



Integrating High-Resolution Coastal Acidification Monitoring Data Across Seven United States Estuaries

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Beginning in 2015, the United States Environmental Protection Agency's (EPA's) National Estuary Program (NEP) started a collaboration with partners in seven estuaries along the East Coast (Barnegat Bay; Casco Bay), West Coast (Santa Monica Bay; San Francisco Bay; Tillamook Bay), and the Gulf of Mexico (GOM) Coast (Tampa Bay; Mission-Aransas Estuary) of the United States to expand the use of autonomous monitoring of partial pressure of carbon dioxide (pCO₂) and pH. Analysis of high-frequency (hourly to sub-hourly) coastal acidification data including pCO2, pH, temperature, salinity, and dissolved oxygen (DO) indicate that the sensors effectively captured key parameter measurements under challenging environmental conditions, allowing for an initial characterization of daily to seasonal trends in carbonate chemistry across a range of estuarine settings. Multi-year monitoring showed that across all water bodies temperature and pCO_2 covaried, suggesting that pCO_2 variability was governed, in part, by seasonal temperature changes with average pCO₂ being lower in cooler, winter months and higher in warmer, summer months. Furthermore, the timing of seasonal shifts towards increasing (or decreasing) pCO2 varied by location and appears to be related to regional climate conditions. Specifically, pCO2 increases began earlier in the year in warmer water, lower latitude water bodies in the GOM (Tampa Bay; Mission-Aransas Estuary) as compared with cooler water, higher latitude water bodies in the northeast (Barnegat Bay; Casco Bay), and upwelling-influenced West Coast water bodies (Tillamook Bay; Santa Monica Bay; San Francisco Bay). Results suggest that both thermal and non-thermal influences are important drivers of pCO₂ in Tampa Bay

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and Mission-Aransas Estuary. Conversely, non-thermal processes, most notably the biogeochemical structure of coastal upwelling, appear to be largely responsible for the observed pCO_2 values in West Coast water bodies. The co-occurrence of high salinity, high pCO_2 , low DO, and low temperature water in Santa Monica Bay and San Francisco Bay characterize the coastal upwelling paradigm that is also evident in Tillamook Bay when upwelling dominates freshwater runoff and local processes. These data demonstrate that high-quality carbonate chemistry observations can be recorded from estuarine environments using autonomous sensors originally designed for open-ocean settings.

Keywords: coastal acidification, ocean acidfication, estuary, autonomous sensor, carbon dioxide, pH, dissolved oxygen, National Estuary Program

INTRODUCTION

An increase in the anthropogenic concentration of carbon dioxide (CO₂) dissolved in marine waters is putting stress on marine systems. This process, known as ocean acidification, refers to the changes in carbonate chemistry to the ocean that result from the absorption of increasing atmospheric CO₂, primarily from human fossil fuel combustion (Doney et al., 2009). The chemical effects of CO₂ absorption (e.g., reduced pH) are naturally buffered by the ocean's carbonate system via reaction with carbonate ions (CO₃²⁻) and bicarbonate ions (HCO₃⁻). With the continued uptake of CO₂ by the oceans, this buffering results in an increase in the partial pressure of CO₂ in water (pCO_2) and a decrease in pH, dissolved CO₃²⁻ and HCO₃⁻ concentrations, and associated carbonate mineral saturation states.

In coastal and estuarine environments, understanding carbonate chemistry is further complicated by the interaction and dynamism of multiple co-occurring chemical, biological, and physical processes, relative to the open ocean, operating at various spatial and temporal scales (Hofmann et al., 2011; Waldbusser and Salisbury, 2014). Coastal and estuarine systems are susceptible to local and regional acidification due to eutrophication, air-water flux of CO₂, coastal upwelling, changes in freshwater inflow, stratification, and other factors. Changes to the carbonate chemistry of marine waters can adversely affect the ability of shellfish and other calcifying organisms to build or maintain their calcium carbonate (CaCO₃) shells and skeletons (Talmage and Gobler, 2010; Gazeau et al., 2013; Waldbusser et al., 2015; Feely et al., 2016). Changes in pCO_2 and pH have also been shown to impact other physiological processes, including species growth, survival, fertilization, embryonic/larval development, and behavior (Fabry et al., 2008; Pörtner, 2008; Doney et al., 2009; Kroeker et al., 2013; Gledhill et al., 2015). Impairments to fish and shellfish physiology can lead to adverse impacts to the ecology of marine and estuarine systems. The effects of ocean and coastal acidification are already being seen in fish and shellfish aquaculture across the country (e.g., Barton et al., 2015; Mabardy et al., 2015), potentially threatening an important industry in many coastal communities (Barton et al., 2012; Ekstrom et al., 2015; Clements and Chopin, 2017).

High-resolution, multiparameter monitoring via autonomous sensors is important for characterizing the carbonate chemistry

of estuarine waters and distinguishing the drivers of coastal acidification. These monitoring data can shed light on the vulnerability of these systems to acidification and guide specific mitigation and adaptation strategies such as using seagrass to decrease pCO₂, improving aquaculture techniques to buffer hatchery systems, and adapting nutrient management plans (Kelly et al., 2011; Strong et al., 2014; Chan et al., 2016). Autonomous sensors have been used extensively in the open ocean to monitor ocean acidification; however, their deployment in coastal and estuarine waters is challenging due to rapid variation over large ranges in temperature, salinity and chemical composition, biofouling, sensor drift, and other factors (Sastri et al., 2019). The United States Environmental Protection Agency's (EPA's) National Estuary Program (NEP)¹ has expanded the use of autonomous carbonate chemistry sensors in estuarine environments. The NEP is a place-based program to protect and restore the water quality and ecological integrity of estuaries of national significance. The 28 estuaries that make up the NEP are located along the Atlantic, Gulf, and Pacific coasts, and in Puerto Rico. The programs are in a variety of institutional settings, including federal, state, and local agencies, universities, and individual nonprofit organizations. Beginning in 2015, EPA funded nine NEPs to purchase autonomous pCO_2 and pH sensors to characterize carbonate chemistry conditions and form a more mechanistic understanding of coastal acidification in their estuaries (Figure 1). EPA's Office of Research and Development (ORD) Pacific Ecological Systems Division conducted the monitoring in Tillamook Bay. Ten NEPs and their partners have been collecting hourly and sub-hourly in situ pCO₂ and pH data and have worked to optimize monitoring methods and data analysis procedures (EPA, 2021). Data from seven of these NEPs are discussed here.

The objective of this study is to describe and compare coastal acidification data from seven coastal systems around the United States in terms of typical values (e.g., median, range), seasonality and co-variability to begin to understand natural variability in carbonate chemistry parameters within and across a range of environmental settings. Continued monitoring and collection of long-term, high-resolution data are needed to detect real trends in coastal acidification, distinguish these from

¹https://www.epa.gov/nep



background variability, and determine the relative influence of the drivers and impact of acidification in these different systems.

NEP COASTAL ACIDIFICATION MONITORING PROGRAM

Water Body Characteristics

The seven NEPs and their partners conducting coastal acidification monitoring that are discussed in this report include two NEPs on the East Coast (Casco Bay Estuary Partnership, ME, United States; Barnegat Bay Partnership, NJ, United States), two NEPs in the Gulf of Mexico [GOM (Tampa Bay Estuary Program, FL, United States; Coastal Bend Bays and Estuaries Program/Mission-Aransas Estuary, TX, United States)] and three NEPs on the West Coast (San Francisco Bay Estuary Partnership, CA, United States; Santa Monica Bay NEP, CA, United States; Tillamook Estuaries Partnership, OR, United States; Figures 1, 2). The NEPs of Long Island Sound (Long Island Sound Study), Massachusetts Bay (Massachusetts Bay NEP) and Mobile Bay (Mobile Bay NEP) also conduct coastal acidification monitoring with autonomous pCO₂ and pH sensors; however, data are unavailable, and therefore, are not the subject of this study. In Massachusetts Bay and Mobile Bay, sensors had yet to be deployed prior to beginning this analysis so data was unavailable for inclusion. In Long Island Sound, researchers were already conducting an in-depth analysis of their monitoring data prior to initiation of this report and wanted to complete their analysis before making the data available for inclusion in a multi-estuary synthesis. The water bodies studied vary in geographic location, size, environmental stressors,

coastal dynamics and processes, and local economic interests (**Supplementary Table 1**). Watershed size ranges from 1,428 km² (Tillamook Bay) to 5,698 km² (Tampa Bay). Population size ranges from \sim 7,500 individuals in the Tillamook Bay watershed to \sim 7,000,000 individuals, in the San Francisco Bay watershed, and land use ranges from more urbanized (Santa Monica Bay, Tampa Bay) to more undeveloped and forested (Tillamook Bay). Santa Monica Bay has the greatest average water depth at 95 meters (m), while Tillamook Bay has an average depth of \sim 1.4 m. Tidal height ranges from 0.15 m in Barnegat Bay to \sim 3 m in Casco Bay.

The NEPs on the East Coast are characterized by cool waters with some coastal upwelling. Casco Bay is fed by twelve significant lake and river systems and also has a large tidal influence. The Bay is dotted with roughly 785 islands, islets, and exposed ledges. Casco Bay has an important shellfish restoration and aquaculture industry, including lobster and clam fisheries. The Barnegat Bay estuarine system is composed of three shallow, micro-tidal bays: Barnegat Bay, Manahawkin Bay, and Little Egg Harbor. A nearly continuous barrier island complex runs along the eastern edge of Barnegat Bay, separating it from the Atlantic Ocean. The estuary has an upwelling center off Little Egg Inlet, and it is also fed by low pH and alkalinity freshwater. It is located in an urban watershed and the Upper Barnegat Bay is highly eutrophic. There are a number of shellfish aquaculture and restoration projects throughout the watershed, in addition to the historic hard clam fishery.

