



1 **Temporal variability and driving factors of the carbonate system in the Aransas**

2 **Ship Channel, TX: A time-series study**

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

5 ¹ Harte 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

10

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

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15 **Keywords:** *p*CO₂, acidification, diel variability, seasonal variability, autonomous sensors



16 **Abstract**

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



39 on shorter timescales, discrete sampling was generally representative of the average
40 carbonate system and average air-sea CO₂ flux on a seasonal and annual basis based on
41 comparison with sensor data.

42 **1. Introduction**

43 Estuaries, the dynamic environments where the coast and freshwater inflows meet
44 the ocean, are economically and ecologically important because they are biological
45 hotspots, but they are also heavily influenced by anthropogenic activity. Because of the
46 diversity of co-occurring biogeochemical processes, estuaries experience substantial
47 spatial and temporal heterogeneity in water quality and chemistry, including carbonate
48 chemistry parameters such as pH and partial pressure of CO₂ ($p\text{CO}_2$) (Hofmann et al.,
49 2011; Waldbusser and Salisbury, 2014). Carbonate chemistry, or the speciation of
50 inorganic carbon in seawater, is important for two main reasons. First, CO₂ acidifies
51 seawater, whether it is a result of uptake from the atmosphere (generally acknowledged
52 as ocean acidification, or OA) or it is produced by biogeochemical processes in the water
53 (that may intensify or alleviate the effects of OA). This is problematic because
54 acidification can negatively affect marine organisms, especially those that construct
55 calcium carbonate shells and skeletons (Barton et al., 2015; Bednaršek et al., 2012;
56 Ekstrom et al., 2015; Gazeau et al., 2007; Gobler and Talmage, 2014). Second, the ocean
57 contributes substantially to the global carbon budget, which is important to understand
58 because of climate change implications. Despite the small surface area of coastal waters
59 relative to the global ocean, coastal waters are recognized as important contributors in
60 global carbon cycling (Borges, 2005; Cai, 2011; Laruelle et al., 2018).



61 While open ocean environments are relatively well studied and understood
62 regarding carbonate chemistry, acidification, and air-sea CO₂ fluxes, large uncertainties
63 remain in estuarine environments. Estuaries are challenging to fully understand because
64 of the heterogeneity between and within estuaries that is driven by diverse processes
65 operating on different time scales such as river discharge, nutrient and organic matter
66 loading, stratification, and coastal upwelling (Jiang et al., 2013; Mathis et al., 2012). The
67 traditional sampling method for carbonate system characterization involving discrete
68 water sample collection and laboratory analysis is known to lead to biases in average
69 *p*CO₂ and CO₂ flux calculations due to daytime sampling that neglects to capture diel
70 variability (Li et al., 2018). Mean diel ranges in pH can exceed 0.1 unit and single day
71 ranges can exceed 1 pH unit, with especially high diel variability in biologically
72 productive areas or areas with higher mean *p*CO₂ (Challener et al., 2015; Cyronak et al.,
73 2018; Schulz and Riebesell, 2013; Semesi et al., 2009; Yates et al., 2007). These diel
74 ranges can far surpass the magnitude of the changes in open ocean surface waters that
75 have occurred since the start of the industrial revolution and rival spatial variability in
76 productive systems, indicating their importance for a full understanding of the carbonate
77 system.

78 Despite the need for high-frequency measurements, sensor deployments have
79 been limited in estuarine environments (especially compared to their extensive use in the
80 open ocean) because of the challenges associated with varying conditions, biofouling,
81 and sensor drift (Sastri et al., 2019). Carbonate chemistry monitoring in the Gulf of
82 Mexico (GOM), and especially its estuaries, has been relatively minimal compared to the
83 United States east and west coasts. The GOM estuaries, where this study takes place,



84 currently have less exposure to concerning levels of acidification than other estuaries
85 because of their high temperatures (causing water to hold less CO₂ and support high
86 productivity year-round) and often suitable river chemistries (i.e. relatively high buffer
87 capacity) (McCutcheon et al., 2019; Yao et al., 2020). However, respiration-induced
88 acidification is present in both the open GOM (e. g., subsurface water influenced by the
89 Mississippi River Plume and outer shelf region near the Flower Garden Banks National
90 Marine Sanctuary) and GOM estuaries, and most estuaries in the northwestern GOM
91 have also experienced long-term acidification (Cai et al., 2011; Hu et al., 2018, 2015;
92 Kealoha et al., 2020; McCutcheon et al., 2019; Robbins and Lisle, 2018). This known
93 acidification as well as the relatively high CO₂ fluxes from the estuaries of the northwest
94 GOM (which may change our understanding of global estuarine contribution to the
95 carbon budget) illustrates the necessity to study the baseline variability and driving
96 factors of carbonate chemistry in the region. In this study, we explored temporal
97 variability in the carbonate system in Aransas Ship Channel—a tidal inlet in a semi-arid
98 region of the northwestern GOM—using both discrete bottle sample collection and
99 hourly sensor measurements, and we explored the driving factors of that variability using
100 data from other co-located environmental sensors.

101 **2. Materials and Methods**

102 *2.1 Location*

103 Autonomous sensor monitoring and discrete water sample collections for
104 laboratory analysis of carbonate system parameters were performed in the Aransas Ship
105 Channel (27°50'17"N, 97°3'1"W). The Aransas Ship Channel is one of the few permanent
106 tidal inlets that intersect a string of barrier islands and connect the GOM coastal waters



107 with the lagoonal estuaries in the northwest GOM (Fig. 1). The Aransas Ship Channel
108 provides the direct connection between the nwGOM and the Mission-Aransas Estuary
109 (Copano and Aransas Bays) to the north and Nueces Estuary (Nueces and Corpus Christi
110 Bays) to the south (Fig. 1). The tidal range in the region is small, with around 0.6 m tides
111 on the open coast and less than 0.3 m in the estuaries (Montagna et al., 2011). Mission-
112 Aransas Estuary (MAE) is fed by two small rivers, the Mission (1787 km² drainage
113 basin) and Aransas (640 km² drainage basin) Rivers (<http://waterdata.usgs.gov/>), which
114 both experience low base flows punctuated by periodic high flows during storm events.
115 MAE has an average residence time of one year (Solis and Powell, 1999), so there is a
116 substantial lag between time of rainfall and riverine delivery to the Aransas Ship Channel
117 in the lower estuary. A significant portion of riverine water flowing into Aransas Bay
118 originates from the larger rivers further northeast on the Texas coast via the Intracoastal
119 Waterway (i.e. Guadalupe River (26,625 km² drainage basin) feeds San Antonio Bay and

120 has a much shorter residence time of nearly 50 days) (Solis and Powell, 1999; USGS, 2001).

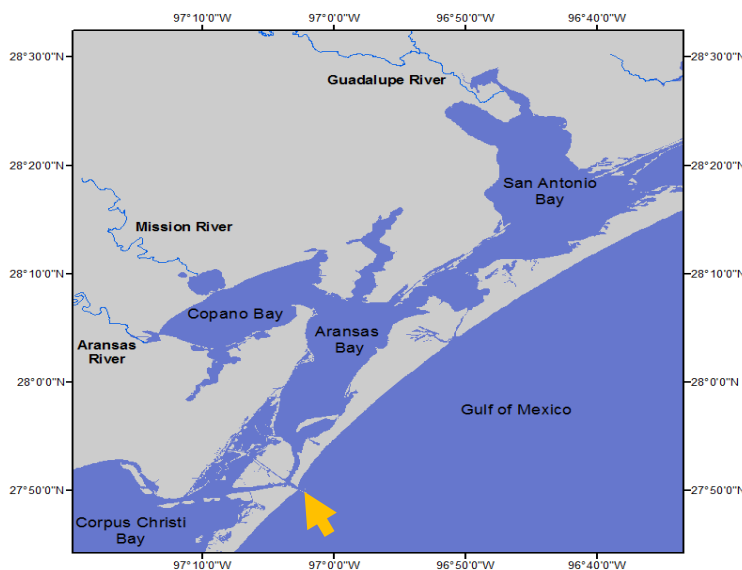


Fig 1. Location of Aransas Ship Channel where this study took place (arrow) and surrounding bay systems.



133 2.2 *Continuous Monitoring*

134 Autonomous sensor monitoring (referred to throughout as continuous monitoring)
135 of pH and $p\text{CO}_2$ was conducted from Nov. 8, 2016 to Aug. 23, 2017 at the University of
136 Texas Marine Science Institute's research pier in the Aransas Ship Channel. The sensor
137 deployment was shorter than intended because the pier where the sensors were deployed
138 was destroyed in the aftermath of Hurricane Harvey in 2017. The pH data were collected
139 using an SATlantic[®] SeaFET pH sensor (on total pH scale) and $p\text{CO}_2$ data were collected
140 using a Sunburst[®] SAMI-CO₂. Temperature and salinity data were measured by a YSI[®]
141 600OMS V2 sonde. Hourly data collected by all sensors (pH, $p\text{CO}_2$, salinity, and
142 temperature) were saved in the onboard data loggers and downloaded during service trips
143 to the field site. Sensor failures or pump failures occurred for short periods of time
144 throughout the deployment. Measurements were recorded on 262 individual days, with
145 176 of those days having the full set of 24 (hourly) measurements.

146 Ideally, *in-situ* sensors should be deployed under the sea surface. However, to
147 reduce the maintenance cost and effort for sensors deployed in warm water that
148 experiences intense biofouling, the sensors were set up to measure pH and $p\text{CO}_2$ from an
149 *ex situ* position using *in situ* seawater. Water was pumped from ~1 m below the sea
150 surface into the bottom spigot of a 100-Qt cooler that housed the SAMI-CO₂ and SeaFET
151 sensors. To allow water outflow, a 1" hole was drilled at the opposite side of the spigot
152 near the top of the cooler rim, allowing water to flow back to the sea surface. The pump
153 was programmed to turn on 20 minutes before each hour, pumping more than enough
154 water to fully flush the cooler, and sensors recorded measurements on the hour. The YSI



155 sonde was deployed directly into the Aransas Ship Channel inside a 2” PVC pipe at ~1 m
156 below the sea surface.

157 Visits to the field site were conducted every two weeks to service all sensors and
158 clean the cooler. Additionally, duplicate water samples were collected on the hour during
159 service trips for quality assurance of sensor data and to check that surface water and
160 cooler chemistries aligned. Samples were collected from the channel near the pump inlet
161 and from the cooler that housed the sensors, and water temperature and salinity were
162 measured using a handheld YSI data sonde. The average difference between sensor pH
163 and laboratory pH (from the cooler) was used to establish a correction of -0.05 to the
164 final *in-situ* pH data since the SeaFET may experience drift. The difference between the
165 sensor $p\text{CO}_2$ and calculated $p\text{CO}_2$ is reported, but it is not used for a correction since the
166 spectrophotometric measurements of the SAMI-CO₂ should not experience drift. Sensor
167 data were discarded from analysis during known periods of pump failure when cooler
168 chemistry separated from that of the Aransas Ship Channel.

169 *2.3 Discrete Sample Collection and Analysis*

170 In addition to the discrete sample collections that occurred for quality assurance
171 during sensor servicing visits, long-term monitoring via discrete water sample collection
172 was conducted at the Aransas Ship Channel from May 2, 2014 to February 25, 2020.
173 Sampling was conducted from a small vessel at a station very near to the sensor
174 deployment every two weeks during the summer months and monthly during the winter
175 months. Water sample collection followed standard protocol for ocean carbonate
176 chemistry studies (Dickson et al., 2007). Ground glass borosilicate bottles (250 mL) were
177 filled with surface water and preserved with 100 μL saturated mercury chloride (HgCl_2).



178 Apiezon[®] grease was applied to the bottle stopper, which was then secured to the bottle
179 using a rubber band and a nylon hose clamp.

180 These samples were used for laboratory dissolved inorganic carbon (DIC) and pH
181 measurements. DIC was measured by injecting 0.5 mL of sample into 1 ml 10% H₃PO₄
182 (balanced by 0.5 M NaCl) with a high-precision Klotz syringe pump. The CO₂ gas
183 produced through sample acidification was then stripped using high-purity nitrogen gas
184 and carried into a Li-Cor infrared gas detector. DIC analyses had a precision of 0.1%.
185 Certified Reference Material (CRM) was used to ensure the accuracy of the analysis
186 (Dickson et al. 2003). For samples with salinity >20, pH was measured using a
187 spectrophotometric method at 25 ± 0.1°C (Carter et al. 2003) and the Douglas and Byrne
188 (2017) equation. Analytical precision of the spectrophotometric method for pH
189 measurement was ±0.0004 pH units. A calibrated Orion Ross glass pH electrode was
190 used to measure pH at 25 ± 0.1°C for samples with salinity <20, and analytical precision
191 was ±0.01 pH units. All pH values obtained using the potentiometric method were
192 converted to total scale at *in situ* temperature (Millero 2001). Salinity of the discrete
193 samples was measured using a benchtop salinometer calibrated by MilliQ water and a
194 known salinity CRM.

