

1 **Temporal variability and driving factors of the carbonate system in the Aransas**
2 **Ship Channel, TX, USA: A time-series study**

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4 **Melissa R. McCutcheon¹, Hongming Yao^{1,#}, Cory J. Staryk¹, Xinping Hu¹**

5 ¹Harte Research Institute for Gulf of Mexico Studies, Texas A&M University – Corpus
6 Christi, TX 78412, USA

7 [#] current address: Shenzhen Engineering Laboratory of Ocean Environmental Big Data
8 Analysis and Application, Shenzhen Institute of Advanced Technology, Chinese
9 Academy of Sciences, Shenzhen 518055, China

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12 *Correspondence to:* Melissa R. McCutcheon (melissa.mccutcheon@tamucc.edu)

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16 **Abstract**

17 The coastal ocean is affected by an array of co-occurring biogeochemical and
18 anthropogenic processes, resulting in substantial heterogeneity in water chemistry,
19 including carbonate chemistry parameters such as pH and partial pressure of CO₂ (*p*CO₂).
20 To better understand coastal and estuarine acidification and air-sea CO₂ fluxes, it is
21 important to study baseline variability and driving factors of carbonate chemistry. Using
22 both discrete bottle sample collection (2014-2020) and hourly sensor measurements
23 (2016-2017), we explored temporal variability, from diel to interannual scales, in the
24 carbonate system (specifically pH and *p*CO₂) at the Aransas Ship Channel located in
25 northwestern Gulf of Mexico. Using other co-located environmental sensors, we also
26 explored the driving factors of that variability. Both sampling methods demonstrated
27 significant seasonal variability at the location, with highest pH (lowest *p*CO₂) in the
28 winter and lowest pH (highest *p*CO₂) in the summer. Significant diel variability was also
29 evident from sensor data, but the time of day with elevated *p*CO₂/depressed pH was not
30 consistent across the entire monitoring period, sometimes reversing from what would be
31 expected from a biological signal. Though seasonal and diel fluctuations were smaller
32 than many other areas previously studied, carbonate chemistry parameters were among
33 the most important environmental parameters to distinguish between time of day and
34 between seasons. It is evident that temperature, biological activity, freshwater inflow, and
35 tide level (despite the small tidal range) are all important controls on the system, with
36 different controls dominating at different time scales. The results suggest that the
37 controlling factors of the carbonate system may not be exerted equally on both pH and
38 *p*CO₂ on diel timescales, causing separation of their diel or tidal relationships during

39 certain seasons. Despite known temporal variability on shorter timescales, discrete
40 sampling was generally representative of the average carbonate system and average air-
41 sea CO₂ flux on a seasonal and annual basis when compared with sensor data.

42 **1. Introduction**

43 Coastal waters, especially estuaries, experience substantial spatial and temporal
44 heterogeneity in water chemistry—including carbonate chemistry parameters such as pH
45 and partial pressure of CO₂ (*p*CO₂)—due to the diversity of co-occurring biogeochemical
46 and anthropogenic processes (Hofmann et al., 2011; Waldbusser and Salisbury, 2014).
47 Carbonate chemistry is important because an addition of CO₂ acidifies seawater, and
48 acidification can negatively affect marine organisms (Barton et al., 2015; Bednaršek et
49 al., 2012; Ekstrom et al., 2015; Gazeau et al., 2007; Gobler and Talmage, 2014).
50 Additionally, despite the small surface area of coastal waters relative to the global ocean,
51 coastal waters are recognized as important contributors in global carbon cycling (Borges,
52 2005; Cai, 2011; Laruelle et al., 2018).

53 While carbonate chemistry, acidification, and air-sea CO₂ fluxes are relatively
54 well studied and understood in open ocean environments, large uncertainties remain in
55 coastal environments. Estuaries are especially challenging to fully understand because of
56 the heterogeneity between and within estuaries that is driven by diverse processes
57 operating on different time scales such as river discharge, nutrient and organic matter
58 loading, stratification, and coastal upwelling (Jiang et al., 2013; Mathis et al., 2012). The
59 traditional sampling method for carbonate system characterization involving discrete
60 water sample collection and laboratory analysis is known to lead to biases in average
61 *p*CO₂ and CO₂ flux calculations due to daytime sampling that neglects to capture diel

62 variability (Li et al., 2018). Mean diel ranges in pH can exceed 0.1 unit in many coastal
63 environments, and especially high diel ranges (even exceeding 1 pH unit) have been
64 reported in biologically productive areas or areas with higher mean $p\text{CO}_2$ (Challener et
65 al., 2016; Cyronak et al., 2018; Schulz and Riebesell, 2013; Semesi et al., 2009; Yates et
66 al., 2007). These diel ranges can far surpass the magnitude of the changes in open ocean
67 surface waters that have occurred since the start of the industrial revolution and rival
68 spatial variability in productive systems, indicating their importance for a full
69 understanding of the carbonate system.

70 Despite the need for high-frequency measurements, sensor deployments have
71 been limited in estuarine environments (especially compared to their extensive use in the
72 open ocean) because of the challenges associated with ~~varying conditions~~highly variable
73 salinities, biofouling, and sensor drift (Sastri et al., 2019). Carbonate chemistry
74 monitoring in the Gulf of Mexico (GOM), has been relatively minimal compared to the
75 United States east and west coasts. The GOM estuaries currently have less exposure to
76 concerning levels of acidification than other estuaries because of their high temperatures
77 (causing water to hold less CO_2 and support high productivity year-round) and often
78 suitable river chemistries (i.e., relatively high buffer capacity) (McCutcheon et al., 2019;
79 Yao et al., 2020). However, respiration-induced acidification is present in both the open
80 GOM (e. g., subsurface water influenced by the Mississippi River Plume and outer shelf
81 region near the Flower Garden Banks National Marine Sanctuary) and GOM estuaries,
82 and most estuaries in the northwestern GOM have also experienced long-term
83 acidification (Cai et al., 2011; Hu et al., 2018, 2015; Kealoha et al., 2020; McCutcheon et
84 al., 2019; Robbins and Lisle, 2018). This ~~known acidification~~evidence of acidification as

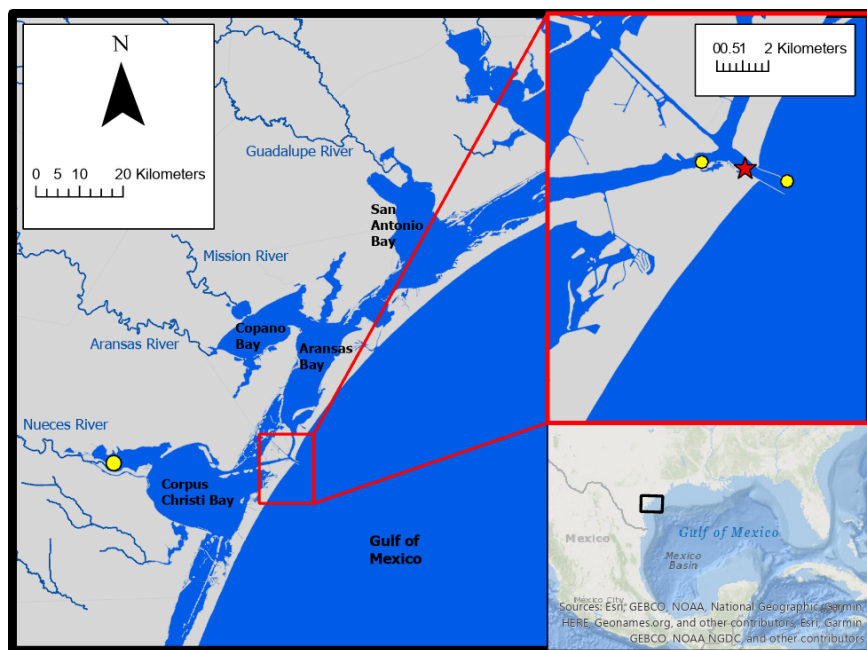
85 well as the relatively high CO₂ efflux from the estuaries of the northwest GOM illustrates
86 the necessity to study the baseline variability and driving factors of carbonate chemistry
87 in the region. In this study, we explored temporal variability in the carbonate system in
88 Aransas Ship Channel (ASC)—a tidal inlet where the lagoonal estuaries meet the coastal
89 waters in a semi-arid region of the northwestern GOM—using both discrete bottle sample
90 collection and hourly sensor measurements, and we explored the driving factors of that
91 variability using data from other co-located environmental sensors. The characterization
92 of carbonate chemistry and consideration of regional drivers can provide context to
93 acidification and its impacts and improved estimates of air-sea CO₂ fluxes.