The NEPs in the GOM region are in a transition zone between warm-temperate and tropical biogeographic provinces, and are characterized by warm, productive waters. Tampa Bay is a large, shallow, open-water estuary stretching 1,030 km²



at high tide. It is influenced by the mixing of GOM waters with freshwater flow from more than 100 tributaries, dozens of meandering, brackish-water creeks and four major rivers. On average, Tampa Bay is only ~ 4 m deep; however, manmade shipping channels have been dredged to allow large ships safe passage to the Port of Tampa and other bay harbors. It has economically important shellfish and finfish populations. Tampa Bay is specifically examining the role of seagrass in protecting marine species from the harmful effects of coastal acidification. The monitoring location in the Mission Aransas Estuary in the Port Aransas Ship Channel (i.e., Aransas Pass tidal inlet) connects the Gulf coastal water with the Aransas, Corpus Christi, and Redfish bays. As secondary bays to Corpus Christi and Aransas bays, Nueces and Copano bays receive freshwater input from the Nueces River and Mission/Aransas rivers, respectively. The high alkalinity of freshwater flows into Mission Aransas Estuary are an important characteristic of the bay (Hu et al., 2015).

The NEPs in the West Coast region are characterized as having cooler, deeper waters with prominent coastal upwelling (Leinweber and Gruber, 2013; Raimonet and Cloern, 2017). Santa Monica Bay is influenced by both freshwater inflows, primarily from the Ballona Creek, Malibu Creek, and Topanga Creek watersheds, and coastal upwelling. Santa Monica Bay is especially unique amongst these seven estuaries because it is more of an open, deep coastal site on a narrow continental shelf rather than an enclosed estuary. The Santa Monica Bay Foundation is specifically looking at the feasibility of restored kelp forests as a pH refuge for marine life from the harmful effects of ocean acidification. The San Francisco Bay monitoring sites are located within a tidal excursion of the mouth of the bay, at the interface between Central Bay (outer embayment) and San Pablo Bay (North Bay). There is a high range of salinity at this location in which at low tide, there is an estuarine water signal influenced by freshwater flows, while at high tide, there is an ocean water signal. There is no commercial shellfish production in San Francisco Bay due to historic water quality issues. The herring fishery is the only commercial fishery inside San Francisco Bay; however, there is an extensive nursery habitat for the economically important Dungeness crab fishery, restoration efforts for the native Olympic oyster, and concern for migrating salmonid and other endangered species in the upper estuary. The Tillamook Estuary is a relatively shallow estuary with maintained jetties and channels less than 6.7 m. It is located in a more rural watershed with high nutrient inputs from dairy and timber farms and wastewater treatment plants. The estuary is influenced by periods of high river discharge. Tillamook Estuary

has commercial shellfish aquaculture (including oysters and bay clams) as well as extensive recreational fisheries. The monitoring site in Tillamook Estuary is located at the Port of Garibaldi, which is a commercial fish offloading dock near the mouth of the estuary as well as in the vicinity of a wastewater treatment outfall.

Monitoring Timeline

Casco Bay was the first program to begin coastal acidification monitoring in 2015. The other NEPs began monitoring in 2016 or 2017. **Figure 2** shows the time periods in which monitoring data were collected in each NEP as of publication of this study. Most continue to collect data as of June 2020.

Deployments and Measurements

The monitoring approaches used by each program varied, including deployment methods, types of equipment, and discrete sampling methods, and were driven largely by existing capacities as well as regional influences and scientific interests. Coastal acidification monitoring equipment was deployed on fixed, land-based structures (e.g., docks, piers, and pilings), as well as water-based moorings (Supplementary Table 2, Figure 3). Differences in deployment locations were undoubtedly reflected in the recorded data. For example, shallower, nearshore sites were expected to preserve a stronger diel biological signal and would be more sensitive to freshwater runoff or land-based pollutants as compared to further offshore sites like deeper water Santa Monica Bay and San Francisco Bay deployments. Each of the seven NEPs collected in situ measurements of pCO₂, pH, temperature, salinity, and DO (Tables 1, 2). However, DO data for Mission-Aransas Estuary were collected at a different depth in the water column and therefore were not evaluated.

Instruments

Several different autonomous sensors were used in the estuaries. Specifications of the various instruments, including instrument resolution, precision, and operating ranges, are shown in Table 2. The sensors used for measuring pH include the Sea-Bird SeaFET and Sea-Bird SeapHOx. The SeaFET pH sensor is an ion-sensitive field effect transistor (ISFET), which is more precise and stable over time compared to pH sensors that use glass electrodes. The pH range for the SeaFET is 6.5 to 9.0. The SeapHOx integrates a SeaFET pH sensor with additional sensors that measure temperature, salinity, pressure, and DO. The SeapHOx also includes an internal water pump and anti-fouling technology. Both the SeaFET and SeapHOx have internal battery power and data logging capabilities. For measurement of pCO₂, four of the NEPs used the Sunburst SAMI-CO2 and the remaining NEPs used Pro-Oceanus CO2-Pro or Moored Autonomous pCO₂ (MAPCO₂) systems. The Sunburst SAMI-CO2 uses a colorimetric method to measure the partial pressure of CO₂ from 200 to 600 µatm typically, although extended range calibrations are available by request through the manufacturer. The Pro-Oceanus CO2-Pro and MAPCO₂ measure pCO_2 using an infrared CO₂ detector. See Sutton et al. (2014) for additional details on the MAPCO₂ system. Supporting data (e.g., temperature, salinity, pressure, DO), are measured using a variety of Sea-Bird, Yellow Springs Instrument (YSI), or Aanderaa instruments. All pH data reported by the NEPs are on the total pH scale, (pH_T) allowing direct comparison.

METHODS

Data QA/QC and Validation

The protocols for assessing data quality and validating *insitu* measurements varied across the NEPs. Because they are all federally funded programs, all NEPs incorporate comparably stringent Quality Assurance/Quality Control (QA/QC) procedures in their data screening and validation protocols as outlined in a Quality Assurance Project Plan (QAPP)². This includes collecting data that comply with EPA rules for surface-water-quality monitoring programs and water-quality assessments to support decisions related to mission objectives.

Reporting Limits

Instrument reporting limits were determined by the range and precision of the sensors being used as provided by the manufacturers of the instrument sensors. This information is summarized in **Table 2**.

Accuracy

Accuracy measures how close results are to a true or expected value and can be determined by comparing a standard or reference sample to its actual value. According to the specifications of the SeapHOx and the CO2-Pro CV, the pH and pCO_2 sensors have accuracies of \pm 0.05 and 0.5% of the measured value, respectively. The Sea-Bird SeaFET and the SunBurst SAMI-CO₂ sensors have accuracies of \pm 0.05 and \pm 0.3 µatm, respectively (Table 2). Due to the potential for signal drift and biofouling, the NEPs used laboratorygrade instruments to perform validations of autonomous data through measurement of discrete water samples collected from the instrument deployment sites. For example, in Tampa Bay, instrument sensors underwent laboratory validation in an instrument test tank which was cross-validated with discrete measurement of parameters. Accuracy of the measurements was ensured by using Certified Reference Materials for analysis of dissolved inorganic carbon (DIC) and total alkalinity (TA). Similar protocols were employed by the other NEPs. Discrete samples were typically collected every 1-6 weeks (see section "Discrete Sampling" for details).

Precision

The precision of data is a measure of the reproducibility of a measurement and includes components of random error. Precision is strictly defined as a measure of the closeness with which multiple analyses of a given sample agree with each other. For this reason, the most common method used to collect real-time data from pH and pCO_2 sensors

²https://www.epa.gov/quality/quality-assurance-project-plan-development-tool



TABLE 1 Autonomous sensors used to monitor water chemistry in seven of United States Environmental Protection Agency's (EPA's) National Estuary Programs (NEPs).

Water Body	Autonomous Deployment Instruments						
	CTD ^a	pCO ₂	рН	DO			
Santa Monica Bay	Sea-Bird SeapHOx	Sunburst SAMI-CO ₂	Sea-Bird SeapHOx	Sea-Bird SeapHOx			
San Francisco Bay	Sea-Bird SeapHOx	MAPCO ₂	Sea-Bird SeapHOx and Sea-Bird SeaFET	Sea-Bird SeapHOx			
Tillamook Bay	Sea-Bird SeapHOx and YSI	SunBurst SAMI-CO ₂	Sea-Bird SeapHOx and Sea-Bird SeaFET	Sea-Bird SeapHOx and YSI			
Barnegat Bay	YSI	Pro-Oceanus CO ₂ -Pro CV	Sea-Bird SeaFET	YSI			
Casco Bay	Sea-Bird CTD	SunBurst SAMI-CO ₂	Sea-Bird SeaFET	Aanderaa Oxygen Optode			
Mission-Aransas Estuary	YSI	SunBurst SAMI-CO ₂	Sea-Bird SeaFET	YSI			
Tampa Bay	Sea-Bird SeapHOx	Pro-Oceanus CO ₂ -Pro CV	Sea-Bird SeapHOx	Sea-Bird SeapHOx			

pCO₂, partial pressure of carbon dioxide; DO, dissolved oxygen.

^aA variety of electronic instrument packages were used by the NEPs to measure conductivity, temperature, and depth (CTD).

in the field by the NEPs was to average a minimum of five consecutive measurements made by the sensors at their respective maximum sampling frequencies. Should extreme variations occur, the sensors were checked to verify their proper functionality, and service was conducted depending on the nature of the problem.