195 2.4 Data Processing and Statistical Analyses

196 For the discrete samples, *p*CO₂ was calculated using CO2Sys for Excel.
197 Carbonate speciation calculations were done using Millero (2010) carbonic acid
198 dissociation constants (K₁ and K₂), Dickson (1990) bisulfate dissociation constant, and
199 Uppström (1974) borate concentration. Temporal variability was investigated in the form
200 of seasonal and diel variability (Tables 1-2). For seasonal analysis, December to February



201 was considered winter, March to May was considered spring, June to August was
202 considered summer, and September to November was considered fall. Two-way
203 ANOVAs were used to examine differences in parameter means between seasons, using
204 differences between monitoring methods as the second factor (as differences between
205 seasons may not be the same between monitoring methods, Table 3). Since there was a
206 significant interaction in the two-way ANOVA, the differences between seasons were
207 investigated within each monitoring method. Post-hoc multiple comparisons (between
208 seasons within sampling types) were conducted using the Westfall adjustment (Westfall,
209 1997). For diel comparisons, daytime and nighttime variables were defined as 09:00-
210 15:00 local standard time and 21:00-03:00 local standard time, respectively, based on the
211 6-hour periods with highest and lowest photosynthetically active radiation (PAR; data
212 obtained from the Mission-Aransas National Estuarine Research Reserve (MANERR) at
213 <https://missionaransas.org/science/download-data>. Paired *t*-tests, comparing the daytime
214 mean with the nighttime mean on respective days, were used to look for significant
215 differences between daytime and nighttime parameter values across the full sampling
216 period and within each season (Table 2).

217 Equation 1 was used for air-water CO₂ flux calculations (Wanninkhof, 1992;
218 Wanninkhof et al., 2009). Positive flux values indicate CO₂ emission from the water into
219 the atmosphere (the estuary acting as a source of CO₂), and negative flux values indicate
220 CO₂ uptake by the water (the estuary acting as a sink for CO₂).

$$221 \quad F = k K_0 (p\text{CO}_{2,w} - p\text{CO}_{2,a}) \quad (1)$$



222 where k is the gas transfer velocity (in m d^{-1}), K_0 (in $\text{mol l}^{-1} \text{atm}^{-1}$) is the solubility
223 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
224 μatm) in the water and air, respectively.

225 We used the wind speed parameterization for gas transfer velocity (k) from Jiang
226 et al. (2008) converted from cm h^{-1} to m d^{-1} , which is thought to be the best estuarine
227 parameterization at this time (Crosswell et al., 2017) as it is a composite of k over several
228 estuaries. The calculation of k requires a windspeed at 10 m above the surface, so
229 windspeeds measured at 3 m above the surface were converted using the power law wind
230 profile (Hsu, 1994; Yao and Hu, 2017). To assess uncertainty, other parameterizations
231 with direct applications to estuaries in the literature were also used to calculate CO_2 flux
232 (Raymond and Cole 2001; Ho et al. 2006). We note that parameterization of k based on
233 solely windspeed is flawed because several additional parameters can contribute to
234 turbulence including turbidity, bottom-driven turbulence, water-side thermal convection,
235 tidal currents, and fetch (Wanninkhof 1992, Abril et al., 2009, Ho et al., 2104, Andersson
236 et al., 2017), however it is currently the best option for this system given the limited
237 investigations of CO_2 flux and contributing factors in estuaries.

238 Hourly windspeed data used in calculations were retrieved from the NOAA-
239 controlled Texas Coastal Ocean Observation Network (TCOON;
240 <https://tidesandcurrents.noaa.gov/tcoon.html>). The closest station with windspeed data,
241 Port Aransas Station, was located directly in the Aransas Ship Channel (further inshore
242 than our monitoring location), however there were several long periods of missing data.
243 To fill in the data gaps, wind speed data from nearby Aransas Pass Station were used, and
244 for all subsequent gaps, data from nearby Nueces Bay Station were used. For continuous



245 monitoring data, TCOON's measured hourly windspeed at each time point was used in
246 flux calculations. For biweekly discrete samples, averaged daily windspeeds were
247 calculated from TCOON's measured hourly windspeeds and used in flux calculations for
248 the respective day. Monthly mean atmospheric $x\text{CO}_2$ data (later converted to $p\text{CO}_2$) were
249 obtained from NOAA's flask sampling network of the Global Monitoring Division of the
250 Earth System Research Laboratory at the Key Biscayne (FL, USA) station, when
251 available

252 ([https://www.esrl.noaa.gov/gmd/dv/data/index.php?site=KEY¶meter_name=Carbon](https://www.esrl.noaa.gov/gmd/dv/data/index.php?site=KEY¶meter_name=Carbon%2BDioxide)
253 [%2BDioxide](https://www.esrl.noaa.gov/gmd/dv/data/index.php?site=KEY¶meter_name=Carbon%2BDioxide)). For 2019 and 2020, when $x\text{CO}_2$ data from Key Biscayne were
254 unavailable, monthly global average values were used
255 (ftp://aftp.cmdl.noaa.gov/products/trends/co2/co2_mm_mlo.txt).

256 *Factors controlling the carbonate system parameters*

257 Thermal versus non-thermal controls on $p\text{CO}_2$ were investigated following
258 Takahashi et al. (2002) over annual, seasonal, and daily time scales (Table 4). When
259 calculating annual T/B values with discrete data, only complete years (sampling from
260 January to December) were included. When calculating daily T/B values with continuous
261 data, only complete days (24 hourly measurements) were included.

$$262 \quad p\text{CO}_{2,\text{thermal}} = p\text{CO}_{2,\text{mean}} \times \exp[\delta \times (T_{\text{obs}} - T_{\text{mean}})] \quad (2)$$

$$263 \quad p\text{CO}_{2,\text{nonthermal}} = p\text{CO}_{2,\text{obs}} \times \exp[\delta \times (T_{\text{mean}} - T_{\text{obs}})] \quad (3)$$

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



$$268 \quad T/B = \frac{\max(pCO_{2,thermal}) - \min(pCO_{2,thermal})}{\max(pCO_{2,non-thermal}) - \min(pCO_{2,non-thermal})} \quad (4)$$

269 Where a T/B greater than one indicates that temperature's control on pCO_2 is greater than
270 the control from non-thermal factors (i.e. physical and biological processes) and a T/B
271 less than one indicates that non-thermal factors' control on pCO_2 is greater than the
272 control from temperature.

273 Tidal control on parameters was investigated using only the continuous
274 monitoring data (Table 5). Hourly measurements of water level immediately offshore
275 from the Aransas Ship Channel were obtained from NOAA's Tides and Currents Aransas
276 Pass Station
277 [https://tidesandcurrents.noaa.gov/waterlevels.html?id=8775241&name=Aransas,%20Ara](https://tidesandcurrents.noaa.gov/waterlevels.html?id=8775241&name=Aransas,%20Aransas%20Pass&state=TX)
278 [nsas%20Pass&state=TX](https://tidesandcurrents.noaa.gov/waterlevels.html?id=8775241&name=Aransas,%20Aransas%20Pass&state=TX). Tide data were merged with our sensor data by date and hour;
279 given that there were gaps in available water level measurements (and no measurements
280 prior to December 20, 2016), the usable dataset was reduced from 6088 observations to
281 5121 observations and fall was omitted from analyses. To examine differences between
282 parameters during high tide and low tide, we defined high tide as tide level greater than
283 the third quartile tide level value and low tide as a tide level less than the first quartile
284 tide level value. The difference between high and low tide for each parameter was
285 examined within each season (using t-tests) because of a significant interaction (based on
286 $\alpha=0.05$) between the season and high/low tide factors in a two-way ANOVA.

287 **3. Results**

288 *3.1 Continuous monitoring results*

289 Over the 10-month continuous monitoring period, all sensor-measured parameters
290 showed substantial temporal variability on seasonal and diel time scales (Fig. 2, Tables 1-



291 3). Mean values of sensor-measured parameters over the entire monitoring period were:
 292 temperature - $23.1^{\circ}\text{C} \pm 5.3^{\circ}\text{C}$, ranging from 9.4°C to 31.7°C ; salinity - 30.8 ± 3.7 ,
 293 ranging from 18.3 to 38.9; pH - 8.12 ± 0.10 , ranging from 7.79 to 8.45; and $p\text{CO}_2$ - $416 \pm$
 294 $60 \mu\text{atm}$, ranging from $251 \mu\text{atm}$ to $620 \mu\text{atm}$ (Table 1). Temperature was significantly
 295 different between each season (Table 3), with the highest being summer and the lowest
 296 being winter (Table 1). Salinity was highest in the summer and lowest in the fall, and
 297 salinity differed between all seasons except from spring and winter (Tables 1 and 3). pH
 298 and $p\text{CO}_2$ were both significantly different between all seasons (Table 3). Winter had
 299 both the highest seasonal pH (8.19 ± 0.08) and lowest seasonal $p\text{CO}_2$ ($365 \pm 44 \mu\text{atm}$)
 300 and summer had both the lowest seasonal pH (8.05 ± 0.06) and highest seasonal $p\text{CO}_2$
 301 ($463 \pm 48 \mu\text{atm}$) (Table 1, Fig. 2- 3).

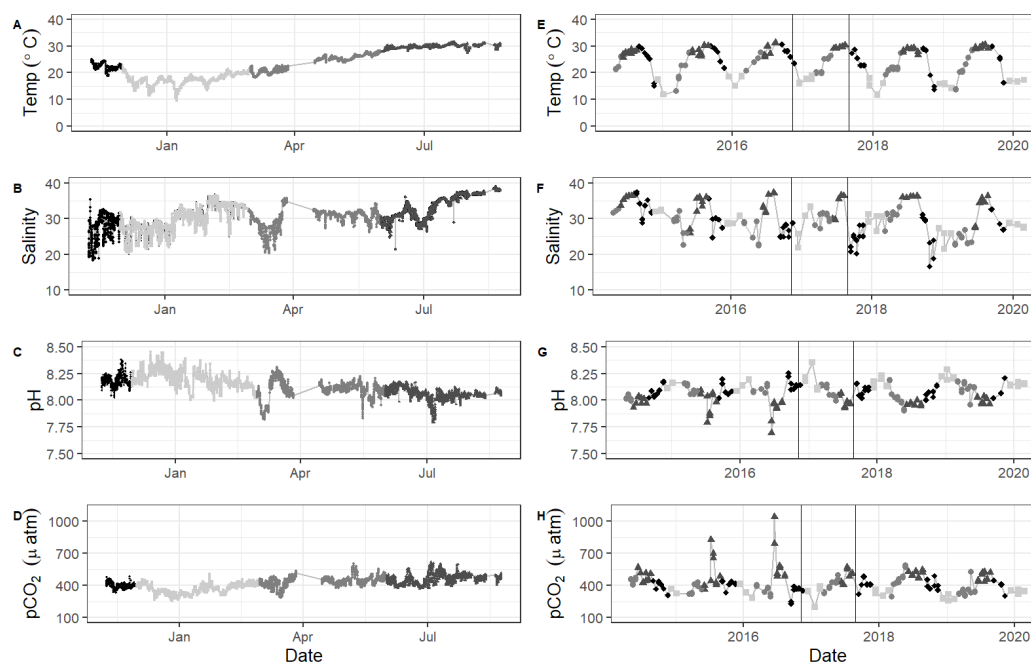
302 Table 1. Mean and standard deviation of annual and seasonal temperature, salinity, pH,
 303 $p\text{CO}_2$, and CO_2 flux from continuous monitoring, discrete sampling over the continuous
 304 monitoring period, and discrete sampling over the entire sampled period. Reported annual
 305 means are seasonally weighted to account for disproportional sampling between seasons
 306 (however, reported annual standard deviation is associated with the un-weighted,
 307 arithmetic mean). CO_2 fluxes were calculated using the Jiang et al. (2008) wind speed
 308 parameterization for gas transfer velocity, and ranges of CO_2 flux that are given in
 309 brackets represent means calculated using parameterizations from Ho et al. (2006) and
 310 Raymond and Cole (2001), respectively.

