



Decadal-Scale Acidification Trends in Adjacent North Carolina Estuaries: Competing Role of Anthropogenic CO₂ and Riverine Alkalinity Loads

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Decadal-scale pH trends for the open ocean are largely monotonic and controlled by anthropogenic CO₂ invasion. In estuaries, though, such long-term pH trends are often obscured by a variety of other factors, including changes in net metabolism, temperature, estuarine mixing, and riverine hydrogeochemistry. In this study, we mine an extensive biogeochemical database in two North Carolina estuaries, the Neuse River estuary (NeuseRE) and New River estuary (NewRE), in an effort to deconvolute decadal-scale trends in pH and associated processes. By applying a Generalized Additive Mixed Model (GAMM), we show that temporal changes in NewRE pH were insignificant, while pH decreased significantly throughout much of the NeuseRE. In both estuaries, variations in pH were accompanied by increasing river discharge, and were independent of rising temperature. Decreases in bottom-water pH in the NeuseRE coincided with elevated primary production in surface waters, highlighting the importance of eutrophication on long-term acidification trends. Next, we used a simple mixing model to illustrate the impact of changing river discharge on estuarine carbonate chemistry. We found that increased riverine alkalinity loads to the NewRE likely buffered the impact of CO₂-intrusion-induced acidification. In the NeuseRE, however, elevated dissolved inorganic carbon loads further decreased the buffering capacity, exacerbating the effects of CO₂-intrusion-driven acidification. Taken together, the findings of this study show that future trajectories in estuarine pH will be shaped by complex interactions among global-scale changes in climate, regional-scale changes in precipitation patterns, and local-scale changes in estuarine biogeochemistry.

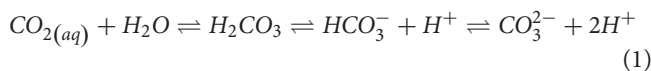
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INTRODUCTION

Anthropogenic activities have caused dramatic increases in atmospheric CO₂ levels, from a pre-industrial level of <300 ppm to the current value of 407 ppm (Keeling and Keeling Charles, 2017). Much of the open ocean is under-saturated in CO₂ with respect to the atmosphere, causing a net uptake of CO₂, mitigating approximately 30% of anthropogenic emissions (Le Quéré et al., 2016).

This absorption of atmospheric CO₂ is leading to the increase in aqueous CO₂ concentration {[CO_{2(aq)}]}, which affects the speciation of inorganic carbon in the ocean, shifting the following reaction (Eq. 1) toward the right, favoring the formation of H⁺ and lowering the pH. Accordingly, anthropogenic CO₂ emissions have caused ocean pH to fall at a rate of approximately 0.001–0.0025 yr⁻¹ (Doney et al., 2009; Bates et al., 2014; Takahashi et al., 2014).

Excess H⁺ produced by the dissolution of CO_{2(aq)} pairs with CO₃²⁻ to form HCO₃⁻, resulting in a net decrease in [CO₃²⁻] and saturation states (Ω) of various carbonate minerals, adding stress to marine calcifying organisms. This process is collectively known as ocean acidification, and may directly damage calcifying organisms, further decrease the fitness of commercially valuable groups, by directly damaging shells or compromising early development and survival.



Over very long time scales (i.e., >100,000 years), it is expected that increased continental weathering and subsequent delivery of alkalinity via rivers will partially mitigate the coastal and open ocean acidification signal (Doney and Schimel, 2007; Müller et al., 2016). Over shorter time scales, though, riverine inputs of freshwater and related solutes may cause varied effects on the carbonate system in estuaries, resulting in contrasting pH trajectories (Aufdenkampe et al., 2011; Hu and Cai, 2013; Hu et al., 2015; Müller et al., 2016). For example, changing local water budgets resulted in a range of temporal trajectories in riverine alkalinity loading for a set of Northwest Gulf of Mexico estuaries, where estuaries experiencing decreasing alkalinity loads also became acidified (Hu et al., 2015). Similarly, long-term alkalinity increases in the Baltic sea have partially or wholly counteracted anthropogenic CO₂-induced acidification in that system (Müller et al., 2016). However, the impact of short-term variations in river discharge on the estuarine carbonate system is less clear (Mote et al., 2008; Philips et al., 2011; Couldrey et al., 2016; Baumann and Smith, 2017; Siam and Eltahir, 2017).

A variety of biogeochemical processes will interact with these watershed-scale drivers, either enhancing or counteracting acidification trends in estuaries. For example, nutrient-enhanced primary production has been linked with elevated pH in well-mixed mesocosm experiments (Nixon et al., 2015), but net production and respiration are decoupled in stratified estuaries, where the latter process decreases bottom water pH (Feely et al., 2010; Cai et al., 2011; Waldbusser and Salisbury, 2014). Still, net ecosystem metabolism in productive coastal systems like seagrass beds has been suggested to lessen long-term coastal ocean acidification (Unsworth et al., 2012), but increase the short-term extremes in pH (Cyronak et al., 2018; Pacella et al., 2018). H₂S oxidation and subsequent H₂SO₄ additions may exacerbate acidification near the pycnocline of stratified systems (Cai et al., 2017). Anaerobic production of CO₂ and alkalinity in wetlands can act to simultaneously alkalize and acidify pore-water, leading to variable impacts on estuarine pH (Hu and Cai, 2011; Wang Z.A. et al., 2016). At the same time, estuarine waters are mixing with oceanic water