94 **2. Materials and Methods**

95 *2.1 Location*

96 Autonomous sensor monitoring and discrete water sample collections for
97 laboratory analysis of carbonate system parameters were performed in ASC (located at
98 27°50'17"N, 97°3'1"W). ASC is one of the few permanent tidal inlets that intersect a
99 string of barrier islands and connect the GOM coastal waters with the lagoonal estuaries
100 in the northwest GOM (Fig. 1). ASC provides the direct connection between the
101 northwestern GOM and the Mission-Aransas Estuary (Copano and Aransas Bays) to the
102 north and Nueces Estuary (Nueces and Corpus Christi Bays) to the south (Fig. 1). The
103 region is microtidal, with a small tidal range relative to many other estuaries, ranging
104 from ~ 0.6 m tides on the open coast to less than 0.3 m in upper estuaries (Montagna et
105 al., 2011). Mission-Aransas Estuary (MAE) is fed by two small rivers, the Mission (1787
106 km² drainage basin) and Aransas (640 km² drainage basin) Rivers
107 (<http://waterdata.usgs.gov/>), which both experience low base flows punctuated by

108 periodic high flows during storm events. MAE has an average residence time of one year
109 (Solis and Powell, 1999), so there is a substantial lag between time of rainfall and
110 riverine delivery to ASC in the lower estuary. A significant portion of riverine water
111 flowing into Aransas Bay originates from the larger rivers further northeast on the Texas
112 coast via the Intracoastal Waterway (i.e., Guadalupe River (26,625 km² drainage basin)
113 feeds San Antonio Bay and has a much shorter residence time of nearly 50 days) (Solis
114 and Powell, 1999; USGS, 2001).



115
116 **Figure 1.** Study area. The location of monitoring in the Aransas Ship Channel (red star)
117 and the locations of NOAA stations used for wind data (yellow circles) are shown.

118
119 *2.2 Continuous Monitoring*

120 Autonomous sensor monitoring (referred to throughout as continuous monitoring)
121 of pH and $p\text{CO}_2$ was conducted from Nov. 8, 2016 to Aug. 23, 2017 at the University of

122 Texas Marine Science Institute's research pier in ASC. Hourly pH data were collected
123 using an SAtlantic® SeaFET pH sensor (on total pH scale) and hourly $p\text{CO}_2$ data were
124 collected using a Sunburst® SAMI-CO₂. The pH and $p\text{CO}_2$ sensors were placed in a
125 flowthrough system that received surface water from ASC using a time-controlled
126 diaphragm pump -prior to each measurement. Hourly temperature and salinity data were
127 measured by a YSI® 600OMS V2 sonde. All hourly data were single measurements taken
128 on the hour. The average difference between sensor pH and discrete quality assurance
129 samples measured spectrophotometrically in the lab was used to establish a correction
130 factor (-0.05) across the entire sensor pH dataset. Note, this correction scheme was not
131 ideal (Bresnahan et al., 2014) although less rigorous correction based on sensor and
132 discrete pH values has also been used (Shadwick et al. 2019). Nevertheless, the overall
133 good agreement between discrete and corresponding sensor pH values during the
134 deployment period suggested that the SeaFET sensor remained stable. It is also worth
135 noting that our monitoring setup remained free from biofouling during the 10-month
136 period. We attribute this to the deployment design in which the high frequency movement
137 of the pumping mechanisms in the diaphragm pump must have eliminated the influence
138 of animal larvae. The average difference between sensor pH and discrete quality
139 assurance samples measured spectrophotometrically in the lab was used to establish a
140 correction (-0.05) based on a single calibration point across the entire sensor pH dataset
141 (Bresnahan et al., 2014). See supplemental materials for additional sensor deployment
142 and quality assurance information.

143 *2.3 Discrete Sample Collection and Sample Analysis*

144 Long-term monitoring via discrete water sample collection was conducted at ASC
145 from May 2, 2014 to February 25, 2020 (in addition to the discrete, quality assurance
146 sample collections). A single, discrete, surface water sample was collected every two
147 weeks during the summer months and monthly during the winter months from a small
148 vessel at a station near (<20 m from) the sensor deployment. Water sample collection
149 followed standard protocol for ocean carbonate chemistry studies (Dickson et al., 2007).
150 Ground glass borosilicate bottles (250 mL) were filled with surface water and preserved
151 with 100 μ L saturated mercury chloride (HgCl_2). Apiezon[®] grease was applied to the
152 bottle stopper, which was then secured to the bottle using a rubber band and a nylon hose
153 clamp.

154 These samples were used for laboratory dissolved inorganic carbon (DIC) and pH
155 measurements. DIC was measured by injecting 0.5 mL of sample into 1 ml 10% H_3PO_4
156 (balanced by 0.5 M NaCl) with a high-precision Kloehe syringe pump. The CO_2 gas
157 produced through sample acidification was then stripped using high-purity nitrogen gas
158 and carried into a Li-Cor infrared gas detector. DIC analyses had a precision of 0.1%.
159 Certified Reference Material (CRM) was used to ensure the accuracy of the analysis
160 (Dickson et al. 2003). For samples with salinity>20, pH was measured using a
161 spectrophotometric method at $25 \pm 0.1^\circ\text{C}$ (Carter et al. 2003) and the Douglas and Byrne
162 (2017) equation. Analytical precision of the spectrophotometric method for pH
163 measurement was ± 0.0004 pH units. A calibrated Orion Ross glass pH electrode was
164 used to measure pH at $25 \pm 0.1^\circ\text{C}$ for samples with salinity<20, and analytical precision
165 was ± 0.01 pH units. All pH values obtained using the potentiometric method were

166 converted to total scale at *in situ* temperature (Millero 2001). Salinity of the discrete
167 samples was measured using a benchtop salinometer calibrated by MilliQ water and a
168 known salinity CRM. For discrete samples, $p\text{CO}_2$ was calculated in CO2Sys for Excel
169 using laboratory-measured salinity, DIC, pH, and *in situ* temperature for calculations.
170 Carbonate speciation calculations were done using Millero (2010) carbonic acid
171 dissociation constants (K_1 and K_2), Dickson (1990) bisulfate dissociation constant, and
172 Uppström (1974) borate concentration.

173 2.4 Calculation of CO_2 fluxes

174 Equation 1 was used for air-water CO_2 flux calculations (Wanninkhof, 1992;
175 Wanninkhof et al., 2009). Positive flux values indicate CO_2 emission from the water into
176 the atmosphere (the estuary acting as a source of CO_2), and negative flux values indicate
177 CO_2 uptake by the water (the estuary acting as a sink for CO_2).

$$178 F = k K_0 (p\text{CO}_{2,w} - p\text{CO}_{2,a}) \quad (1)$$

179 where k is the gas transfer velocity (in m d^{-1}), K_0 (in $\text{mol l}^{-1} \text{atm}^{-1}$) is the solubility
180 constant of CO_2 (Weiss, 1974), and $p\text{CO}_{2,w}$ and $p\text{CO}_{2,a}$ are the partial pressure of CO_2 (in
181 μatm) in the water and air, respectively.

182 We used the wind speed parameterization for gas transfer velocity (k) from Jiang
183 et al. (2008) converted from cm h^{-1} to m d^{-1} , which is thought to be the best estuarine
184 parameterization at this time (Crosswell et al., 2017), as it is a composite of k over
185 several estuaries. The calculation of k requires a windspeed at 10 m above the surface, so
186 windspeeds measured at 3 m above the surface were converted using the power law wind
187 profile (Hsu, 1994; Yao and Hu, 2017). To assess uncertainty, other parameterizations
188 with direct applications to estuaries in the literature were also used to calculate CO_2 flux

189 (Raymond and Cole 2001; Ho et al. 2006). We note that parameterization of k based on
190 solely windspeed is flawed because several additional parameters can contribute to
191 turbulence including turbidity, bottom-driven turbulence, water-side thermal convection,
192 tidal currents, and fetch (Wanninkhof 1992, Abril et al., 2009, Ho et al., 2104, Andersson
193 et al., 2017), however it is currently the best option for this system given the limited
194 investigations of CO₂ flux and contributing factors in estuaries.

195 Hourly averaged windspeed data for use in CO₂ flux calculations were retrieved
196 from the NOAA-controlled Texas Coastal Ocean Observation Network (TCOON;
197 <https://tidesandcurrents.noaa.gov/tcoon.html>). Windspeed data from the nearest TCOON
198 station (Port Aransas Station – located directly in ASC, < 2 km inshore from our
199 monitoring location) was prioritized when data were available. During periods of missing
200 windspeed data at the Port Aransas Station, wind speed data from TCOON’s Aransas
201 Pass Station (< 2 km offshore from monitoring location) were next used, and for all
202 subsequent gaps, data from TCOON’s Nueces Bay Station (~ 40 km away) were used
203 (Fig. 1; additional discussion of flux calculation and windspeed data can be found in
204 supplementary materials). For flux calculations from continuous monitoring data, each
205 hourly measurement of $p\text{CO}_2$ was paired with the corresponding hourly averaged
206 windspeed. For flux calculations from discrete sample data, the $p\text{CO}_2$ calculated for each
207 sampled day was paired with the corresponding daily averaged windspeed (calculated
208 from the retrieved hourly averaged windspeeds).