Parameter		Continuous Monitoring		Discrete Sampling	
		Time Period	Nov. 8 2016 – Aug 23, 2017	Nov. 8 2016 – Aug 23, 2017	May 2, 2014- Feb. 25, 2020
Temperature (°C)	Annual		23.1 ± 5.3	23.5 ± 5.0	24.1 ± 5.3
	Winter		17.3 ± 2.1	17.3 ± 1.1	16.2 ± 2.0
	Spring		23.8 ± 2.8	23.4 ± 2.9	22.6 ± 3.7
	Summer		29.7 ± 0.8	29.6 ± 0.5	28.7 ± 1.4
	Fall		22.5 ± 2.1	23.6 ± 0.1	25.5 ± 4.5
Salinity	Annual		30.8 ± 3.7	30.4 ± 3.5	30.1 ± 4.4
	Winter		30.0 ± 3.7	29.3 ± 4.6	28.9 ± 2.9
	Spring		30.2 ± 2.6	30.0 ± 1.7	28.7 ± 3.4
	Summer		33.3 ± 3.2	33.6 ± 3.2	34.6 ± 2.8
	Fall		27.6 ± 3.7	28.8 ± 0.1	28.4 ± 4.5
pH	Annual		8.12 ± 0.10	8.092 ± 0.078	8.079 ± 0.092
	Winter		8.19 ± 0.08	8.157 ± 0.041	8.162 ± 0.065
	Spring		8.09 ± 0.09	8.078 ± 0.056	8.077 ± 0.066



	Summer	8.05 ± 0.06	7.999 ± 0.051	7.975 ± 0.046
	Fall	8.18 ± 0.05	8.136 ± 0.001	8.100 ± 0.071
$p\text{CO}_2$ (μatm)	Annual	416 ± 60	400 ± 71	406 ± 100
	Winter	365 ± 44	349 ± 31	331 ± 39
	Spring	436 ± 45	413 ± 54	396 ± 67
	Summer	463 ± 48	480 ± 59	511 ± 108
	Fall	400 ± 25	357 ± 2	386 ± 62
CO_2 Flux ($\text{mmol m}^{-2} \text{d}^{-1}$)	Annual	0.2 ± 23.7 [0.1 – (-87.6)]	-1.5 ± 9.2 [(-2.6) – (-4.5)]	$(-0.8) \pm 18.7$ [(-0.7) – (-5.3)]
	Winter	$(-16.9) \pm 29.2$ [(-14.6) – (-444.0)]	$(-9.9) \pm 5.2$ [(-8.3) – (-16.2)]	$(-13.0) \pm 13.5$ [(-10.6) – (-25.6)]
	Spring	7.6 ± 15.0 [6.5 – 109.0]	1.0 ± 7.1 [1.0 – 3.3]	$(-6.5) \pm 12.2$ [(-5.5) – (-18.0)]
	Summer	10.8 ± 13.3 [9.1 – 28.9]	10.5 ± 7.8 [8.6 – 16.3]	18.3 ± 19.6 [15.3 – 65.5]
	Fall	$(-0.9) \pm 7.7$ [(-0.7) – (-44.0)]	(-7.5) [(-6.2) – (-11.4)]	$(-2.3) \pm 13.7$ [(-1.9) – (-0.9)]

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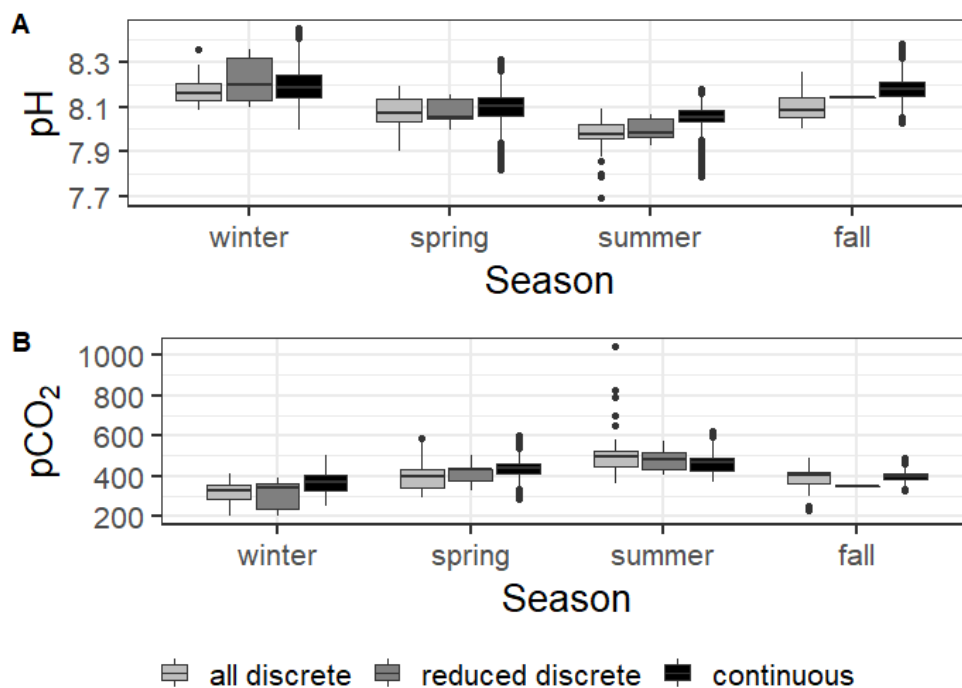
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Fig 2. Time series data from continuous monitoring (A-D, Nov 8, 2016 to Aug 3, 2017) and discrete sample analysis (E-H, Nov 8, 2016 to Aug 3, 2017) at the Aransas Ship Channel. Gray scale (and shape) in the datapoints represents divisions between the four seasons. Vertical lines in (E-H) denote the time period of continuous monitoring.



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Figure 3. Boxplots of seasonal variability in pH and $p\text{CO}_2$ using all discrete data (May 2, 2014- Feb. 25, 2020), reduced discrete data (Nov. 8 2016 – Aug 23, 2017, to overlap with continuous monitoring), and continuous sensor data (Nov. 8 2016 – Aug 23, 2017)

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Table 2. Diel variability in system parameters from continuous sensor data (Nov 8, 2016 – Aug 23, 2017). The p-values reported are from a paired- t test comparing the means of each day (9am-3pm LST) with the mean of the same night (9pm – 3am LST); all significant results based on $\alpha=0.05$ are bolded. Diel range calculations were done using only days with the full 24 hours of hourly measurements (176 out of 262 measured) to ensure that data gaps did not influence the calculations. Reported fluxes use the Jiang et al. (2008) gas transfer velocity parameterization. Note that the Fall season had much fewer observations than other seasons because of the timing of sensor deployment.

Parameter	Time Period	Daytime Mean	Nighttime Mean	Day versus Night p-value	Mean Diel Range	Minimum Diel Range	Maximum Diel Range
Temperature (°C)	Full Sampling Period	23.0 ± 5.3	23.2 ± 5.4	<0.0001	1.3 ± 0.8	0.30	3.93
	Winter	17.2 ± 2.1	17.4 ± 2.1	0.2055	1.5 ± 0.8	0.3	3.8
	Spring	23.7 ± 2.7	23.8 ± 2.9	0.5579	1.2 ± 0.6	0.3	3.0
	Summer	29.6 ± 0.7	29.9 ± 0.8	<0.0001	1.0 ± 0.6	0.3	3.8
	Fall	22.0 ± 1.19	23.0 ± 1.0	<0.0001	1.8 ± 0.9	0.8	3.9



Salinity	Full	30.5 ± 4.1	31.0 ± 3.3	0.0004	3.4 ± 2.7	0.250	15.870
	Sampling Period						
	Winter	29.6 ± 4.2	30.4 ± 3.1	0.0051	3.8 ± 2.2	0.25	9.48
	Spring	30.1 ± 2.6	30.2 ± 2.6	0.5604	2.5 ± 2	0.4	8.17
	Summer	33.4 ± 3.2	33.1 ± 3.3	0.0550	2.0 ± 1.7	0.3	9.73
	Fall	25.9 ± 3.9	29.0 ± 3.2	<0.0001	7.7 ± 3.6	1.2	15.87
pH	Full	8.12 ± 0.10	8.13 ± 0.09	<0.0001	0.09 ± 0.05	0.02	0.28
	Sampling Period						
	Winter	8.18 ± 0.08	8.20 ± 0.07	0.0108	0.10 ± 0.05	0.02	0.28
	Spring	8.09 ± 0.09	8.10 ± 0.08	0.3286	0.08 ± 0.03	0.03	0.18
	Summer	8.04 ± 0.06	8.07 ± 0.05	<0.0001	0.08 ± 0.04	0.03	0.19
	Fall	8.20 ± 0.05	8.17 ± 0.05	0.0038	0.12 ± 0.04	0.03	0.20
pCO₂ (µatm)	Full	417 ± 54	416 ± 65	0.7065	58 ± 33	12.6	211.3
	Sampling Period						
	Winter	374 ± 44	358 ± 43	<0.0001	43 ± 21	12.6	121.1
	Spring	438 ± 42	437 ± 48	0.7237	61 ± 31	20.5	152.8
	Summer	452 ± 44	471 ± 51	0.0003	74 ± 42	23.6	211.3
	Fall	406 ± 24	399 ± 27	0.0545	56 ± 18	22	92.2
CO₂ Flux (mmol m⁻² d⁻¹)	Full	0.0 ± 6.3	-1.3 ± 5.9	0.3028	34.1 ± 29.0	2.7	189.0
	Sampling Period						
	Winter	-14.9 ± 8.4	-19.1 ± 7.7	0.0676	46.6 ± 38.9	2.7	189.0
	Spring	7.6 ± 5.2	7.0 ± 5.2	0.6680	27.5 ± 18.5	4.9	115.0
	Summer	9.4 ± 5.6	11.7 ± 5.2	0.1167	32.3 ± 22.9	4.5	111.0
	Fall	0.1 ± 3.8	-0.3 ± 3.5	0.7449	17.0 ± 10.2	3.9	40.1

333

334 Table 3. Tests examining differences in mean carbonate system parameters between
 335 seasons and between types of sampling (continuous monitoring with sensors Nov. 8 2016
 336 – Aug 23, 2017, discrete sample collection and laboratory measurement during only the
 337 continuous monitoring period Nov. 8 2016 – Aug 23, 2017, and discrete sample
 338 collection and laboratory measurement during the entire sampling period May 2, 2014-
 339 Feb. 25, 2020). For both the two-way ANOVA and associated one-way ANOVAs, p-
 340 values are listed. All significant results based on $\alpha=0.05$ are bolded, and the F statistic is
 341 in parentheses. Since all two-way ANOVAs had a significant interaction between factors,
 342 individual one-way ANOVAs were conducted for each level of the other factor.
 343 Following significant one-way ANOVAs, multiple comparisons using the Westfall
 344 adjustment (Westfall, 1997) were conducted; individual comparisons with significantly
 345 different means (based on $\alpha=0.05$) are listed as unequal beneath the one-way ANOVA
 346 results (All \neq indicates that every individual comparison between levels had significantly
 347 different means. W = winter, Sp = spring, Su = summer, F = fall; C = continuous sensor
 348 data, D = discrete sample data over the entire discrete monitoring period, D_C = Discrete
 349 sample data during only the period of continuous monitoring).

