An assessment of ocean alkalinity enhancement using aqueous hydroxides: kinetics, efficiency, and precipitation thresholds

3

Mallory C. Ringham¹, Nathan Hirtle¹, Cody Shaw¹, Xi Lu¹, Julian Herndon^{2,3}, Brendan R. Carter^{2,3},
Matthew D. Eisaman^{4,5}

5 Matthew D. Eisaman ^{4,5}
6
7 ¹Stony Brook University, Stony Brook, NY, USA

8 ²Cooperative Institute for Climate Ocean and Ecosystem Studies, University of Washington, Seattle, USA

9 ³Pacific Marine Environmental Laboratory, National Oceanic and Atmospheric Administration, Seattle, WA, USA*

10 ⁴ Department of Earth & Planetary Sciences, Yale University, New Haven, CT, USA

⁵ Yale Center for Natural Carbon Capture, Yale University, New Haven, CT, USA

12

13 Correspondence to: Mallory Ringham (mallory.ringham@stonybrook.edu); Current address: Ebb Carbon Inc., San

14 Carlos, CA, USA

15 Abstract

16 Ocean alkalinity enhancement (OAE) is a promising approach to marine carbon dioxide removal (mCDR) that

17 leverages the large surface area and carbon storage capacity of the oceans to sequester atmospheric CO₂ as dissolved

18 bicarbonate (HCO₃⁻). One OAE method involves the conversion of salt in seawater into aqueous alkalinity (NaOH),

19 which is returned to the ocean. The resulting increase in seawater pH and alkalinity causes a shift in dissolved

inorganic carbon (DIC) speciation toward carbonate and a decrease in the surface-ocean pCO_2 . The shift in the pCO_2

21 results in enhanced uptake of atmospheric CO_2 uptake by the seawater due to gas exchange. In this study, we

systematically test the efficiency of CO_2 uptake in seawater treated with NaOH at aquaria (15L) and tank (6000L) cales to establish operational boundaries for safety and efficiency in advance of scaling up to field experiments.

scales to establish operational boundaries for safety and efficiency in advance of scaling up to field experiments.
 CO₂ equilibration occurred on order of weeks to months, depending on circulation, air forcing, and air bubbling

25 conditions within the test tanks. An increase of ~0.7-0.9 mol DIC/ mol added alkalinity (in the form of NaOH) was

26 observed through analysis of seawater bottle samples and pH sensor data, consistent with the value expected given

27 the values of the carbonate system equilibrium calculations for the range of salinities and temperatures tested.

- 28 Mineral precipitation occurred when the bulk seawater pH exceeded 10.0 and $\Omega_{aragonite}$ exceeded 30.0. This
- 29 precipitation was dominated by Mg(OH)₂ over hours to 1 day before shifting to CaCO_{3, aragonite} precipitation. These
- 30 data, combined with models of the dilution and advection of alkaline plumes, will allow for estimation of the
- 31 amount of carbon dioxide removal expected from OAE pilot studies. Future experiments should better approximate
- 32 field conditions including sediment interactions, biological activity, ocean circulation, air-sea gas exchange rates,
- **33** and mixing-zone dynamics.

34 Keywords

Ocean Alkalinity Enhancement (OAE); marine carbon dioxide removal (mCDR); ocean carbon dioxide removal
 (ocean CDR)

37

38 1 Introduction

39 The Sixth Assessment Report of the Intergovernmental Panel on Climate Change reported that in addition to a

 $40 \qquad \text{drastic decrease in CO}_2 \text{ emissions, active removal of 5-15 Gt of atmospheric CO}_2 \text{ per year by 2100 is necessary to}$

41 constrain average global warming to less than 1.5 - 2 °C (noting that the magnitude of carbon removals varies by

- 42 climate scenario: IPCC, 2022; Rogelj, 2018). A wide variety of negative emissions technologies (NETs) are under
- 43 development to meet this enormous challenge (Minx et al., 2018; NASEM, 2019; NASEM, 2021; Rueda et al.,
- 44 2021; Vitillo et al., 2022).
- 45 A suite of promising approaches to CO₂ removal termed ocean or marine carbon dioxide removal (ocean CDR or
- 46 mCDR, respectively) leverage the enormous surface area and carbon storage capacity of the ocean (Boettcher et al.,
- 47 2019; NASEM, 2021). Ocean alkalinity enhancement (OAE) is an mCDR method that aims to store atmospheric
- 48 CO_2 in a dissolved phase in the ocean as bicarbonate ions (HCO₃⁻), thereby accelerating a natural planetary CO_2
- 49 regulation mechanism, the carbonate-silicate cycle (Berner, 1983; Isson et al., 2020). OAE has the potential to scale 50 to gigatons of CO₂ removal per year (He and Tyka, 2023), but development of this approach requires careful
- 51 consideration of: the methods and materials used to source and process alkalinity; the form and method of delivery
- 52 of alkalinity to the surface ocean (for example, aqueous or solid phase); and selection of appropriate geographic sites
- 53 for alkalinity dispersal (Oschlies et al., 2023). OAE methods under exploration include: mining and crushing
- 54 alkaline minerals (e.g., olivine, basalts) to be spread via ship or in coastal environments (e.g., beach restoration, or
- 55 salt marsh distribution) (Feng et al., 2017; Köhler, Hartmann, and Wolf-Gladrow, 2010; Monserrat et al., 2018;
- 56 Rigopoulos et al., 2018); the mining or industrial production of $Mg(OH)_2$ or mining CaCO₃ and calcining it to CaO
- 57 or Ca(OH)₂, with the Mg(OH)₂ or Ca(OH)₂ spread via ship or coastal outfall pipe (Harvey, 2008; Ilyina et al., 2013;
- 58 Kheshgi, 1995; La Plante, 2023; Moras et al., 2022; Nduagu, 2012; Rau, 2008; Renforth and Henderson, 2017; 59 Shaw, 2022); and the electrochemical conversion of saltwater into aqueous hydroxides and dispersal via coastal
- 60
- outfalls (de Lannoy et al., 2018; Eisaman et al., 2018; Lu et al., 2022; Tyka, Van Arsdale, and Platt, 2022; Eisaman
- 61 et al., 2023).
- 62 Many of these approaches and technologies are at a nascent stage. We must move quickly to quantitatively test and

63 characterize their performance to determine which, if any, justify larger-scale deployment. The electrochemical

64 conversion of salt (NaCl) into aqueous alkalinity (NaOH) has many potential advantages in scaling considerations,

- 65 including simplified distribution of a liquid product to the ocean, avoidance of mining and the transportation of the
- 66 alkalinity source over long distances, and avoidance of potentially harmful impurities present in mined alkalinity
- 67 sources (NASEM, 2021; Caserini, Storni, and Grosso, 2022).

68 Total alkalinity (TA) is defined as the excess of proton acceptors over proton donors in an aqueous solution (Eq. 1),

- 69 where ellipses represent neglected acids and bases (Dickson 1981; Dickson 1992; Wolf-Gladrow et al., 2007). A
- 70 higher TA value for a seawater sample indicates that it has a higher buffering capacity than a sample with a lower
- 71 TA value. That is, for sample with a higher TA value, the addition of a given amount of acid to the sample will
- 72 decrease its pH less than for a sample with a lower TA value.

73
$$TA = [HCO_{3}^{-}] + 2 [CO_{3}^{2-}] + [B(OH)_{4}^{-}] + [OH^{-}] + [HPO_{4}^{2-}] + 2 [PO_{4}^{3-}] + \dots - [H^{+}] - [HSO_{4}^{-}] - \dots (1)$$

74 From Eq. (1), we see the increased OH⁻ concentration in a treated seawater solution corresponds to a salt solution

75 with increased alkalinity relative to the starting salt solution. This increase in OH⁻ ion concentration rapidly

76 increases the seawater pH upon mixing, resulting in a shift of the dissolved inorganic carbon (DIC) speciation

77 towards carbonate (Eisaman et al., 2023):

78

$$OH^- + HCO_3^- \rightarrow H_2O + CO_3^{2-}$$
 (2a)

 79
 $OH^- + CO_{2,aq} \rightarrow HCO_3^-$
 (2b)

- 80 The concentration of dissolved CO_2 gas ($CO_{2,ac}$) in this alkalinity-enhanced seawater is less than it would be if it
- 81 were in equilibrium with atmospheric CO₂ (Equation 2b). Over the longer timescale required for air-sea gas
- 82 exchange - weeks to months (Wang et al., 2023) or months to years (He and Tyka, 2023) depending on location - the
- 83 disequilibrium in the surface ocean resulting from the alkalinity addition drives the invasion of atmospheric CO_2
- 84 into seawater (or lessens the outgassing of CO_2 from the surface ocean to the atmosphere), where it reacts with

carbonate and is stored primarily in the stable bicarbonate phase (Jones et al., 2014; Bach et al., 2023; Renforth and
Henderson, 2017; Eisaman et al., 2023).

87
$$CO_{2,g} \to CO_{2,aq}$$
 (3a)
88 $CO_{2,aq} + CO_{3^{2^{-}}} + H_2O \to 2HCO_{3^{-}}$ (3b)

89 Under typical ocean conditions, after equilibrium has been reached, OAE results in an increase in the DIC in

seawater on the order of 0.7-0.9 moles of DIC per mole of NaOH added, with a slightly increased pH relative to theinitial value (He and Tyka, 2023).

92 It is possible that air-sea gas exchange will not completely drive the seawater pCO_2 to the initial unperturbed value 93 before the seawater sinks into the ocean interior and loses contact with the atmosphere for hundreds to thousands of 94 years. Therefore, the DIC anomaly relative to the alkalinity anomaly present when the seawater sinks into the ocean 95 interior may be used to assess the effective impact of the OAE for capturing atmospheric CO₂ on the 0-100 year

96 timescales that are most important for climate interventions.

97 In addition to the storage of atmospheric CO_2 in the form of DIC, this process may have the potential to locally and 98 transiently mitigate the elevated pCO_2 associated with ocean acidification (NASEM, 2021; Cross et al., 2023; 99 Butenschön et al., 2021). In a water body with a finite seawater exchange rate with the ocean, such as a semi-100 protected estuary or bay, alkalinity could be added in a controlled manner such that the combination of the rapid 101 reactions described by Eq.(1) and the exchange/flushing rate with the open ocean result in the bay being held in 102 steady-state at a target pH or aragonite saturation state value that is higher than its equilibrium value under 103 conditions of ocean acidification. As this added alkalinity diffuses through the bay and makes its way to the open 104 ocean, CO₂ removal and storage as DIC would occur. By metering the rate of alkalinity addition to the bay to match 105 the flushing rate, the pH or saturation state of the bay can be held at a constant target value. Even once equilibrium 106 has been achieved in the open ocean, the pH and the carbonate ion concentration in the open ocean remains slightly 107 higher than before the alkaline discharge. However, the absolute value of this pH increase after equilibrium has been 108 reached is sufficiently small relative to the alkalinity and DIC increase that mitigating ocean acidification on a 109 global scale with this method is unfeasible. For example, increasing the equilibrium pH value from 8.0 to 8.1 at a 110 fixed pCO_2 of 400 uatm (at 20 C and 35 salinity with no macronutrients) requires a TA increase of around ~620 111 µmol/kg-sw and a DIC increase of around 520 µmol/kg-sw. Using these numbers, mitigating OA over the entire 360 112 million $\rm km^2$ surface of the ocean to a depth of 100 meters would require around 487 gigatons of cumulative CO₂ 113 removal. Deploying SEA MATE in the ocean or coastal waters will require an understanding of carbonate chemistry 114 in seawater in the ocean volume under consideration, as well as thresholds for safe operation. For example, at the 115 point of alkaline dispersal where there is the maximum change in seawater chemistry, SEA MATE must control the 116 rate of alkalinity addition relative to the rate of mixing and dilution in the ocean to avoid the precipitation of 117 Mg(OH)₂ or CaCO₃ (Hartmann et al., 2023; Moras et al., 2022). While Mg(OH)₂ readily redissolves, an increase in 118 turbidity due to precipitation may negatively affect marine organisms (Bainbridge et al., 2018; Broderson et al., 119 2017). By contrast, CaCO₃ will generally not redissolve in the surface ocean without biological mediation, and 120 runaway precipitation, where alkalinity removed by precipitation exceeds that added by the OAE treatment, can 121 occur under conditions of increased aragonite saturation state and increased nucleation sites in the water column 122 (Moras et al., 2022). CaCO₃ precipitation could counteract the intended effect of the OAE intervention by removing 123 alkalinity from the surface ocean and releasing CO₂ gas via Eq. 5 (Zeebe and Wolf-Gladrow, 2001):