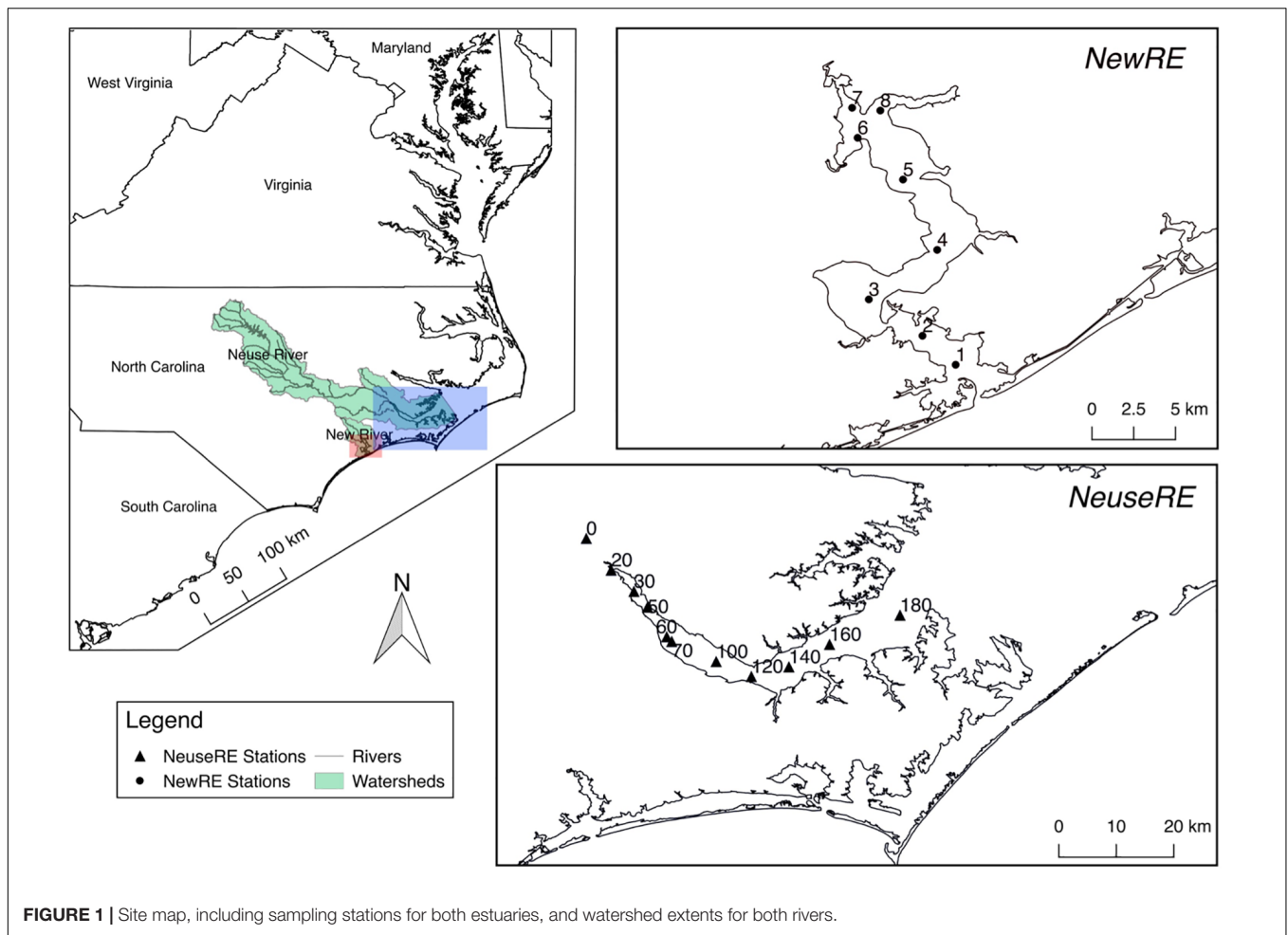
that is also becoming enriched in CO₂ due to anthropogenic CO₂ emissions.

The above natural and anthropogenic processes will combine non-linearly to affect pH, to an extent that may be exacerbated by the typically poor buffering capacity of estuarine water (Feely et al., 2010; Cai et al., 2011; Hu and Cai, 2013; Jeffrey et al., 2016; Van Dam et al., 2018a). Future trajectories in estuarine acidification will depend on the interactions among a variety of complex drivers, including the invasion of anthropogenic CO₂, changes in freshwater delivery, and changes in net ecosystem metabolism. However, we do not yet know how these factors will interact over decadal time scales, or across different estuaries. The estuaries of eastern North Carolina, United States, offer a unique opportunity to compare the impacts of these factors on decadal-scale acidification trends. Both New and Neuse River estuaries, NC, experience the same climate, and have seen increases in river discharge that are of a similar magnitude over the past 10 years, but contrasting watersheds properties cause dissolved inorganic carbon (DIC) and total alkalinity (TA) loading to differ (Van Dam et al., 2018a). In this study, we investigate decadal trends in pH and associated parameters in these two adjacent estuaries, then use a simple modeling approach to examine the sensitivity of estuarine acidification to above three factors.

MATERIALS AND METHODS

Site Description

The Neuse River Estuary (NeuseRE) and New River Estuary (NewRE) are microtidal (tidal amplitude < 1 m), lagoonal systems located in the eastern North Carolina coastal plain (**Figure 1**), both of which have relatively long fresh water residence times [average 58 (±34) and 46 (±27) days, respectively (Van Dam et al., 2018a)]. Both estuaries are shallow, with average water depths of 1.8 and 2.7 m in the NewRE and NeuseRE, respectively. While the NewRE is connected directly with the coastal ocean through the New River inlet, a string of barrier islands (Outer Banks, NC) with narrow tidal inlets restricts water exchange between the NewRE and the ocean. The long residence time and limited tidal exchange has interacted with a history of anthropogenic nutrient enrichment to drive both estuaries toward eutrophy. Historically, average primary productivity (PPR) has been approximately 200–500 g C m⁻² yr⁻¹ in the NeuseRE (Boyer et al., 1993; Mallin et al., 1993) and <250 gC m⁻² yr⁻¹ in the NewRE (Mallin et al., 2005). A key difference between these two microtidal, lagoonal estuaries is their size; the surface area of the NeuseRE (352 km²) is approximately 5 times that of the NewRE (79 km²), while the Neuse River watershed (15,700 km²) is over 15 times larger than that of the New River (1,024 km²). In contrast with the heavily agricultural Neuse River watershed, much of the New River watershed contains impervious surfaces, causing river discharge in this system to be relatively ‘flashy’ in response to storm events (Hall et al., 2012; Peierls et al., 2012). Submarine groundwater discharge is a very small contributor to total freshwater loads in both



NewRE (Crosswell et al., 2017) and NeuseRE (Fear et al., 2007; Null et al., 2011).