Parameter	Two-way ANOVA			One-way ANOVA and post-hoc multiple comparison results for differences between types of sampling				One-way ANOVA and post-hoc multiple comparison results for difference between seasons		
	Interaction	Season	Sampling type	winter	spring	summer	fall	Continuous	Discrete (Continuous Period)	Discrete (Entire Period)



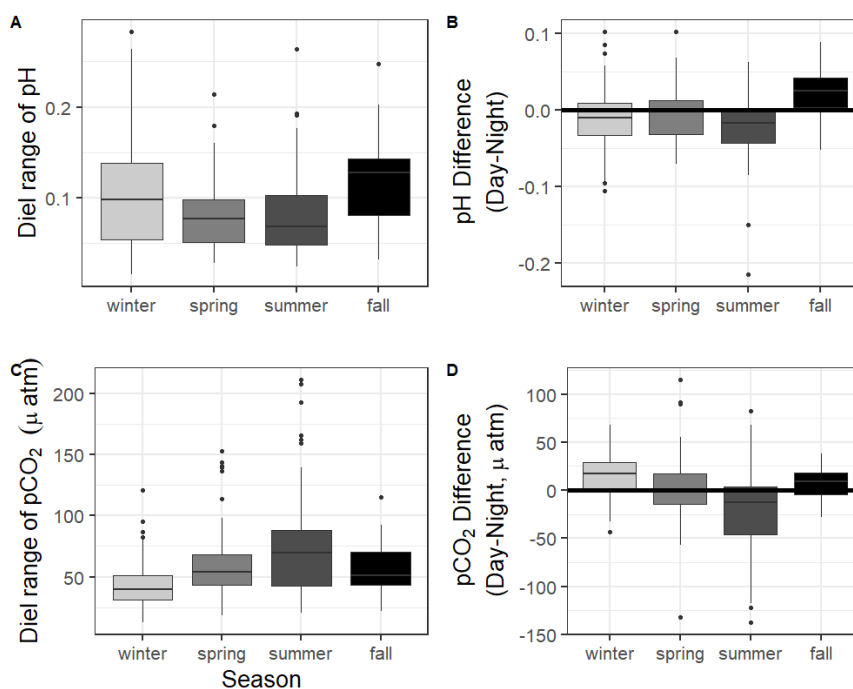
Temp (°C)	<0.0001 (15.8)	<0.0001 (12369.7)	0.7346 (0.3)	0.0710 (2.6)	0.1052 (2.3)	<0.0001 (19.6)	<0.0001 (61.4)	<0.0001 (12559)	<0.0001 1 (22.8)	<0.0001 (58.2)
						D≠C	D≠C	All≠	W≠Su; W≠Sp; W≠F; Su≠Sp; Su≠F	All≠
Salinity	0.0141 (2.7)	<0.0001 (598.7)	0.6509 (0.4)	0.1716 (1.8)	0.0013 (6.7)	0.1921 (1.7)	0.7007 (0.4)	<0.0001 (580.0)	0.2516 (1.6)	<0.0001 (17.5)
					D≠C			W≠Su; W≠F; Su≠Sp; Su≠F; Sp≠F		W≠Su; Su≠Sp; Su≠F
pH	0.0013 (3.7)	<0.0001 (1412.3)	<0.0001 (24.0)	0.4026 (0.9)	0.9238 (0.1)	<0.0001 (24.1)	<0.0001 (33.2)	<0.0001 (1381.2)	0.0152 (5.7)	<0.0001 (35.3)
						D≠C C≠Dc	D≠C	All≠	W≠Su	W≠Su; W≠Sp; W≠F; Su≠Sp; Su≠F
pCO₂ (µatm)	<0.0001 (10.4)	<0.0001 (1747.3)	0.0147 (4.2)	0.0018 (6.4)	<0.0001 1 (17.4)	0.0002 (8.4)	0.0398 (3.2)	<0.0001 (1737.6)	0.0407 (4.0)	<0.0001 (8.4)
				D≠C	D≠C	D≠C	All=	All≠	W≠Su	W≠Su; W≠Sp; W≠F; Su≠Sp; Su≠F
CO₂ Flux (mmol m⁻² d⁻¹)	0.0144 (2.6)	<0.0001 (738.1)	0.6739 (0.4)	0.9140 (0.1)	<0.0001 (11.8)	0.0214 (3.9)	0.5849 (0.5)	<0.0001 (725.9)	0.0299 (4.5)	<0.0001 (19.2)
					D≠C	D≠C		All≠	W≠Su	W≠Su; W≠F; Su≠Sp; Su≠F

350

351 There was substantial diel variability in parameters (Table 2, Fig. 4). Over the 10-
 352 month in-situ monitoring period, temperature had a mean diel range (daily maximum
 353 minus daily minimum) of $1.3 \pm 0.8^\circ\text{C}$ (Table 2). Daytime and nighttime temperature
 354 differed significantly during the summer and fall months, with higher temperatures at
 355 night for both seasons (Table 2). The mean diel range of salinity was 3.4 ± 2.7 (Table 2).
 356 Daytime and nighttime salinity differed significantly during the winter and fall months,
 357 with higher salinities at night for both seasons. The mean diel range of pH was $0.09 \pm$



358 0.05 (Table 2). Daytime and nighttime pH differed significantly during the winter,
359 summer, and fall months; nighttime pH was significantly higher than that of the daytime
360 during the summer and winter months, and daytime pH was significantly higher during
361 the fall (Table 2, Fig. 4). The mean diel range of $p\text{CO}_2$ was $58 \pm 33 \mu\text{atm}$ (Table 2, Fig.
362 4). Daytime and nighttime $p\text{CO}_2$ differed significantly during the winter and summer
363 months; nighttime $p\text{CO}_2$ was significantly higher than that of the daytime during the
364 summer and daytime $p\text{CO}_2$ was significantly higher during the winter (Table 2, Fig. 4).

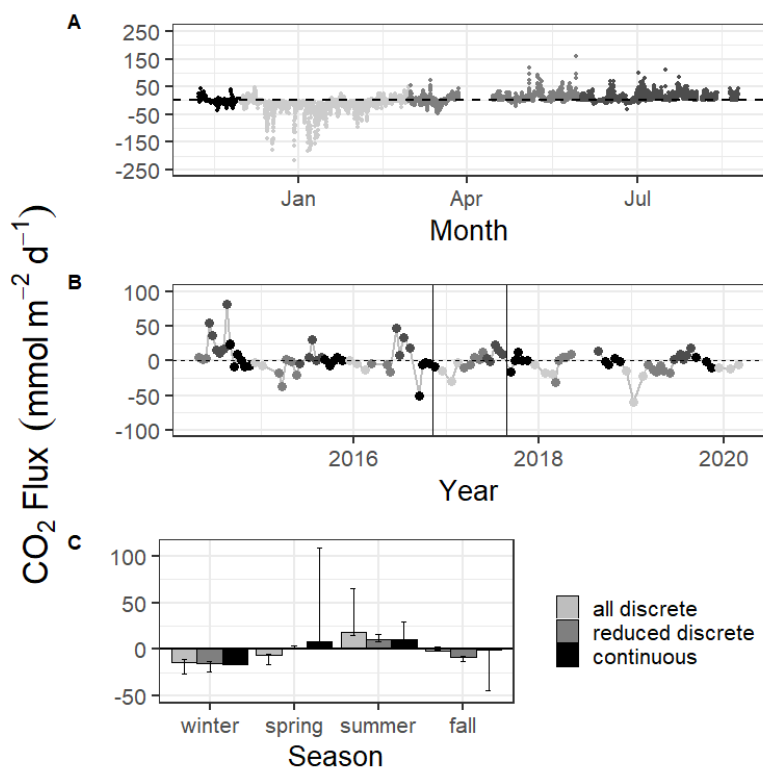


365
366 Figure 4. Boxplots showing the diel range (maximum minus minimum) and difference in
367 daily parameter mean daytime minus nighttime measurements for pH and $p\text{CO}_2$ from
368 continuous sensor data.
369

370 The seasonally weighted mean CO_2 flux calculated from sensor data across the
371 entire monitoring period was $0.2 \pm 23.7 \text{ mmol m}^{-2} \text{ d}^{-1}$ (Table 1). Mean CO_2 flux differed
372 by season (Table 3). Winter and fall both had net negative CO_2 flux (winter was most



373 negative), and summer and spring both had a net positive CO₂ flux (summer was most
374 positive) (Table 1, Fig. 5). CO₂ flux also fluctuated on a daily scale, with the mean diel
375 range (daily maximum – minimum) over the entire monitoring period being 34.1 ± 29.0
376 $\text{mmol m}^{-2} \text{d}^{-1}$ (Table 2). However, there was not a significant difference in CO₂ flux
377 calculated for daytime versus nighttime hours for the entire monitoring period or any
378 individual season based on $\alpha=0.05$ (paired t-test, Table 2).



379 **Figure 5.** CO₂ flux calculated over the sampling periods from continuous (A) and
380 discrete (B) data using the Jiang et al. (2008) wind speed parameterization. Gray scale in
381 (A) and (B) denote different seasons. Vertical lines in (B) denote the time period of
382 continuous monitoring. (C) shows the seasonal mean CO₂ flux calculated using the Jiang
383 et al. (2008) gas transfer velocity parameterization and error bars representing mean CO₂
384 flux calculation using Ho and Raymond and Cole windspeed parameterizations. The
385 different color bars within each season represent all discrete data (May 2, 2014- Feb. 25,
386 2020), reduced discrete data (Nov. 8 2016 – Aug 23, 2017, to overlap with continuous
387 monitoring), and continuous sensor data (Nov. 8 2016 – Aug 23, 2017).
388
389

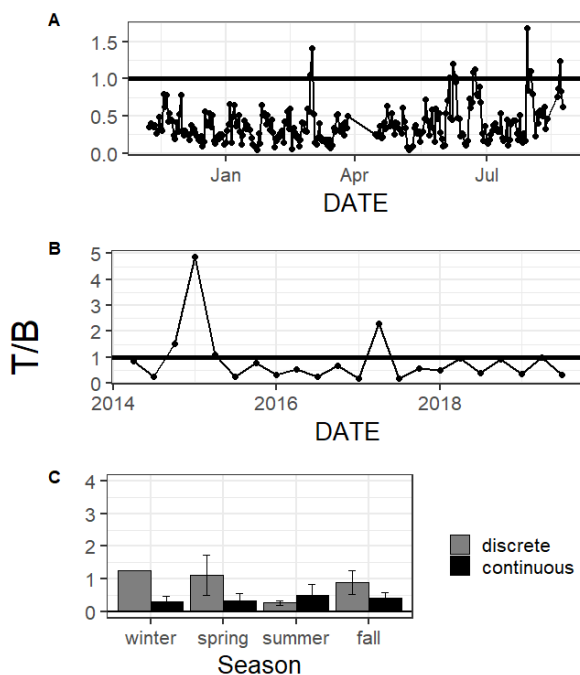


390 The relative influence of thermal and nonthermal factors (T/B) in controlling
 391 $p\text{CO}_2$ varied over different time scales (Table 4, Fig. 6). T/B calculated from sensor data
 392 for the entire period was 0.98, indicating that the magnitude of control of non-thermal
 393 processes on $p\text{CO}_2$ was slightly greater than that of temperature. Within seasons, T/B
 394 calculated from sensor data ranged from 0.51 in the winter to 0.69 in the spring, showing
 395 that non-thermal processes exert more control on $p\text{CO}_2$ within each individual season
 396 (Table 4). On a daily scale, only 11 of the 178 days with measurements for all 24 hours
 397 had temperature control of $p\text{CO}_2$ exceeding the non-thermal control (Table 4, Fig. 6).

398 **Table 4.** Thermal versus non-thermal control on $p\text{CO}_2$ (Takahashi et al. 2002) over
 399 different time scales using both continuous sensor data and discrete sample data
 400 (indicated as Sampling Type C and D, respectively). If more than one segment of time is
 401 being considered ($n > 1$), $\Delta p\text{CO}_2$ values are the mean \pm standard deviation of all segments,
 402 the T/B values are the minimum and maximum T/B, and the number out of n with T/B > 1
 403 (indicating greater control of $p\text{CO}_2$ by temperature than other processes) is recorded. The
 404 summary of annual T/B values from discrete data includes only 2015–2019 ($n = 5$ years;
 405 2014 and 2020 were omitted since monitoring did not occur throughout the entire year).
 406 Daily values from continuous data were only reported for those days with all 24
 407 measurements.

Time Period / Scale	Sampling type	n	$\Delta p\text{CO}_2, \text{thermal}$ (μatm)	$\Delta p\text{CO}_2, \text{nonthermal}$ (μatm)	T/B	Number out of n with T/B > 1
Full Monitoring Period (May 2, 2014– Feb. 25, 2020)	D	1	301.9	537.8	0.56	
Annual	D	5	259.3 ± 16.0	319.1 ± 130.9	0.48 – 1.17	2/5
Continuous Monitoring Period (Nov 2016 – August 2017)	C	1	355.0	360.7	0.98	
Winter	C	1	168.2	328.4	0.51	
	D	6	42.2 ± 23.4	101.7 ± 78.7	0.20 – 4.90	1/6
Spring	C	1	171.4	246.9	0.69	
	D	6	142.3 ± 53.7	147.8 ± 67.3	0.59 – 2.42	2/6
Summer	C	1	100.2	179.9	0.56	
	D	6	46.9 ± 26.6	176.9 ± 108.3	0.21 – 0.35	0/6
Fall	C	1	105.9	181.6	0.58	
	D	6	179.8 ± 59.5	176.6 ± 78.1	0.59 – 3.06	2/6
Daily	C	178	21.8 ± 11.8	63.8 ± 30.3	0.05 – 1.68	11/178

408



409
410 **Figure 6.** T/B (thermal $p\text{CO}_2$ / non-thermal $p\text{CO}_2$) calculated for each day from
411 continuous (A) and each season from discrete (B) data. Bar graphs showing the seasonal
412 mean and standard deviation of T/B from both discrete and continuous data (C).
413

414 Tidal fluctuations had a significant effect on carbonate system parameters (Table
415 5). Both temperature and salinity were higher at low tide during the winter and summer
416 months and higher at high tide during the spring. pH was higher at high tide during the
417 winter and summer and higher at low tide during the spring, and $p\text{CO}_2$ was higher during
418 low tide during winter, spring, and summer (Table 5). CO_2 flux also varied with tidal
419 fluctuations. CO_2 flux was higher in the low tide condition for all season with tide data;
420 the location was less of a CO_2 sink during low tide conditions in the winter and more of a
421 CO_2 source during low tide conditions in the spring and summer.