Bias

Bias (or drift) is a measurement of correctness and includes components of systematic error. A measurement is considered unbiased when the value reported does not differ from the true value. For the SunBurst SAMI-CO₂, internal periodic blanks were automatically run to correct for drift of the electro-optical

TABLE 2 | Sensor Specifications.

Instrument	Parameter	Accuracy	Precision	Resolution	Range
SunBurst SAMI-CO ₂	pCO ₂	+/- 3 μatm	± 0.5–1 μatm		150–700 ^a
Pro-Oceanus CO ₂ -Pro CV	pCO ₂	\pm 0.5% of meas. val.	0.01 ppm		0-10,000
MAPCO2 ^b	pCO ₂	< 2 µatm	0.7 ppm		0–800
Sea-Bird SeapHOx	рН	± 0.05		±0.004	6.5–9
	DO	\pm 0.1 mg L ⁻¹		$0.2 \ \mu mol \ kg^{-1}$	120% of surf. sat.
	Temp	\pm 0.002°C ^c \pm 0.01°C ^d		0.0001°C	-5 to 45°C
Satlantic SeaFET	рН	± 0.05	± 0.004		6.5–9
Aanderaa Oxygen Optode	DO	$< 8 \ \mu M$		$<0.1 \ \mu M$	0–1,000 μM

^ainstrument can be calibrated for extended ranges.

^bLiCOr LI-820 CO₂ gas analyzer (Sutton et al., 2014).

^cTemperature range: –5 to 35°C.

^dTemperature range: 35 to 45°C.

system, while reference measurements of the light emitting diodes (LEDs) correct for interim deviations. Instruments were inspected, maintained, and cleaned periodically to contain drift caused by biofouling or other factors.

For Tampa Bay, the data were deemed biased if the sensorcollected values deviated from \pm 0.05 pH unit for pH and \pm 0.5 percent for *p*CO₂. In Mission-Aransas Estuary, the data was deemed biased if the sensor collected values deviated from the specified accuracy values (i.e., \pm 0.02 pH unit for pH and \pm 3 µatm for *p*CO₂).

Discrete Sampling

In addition to *in situ* sensor measurements, discrete water samples were collected by the NEPs and their partners to validate the sensor measurements and provide additional analytical data necessary to characterize the water chemistry. These data are not included in the analysis for this study but will be the focus of forthcoming estuary-specific reports. The NEPs most often collected and analyzed discrete samples for pH, DIC, and TA. The frequency of discrete sample collection ranged from weekly (e.g., Barnegat Bay), to bi-weekly to monthly (e.g., Tillamook Bay; Tampa Bay; Mission-Aransas Estuary) to every 4–6 weeks (Casco Bay; San Francisco Bay) to quarterly (e.g., Santa Monica Bay) and was often timed to coincide with sensor cleaning, other maintenance, and data downloading.

Some NEPs also used discrete or *in situ* measurements collected by other research programs to cross-calibrate their sensor data. For example, Santa Monica Bay used conductivity, temperature, depth (CTD) profile data, collected quarterly by the Los Angeles County Sanitation District (LACSD) at nearby stations, to evaluate the comparability between those CTD measurements and the acidification mooring sensors.

In summary, sensor data collected across the NEPs was assured through factory and laboratory instrument calibrations, validation to water quality measurements (discrete samples), inspection of the data record for anomalous data, proper maintenance and examination for bias and fouling, checks against oceanographic properties and other programs' data and electronic validation and verification. Only data obtained from *in situ* instrument sensors which were supported by appropriate quality control data and met the measurement performance specifications defined here were considered acceptable and used in this analysis.

In this manuscript, we focus on data QA/QC and validation derived from instrument resolution and limitations as provided by the manufacturers of the instrument sensors (**Table 2**) along with internal trend and correlation analysis (see section "Overall Patterns of pCO_2 and pH"). Additional validation and continued refinement of protocols are underway across the NEPs as more data continues to be collected and will be the focus of their own forthcoming reports and publications.

Descriptive Statistics

Statistical analyses of the data³ were performed using R Statistical Software (version 3.6.2) and Tableau Desktop (version 2020.2) data visualization software. As an initial assessment, the site-specific range, mean, and standard deviation of each measured parameter (pCO_2 , pH, temperature, salinity, and DO) were calculated for each northern hemisphere meteorological season: winter (December-February), spring (March-May), summer (June-August), fall (September-November). Box plots of monthly-binned data, multi-parameter time series plots, and cross-plots of measured parameters were prepared to visualize relationships among coastal carbonate system parameters within, and across the water bodies at short (diel) and longer (seasonal) time scales.

Thermal and Nonthermal Controls on pCO_2

Physical factors (e.g., temperature-dependent solubility, transport and mixing [including coastal upwelling and freshwater outflow]) and biological factors (e.g., primary production and respiration) are important controls on pCO_2 changes in coastal and estuarine environments (DeGrandpre et al., 2002; Feely et al., 2008, 2018; Huang et al., 2015; Wanninkhof et al., 2015). To separate the solubility effect due to water temperature change on observed pCO_2 (pCO_{2obs}) and to contrast the effect of temperature with the cumulative influence of other physical and biological factors, we calculated (1) non-temperature controlled pCO_2 values ($pCO_{2bio/hvdro}$) and (2) temperature-controlled

³https://doi.org/10.25921/xg33-1n83

 $pCO_{2thermal}$ values ($pCO_{2thermal}$) following the approach of Takahashi et al. (2002). It is acknowledged that Takahashi et al. (2002) represents an open-ocean study with minimal salinity variance. The coastal systems discussed in this study experience large salinity swings, which will slightly impact Henry's Law constant (K_H), and, by extension, the dissolution of CO₂ as described by Henry's Law: ($[CO_{2(aq)}] = K_H * pCO_2$). $pCO_{2bio/hydro}$ values were calculated for each water body on a site-specific basis by normalizing pCO_{2obs} values to the site-specific mean water temperature as calculated over the course of monitoring [**Eq. (1**)]:

$$pCO_{2\ bio/hydro} = pCO_{2\ obs} \times e^{0.0423(T_{mean} - T_{obs})}$$
(1)

where pCO_{2obs} is the pCO_2 at *in situ* temperature in degrees Celsius (°C), T_{mean} is the site-specific mean temperature, which ranged from 11.4°C in Tillamook to 24.2°C in Mission-Aransas, and T_{obs} is the site-specific *in situ* temperature in degrees Celsius. Changes in $pCO_{2bio/hydro}$ primarily represent changes in pCO_2 due to the combined influences of respiration and production, alkalinity, upwelling, diffusion of CO₂ between the ocean and atmosphere, and advection past the sensor suite by tides and currents.

In contrast, to calculate the effect that only temperature would have on pCO_{2obs} , we again followed the method of Takahashi et al. (2002):

$$pCO_{2 thermal} = pCO_{2 mean} \times e^{0.0423(T_{obs} - T_{mean})}$$
(2)

RESULTS

Overall Patterns of pCO₂ and pH

Variability in observed pCO_{20bs} and pH are shown in Figures 4, 5. The mean, range, and standard deviation of the mean for each measured parameter on a seasonal basis are shown in Supplementary Table 3. Box plots with all underlying data shown illustrate the variability of pCO_{20bs} and pH data that were collected within and across the seven water bodies (Figures 4, 5). Variability discussed here refers to the difference between the upper and lower whiskers (1.5 times the interquartile range [IQR]) of the box plots. This was done to facilitate comparison of data across a range of systems with variable sample sizes and monitoring timelines and reduce the risk of including potentially short-lived, albeit likely real, extreme events. Note that all data included in the results and discussion passed Quality Assurance/Quality Control (QA/QC) checks and therefore are assumed to represent real data. The term "IQR pCO2" and "IQR pH" used in the following sections refers to the variability in pCO₂ and pH values, respectively, that fall within 1.5 times the IQR.

The largest ranges of pCO_2 values, based on the difference between the upper and lower whiskers of the box plots, were observed in Casco Bay (191 – 1,024 µatm) and Barnegat Bay (438 – 1,180 µatm; **Figure 4**). When all data are considered, Tillamook Bay has both the lowest (143 µatm) and highest (1,405 µatm) measured pCO_2 values. In Casco Bay, variability in IQR pCO₂ was associated with the correspondingly largest range in IQR pH (7.39 - 8.32) measured across all of the water bodies and highly variable salinity (Supplementary Figure 1). Similarly, in Barnegat Bay the variability in IQR pCO₂ values was associated with a correspondingly large range in IQR pH (7.61 - 8.23) but smaller range of salinity values (Figures 4, 5, Supplementary Figure 1; Supplementary Table 3). The smallest range in IQR pCO_2 was observed in Mission-Aransas Estuary (287 - 568 µatm), where IQR pH values ranged from 7.86 to 8.37 (Figures 4, 5). Ranges of IQR pCO₂ (374 - 860 µatm) and pH (7.70 - 8.13) in Tampa Bay were similar to those in Mission-Aransas. The two Santa Monica Bay deployments exhibited the smallest ranges of temperature, salinity, and DO, with deep waters being the least variable (Figures 4, 5, Supplementary Figure 1; Supplementary Table 3). Deep waters in Santa Monica Bay also exhibited the smallest range of IQR pH values (7.61 -8.01), although IQR pCO2 values were more variable (382 -1,028 µatm; Figures 4, 5, Supplementary Table 3). Salinity values in both Santa Monica Bay deployments fell in a very narrow range (Supplementary Figure 1). Variability in IQR pCO₂ and pH was greater in nearshore waters of Santa Monica Bay (**Figures 4**, **5**). Variability in San Francisco Bay IQR pCO_2 values (372 - 1,018 µatm) were similar to those observed in Santa Monica Bay. Deeper water Santa Monica Bay and San Francisco Bay showed a bimodal distribution of pCO₂ values (Supplementary Figure 2). Time series plots of all measured parameters from all sites are shown in Supplementary Figures 2-6 and relationships between parameters are shown as cross-plots in Supplementary Figures 7-15. Frequency distribution plots for pCO₂ and pH data from each water body are shown in Supplementary Figures 16-17.