Long-Term Data Collection

Long-term data were acquired from two monitoring programs established in each estuary from 2005 to 2017. Associated with the Defense Coastal Estuarine Research Program (DCERP), monthly sample collection began in the NewRE in the Fall of 2007. The goal of this project was to assess the impacts of land use in the watershed on water quality in the NewRE. Likewise, bi-weekly to monthly sample collection occurred in the NeuseRE beginning in 1999, as a part of the Neuse River Estuary Modeling and Monitoring Project (ModMon). ModMon is a collaborative effort of both universities and the state of North Carolina, aiming at the collection of water-quality compliance data. For DCERP and ModMon programs, surface (~0.2 m) and bottom-water (~0.5 m above bottom) samples were collected at each station using a diaphragm pump, and were immediately stored in a cooler. For each surface sample, an estimate of phytoplankton PPR was made using the ^{14}C uptake method, and *in-vitro* Chlorophyll-*a* fluorescence (Chl-*a* IV) was determined using acetone extraction (Paerl et al., 1998; Hall et al., 2012). The ^{14}C uptake method generates PPR estimates that lie somewhere between gross and

net PPR. This is because a fraction of the ^{14}C incorporated by phytoplankton is respired during the incubation period, and lost as CO_2 before the final scintillation count. However, this method has remained stable over time in both ModMon and DCERP programs, allowing us to reliably assess trends in PPR over time.

At each DCERP and ModMon station, vertical profiles of pH, dissolved oxygen (DO), salinity (Sal), and temperature (Temp) were collected at a vertical resolution of 0.5 m (YSI 6600 multi-parameter sonde, Yellow Springs Inc, Yellow Springs, OH). The pH sensor (YSI model 6561) was calibrated against NIST-traceable NBS buffers, and has a manufacturer's suggested accuracy of ± 0.2 units, which is far less than the typical spatial variability in pH (Van Dam et al., 2018a). Once water samples were returned from the field, a small aliquot was immediately stored, unpreserved, in a 20 mL scintillation vial with no headspace, and was analyzed for DIC within 24 h using a Shimadzu TOC-5000A in inorganic carbon mode. Because DIC in these unpreserved water samples is often lower than when preserved with HgCl_2 , or by filtration, we applied a correction factor (Crosswell et al., 2012) to DIC values obtained from the long-term monitoring dataset ($\text{DIC} = 7.18 + 1.16 \times \text{DIC}_{\text{unpreserved}}$). We then calculated TA from this corrected DIC value, pH, Temp, and Sal using CO2SYS

(Lewis and Wallace, 1998), with the carbonic acid dissociation constants of Millero (2010), and the NBS scale for pH.

Statistical Approach

We calculated trends in pH/PPR/Chl-a using a Generalized Additive Mixed Modeling (GAMM, Eq. 2), a method which has been previously applied to calculate multi-decadal $f\text{CO}_2$ trend in various coastal environments (Wang H. et al., 2016, 2017; Reimer et al., 2017). Briefly, a penalized spline was first used to fit the seasonal cycle, and a harmonic function was adopted to fit daily cycles, then linear regressions were used to model the variability resulting from salinity and DO change. Sampling date was also included in GAMM as a linear effect, and its coefficient represents the pH/PPR/Chl-a long-term change. The method also weighted the observations using explicit models for heteroscedasticity to account for an observed unequal variance in salinity.

$$\text{pH/PPR/Chl} - a = f(\text{seasonal cycle}) + f(\text{daily cycle}) + f(\text{DO, Sal}) + f(\text{long-term trend}) + \text{error} \quad (2)$$

Note, the “best” model was selected based on smallest AICc values (Supplementary Table S1 and Supplementary Figure S1). Trends were assessed for the time period 2005–2017 in the NeuseRE, and for 2007–2017 in the NewRE. Salinity and temperature trends were calculated after removing the seasonal cycle only. Pre-2005 data were not used for the NeuseRE because: (1) residuals were not normally distributed before 2005, (2) different pH probe was used before 2005, and data were more scattered, (3) surveys were conducted in early morning, causing pH measurements to be biased toward lower values (based on typical diel variations in these estuaries), and (4) data do not exist for the NewRE before 2007.

Box Model

In this study, we used a simple box model approach to address the relative importance of the following three factors on long term estuarine acidification trends: (1) continuous anthropogenic CO_2 invasion, (2) varying PPR, and (3) changing river discharge (and TA/DIC loads). This model is intended to represent average conditions over yearly to decadal-time scales, rather than shorter-term variations in pH, which can be very difficult to predict. As future changes in river discharge are very uncertain, we estimated a range of river discharge (Q_R) by estimating the average discharge (Q_R) from Supplementary Table S2 over time, extrapolating forward 70 years, and backward 20 years. This range of Q_R (New River: $1.3\text{--}8.9 \text{ m}^3 \text{ s}^{-1}$, Neuse River: $82\text{--}221 \text{ m}^3 \text{ s}^{-1}$) is intended to include current conditions, and what could be expected for flood and drought years in the next century. The riverine end-member concentrations of DIC (DIC_R) was set based on the empirical relationships with Q_R determined in Van Dam et al., 2018b (Supplementary Table S2). We set the riverine TA (TA_R) value based on the linear relationship between measured DIC and calculated TA for the station closest to the river mouth, when salinity was <5 for the NewRE, or <2 for the NeuseRE. Those equations are also shown in Supplementary Table S2. The r^2 for these

relationships were 0.96 for the NeuseRE, and 0.97 for the NewRE. We assumed a constant oceanic TA end-member (TA_O) of $2500 \mu\text{mol kg}^{-1}$, and increasing oceanic DIC (DIC_O) as a function of increasing atmospheric CO_2 (i.e., $\text{DIC}_O \sim p\text{CO}_2$, TA). These end-member DIC and TA values were mixed within the estuary, allowing us to calculate pH under different atmospheric CO_2 levels.