422 **Table 5.** Differences in temperature, salinity and mean carbonate system parameters from
423 continuous sensor data between high tide and low tide. High tide was defined as a tide
424 level greater than Q3 and low tide was defined as a tide level less than Q1. Seasons were
425 examined separately with t-tests because of a significant interaction (based on $\alpha=0.05$)



426 between the season and high/low tide factors in a two-way ANOVA. Fall was omitted
 427 from the analysis because tide data was only available at the location beginning
 428 December 20, 2016.
 429

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

430

431 3.2 Discrete sampling results

432 All results reported here are for the entire 5+ years of monitoring; the subset of
 433 discrete sample data that overlaps with the continuous monitoring period will be
 434 addressed only in the discussion for method comparisons. All reported discrete sampling
 435 parameters showed substantial temporal variability over the 5+ years of monitoring (Fig.
 436 2E-H). The mean temperature was 24.1 ± 5.3°C, ranging from 11.8 – 31.2°C; the mean
 437 salinity was 30.1 ± 4.4, ranging from 16.7 – 37.5; the mean pH was 8.079 ± 0.092,
 438 ranging from 7.693 to 8.354; and the mean pCO₂ was 406 ± 100 µatm, ranging from 199
 439 to 1043 (Table 1). These parameters all experienced significant seasonal variability
 440 (Tables 1 and 3). Temperature was significantly different between each season, highest in
 441 summer and lowest in winter (Tables 1 and 3). Salinity was highest during the summer
 442 months and was not significantly different between other seasons (Tables 1 and 3). pH
 443 and pCO₂ were both significantly different between all seasons with the exception of



444 spring and fall (Table 3). Winter had both the highest seasonal pH (8.162 ± 0.065) and
445 lowest seasonal $p\text{CO}_2$ ($331 \pm 39 \mu\text{atm}$), and summer had both the lowest seasonal pH
446 (7.975 ± 0.046) and highest seasonal $p\text{CO}_2$ (511 ± 108) (Tables 1 and 3, Fig. 3).

447 Average annual CO_2 flux calculated with discrete sample data was slightly
448 negative ($-0.9 \pm 18.7 \text{ mmol m}^{-2} \text{ d}^{-1}$, Table 1). CO_2 flux varied greatly by season. Summer,
449 the only season with a net positive CO_2 flux over the 5+ year period, had significantly
450 higher flux than all other seasons; winter had the lowest calculated flux, but it was not
451 significantly different from spring (Tables 1 and 3).

452 As with the continuous data, T/B calculated from the discrete data varied over
453 different time scales (Table 4, Fig. 6). For the entire period, T/B was 0.56, indicating that
454 non-thermal processes exerted more control than temperature on $p\text{CO}_2$. The annual T/B
455 varied from 0.48 to 1.17, with two of the five sampled years having T/B greater than one
456 (i.e. more thermal influence). While the majority of individual seasons that were sampled
457 experienced stronger non-thermal control on $p\text{CO}_2$ (T/B < 1), the only season that never
458 experienced stronger thermal control was summer, with summer T/B values ranging from
459 0.21 – 0.35 for the 6 sampled years (Table 4).

460 Discussion

461 4.1 Factors controlling temporal variability in carbonate system parameters

462 4.1.1 Thermal versus non-thermal control of $p\text{CO}_2$

463 Substantial variability in the carbonate system was observed at the study site over
464 multiple time scales including diel, seasonal, and interannual. Many physical and
465 biological factors (e.g., temperature, currents, tides, wind speed, net ecosystem
466 metabolism, etc.) can exert control on $p\text{CO}_2$ and subsequently exert control on other



467 carbonate system parameters. Using the thermal versus non-thermal analysis of control
468 on $p\text{CO}_2$ from Takahashi et al. (2002), we were able to determine that non-thermal
469 processes generally exert more control on the $p\text{CO}_2$ in the Aransas Ship Channel relative
470 to temperature over multiple time scales (Table 4, $T/B < 1$). Only five of 24 seasons (one
471 winter, two spring, and two fall) throughout the years of discrete sampling had greater
472 variability in $p\text{CO}_2$ attributed to temperature ($\Delta p\text{CO}_{2, \text{thermal}}$) than other processes ($\Delta p\text{CO}_{2, \text{nonthermal}}$)
473 (nonthermal) (Table 4). The magnitude of $p\text{CO}_2$ variation attributed to non-thermal processes
474 varied greatly over multiple time scales (i.e. $\Delta p\text{CO}_{2, \text{nonthermal}}$ had large standard
475 deviations, Table 4). For example, in 2016 $p\text{CO}_2$ had the strongest non-thermal control of
476 any year, with a $\Delta p\text{CO}_{2, \text{nonthermal}}$ of 538 μatm , while 2019 had the weakest control from
477 non-thermal processes of any year, with a $\Delta p\text{CO}_{2, \text{nonthermal}}$ of 208. Conversely, the
478 magnitude of $p\text{CO}_2$ variation attributed to temperature was consistent across time scales.
479 For example, in 2015 $p\text{CO}_2$ had the strongest thermal control of any year, with a $\Delta p\text{CO}_{2, \text{thermal}}$
480 of 276 μatm , while 2019 had the weakest thermal control of any year, with a
481 $\Delta p\text{CO}_{2, \text{thermal}}$ of 243 μatm .

482 The difference in T/B between sampling methods is relatively small over the 10-
483 month sensor deployment period (Table 4). Each method suggested temperature and
484 nonthermal processes exert a relatively similar control on $p\text{CO}_2$, but continuous
485 monitoring demonstrated a greater magnitude of fluctuation resulting from both
486 temperature and non-thermal processes (i.e. greater $\Delta p\text{CO}_{2, \text{thermal}}$ and $\Delta p\text{CO}_{2, \text{nonthermal}}$).
487 Over shorter time scales, like individual seasons, the calculated T/B did not align well
488 between sampling methods. The T/B (calculated using only data from Nov 2016 –
489 August 2017) for discrete versus continuous sampling was respectively 0.17 versus 0.51



490 for winter, 2.37 versus 0.69 for spring, and 0.20 versus 0.56 for summer (Table 4, Fall
491 was omitted since there was only one discrete sampling event during the fall of the
492 continuous monitoring period). Sampling bias due to the small number of within-season
493 sampling events for the discrete monitoring likely resulted in this difference. From both
494 continuous and discrete data, summers always had stronger control exerted on $p\text{CO}_2$ from
495 nonthermal processes than temperature. While temperatures were high during the
496 summer months, the within-season variability in temperature was the lowest (Table 1);
497 less of a temperature swing resulted in less thermal control on the system. Conversely,
498 spring and fall seasons, which experienced the greatest temperature swings (Table 1), had
499 greater relative temperature control exerted on $p\text{CO}_2$ (Table 4). The differences in
500 $\Delta p\text{CO}_{2, \text{thermal}}$ and $\Delta p\text{CO}_{2, \text{nonthermal}}$ between monitoring methods illustrate that there is
501 information that is missed when only sampling bimonthly/monthly and during the
502 daytime. Generally, both $\Delta p\text{CO}_{2, \text{thermal}}$ and $\Delta p\text{CO}_{2, \text{nonthermal}}$ are higher when calculated
503 from sensors than discrete sampling, indicating that the extremes are generally not
504 captured by the discrete sampling and sensor data would provide a better understanding
505 of system controls.

506 The relative importance of thermal versus non-thermal controls may be modulated
507 by tide level. The influence of tides can be removed from the calculated non-thermal
508 $p\text{CO}_2$ term, leaving only biological processes and other physical controls in the non-
509 thermal term, by examining periods of high tide and low tide separately. Using our sensor
510 data and the same water level data used for the tide analysis, we found that T/B is higher
511 during the high tide condition within each season. T/B for high tide and low tide,
512 respectively, was 0.60 and 0.52 for winter, 0.84 and 0.66 for spring, and 0.62 and 0.58 for



513 summer. The higher control exerted by nonthermal processes during low tide seems
514 intuitive given that there is less volume of water for the end products of biological
515 processes to build up in. The difference in T/B between high tide and low tide conditions
516 was greatest in the spring, likely due to a combination of elevated spring-time
517 productivity and larger tidal ranges in the spring.

518 Using data from the first year of our discrete sampling (May 2014 – April 2015),
519 Yao and Hu (2017) reported that the Aransas Ship Channel T/B was 1.53 during drought
520 and 1.79 during a period of flooding, both of which are significantly higher than what we
521 found over most timescales (the exception being certain individual seasons, mostly
522 during that first year of sampling, Table 4, Fig. 6B). Yao and Hu (2017) also found that
523 locations in the upper estuary experienced lower T/B during flooding conditions than
524 drought conditions, but the opposite was found for the Aransas Ship Channel location,
525 where the flooding conditions had higher T/B. It is likely that the high T/B calculated by
526 Yao and Hu (2017) was a result of the drought condition at the beginning of their
527 sampling; given the long residence time of MAE, the Aransas Ship Channel may not
528 have experienced the influence of the freshwater inflow by the end of the Yao and Hu
529 (2017) study. Once the freshwater reached the Aransas Ship Channel location, it would
530 likely experience a reduced T/B as did the upper parts of the system. Since then, there has
531 not been another significant drought in the system, so it seems as though the non-thermal
532 controls on $p\text{CO}_2$ are more important at this location under normal freshwater inflow
533 conditions.



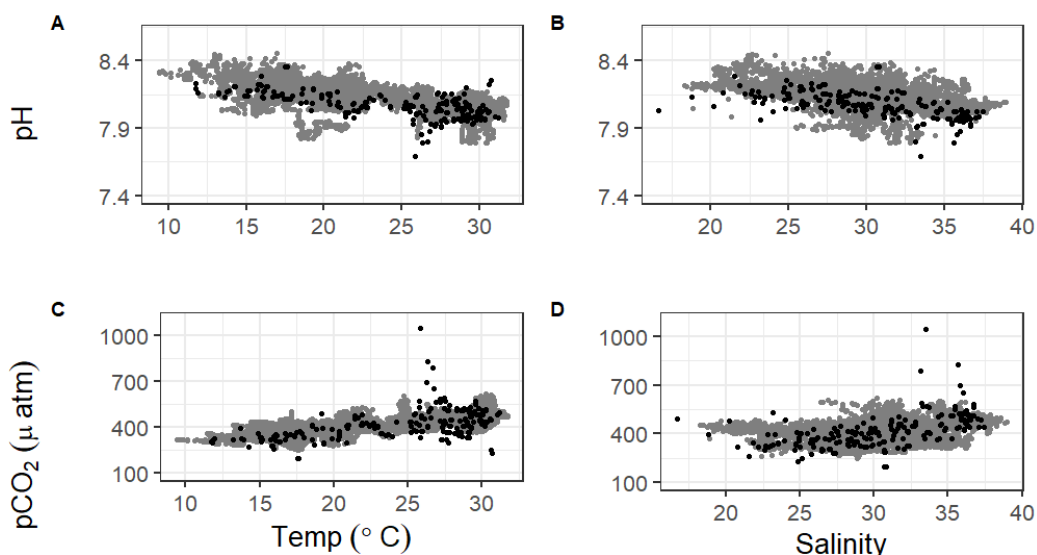
534 *4.1.2 Investigating controls on the carbonate system using relationships between*
535 *carbonate system parameters and other environmental parameters*

536 We further investigated controls on the carbonate system using tide and
537 windspeed data (obtained from NOAA's Aransas Pass station at
538 <https://tidesandcurrents.noaa.gov/>) and dissolved oxygen, PAR, turbidity, and chlorophyll
539 fluorescence data (obtained from the MANERR at
540 <https://missionaransas.org/science/download-data>) along with our continuous and discrete
541 data. All investigations of relationships between environmental parameters discussed
542 below included only the observations with no significant water column stratification
543 (defined as a salinity difference of less than 3 between surface water from our YSI and
544 bottom water (>5 m) from the MANERR's YSI). This omission of stratified water was
545 intended to omit instances of substantial differences in chemical parameters between the
546 surface and bottom water since all MANERR environmental data used in our analysis
547 were measured at depth while our sensors measured surface water. Omitting stratified
548 water reduced our continuous dataset from 6088 to 5524 observations, and omitting
549 observations where there were no MANERR data to determine stratification further
550 reduced the dataset to 4112 observations. Similarly, removing instances of stratification
551 reduced discrete sample data from 104 to 89 surface water observations.