124
$$Ca^+ + 2HCO_3^- \rightarrow CaCO_3(s) + CO_2(g) + H_2O$$
 (5)

125 Upon dispersal to the ocean through a coastal outfall pipe, the added alkalinity is advected and diffuses away from 126 the point source, becoming increasingly diluted through the mixing zone. Because the timescale for air-sea gas

127 exchange and re-equilibration described by Eq. (2) is longer than the characteristic timescale for dilution driven by

- 128 tides, currents, and weather, most of the CO_2 removal occurs far from the mixing zone. Dilution will spread the
- impacts over a broad area, to an extent that it is unlikely that the impacts on the DIC distribution can be quantified
- 130 using only direct measurements, given current instrument resolution and the typical dynamic range of natural
- variability (Wang et al., 2023). In general, options for measurement, reporting, and verification (MRV) of OAE will
- therefore rely on (Ho et al., 2023): experimentation in laboratory and mesocosm settings, such as the work we
- describe here, to establish CO₂ removal dynamics under conditions of OAE; direct monitoring of the rate and
- 134 characteristics of alkalinity addition into seawater; monitoring the seawater carbonate and environmental chemistry135 in the immediate vicinity of the outfall via sensors and sampling (Cyronak et al., 2023; Schulz et al., 2023); and
- 136 ocean modeling to estimate CDR beyond the range of direct detection (Fennel et al., 2023).
- 137 While some work has investigated various aspects of NaOH-based ocean alkalinity enhancement in microcosms
- (Ferderer et al., 2022; Hartmann et al., 2023), and mesocosms (Groen et al., 2023), and other work has studied the
 release of NaOH over natural coral reefs as a method of local ocean acidification mitigation (Albright et al., 2016), a
- 140 systematic characterization of the efficiency and kinetics of OAE as a function of key process parameters has not yet
- been performed. Here we report the first tank-scale tests of OAE that use aqueous hydroxide (NaOH) to enhance the
- 142 alkalinity of natural seawater, a process that mimics OAE via the electrochemical brine-to-alkalinity conversion
- 143 used in the SEA MATE process. Our experiments, conducted in 6,000 liter tanks using seawater pumped from Flax
- Pond on Long Island Sound in Stony Brook NY, quantify the magnitude and timescale of the CO₂ removal from the
- 145 air and storage as seawater DIC by monitoring the air-seawater re-equilibration after an initial alkalinity
- 146 perturbation. In addition, our use of both laboratory-processed bottle samples and field-deployable sensors to
- 147 measure and over-constrain the carbonate chemistry response allows us to assess the suitability of certain sensing
- 148 platforms for MRV. Finally, we investigate safe thresholds for the rate and concentration of alkalinity addition to
- avoid: (1) the precipitation and redissolution of Mg(OH)₂ that can lead to local, temporary increases in turbidity; and
- 150 (2) the precipitation of CaCO₃, which partially reverses the intended OAE effect by removing alkalinity from, and
- 151 releasing CO₂ gas into, the surrounding seawater.
- 152 Using this approach, we address the following key questions:
- (1) How much additional atmospheric CO₂ is stored in seawater as DIC in response to a given alkalinityperturbation?
- (2) What is the timescale for CO₂ removal from the air, and how does it depend on pH and the magnitude ofalkalinity enhancement?
- (3) What are the conditions for Mg(OH)₂ precipitation upon addition of NaOH to seawater?
- (4) What are the threshold values for pH and aragonite saturation state beyond which undesired CaCO₃ precipitationwill occur?
- 160 Answering these questions is key to assessing the viability of this approach and to optimizing its eventual
- 161 deployment. The large tank experiments presented in this manuscript provide a stepping stone between bench-scale
- 162 experiments and in-situ mesocosms or field pilots. Even if these experiments simply confirm stoichiometric and
- 163 modeled expectations, this is critical information in the design and implementation of OAE deployments. This work
- 164 is a necessary part of the growing scientific body that will allow for field trials to progress.
- 165 2. Methods
- 166 2.1 Experimental procedure

- 167 We investigated the carbonate chemistry changes resulting from the addition of NaOH_(aq) to natural seawater over
- 168 timescales ranging from 2 weeks to 2 months in a series of experiments at two scales: (1) two large (~6200 L)
- 169 indoor tanks, and (2) multiple 15 L aquaria (Fig. 1). The large volume of tank experiments allowed for precise
- 170 measurement of the seawater carbonate chemistry via bottle sampling (1L each, sent to NOAA/PMEL for analysis) 171
- with high sampling frequency. To compliment these measurements, we also performed a series of experiments in 172
- smaller aquaria (15 L each), which enabled a larger number of replicates and a faster time to equilibrium when
- 173 bubbled with air.



175	Figure 1: (<i>a</i> , <i>b</i>) Flax Pond Marine Laboratory is located on Long Island Sound, New York, USA (© Google Earth
176	2022). (c) The ~6000 L control and experiment tanks were instrumented with a series of oceanographic sensors and
177	sampled routinely for DIC/ TA analyses to allow for measurement of carbon uptake over time following an addition
178	of alkalinity in the form of NaOH. The ~15 L aquaria were instrumented with standard pH electrodes and monitored
179	with routine TA analyses. The Forced air* and Ambient air streams* indicate their use in some but not all
180	experiments, as noted in later sections.

- 181 This study was conducted at the Flax Pond Marine Laboratory at Stony Brook University, NY. All experiments used 182 natural seawater collected from Flax Pond, part of a 128-acre salt marsh tidal wetlands connected to the Long Island
- 183 Sound. The surface areas of the tanks and aquaria were ~4.6 m^2 and ~0.1 m^2 , respectively. The tanks had a diameter
- 184 of 2.4 m, a total height of 1.52 m, and were typically filled to a height of ~1.35 m, allowing for a corresponding
- 185 seawater volume of 6185 L. The aquaria had a diameter of 0.3 m and were typically filled to a height of ~0.23 m, for
- 186 a total seawater volume of 15 L. The large tank volumes were chosen to limit interactions with walls while
- 187 increasing the air-seawater boundary, and to lose a smaller fraction of their volume to evaporation. These tanks
- 188 allow for in-situ oceanographic sensor deployment and frequent bottle sampling while retaining semi-controlled 189
- temperature, mixing, filtration, and biological control. The inherent limitations of these tank tests include limited air-190 sea interaction, unrealistic light levels and circulation, and biological responses that are not a perfect representation

- 191 of natural seawater in the ocean, but serve as a stepping stone to mesocosm and eventual field experiments. On
- average, the large (~6,000 L) tank experiments took ~6.5 weeks after dosing with NaOH to reach 90% of the
- 193 calculated or extrapolated asymptotic Δ DIC/TA addition ratio indicative of full air-seawater equilibrium, as will be
- discussed in Section 3. Therefore, in addition to the large tank tests, we conducted a series of smaller aquaria alkalinity additions to increase our capacity for experimental test cases. The limitations of the aquaria include
- alkalinity additions to increase our capacity for experimental test cases. The limitations of the aquaria include
 limited sensor options, unrealistic circulation, and limited biological control. While it is expected that equilibration
- 197 occurs more rapidly in the small aquaria than in the large tanks, the results from these cases should be similar as
- 198 CO₂ equilibrates across the air-sea boundary. However, we note that some variation is expected due to limited
- sensing and sampling options in the smaller aquaria and the greater potential for biological growth in the large tanks
- 200 over longer timescales.

201 2.1.1 Tank experiments

- 202 Seawater was pumped into the tanks at high tide through a series of sock filters to exclude macroscopic biology. The
- tanks were then dosed to 40 ppm bleach (sodium hypochlorite) and the shock-treated seawater was allowed tocirculate through the tanks for ~1 day to limit biological growth. The seawater was then circulated through UV light
- arrays to break down the bleach over ~1-2 weeks, as assessed by a standard Hach test kit for free chlorine. During
- this period, seawater was pumped between the two large test tanks (~25 L/min) to increase mixing of the bleach and
- to homogenize the tanks to similar initial conditions. For the remainder of each experiment, the seawater was
- 208 continually pumped through the UV sterilizers. Measurements of total alkalinity showed no significant differences
- in the bulk seawater TA before and after the bleaching process in any experiment or control tank. In an early
- 210 experiment (in which bulk pH_T was raised from the initial condition to 8.7, as described below), the initial pH_T and
- 211 DIC varied between the control and experiment tanks by 0.17 and 77μ mol kg⁻¹, respectively. This was because
- seawater was pumped from Flax Pond into multiple reservoirs and was then unevenly distributed between the tanks.The experiments were subsequently refined to allow for several days of cross-pumping between tanks to
- homogenize the control and experiment seawater before NaOH was added at the start of an experiment. More details
- 215 on experimental variations and a larger summary table are available in the Supplementary Materials.
- 216 Oceanographic sensors and discrete daily bottle sampling, as described in Sections 2.2 and 2.3, respectively, were 217 deployed for carbonate chemistry analysis for several days prior to the alkalinity addition to understand the initial 218 baseline conditions in both tanks. Two submerged pumps were used for water circulation within each tank: the first 219 pump (Current eFlux DC Flow Pump, 210 GPH) cycled seawater through the UV arrays with an estimated 220 overturning time of the bulk tank on order of 1 day, and a second (Kedsum Submersible pump, 260 GPH), mounted 221 at an angle halfway down the tank wall, allowed for subsurface circulation within the tank to reduce the occurrence 222 of unmixed 'dead zones' and subsequent non-homogenous biological growth, as assessed visually on the surface of 223 the water and/or tank lining. Initial tank experiments were conducted with a still surface condition, i.e., with no 224 visible water movement across the surface of each tank. As experiments progressed, forced air movement was added 225 across the surface of each tank using a stationary fan with a wind speed of ~5 kph. This was done to control for 226 potential variations in the laboratory HVAC system and to potentially reduce the time to equilibration for the 227 experiments by increasing the rate of air-sea CO₂ equilibration. In later experiments, air was bubbled into the bottom 228 of each tank at a rate of ~ 30 L min⁻¹ with an estimated surface area of ~ 0.3 m², with a goal of further increasing the 229 rate of equilibration to allow for more rapid throughput of experiments. These variations are further discussed in 230 Section 2.4.
- After baselining, one tank (referred to as the "experimental tank") was dosed with enough 0.5 M NaOH (see
- 232 Supplementary Materials) to raise the bulk seawater pH to the target pH of interest for a given experiment, and the
- same volume of DI water was added to the other tank (referred to as the "control tank"). NaOH additions were
- typically dosed into the tank via peristaltic pump at a low enough rate (~50 mL/min) that a steady increase in bulk
- tank pH was observed, but local pH measured just below the NaOH introduction never exceeded a pH of 9.0. A

- pump (~25 L/min) was placed just below the NaOH stream to speed the mixing of NaOH into the bulk tank,
- 237 increase dilution from the point source, and to prevent the immediate precipitation of Mg(OH)₂ upon contact of the
- 238 NaOH with seawater. This pump was removed after the full volume of NaOH was mixed into the tank.
- After the alkalinity addition, the tanks were left to equilibrate with the atmosphere and were monitored by sensors
- and sampling as described in Sections 2.2 and 2.3. The tanks were indoors in the wet laboratory at Flax Pond Marine
- Lab, such that temperature and CO_2 concentration were moderated by the building's HVAC system, but varied
- throughout days and seasons depending on other uses of the lab space. The experiments were concluded when the observed pH or DIC (calculated from daily pH and frequent TA measurements) appeared to stabilize (e.g., Δ pH
- $\pm 0.05\%$ or $\Delta DIC \pm 10 \ \mu mol \ kg^{-1}$ per day) over several days. The continuous improvement of experimental methods
- 245 during this study resulted in some minor variations among the methods used for each experiment, including methods
- of NaOH dosing, tank circulation, and biological control, as discussed where necessary in Section 3 and in the
- 247 Supplementary Materials.