209 Monthly mean atmospheric $x\text{CO}_2$ data (later converted to $p\text{CO}_2$) for flux
210 calculations were obtained from NOAA’s flask sampling network of the Global
211 Monitoring Division of the Earth System Research Laboratory at the Key Biscayne (FL,

212 USA) station. Global averages of atmospheric $x\text{CO}_2$ were used when Key Biscayne data
213 were unavailable. Each $p\text{CO}_2$ observation (whether using continuous or discrete data)
214 was paired with the corresponding monthly averaged $x\text{CO}_2$ for flux calculations.
215 Additional information and justification are available in supplemental materials.

216 *2.5 Additional data retrieval and data processing to investigate carbonate system*
217 *variability and controls*

218 All reported annual mean values are seasonally weighted to account for
219 disproportional sampling between seasons. However, reported annual standard deviation
220 is associated with the un-weighted, arithmetic mean (Table S1). Temporal variability was
221 investigated in the form of seasonal and diel variability (Tables S1, S2, S3). For seasonal
222 analysis, December to February was considered winter, March to May was considered
223 spring, June to August was considered summer, and September to November was
224 considered fall. It is important to note that the Fall season had much fewer continuous
225 sensor observations than other seasons because of the timing of sensor deployment. For
226 diel comparisons, daytime and nighttime variables were defined as 09:00-15:00 local
227 standard time and 21:00-03:00 local standard time, respectively, based on the 6-hour
228 periods with highest and lowest photosynthetically active radiation (PAR; data from co-
229 located sensor, obtained from the Mission-Aransas National Estuarine Research Reserve
230 (MANERR) at <https://missionaransas.org/science/download-data>). Diel ranges in
231 parameters were calculated (daily maximum minus daily minimum) and only reported for
232 days with the full 24 hours of hourly measurements (176 out of 262 measured days) to
233 ensure that data gaps did not influence the diel ranges (Table S3).

234 Controls on $p\text{CO}_2$ from thermal and non-thermal (i.e., combination of physical
 235 and biological) processes were investigated following Takahashi et al. (2002) over
 236 annual, seasonal, and daily time scales using both continuous and discrete data. Over any
 237 given time period, this method uses the ratio of the ranges of temperature-normalized
 238 $p\text{CO}_2$ ($p\text{CO}_{2,\text{nt}}$, Eq. 2) and the mean annual $p\text{CO}_2$ perturbed by the difference between
 239 mean and observed temperature ($p\text{CO}_{2,\text{t}}$, Eq. 3) to calculate the relative influence of non-
 240 thermal and thermal effects on $p\text{CO}_2$ (T/B, Eq. 4). When calculating annual T/B values
 241 with discrete data, only complete years (sampling from January to December) were
 242 included (2014 and 2020 were omitted). When calculating daily T/B values with
 243 continuous data, only complete days (24 hourly measurements) were included.

$$244 \quad p\text{CO}_{2,\text{nt}} = p\text{CO}_{2,\text{obs}} \times \exp[\delta \times (T_{\text{mean}} - T_{\text{obs}})] \quad (2)$$

$$245 \quad p\text{CO}_{2,\text{t}} = p\text{CO}_{2,\text{mean}} \times \exp[\delta \times (T_{\text{obs}} - T_{\text{mean}})] \quad (3)$$

246 where the value for δ ($0.0411 \text{ } ^\circ\text{C}^{-1}$), which represents average $[\partial \ln p\text{CO}_2 / \partial$
 247 Temperature] from field observations, was taken directly from Yao and Hu (2017), T_{obs} is
 248 the observed temperature, and T_{mean} is the mean temperature over the investigated time
 249 period.

$$250 \quad T/B = \frac{\max(p\text{CO}_{2,\text{thermal}}) - \min(p\text{CO}_{2,\text{thermal}})}{\max(p\text{CO}_{2,\text{non-thermal}}) - \min(p\text{CO}_{2,\text{non-thermal}})} \quad (4)$$

251 Where a T/B greater than one indicates that temperature's control on $p\text{CO}_2$ is greater than
 252 the control from non-thermal factors and a T/B less than one indicates that non-thermal
 253 factors' control on $p\text{CO}_2$ is greater than the control from temperature.

254 Tidal control on parameters was investigated using our continuous monitoring
 255 data and tide level data obtained from NOAA's Aransas Pass Station (the Aransas Pass

256 Station used for windspeed data, < 2 km offshore from monitoring location, Fig. 1) at
257 <https://tidesandcurrents.noaa.gov/waterlevels.html?id=8775241&name=Aransas.%20Aransas%20Pass&state=TX>. Hourly measurements of water level were merged with our
258 sensor data by date and hour. Given that there were gaps in available water level
259 measurements (and no measurements prior to December 20, 2016), the usable dataset was
260 reduced from 6088 observations to 5121 observations and fall was omitted from analyses.
261 To examine differences between parameters during high tide and low tide, we defined
262 high tide as tide level greater than the third quartile tide level value and low tide as a tide
263 level less than the first quartile tide level value.
264

265 Other factors that may exert control on the carbonate system were investigated
266 through parameter relationships. In addition to previously discussed tide and windspeed
267 data, we obtained dissolved oxygen (DO), PAR, turbidity, and chlorophyll fluorescence
268 data from MANERR-deployed environmental sensors that were co-located at our
269 monitoring location (obtained from <https://missionaransas.org/science/download-data>).
270 Given that MANERR data are all measured in the bottom water (>5 m) while our sensors
271 were measuring surface waters, we excluded the observations with significant water
272 column stratification (defined as a salinity difference > 3 between surface water and
273 bottom water) from analyses. Omitting stratified water reduced our continuous dataset
274 from 6088 to 5524 observations (removing 260 winter, 133 spring, 51 summer, and 120
275 fall observations), and omitting observations where there were no MANERR data to
276 determine stratification further reduced the dataset to 4112 observations. Similarly,
277 removing instances of stratification reduced discrete sample data from 104 to 89 surface
278 water observations.

279 *2.6 Statistical Analyses*

280 All statistical analyses were performed in R, version 4.0.3 (R Core Team, 2020).
281 To investigate differences between daytime and nighttime parameter values (temperature,
282 salinity, pH, $p\text{CO}_2$, and CO_2 flux) using continuous monitoring data across the full
283 sampling period and within each season, paired t -tests were used, pairing each respective
284 day's daytime and nighttime values (Table S3). We also used loess models (locally
285 weighted polynomial regression) to identify changes in diel patterns over the course of
286 our monitoring period.

287 Two-way ANOVAs were used to examine differences in parameter means
288 between seasons and between monitoring methods (Table S2). Since there were
289 significant interactions (between season and sampling type factors) in the two-way
290 ANOVAs for each individual parameter (Table S2), differences between seasons were
291 investigated within each monitoring method (one-way ANOVAs) and the differences
292 between monitoring methods were investigated within each season (one-way ANOVAs).
293 For the comparison of monitoring methods, we included both the full discrete sampling
294 data as well as a subset of the discrete sampling data to overlap with the continuous
295 monitoring period (referred to throughout as reduced discrete data or D_C) along with the
296 continuous data. To interpret differences between monitoring methods, a difference in
297 means between the continuous monitoring and discrete monitoring datasets would only
298 indicate that the 10-month period of continuous monitoring was not representative of the
299 5+ year period that discrete samples have been collected, but a difference in means
300 between the continuous data and discrete sample data collected during the continuous
301 monitoring period represents discrepancies between types of monitoring. Post-hoc

302 multiple comparisons (between seasons within sampling types and between sampling
303 types within seasons) were conducted using the Westfall adjustment (Westfall, 1997).

304 Differences in parameters between high tide and low tide conditions were
305 investigated using a two-way ANOVA to model parameters based on tide level and
306 season. In models for each parameter, there was a significant interaction between tide
307 level and season factors (based on $\alpha=0.05$, results not shown), thus t-tests were used
308 (within each season) to examine differences in parameters between high and low tide
309 conditions. Note that fall was omitted from this analysis because tide data were only
310 available at the location beginning December 20, 2016. Sample sizes were the same for
311 each parameter (High tide – winter: 354, spring: 569, summer: 350; Low tide – winter:
312 543, spring: 318, summer: 415).

313 Additionally, to gain insight to carbonate system controls through correlations, we
314 conducted Pearson correlation analyses to examine individual correlations of pH and
315 $p\text{CO}_2$ (both continuous and discrete) with other environmental parameters (Table S5).