After setting the end-member chemistry, we assumed that the main factors affecting DIC concentration in surface and bottom water was net ecosystem metabolism. The NewRE and NeuseRE are both stratified estuaries, where net ecosystem production and respiration are vertically decoupled, meaning that some fraction of net production in the surface is exported to bottom waters where it is respired. Hence, we used average net ecosystem productivity ($\overline{\text{NEP}}$, $\text{mol m}^{-2} \text{ yr}^{-1}$) from Herrmann et al. (2015), and assumed that a certain fraction of net production was respired in bottom waters (R). Note that the sign of $\overline{\text{NEP}}$ was reversed from that given in Herrmann et al. (2015), such that positive values indicate net heterotrophy. Given the paucity of data regarding bottom-water respiration in these systems, we set R at 40% ($R = 0.4 \times \overline{\text{NEP}}$). Variations in R do not affect the general trend of modeled pH with variations in Q_R or $p\text{CO}_2$, but does cause moderate changes in the absolute value of modeled pH. As an example of these variations, modeled pH results are shown for R values of 0.25 (a), 0.5 (b), and 0.75 (c) in Supplementary Figure S4.

Areal $\overline{\text{NEP}}$ was converted into a concentration change by the following equation where z_{ave} is the average water depth, τ_{FW} is the residence time (years), and F_{mix} is the mixing fraction, given by the ratio of $\text{Sal}_E/\text{Sal}_O$ (i.e., $F_{\text{mix}} = 10/35 = 0.29$ for oligohaline region). Multiplying by F_{mix} scales the DIC loss due to air-water exchange from the whole estuary down to the region of interest.

$$\text{NEP}_E (\mu\text{M}) = 10^3 \times \left(\frac{\overline{\text{NEP}} \times \tau_{\text{FW}} \times F_{\text{Mix}}}{z_{\text{ave}}} \right) \quad (3)$$

With DIC consumption/production terms defined, surface and bottom water DIC in the estuary (DIC_S and DIC_B respectively) was calculated assuming conservative mixing, with the addition of net metabolism and air-sea exchange:

$$\text{DIC}_S = (\text{DIC}_O \times F_{\text{Mix}}) + (\text{DIC}_R \times (1 - F_{\text{Mix}})) - \text{NEP}_E \quad (4)$$

$$\text{DIC} = (\text{DIC}_O \times F_{\text{Mix}}) + \text{DIC}_R \times (1 - F_{\text{Mix}}) + (R) \quad (5)$$

Total alkalinity in the estuary (TA_E) was determined by conservative mixing of river and ocean end-members. We also accounted for the small amount of TA production (consumption) that occurs during the production (remineralization) of Redfield organic matter, by assuming TA changes with DIC at a ratio of $-17/106$.

From these modeled TA and DIC values, estuarine pH (pH_{est}) was calculated in CO2SYS using the carbonic acid dissociation constants of Millero (2010), at a constant temperature of 20°C , very close to the median temperature in the NeuseRE (20.5°C)