248 2.1.2 Aquaria experiments

- 249 A series of polycarbonate aquaria were filled with 15 L of seawater taken from the large control tank just after the 250 described bleaching and bleach breakdown procedure was completed. NaOH was dosed into each aquarium to reach 251 a targeted bulk pH_T, with a corresponding volume of DI H₂O added to the control aquarium, and then the seawater 252 was allowed to equilibrate with atmospheric pCO_2 over days to weeks. The aquaria did not have either UV light 253 arrays for biological control or aquarium pumps for internal circulation. With the exception of a single target $pH_T 8.5$ 254 experiment, all aquaria were bubbled with ambient air (~4 L min⁻¹) via a standard aquarium bubbling bar spanning 255 the center diameter of each aquarium, allowing for rapid CO₂ exchange. There was no fine control on air bubbling, 256 but the surface area of all air bubbles in a given aquarium at any point in time was estimated at ~0.01 m². No sensors 257 were deployed in the aquaria due to their limited size, and seawater chemistry was established via discrete pH_T and 258 TA measurements (Sect. 2.2). An optically clear lid was placed on each aquarium to reduce evaporation and 259 splashing onto nearby equipment. Some evaporation was evident from the rising TA throughout these experiments, 260 but was not resolvable within the resolution of a handheld salinometer used for these experiments. Temperature was 261 discretely recorded from a combination Ross pH electrode.
- 262 As shown in Eq. (6), we define the dimensionless 'Carbon-to-Alkalinity Ratio' (CAR) for our experiments as the 263 molar ratio of the increase in *n*DIC (in units of μ mol/kg, normalized to the system's initial salinity to account for 264 evaporation) to the magnitude of the TA increase (ΔTA , in units of μ mol/kg). *n*DIC_{eou} is the measured (via direct 265 titration) or calculated (via CO2SYS using measured TA and pH_T) DIC value that the system reached at the end of 266 an experiment (Pierrot et al., 2006; Van Heuven et al., 2011). Some experiments were left long enough to achieve 267 equilibration with atmospheric CO₂, but others were halted early. In these cases, a CO2SYS calculation was used to 268 estimate the DIC increase expected at equilibration given initial seawater conditions, and the difference between this 269 value and the final recorded *n*DIC_{equ} was used to estimate the overall percent equilibration for a given experiment. 270 Depending on experimental constraints described in later sections, $nDIC_i$ may represent either: (1) the final nDIC271 measured (via titration of bottle samples) or calculated (via CO2SYS using seawater TA and pH) in the control tank, 272 or (2) the 'baseline' *n*DIC before the addition of NaOH to a given aquaria experiment, for cases where a 273 corresponding control case may not be available. Note that because we are reporting CAR values where the 274 measured DIC has reached or has been estimated at equilibrium, the CAR values we measure and report reflect the 275 ratio of Δ DIC to Δ TA that would be expected given sufficient time for air-sea exchange to reach equilibrium, and so 276 are equivalent to directly measuring the value of the "TA addition potential impact ratio" as defined by Wang et al., 277 2023.
- 278 Carbon-to-Alkalinity Ratio (CAR) = $(nDIC_{equ} nDIC_i) / \Delta TA$ (6)

279 2.2 Oceanographic sensors

- 280 Each tank was instrumented with a series of sensors placed halfway down the wall of the tank near the inlet of the
- 281 UV circulation pump. A Seabird 38 Digital Oceanographic Thermometer and Seabird 45 MicroTSG
- Thermosalinograph continuously monitored seawater temperature and salinity, respectively. Dissolved oxygen was
 measured by a PME miniDOT Logger at 10 min resolution. pH_T was monitored daily by a SAMI-pH (manufacture)
- 283 measured by a PME miniDOT Logger at 10 min resolution. pH_T was monitored daily by a SAMI-pH (manufacturer 284 specified accuracy/precision ~ 0.003/0.001, though this accuracy is likely an underestimate of the uncertainty given
- 285 known challenges for the calibration of the pH_T measurements) and by a semi-automated spectrophotometric (spec-
- pH) pH unit (~± 0.0055/0.0004) as described by Carter et al. (2013). CRM measurements were taken by each pH
- system at the beginning and end of each experiment and were used alongside discrete samples of DIC and TA as
- described in Section 2.3 to constrain the stability of each sensor. The SAMI-pH measurements were recorded at
 ambient seawater temperature and corrected for in-situ salinity as recorded by the Seabird Thermosalinograph
- following best practices from the manufacturer. The spec-pH analyses occurred in a jacketed cuvette held at 20 °C
- (regulated via water bath) and were corrected to the in-situ bulk tank temperature and salinity as recorded by theSeabird Thermometer and Thermosalinograph. Both the SAMI-pH and spec-pH rely on spectrophotometric analysis
- 293 of metacresol purple indicator dye, which allows for pH measurement within the pH_T range of approximately 7 to 9.
- For experiments in which enough NaOH was dosed into seawater to raise pH above these limits, a Thermo Scientific
- 295 Orion ROSS Ultra pH/ATC Triode combination electrode (8157BNUMD) was used to monitor pH_{NBS} at the surface
- 296 of the tank (± 0.01 precision), which was then converted to pH_T for comparison with the other pH sensor systems.
- A LiCOR LI-850 sensor was used to analyze atmospheric pCO_2 (±1.5% accuracy) above the tanks. The inlet to this sensor was periodically moved between tanks to ensure that atmospheric pCO_2 in the vicinity of the control and experiment tanks was the same. AcuRite Iris weather stations were mounted on the side of each tank to monitor air temperature (±2 °C), relative humidity (±3%), and air speed (±0.8 m s⁻¹). All data were compiled on an hourly basis in a custom R package.

302 2.3 Discrete sampling

- 303 Two types of discrete sampling were used to constrain carbonate chemistry throughout these experiments. First, 500
- 304 mL of seawater was collected and preserved from each tank, typically on a daily basis, and as frequently as hourly
- during the addition of NaOH, following best practices laid out by Dickson (2007) including overflowing of the
- 306 sample bottles during collection and addition of 0.2 mL of saturated mercuric chloride (HgCl₂) as a preservative.
- 307 These bottle samples were analyzed for DIC and TA at NOAA Pacific Marine Environmental Laboratory
- 308 (NOAA/PMEL). DIC concentrations were measured using a coulometer (UIC Inc.) and Single Operator
- 309 Multiparameter Metabolic Analyzer (SOMMA) (Johnson et al., 1985). TA was determined by an open-cell
- acidimetric titration (Dickson et al. SOP 3b, 2007). The accuracy of DIC and TA measurements was assessed with
- 311 Certified Reference Materials (CRMs, supplied by the Dickson laboratory at Scripps Institution of Oceanography),
- 312 and overall uncertainty for both DIC and TA was typically $\pm 0.1\%$ (~2 µmol/kg).
- 313 In addition, discrete seawater samples were analyzed for TA via open-cell potentiometric titration at Stony Brook
- 314 University. A Thermo Scientific Orion ROSS Ultra pH/ATC Triode combination electrode (8157BNUMD),
- 316 seawater sample with a dilute HCl solution (~0.1 M in 0.7 M NaCl, calibrated daily with CRM or a secondary
- seawater standard) following a modified Gran titration procedure using a Kloehn digital syringe pump (Song et al.,
- 318 2020; Wang and Cai, 2004). The precision of TA measurements was ~±5-10 µmol/kg. This TA data was corrected
- to that of the bottle samples analyzed via titration at NOAA PMEL where available (see Supplementary Materials).
- 320 There are several differences between the aquaria experiments and the larger tank experiments. First, the aquaria
- 321 experiments were monitored daily to every few days by discrete measurement of TA at Stony Brook University and

- 322 pH_{NBS} via Thermo Scientific Orion ROSS Ultra pH/ATC Triode combination electrode (8157BNUMD) (±0.01
- **323** precision), which was then converted to pH_T and corrected against the other pH sensor systems via occasional bottle
- 324 samples for DIC and TA analysis at NOAA PMEL. Variations between these experiments are noted in Section 3
- 325 where necessary and in the Supplementary Materials.
- 326 In either tank or aquaria cases where mineral precipitation was observed, 0.5 1 L of seawater was vacuum filtered
- through a 0.45 μm Whatman GF/F filter via vacuum pump and the solids were rinsed with DI water 3 times to
- remove NaCl. The precipitate was dried in an oven at 90 °C, then crushed into a uniform powder via mortar and
 pestle. Samples were analyzed via Hitachi 4800 Scanning Electron Microscopy (SEM) (5 kV) and Rigaku SmartLab
- pestle. Samples were analyzed via Hitachi 4800 Scanning Electron Microscopy (SEM) (5 kV) and Rigaku SmartLab X-ray Diffraction (XRD) (Cu Kα, 1.5406 Å, 10 - 100° 2 θ at 4°/min) at Brookhaven National Laboratory at the
- 331 Materials Synthesis and Characterization Facility of the Center for Functional Nanomaterials.

332 2.4 Evaluation of CO₂ uptake by seawater in response to NaOH perturbation

- 333 Seawater carbonate chemistry measurements were used to analyze the uptake of CO₂ in each tank, primarily relying
- 334 on calculations from the NOAA/PMEL DIC and TA analyses of bottle samples when available and using sensor pH
- and Stony Brook TA measurements for cross-verification or to fill in between discrete DIC samples. DIC and TA
- data were normalized to the salinity at the start of a given experiment to account for evaporation (Friis et al., 2003).
- 337 Carbonate chemistry calculations were then performed using CO2SYS (Lewis and Wallace, 1998), with Lueker et
- al. (2000) carbonate constants, Dickson (1990) for KSO₄, and Lee et al. (2010) for total boron. Wherever possible, a
- 339 combination of CRM analyses and comparisons between simultaneous pH sensor and NOAA PMEL bottle samples
- 340 were used to correct SAMI-pH and spectrophotometric pH sensor data for drift.
- 341 Changes in the seawater carbonate chemistry over time were analyzed with respect to shifts away from the baseline
- 342 within a single control or experiment tank, as well as with respect to the differences between the control and
- 343 experimental tanks.

344 3 Results and Discussion

345 3.1 Large tank experiments

346 A summary of the range of oceanographic variables measured by sensors and bottle samples, calculated via 347 CO2SYS, or extrapolated to equilibration conditions during the large tank experiments is provided in Table 1. This 348 summary includes 6 experiments including 3 targeting pH_T 8.5 (still surface water, with forced air, and with forced 349 air and air bubbling) and one (each) targeting pH_T values of 8.7 (still surface water), 9.5 (with forced air and air 350 bubbling), and 10.3 (still surface water). While the initial seawater conditions were similar between the control and 351 experiment tanks, we note that these cases are not entirely comparable after the termination of cross-pumping 352 between tanks and the subsequent addition of alkalinity. While tanks were initially bleached, eventually some 353 biological growth was noted in each tank with potential differences in spatial and temporal distribution as well as 354 species and community differences. Herein, we assume that differences between the control and experiment cases 355 are due to the addition of alkalinity alone, but we note that characterization of other potential confounding factors is 356 a subject for future work.