316 To better understand overall system variability over different time scales, we used
317 a linear discriminant analysis (LDA), a multivariate statistic that allows dimensional
318 reduction, to determine the linear combination of environmental parameters (individual
319 parameters reduced into linear discriminants, LDs) that allow the best differentiation
320 between day and night as well as between seasons. We included $p\text{CO}_2$, pH, temperature,
321 salinity, tide level, wind speed, total PAR, DO, turbidity, and fluorescent chlorophyll in
322 this analysis. All variables were centered and scaled to allow direct comparison of their
323 contribution to the system variability. The magnitude (absolute value) of coefficients of
324 the LDs (Table 1) represents the relative importance of each individual environmental

325 parameter in the best discrimination between day and night and between seasons, i.e., the
326 greater the absolute value of the coefficient, the more information the associated
327 parameter can provide about whether the sample came from day or night (or winter,
328 spring, or summer). Only one LD could be created for the diel variability (since there are
329 only two classes to discriminate between – day and night). Two LDs could be created for
330 the seasonal variability (since there were three classes to discriminate between – fall was
331 omitted because of the lack of tidal data), but we chose to only report the coefficients for
332 LD1 given that LD1 captured 95.64% of the seasonal variability.

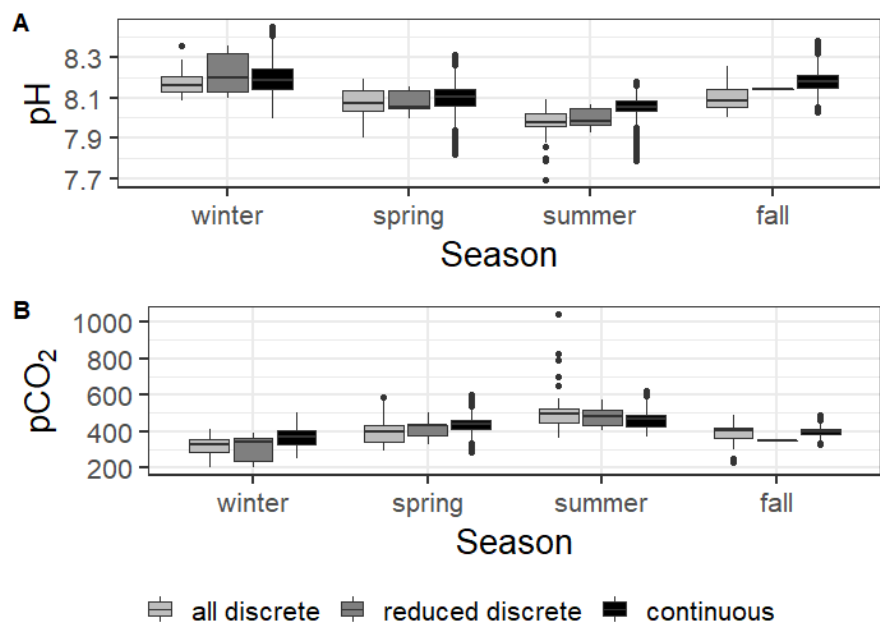
333

334 **3. Results**

335 *3.1 Seasonal variability*

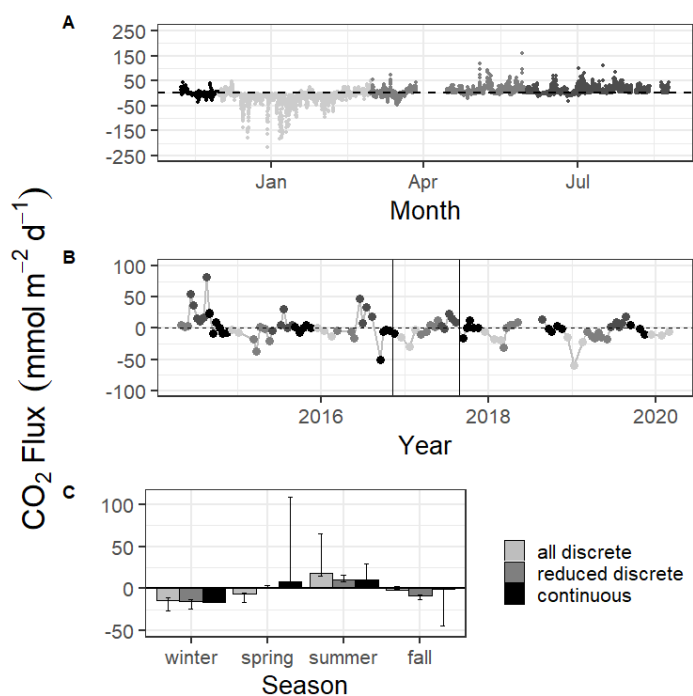
336 Both the continuous and discrete data showed substantial seasonal variability for
337 all parameters (Fig. 2, Tables S1 and S2). All discrete sample results reported here are for
338 the entire 5+ years of monitoring; the subset of discrete sample data that overlaps with
339 the continuous monitoring period will be addressed only in the discussion for method
340 comparisons (Section 4.1.1). Both continuous and discrete data demonstrate significant
341 differences in temperature between each season, with the highest temperature in summer
342 and the lowest in winter (Tables S1 and S2). Mean salinity during sampling periods was
343 highest in the summer and lowest in the fall (Table S1). Significant differences in
344 seasonal salinity occurred between all seasons except spring and winter for continuous
345 data, but only summer differed from other seasons based on discrete data (Tables S1 and
346 S2).

347 Carbonate system parameters also varied seasonally (Fig. 2). For both continuous
 348 and discrete data, winter had the highest seasonal pH (8.19 ± 0.08 and 8.162 ± 0.065 ,
 349 respectively) and lowest seasonal $p\text{CO}_2$ ($365 \pm 44 \mu\text{atm}$ and $331 \pm 39 \mu\text{atm}$,
 350 respectively), while summer had the lowest seasonal pH (8.05 ± 0.06 and 7.975 ± 0.046 ,
 351 respectively) and highest seasonal $p\text{CO}_2$ ($463 \pm 48 \mu\text{atm}$ and 511 ± 108 , respectively)
 352 (Fig. 2, Table S1). All seasonal differences in pH and $p\text{CO}_2$ were significant, except for
 353 the discrete data spring versus fall for both parameters (Table S2).
 354
 355



356
 357 **Figure 2.** Boxplots of seasonal variability in pH and $p\text{CO}_2$ using all discrete data,
 358 reduced discrete data (to overlap with continuous monitoring, Nov. 8 2016 – Aug 23,
 359 2017), and continuous sensor data.
 360

361 Mean CO₂ flux differed by season (Fig. 3, Tables S1 and S2). Both continuous
 362 and discrete data records resulted in net negative CO₂ fluxes during fall and winter
 363 months, with winter being most negative. Both methods reported a net positive flux for
 364 summer, while spring fluxes were positive according to continuous data and negative
 365 according to the 5+ years of discrete data (Fig. 3, Table S1). Annual net CO₂ fluxes were
 366 near zero (Table S1).
 367



368 **Figure 3.** CO₂ flux calculated over the sampling periods from continuous (A) and
 369 discrete (B) data. Gray scale in (A) and (B) denote different seasons. Vertical lines in (B)
 370 denote the time period of continuous monitoring. (C) shows the seasonal mean CO₂ flux.
 371 Error bars represent mean CO₂ flux using Ho (2006) and Raymond and Cole (2001)
 372 windspeed parameterizations.
 373
 374

375 Results of the LDA incorporated carbonate system parameters along with
 376 additional environmental parameters to get a full picture of system variability over
 377 seasonal timescales (Table 1). The most important parameter in system variability that
 378 allowed differentiation between seasons was temperature (Table 1, Seasonal LD1), as
 379 would be expected with the clear seasonal temperature fluctuations (Fig. S1E). The
 380 second most important parameter for seasonal differentiation was chlorophyll, likely
 381 indicating clear seasonal phytoplankton blooms. The carbonate chemistry also played a
 382 critical role in seasonal differentiation, as $p\text{CO}_2$ was the third most important factor
 383 (Table 1).

384 **Table 1.** Coefficients of linear discriminants (LD) from LDA using continuous sensor
 385 data and other environmental parameters. Discriminants for both diel and seasonal
 386 variability shown.