Seasonal Patterns in pCO₂ and pH

Seasonal patterns in pCO₂, pH, temperature, salinity, and DO were observed across the water bodies. In general, pCO_2 values were higher and more variable in warmer, summer months and lower and less variable during cooler, winter months (Figures 6-11; Supplementary Figure. 2). The major exception to this pattern is Santa Monica Bay, where the highest pCO_2 values were observed during the spring (Figures 6, 8; Supplementary Figure 2). Both Mission-Aransas Estuary and Tampa Bay displayed a pattern of increasing pCO_2 from cooler, winter months into warmer, summer months with yearly pCO_2 values reaching their peak in July and August (Figures 6, 9). Overall, the magnitude of these seasonal changes in pCO_2 is significantly muted in Mission-Aransas Estuary and Tampa Bay as compared to the other water bodies. Salinity was more variable throughout the year in Mission-Aransas than in Tampa Bay (Supplementary Figure 18). Similar seasonal patterns of pCO_2 variability were observed in Casco Bay and Tillamook Bay where data from seasons over consecutive years are available (Figures 6, 10, 11, Supplementary Figures 2, 3). In these water bodies, however, a shift towards a trend of increasing pCO_2 values began earlier in the year- in late winter (February) in Tillamook Bay and in mid-spring (April) in Casco Bay- with maximum pCO2 values generally occurring in late summer-early fall. A shift



back towards decreasing pCO₂ values occurred in mid-summer (August) in Tillamook Bay to mid-fall (October) in Casco Bay. These seasonal patterns and the relative magnitude of seasonal variability are comparable across consecutive years [i.e., 2015-2018 (Casco Bay); 2017-2019 (Tillamook Bay)]; although, 2019 winter – early Spring pCO₂ values in Tillamook Bay were slightly elevated as compared to 2018 winter values. In Santa Monica Bay, pCO_2 values increased from the winter into the spring with values reaching their maximum in May before steadily decreasing through the fall and reaching their lowest values during the winter (Figures 6, 8). High pCO_2 values in the spring-early summer in Santa Monica Bay are associated with high salinity, low temperature, low DO water (Supplementary Figure 18; Figure 8), indicative of waters that have upwelled during the spring and early summer. Similar patterns of seasonal pCO₂ variability were observed in San Francisco Bay; however, the shift from decreasing to increasing pCO_2 values occurred later in spring (~April). During the summer, pCO₂ values were highest and remained relatively high (monthly median \sim 720–750 ppm) through December. pCO₂ data for Barnegat Bay is relatively sparse compared to other water bodies and there is variability in the data; however, there are indications of a similar trend of increasing pCO_2 from winter months into the spring; however, there is variability in the data (Figure 6).

In general, seasonal trends in pH were opposite of those observed in pCO_2 throughout the year with rises and falls in pCO_2 associated with falls and rises, respectively, in pH

(Figure 7). Significant variability (i.e., large range of values) in pH was observed throughout the year in all water bodies, but this variability was particularly large in summer and fall months where it was observed across multiple connective years in Tillamook Bay and Casco Bay, respectively (Figures 7, 10, 11). Some of the lowest recorded pH values across all of the water bodies were associated with these periods of high variability in pH and salinity values in Tillamook and Casco bays (Figure 7; Supplementary Figures 3, 18). Deeper water in Santa Monica Bay and San Francisco Bay showed similarly low pH values in the fall and summer, respectively (Figure 7). Overall, Mission-Aransas Estuary, Tampa Bay, and Barnegat Bay share similar patterns of seasonal pH variability with pH values generally falling from the winter into the summer followed by a trend towards higher values in the fall and back into the winter (Figure 7). The deep and shallow records from Santa Monica Bay show similar seasonal patterns of pH with values decreasing from the winter into the spring and then generally increasing into the fall, a pattern opposite of that observed for pCO_2 (Figures 6, 7). Although pCO_2 variability in Santa Monica Bay and San Francisco Bay was quite similar, seasonal patterns of pH variability within these two water bodies were very different. In San Francisco Bay, pH values were highest in spring followed by an overall decrease in pH values through the fall before a return to slightly more positive values in the winter (Figure 7). This pattern was particularly clear at the 17 m sensor in San Francisco Bay, where a more complete dataset exists.



partial pressure of carbon dioxide (pCO_2) (Mission-Aransas Estuary) to highest median pCO_2 (Barnegat Bay) as calculated from all available data. Whiskers extend to data within 1.5 times the interquartile range; * = multi-year records. SMB. SMB_15 = Santa Monica Bay (15 m deployment); SMB_60 = Santa Monica Bay (60 m deployment); SFB_1 = San Francisco Bay (1 m deployment); SFB_17 = San Francisco Bay (17 m deployment).

Thermal and Nonthermal *p*CO₂ Calculations

Calculations of thermal and nonthermal controls on pCO₂, referred to as pCO_{2thermal} and pCO_{2bio/hydro}, respectively, were made in order to contrast the influence of water temperature on pCO_2 variation and the aggregate of other processes such as primary productivity, respiration, and mixing. These theoretical pCO₂ values and *in situ*, observed pCO₂ values (pCO_{2obs}) are shown in Figure 12. Plots of all pCO_{20bs}, pCO_{2thermal} and pCO_{2bio/hvdro} data are in Supplementary Figures 19-26. Relationships and patterns of pCO_{2obs}, pCO_{2thermal} and pCO_{2bio/hydro} are similar in Tampa Bay and Mission-Aransas Estuary. A trend towards increasingly higher pCO_{2thermal} values was observed from the winter through summer months in these GOM water bodies, with peak pCO_{2thermal} values occurring in July (Figure 12). In general, seasonal patterns of *p*CO_{2*bio/hydro*} are opposite of those seen in pCO_{2thermal}, with pCO_{2bio/hydro} values falling from the winter through summer months and reaching their lowest values in the summer (Figure 12).

In Casco Bay, pCO_{2obs} , $pCO_{2thermal}$, and $pCO_{2bio/hydro}$ showed broadly similar patterns throughout the year, and seasonally across consecutive years (**Figure 12**, **Supplementary Figure 24**). During the spring, pCO_{2obs} values were well below what would

be expected from temperature alone ($pCO_{2thermal}$). pCO_{2obs} values generally increased from spring into the fall, reaching their highest values around October, before beginning to decrease into the cooler, fall and winter months. $pCO_{2bio/hydro}$ values continued to increase into the late fall, reaching their maximum around October, while $pCO_{2thermal}$ values reached their maximum around August before beginning to decrease into the cooler, winter months. Only a few months of data are available for Barnegat Bay, therefore seasonal patterns were not distinguished (**Figure 12**).

Relationships among pCO_{2obs} , $pCO_{2thermal}$, and $pCO_{2bio/hydro}$ and seasonal patterns of pCO_2 variability are very different in West Coast water bodies. In Santa Monica and Tillamook bays, $pCO_{2bio/hydro}$ was more closely associated with pCO_{2obs} while seasonal undulations in $pCO_{2thermal}$ were less extreme, as compared to the patterns in Mission-Aransas Estuary and Tampa Bay (**Figure 12**; **Supplementary Figures 19**, **20**, **22**). Although muted, $pCO_{2thermal}$ values in Santa Monica Bay (both depths/records), generally decreased from winter through spring. This is followed by an upward trend towards more positive $pCO_{2thermal}$ values from spring into summer. Seasonal patterns in $pCO_{2bio/hydro}$ were similar across these three water bodies (**Figure 12**). At both depths in Santa Monica Bay and in Tillamook Bay, $pCO_{2bio/hydro}$ and pCO_{2obs} tracked



one another very closely throughout the year (i.e., there was little influence of changes in solubility associated with changes in temperature). In Tillamook, this pattern was observed over multiple consecutive years.

Diel Patterns in pCO₂

Daily ranges of monthly binned nonthermal pCO_2 (pCO_{2bio/hydro}) values binned by month are expressed in box plots and highlight the extent to which diel ranges vary throughout the year within, and across the water bodies (Figure 13). In general, with the exception of Santa Monica Bay and San Francisco Bay, overall patterns are similar to those observed in the monthly binned data in that there is greater diel variability in warmer months. Some of the greatest diel fluctuations in pCO_{2bio/hydro} were observed in Tillamook Bay during the late summer-early fall with values increasing from the spring into fall (Figure 13C). Large diel ranges in pCO_{2bio/hydro} were observed at nearshore Santa Monica Bay and near-surface San Francisco Bay where diel variability increased through spring and summer and the largest ranges were found in early summer (Figures 13B,G). Consistent and relatively small diel ranges in

 $pCO_{2bio/hydro}$ were observed in deeper waters in Santa Monica Bay, which is below the euphotic zone (Figure 13F). Large diel ranges were observed in Barnegat Bay as well; however, data are only available for January–April (Figure 13H). Overall, diel variability in $pCO_{2bio/hydro}$ in Tampa Bay and Mission-Aransas Estuary was smaller compared to the other water bodies (Figures 13A,D). Median ranges were typically < 100 µatm in Mission-Aransas Bay and Tampa Bay throughout the year.