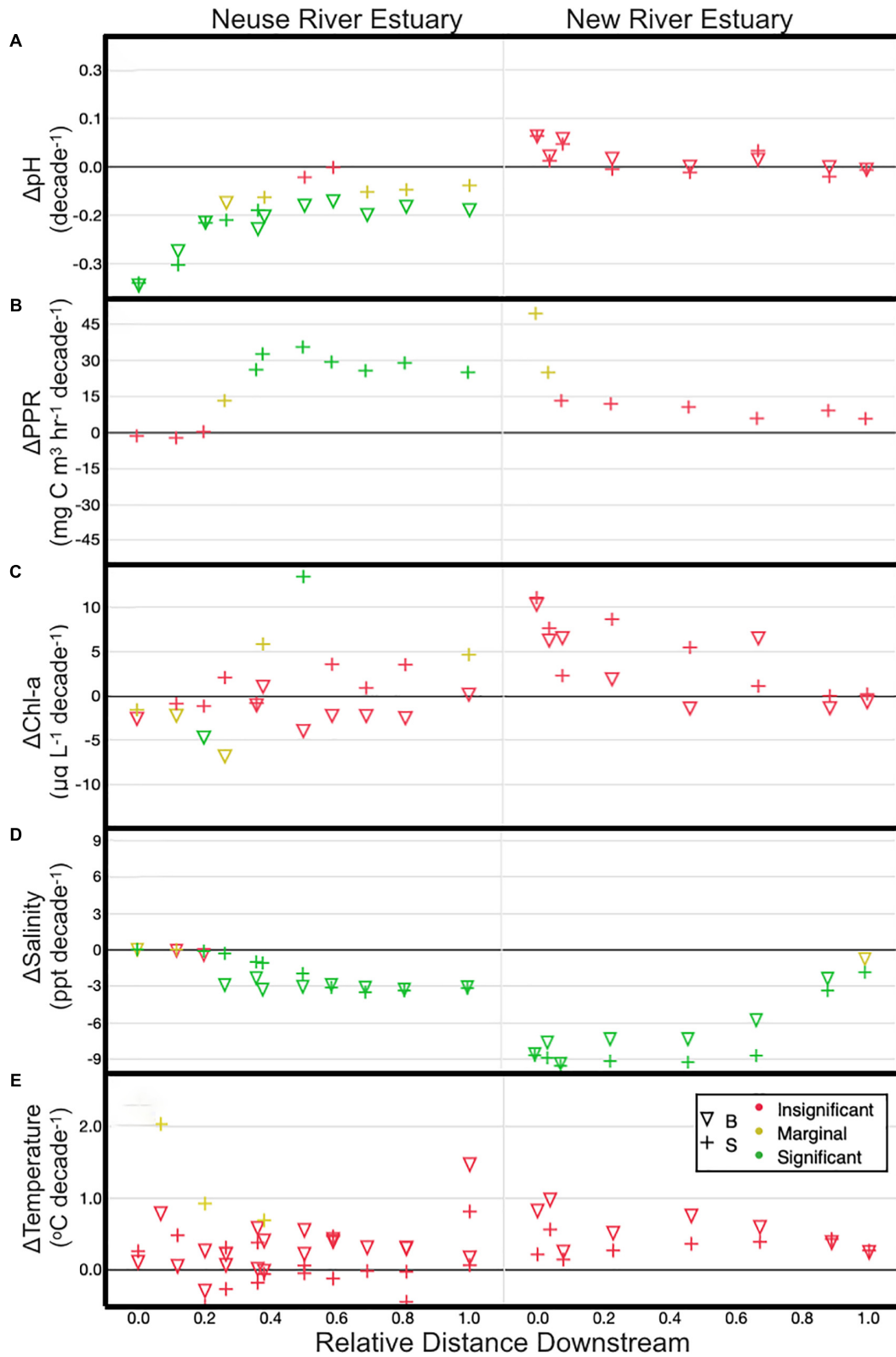


FIGURE 2 | Post-2005 trends for pH (A), PPR (B), Chl-a (C), Salinity (D), and Temperature (E) for surface (+) and bottom water (▽). Trends are shown relative to distance downstream, which is the distance from the head of the estuary divided by the total length of the estuary. Colors indicate the *p*-value for a *t*-test on the slope of the trend line (green: $p < 0.01$; yellow: $0.01 < p < 0.1$; red: $p > 0.1$).

and NewRE (20.8°C). A constant output temperature was used to control for the effect of changing temperature on pH and pCO₂.

RESULTS AND DISCUSSION

Physical Condition Change

From 1996 to 2017, Neuse river discharge at Ft Barnwell increased at an average rate of 1.55 m³ s⁻² yr⁻¹, or 2.8% yr⁻¹, while New River discharge at Gum Branch increased at 0.084 m³ s⁻² yr⁻¹ (1.4% yr⁻¹) between 2007 and 2017. Variations in monthly averaged discharge during the Spring and Winter were significantly correlated with El Nino Southern Oscillation (ENSO) patterns, such that positive ENSO phase was associated with increases in discharge (**Supplementary Figure S2**). However, ENSO phase could not explain changes in discharge during the

Fall and Summer. The observed increases in river discharge were reflected in a large and significant freshening trend in both estuaries. Averaged across all surface stations, salinity decreased at a rate of 0.002 to 0.35 yr⁻¹ in the NeuseRE, and 0.18 to 0.96 yr⁻¹ in the NewRE (**Figure 2D**). In both estuaries, trends in bottom water salinity followed those from surface water.

Despite generally non-significant changes in Chl-a (**Figure 2C**), PPR in the middle and lower NeuseRE increased significantly from 2005 to 2017, at a rate of approximately 30 mg C m⁻³ hr⁻¹ decade⁻¹. Elevated river discharge has been shown to drive a seaward shift in the location of peak phytoplankton biomass (Peierls et al., 2012), so it is not surprising that increased river inputs were accompanied with elevated PPR in the lower NeuseRE (**Figure 2B**). Coinciding with this increasing surface water PPR in the NeuseRE, bottom water pH fell significantly at approximately ~0.02 yr⁻¹ (0.2 decade⁻¹), while surface