	Seasonal	Diel
	LD1	LD1
Temperature (°C)	-3.53	0.54
Salinity	0.04	0.15
$p\text{CO}_2$ (μatm)	-0.29	-0.16
pH	0.10	0.06
Tide Level (m)	-0.24	0.10
Wind speed (ms^{-1})	0.05	-0.00
Total PAR	-0.07	-2.29
DO (mg L^{-1})	0.09	-0.08
Turbidity	0.15	-0.06
Fluor. Chlorophyll	-0.40	0.14

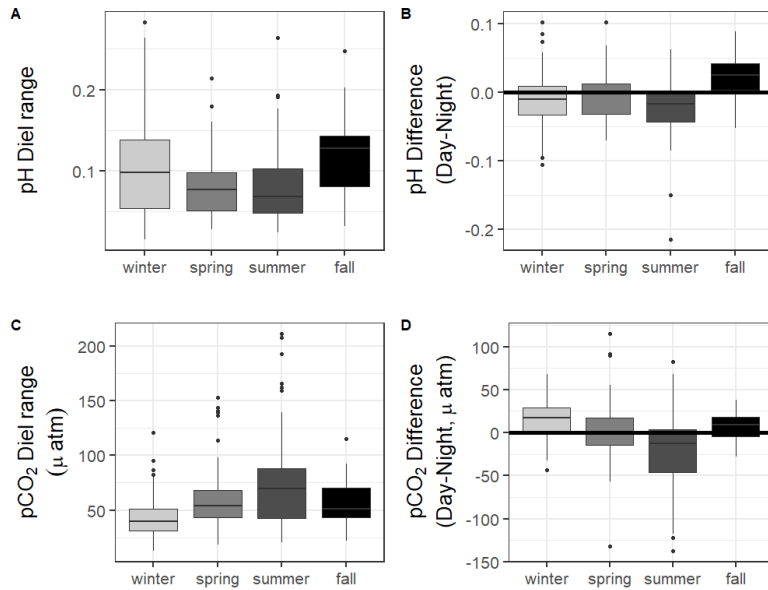
387
 388 *3.2 Diel variability*

389 The 10 months of in-situ continuous monitoring revealed that there was
 390 substantial diel variability in measured parameters (Fig. 4, Table S3). Temperature had a
 391 mean diel range of $1.3 \pm 0.8^\circ\text{C}$ (Table S3). Daytime and nighttime temperature differed
 392 significantly during the summer and fall months, with higher temperatures at night for
 393 both seasons (Table S3). The mean diel range of salinity was 3.4 ± 2.7 (Table S3).
 394 Daytime and nighttime salinity differed significantly during the winter and fall months,

395 with higher salinities at night for both seasons. The mean diel range of pH was $0.09 \pm$
396 0.05 (Table S3). Daytime and nighttime pH differed significantly during the winter,
397 summer, and fall, with nighttime pH significantly higher during summer and winter and
398 lower during fall (Fig. 4, Table S3). The mean diel range of $p\text{CO}_2$ was $58 \pm 33 \mu\text{atm}$ (Fig.
399 4, Table S3). Daytime and nighttime $p\text{CO}_2$ differed significantly during the winter and
400 summer months, with nighttime $p\text{CO}_2$ significantly higher during the summer and lower
401 during the winter (Fig. 4, Table S3). No significant difference in daytime and nighttime
402 DO were observed during any season (Fig. 5F; paired t-tests, winter $p = 0.1573$, spring p
403 $= 0.4877$, summer $p = 0.794$).

404 Loess models that investigated the evolution of day-night difference in parameters
405 revealed that other environmental parameters, including salinity, temperature, and tide
406 level, also had diel patterns that varied over the duration of our continuous monitoring
407 (Fig. 5).

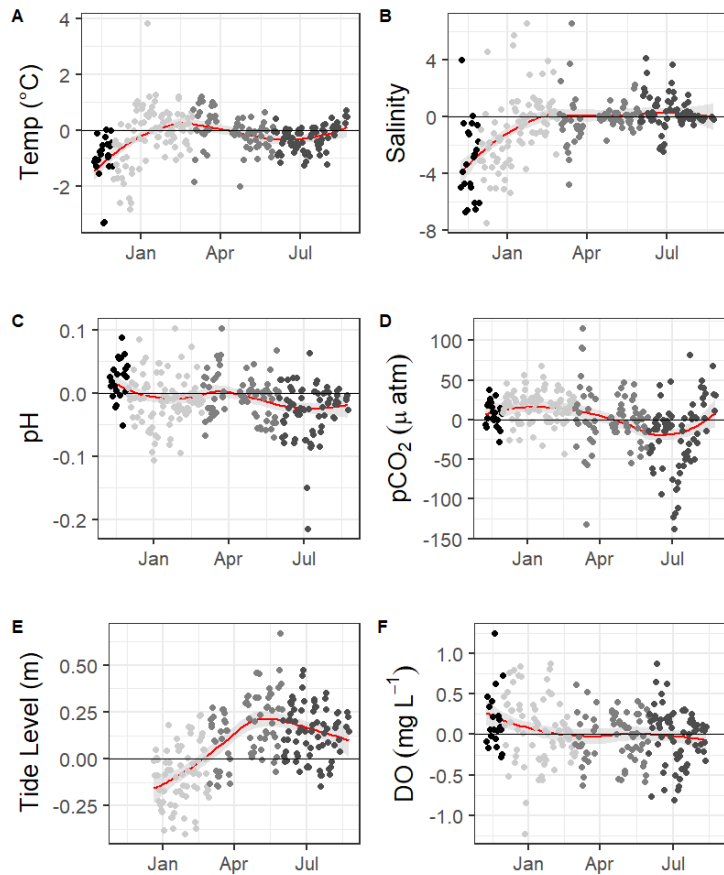
408



409 **Figure 4.** Boxplots of the diel range (maximum minus minimum) and difference in daily
 410 parameter mean daytime minus nighttime measurements for pH and $p\text{CO}_2$ from
 411 continuous sensor data.
 412

413
 414 CO_2 flux also fluctuated on a daily scale, with a mean diel range of 34.1 ± 29.0
 415 $\text{mmol m}^{-2} \text{d}^{-1}$ (Table S3). However, there was not a significant difference in CO_2 flux of
 416 daytime versus nighttime hours for the entire monitoring period or any individual season
 417 based on $\alpha=0.05$ (paired t-test, Table S3).

418
 419



420
 421 **Figure 5.** Loess models (red line) and their confidence intervals (gray bands) showing the
 422 difference in daily daytime mean minus nighttime mean measurements. The gray scale of
 423 the data points represents the four seasons over which data were collected. Data span
 424 from Nov 8, 2016 to Aug 3, 2017, except for the tide data, which began December 20,
 425 2016.
 426

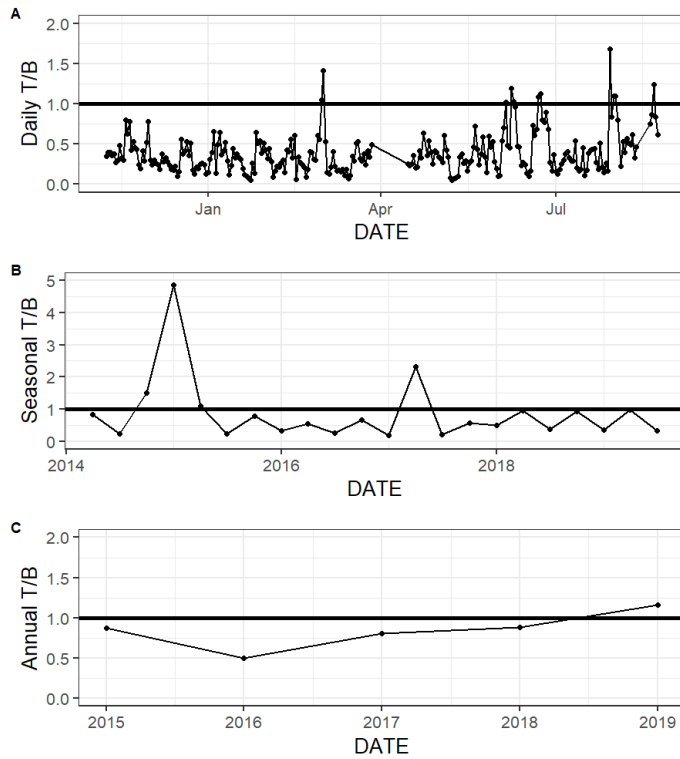
427 Results of the LDA for differentiation between daytime and nighttime conditions
 428 revealed that the most important factor was PAR, as would be expected (Table 1, Diel
 429 LD1). Temperature was the second most important factor to differentiate between day
 430 and night. The carbonate chemistry also played a critical role in day/night differentiation,

431 as $p\text{CO}_2$ was the third most important parameter, providing more evidence for
432 differentiation between day and night than other parameters that would be expected to
433 vary on a diel timescale (e.g., chlorophyll and DO) (Table 1).

434 *3.3 Controlling factors and correlates*

435 The relative influence of thermal and non-thermal factors (T/B) in controlling
436 $p\text{CO}_2$ varied over different time scales (Fig. 6, Table S4). Based on continuous data, non-
437 thermal processes generally exerted more control than thermal processes ($T/B < 1$) over
438 the entire 5+ years of discrete monitoring, within each season, and over most (167/178)
439 days (Fig. 6, Table S4). Annual T/B from discrete data ranged from 0.50 to 1.16, with
440 only one of the five sampled years having T/B greater than one (i.e., more thermal
441 influence; Table S4). While most individual seasons that were sampled experienced
442 stronger non-thermal control on $p\text{CO}_2$ ($T/B < 1$), the only season that never experienced
443 stronger thermal control was summer, with summer T/B values ranging from 0.21 – 0.35
444 for the 6 sampled years (Table S4).