- 357 The initial pH_T, TA, and DIC varied across experiments as seawater was collected between March 2022 and May
- **358** 2023, ranging from pH 7.66 (December 2022) 7.95 (May 2023), TA 2001 (May 2023) 2176 (March 2023)
- μmol/kg, and DIC 1847 (May 2023) 2021 (March 2023) μmol/kg. Both measured and CO2SYS -calculated DIC
- and TA values were normalized to salinity to account for evaporation, which drove salinity increases ranging from
- $361 \qquad 0.2 7.1 \text{ across these experiments.}$

- 362 After the addition of NaOH, the control and experiment tanks were allowed to equilibrate with atmospheric CO₂.
- 363 While refinements in the experimental design allowed for complete or near-complete equilibration in later
- experiments, as determined by the stabilization of nDIC at some asymptotic value, early experiments were
- terminated before full equilibration. In all experiments, the absorption of atmospheric CO₂ began immediately after the NaOH addition, as determined by decreasing pH and Ω_{arag} and increasing DIC and seawater *p*CO₂. nTA was
- fairly stable or increasing (+10 60 μ mol kg⁻¹) after the NaOH addition in all cases except the pH_T = 10.3
- **368** experiment, where nTA and DIC rapidly decreased due to runaway CaCO₃ precipitation. A stable TA value is an
- indicator that no significant persistent mineral precipitation (e.g., Mg(OH)₂ or CaCO₃) has occurred. In the absence
- of active mixing or bubbling, Mg(OH)₂ precipitation occurred immediately upon the introduction of NaOH to
- 371 seawater, however the precipitation can be rapidly dissolved by turbulence (i.e., pumping NaOH directly above a
- 372 strong circulation pump and/or stream of air bubbles). No CaCO₃ precipitation was observed in the tanks or aquaria
- 373 for which the bulk seawater pH_T was <10.0. The $pH_T = 10.3$ experiment was designed to induce CaCO₃ runaway
- **374** precipitation, as described in Section 3.3.
- 375 Ω_{arag} ranged from 1.4 2.5 in the control tanks with minimal variation over the course of any given experiment.
- **376** During the three experiments in which bulk pH_T was increased to ~8.5, Ω_{arag} increased immediately to 6.0 6.3 at the
- peak of the experiments, before slowly decreasing to 2.8 3.0 as the seawater equilibrated with atmospheric CO₂.
- **378** For the bulk pH_T 9.5 experiment, Ω_{arag} increased to 20.2 and slowly decreased to 5.0 when the experiment was
- 379 ended at full equilibration. Mineral precipitation was observed in the bulk pH_T 10.3 experiment, where Ω_{arag} was
- **380** increased to 30.3 and rapidly (<1 week) fell to 5.2 after the addition of NaOH.
- 381 The results of one representative set of time-series measurements from the control and experiment tanks are shown
- in Figure 2 for the case where pH_T of the bulk experiment tank was raised to 8.5 then allowed to relax into
- again the atmosphere without the addition of surface air forcing or bubbling. Time-series plots for the
- 384 other tank-scale experiments are available in the Supplementary Materials.





- **Figure 2:** Time-series data for the case where pH_T of the bulk experiment tank was raised to 8.5 with no forced air
- 388 flow and no bubbling (still surface) for control (blue, solid) and experiment (red, dashed) tanks: (a) continuously
- 389 measured air temperature, (*b*) atmospheric *p*CO₂, (*c*) seawater temperature, (*d*) salinity, and (*e*) dissolved oxygen; (*f*)
- 390 pH_T measured by the SAMI-pH (*circles*) and interpolated from the spec-pH (*line*), corrected to bottle sample and
- **391** CRM data; (g) NOAA/PMEL-measured TA and (h) DIC from bottle samples and normalized to salinity; (i)
- seawater pCO_2 and (*j*) saturation state of aragonite (Ω_{arag}) calculated from interpolated nDIC and nTA data via
- **393** CO2SYS; (*k*) the observed carbon uptake ratio (CAR) as $(n\text{DIC}_{exp} n\text{DIC}_{control}) / \Delta TA_{NaOH addition}$ (*solid*) and the
- theoretical CAR (*dashed*) from a CO2SYS calculation using measured TA and the average $pCO2_{atm}$ to estimate the equilibrium change in DIC (*dashed*); (*l*) the percent equilibration estimated between the observed and theoretical
- 396 CAR. Data gaps in panels a, b, and e are due to connectivity issues while offloading sensor data.
- **397** The Δ nTA and Δ nDIC values calculated between the control and experiment tanks are summarized in Figure 3
- 398 where nTA and nDIC were interpolated between bottle samples measured at NOAA-PMEL, and/or were calculated
- via CO2SYS using sensor pH_T and TA measured at Stony Brook University corrected to less frequent NOAA-
- 400 PMEL TA and DIC bottle samples. The ratio of the Δ nDIC to the addition of alkalinity in the form of NaOH, or
- 401 ΔnTA , is included in Figure 3 for all experiments except that of the bulk pH_T increase to 10.3. Neglecting
- 402 experiments that were terminated before full equilibration, the final observed CAR ranged from 0.75 ± 0.04 to 0.87
- 403 ± 0.08 (Table 1).
- 404 An anomalous event was noted in both the experiment and control cases for the target pH_T 8.5 experiment with
- 405 forced air movement across the surface of the tank, wherein an increase in TA and DIC was noted around day 30 of
- 406 the experiment. The cause of this event is unclear but could include biological changes in both tanks, the
- 407 introduction of alkalinity from environmental contaminants, or the anomalous delayed release of alkalinity from
- 408 suspended solids. This event was not observed in any other case, and highlights the importance of using controls to
- 409 understand complex interactions in these experiments. A time-series including this event is available in the
- 410 Supplementary Materials.
- 411 Henry's law and CO2SYS calculations were used to estimate the initial and final equilibration condition of each
- tank experiment. LiCOR $pCO_{2,atm}$ measurements were averaged across experiments to a representative value of 421
- 413 \pm 14 ppm, which was used with the initial seawater temperature and salinity to estimate $pCO_{2,seawater}$ at the beginning
- 414 of each experiment. This initial $pCO_{2,seawater}$ was in all cases greater than the atmospheric $pCO_{2,seawater}$, indicating 415 that the seawater was not fully equilibrated with the atmosphere at the time when NaOH was added, likely due to
- 415 that the seawater was not fully equilibrated with the atmosphere at the time when NaOH was added, likely due to 416 respiration and decomposition of organic material (Section 2.1), and as such, the tanks should outgas CO₂. The
- initial equilibrium DIC was estimated from a CO2SYS calculation using the $pCO_{2,seawater}$ and nTA_i , which in all
- 418 cases was less than the initial *n*DIC measured or calculated from nTA_i and $pH_{T,i}$ (by 29 108 µmol kg⁻¹). These
- 419 observations underscore the importance of having a control tank to capture natural dynamics of CO₂ ingassing and
- 419 observations underscore the importance of naving a control tank to capture natural dynamics of CO₂ ingassing420 outgassing to ensure that changes in DIC attributed to OAE are correctly accounted for.
- 421 The final equilibrium *n*DIC was estimated from a CO2SYS calculation using the same $pCO_{2,seawater}$ and the *n*TA
- 422 measured just after the NaOH addition, corrected for the linear increase in salinity over the course of the experiment.
- 423 The ratio of the expected $\Delta nDIC$ calculated at equilibrium with the atmosphere to the addition of alkalinity provides
- 424 a simple estimate of the expected CO_2 storage capacity for a given experiment. The percent equilibration for each 425 experiment was then estimated from the measured and expected values for CAR. Within the series of experiments
- 425 experiment was then estimated from the measured and expected values for CAR. Within the series of experiments 426 with a targeted pH_T of 8.5, the timeline to reach an estimated 90% CO₂ equilibration decreased from 65 days (with
- 426 with a targeted pH_T of 8.5, the timeline to reach an estimated 90% CO₂ equilibration decreased from 65 days (with 427 internal circulation but still water at the surface of the tank), to 50 days (with the addition of forced air movement
- 428 across the surface of the tank) to 22 days (with the addition of air bubbling). We note that only the two cases
- 429 (targeted pH_T of 8.5 and 9.5) with the addition of air bubbling reached full equilibration with the atmosphere.

430 Table 1: Range of variables measured, calculated, or extrapolated in large tank experiments, where M denotes direct

431 measurement, C denotes calculation via CO2SYS, and E denotes extrapolation to equilibrium conditions. Subscripts

432 *i* and *f* refer to initial and final conditions, and 'peak' refers to the time point immediately after the addition of

433 NaOH.

pH target -		8.5		8.5		8.5		8.7		9.5		10.3	
Surface condition	-	Still		Forced Air		Forced Air and Air Bubbles		Still		Forced Air and Air Bubbles		Still	
Tank (C = control, E = experiment)	-	С	Е	С	Е	С	Е	С	Е	С	Е	С	Е
$\Delta TA = NaOH$ addition (± 10 μ mol/kg)	М	0	409	0	462	0	375	0	626	0	1406	0	3305
Salinity _i (g/kg)	М	28.7	28.7	30.2	30.2	30.4	30.4	26.9	26.8	26.9	26.9	28.5	28.4
Salinity _f (g/kg)	М	30.5	30.2	37.3	36.6	34.7	33.7	27.6	27.6	29.0	29.2	28.6	28.6
$pH_{T,i} (\pm 0.005)$	М	7.76	7.76	7.73	7.73	7.93	7.93	7.92	7.75	7.95	7.95	7.70	7.75
pH _{T,peak} (± 0.005)	М	-	8.54	-	8.58	-	8.49	-	8.68	-	9.51	-	10.10
$pH_{T,f} (\pm 0.005)$	М	7.88	8.05	7.85	7.99	7.99	8.01	7.84	8.26	8.01	8.21	7.75	9.52
$nTA_i (\pm 10 \ \mu mol/kg)$	М	2049	2049	2069	2069	2248	2248	2075	2075	2007	2007	2023	2025
$nTA_{peak} (\pm 10 \ \mu mol/kg)$	М	-	2458	-	2531	-	2623	-	2701	-	3414	-	5330
$nTA_{f} (\pm 10 \ \mu mol/kg)$	М	2080	2528	2235	2674	2246	2624	2095	2696	2014	3363	2041	1253
nDIC _i (µmol/kg)	М	1944	1947	1957	1996	2082	2087	1897	1975	1852	1852	1928	1938
nDIC _f (µmol/kg)	М	1908	2280	2084	2433	2027	2365	1937	2336	1832	2977	1947	720
$\Omega_{aragonite,i}$	С	1.39	1.37	1.4	1.1	2.0	2.0	2.4	2.4	1.9	1.9	1.4	1.3
$\Omega_{\mathrm{aragonite,peak}}$	С	-	5.9	-	6.0	-	6.2	-	8.8	-	19.3	-	30.3
$\Omega_{\mathrm{aragonite,f}}$	С	2.0	3.0	1.7	2.8	2.5	3.0	1.9	4.4	2.1	4.9	1.4	5.2
CAR _f	С	-	0.85 ± 0.04	-	0.75 ± 0.04	-	$\begin{array}{c} 0.87 \pm \\ 0.08 \end{array}$	-	0.52 ± 0.07	-	0.82 ± 0.09	-	-
CAR _{equilibrium}	Е	-	0.89	-	0.85	-	0.85	-	0.84	-	0.81	-	-
% equilibration (time elapsed in days)	Е	-	95 ± 10 (67)	-	92 ± 10 (63)	-	102 ± 12 (45)	-	79 ± 6 (42)	-	104 ± 7 (74)	-	(13)







436 Figure 3: Results of 5 tank-scale experiments in which enough NaOH was added to each tank to raise the bulk pH_T 437 to 8.3 - 9.7. pH_T decreased rapidly in all cases in which air bubbling sped equilibration with atmospheric CO₂. 438 Results include: (a) measured pH_T, (b) measured nTA, (c) measured nDIC or CO2SYS calculated (for pH_T 9.5 case 439 only), (d) CO2SYS -calculated pCO2, (e) the observed carbon uptake ratio (CAR) as (nDICexp - nDICcontrol) / 440 Δ nTA_{NaOH addition} with horizontal dashed lines representing the expected range of 0.7-0.9 mol CO₂ uptake / mol NaOH 441 added to seawater, the change in (f) nTA and (g) nDIC compared to the baseline measurements before the addition

442 of NaOH, and the percent equilibration estimated between the observed and theoretical CAR.