552 To extend upon the above discussion of thermal versus non-thermal controls on
553 $p\text{CO}_2$, the extent of thermal control on both pH and $p\text{CO}_2$ can be investigated based on
554 relationships between parameters. There is a strong negative correlation between pH and
555 temperature and a strong positive correlation between $p\text{CO}_2$ and temperature (Table 6,
556 Fig. 7). The direction of these relationships (sign of the correlation coefficient) at the



557 Aransas Ship Channel was the same as in open ocean waters despite these relationships
558 not being consistent across different estuarine environments (N. Rosenau, personal
559 communications). The strong correlations with temperature support our findings that
560 thermal controls on $p\text{CO}_2$ can be important over multiple time scales. Significantly
561 warmer water temperatures were observed during the nighttime in both summer and fall
562 (Table 2, Fig. 8), indicating that temperature could exert a slight control on the carbonate
563 system over a diel time scale. More substantial temperature swings between seasons
564 indicate that temperature is more important over seasonal time scales (Table 1). In
565 addition to direct thermal control at our site, the strong correlations with temperature are
566 likely derived from changes in net community metabolism associated with temperature
567 (Caffrey, 2004). For example, the strong negative correlation between nonthermal $p\text{CO}_2$
568 and temperature (Table 6) is likely indicative of enhanced primary productivity in
569 warmer waters.



• continuous • discrete

570



571 Figure 7. Correlations of pH and $p\text{CO}_2$ with temperature and salinity from continuous
 572 sensor data (gray) and all discrete data (black).
 573

574 Table 6. Pearson correlation coefficients between surface water carbonate system
 575 parameters and other water quality and environmental parameters for both continuous
 576 sensor data and discrete sample data (entire sampling period). Only observations without
 577 significant stratification in the water column were included in these analyses. Parameter
 578 pairs with a significant correlation based on $\alpha=0.05$ have a correlation coefficient
 579 reported. Asterixis are used to indicate the level of significance of the correlation, *
 580 $p<0.05$, ** $p<0.01$, *** $p<0.0001$. The correlation coefficient is listed as 0 if the
 581 relationship was not significant. N/A is listed when the analysis was omitted because the
 582 environmental parameter did not have observations corresponding to the date and time of
 583 at least half of our discrete sample measurements (45 observations).

	pH		$p\text{CO}_2$		$p\text{CO}_2$, nonthermal	
	Continuous	Discrete	Continuous	Discrete	Continuous	Discrete
Temperature (°C)	-0.55 ***	-0.59 ***	0.75 ***	0.53 ***	-0.73 ***	-0.45 ***
Salinity	-0.47 ***	-0.74 ***	0.53 ***	0.69 ***	-0.28 ***	0.35 **
Wind Speed (m s^{-1})	-0.04 **	N/A	0.15 ***	N/A	0	N/A
Dissolved Oxygen (mg L^{-1})	0.55 ***	0	-0.81 ***	0	0.45 ***	0
Tide Level (m)	0	0	-0.15 ***	0	-0.15 ***	-0.55 **
Turbidity	-0.08 ***	N/A	-0.14 ***	N/A	-0.28 ***	N/A
Fluor. Chlorophyll	0.12 ***	N/A	-0.22 ***	N/A	0.34 ***	N/A

584
 585 Though annual average $p\text{CO}_2$ and CO_2 flux are higher in the upper estuary and
 586 lower offshore than at our study site, the same seasonal pattern of elevated $p\text{CO}_2$ and
 587 positive CO_2 flux in the summer and depressed $p\text{CO}_2$ and negative CO_2 flux during the
 588 winter observed at our site has also been observed throughout the entire MAE and in the
 589 open Gulf of Mexico (Hu et al., 2018; Yao and Hu, 2017). Seasonal fluctuations in pH
 590 and $p\text{CO}_2$ are low at our study site relative to other systems that have been studied to date
 591 (Carstensen et al., 2018; Yao and Hu, 2017), which may be in part due to the relatively
 592 small seasonal temperature changes (Table 1) in this warm, semiarid environment.
 593 Despite substantial seasonal thermal control at our site, simple linear regressions indicate
 594 that temperature had substantially higher explanatory value for pH and $p\text{CO}_2$ in offshore
 595 GOM waters ($R^2 = 0.81$ and 0.78 , respectively (Hu et al., 2018)) than at our site ($R^2 =$



596 0.30 and 0.52, respectively, for sensor data and $R^2 = 0.38$ and 0.25, respectively, for
597 discrete data).

598 Other physical factors that may exert control on the carbonate system (including
599 windspeed, salinity, tide level, and turbidity) can also be investigated through parameter
600 relationships. We investigated wind speed as a possible control on the carbonate system
601 to gain insight into the effect of wind-driven CO_2 fluxes on the inventory of CO_2 in the
602 water column (and subsequent impacts to the entire carbonate system). The Texas coast
603 has relatively high wind speeds, with the mean wind speed observed during our
604 continuous monitoring period being 5.8 m s^{-1} . While this results in relatively high
605 calculated CO_2 fluxes (Fig. 5), the seasonal relationship between $p\text{CO}_2$ and windspeed
606 does not support a change in inventory with higher winds. Linear regression analysis
607 within each season reveals that winter, spring, and fall all experience increases in $p\text{CO}_2$
608 with increasing wind, while there is not a significant relationship in summer. Since spring
609 and summer both have a mean estuarine $p\text{CO}_2$ greater than atmospheric level (and
610 positive CO_2 flux, Table 1) a negative relationship between windspeed and $p\text{CO}_2$ would
611 be necessary to support this hypothesis.

612 Previous studies have indicated that freshwater inflow may exert a primary
613 control on the carbonate system in the estuaries of the northwestern GOM (Hu et al.,
614 2015; Yao et al., 2020; Yao and Hu, 2017). Increased freshwater inflow resulting from
615 storms has also been shown to increase community respiration, which would
616 subsequently increase $p\text{CO}_2$, in the upper reaches of the MAE (Bruesewitz et al., 2013).
617 MAE is also known to experience large swings in the chemistry of its freshwater inputs,
618 with relatively high levels of dissolved inorganic carbon and total alkalinity during base



619 flows but much lower levels due to dilution during intense flooding (Yao et al., 2020).
620 Given the location of our sampling in the lower portion of the estuary and the long
621 residence time in the system, we will not directly address freshwater inflows as a
622 controlling factor, but the influence of freshwater inflow may be evident in the response
623 of the system to changes in salinity. Carbonate system variability is much lower at our
624 study site than it is in the more upper reaches of MAE, likely due to the lesser influence
625 of freshwater inflow and its associated changes in biological activity at the Aransas Ship
626 Channel (Yao and Hu, 2017). Salinity from both sensor and discrete monitoring was
627 strongly correlated with pH and $p\text{CO}_2$, with correlation coefficients nearing (continuous)
628 or surpassing (discrete) that of the correlations with temperature (Fig. 7; Table 6). Periods
629 of lower salinity had higher pH and lower $p\text{CO}_2$, likely due to enhanced freshwater
630 influence and subsequent elevated primary productivity at the study site. Fluctuating
631 salinity at the Aransas Ship Channel may also result from direct precipitation,
632 stratification, and tidal fluctuations. Based on the simple linear regression of salinity with
633 tide level, there is a significant ($p < 0.0001$) relationship between tide level and salinity,
634 but the amount of variability in salinity that tides can explain (based on model R^2) is only
635 about 2%.

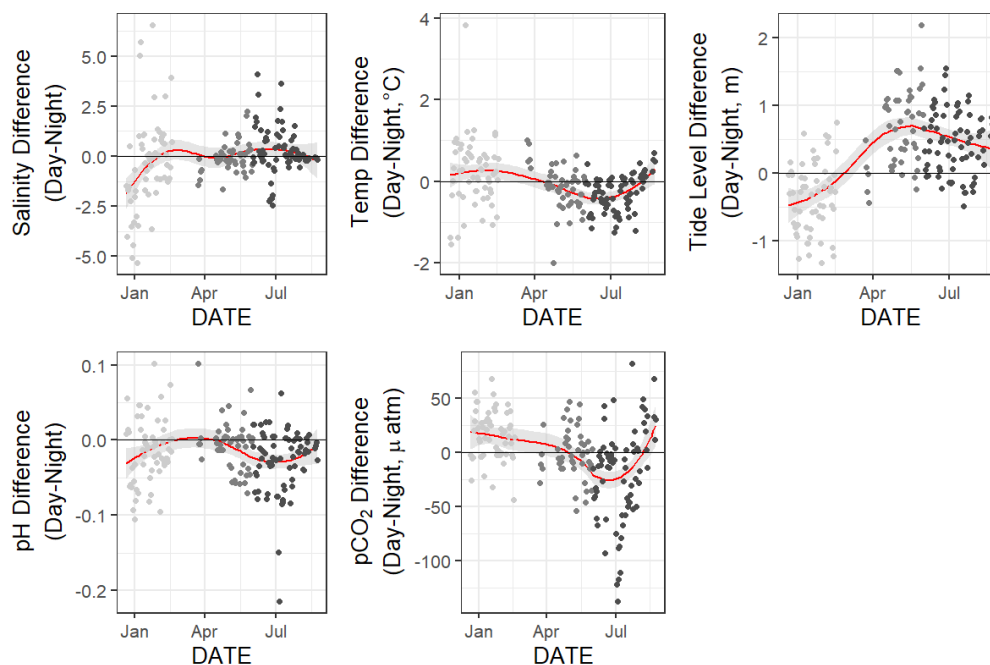
636 Tidal fluctuations were clearly important to carbonate system variability at the
637 Aransas Ship Channel (Table 5). While the northwestern GOM estuaries are generally
638 microtidal, the constricted tidal inlets such as the Aransas Ship Channel may experience
639 relatively large tidal fluctuations. The water level data used in this analysis came from a
640 location directly offshore from our study site, and water level had a range of 1.30 m
641 (maximum – minimum recorded water level) over the 10-month continuous monitoring



642 period. Mean water level varied between all seasons; mean spring (highest) water levels
643 were on average 0.08 m higher than winter (lowest) water levels (ANOVA $p < 0.0001$, fall
644 was not considered because of a lack of water level data). Tidal influence on pH was less
645 clear. Data from continuous monitoring did not show a significant correlation between
646 pH and tide level across the entire monitoring period (Table 6). Significant differences in
647 mean pH between tide levels were recorded during each season; pH was higher at high
648 tide (corresponding with the lower $p\text{CO}_2$) during the winter and summer, but pH was
649 lower at high tide (conflicting with the lower $p\text{CO}_2$) in the spring (Table 5). This
650 separation between water level correlation with pH and $p\text{CO}_2$ suggests that different
651 controlling factors of the carbonate system may not be exerted equally on both $p\text{CO}_2$ and
652 over different timescales. Similar to pH, both temperature and salinity experienced
653 seasonally dependent reversals in their difference between tide levels during the spring;
654 each were higher at low tide during winter and summer and higher at high tide during
655 spring (Table 5). Given the negative relationship of both temperature and salinity with
656 pH, it is likely these parameters became important controls on pH in the spring.
657 To help examine controls on the carbonate system on a diel time scale, we used loess
658 models (locally weighted polynomial regression) to identify changes in diel patterns over
659 the course of our monitoring period (Fig. 8). Both tidal and biological controls on the
660 carbonate system can operate on a diel time scale. The GOM is one of the few places in
661 the world that experiences diurnal tides (Seim et al., 1987; Thurman, 1994), so
662 theoretically, the fluctuations in $p\text{CO}_2$ associated with tides may align to either amplify or
663 reduce/reverse the fluctuations that would result from diel variability in net community
664 metabolism. The mean daily tidal fluctuation during our continuous monitoring period



665 was $0.39 \text{ m} \pm 0.13 \text{ m}$, which did not significantly differ between seasons (ANOVA
666 $p=0.739$). However, diel patterns in tidal fluctuations exhibited a strong seasonal pattern
667 during the continuous monitoring period, with spring and summer having higher tide
668 level during the daytime and winter having higher tide level during the nighttime (Fig. 8).
669 This same seasonal pattern in diel tidal fluctuations is exhibited from Dec 20, 2016 (when
670 the tide data is first available) through the rest of our discrete monitoring period (Feb 25,
671 2020), indicating that tidal control on diel variability of carbonate system parameters was
672 likely consistent throughout this time period.