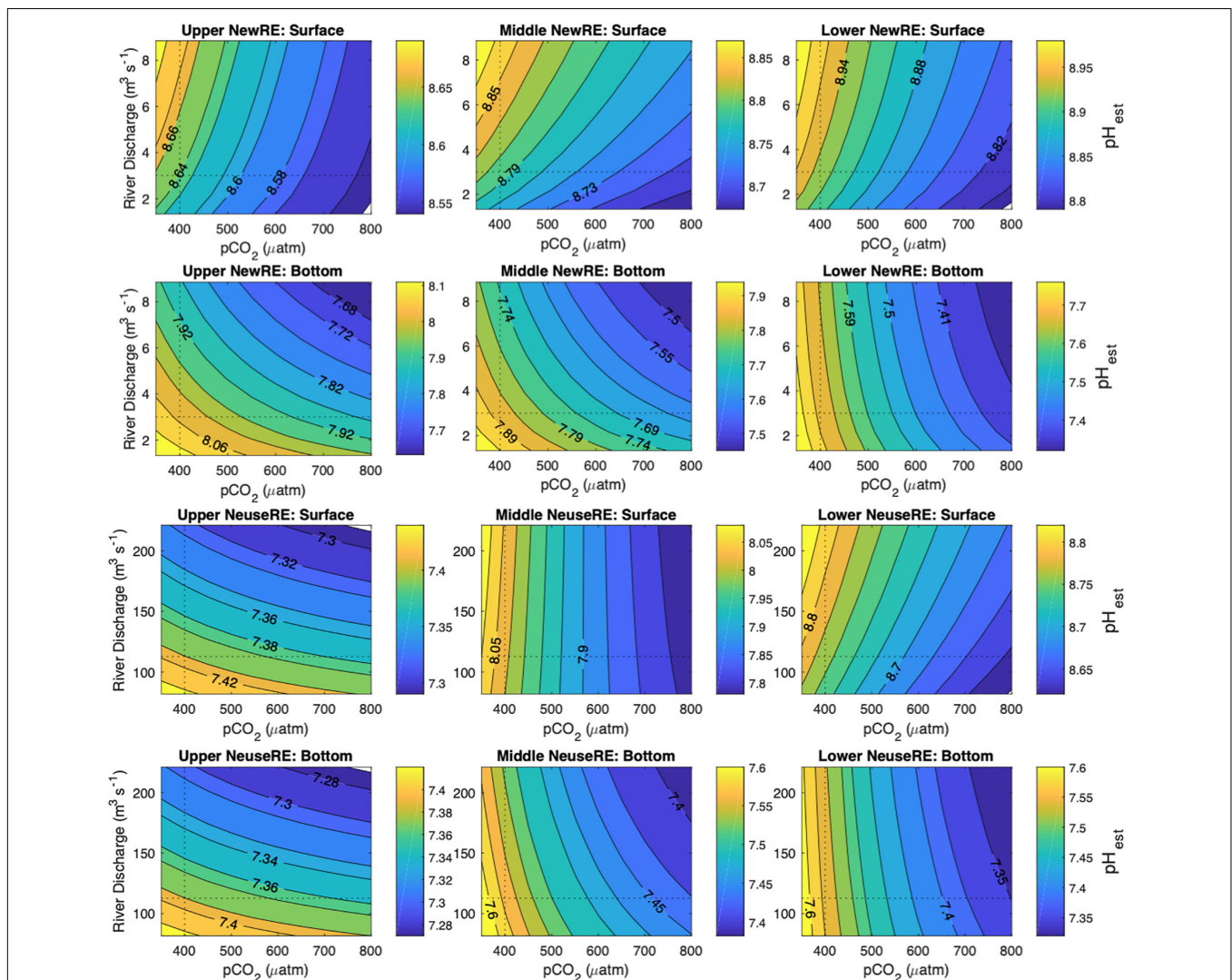


FIGURE 3 | Model pH_{est} as a function of atmospheric pCO₂ (μatm) and river discharge (m³ s⁻¹), where pH_{est} values are shown by the colored contours. Model results are shown for surface and bottom estuarine boxes, for the regions described in the **Supplementary Material**.

water pH trends were generally non-significant. Trends in bottom water DO were not significantly different from zero in the NeuseRE (**Supplementary Figure S3**), indicating that the observed decreases in pH may have been accelerated by factors beyond aerobic decomposition.

Trends in both PPR and pH for the NewRE were not significant (**Figures 2A,B**). Meanwhile, changes in temperature over the study period were generally insignificant and variable, between -0.09 and $0.28^{\circ}\text{C yr}^{-1}$ (average = $0.03 \pm 0.06^{\circ}\text{C yr}^{-1}$), similar to the trends observed in the coastal ocean (Reimer et al., 2017). Because pH decreases with temperature (holding DIC and TA constant), it is possible to assess the thermal effect on pH changes. At a TA and DIC of 900 and 1000 $\mu\text{mol kg}^{-1}$ respectively, an average temperature increase of $0.03^{\circ}\text{C yr}^{-1}$ results in a calculated pH decrease of -0.00024 yr^{-1} . This is approximately ~ 2 orders of magnitude below the observed trends in pH (**Figure 2A**), indicating that the effect of warming waters on acidification was negligible, relative to other factors.

Similar to the observed trend of acidification in the NeuseRE, which was accompanied by consistent DO, pH in the NewRE was stable (**Figure 2A**), while bottom water DO in the upper NewRE showed significant decreases (**Supplementary Figure S3**). This discrepancy between long term trends in DO and pH may be partially explained by the difference in air-water exchange for O_2 and CO_2 , which is more rapid for O_2 than for CO_2 . These counterintuitive trends in long-term acidification may be explained by changes in chemical buffering that served to accelerate acidification in the NeuseRE and to buffer acidification stress in the NewRE. Further explanation for this is given in the following section.

Box Model Results

In both estuaries, a doubling of oceanic pCO_2 from 400 to 800 μatm (reflecting estimated increases in atmospheric CO_2 over the next century) caused modeled surface water pH_{est} to fall by 0.094–0.14 in the NewRE, and by 0.037–0.23 in the NeuseRE. This modeled future decrease in pH over the next century corresponds to a rate of change of 0.004–0.023 decade^{-1} , which is within the range expected for the open ocean, but is an order of magnitude slower than the observed trend for the NeuseRE (**Figure 3**). This suggests that future trends in acidification will be driven by multiple interacting factors including the invasion of anthropogenic CO_2 , and other factors including changing riverine TA loads or NEM. In the upper NeuseRE, elevated discharge was associated with decreasing surface water pH_{est} (pH_{est} at high Q_R – pH_{est} at low Q_R = $\Delta\text{pH}_{\text{est}}$ = -0.086). Throughout the rest of the NeuseRE, increases in Q_R corresponded with increasing surface pH_{est} , such that $\Delta\text{pH}_{\text{est}}$ was 0.019 and 0.045 across the forecasted range in Q_R , in the middle and lower segments, respectively. In the NewRE, surface pH_{est} always increased with Q_R , with $\Delta\text{pH}_{\text{est}}$ between 0.037 and 0.14.

To summarize, the invasion of anthropogenic CO_2 resulted in moderate decreases in modeled pH_{est} , but this acidification was buffered when accompanied by elevated river discharge. The buffering effect of riverine inputs was greatest in surface waters

and appears to be larger in the NewRE than in the NeuseRE. This difference between estuaries is particularly pronounced in the upper, oligohaline regions of both estuaries, where increases in Q_R were accompanied by rising pH in the NewRE but decreasing pH in the NeuseRE. It is important to note that this difference in pH change between surface and bottom waters may be overestimated by the box model, which does not consider mixing between these two layers. However, this model does provide an indication that the pH in both estuaries are sensitive to variations in fresh water delivery.