443 **3.2 Aquaria experiments**

444 Table 2 provides a summary of the range of oceanographic variables quantified for the aquaria experiments.

445 Table 2: Range of variables measured, calculated, or extrapolated in aquaria experiments, where M denotes direct

446 measurement, C denotes calculation via CO2SYS, and E denotes estimation within specified equilibration

447 conditions. Subscripts i and f refer to initial and final conditions, and 'peak' refers to the time point immediately 448 after the addition of NaOH.

pH target	-	0 Control	8.3	8.5	8.5 Without air bubbles	8.7	9.3	9.5	9.7	9.9	10.0	10.1	10.2	10.3
$\Delta TA = NaOH$ addition (± 10 μ mol/kg)	М	0	187	331	362	543	1409	1679	2037	2216	2276	2504	2796	3829
$pH_{T,i} \left(\pm \ 0.1\right)$	М	7.94	7.97	7.90	7.86	7.95	7.98	7.98	7.98	8.06	8.04	8.04	8.04	7.95
pH_{T} , _{peak} (± 0.1)	М	-	8.28	8.41	8.40	8.63	9.22	9.43	9.64	9.83	9.91	10.23	10.32	10.20
$pH_{T,f}(\pm0.1)$	М	8.06	8.03	8.07	8.11	8.08	8.21	8.20	8.23	8.65	8.96	8.72	9.46	7.99
$\frac{TA_i (\pm 10)}{\mu mol/kg}$	М	2265	2262	2250	2250	2250	2393	2393	2393	2531	2531	2531	2531	2250
$\begin{array}{l} TA_{peak} \left(\pm 10 \right. \\ \mu mol/kg \right) \end{array}$	М	-	2449	2582	2611	2793	3801	4072	4430	4748	-	-	-	4608
$\frac{TA_{f} (\pm 10)}{\mu mol/kg}$	М	2323	2476	2640	2645	2822	3837	4110	4420	4462	1702	1835	1537	2202
DIC _i (µmol/kg)	С	2089	2073	2091	2107	2070	2192	2192	2192	2282	2287	2287	2287	2067

DIC_{f} (µmol/kg)	С	2113	2246	2377	2382	2540	3372	3486	3877	3389	992	1244	671	2003
$\Omega_{aragonite,i}$	С	2.1	2.2	1.9	1.8	2.1	2.34	2.4	2.4	2.9	2.8	2.8	2.8	2.1
$\Omega_{aragonite,peak}$	С	-	4.2	5.5	5.5	8.1	19.5	23.1	27.0	29.8	30.2	30.9	32.4	38.9
$\Omega_{aragonite,f}$	С	2.4	2.7	3.1	3.1	3.4	5.9	7.9	7.1	13.7	6.5	5.7	7.0	2.2
CAR _f	С	-	$\begin{array}{c} 0.92 \pm \\ 0.10 \end{array}$	0.87 ± 0.06	$\begin{array}{c} 0.76 \pm \\ 0.05 \end{array}$	$\begin{array}{c} 0.87 \pm \\ 0.04 \end{array}$	$\begin{array}{c} 0.84 \pm \\ 0.02 \end{array}$	$\begin{array}{c} 0.86 \pm \\ 0.02 \end{array}$	$\begin{array}{c} 0.84 \pm \\ 0.02 \end{array}$	0.50	-	-	-	-
$\mathrm{CAR}_{\mathrm{equilibrium}}$	Е	-	0.69	0.67	0.64	0.77	0.80	0.80	0.80	0.81	-	-	-	-
% equilibration (time elapsed in days)	E	(40)	130 (16)	126 (18)	116 (40)	111 (16)	104 (18)	106 (18)	104 (18)	62 (1)	(1)	(1)	(1)	(16)
CaCO ₃ precipitation?	Μ	-	No	No	No	No	No	No	No	No	Yes	Yes	Yes	Yes

449 The aquaria experiments are not directly comparable to the control stated in Table 2. Seawater for one control

450 aquarium was collected in March 2023, and was monitored for pH_T and TA changes through May 2023. Seawater

451 for the experimental aquaria was collected in three batches between March, April, and May 2023, with only 4-6

452 aquaria experiments running in parallel within each set of experiments due to space and analytical throughput

453 constraints. Because of this, the experiments started in March 2023 could be compared directly to the control (target

454 pHT 8.3, 8.5, 8.5 still, and 8.7), but the rest of the experiments used different initial seawater than the control

455 aquarium. The CAR for each aquaria experiment was therefore calculated from changes in DIC and TA between the

456 initial 'baseline' condition and after the NaOH was added within a given aquarium, rather than between the

457 experiment and control cases. The CAR ranged between 0.76 ± 0.05 and 0.92 ± 0.10 , excluding cases where mineral 458 precipitation was evident and for the pH_T 9.9 case where the experiment ended after one day due to a sensor logging

459 failure. This wide range in $\Delta DIC / \Delta TA$ is likely due to the limited number of TA samples collected throughout these

460 experiments (daily at best with no duplicates due to the limited volume), and the imprecision of electrode-based pH_T

461 measurements relative to the SAMI-pH and spec-pH based measurements used in the large tank experiments.

462 No significant changes in salinity were recorded during these experiments as measured by a handheld salinometer
463 with a range of 30 - 31. Therefore, DIC and TA were not normalized to salinity. Temperature values ranged from 19
464 21 °C during the superimente.

464 - 21 °C during the experiments.

465 Similar to the large tank experiments, we used Henry's law and CO2SYS calculations to estimate the initial and 466 final equilibration condition of each aquaria experiment. The same average $pCO_{2,atm}$ of 421 ± 14 ppm was used with

467 the initial seawater temperature and salinity to estimate $pCO_{2,seawater}$ at the beginning of each experiment. The initial

- 468 equilibrium DIC was estimated from a CO2SYS calculation using this *p*CO_{2,seawater} and TA_i, which in all cases was
- 469 less than the initial DIC calculated from TA_i and $pH_{T,i}$ (by 16 36 µmol kg⁻¹). This indicates that the seawater was
- 470 not fully equilibrated with the atmosphere at the time when NaOH was added, likely due to respiration and

471 decomposition of organic material (Section 2.1), and as such, the aquaria should outgas CO₂. The final equilibrium

472 DIC was estimated from a CO2SYS calculation using the same $pCO_{2,seawater}$ and the TA measured just after the

- 473 NaOH addition. The percent equilibration for each experiment was then estimated between the measured and
- 474 predicted values for $\Delta DIC/\Delta TA$. Due to the air bubbling, most experiments approached equilibrium with the
- 475 atmosphere within 1-7 days, with the exception of the non-bubbled $pH_T 8.5$ experiment that took ~20 days. The
- 476 surface water of this non-bubbled experiment was stagnant, and the water was only mixed via stirring just before
 477 taking pH and TA samples. Absorption of atmospheric CO₂ began immediately after the NaOH addition, as
- 478 determined by decreasing pH_T. We note that there are significant uncertainties in these equilibrium estimates leading

479 to estimates of >100% equilibration. These estimates would be better constrained with more continuous carbonate

- 480 chemistry measurements, particularly seawater and atmosphere pCO_2 throughout the experiments that would allow
- for more direct calculation of air-sea CO_2 flux and equilibration, and finer control of bubbling and diffusion rates are
- 482 necessary to define the timeline for equilibration within the aquaria.

- 483 Each aquarium was gently stirred during the addition of NaOH to prevent and/or redissolve Mg(OH)₂ precipitation.
- 484 No CaCO₃ precipitation was observed in the tanks below a bulk seawater pH_T of 10.0, and TA remained stable in
- 485 each of these experiments with the exception of some increase driven by minor evaporation on the order of +2
- 486 umol/kg per day. Experiments where CaCO₃ precipitation was induced by increasing the starting pH to values above
- 487 10 are discussed in Section 3.3.



488 The aquaria experiments with target pH_T from 8.3 – 9.9 are summarized in Figure 4.

490 Figure 4: Results of 9 aquaria experiments in which enough NaOH was added to each aquaria to raise the bulk pH_T 491 to 8.3 - 9.9, pH_T decreased rapidly in all cases in which air bubbling sped equilibration with atmospheric CO₂. 492 Results include: (a) measured pH_T, (b) measured TA, (c) CO2SYS-calculated DIC, (d) CO2SYS-calculated pCO₂,

493 (e) the observed carbon uptake ratio (CAR) as $(DIC_{exp} - DIC_{baseline}) / \Delta TA_{NaOH addition}$ with horizontal dashed lines

- 494 representing the expected range of $0.7-0.9 \text{ mol CO}_2$ uptake / mol NaOH added to seawater, the change in (f) TA and
- 495 (g) DIC compared to the baseline measurements before the addition of NaOH, and the percent equilibration
- 496 estimated between the observed and theoretical CAR.

497 In general, the large tanks and aquaria showed reasonable agreement in achieving values for CAR within the

- 498 expected range of 0.7-0.9 (He and Tyka, 2023; Burt et al., 2021; Wang et al., 2023). While the use of aquaria
- 499 bubbled with air to speed equilibration allowed for a greater range of data collection within a constrained experiment 500 timeline, the quality of this data is limited by the lack of appropriate sensors to fit into these small 15 L aquaria and
- 501 fewer bottle samples due to the reduced quantity of seawater. However, while the large tanks allow for a larger
- 502 range of oceanographic sampling and sensing techniques, it is more challenging to fully quantify mixing and
- 503 circulation rates in the current large tank experimental setup.
- 504 Figure 5 shows the dependence of the equilibrium values of Δ DIC, CAR, and Δ pH_T = (pH_{final} – pH_{initial}) as a function 505 of the alkalinity addition for both tank and aquaria experiments in which the final percent equilibration for CO₂ was 506
- estimated at greater than 90%. Results for tank and aquaria experiments aligned well, with increasing Δ DIC for
- 507 increasing alkalinity additions. The CAR was observed for all experiments to fall within the range expected for 508
- seawater with the temperature and salinity values used in these tests. As expected from calculations of the response 509
- of the seawater carbonate buffer system to additions of alkalinity, the pH_T at equilibrium exceeded the initial pH_T 510 value prior to the addition of alkalinity. That is, even once equilibrium in the alkalinity enhanced experiment tank
- 511 had been reached, the ending pH value was slightly elevated relative to the starting pH value. This finding warrants

512 further investigation on the potential of OAE to mitigate some acidification impacts in controlled field trials by 513 metering the discharge of alkalinity to semi-protected water body.



514

Figure 5: (a) The change in final CO2SYS-predicted DIC relative to the initial conditions for tank, aquaria, and control experiments increases with increasing NaOH additions for cases where the air-sea CO_2 equilibration was estimated at >90% at the termination of each experiment. (b) CO2SYS-predicted CAR (Δ DIC / Alkalinity addition) at air-sea equilibrium conditions for tank, aquaria, and control experiments, with horizontal dashed lines

519 representing the expected range of 0.7-0.9 mol CO_2 uptake / mol NaOH added to seawater. (c) The measured ΔpH_T

alkalinity addition for both tank and aquaria experiments.

522 3.3 Experiments exceeding the CaCO₃ precipitation threshold

523 While $Mg(OH)_2$ precipitation occurs immediately upon introduction of concentrated (i.e., ~0.5 M) NaOH to still 524 seawater, it may be rapidly dissolved or avoided entirely by gentle mixing, including via the use of stirrers, 525 circulation pumps, or air bubblers. This precipitation and redissolution happened rapidly enough that it was not 526 identified in any TA or other variables measured in the aquaria and tank tests. However, in cases where enough 527 NaOH was added to raise the bulk seawater pH_T to greater than 10.0 (i.e. in one large tank test with a target pH_T of 528 10.3, and in 4 aquaria experiments ranging from pH_T 10.0-10.3), runaway precipitation of Mg(OH)₂ and CaCO₃ was 529 observed. This was characterized by a sharp reduction in both TA and DIC and an increase in turbidity. Runaway

530 precipitation has been described as a condition in which more alkalinity is removed from seawater by mineral

- 531 precipitation than was initially added until a new steady state is achieved (Moras et al., 2022; Hartmann et al., 2023;
- 532 Suitner et al., 2023). This can significantly impact the efficiency of OAE, and has implications for biological

533 productivity, as increased turbidity may impact photosynthesis or predator-prey interactions.