673 Figure 8. Loess models (red line) and their confidence intervals (gray band around red
674 line) showing the difference in daily parameter mean daytime minus nighttime
675 measurements. The gray scale of the data points represents the four seasons over which
676 data were collected.
677
678

679 Based on diel tidal fluctuations at this site, tidal control should amplify the
680 biological control signal (nighttime $p\text{CO}_2 >$ daytime $p\text{CO}_2$) during spring and summer



681 and reduce or reverse the biological control signal during the winter. This was supported
682 by our $p\text{CO}_2$ data, which showed nighttime $p\text{CO}_2$ significantly greater than daytime $p\text{CO}_2$
683 in the summer (Table 2). The full reversal of the biological signal in the winter (Table 2,
684 nighttime $p\text{CO}_2 < \text{daytime } p\text{CO}_2$) indicated that biological activity was not the strongest
685 controlling factor on the diel time scale and was likely exceeded by tidal control. Winter
686 also had higher daytime temperature (Table 3), which could also contribute to the higher
687 daytime $p\text{CO}_2$, while summer diel temperature and tides would act to amplify the
688 biological signal.

689 Again, the diel variability in pH did not mirror $p\text{CO}_2$ as would be expected. The
690 loess models show that daily variability in pH closely mirrors that of temperature while
691 the daily variability in $p\text{CO}_2$ much closer reflects the tide level (Fig. 8), indicating that
692 controlling factors of the carbonate system may not be exerted equally on both pH and
693 $p\text{CO}_2$.

694 The extent of biological control on the system can also be investigated based on
695 correlations between carbonate system parameters and dissolved oxygen (DO).
696 Respiration-driven acidification is one of the most important local to regional
697 contributors to acidification in coastal waters, with acidification closely linked to the
698 widespread issue of deoxygenation (Rabalais et al., 2014; Strong et al., 2014). There were
699 no observations of hypoxia at our study site during our monitoring, with minimum DO
700 levels of 3.9 mg L^{-1} and 4.0 mg L^{-1} for our continuous monitoring period and our discrete
701 sampling period, respectively. Despite the lack of hypoxia, there was a strong
702 relationship between the carbonate system parameters and DO (Table 6), suggesting that



703 net ecosystem metabolism may exert an important control on the carbonate system on
704 certain time scales.

705 There was no significant difference in daytime and nighttime DO during any
706 season (paired t-tests, winter $p = 0.1573$, spring $p = 0.4877$, summer $p = 0.794$) despite
707 the significant differences in pH and $p\text{CO}_2$ between daytime and nighttime (Table 2).
708 This suggests that net community metabolism is likely not a strong controlling factor of
709 carbonate system parameters at this site on a diel time scale. The control exerted on the
710 carbonate system by biological processes is likely much greater on the seasonal scale
711 than the diel scale. The correlation between continuous $p\text{CO}_2$ and DO is stronger than
712 $p\text{CO}_2$ and temperature, which suggests strong biological control and supports the
713 indication by T/B values that non-thermal processes exert more control on $p\text{CO}_2$ than
714 temperature. Both types of sampling (i.e., continuous and discrete) demonstrate that pH is
715 generally highest in the winter and lowest in the summer and $p\text{CO}_2$ is highest in the
716 summer and lowest in the winter (Figs. 2, and 3, Table 1). Though this seasonal pattern
717 corresponds with the directional response from temperature fluctuations, it can also be
718 explained by biological activity.

719 Given that this sampling location is in a ship channel where boat traffic (including
720 large oil tankers) is relatively heavy, there is potential for atmospheric deposition of acids
721 (SO_x and NO_x) to play a role in the carbonate system variability (Doney et al. 2007,
722 Hunter et al. 2011). To try to understand this control, we deployed air samplers at our
723 study site for eight 2-week periods. The levels of atmospheric NO_2 and SO_2 did not vary
724 widely over the time period. NO_2 was ranged 5.45 to 6.99 ppb (6.13 ± 0.63 ppb), and SO_2
725 ranged 1.15 to 1.18 ppb (1.43 ± 0.35 ppb) over the sampling dates (J. D. Felix, personal



726 communications). There was no apparent correlation between these values and the pH or
727 $p\text{CO}_2$ levels over the 2-week sampling periods.

728 Co-locating our pH and $p\text{CO}_2$ sensors with other coastal environmental
729 monitoring sensors allowed insight into correlated environmental parameters and
730 potential driving forces of carbonate chemistry on diel and seasonal time scales. The
731 results of this study provide strong support for the continued implementation of carbonate
732 chemistry monitoring in conjunction with preexisting coastal environmental monitoring
733 infrastructure. Our understanding of any estuarine system could benefit from long-term
734 effective deployments of these monitoring tools. Strategically locating carbonate
735 chemistry sensors at estuarine sites that are subject to local OA drivers or support large
736 biodiversity or commercially important species may be the most crucial in guiding future
737 mitigation and adaptation strategies for natural systems and aquaculture facilities (Chan
738 et al., 2013; Strong et al., 2014).

739 *4.2 Carbonate chemistry as a component of overall estuarine system variability*

740 Estuaries and coastal areas are dynamic systems with human influence, riverine
741 influence, and influence from an array of biogeochemical processes, resulting in highly
742 variable chemical and environmental conditions. To better understand overall system
743 variability over different time scales, we used a linear discriminant (LD) analysis, a
744 multivariate statistic that allows dimensional reduction, to determine the linear
745 combination of environmental parameters (individual parameters reduced into linear
746 discriminants, LDs) that allow the best differentiation between day and night as well as
747 between seasons. This used the same suite of environmental data and data sources as
748 Sect. 4.1.2.



749 All variables were centered and scaled to allow direct comparison of their
750 contribution to the system variability. The magnitude (absolute value) of coefficients of
751 the LDs (Table 7) represents the relative importance of each individual environmental
752 parameter in the best discrimination between day and night and between seasons, i.e., the
753 greater the absolute value of the coefficient, the more information the associated
754 parameter can provide about whether the sample came from day or night (or winter,
755 spring, or summer). Only one LD could be created for the diel variability (since there are
756 only two classes to discriminate between – day and night). Two LDs could be created for
757 the seasonal variability (since there were three classes to discriminate between – fall was
758 omitted because of the lack of tidal data), but only the coefficients for LD1 are reported
759 (Table 7) given that LD1 captured 95.64% of the seasonal variability.

760 Table 7. Coefficients of linear discriminants (LD) from discriminant function analysis
761 (DFA) using continuous sensor data and other environmental parameters. Results for
762 discriminants for both diel and seasonal variability shown. All variables were centered
763 and scaled. For the seasonal analysis, only LD1 is given since it was captured 95.64% of
764 the variability (for the diel analysis, there is only one). Given that many of the water
765 quality parameters were measured in bottom waters and our sensors were measuring
766 surface waters, only those observations without significant stratification in the water
767 column (a salinity difference of less than 3 between surface and bottom) were included in
768 these analyses.

	Diel	Seasonal
	LD1	LD1
Temperature (°C)	0.5406	-3.5279
Salinity	0.1473	0.0432
pCO ₂ (µatm)	-0.1612	-0.2928
pH	0.0593	0.0991
Tide Level (m)	0.0968	-0.2389
Wind speed (ms ⁻¹)	-0.0009	0.0504
Total PAR	-2.2878	-0.0676
DO (mg L ⁻¹)	-0.0839	0.0859
Turbidity	-0.0561	0.1455
Fluor. Chlorophyll	0.1397	-0.4040

769

770 As would be expected, we found that PAR provided the most differentiation
771 between daytime and nighttime conditions (based on the largest coefficient associated



772 with Diel LD1, Table 7). Temperature was the second most important factor in
773 differentiating between day and night; this corresponds to the diel variability that we
774 detected where both summer and fall had clear separation of mean temperature between
775 day and night, with nighttime temperatures being 0.3 and 1.0 higher, respectively (Table
776 3). The next most important parameter in differentiating between day and night in this
777 system was $p\text{CO}_2$, providing more evidence for differentiation between day and night
778 than other parameters that would be expected to vary on a diel timescale (e.g. chlorophyll
779 and DO). As for system variability that allowed differentiation between the four seasons,
780 the most important parameter in system variability was temperature (Table 7, Seasonal
781 LD1), as would be expected with the clear seasonal temperature fluctuations (Fig. 2E).
782 The second most important parameter in contributing to seasonal variability was
783 chlorophyll, likely indicating clear seasonal blooms. The third most important parameter
784 for seasonal differentiation was $p\text{CO}_2$; therefore $p\text{CO}_2$ variability seems to be more
785 closely tied to seasons than variability in tide level, DO, or the array of other parameters
786 (Table 7).

787 The contribution of pH to discriminating along diel or seasonal scales was less
788 than $p\text{CO}_2$ despite the same seasonal differences that were identified by ANOVA (Table
789 3) and more seasons with significant diel differences (Table 2). However, pH still seemed
790 to be relatively important on seasonal scales, having clearer contribution to seasonal
791 system variability than several other parameters including DO and salinity.

792 We can conclude that carbonate chemistry parameters are among the most
793 important of variants on both daily and seasonal time scales in this coastal setting.
794 Compared to six other estuaries around the United States with similar sensor deployments



795 for carbonate chemistry characterization, our study site has a relatively small range of pH
796 and $p\text{CO}_2$ on both diel and seasonal scales (N. Rosenau, personal communication). While
797 we do not have the same suite of environmental data for these other systems, this
798 suggests that the relative amount of system variability contributed by carbonate chemistry
799 may be even greater in other estuarine systems. The relatively small fluctuations in pH
800 and $p\text{CO}_2$ that are seen on a daily scale at the Aransas Ship Channel is likely due to the
801 subtropical setting with little ocean upwelling influence and the lower estuary position of
802 our monitoring (further removed from the already small freshwater influence), but it may
803 also be tied to the system's relatively high buffer capacity. Just as the extent of hypoxia-
804 induced acidification was relatively low in Corpus Christi Bay compared to other systems
805 because of the bay's high buffer capacity (McCutcheon et al., 2019), the extent of pH
806 fluctuation on a daily scale from biological activity would also be modulated by the
807 intrinsic buffer capacity, which is likely also high in this system due to high alkalinity in
808 the freshwater endmembers (Yao et al., 2020).

809 *4.3 Comparing continuous monitoring and discrete sampling*

810 *4.3.1 Representative sampling in a temporally variable environment*

811 Discrete water sample collection and analysis is the most common method that
812 has been employed to attempt to understand the carbonate system of estuaries. However,
813 it is difficult to know if these samples are representative of the spatial and temporal
814 variability in carbonate system parameters. While this time-series study cannot conclude
815 whether our broader sampling efforts in the MAE are representative of the spatial
816 variability in the estuary, it can investigate how representative our bimonthly to monthly



817 sampling is of the more high-frequency temporal variability that the Aransas Ship
818 Channel experiences.

819 One-way ANOVAs were conducted to compare between monitoring methods
820 (separate one-way ANOVAs within each season because of the significant interaction
821 between these factors in an initial two-way ANOVA). There were three levels of
822 monitoring method included in the comparison of means: continuous monitoring, discrete
823 monitoring during only the continuous monitoring period, and discrete monitoring over
824 the entire period (C, D_c, and D, respectively, in Table 3). To interpret the results, a
825 difference in means between the continuous monitoring and discrete monitoring datasets
826 would only indicate that the 10-month period of continuous monitoring was not
827 representative of the 5+ year period that discrete samples have been collected, but a
828 difference in means between the continuous data and discrete sample data collected
829 during the continuous monitoring period represents discrepancies between types of
830 monitoring.

831 There were several instances where seasonal parameter means significantly
832 differed between the 10-month continuous monitoring period and the 5+ year discrete
833 sampling period (Table 3, $C \neq D$ or $D_c \neq D$) including temperature in the summer and fall,
834 salinity in the spring, pH in the summer and fall, and $p\text{CO}_2$ in winter, spring, and summer.
835 While clear seasonal variability was demonstrated for most parameters (using both
836 continuous and discrete data for the entire period), these differences between the 10-
837 month continuous monitoring period and our 5+ year monitoring period illustrate that
838 there is also interannual variability in the system. Therefore, short periods of monitoring
839 are unable to fully capture current baseline conditions.



840 During the continuous monitoring period (2016-2017), we found no significant
841 difference between sampling methods in the seasonal mean temperature, salinity, or
842 *p*CO₂. The two sampling methods also resulted in the same mean pH for all seasons
843 except for summer, when the sensor data recorded a higher mean pH than discrete
844 samples (Tables 1 and 3). During this case, we can conclude that discrete monitoring did
845 not accurately represent the system variability that was able to be captured by the sensor
846 monitoring. However, given that most seasons did not show differences in pH or *p*CO₂
847 between sampling methods, the descriptive statistics associated with the discrete
848 monitoring did a fair job of representing system means. This is evidence that long-term
849 discrete monitoring efforts, which are much more widespread in estuarine systems than
850 sensor deployments, can be generally representative of the system despite known
851 temporal variability on shorter time scales.