Inputs of TA from the watershed, in excess of DIC loads, may help buffer CO_2 -driven acidification. Hence, variations in the ratio of TA:DIC have been used to indicate changes in the ability of water to buffer changes in pH (Hu and Cai, 2011). As shown in **Figure 4**, the TA:DIC ratio decreases exponentially with Q_R in both rivers. This suggests that both DIC and TA concentration in the river is primarily driven by weathering inputs, which are progressively diluted by surface water as Q_R increases. However, TA appears to decrease more rapidly than DIC as Q_R increases, causing the ratio of TA:DIC to decrease with rising Q_R , converging on a value of ~ 0.5 at high discharge (**Figure 4**). At low Q_R , TA:DIC was relatively high in the New River, occasionally exceeding 1:1. TA:DIC was always below 1:1 in the Neuse River. Decadal trends in the TA:DIC ratio were stable in the NewRE, but were significantly negative (decreasing trend) in much of the upper NeuseRE (**Supplementary Figure S3**), corresponding with the decreasing pH trend in that estuary.

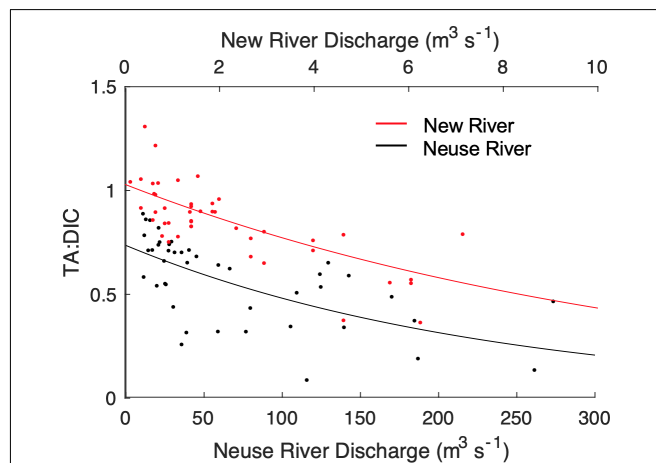


FIGURE 4 | Relationship between river endmember TA:DIC and discharge for the New (red points) and Neuse (black points) rivers. Exponential fits are of the form $f(x) = a \cdot \exp(b \cdot x)$, where the coefficients are $a = 0.74$ and $b = -0.0043$ for the Neuse River, and $a = 1.03$ and $b = -0.086$ for the New River. Water quality data (TA, pH, Temperature, Salinity) were obtained from STORET for the North Carolina Dept. of Environmental Quality site J7850000 (Neuse River near Ft Barnwell), and from sites 21NC01WQ-P0600000 and 21NC01WQ-P1200000 in the upper New River. Only matching pairs of TA, pH, temp, and salinity were used, yielding a relatively small set of data from 1990 to 1993. DIC was calculated from measured TA, pH, Temperature, and Salinity with CO2SYS. Matching discharge data were obtained from the USGS gage at Gum Branch for the New River (2093000), and the USGS gage at Kinston for the Neuse River (2089500).

To better understand the effect of changing river end-member TA:DIC on buffering within the estuary, we calculated pH across the estuarine salinity gradient (with CO2SYS) for a variety of river end-member TA:DIC ratios that fall within the range of **Figure 5A**. This assessment is intended to indicate potential pH values for an idealized estuary, and does not include DIC/TA production or consumption by net metabolic processes or air-water CO₂ exchange. When salinity is greater than ~20, carbonate chemistry is heavily controlled by mixing with well-buffered ocean water, thus, the decreasing riverine TA:DIC have limited effect on pH. However, when salinity falls below ~10, the impact of variable riverine TA:DIC ratios becomes more apparent, causing large changes in calculated pH (**Figure 5A**). Furthermore, because the ratio of TA:DIC in the coastal ocean is greater than 1 (at least in this case), scenarios where riverine TA:DIC is less than 1 result in a point where TA:DIC passes through 1:1 zone, where buffering is locally minimized (Hu and Cai, 2013). To more precisely quantify this process, we calculate a buffer factor (β_{DIC}), which can be expressed in partial differential form as in Egleston et al. (2010):

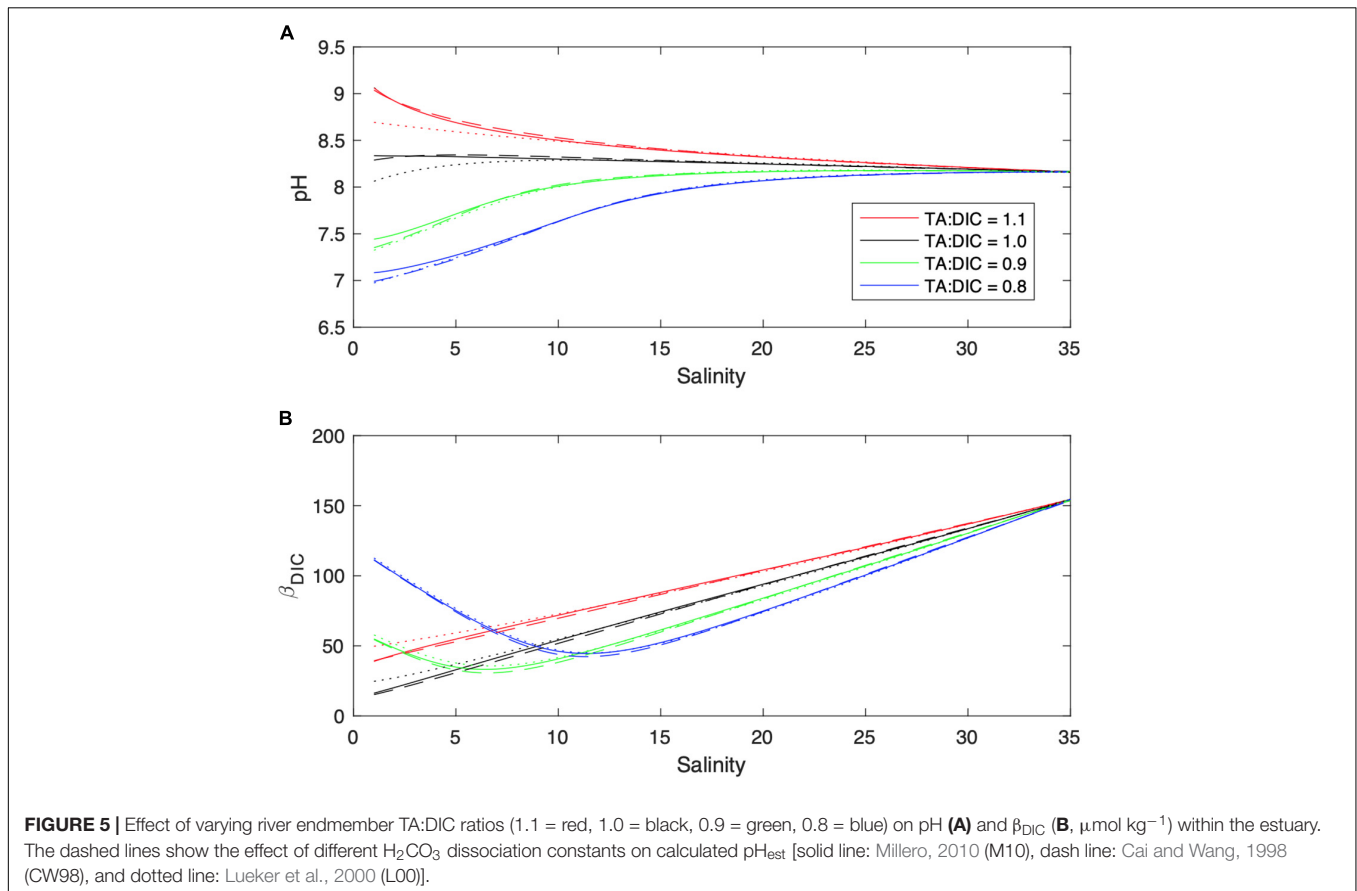
$$\beta_{DIC} = \left(\partial \ln [H^+] / \partial DIC \right)^{-1} \quad (6)$$