534 In both the tank and aquarium $pH_T 10.3$ cases, discrete samples of the precipitate were collected at seven different 535 times after the bulk pH_T value reached 10.3 (0h, 3h, 24h, 49h, 71h, 145h, 167h - see Fig. 6) for XRD and SEM 536 analysis. At each timepoint, 0.5 - 1 L seawater was collected from the tank sampling port or from the center of the 537 aquaria. In cases where precipitation had visibly settled at the bottom of the aquaria, this material was stirred into the 538 water column before sampling. We note that material that settled to the bottom of the large tanks was not directly 539 collected, and that only a subset of precipitation was collected at each time point, such that later timepoints may 540 include solids that had precipitated at the beginning of the experiment. The filtered seawater was immediately 541 analyzed for TA and pH via Ross electrode because the heightened pH was out of the range of spectrophotometric 542 methods. Bottle samples of filtered seawater were not able to be analyzed at NOAA PMEL due to the continued 543 precipitation of CaCO₃ after filtration and preservation. Both XRD and SEM results showed the dominance of 544 Mg(OH)₂ precipitation immediately after the alkalinity addition and the corresponding increase in pH and $\Omega_{aragonite}$ 545 (to a value of around 30). The $Mg(OH)_2$ precipitation at this stage was thick, slurry-like, and difficult to 546 appropriately rinse. Broad peaks associated with brucite in the 0 and 3 hr time points may reflect that these signals

547 were partially obscured by the presence of other salts, and a sharp peak in the 0 hr time point $\sim 27^{\circ} 2\theta$ is likely

 $^{520 \}qquad = (pH_{final} - pH_{initial}) \text{ increases with}$

- sociated with NaCl. Within hours of the NaOH addition, the runaway CaCO₃ precipitation began, characterized by
- 549 fine, light particulates in the water column and a sharp increase in turbidity. Within \sim 24 hours of the NaOH addition, 550 most Mg(OH)₂ signals had disappeared, leaving only aragonite and calcite peaks in the XRD. The results of the
- 551 XRD analysis for the tank experiment are summarized in Figure 7, and the aquarium experiment showed similar
- 552 results (see Supplementary Materials). TA decreased throughout the precipitation of Mg(OH)₂ and CaCO₃, and was
- below that of the initial seawater within 24 hours of the NaOH addition. In the tank experiment, the initial TA (2025
- 554 μmol/kg) was raised by 3305 μmol/kg. Within 3 days the TA had decreased to 1583 μmol/kg and continued to
- $\label{eq:constraint} 555 \qquad \text{decrease through the termination of the experiment to 1253 μmol/kg 10 days after the addition of NaOH. The DIC,}$
- 556 which was initially measured at 1938 μmol/kg, decreased to 720 μmol/kg by the end of the experiment. This
- speriment shows that runaway CaCO₃ can result in a significant loss of both efficiency of alkalinity dosing for
- 558 OAE projects and of storage of carbon in the form of DIC. A figure of time-series data collected during the tank
- 559 experiment is available in the Supplementary Materials.





- 570 In summary, the presence and duration of brucite precipitation upon addition of 0.5 M aqueous NaOH depends on
- the ratio of the NaOH addition rate to the local dilution rate in the receiving waters. Future research using flow
- through tanks could help identify thresholds below which brucite precipitation can be avoided or limited, and the
- 573 immediate formation of Mg(OH)₂ may be reversible, as also noted by Suitner et al. (2023) and Cyronak et al. 574 (2023). At the given initial seawater conditions, the threshold for aragonite precipitation began at an Ω_{arag} of 30,
- 574 (2025). At the given initial seawater conditions, the threshold for aragonite precipitation began at an Ω_{arag} of 50, 575 corresponding to pH_T > 10.0, and continued as Ω_{arag} decreased to ~5.2. This threshold corresponded to an increase in
- 576 TA of >2270 μ mol/kg. The potential for runaway aragonite precipitation may be reduced by active mixing at the
- 577 point of NaOH introduction, maintaining a mixing volume below bulk seawater pH_T of 10.0, and allowing for
- 578 appropriate dilution in flow-through conditions, particularly on timescales of hours after alkalinity addition.

579 Additional characterization of runaway precipitation thresholds at varying temperatures, salinities, and suspended 580 particulate conditions will allow for OAE implementation designs that best avoid this potential risk to OAE 581 efficiency and ecosystem perturbation. We note that these results are only valid for experiments that are open to the 582 atmosphere allowing for exchange of CO_2 across the air-sea boundary using an aqueous hydroxide feedstock for 583 alkalinity, and are not comparable to experiments such as closed bottle incubations, where sustained conditions of 584 high Ω_{arag} may result in precipitation at different thresholds. We also note that we do not assume zero aragonite 585 precipitation at conditions below the stated thresholds, but that potential precipitation is not readily detectable with 586 our experimental setup. For example, heterogeneous CaCO₃ precipitation events, such as may occur on suspended 587 sediments in the water column, have been suggested through characteristic changes in seawater TA/DIC ratios in 588 cases of riverine inputs and bottom sediment resuspension (Bustos-Serrano et al., 2009; Wurgaft et al., 2016; 2021). 589 Suspended sediments in the context of OAE project sites could influence OAE efficiency and the potential for 590 runaway precipitation and should be included in future studies (Bach, 2023). The thresholds determined in this study 591 are significantly higher than those of some mineral-based OAE studies, including precipitation after an increase in 592 TA of ~500 µmol/kg using CaO and Ca(OH)₂ mineral additions (Moras et al., 2022). Hartmann et al. (2022) noted 593 precipitation resulting from alkalinity additions of >600 µmol/kg Mg(OH)₂, and found that alkaline solutions 594 avoided carbonate precipitation better than reactive alkaline particle additions to seawater. Suitner et al. (2023) 595 suggested that alkalinity additions $> 2000 \mu mol/kg$ could be achievable given sufficient dilution to avoid runaway 596 precipitation. Together, these studies highlight the need to expand research into runaway precipitation to

- 597 characterize potential inefficiencies in OAE, particularly in in-situ experiments to establish relationships applicable
- to ocean environments.

599 5 Summary and Future Work

600 These results demonstrate that ocean alkalinity enhancement using aqueous sodium hydroxide in seawater results in 601 CO_2 removal from air at an efficiency of 0.75 (\pm 0.04) – 0.92 (\pm 0.10), with 90% equilibration typically achieved 602 within 7 - 9 weeks (still surface water with ~16 L/min subsurface circulation through UV arrays) to 3 - 5 weeks 603 (with the addition of ambient air bubbling into the bottom of each tank) of the initial addition when performed in 604 ~6000 L tanks with seawater-air contact areas of around 4.6 m². These results are in general agreement with ratios 605 noted in Burt et al. (2021), He and Tyka (2023), and Wang et al., (2023). Here, uncertainties are driven by sensor 606 precision and temporal resolution in discrete DIC and TA sampling, the limited number of experiments with 607 minimal opportunities for duplicates or replicates, and poorly constrained data on mixing, circulation, and air 608 bubbling rates. Ongoing experiments seek to improve each of these conditions and should particularly focus on 609 constraining the movement of water within a given tank to improve air-sea equilibration estimates and to allow for 610 better extrapolation from tank to field experiments. In addition, a focus of ongoing and future work is to provide rate 611 estimates for the uptake of atmospheric CO₂ in response to an NaOH addition, allowing for fitting and extrapolation

612 of a shortened experiment to equilibration with the atmosphere.

613 We relied on several methods to constrain seawater carbonate chemistry. The tank-scale experiments primarily 614 relied on discrete (at most daily) DIC and TA sampling (NOAA PMEL), paired with daily measurements from 615 spectrophotometric pH systems (SAMI-pH and a semi-automated benchtop spec-pH system following Carter et al. 616 (2013)) and local TA measurements. With appropriate calibration or correction of the spec-pH systems relative to 617 CRM, there was no significant difference in carbonate calculations using the NOAA PMEL DIC-TA or spec-pH-618 local TA pairings, though the latter case typically produced larger uncertainties. Aquaria experiments relied on a 619 standard glass pH electrode (at most daily, corrected to spectrophotometric systems) with discrete (at most daily) TA 620 measurements, which provided reasonable data relative to the tank experiments. As a result, ongoing tank-scale 621 experiments have limited the volume of discrete DIC and TA samples collected for analysis at NOAA PMEL to 622 allow for faster and less expensive monitoring via spec-pH and local TA titrations. However, we note that the major 623 limitation in this measurement pathway lies in the spec-pH method, which is typically limited to pH_T measurements 624 ranging from 7-9 for the meta-cresol purple indicator dye used. While our measurements retained some sensitivity

- 625 up to pH_T 9.5, such a method should typically be considered unreliable at these pH_T values, and we relied on
- 626 frequent correction to CRM and comparison with DIC/TA samples. Thymol blue is an alternative
- 627 spectrophotometric pH_T indicator dye with sensitivity over the higher pH_T conditions observed during these initial
- trials and will be assessed for future experiments (Zhang and Byrne, 1996; Liu et al., 2006).

629 Aqueous NaOH with concentrations as high as 0.5 M can be added directly to turbulent seawater with only limited 630 observable precipitation of Mg(OH)₂. In these conditions this precipitated mineral rapidly redissolves on the 631 timescales of minutes to seconds. Improved control over the NaOH dosing rate (in our tank experiments, ~50 mL 632 NaOH/min) and the turbulence of the receiving water through metered flow through experiments will be valuable in 633 extrapolating to field conditions. This precipitation is detectable both visually and through turbidity measurements 634 and implies that straightforward measurement of pH and turbidity at the dispersal site can be used to adjust the 635 alkalinity dispersal rate according to local mixing conditions such that Mg(OH)₂ precipitation is avoided and/or 636 redissolves when it occurs. No significant CaCO₃ precipitation was observed at pH <10.0 or $\Omega_{aragonite}$ < 30.0. 637 Runaway CaCO₃ precipitation was observed above these thresholds, where a massive precipitation and settling of 638 $Mg(OH)_2$ and CaCO₃ solids results in less alkalinity in the overlying water than at the starting condition. pH and 639 turbidity sensing combined with discrete TA measurements could be used as a feedback signal for alkalinity dosing 640 into seawater to ensure that the local maximum thresholds at the dispersal location do not approach or exceed 641 conditions that promote significant CaCO₃ precipitation. We note that future investigations seeking to better 642 approximate field conditions should take into account seasonal and tidal shifts in temperature and salinity, and 643 varying conditions of suspended sediment in the water column, including that of aerial dust, terrestrial runoff, and 644 resuspended bottom sediments.