445



446

447 **Figure 6.** Thermal versus non-thermal control on $p\text{CO}_2$ daily (A), seasonal (B), and
 448 annual (C) time scales using both continuous sensor data (daily, from Nov 8, 2016 to Aug
 449 3, 2017) and discrete sample data (seasonal and annual, from May 2, 2014- Feb. 25, 2020).
 450

451 Tidal fluctuations seemed to have a significant effect on carbonate system
 452 parameters (Table 2). Both temperature and salinity were higher at low tide during the
 453 winter and summer months and higher at high tide during the spring. $p\text{CO}_2$ was higher
 454 during low tide during all seasons. pH was higher during high tide during the winter and
 455 summer, but this reversed during the spring, when pH was higher at low tide. CO_2 flux
 456 also varied with tidal fluctuations. CO_2 flux was higher (more positive or less negative) in
 457 the low tide condition for all seasons (though the difference was not significant in

458 spring), i.e., the location was less of a CO₂ sink during low tide conditions in the winter
 459 and more of a CO₂ source during low tide conditions in the summer.

460

461 **Table 2.** Mean and standard deviation of temperature, salinity, pH, pCO₂, and calculated
 462 CO₂ flux (from continuous sensor measurements) during high and low tide conditions.

463

Parameter	Season	High Tide Mean	Low Tide Mean	Difference between tide levels, t-test p-value
Temperature (°C)	Winter	16.7 ± 1.7	17.6 ± 2.0	<0.0001
	Spring	24.4 ± 2.7	23.6 ± 2.7	<0.0001
	Summer	29.3 ± 0.5	30.1 ± 0.7	<0.0001
Salinity	Winter	30.2 ± 2.5	31.3 ± 2.9	<0.0001
	Spring	30.4 ± 1.9	30.0 ± 2.7	0.0071
	Summer	30.5 ± 2.4	34.5 ± 3.0	<0.0001
pH	Winter	8.20 ± 0.08	8.15 ± 0.06	<0.0001
	Spring	8.07 ± 0.09	8.10 ± 0.07	<0.0001
	Summer	8.08 ± 0.04	8.04 ± 0.06	<0.0001
pCO₂ (µatm)	Winter	331 ± 40	378 ± 42	<0.0001
	Spring	435 ± 33	443 ± 50	0.0154
	Summer	419 ± 30	482 ± 48	<0.0001
CO₂ Flux (mmol m⁻² d⁻¹)	Winter	-33.0 ± 38.1	-11.7 ± 21.8	<0.0001
	Spring	7.4 ± 14.0	8.7 ± 14.8	0.2248
	Summer	1.8 ± 6.3	16.0 ± 14.5	<0.0001

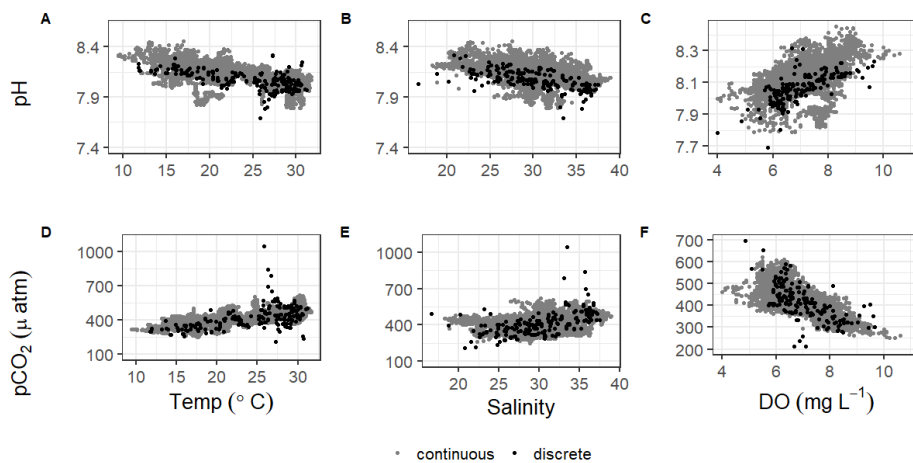
464

465 Mean water level varied between all seasons; mean spring (highest) water levels
 466 were on average 0.08 m higher than winter (lowest) water levels (ANOVA p<0.0001, fall
 467 was not considered because of a lack of water level data). The mean daily tidal range
 468 during our continuous monitoring period was 0.39 m ± 0.13 m, which did not
 469 significantly differ between seasons (ANOVA p=0.739). However, the day-night
 470 difference in tide level exhibited a strong seasonality, with spring and summer having
 471 higher tide level during the daytime and winter having higher tide level during the
 472 nighttime (Fig. 5).

473 There were significant correlations between carbonate system parameters (pH and
 474 pCO₂) and many of the other environmental parameters, including windspeed, DO,

475 turbidity, and fluorescent chlorophyll (Figure 7, Table S5). Both the continuous and
476 discrete sampling types indicate that pH has a significant negative relationship with both
477 temperature and salinity and $p\text{CO}_2$ has a significant positive relationship with both
478 temperature and salinity (Fig. 7). However, correlations with temperature were stronger
479 for continuous data and correlations with salinity were stronger for discrete data (Table
480 S5). The strongest correlations between continuous carbonate system data and all
481 investigated environmental parameters were with DO (positive correlation with pH and
482 negative correlation with $p\text{CO}_2$; Table S5). It is worth noting that there were no
483 observations of hypoxia at our study site during our monitoring, with minimum DO
484 levels of 3.9 mg L^{-1} and 4.0 mg L^{-1} for our continuous monitoring period and our discrete
485 sampling period, respectively.

486
487



488
489 **Figure 7.** Correlations of pH and $p\text{CO}_2$ with temperature, salinity, and DO from
490 continuous sensor data (gray) and all discrete data (black).
491

492 **Discussion**

493 *4.1 Comparing continuous monitoring and discrete sampling: Representative sampling in*
494 *a temporally variable environment*

495 Discrete water sample collection and analysis is the most common method that
496 has been employed to attempt to understand the carbonate system of estuaries. However,
497 it is difficult to know if these samples are representative of the spatial and temporal
498 variability in carbonate system parameters. While this time-series study cannot conclude
499 whether our broader sampling efforts in the MAE are representative of the spatial
500 variability in the estuary, it can investigate how representative our bimonthly to monthly
501 sampling is of the more high-frequency temporal variability that ASC experiences.

502 There were several instances where seasonal parameter means significantly
503 differed between the 10-month continuous monitoring period and the 5+ year discrete
504 sampling period (Table S2, $C \neq D$ or $D_c \neq D$) including temperature in the summer and
505 fall, salinity in the spring, pH in the summer and fall, and $p\text{CO}_2$ in winter, spring, and
506 summer. While clear seasonal variability was demonstrated for most parameters (using
507 both continuous and discrete data for the entire period), these differences between the 10-
508 month continuous monitoring period and our 5+ year monitoring period illustrate that
509 there is also interannual variability in the system. Therefore, short periods of monitoring
510 are unable to fully capture current baseline conditions.

511 During the continuous monitoring period (2016-2017), we found no significant
512 difference between sampling methods in the seasonal mean temperature, salinity, or
513 $p\text{CO}_2$. The two sampling methods also resulted in the same mean pH for all seasons
514 except for summer, when the sensor data recorded a higher mean pH than discrete

515 samples (Tables S1 and S2). During this case, we can conclude that discrete monitoring
516 did not accurately represent the system variability that was able to be captured by the
517 sensor monitoring. However, given that most seasons did not show differences in pH or
518 $p\text{CO}_2$ between sampling methods, the descriptive statistics associated with the discrete
519 monitoring did a fair job of representing system means. This is evidence that long-term
520 discrete monitoring efforts, which are much more widespread in estuarine systems than
521 sensor deployments, can be generally representative of the system despite known
522 temporal variability on shorter time scales. However, further study would be needed to
523 determine if this applies throughout the system, as the upper estuary generally
524 experiences greater variability.

525 Understanding the relationships of pH and $p\text{CO}_2$ with temperature and salinity is
526 important in a system (Fig. 7). Based on the results of an Analysis of Covariance
527 (ANCOVA), the relationship (slope) of pH with both temperature and salinity and of
528 $p\text{CO}_2$ with salinity were not significantly different between types of monitoring
529 (considering the sensor deployment period only), supporting the effectiveness of long-
530 term discrete monitoring programs when sensors are unable to be deployed. However,
531 ANCOVA did reveal the relationship of $p\text{CO}_2$ with temperature is significantly different
532 (method:temp $p=0.0062$) between monitoring methods.