DISCUSSION

Estuaries and coastal waters are highly vulnerable to the impacts of acidification. Yet, little is known about the extent of this vulnerability and the relative influence of estuary-specific drivers that contribute to acidification, such as intrusion of CO_2 -rich seawater, nutrient enrichment, locally elevated atmospheric CO_2 from urban and agricultural activities, net ecosystem production and respiration, and freshwater flows (Salisbury et al., 2008; Feely et al., 2008, 2010, 2018; Gledhill et al., 2015; Northcott et al., 2019; Rheuban et al., 2019). This multitude of influences leads



to greater pH and pCO₂ variability and more acidic conditions as compared to open ocean values (Cai et al., 2021). For example, pH values ranged from 7.16 to 8.32 in Casco Bay and from 7.21 to 8.31 and in Tillamook Bay whereas the global range of surface ocean pH values is \sim 7.7 to 8.7 (Takahashi et al., 2014; Bakker et al., 2016; Jiang et al., 2019). Comparably large pH ranges, up to 1.2 within a single month, have been observed at other coastal sites with some of the most extreme pH values showing large interannual variability (e.g., Dorey et al., 2013; Sutton et al., 2016; Pecquet et al., 2017). Autonomous surface deployments in coastal settings have recorded seasonal (wintersummer) CO₂ differences of up to 330 µatm compared to difference up to 70 µatm in open ocean settings (Shadwick et al., 2015; Sutton et al., 2019). For comparison, seasonal pCO₂ differences observed in the coastal systems described here are \sim 2-4x as large (e.g., Casco Bay – 833 µatm; Barnegat Bay – 697 µatm; Santa Monica Bay - 646 µatm), the largest being observed in Tillamook Bay (1,263 µatm). In some cases (e.g., Tillamook Bay, Casco Bay, Santa Monica Bay_15 m), diel amplitudes in

 pCO_2 and pH are comparable to those observed seasonally. Similar observations have been made in other coastal systems (Torres et al., 2021), albeit the amplitudes were not as extreme as those observed in this study. Given the variability in estuarine systems, multi-year to decadal deployments will be needed to more fully understand the natural variability and important processes driving acidification in these systems. For this initial data synthesis, analyses were restricted to shorter (e.g., seasonal) time scales that are appropriate for existing datasets.

An important goal of the NEP coastal acidification monitoring program is to inform longer-term management decisions as more data are collected. For example, aragonite saturation state (Ω Ar) is an important calculated measure that is most closely related to organism health for calcifiers and can be used to understand the vulnerability of water bodies to acidification. This comparative study did not calculate Ω Ar for each water body because additional parameterization of the carbonate chemistry, including validation data, is necessary and was not available on a consistent basis across the water bodies. Specifically, calculating



2018–February 2019.



aragonite saturation requires that, in addition to temperature and salinity, at least two of the carbonate parameters (pCO_2 , total alkalinity, DIC, pH) be known. However, pCO_2 and pH are not an ideal set of input parameters for calculating Ω Ar because they carry the largest uncertainties compared to other input pairs (Millero, 1995; Orr et al., 2018) and the controls on these parameters (e.g., pH and Ω Ar) are not entirely consistent (Xue et al., 2021). Such uncertainties may be even larger for estuarine sites with highly variable salinity as disassociation constants are poorly constrained compared with ocean salinity. Ideally a better constrained Ω Ar dataset should be independently verified through inclusion of additional CO₂ system parameters (e.g., DIC and TA). However, these data were not collected across all water bodies, and as such, the calculated Ω Ar results would be based on inconsistent methodologies (i.e., using variable input parameters).

Rather, this study's focus is on patterns of pCO_2 and pH variability and the relationship of these parameters to temperature, salinity, and DO. A similar preliminary analysis of high-frequency monitoring data from the National Estuarine





Research Reserve System has shown concurrent fluctuations in pH and DO to be characteristic of nearshore habitats and explored how pH data paired with DO data could be recording the effects of enhanced coastal acidification (Baumann and Smith, 2017).

Patterns of pCO₂ and pH

In general, across all water bodies, temperature and pCO_2 were lower in cooler, winter months and higher in warmer, summer months. A similar pattern has been observed in other coastal systems (Wallace et al., 2014).

Higher-latitude, cooler water bodies like Tillamook Bay, Casco Bay, and Barnegat Bay and the deeper waters of California coastal water bodies have slightly higher pCO_2 and lower pH than warmer GOM water bodies. This is likely due in part to the large annual temperature variability that typifies these higher latitude systems as compared to subtropical, warmer GOM systems (Tampa Bay and Mission-Aransas). For example, in Barnegat Bay in the northeastern United States, there was a 31°C range in annual water temperature (-1.9°C to 29.3°C) while in Tampa Bay in the GOM, the annual temperature range was 19°C (13.1°C to 31.8°C). The higher *p*CO₂ values observed in Santa Monica Bay and San Francisco Bay are best explained by the upwelling of cold, high *p*CO₂, low pH water rather than a large annual temperature range; annual temperatures in the deeper water Santa Monica Bay and San Francisco Bay deployments only vary by ~11.6°C. Furthermore, the timing of shifts toward increasing (or decreasing) *p*CO₂ values varied by location and appear to be



largely dictated by local climate conditions, leading to increases in pCO_2 that began earlier in the year in the warmer water, lower latitude GOM water bodies of Tampa Bay and Mission-Aransas Estuary as compared to cooler water, higher latitude water bodies such as Tillamook Bay and Casco Bay. Thermally driven changes are also inferred to be largely responsible for the lower variability of pCO_2 values observed in lower latitude, warmer water bodies as compared to higher latitude, cooler water bodies that have larger ranges of annual temperature variability.

Strong seasonal variations in pCO_2 have been observed in other coastal systems (see Jiang et al., 2021 for a recent compilation of coastal data sets). For example, in the Delaware Bay, during the warmer summer months, pCO_2 values of ~400– 500 µatm, have been recorded, with values reaching 200 to 350 µatm in the mid- and lower bay regions, while in the winter (December) pCO_2 measured across the bay ranged from 500– 650 µatm (Joesoef et al., 2015). Biological and thermal controls as well as variable mixing and stratification and variability in river discharge have been inferred as the dominant controls responsible for this variability (Joesoef et al., 2015).

Other coastal systems in the northeastern United States (e.g., Long Island Sound, NY, United States; Narraganset Bay, RI, United States; Jamaica Bay, NY, United States; Hempstead Bay, NY, United States) are characterized by high pCO_2 (>1000 μ atm) coupled with low DO and low pH (<7.4) during warmer summer months (Wallace et al., 2014). These patterns have been largely attributed to high rates of microbial respiration driven by enhanced nutrient loadings coupled with water column stratification with changes in summer temperature accounting for < 5% of the pCO_2 increases (Wallace et al., 2014).

Sensor deployment location in the estuary (nearshore, shallower water vs. offshore, deeper water) surely impacted the data collected during this study via the extent to which various processes are at play (e.g., freshwater runoff, biological activity, tides, heat dissipation [shallow vs. deep]) in these different

locations. This will be and important consideration in future efforts to tease out drivers of acidification in these water bodies and to facilitate comparison with other data sets.

Thermal and Nonthermal *p*CO₂ Calculations

In addition to seasonal changes in temperature, the pCO_2 in nearsurface waters that exchange CO_2 directly with the atmosphere is affected by DIC, pH and alkalinity. Water temperature is primarily regulated by physical processes (i.e., solar energy input, ocean-atmosphere heat exchange, and upwelling). DIC concentration and alkalinity, however, are controlled by a combination of physical (transport and mixing of different water masses, air-sea gas exchange) and biological processes (i.e., photosynthesis and respiration; carbonate precipitation and dissolution) (Feely et al., 2008, 2018; Cai et al., 2020).

To better understand the relative influence of water temperature with respect to other physical and biological processes, theoretical pCO_{2thermal} and non-thermal pCO_{2bio/hydro} values were calculated. Respectively, these values reflect the predicted pCO_2 values if temperature were the sole control on pCO_2 and the predicted pCO_2 in the absence of temperature change. Changes in pCO_{2bio/hydro} primarily represent changes in total CO2 due to the combined influences of biology and diffusion of CO₂ between the ocean and atmosphere, as well as advection past the sensor suite by tides and currents in the absence of temperature changes. Changes in pCO_{2thermal} are theoretical changes in pCO_2 as a result of the decreasing solubility of CO₂ with increasing temperature. Therefore, the relative magnitudes of $pCO_{2thermal}$ ($\Delta pCO_{2thermal}$) and $pCO_{2bio/hydro}$ ($\Delta pCO_{2bio/hydro}$) should reflect the dominant factor (i.e., temperature vs. other parameters) in controlling the observed pCO₂ changes throughout these individual monitoring periods. The overall pattern across the estuaries



is that $\Delta p \text{CO}_{2thermal}$ is smaller compared to the magnitude of $p \text{CO}_{2bio/hydro}$ except in Mission-Aransas Estuary and Tampa Bay. This is consistent with a previously well-described pattern of temperature controlling open-ocean surface-water seasonal $p \text{CO}_2$ variations at subtropical and lower temperate latitudes and biophysical processes exerting more significant control at moderate and higher latitudes (Takahashi et al., 2002).

Collectively, the observed differences among $pCO_{2thermal}$, $pCO_{2bio/hydro}$, and pCO_{2obs} values described here indicate that temperature and the aggregate of other factors, including seasonal net community productivity, tidal fluctuations, and freshwater inflows are equally important in controlling pCO_2 variability in Mission-Aransas and Tampa Bay with the degree to which these processes impact pCO_{2obs} varying throughout the year. In Tampa Bay, strong tidal control and the location of the sensor package near the mouth of the bay may have resulted in capturing of variable and complex signatures due to both water mass exchange between Tampa Bay and the GOM and associated salinity and DO differences and biological productivity.