- 645 In these experiments, the seawater was filtered and bleach treated prior to experiments to limit biological growth,
- and both tank and aquaria experiments were conducted indoors with limited light. Nevertheless, in most
- 647 experiments, biological growth was observed after a few weeks, including cyanobacteria and coccolithophores. A
- $648 \qquad \text{series of experiments are underway to test the difference in CO_2 removal efficiency for two side-by-side tanks, both}$
- 649 of which are dosed with NaOH, but only one of which was bleached. Preliminary results show minimal difference
- between the bleached and unbleached tanks, indicating these experiments are applicable to real-world conditions, at
- least for regions with biological communities similar to that of Long Island Sound, but further investigation is
- warranted.
- 653 A focus of future work is to consider the potential impact of the SEAMATE process on local ocean acidification 654 mitigation efforts. We note that in each constrained tank and aquarium experiment, the pH_T at equilibrium exceeds
- 654 mitigation efforts. We note that in each constrained tank and aquarium experiment, the pH_T at equilibrium exceeds 655 the initial pH_T value prior to the addition of alkalinity. A controlled release of alkalinity could theoretically be
- 655 the initial pH_T value prior to the addition of alkalinity. A controlled release of alkalinity could theoretically be 656 configured to maintain a locally elevated pH_T value relative to pre-alkaline conditions, with potential uses in
- aquaculture and hatchery environments.
- 658 These results provide clear and practical guidelines for MRV for OAE implementations employing aqueous 659 alkalinity. First, carbonate chemistry and turbidity measurements at the alkalinity dispersal location can ensure that 660 seawater parameters such as pH and $\Omega_{aragonite}$ remain within pre-determined safe bounds and that unwanted 661 precipitation is avoided. Second, for a given OAE deployment, where ocean models provide a reasonable certainty 662 about the fraction of the alkalinity plume remaining in the surface over weeks to months, the CO₂ removal efficiency 663 and timescale for air-seawater equilibration provided by our experiments can place a lower bound on the amount of 664 CO₂ removal expected from that OAE intervention. Expanding these studies from tank scale to mesocosm and field 665 experiments will be crucial to understanding biological impacts and constraining realistic air-sea interactions in 666 response to this type of OAE (Oschlies et al., 2023).

667 Data availability

- 668 Data are described in the manuscript and provided Supplementary Materials, which includes a .csv file with
- 669 processed sensor and sample time-series data at hourly resolution.

670 Author contribution

- 671 MDE and BRC designed the experiments and MCR carried them out with support from NH, CS, and XL. JH
- provided support on experimental setup and instrumentation. MCR prepared the manuscript with contributions fromall co-authors.

674 Competing interests

675 MCR is Lead Oceanographer and Head of MRV at Ebb Carbon, Inc. MDE is Co-Founder and Chief Scientific676 Advisor at Ebb Carbon, Inc.

677 Acknowledgements

- 678 We would like to than Stephen Abrams and Thomas Wilson at Stony Brook University Flax Pond Marine Lab for
- technical assistance in experiment setup. We thank Chris Ikeda and Susan Curless of NOAA PMEL for support in
- discrete sample analysis. We thank Mike Tyka for productive discussions. We thank Eyal Wurgaft for assistance in
- **681** TA titrations.

682 Funding

- 683 We acknowledge funding from The Grantham Foundation for the Protection of the Environment under the SEA
- 684 MATE (Safe Elevation of Alkalinity for the Mitigation of Acidification Through Electrochemistry) grant. In
- addition, this research used the <u>XRD</u> facility of the Center for Functional Nanomaterials (CFN), which is a U.S.
- 686 Department of Energy Office of Science User Facility, at Brookhaven National Laboratory under Contract No. DE-
- 687 SC0012704. BRC and JH were funded through the Cooperative Institute for Climate, Ocean, and Ecosystem
- 688 Studies (CICOES) under NOAA Cooperative Agreement NA200AR4320271 and supported by NOAA's PMEL.

689 References 690

- Albright, R., Caldeira, L., Hosfelt, J., Kwiatkowski. L., Maclaren, J.K., Mason, B.M., Nebuchina, Y. et al.: Reversal of ocean acidification enhances net coral reef calcification. Nature, 531, no. 7594: 362-365, 2016.
- **693** Bach, L.T.: The additionality problem of ocean alkalinity enhancement. Biogeosciences, 21, 261-277, 2024.
- Bach, L. T., Gill, S.J., Rickaby, R.E.M., Gore, S., and Renforth, P.: CO₂ removal with enhanced weathering and
 ocean alkalinity enhancement: potential risks and co-benefits for marine pelagic ecosystems. Frontiers in
 Climate, 1, 7, 2019.
- Bainbridge, Z.,Lewis, S., Bartley, R., Fabricius, K., Collier, C., Waterhouse, J., Garzon-Garcia, A., Robson, B.,
 Burton, J., Wenger, A., and Brodie, J: Fine sediment and particulate organic matter: A review and case study on
 ridge-to-reef transport, transformations, fates, and impacts on marine ecosystems. Marine Pollution Bulletin
 135, pp. 1205-1220. 2018
- Berner, R. A., Lasaga, A.C., and Garrels, R.M.: Carbonate-silicate geochemical cycle and its effect on atmospheric carbon dioxide over the past 100 million years. Am. J. Sci.;(United States) 283, no. 7, 1983.
- Boettcher, M., Chai, F. Cullen, J., Goeschl, T., Lampitt, R., Lenton, A., Oschlies, A. et al.: High level review of a
 wide range of proposed marine geoengineering techniques. GESAMP Working Group Reports and Studies, 41,
 2019.
- Broderson, K.E., Hammer, K.J., Schrameyer, V., Floytrup, A., Rasheed, M.A., Ralph, P.J., Kühl, M., and Pederson,
 O.: Sediment resuspension and deposition on seagrass leaves impedes internal plant aeration and promotes
 phytotoxic Hh2S intrusion. Frontiers in Plant Science 8. 2017
- Burt, D.J., Fröb, F., & Ilyina, T.: The sensitivity of the marine carbonate system to regional ocean alkalinity
 enhancement. Frontiers in Climate 3, 624075. 2021

- Bustos-Serrano, H., Morse, J.W., & Millero, F.J.: The formation of whitings on the Little Bahama Bank. Marine
 Chemistry 113, no. 1-2, pp. 1-8. 2009
- Butenschön, M., Lovato, T., Masina, S., Caserini, S., and Grosso, M.: Alkalinization scenarios in the Mediterranean
 Sea for efficient removal of atmospheric CO2 and the mitigation of ocean acidification. Frontiers in Climate 3,
 614537. 2021
- Carter, B. R., J. A. Radich, H. L. Doyle, and A. G. Dickson.: An automated system for spectrophotometric seawater
 pH measurements. Limnology and Oceanography: Methods, 11, no. 1: 16-27, 2013.
- Caserini, S., Storni, N., & Grosso, M.: The availability of limestone and other raw materials for ocean alkalinity
 enhancement. Global Biogeochemical Cycles, 36, e2021GB007246. https://doi. org/10.1029/2021GB007246,
 2022.
- 721 Cross, J.N., Sweeney, C., Jewett, E.B., Feely, R.A., McElhany, P., Carter, B., Stein, T., Kitch, G.D., and Gledhill,
 722 D..: Strategy for NOAA carbon dioxide removal research: A white paper documenting a potential NOAA CDR
 723 science strategy as an element of NOAA's Climate Interventions Portfolio. NOAA Special Report. NOAA,
 724 Washington, DC. DOI: 10.25923/gzke-8730. 2023Cyronak, T., Albright, R., and Bach, L.: Field experiments in
 725 ocean alkalinity enhancement research. *State of the Planet*, 2-oae2023 (7). 2023.
- de Lannoy, C.-F., Eisaman, M.D., Jose, A., Karnitz, S.D., DeVaul, R.W., Hannun, K., and Rivest, J.L.B.: Indirect
 ocean capture of atmospheric CO₂: Part I. Prototype of a negative emissions technology. International journal of
 greenhouse gas control, 70: 243-253, 2018.
- Dickson, A. G.: An exact definition of total alkalinity and a procedure for the estimation of alkalinity and total
 inorganic carbon from titration data. Deep Sea Research Part A. Oceanographic Research Papers, 28(6), 609–623, 1981.
- 733 Dickson, A. G.: The development of the alkalinity concept in marine chemistry. Marine Chemistry, 40(1–2), 49–63, 1992.
- Dickson, A.G.: Thermodynamics of the dissociation of boric acid in synthetic seawater from 273.15 to 318.15 K.
 Deep Sea Research Part A. Oceanographic Research Papers, 37, no. 5: 755-766, 1990.
- 737 Dickson, A.G., Sabine, C.L., and Christian, J.R.: Guide to best practices for ocean CO₂ measurements. North Pacific
 738 Marine Science Organization, 2007.Eisaman, M. D., Parajuly, K., Tuganov, A., Eldershaw, C., Chang, N.,
 739 Littau, K. A. CO₂ Extraction from Seawater Using Bipolar Membrane Electrodialysis, Energy Environ. Sci., 5:
 740 7346. https://doi.org/10.1039/c2ee03393c, 2012.
- Fisaman, M. D.; Rivest, J. L. B.; Karnitz, S. D.; De Lannoy, C.-F.; Jose, A.; DeVaul, R. W.; Hannun, K. Indirect
 Ocean Capture of Atmospheric CO₂: Part II. Understanding the Cost of Negative Emissions. International
 Journal of Greenhouse Gas Control, 70: 254–261, https://doi.org/10.1016/j.ijggc.2018.02.020, 2018.
- Eisaman, M. D., Geilert, S., Renforth, P., Bastianini, L., Campbell, J., Dale, A. W., Foteinis, S., Grasse, P., Hawrot,
 O., Löscher, C. R., Rau, G. H., and Rønning, J.: Assessing the technical aspects of ocean-alkalinityenhancement approaches, in: Guide to Best Practices in Ocean Alkalinity Enhancement Research, edited by:
 Oschlies, A., Stevenson, A., Bach, L. T., Fennel, K., Rickaby, R. E. M., Satterfield, T., Webb, R., and Gattuso,
 J.-P., Copernicus Publications, State Planet, 2-oae2023, 3, <u>https://doi.org/10.5194/sp-2-oae2023-3-2023</u>, 2023.
- Feely, R.A., Alin, S., Carter, B., Bednaršek, N., Hales, B., Chan, F., Hill, T.M., Gaylord, B., Sanford, E., Byrne,
 R.H., Sabine, C.L., Greeley, D., Juranek, L., Chemical and biological impacts of ocean acidification along the
 west coast of North America, Estuarine, Coastal and Shelf Science, doi: 10.1016/j.ecss.2016.08.043, 2016.
- west coast of North America, Estuarine, Coastal and Shelf Science, doi: 10.1016/j.ecss.2016.08.043, 2016.
 Feng, E. Y., Koeve, W., Keller, D.P., and Oschlies, A.: Model-Based Assessment of the CO₂ Sequestration Potential of Coastal Ocean Alkalinization. Earth's Future, 5, no. 12: 1252-1266, 2017.
- Fennel, K., Long, M.C., Algar, C., Carter, B., Keller, D., Laurent, A., Mattern, J.P., Musgrave, R., Oschlies, A.,
 Ostiguy, J., Palter, J.B., and Whitt, D.B.: Modeling considerations for research on Ocean Alkalinity
 Enhancement (OAE). *State of the Planet*, 2-oae2023 (9). 2023. Ferderer, A., Chase, Z., Kennedy, F., Schulz,
 K.G., and Bach, L.T.: Assessing the influence of ocean alkalinity enhancement on a coastal phytoplankton
 community. Biogeosciences 19, no. 23: 5375-5399, 2022.
- Friis, K.; Körtzinger, A.; Wallace, D. W. R. The Salinity Normalization of Marine Inorganic Carbon Chemistry Data. Geophys. Res. Lett., 30 (2). <u>https://doi.org/10.1029/2002GL015898</u>, 2003.
- Groen, A., Kittu, L., Ortiz Cortes, J., Schulz, K., and Riebesell, U.: Assessing the response of particulate organic
 matter stoichiometry to ocean alkalinisation. Ocean Visions Summit, Atlanta, Georgia, USA,
 2023ocvi.conf27171G. 4-6 April 2023
- Hartmann, J., Suitner, N., Lim, C., Schneider, J., Marín-Samper, L., Arístegui, J., Renforth, P., Taucher, J., and Riebesell, U.: Stability of alkalinity in ocean alkalinity enhancement (OAE) approaches–consequences for durability of CO₂ storage. Biogeosciences 20, no. 4: 781-802, 2023.