533 The high temporal resolution of sensor data is presumably better for estimating
534 CO_2 flux at a given location than discrete sampling. Previous studies have pointed out
535 that discrete sampling methods, which generally involve only daytime sampling, do not
536 adequately capture the diel variability in the carbonate system and may therefore lead to
537 biased CO_2 fluxes (Crosswell et al., 2017; Liu et al., 2016). However, we found no

538 significant difference (within any season) between CO₂ flux values calculated with
539 hourly sensor data versus single, discrete samples collected monthly to twice monthly
540 (Table S2, Fig. 3). Calculated CO₂ fluxes also did not significantly differ between day
541 and night during any season, despite some differences in *p*CO₂ (Table S3), likely due to
542 the large error associated with the calculation of CO₂ flux (Table S1, Fig. 3) which will
543 be further discussed below. Therefore, the expected underestimation of CO₂ flux based
544 on diel variability of *p*CO₂ was not encountered at our study site, validating the use of
545 discrete samples for quantification of CO₂ fluxes (until methods with less associated error
546 are available). Even given less error in calculated flux, estimated fluxes would likely not
547 differ between methods on an annual scale (as *p*CO₂ did not), but CO₂ fluxes may differ
548 on a seasonal scale since the differences between daytime and nighttime *p*CO₂ were not
549 consistent across seasons (Table S3, Fig. 4).

550 There are many factors contributing to error associated with CO₂ flux. There is
551 still large error associated with estimates of estuarine CO₂ flux because turbulent mixing
552 is difficult to model and turbulence is the main control on CO₂ gas transfer velocity, *k*, in
553 shallow water environments. Thus, our wind speed parameterization of *k* is imperfect and
554 likely the greatest source of error (Borges and Abril, 2011; Van Dam et al., 2019). Other
555 notable sources of error include the data treatment. For example, we chose to seasonally
556 weight the individual calculated flux values in the calculation of annual flux to account
557 for differences in sampling frequency between seasons. From continuous data, the
558 weighted average flux was 0.2 mmol m⁻² d⁻¹, although choosing not to seasonally weight
559 and simply look at the arithmetic mean of fluxes calculated directly from sampling dates
560 would have resulted in an annual CO₂ flux of -0.7 mmol m⁻² d⁻¹ for the same period.

561 Similarly, the weighted average flux from all 5+ years of discrete data was -0.9 mmol m^{-2}
562 d^{-1} , but the arithmetic mean of fluxes would have resulted in an annual CO_2 flux of 0.2
563 $\text{mmol m}^{-2} \text{ d}^{-1}$ for the same period. Another source of error that could be associated with
564 the calculation of flux from the discrete data is the way in which wind speed data are
565 aggregated to be used in the windspeed parameterization. We decided to use daily
566 averages of the windspeed for calculations. Using the windspeed measured for the closest
567 time to our sampling time or the monthly averaged wind speed may have resulted in very
568 different flux values.

569

570 *4.2 Factors controlling temporal variability in carbonate system parameters*

571 Our study site had a relatively small range of pH and $p\text{CO}_2$ on both diel and
572 seasonal scales compared to other coastal regions (Challener et al., 2016; Yates et al.,
573 2007). This small variability is likely tied to a combination of the subtropical setting
574 (small temperature variability), the lower estuary position of our monitoring (further
575 removed from the already small freshwater influence), little ocean upwelling influence,
576 and the system's relatively high buffer capacity that results from the high alkalinity of the
577 freshwater endmembers (Yao et al., 2020). Just as the extent of hypoxia-induced
578 acidification was relatively low in Corpus Christi Bay because of the bay's high buffer
579 capacity (McCutcheon et al., 2019), the extent of pH fluctuation resulting from all
580 controlling factors at ASC would also be modulated by the region's high intrinsic buffer
581 capacity.

582 *4.2.1 Thermal and biological controls on carbonate chemistry*

583 We demonstrated that both temperature and non-thermal processes exert control
584 on $p\text{CO}_2$, but non-thermal control generally surpasses thermal control in ASC over

585 multiple time scales (Fig. 6, Table S4, $T/B < 1$). The magnitude of $p\text{CO}_2$ variation
586 attributed to non-thermal processes varied greatly (i.e., $\Delta p\text{CO}_{2,\text{nt}}$ had large standard
587 deviations, Table S4). For example, during the year of strongest non-thermal control
588 (2016), $\Delta p\text{CO}_{2,\text{nt}}$ was 534 μatm versus $\Delta p\text{CO}_{2,\text{nt}}$ of 209 μatm in the year of weakest
589 thermal control (2019). Conversely, the magnitude of $p\text{CO}_2$ variation attributed to
590 temperature was consistent across time scales. For example, during the year of strongest
591 thermal control (2015), $\Delta p\text{CO}_{2,\text{t}}$ was 276 μatm versus $\Delta p\text{CO}_{2,\text{t}}$ of 242 μatm in the year of
592 weakest thermal control (2017). Spring and fall seasons, which experienced the greatest
593 temperature swings (Table S1), had greater relative temperature control exerted on $p\text{CO}_2$
594 out of all seasons (Fig. 6, Table S4). The difference in T/B between sampling methods is
595 relatively small over the 10-month sensor deployment period, but it is worth noting that
596 T/B did not align over shorter seasonal time scales sampling methods (Fig. 6, Table S4).
597 Continuous monitoring demonstrated a greater magnitude of fluctuation resulting from
598 both temperature and non-thermal processes (i.e., greater $\Delta p\text{CO}_{2,\text{t}}$ and $\Delta p\text{CO}_{2,\text{nt}}$),
599 indicating that the extremes are generally not captured by the discrete, daytime sampling,
600 and sensor data would provide a better understanding of system controls.

601 The greater influence of non-thermal controls that we report conflicts with Yao
602 and Hu (2017), who found that ASC was primarily thermally controlled (T/B 1.53 – 1.79)
603 from May 2014 to April 2015. Yao and Hu (2017) also found that locations in the upper
604 estuary experienced lower T/B during flooding conditions than drought conditions.
605 Although the opposite was found at ASC, it is likely that the high T/B calculated at ASC
606 by Yao and Hu (2017) was still a result of the drought condition due to the long residence
607 time of the estuary. Since 2015, there has not been another significant drought in the

608 system, so it seems that non-thermal controls on $p\text{CO}_2$ are more important at this location
609 under normal freshwater inflow conditions.

610 Significantly warmer water temperatures were observed during the nighttime in
611 both summer and fall (Fig. 5), indicating that temperature could exert a slight control on
612 the carbonate system over a diel time scale. We note that significant differences in day
613 and night temperature within seasons do not indicate that diel differences were observed
614 on all days within the season, as large standard deviations in both daytime and nighttime
615 values result in considerable overlap. More substantial temperature swings between
616 seasons would result in more temperature control over a seasonal timescale. ASC seems
617 to have less thermal control of the carbonate system than offshore GOM waters, as
618 temperature had substantially higher explanatory value for pH and $p\text{CO}_2$ based on simple
619 linear regressions in offshore GOM waters ($R^2 = 0.81$ and 0.78 , respectively (Hu et al.,
620 2018)) than at ASC ($R^2 = 0.30$ and 0.52 , respectively, for sensor data and $R^2 = 0.38$ and
621 0.25 , respectively, for discrete data).

622 Though annual average $p\text{CO}_2$ (and CO_2 flux) are higher in the upper MAE and
623 lower offshore than at our study site, the same seasonal patterns that we observed (i.e.,
624 elevated $p\text{CO}_2$ and positive CO_2 flux in the summer and depressed $p\text{CO}_2$ and negative
625 CO_2 flux during the winter, Table S1, Fig. S1) has also been observed throughout the
626 entire MAE and the open Gulf of Mexico (Hu et al., 2018; Yao and Hu, 2017). These
627 seasonal patterns correspond with both the directional response of the system to
628 temperature and net community metabolism response to changing temperature, i.e.,
629 elevated respiration in summer months (Caffrey, 2004). Despite that there were no
630 observations of hypoxia, there was a strong relationship between the carbonate system

631 parameters and DO (Fig. 7, Table S5), suggesting that net ecosystem metabolism may
632 exert an important control on the carbonate system on seasonal time scales. The lack of
633 day-night difference in DO (Fig. 5F) despite the significant day-night difference in both
634 pH and $p\text{CO}_2$ suggests that net community metabolism is likely not a strong controlling
635 factor on diel time scales. Biological control likely becomes more important over
636 seasonal timescales.

637 *4.2.2 Tidal control on carbonate chemistry*

638 While the tidal range in the northwestern GOM is relatively small (1.30 m over
639 our 10-month continuous monitoring period), the tidal inlet location of our study site
640 results in proportionally more “coastal water” during high tide and proportionally more
641 “estuarine water” during low tide. The carbonate chemistry signal of these different water
642 masses was seen in the differences between high tide and low tide conditions at ASC
643 (i.e., high tide having lower $p\text{CO}_2$ because coastal waters are less heterotrophic than
644 estuarine waters, Table 2). Consequently, the relative importance of thermal versus non-
645 thermal controls may be modulated by tide level. We calculated the thermal and non-
646 thermal $p\text{CO}_2$ terms separately during high tide and low tide periods and found that non-
647 thermal control is more important during low tide conditions (within each season T/B is
648 0.10 ± 0.07 lower during the low tide than high tide). This is likely because low tide has
649 proportionally more “estuarine water” at the location and because there is less volume of
650 water for the end products of biological processes to accumulate. The difference in T/B
651 between high tide and low tide conditions was greatest in the spring, likely due to a
652 combination of elevated spring-time productivity and larger tidal ranges in the spring.