Multidecadal decreases in both total alkalinity and pH have been observed in Mission-Aransas Estuary (Hu et al., 2015). The rivers feeding Mission-Aransas Estuary are highly alkaline and the long-term acidification has consequently been attributed to decreased freshwater inflow (and a resulting decreased alkalinity delivery) to the estuary due to freshwater diversions (Hu et al., 2015). The relative importance of thermal and biological controls at different times appears to vary with variations in freshwater inflow (Yao and Hu, 2017), highlighting the importance of freshwater inflow variability in this semiarid system. While the Mission-Aransas Estuary is microtidal, the tidal fluctuations at the Aransas Ship Channel, where the autonomous sensors were located, are relatively large compared to the upper estuary and appear to exert an important control on the daily fluctuations of pH and pCO₂ causing expected differences between day and night observations (based on biological activity) to reverse for portions of the year.

The co-occurrence of high pCO_2 , low pH, low DO, high salinity, and low temperature water is consistent with

a coastal upwelling signature which is known to occur in the vicinity of Santa Monica Bay, San Francisco Bay and Tillamook Bay (Leinweber and Gruber, 2013; Raimonet and Cloern, 2017). In Tillamook Bay, this pattern was observed across consecutive years. These relationships are particularly clear in deep waters in Santa Monica Bay, while the relationships are more variable in nearshore shallower waters. The upwelling signal in West Coast water bodies was also recorded in the seasonal patterns of pCO_{20bs}, pCO_{2thermal} and pCO_{2bio/hydro}. This signal was particularly evident in Santa Monica Bay and San Francisco Bay, but was also observed in Tillamook Bay (Figures 8, 13; Supplementary Figures 19 – 22). Overall, the variability in $pCO_{2thermal}$ within these West Coast water bodies is diminished compared to seasonal pCO_{2thermal} variability in GOM and East Coast water bodies due to relatively small changes in water temperature. In general, pCO_{2bio/hydro} values in Santa Monica Bay and Tillamook Bay were more closely aligned with pCO_{2obs} over the course of the year. In Santa Monica Bay (both deployments), nonthermal controls were particularly important in late winter to early spring. During this period, pCO_{2thermal} values were falling, reaching their lowest annual value well below pCO_{20bs}, presumably due to cold upwelled water. Then, from the summer into the fall, pCO_{2thermal} values gradually increased while pCO_{20bs} continually decreased and tracked more closely with pCO_{2bio/hvdro}. Collectively, these patterns are consistent with a prominent coastal upwelling signature that is rich in CO₂ peaking in the spring. Other studies have found that Santa Monica Bay has maximum pCO_2 and minimum pH in April and May, coinciding with peak upwelling (Leinweber and Gruber, 2013; McLaughlin et al., 2018).

While the seasonal pattern of $pCO_{2thermal}$ is similar across these West Coast sites, the relationship and timing among pCO_{2thermal}, pCO_{2obs}, and pCO_{2bio/hydro} are different. In Tillamook Bay, pCO_{2obs} increased from the winter into the summer, reaching its maximum in late summer-early fall before decreasing, reaching minimum values in the winter (around February). Additionally, instead of pCO_{2thermal} values reaching their minimum when pCO_{20bs} and pCO_{2thermal} were near their maximum, in Tillamook, pCO_{20bs}, pCO_{2bio/hydro}, and pCO_{2thermal} all reach their maximum around the same time, in late summer-early fall. Large variability in pCO_2 , pH, and salinity values were observed in Tillamook Bay, including the lowest and highest pCO_2 values recorded across all the water bodies (all measurements considered). This may be at least in part related to strong tidal forcing in the estuary which could cause extreme mixing. Seasonal patterns were also evident, and Tillamook Bay is also known to experience periods of coastal upwelling, particularly during spring and summer months (Colbert and McManus, 2003). Upwelling of cold, saline rich waters during the summer have also been observed in the nearby Salish Sea (Cai et al., 2021). The upwelling signal in Tillamook Bay was attenuated as compared to Santa Monica Bay and San Francisco Bay. At least some of this attenuation was likely due to the location of the sensor packages in the estuary (close to shore/shallower in Tillamook vs. further offshore, deeper water in Santa Monica

Bay). Tillamook Bay has high freshwater inflow relative to estuarine volume and as a result, the upwelling signal is modified by riverine inputs, *in situ* biological uptake and regeneration processes within the bay (Colbert and McManus, 2003). This is supported by observations of diminished freshwater flow in the summer and increased freshwater flow in the winter driven by seasonal variations in precipitation. As such, coastal upwelling along with freshwater inputs may be playing particularly important roles in controlling pCO_2 variability during these months.

In Casco Bay, peak $pCO_{2thermal}$ values occurred in August and then continuously decreased until January. This pattern was consistent across consecutive years of data collection. pCO_{2obs} values continued to increase over this same period as $pCO_{2thermal}$ was decreasing, reaching their maximum a couple of months later, in October. $pCO_{2bio/hydro}$ values continued to increase from August to October, similar to the pattern of pCO_{2obs} . Collectively, these patterns indicate a significant nonthermal control on pCO_{2obs} characterized by enhanced biological activity in the spring resulting in enhanced removal of CO_2 from the water column and then the return of CO_2 to the water column in the late summer and fall. Only a few months of data are available for Barnegat Bay, therefore seasonal patterns among pCO_2 and the various transformations were difficult to distinguish.

Recent work has shown that observed amplitudes of seasonal pH and pCO2 cycles increase moving landward from coastal ocean to nearshore and estuarine monitoring stations (Fassbender et al., 2018). The dynamic seasonal cycles observed in the estuarine pH and pCO₂ monitoring data discussed here are consistent with this pattern, as well as with other published estuarine datasets (e.g., Hofmann et al., 2011; Wallace et al., 2014; Joesoef et al., 2015; Fairchild and Hales, 2021; Jiang et al., 2021; Torres et al., 2021). This highlights the relatively extreme carbonate chemistry experienced in estuaries compared to open ocean environments, and the continuing need to consider how ongoing coastal acidification will modify the seasonality of carbonate chemistry cycles (Fassbender et al., 2018). This work will be critical to inform our understanding of how coastal acidification manifests in nearshore and estuarine habitats, including how changing seasonality may control the timing of exceedance of water quality and biological thresholds (Pacella et al., 2018; Cai et al., 2021).

Diel Variability in pCO₂ and pH

Shorter-term, diel trends observed in the data are superimposed on the longer-term trends (**Figure 13**). These trends collectively reflect the influence of weather, tides, and biological activity (e.g., productivity, respiration), drivers that have been documented in other coastal systems (e.g., Yates et al., 2007; Dai et al., 2009; Pacella et al., 2018; Shen et al., 2019). For example, DO and pCO_2 relationships can be indicative of photosynthesis and respiration processes and large ranges of pH and DO indicate more photosynthetic activity and potentially nutrient enrichment. When deviations in DO-pH relationships are observed, other factors may be controlling dynamics, such as freshwater input and upwelling. The influence of tidal fluctuations on seawater chemistry is evident in the regular pattern of salinity variations. For example, in Casco Bay and Tampa Bay large diel ranges in salinity (~4 Practical Salinity Units [PSU]) are observed which are likely related to tidal cycles. As expected, in the deeper Santa Monica Bay deployment, the pattern is irregular, and the salinity signal was muted with daily variability on the order of ~ 0.2 PSU; this is because this is an open coast site. Daily patterns of DO and pCO_2 were also evident wherein increases in pCO_2 corresponded to decreases in DO concentrations, and vice versa. This could be related to the relative influence of productivity and respiration throughout the day. The timing of these day/night patterns (intervals) was quite regular in the shallower Casco Bay, Tampa Bay, and Santa Monica Bay (15 m) deployments but less so in the deeper Santa Monica Bay (60 m) deployment. More detailed analyses outside the scope of this study are needed to separate the day-night signal from the diurnal and semidiurnal tidal signal.

CONCLUSION

The United States EPA's National Estuary Programs (NEP) along with their partnering institutions and agencies are expanding the use of autonomous monitoring pCO_2 and pH sensors to evaluate carbonate chemistry in estuarine environments. Analysis of year to multi-year data from seven NEPs indicate that the sensors effectively captured key parameter measurements enabling an initial characterization of daily to seasonal trends in carbonate chemistry across a range of environmental settings. This study documented extreme variability in coastal acidification data within, and across, a range of coastal systems. Interestuary annual pCO2 and pH variability up to 1,263 µatm (Tillamook Bay) and 1.16 pH units (Casco Bay) respectively were observed. Across all water bodies, temperature and pCO_2 were, in general, lower in cooler, winter months and higher in warmer, summer months; pCO2 and pH values were also more variable in warmer months. As expected, clear coastal upwelling signals were observed in the monitoring data from Santa Monica Bay, San Francisco Bay, and Tillamook Bay. Evidence of both thermal and nonthermal controls on pCO_2 were observed in these systems (i.e., seasonal net community productivity, tidal fluctuations, and freshwater inflows). Evidence of tidal control and day-night changes (e.g., daily temperature variability and diurnal variability in the relative contribution of photosynthesis and respiration) were observed. The relative influence of these drivers should be evaluated as more data becomes available.

Acidification is occurring globally but is a particular concern in coastal zones where local amplifiers such as upwelling, eutrophication, and freshwater inflows combined with human activities exist. The buffering capacity in coastal systems varies as a result of multiple environmental factors. Continued high-resolution monitoring will be critical for understanding how these coastal systems are changing and for identifying and quantifying the factors that contribute to coastal acidification, classifying the susceptibility of these water bodies to the impacts of acidification, and determining which mitigation and adaptive management strategies could be most impactful.