- Harvey, L.: Mitigating the atmospheric CO₂ increase and ocean acidification by adding limestone powder to upwelling regions, Journal of 640 Geophysical Research: Oceans, 113, 2008.
- He, J. and Tyka, M. D.: Limits and CO₂ equilibration of near-coast alkalinity enhancement, Biogeosciences, 20, 27–
 43, https://doi.org/10.5194/bg-20-27-2023, 2023.
- Ho, D. T., Bopp, L., Palter, J.B., Long, M.C., Boyd, P.W., Neukermans, G., and Bach, L.T.: Monitoring, reporting, and verification for ocean alkalinity enhancement. *State of the Planet*, 2-oae2023 (12). 2023. Ilyina, T., Wolf-Gladrow, D., Munhoven, G., and Heinze, C.: Assessing the potential of calcium-based artificial ocean alkalinization to mitigate rising atmospheric CO₂ and ocean acidification, Geophysical Research Letters, 40, 5909-5914, 2013.
- IPCC: Summary for Policymakers. In: Climate Change 2021: The Physical Science Basis, Contribution of Working
 Group I to the Sixth Assessment Report of the Intergovernmental Panel on Climate Change, edited by: MassonDelmotte, V., Zhai, P., Pirani, A., Connors, S. L., Péan, C., Berger, S., Caud, N., Chen, Y., Goldfarb, L., Gomis,
 M. I., Huang, M., Leitzell, K., Lonnoy, E., Matthews, J. B. R., Maycock, T. K., Waterfield, T., Yelekçi, O., Yu,
 R., and Zhou, B.: Cambridge University Press, Cambridge, United Kingdom and New York, NY, USA, 3–32,
 https://doi.org/10.1017/9781009157896.001, 2022.
- Isson, T. T., Planavsky, N. J., Coogan, L. A., Stewart, E. M., Ague, J. J., Bolton, E. W., et al.: Evolution of the global carbon cycle and climate regulation on earth. Global Biogeochemical Cycles, 34, e2018GB006061. https://doi.org/10.1029/2018GB006061, 2020.
- Johnson, K.M., King, A.E., and Sieburth, J.M.: Coulometric TCO2 analyses for marine studies: An introduction.
 Marine Chemistry 16, pp. 61-82. 1985.
- Jones, D.C., Ito, T., Takano, Y., and C.-W Hsu, C.-W.: Spatial and seasonal variability of the air-sea equilibration timescale of carbon dioxide. Global Biogeochemical Cycles, 28(11), 1163–1178, https://doi.org/10.1002/2014GB004813, 2014.
- 790 Kheshgi, H. S.: Sequestering atmospheric carbon dioxide by increasing ocean alkalinity, Energy, 20, 915-922, 1995.
- Köhler, P., Hartmann, J., and Wolf-Gladrow, D.A.: Geoengineering potential of artificially enhanced silicate
 weathering of olivine. Proceedings of the National Academy of Sciences 107, no. 47: 20228-20233, 2010.
- La Plante, E., Chen, X., Bustillos, S., Bouissonnie, A., Traynor, T., Jassby, D., Corsini, L., Simonetti, D., and Sant,
 G.: Electrolytic seawater mineralization and the mass balances that demonstrate carbon dioxide removal. ACS
 EST Engg. <u>https://doi.org/10.1021/acsestengg.3c00004</u>, 2023.
- Lee, K., Kim, T.-W., Byrne, R.H., Millero, F.J., Feely, R.A., and Liu, Y.-M.: The universal ratio of boron to
 chlorinity for the North Pacific and North Atlantic oceans. Geochimica et Cosmochimica Acta 74, no. 6: 18011811, 2010.
- Lewis, E., Wallace, D., & Allison, L. J.: Program developed for CO₂ system calculations.
 <u>https://doi.org/10.2172/639712</u>, 1998.
- Liu, X., Wang, Z.A., Byrne, R.H., Kaltenbacher, E.A., and Bernstein, R.E.: Spectrophotometric measurements of pH in-situ: laboratory and field evaluations of instrumental performance. Environmental Science & Technology 40, no. 16, 5026-5044. 2006
- Lu, X., Ringham, M., Hirtle, N., Hillis, K., Shaw, C., Herndon, J., Carter, B.R., and Eisaman, M.D.:
 Characterization of an Electrochemical Approach to Ocean Alkalinity Enhancement. In AGU Fall Meeting
 Abstracts, vol. 2022, pp. GC31C-01. 2022.
- Lueker, T.J., Dickson, A.G., and Keeling, C.D.: Ocean pCO₂ calculated from dissolved inorganic carbon, alkalinity, and equations for K1 and K2: validation based on laboratory measurements of CO₂ in gas and seawater at equilibrium. Marine chemistry 70, no. 1-3: 105-119, 2000.
- Minx, J.C., Lamb, W.F., Callaghan, M.W., Fuss, S., Hilaire, J., Creutzig, F., Amann, T., et al.: Negative
 emissions—Part 1: Research landscape and synthesis. Environmental Research Letters 13, no. 6: 063001, 2018.
- Montserrat, F., Renforth, P., Hartmann, J., Leermakers, M., Knops, P., and Meysman, F.J.R.: Olivine dissolution in seawater: implications for CO₂ sequestration through enhanced weathering in coastal environments.
 Environmental Science & Technology 51, no. 7: 3960-3972, 2017.
- Moras, C.A., Bach, L.T., Cyronak, T., Joannes-Boyau, R., and Schulz., K.G.: Ocean alkalinity enhancement–
 avoiding runaway CaCO₃ precipitation during quick and hydrated lime dissolution. Biogeosciences 19, no. 15:
 3537-3557, 2022.
- 818 National Academies of Sciences, Engineering, and Medicine. A research strategy for ocean-based carbon dioxide
 819 removal and sequestration. 2021.
- National Academies of Sciences, Engineering, and Medicine. Negative Emissions Technologies and Reliable
 Sequestration: A Research Agenda. 2018.

- 822 Nduagu, E. "Production of Mg(OH)₂ from Mg-silicate rock for CO₂ mineral sequestration. Dissertation for Abo
 823 Akademi University, 2012.Oschlies, A., Bach, L., Rickaby, R., Satterfield, T., Webb, R. M., and Gattuso, J.-P.:
 824 Climate targets, carbon dioxide removal and the potential role of Ocean Alkalinity Enhancement. *State of the*825 *Planet*, 2-oae2023 (1). 2023.
- Pierrot, D., Lewis, E., and Wallace, D.W.R.: MS Excel program developed for CO₂ system calculations.
 ORNL/CDIAC-105a. Carbon Dioxide Information Analysis Center, Oak Ridge National Laboratory, U.S.
 Department of Energy, Oak Ridge, Tennessee, 2006.
- Rau, G.H.: Electrochemical splitting of calcium carbonate to increase solution alkalinity: Implications for mitigation
 of carbon dioxide and ocean acidity. Environmental science & technology 42, no. 23: 8935-8940, 2008.
- Renforth, P., and Henderson, G.: Assessing ocean alkalinity for carbon sequestration. Reviews of Geophysics 55, no. 3: 636-674, 2017.
- Rigopoulos, I., Harrison, A.L., Delimitis, A., Ioannou, I., Efstathiou, A.M., Kyratsi, T., and Oelkers, E.H. : Carbon
 sequestration via enhanced weathering of peridotites and basalts in seawater. Applied Geochemistry 91: 197 207, 2018.
- Rueda, O., Mogollón, J.M., Tukker, A., and Scherer, L.: Negative-emissions technology portfolios to meet the 1.5°
 C target. Global Environmental Change 67: 102238, 2021.
- Rogelj, J., Popp, A., Calvin, K. V., Luderer, G., Emmerling, J., Gernaat, D., Fujimori, S., Strefler, J., Hasegawa, T.,
 Marangoni, G., Krey, V., Kriegler, E., Riahi, K., van Vuuren, D. P., Doelman, J., Drouet, L., Edmonds, J.,
 Fricko, O., Harmsen, M., Havlík, P., Humpenöder, F., Stehfest, E., and Tavoni, M.: Scenarios towards limiting
 global mean temperature increase below 1.5 °C, Nat. Clim. Change, 8, 325–332, https://doi.org/10.1038/s41558018-0091-3, 2018.
- Schulz, K. G., Bach, L. T., and Dickson, A. G.: Seawater carbonate system considerations for ocean alkalinity
 enhancement research: theory, measurements, and calculations. *State of the Planet*, 2-oae2023 (2). 2023...
- Shaw, C., Ringham, M.C., Lu, X., Carter, B.R., Eisaman, M.D., and Tyka, M.: Understanding the Kinetics of
 Electrochemically derived Magnesium Hydroxide for Ocean Alkalinity Enhancement. In AGU Fall Meeting
 Abstracts, vol. 2022, pp. GC32I-0713. 2022.
- Song, S., Wang, Z.A., Gonneea, M.E., Kroeger, K.D., Chu, S.N., Li, D., and Liang, H.: An important
 biogeochemical link between organic and inorganic carbon cycling: Effects of organic alkalinity on carbonate
 chemistry in coastal waters influenced by intertidal salt marshes. Geochimica et Cosmochimica Acta 275:123139, 2020.
- 852 Suitner, N., Faucher, G., Lim, C., Schneider, J., Moras, C.A., Riebesell, U., and Hartmann, J.: Ocean alkalinity
 853 enhancement approaches and the predictability of runaway precipitation processes- Results of an experimental
 854 study to determine critical alkalinity ranges for safe and sustainable application scenarios. EGUsphere
 855 [preprint], https://doi.org/10.5194/egusphere-20223-2611, 2023
- Tyka, M.D., Arsdale, C.V., and Platt, J.C.: CO₂ capture by pumping surface acidity to the deep ocean. Energy &
 Environmental Science 15, no. 2: 786-798, 2022.
- 858 Van Heuven, S., Pierrot, D., Rae, J., Lewis, E., & Wallace, D.: MATLAB program developed for CO₂ system
 859 calculations. ORNL/CDIAC-105b, 530, 2011.
- Vitillo, J. G., Eisaman, M.D., Aradóttir, E.S.P., Passarini, F., Wang, T., and Sheehan, S.W.: The role of carbon capture, utilization and storage for economic pathways that limit global warming to below 1.5° C." Iscience: 104237, 2022.
- 863 Wang, H., Pilcher, D. J., Kearney, K. A., Cross, J. N., Shugart, O. M., Eisaman, M. D., & Carter, B. R.: Simulated
- impact of ocean alkalinity enhancement on atmospheric CO2 removal in the Bering Sea. Earth's Future, 11(1),
 e2022EF002816, 2023
- Wang, Z. A. and Cai, W. J.: Carbon dioxide degassing and inorganic carbon export from a marsh-dominated estuary (the Duplin River): A marsh CO₂ pump. Limnol. Oceanogr. 49, 341–354, 2004.
- Wolf-Gladrow, D. A., Zeebe, R. E., Klaas, C., Körtzinger, A., & Dickson, A. G.: Total alkalinity: The explicit conservative expression and its application to biogeochemical processes. Marine Chemistry, 106(1–2), 287–300, 2007.
- Wurgaft, E., Steiner, Z., Luz, B., and Lazar, B.: Evidence for inorganic precipitation of CaCO3 on suspended solids
 in the open water of the Red Sea, Marine Chemistry, 186, pp. 145–155, 2016.
- Wurgaft, E., Wang, Z., Churchill, J., Dellapenna, T., Song, S., Du,J., Ringham, M., Rivlin, T., and Lazar, B.:
 Particle triggered reactions as an important mechanism of alkalinity and inorganic carbon removal in river
- 876 plumes, Geophysical Research. Letters, 48, e2021GL093178, https://doi.org/10.1029/2021GL093178, 2021

- 877 878 879 Zeebe, R.E., and Wolf-Gladrow, D.: CO2 in seawater: equilibrium, kinetics, isotopes. Vol. 65, Gulf Professional Publishing. 2001.
- Zhang, H., and Byrne, R.H.: Spectrophotometric pH measurements of surface seawater at in-situ conditions:
- 880 absorbance and protonation behavior of thymol blue. Marine Chemistry 52, no. 1, pp. 17-25. 1996