653 The GOM is one of the few places in the world that experiences diurnal tides
654 (Seim et al., 1987; Thurman, 1994), so theoretically, the fluctuations in $p\text{CO}_2$ associated
655 with tides may align to either amplify or reduce/reverse the fluctuations that would result
656 from diel variability in net community metabolism. Based on diel tidal fluctuations at this
657 site (i.e., higher tides during the day in the spring and summer and higher tides at night
658 during the winter, Fig. 5E) and the higher $p\text{CO}_2$ associated with low tide (Table 2), tidal
659 control should amplify the biological signal (nighttime $p\text{CO}_2 > \text{daytime } p\text{CO}_2$) during
660 spring and summer and reduce or reverse the biological signal during the winter. This
661 tidal control can explain the diel variability present in our $p\text{CO}_2$ data, which showed the
662 full reversal of the expected biological signal in the winter (Fig. 5C, Table S3, nighttime
663 $p\text{CO}_2 < \text{daytime } p\text{CO}_2$), i.e., the higher nighttime tides in winter brought in enough low
664 CO_2 water from offshore to fully offset any nighttime buildup of CO_2 from the lack of
665 photosynthesis. However, we note that the expected diel, biological control was likely
666 minimal since daytime DO was not consistently higher than nighttime DO (Fig. 5F). The
667 same seasonal pattern diel tide fluctuations were exhibited from Dec 20, 2016 (when the
668 tide data is first available) through the rest of our discrete monitoring period (Feb 25,
669 2020), indicating that tidal control on diel variability of carbonate system parameters was
670 likely consistent throughout this 3+ year period. The diel variability in pH did not mirror
671 $p\text{CO}_2$ as would be expected (Fig. 5). The relationship between pH and tide level more
672 closely mirrored the relationships of salinity and temperature with tide level (versus $p\text{CO}_2$
673 relationship with tide level; Table 2), indicating that controlling factors of the carbonate
674 system may not be exerted equally on both pH and $p\text{CO}_2$ over different time scales.

675 *4.2.3 Salinity and freshwater inflow controls on carbonate chemistry*

676 Previous studies have indicated that freshwater inflow may exert a primary
677 control on the carbonate system in the estuaries of the northwestern GOM (Hu et al.,
678 2015; Yao et al., 2020; Yao and Hu, 2017). Though the river water still has elevated
679 pCO₂ and depressed pH compared to the seawater endmember, the high riverine
680 alkalinity (often higher than the seawater endmember) in the region results in relatively
681 well-buffered estuarine conditions in MAE (Yao and Hu, 2017). Carbonate system
682 variability is much lower at ASC than it is in the more upper reaches of MAE, likely due
683 to the lesser influence of freshwater inflow and its associated changes in biological
684 activity at ASC (Yao and Hu, 2017). -Given the location of our sampling in the lower
685 portion of the estuary and the long residence time in the system, we did not directly
686 address river discharge as a controlling factor, but the influence of freshwater inflow may
687 be evident in the response of the system to changes in salinity. Fluctuating salinity at
688 ASC may also result from direct precipitation, stratification, and tidal fluctuations;
689 however, the low R² (0.02) associated with a simple linear regression between tide level
690 and salinity (p<0.0001) indicates that salinity fluctuations are more indicative of non-tidal
691 factors. Salinity data from both sensor and discrete monitoring were strongly correlated
692 with both pH and pCO₂, with correlation coefficients nearing (continuous) or surpassing
693 (discrete) that of the correlations with temperature (Fig. 7; Table S5). Periods of lower
694 salinity had higher pH and lower pCO₂, likely due to enhanced freshwater influence and
695 subsequent elevated primary productivity at the study site.

696 4.2.4 Windspeed and CO₂ inventory

697 We investigated wind speed as a possible control on the carbonate system to gain
698 insight into the effect of wind-driven CO₂ fluxes on the inventory of CO₂ in the water

699 column (and subsequent impacts to the entire carbonate system). The Texas coast has
700 relatively high wind speeds, with the mean wind speed observed during our continuous
701 monitoring period being 5.8 m s^{-1} . While this results in relatively high calculated CO_2
702 fluxes (Fig. 3), the seasonal relationship between $p\text{CO}_2$ and windspeed does not support a
703 change in inventory with higher winds. Since spring and summer both have a mean
704 estuarine $p\text{CO}_2$ greater than atmospheric level (and positive CO_2 flux, Table S1) a
705 negative relationship between windspeed and $p\text{CO}_2$ would be necessary to support this
706 hypothesis, but winter, spring, and fall all experience increases in $p\text{CO}_2$ with increasing
707 wind based on simple linear regression.

708 *4.3 Carbonate chemistry as a component of overall system variability*

709 Estuaries and coastal areas are dynamic systems with human influence, riverine
710 influence, and influence from an array of biogeochemical processes, resulting in highly
711 variable environmental conditions. Based on an LDA used to assess overall system
712 variability using a suite of environmental parameters compiled at a single location, we
713 can conclude that carbonate chemistry parameters are among the most important of
714 variants on both daily and seasonal time scales in this coastal setting. Of the two
715 carbonate system components that we incorporated (pH and $p\text{CO}_2$), $p\text{CO}_2$ was the most
716 critical in discriminating along diel or seasonal scales despite similar seasonal differences
717 that were identified by ANOVA (Table S2) and more seasons with significant diel
718 differences in pH (Table S3). pH seemed to be a larger component of overall system
719 variability on a seasonal time scale (compared to the very small contribution seen on a
720 diel scale, Table 1). Given that the seasonal and diel variability in carbonate chemistry at
721 this location is relatively small compared to other coastal areas that are in the literature,

722 the high contribution of carbonate chemistry to overall system variability that we detected
723 is likely to be present at other coastal locations around the world.

724 **5. Conclusions**

725 We monitored carbonate chemistry parameters (pH and $p\text{CO}_2$) using both sensor
726 deployments (10 months) and discrete sample collection (5+ years) at the Aransas Ship
727 Channel, TX, to characterize temporal variability. Significant seasonal variability and
728 diel variability in carbonate system parameters were both present at the location. Diel
729 fluctuations were smaller than many other areas previously studied. The difference
730 between daytime and nighttime values of carbonate system parameters varied between
731 seasons, occasionally reversing the expected diel variability due to biological processes.
732 Tide level (despite the small tidal range), temperature, freshwater influence, and
733 biological activity all seem to exert important controls on the carbonate system at the
734 location. The relative importance of the different controls varied with timescale, and
735 controls were not always exerted equally on both pH and $p\text{CO}_2$. Carbonate chemistry
736 (particularly $p\text{CO}_2$) was among the most important environmental parameters to in
737 overall system variability to distinguish between both diel and seasonal environmental
738 conditions.

739 Despite known temporal variability on shorter timescales, discrete sampling was
740 generally representative of the average carbonate system on a seasonal and annual basis
741 based on comparison with our sensor data. Discrete data captured interannual variability,
742 which could not be captured by the shorter-term continuous sensor data. Additionally,
743 there was no difference in CO_2 flux between sampling types. All of these findings

744 support the validity of discrete sample collection for carbonate system characterization at
745 this location.

746 This is one of the first studies that investigates high-temporal frequency data from
747 deployed sensors that measure carbonate system parameters in an estuary-influenced
748 environment. Long-term, effective deployments of these monitoring tools could greatly
749 improve our understanding of estuarine systems. This study's detailed investigation of
750 data from multiple, co-located environmental sensors was able to provide insight into
751 potential driving forces of carbonate chemistry on diel and seasonal time scales; this
752 provides strong support for the implementation of carbonate chemistry monitoring in
753 conjunction with preexisting coastal environmental monitoring infrastructure.

754 Strategically locating such sensors in areas that are subject to local acidification drivers
755 or support large biodiversity or commercially important species may be the most crucial
756 in guiding future mitigation and adaptation strategies for natural systems and aquaculture
757 facilities.

758

759 **Data availability**

760 Continuous sensor data are archived with the National Oceanic and Atmospheric
761 Administration's (NOAA's) National Centers for Environmental Information (NCEI)
762 (<https://doi.org/10.25921/dkg3-1989>). Discrete sample data are available in two separate
763 datasets archived with National Science Foundation's Biological & Chemical
764 Oceanography Data Management Office (BCO-DMO) (doi:10.1575/1912/bco-
765 dmo.784673.1 and doi: 10.26008/1912/bco-dmo.835227.1).

766 **Author Contribution**

767 MM and XH defined the scope of this work. XH received funding for all components of
768 the work. MM, HY, and CJS performed field sampling and laboratory analysis of
769 samples. MM prepared the initial manuscript and all co-authors contributed to revisions.

770 **Competing interests**

771 The authors declare that they have no conflict of interest.

772

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