We encourage the continued collection of long-term coastal acidification monitoring data in the system discussed here and expansion of monitoring to other coastal systems. As additional data become more available and patterns of water quality parameter associations become clearer over time, management bodies may use the information to inform management decisions for their estuaries. Recommended next steps could include a more in-depth analysis of the various drivers of acidification in each of these coastal systems through integration of high-resolution monitoring data with other types of data, such as precipitation data, estimates of nutrient inputs, freshwater flows, and upwelling models to inform acidification drivers.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The original data described in this manuscript can be accessed at NOAA's National Centers for Environmental Information (NCEI) at https://doi.org/10.25921/xg33-1n83. Data generated during this study for Tampa Bay are available as a USGS data release (Yates et al., 2019): https://coastal.er.usgs.gov/data-release/doi-P9BAFC7L/.

AUTHOR CONTRIBUTIONS

NR and HG conceived the research. All authors performed the analyses. NR wrote the manuscript with substantial input from all authors.

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REFERENCES

- Bakker, C. E., Pfeil, B., Landa, C. S., Metzl, N., O'Brien, K. M., Olsen, A., et al. (2016). A multi-decade record of high quality fCO₂ data in version 3 of the Surface Ocean CO₂ Atlas (SOCAT). *Earth Syst. Sci. Data* 8, 383–413. doi: 10.5194/essd-8-383-2016
- Barton, A., Hales, B., Waldbusser, G. G., Langdon, C., and Feely, R. A. (2012). The Pacific oyster, *Crassostrea gigasm* shows negative correlation to naturally elevated carbon dioxide levels: Implications for near-term ocean acidification effects. *Limnol. Oceangr* 57, 698–710. doi: 10.4319/lo.2012.57.3.0698
- Barton, A., Waldbusser, G. G., Feely, R. A., Weisberg, S. B., Newton, J. A., Hales, B., et al. (2015). Impacts of coastal acidification on the Pacific Northwest shellfish industry and adaptation strategies implemented in response. *Oceanography* 28, 146–159. doi: 10.5670/oceanog.2015.38
- Baumann, H., and Smith, E. K. (2017). Quantifying metabolically driven pH and oxygen fluctuations in US nearshore habitats at diel to interannual time scales. *Estuar. Coasts* 41, 1102–1117. doi: 10.1007/s12237-017-0321-3
- Cai, W., Xu, Y., Feely, R. A., Wanninkhof, R., Jönsson, B., Alin, S. R., et al. (2020). Controls on surface water carbonate chemistry along North American ocean margins. *Nat. Commun.* 11:2691. doi: 10.1038/s41467-020-16530-z
- Cai, W., Feely, R. A., Testa, J. M., Li, M., Evans, W., Alin, S. R., et al. (2021). Natural and anthropogenic drivers of acidification in large estuaries. *Ann. Rev. Mar. Sci.* 13, 23–55.
- Chan, F., Boehm, A. B., Barth, J. A., Chornesky, E. A., Dickson, A. G., Feely, R. A., et al. (2016). *The West Coast Ocean Acidification and Hypoxia Science Panel: Major Findings, Recommendations, and Actions.* Oakland, CA: California Ocean Science Trust.
- Clements, J. C., and Chopin, T. (2017). Ocean acidification and marine aquaculture in North America: potential impacts and mitigation strategies. *Rev. Aquacult.* 9, 326–341. doi: 10.1111/raq.12140
- Colbert, D., and McManus, J. (2003). Nutrient biogeochemistry in an upwellinginfluenced estuary of the Pacific Northwest (Tillamook Bay, Oregon, USA). *Estuaries* 26, 1205–1219. doi: 10.1007/bf02803625
- Dai, M., Lu, Z., Zhai, W., Chen, B., Cao, Z., Zhou, K., et al. (2009). Diurnal variations of surface seawater CO2 in contrasting coastal environments. *Limnol. Oceanogr.* 54, 735–745. doi: 10.4319/lo.2009.54.3.0735
- DeGrandpre, M. D., Olbu, G. J., Beatty, C. M., and Hammar, T. R. (2002). Air–sea CO₂ fluxes on the US Middle Atlantic Bight. *Deep. Res. Part II* 49, 4355–4367. doi: 10.1016/s0967-0645(02)00122-4
- Doney, S. C., Fabry, V. J., Feely, R. A., and Kleypas, J. A. (2009). Ocean acidification: the other CO₂ problem. *Annu. Rev. Mar. Sci.*, 1, 169–192.
- Dorey, N., Lancon, P., Thorndyke, M., and Dupont, S. (2013). Assessing physiological tipping point of sea urchin larvae exposed to a broad range of pH. *Glob. Change Biol.* 19, 3355–3367. doi: 10.1111/gcb. 12276
- Ekstrom, J. A., Suatoni, L., Cooley, S. R., Pendleton, L. H., Waldbusser, G. G., Cinner, J. E., et al. (2015). Vulnerability and adaptation of US shellfisheries to ocean acidification. *Nat. Clim. Change* 5, 207–214. doi: 10.1038/nclimate2508
- EPA (2021). Measuring Coastal Acidification Using in Situ Sensors in the National Estuary Program. Washington, DC, Document No. EPA-842-R-21001.

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- Fabry, V. J., Seibel, B. A., Feely, R. A., and Orr, J. C. (2008). Impacts of ocean acidification on marine fauna and ecosystem processes. *ICES J. Mar. Sci* 65, 414–432. doi: 10.1093/icesjms/fsn048
- Fairchild, W., and Hales, B. (2021). High-Resolution Carbonate System Dynamics of Netarts Bay, OR From 2014 to 2019. Front. Mar. Sci 7:590236. doi: 10.3389/ fmars.2020.590236
- Fassbender, A. J., Rodgers, K. B., Palevsky, H. I., and Sabine, C. L. (2018). Seasonal asymmetry in the evolution of surface ocean pCO₂ and pH thermodynamic drivers and the influence on sea-air CO2 flux. *Glob. Biogeochem. Cycles* 32, 1476–1497. doi: 10.1029/2017gb005855
- Feely, R. A., Alin, S. R., Carter, B., Bednarsek, N., Hales, B., Chan, F., et al. (2016). Chemical and biological impacts of ocean acidification along the west coast of North America. *Estuar. Coast Shelf Sci.* 183, 260–270. doi: 10.1016/j.ecss.2016. 08.043
- Feely, R. A., Alin, S. R., Newton, J., Sabine, C. L., Warner, M., Devol, A., et al. (2010). The combined effects of ocean acidification, mixing, and respiration on pH and carbonate saturation in an urbanized estuary. *Estuar. Coast. Shelf Sci.* 88, 442–449. doi: 10.1016/j.ecss.2010.05.004
- Feely, R. A., Okazaki, R. R., Cai, W. J., Bednaršek, N., Alin, S. R., Byrne, R. H., et al. (2018). The combined effects of acidification and hypoxia on pH and aragonite saturation in the coastal waters of the California current ecosystem and the northern Gulf of Mexico. *Cont. Shelf Res.* 152, 50–60. doi: 10.1016/j.csr.2017. 11.002
- Feely, R. A., Sabine, C. L., Hernandez-Ayon, J. M., Ianson, D., and Hales, B. (2008). Evidence for upwelling of corrosive 'acidified' water onto the continental shelf. *Science* 320, 1490–1492. doi: 10.1126/science.1155676
- Gazeau, F., Parker, L. M., Comeau, S., Gattuso, J.-P., O'connor, W. A., Martin, S., et al. (2013). Impacts of ocean acidification on marine shelled molluscs. *Mar. Biol.* 160, 2207–2245. doi: 10.1007/s00227-013-2219-3
- Gledhill, D. K., White, M. M., Salisbury, J., Thomas, H., Mlsna, I., Liebman, C. J., et al. (2015). Ocean and coastal acidification off New England and Nova Scotia. *Oceanography* 8, 182–197. doi: 10.5670/oceanog.2015.41
- Hofmann, G. E., Smith, H. E., Johnson, K. S., Send, W., Levin, L. A., Micheli, F., et al. (2011). High-frequency dynamics of ocean pH: a multi-ecosystem comparison. *PLoS One* 6:e2898. doi: 10.1371/journal.pone.0028983
- Hu, X., Beseres Pollack, J., McCutcheon, M. R., Montagna, P., and Ouyang, Z. (2015). Long-term alkalinity decrease and acidification of estuaries in Northwestern Gulf of Mexico. *Environ. Sci. Technol.* 49, 3401–3409. doi: 10. 1021/es505945p
- Huang, W.-J., Cai, W.-J., Wang, Y., Lohrenz, S. E., and Murrell, M. C. (2015). The carbon dioxide system on the Mississippi River-dominated continental shelf in the northern Gulf of Mexico: 1. Distribution and air-sea CO2 flux. *J. Geophys. Res. Ocean* 120, 1429–1445. doi: 10.1002/2014jc010498
- Jiang, L., Carter, B. R., Feely, R. A., Lauvse, S. K., and Olsen, A. (2019). Surface ocean pH and buffer capacity: past, present and future. Sci. Rep. 9:18624.
- Jiang, L.-Q., Feely, R. A., Wanninkhof, R., Greeley, D., Barbero, L., Alin, S., et al. (2021). Coastal Ocean Data Analysis Product in North America (CODAP-NA) – an internally consistent data product for discrete inorganic carbon, oxygen, and nutrients on the North American ocean margins. *Earth Syst. Sci. Data* 13, 2777–2799. doi: 10.5194/essd-13-2777-2021