Expressed in this manner, increases in β_{DIC} indicate an increased buffering capacity, or resistance to changes in pH with DIC additions. As seen in **Figure 5B**, β_{DIC} exhibits a non-monotonic

relationship with salinity (when TA:DIC is < 1), such that buffering is minimized at a salinity of ~7 (riverine TA:DIC = 0.9) or ~12 (TA:DIC = 0.8). Hence, when riverine TA:DIC is less than 1, estuarine pH is particularly sensitive to additional DIC inputs, which can enter via changes in net metabolism, anthropogenic CO₂ inputs, etc. This is the case for the NeuseRE, where TA:DIC appears to always be less than 1. That the greatest pH decreases in the NeuseRE occurred near the river mouth (-0.3 decade⁻¹, **Figure 2A**) further supports the role of poorly buffered river water on estuarine pH trends. TA:DIC was relatively high in the New River (**Figure 4**), causing downstream pH in the estuary to be less sensitive to Q_R. This concept is further supported by decadal trends in the TA:DIC ratio, which decreased at a significant rate across much of the upper NeuseRE, and was stable across the entire NewRE (**Supplementary Figure S3**). To summarize, it appears that TA delivery from the watershed (in excess of DIC) helped to buffer acidification stress in the NewRE, while decadal decreases in TA relative to DIC served to diminish the buffering capacity of the NeuseRE, allowing significant pH decreases to occur.

Summary of Acidification Drivers

A vast and tangled array of factors contribute to temporal variations in estuarine pH. Given this complexity, we determined in a statistically robust manner, the magnitude and direction of decadal-scale biogeochemical trends for two adjacent estuaries.



Further, we identified which factors were the most important to the observed pH trends and determined if these factors differ between the two estuaries studied. While it is beyond the scope of this study to precisely quantify the many components affecting pH, we shed some light on the diverse drivers of estuarine acidification, a process that has gained interest in recent years.

Increases in river discharge [associated with ENSO activity (**Supplementary Figure S2**) during the Spring and Winter], coincided with a significant trend of acidification in the NeuseRE between 2005 and 2017. Acidification was greatest in bottom waters, concurrent with increased PPR in the surface, highlighting the importance of CO₂ inputs from net ecosystem respiration. Temporal trends in pH across the NewRE were small and not significant, despite moderate, but generally insignificant increases in phytoplankton biomass and productivity. Small, insignificant warming trends in both estuaries could not explain the observed bottom-water pH decrease. However, in the NewRE, this acidification ‘pressure’ was largely tempered by riverine loading of TA, which was higher relative to DIC. Inputs of TA from the Neuse River were smaller, however, leading to a weak buffer capacity that was more sensitive to autochthonous and allochthonous CO₂ inputs (**Figure 5**). While it is beyond the scope of this study to rigorously assess climate teleconnections like those between ENSO and annual-scale biogeochemical change, this topic may be of interest for future research, given the likely variations in ENSO with climate change.

Rising atmospheric CO₂ levels will act as a downward pressure on estuarine pH, but will be modified the net production or consumption of CO₂ by net ecosystem metabolism. Thus, in other stratified estuaries experiencing eutrophication, net respiratory inputs of CO₂ to bottom waters may interact with this anthropogenic CO₂ to geometrically lower pH. Meanwhile, concurrent change in river discharge (due to precipitation or human water consumption) have the potential to either enhance or ameliorate these acid inputs, although to an extent that varies according to differences in underlying lithology and hydrology. Hence, all estuaries experiencing future changes in river discharge may experience similar pH trajectories due to the interaction between atmospheric CO₂ inputs and riverine hydrogeochemistry. Local decision makers and stakeholders should consider these factors when considering management strategies aimed at coping with future estuarine acidification.

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DATA AVAILABILITY

The DCERP (<https://dcerp.rti.org/>) and ModMon (<http://paerllab.web.unc.edu/projects/modmon/products-and-applications/>) datasets are published on the data sharing repository Figshare (https://figshare.com/articles/WQ_Datasets_-_Decadal_Acidification_Trends/7643195).

AUTHOR CONTRIBUTIONS

BVD and HW contributed equally to the conception and design of the study.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/fmars.2019.00136/full#supplementary-material>

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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