852 Understanding the relationships of pH and *p*CO₂ with temperature and salinity is
853 important in a system. Both the continuous and discrete sampling types indicate that pH
854 has a significant negative relationship with both temperature and salinity and *p*CO₂ has a
855 significant positive relationship with both temperature and salinity (Fig. 7). Based on
856 the results of an Analysis of Covariance (ANCOVA), the relationship (slope) of pH with
857 both temperature and salinity and of *p*CO₂ with salinity were not significantly different
858 between types of monitoring (considering the sensor deployment period only), supporting
859 the effectiveness of long-term discrete monitoring programs when sensors are unable to
860 be deployed. However, ANCOVA did reveal the relationship of *p*CO₂ with temperature is
861 significantly different (method:temp *p*=0.0062) between monitoring methods.



862 While *in situ* monitoring is usually lacking good spatial coverage, it is effective in
863 capturing temporal resolution and presumably providing better estimates of average CO₂
864 flux at a given location versus periodic sampling. Previous studies have pointed out that
865 discrete sampling methods, which generally involve only daytime sampling, do not
866 adequately capture the diel variability in the carbonate system and may therefore lead to
867 underestimation of CO₂ fluxes. However, we found no significant difference (within any
868 season) between CO₂ flux values calculated with sensor data versus discrete samples
869 (Table 3). Calculated CO₂ fluxes also did not significantly differ between day and night
870 during any season, despite some differences in *p*CO₂ (Table 2), likely due to the large
871 error associated with the calculation of CO₂ flux (Table 1, Fig. 5) which will be further
872 discussed below. Therefore, the expected underestimation of CO₂ flux based on diel
873 variability of *p*CO₂ was not encountered at our study site, validating the use of discrete
874 samples for quantification of CO₂ fluxes (until methods with less associated error are
875 available). Even given less error in calculated flux, estimated fluxes would likely not
876 differ between methods on an annual scale (as *p*CO₂ did not), but CO₂ fluxes may differ
877 on a seasonal scale since the differences between daytime and nighttime *p*CO₂ were not
878 consistent across seasons (Table 2).

879 There are many factors contributing to error associated with CO₂ flux. There is
880 still large error associated with estimates of estuarine CO₂ flux because turbulent mixing
881 is difficult to model and turbulence is the main control on CO₂ gas transfer velocity, *k*, in
882 shallow water environments. Thus, our wind speed parameterization of *k* is imperfect and
883 likely the greatest source of error. Other notable sources of error include the data
884 treatment. For example, we chose to seasonally weight the individual calculated flux



885 values in the calculation of annual flux to account for differences in sampling frequency
886 between seasons. From continuous data, the weighted average flux was $0.2 \text{ mmol m}^{-2} \text{ d}^{-1}$,
887 although choosing not to seasonally weight and simply look at the arithmetic mean of
888 fluxes calculated directly from sampling dates would have resulted in an annual CO_2 flux
889 of $-0.7 \text{ mmol m}^{-2} \text{ d}^{-1}$ for the same period. Similarly, the weighted average flux from all
890 5+ years of discrete data was $-0.9 \text{ mmol m}^{-2} \text{ d}^{-1}$, but the arithmetic mean of fluxes would
891 have resulted in an annual CO_2 flux of $0.2 \text{ mmol m}^{-2} \text{ d}^{-1}$ for the same period. Another
892 source of error that could be associated with the calculation of flux from the discrete data
893 is the way in which wind speed data are aggregated to be used in the windspeed
894 parameterization. We decided to use daily averages of the windspeed for calculations.
895 Using the windspeed measured for the closest time to our sampling time or the monthly
896 averaged wind speed may have resulted in very different flux values.

897 *4.3.2 Direct agreement of measurement methods and quantified uncertainties associated*
898 *with parameters*

899 Direct comparisons were made between measurements from sensors and
900 laboratory-analyzed bottle samples—including both quality control (QC) samples taken
901 from the cooler that housed the sensors at the time when these sensors took recorded
902 readings and long-term monitoring samples taken from the ship channel near the sensors
903 (within 100 m) that occurred at various times and were compared to sensor measurements
904 of the closest full hour (Table 8). The mean difference between the SeaFET pH
905 measurements and the QC samples (continuous – discrete) prior to sensor data correction
906 was 0.05 ± 0.08 (Table 8, which would reduce to 0.00 ± 0.08 following the correction).
907 The mean difference between the SAMICO₂ pCO₂ measurements and the QC samples



908 (continuous – discrete) was -18 ± 44 (Table 8) when discrete sample $p\text{CO}_2$ was calculated
 909 using Millero (2010) constants. We used several different constants to calculate $p\text{CO}_2$ to
 910 check this offset; all were similar in mean and standard deviation, but the offset could be
 911 slightly reduced using Millero (2002) constants.

912 Table 8. Comparison of discrete and continuous monitoring. The difference between
 913 sampling methods is reported in two different ways: the difference between sensor
 914 measurements and laboratory measurement of quality control (QC) bottle samples taken
 915 directly from the cooler (here the pH difference is prior to the sensor pH correction of
 916 $+0.05$), and the difference between sensor measurements and laboratory measurement of
 917 bottle samples taken from a nearby station for our 5+ year monitoring (here the pH
 918 difference is after the sensor pH correction of $+0.05$, see methods for details). For all
 919 calculated parameters, dissociation constants from Millero 2010 were used. Error—
 920 analytical error for directly measured parameters and propagated error for calculated
 921 parameters (mean \pm standard deviation, calculated in the seacarb package in R—
 922 associated with carbonate system variables is also reported.

	Difference between sampling methods (mean difference \pm standard deviation of the difference)		Error (Analytical or Propagated)	
	Sensor – QC cooler samples (prior to sensor pH correction, n=12)	Sensor – discrete samples (after pH sensor correction, n=13)	Discrete Sampling (n = 104)	Continuous Monitoring (n = 6088)
Temperature (°C)			0.1	0.1
Salinity	-0.16 ± 1.44	0.50 ± 1.69	0.01	0.1
pH	-0.05 ± 0.08	0.01 ± 0.12	0.0004	0.05
$p\text{CO}_2$ (μatm)	-18 ± 44	25 ± 63	7 ± 2	1.0
DIC ($\mu\text{mol kg}^{-1}$)			± 2.5	327.4 ± 63.2
TA ($\mu\text{mol kg}^{-1}$)			7.4 ± 0.9	400.7 ± 81.0
Ω_{Ar}			0.19 ± 0.03	1.08 ± 0.31

923
 924 Given that the analytical accuracy of the SeaFET instrument is 0.05 pH units, the
 925 average offset between sensor and laboratory values of quality control samples
 926 demonstrates fair agreement (Table 8). Given that calculated uncertainty associated with
 927 calculated discrete $p\text{CO}_2$ was 7 ± 2 , we did not see great agreement between SAMICO2
 928 $p\text{CO}_2$ and laboratory-calculated $p\text{CO}_2$ for quality control samples (mean difference of -18
 929 ± 44 , Table 8). Mean offsets and their associated standard deviations were larger when



930 comparing sensor data to samples taken during our long-term discrete monitoring effort.
931 This is not surprising given that the discrete sample collection did not occur at the exact
932 time of the sensor measurement or the exact location of the cooler pump inlet. Greater
933 sensor-laboratory agreement has been achieved for open ocean settings, but this larger
934 standard deviation is likely a result of the temporal variability in the more complex
935 estuarine environment where these instruments have been much less widely deployed to
936 date.

937 Propagation of error associated with computed carbonate system parameters was
938 done using the *seacarb* package in R (Gattuso et al., 2018); the error propagation includes
939 error associated with the measurements of the input pair ($1 \mu\text{atm}$ for $p\text{CO}_2$ from
940 SAMICO2 and 0.05 for pH from SeaFET; 0.0004 for laboratory spectrophotometric pH
941 and $2.5 \mu\text{mol kg}^{-1}$ for laboratory DIC), error associated with *in-situ* temperature ($0.1 \text{ }^\circ\text{C}$)
942 and salinity (0.1 for sensor-measured and 0.01 for laboratory-measured), and error
943 associated with total boron the key dissociation constants (standard recommended error
944 used) (Table 8). While the error associated with calculated parameters from discrete
945 bottle samples was relatively small and likely a result of uncertainties in constants (Orr et
946 al., 2018), we note that the error associated with calculated dissolved inorganic carbon
947 (DIC), total alkalinity (TA), and saturation state of aragonite (Ω_{Ar}), which are other
948 frequently addressed carbonate system parameters, was large when calculated with sensor
949 data. This large error is likely a result of both the relatively low analytical precision
950 associated with the pH sensor and the poor mathematical combination of variables for
951 speciation calculations. Hence, we limited the discussion to pH (which was directly
952 measured for both continuous monitoring and laboratory analysis of discrete samples)



953 and $p\text{CO}_2$ (which was directly measured for continuous monitoring and had relatively
954 low error when calculated with discrete sample DIC and pH, Table 8) and omitted any
955 discussion of the parameters with high propagation error. The high error suggests that it
956 may be important to develop and broadly use autonomous sensors that can measure
957 carbonate system parameters that allow for lower propagated error to have a full picture
958 of estuarine carbonate chemistry on high-frequency time scales.

959 **5. Conclusions**

960 We monitored carbonate chemistry parameters (pH and $p\text{CO}_2$) using both sensor
961 deployments (10 months) and discrete sample collection (5+ years) at the Aransas Ship
962 Channel, TX, to characterize temporal variability and investigate controlling factors.
963 Both sampling methods demonstrated significant seasonal variability at the location, with
964 highest pH (lowest $p\text{CO}_2$) in the winter and lowest pH (highest $p\text{CO}_2$) in the summer.
965 Significant diel variability was also evident from sensor data, though diel fluctuations
966 were smaller than many other areas previously studied. Carbonate chemistry parameters
967 were among the most important environmental parameters to distinguish between both
968 diel and seasonal environmental conditions.

969 The difference between daytime and nighttime values of carbonate system
970 parameters varied between seasons, occasionally reversing the expected diel variability
971 due to biological processes. It was evident that biological activity is not the strongest
972 controlling factor of diel variability at this location, likely surpassed by tidal control
973 despite the small tidal range in the northwestern GOM. Controls on the system also
974 differed over different time scales, with temperature becoming a less important control
975 over shorter time scales.



976 Tides exerted significant control on the carbonate system, and low tide allowed
977 more biological control of the system. Higher mean $p\text{CO}_2$ was reported for low tide
978 versus high tide across all seasons. pH was higher at high tide during winter and summer
979 but deviated from the expected pattern during spring with lower pH during high tide. The
980 results suggest that the controlling factors of the carbonate system may not be exerted
981 equally on both pH and $p\text{CO}_2$ on diel timescales, causing separation of their diel or tidal
982 relationships during certain seasons. The detailed investigation of controlling factors
983 provides strong support for the implementation of carbonate chemistry monitoring in
984 conjunction with preexisting coastal environmental monitoring infrastructure, which has
985 had little application in estuarine environments thus far.

986 Despite known temporal variability on shorter timescales, discrete sampling was
987 generally representative of the average carbonate system on a seasonal and annual basis
988 based on comparison with our sensor data. Additionally, there was no difference in CO_2
989 flux between sampling types supporting the validity of discrete sample collection for
990 carbonate system characterization.

991 **Data availability**

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



998 **Author Contribution**

999 MM and XH defined the scope of this work. XH received funding for all components of
1000 the work. MM, HY, and CJS performed field sampling and laboratory analysis of
1001 samples. MM prepared the initial manuscript and all co-authors contributed to revisions.

1002 **Competing interests**

1003 The authors declare that they have no conflict of interest.

1004

1005 **Acknowledgements**

1006 Funding for autonomous sensors and sensor deployment was provided by the
1007 United States Environmental Protection Agency's National Estuary Program via the
1008 Coastal Bend Bays and Estuaries Program Contract No. 1605 Thanks to Rae Mooney
1009 from Coastal Bend Bays and Estuaries Program for assistance in the initial sensor setup.
1010 Funding for discrete sampling as well MM's dissertation research has been supported by
1011 both NOAA National Center for Coastal Ocean Science (Contract No.
1012 NA15NOS4780185) and NSF Chemical Oceanography Program (OCE-1654232). We
1013 also appreciate the support from the Mission-Aransas National Estuarine Research
1014 Reserve in allowing us the boat-of-opportunity for our ongoing discrete sample
1015 collections and the University of Texas Marine Science Institute for allowing us access to
1016 their research pier for the sensor deployment. A special thanks to Hongjie Wang, Lisette
1017 Alcocer, Allen Dees, and Karen Alvarado for assistance with field work.

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