- Joesoef, A., Huang, W.-J., Gao, Y., and Cai, W.-J. (2015). Air-water fluxes and sources of carbon dioxide in the Delaware Estuary: spatial and seasonal variability. *Biogeosciences* 12, 6085-6101. doi: 10.5194/bg-12-6085-2015
- Kelly, R. P., Foley, M. M., Fisher, W. S., Feely, R. A., Halpern, B. S., Waldbusser, G. G., et al. (2011). Mitigating local causes of ocean acidification with existing laws. *Science* 332, 1036–1037. doi: 10.1126/science.1203815
- Kroeker, K. J., Kordas, R. L., Crim, R., Hendriks, I. E., Ramajo, L., Singh, G. S., et al. (2013). Impacts of ocean acidification on marine organisms: quantifying sensitivities and interaction with warming. *Glob. Change Biol.* 19, 1884–1896. doi: 10.1111/gcb.12179
- Leinweber, A., and Gruber, N. (2013). Variability and trends of ocean acidification in the Southern California Current System: a time series from Santa Monica Bay. J. Geophys. Res. Oceans 118, 3622–3633. doi: 10.1002/jgrc.20259
- Mabardy, R. A., Waldbusser, G. G., Conway, F., and Olsen, C. S. (2015). Perception and response of the U.S. west coast shellfish industry to ocean acidification: the voice of the canaries in the coal mine. *J. Shellfish Res.* 34, 207–214.
- McLaughlin, K., Nezlin, N. P., Weisberg, S. B., Dickson, A. G., Ashley, J., Booth, T., et al. (2018). Seasonal patterns in aragonite saturation state on the southern California continental shelf. *Cont. Shelf Res.* 167, 77–86. doi: 10.1016/j.csr.2018. 07.009
- Millero, F. J. (1995). Thermodynamics of the carbon dioxide system in the oceans. *Geochim. Cosmochim. Acta* 59, 661–677. doi: 10.1016/0016-7037(94)00354-0
- Northcott, D., Sevadjian, J., Sancho-Gallegos, D. A., Wahl, C., Friederich, J., and Chavez, F. P. (2019). Impacts of urban carbon dioxide emissions on sea-air flux and ocean acidification in nearshore waters. *PLoS One* 14:e0214403. doi: 10.1371/journal.pone.0214403
- Orr, J. C., Epitalon, J.-M., Dickson, A. G., and Gattuso, J.-P. (2018). Routine uncertainty propagation for the marine carbon dioxide system. *Mar. Chem.* 207, 84–107. doi: 10.1016/j.marchem.2018.10.006
- Pacella, S. R., Brown, C. A., Waldbusser, G. G., Labiosa, R. G., and Hales, B. (2018). Seagrass habitat metabolism increases short-term extremes and long-term offset of CO2 under future ocean acidification. *Proc. Natl. Acad. Sci. U.S.A.* 115, 3870–3875. doi: 10.1073/pnas.1703445115
- Pecquet, A., Dorey, N., and Chan, K. Y. K. (2017). Ocean acidification increases larval swimming speed and has limited effects on spawning and settlement of a robust fouling bryozoan, Bugula neritina. *Mar. Pollut. Bull.* 124, 903–910. doi: 10.1016/j.marpolbul.2017.02.057
- Pörtner, H.-O. (2008). Ecosystem effects of ocean acidification in times of ocean warming: a physiologist's view. *Mar. Ecol. Prog. Ser* 373, 203–217. doi: 10.3354/ meps07768
- Raimonet, M., and Cloern, J. E. (2017). Estuary–ocean connectivity: fast physics, slow biology. *Glob. Change Biol.* 23, 2345–2357. doi: 10.1111/gcb.13546
- Rheuban, J. E., Doney, S. C., McCorkle, D. C., and Jakuba, R. W. (2019). Quantifying the effects of nutrient enrichment and freshwater mixing on coastal ocean acidification. *J. Geophys. Res. Oceans* 124, 9085–9100. doi: 10.1029/ 2019jc015556
- Salisbury, J., Green, M., Hunt, C., and Campbell, J. (2008). Coastal acidification by rivers: a threat to shellfish? *EOS* 89:513. doi: 10.1029/2008eo500001
- Sastri, A. R., Christin, J. R., Achtenberg, E. P., Atamanchuk, D., Buck, J. J. H., Bresnahan, P., et al. (2019). Perspectives on in situ sensors for ocean acidification research. *Front. Mar. Sci.* 6:653. doi: 10.3389/fmars.2019.00653
- Shadwick, E. H., Trull, T. W., Tilbrook, B., Sutton, A. J., Schulz, E., and Sabine, C. L. (2015). Seasonality of biological and physical controls on surface ocean CO₂ from hourly observations at the Southern Ocean Time Series site south of Australia. *Glob. Biogeochem. Cycles* 29, 223–238. doi: 10.1002/2014GB004906
- Shen, C., Testa, J. M., Li, M., Cai, W.-J., Waldbusser, G. G., Ni, W., et al. (2019). Controls on carbonate system dynamics in a coastal plain estuary: a modeling study. J. Geophys. Res. Biogeosci 124, 61–78. doi: 10.1029/2018jg004802
- Strong, A. L., Kroeker, K. J., Teneva, L. T., Mease, L. A., and Kelly, R. P. (2014). Ocean Acidification 2.0: managing our changing coastal ocean chemistry. *BioScience* 64, 581–592. doi: 10.1093/biosci/biu072
- Sutton, A. J., Feely, R. A., Maenner-Jones, S., Musielwicz, S., Osborne, J., Dietrich, C., et al. (2019). Autonomous seawater pCO₂ and pH time series from 40 surface buoys and the emergence of anthropogenic trends. *Earth Syst. Sci. Data* 11, 421–439. doi: 10.5194/essd-11-421-2019
- Sutton, A. J., Sabine, C. L., Feely, R. A., Cai, W.-J., Cronin, M. F., McPhaden, M. J., et al. (2016). Using present-day observations to detect when anthropogenic change forces surface ocean carbonate chemistry outside preindustrial bounds. *Biogeosciences* 13, 5065–5083. doi: 10.5194/bg-13-5065-2016

- Sutton, A. J., Sabine, C. L., Maenner-Jones, S., Lawrence-Slavas, N., Meinig, C., Feely, R. A., et al. (2014). A high-frequency atmospheric and seawater pCO2 data set from 14 open-ocean sites using a moored autonomous system. *Earth Syst. Sci. Data* 6, 353–366. doi: 10.5194/essd-6-353-2014
- Takahashi, T., Sutherland, S. C., Chipman, D. W., Goddard, J. G., Ho, C., Newberger, T., et al. (2014). Climatological distributions of pH, pCO₂, total CO₂, alkalinity, and CaCO₃ saturation in the global surface ocean, and temporal changes at selected locations. *Mar. Chem* 164, 95–125. doi: 10.1016/j.marchem. 2014.06.004
- Takahashi, T., Sutherland, S. C., Sweeney, C., Poisson, A., Metzl, N., Tilbrook, B., et al. (2002). Global air-sea CO2 flux based on climatological surface ocean pCO2 and seasonal biological and temperature effects. *Deep Sea Res.* 49, 1601–1622. doi: 10.1016/s0967-0645(02)00003-6
- Talmage, S. C., and Gobler, C. J. (2010). Effects of past, present, and future ocean carbon dioxide concentrations on the growth and survival of larval shellfish. *Proc. Natl. Acad. Sci. U.S.A.* 40, 17246–17251. doi: 10.1073/pnas.0913804107
- Torres, O., Kwiatkowski, L., Sutton, A. J., Dorey, N., and Orr, J. C. (2021). Characterizing mean and extreme diurnal variability of ocean CO₂ system variables across marine environments. *Geophys. Res. Lett.* 48:e2020GL090228. doi: 10.1029/2020GL090228
- Waldbusser, G. G., and Salisbury, J. E. (2014). Ocean acidification in the coastal zone from an organism's perspective: multiple system parameters, frequency domains, and habitats. *Annu. Rev. Mar. Sci.* 6, 221–247. doi: 10.1146/annurevmarine-121211-172238
- Waldbusser, G. G., Hales, B., Langdon, C. J., Haley, B. A., Schrader, P., Brunner, E. L., et al. (2015). Ocean acidification has multiple modes of action on bivalve larvae. *PLoS One* 10:e0128376. doi: 10.1371/journal.pone.0128376
- Wallace, R. B., Baumann, H., Grear, J. S., Aller, R. C., and Gobler, C. J. (2014). Coastal ocean acidification: the other eutrophication problem. *Estuar. Coast. Shelf Sci* 148, 1–13. doi: 10.1016/j.ecss.2014.05.027
- Wanninkhof, R., Barbero, L., Byrne, R., Cai, W.-J., Huang, W.-J., Zhang, J.-Z., et al. (2015). Ocean acidification along the Gulf Coast and East Coast of the USA. *Cont. Shelf Res.* 98, 54–71. doi: 10.1016/j.csr.2015. 02.008
- Xue, L., Cai, W.-J., Jiang, L.-Q., and Wei, Q. (2021). Why are surface ocean pH and CaCO3 saturation state often out of phase in spatial patterns and seasonal cycles? *Glob. Biogeochem. Cycles* 35:e2021GB006949.
- Yao, H., and Hu, X. (2017). Responses of carbonate system and CO₂ flux to extended drought and intense flooding in a semiarid subtropical estuary. *Limnol. Oceanogr* 62, S112–S130.
- Yates, K. K., Dufore, C., Smiley, N., Jackson, C., and Halley, R. B. (2007). Diurnal variation of oxygen and carbonate system parameters in Tampa Bay and Florida Bay. *Mar. Chem.* 104, 110–124. doi: 10.1016/j.marchem.2006.12.008
- Yates, K. K., Moore, C. S., and Lemon, M. K. (2019). Time Series Of Autonomous Carbonate System Parameter Measurements In (ver. 2.0, August 2019). Middle Tampa Bay, FL: U.S. Geological Survey data release.

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