TA perturbation was detected relative to the baseline condition with 90% certainty.



Figure S6. Detectability of OAE-induced pCO_2 variation in the North Macapá gateway relative to its background variability. (a)(b). The theoretical shift of pCO_2 (µatm) means and distributions for September 2011 and July 2012 due to the addition of 20 µmol kg⁻¹ of TA at the Amazon River mouth. Black lines indicate the baseline pCO_2 calculated by baseline TA and DIC at the river mouth. The mean and standard deviations of pCO_2 in the perturbation scenarios (dash-dot lines) are calculated from perturbed TA and DIC values. (c)(d). The p-value plots from t-tests simulated between baseline and perturbed pCO_2 induced by TA perturbation scenarios at various sample sizes and TA perturbation strengths for September 2011 and July 2012. Areas in yellow (p < 0.1) indicate conditions where an analyst would conclude that the TA perturbation was detected relative to the baseline condition with 90% certainty.



Figure S7. The residual of empirical pCO_2 relative to the measured pCO_2 at the same salinity across the salinity spectrum of 15 $\leq SSS < 35$ for September 2011 (red circle) and July 2012 (blue circle). Note that $pCO_2^{\text{empirical}}$ _residual = $pCO_2^{\text{empirical}} - \frac{pCO_2^{\text{measured}}}{15} < SSS < 35$. $pCO_2^{\text{empirical}}$ is calculated from cruise-specific pCO_2^{measured} versus SSS regression at a satellitederived salinity (Mu et al., 2021).



Figure S8. Detectability of OAE-induced pCO₂ variation at different salinity ranges of the plume water. (a). The theoretical shift of plume pCO₂ (µatm) distributions at 15 < SSS < 20, as a result of a TA perturbation of 20 µmol kg⁻¹ at the river mouth. Black lines indicate the baseline pCO₂ calculated from the river and ocean carbonate endmembers at SSS = 17.5 (following the mixing curve shown in Figure S3b). The standard deviation of the baseline pCO₂ distribution is calculated from the root-mean-square error of pCO₂ residual values for 15 < SSS < 20, described in Figure S7. The mean of the pCO₂ in the perturbation scenarios (dash-dot lines) are calculated from perturbed TA and DIC values that follow the dash-dot line in Figure S3b. (a). Similar to (a) but for the salinity range of 30 < SSS < 35. (c)(d). The p-value plots from t-tests simulated between baseline and perturbed pCO₂ in the plume, induced by TA perturbation scenarios at various sample sizes and perturbation strengths for 15 < SSS < 20 and 30 < SSS < 35, respectively, in July 2012. Areas in yellow (p < 0.1) indicate condition with 90% certainty.

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