

Acceleration of ocean warming, salinification, deoxygenation and acidification in the surface subtropical North Atlantic Ocean

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Ocean chemical and physical conditions are changing. Here we show decadal variability and recent acceleration of surface warming, salinification, deoxygenation, carbon dioxide (CO₂) and acidification in the subtropical North Atlantic Ocean (Bermuda Atlantic Time-series Study site; 1980s to present). Surface temperatures and salinity exhibited interdecadal variability, increased by -0.85 °C (with recent warming of 1.2 °C) and 0.12, respectively, while dissolved oxygen levels decreased by -8% (-2% per decade). Concurrently, seawater DIC, *f*CO₂ (fugacity of CO₂) and anthropogenic CO₂ increased by -8%, 22%, and 72% respectively. The winter versus summer *f*CO₂ difference increased by 4 to 8 μatm decade⁻¹ due to seasonally divergent thermal and alkalinity changes. Ocean pH declined by 0.07 (-17% increase in acidity) and other acidification indicators by -10%. Over the past nearly forty years, the highest increase in ocean CO₂ and ocean acidification occurred during decades of weakest atmospheric CO₂ growth and vice versa.

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Recent acceleration of changes in ocean physics and chemistry since the 1980s is demonstrated at two sustained open-ocean hydrographic stations in the North Atlantic Ocean near Bermuda. Of these two time-series sites, Hydrostation S (formerly known as the *Panulirus* site) is the oldest, located ~25 km southeast of Bermuda at 32°10'N, 64°30'W, and consists of repeat biweekly hydrographic observations of temperature, salinity and dissolved oxygen (DO) conducted through the water column since 1954^{1,2}. The Bermuda Atlantic Time-series Study (BATS) site is located ~80 km southeast of Bermuda (31°40'N, 64°10'W)³. It consists of a comprehensive monthly sampling of the physics, chemistry and biology of the entire water column commencing in October 1988^{4–7}, and one of only a handful of ship-based biogeochemical time-series sites remaining in the global ocean⁷. The BATS–hydrostation S time-series represent the two longest records in the global open ocean of warming, salinification, deoxygenation and changes in ocean carbon dioxide (CO₂)–carbonate chemistry that drives ocean acidification (OA). Observations over the past 40 years in the subtropical gyre of the North Atlantic Ocean allow for examination of variability and the trends of ocean change which critically underpin future assessment and prediction. BATS–Hydrostation S time-series data also provide partial support for the findings of a recent study indicating that the highest increases in ocean CO₂ and OA occur during decades of weakest atmospheric CO₂ growth and vice versa⁸. Here, we focus on surface/mixed layer changes while examination of variability and trends in deeper waters is the subject of other studies.

The emissions of anthropogenic CO₂ to the atmosphere due to fossil fuel use, cement manufacture and land-use changes⁹ have increased rapidly over the last decade¹⁰. While anthropogenic CO₂ accumulates in the atmosphere, it is also taken up by the terrestrial biosphere and oceans¹¹. The annual global ocean uptake is estimated^{10,12–14} at 1.4–2.6 Pg C yr⁻¹ (Pg = 10¹⁵ g) at present, with annual rates of the ocean CO₂ sink increasing with time¹³, with the cumulative total global ocean uptake of anthropogenic CO₂ since pre-industrial times estimated at 120–150 Pg C^{12–14}. The range of global ocean anthropogenic CO₂ uptake reflects the different year of the study and evolving data-driven knowledge about the oceans. Ocean uptake of anthropogenic CO₂ and seawater chemistry changes such as the reduction in seawater pH, and saturation states for calcium carbonate (CaCO₃) minerals (i.e., Ω) such as calcite (Ω_{calcite}) and aragonite ($\Omega_{\text{aragonite}}$) is termed ocean acidification (OA)¹⁵. It has likely problematic consequences for marine organisms and ecosystems that are, as yet, poorly understood¹⁶.

In addition, the decrease in ocean DO concentration¹⁷, a process termed “ocean deoxygenation (ODO)”^{18,19}, is a global environmental issue that has profound future implications for the status of the oceans^{20,21}. The emerging contemporary narrative about ODO relates to its coupling with other global environmental changes such as ocean warming and OA²¹. Oceanic DO is highly sensitive to solubility effects²², and thus in turn to ocean physics and air–sea exchange, and long-term drivers such as climate change, stratification and ocean primary production^{17–21,23}. A broader environmental context for this study is that there are complex interactions between OA, ODO, ocean biogeochemical cycling, and ocean biology, including microbial pathways²⁴ to zooplankton²⁵, and expansion of oxygen minimum zones^{26,27}. Careful and critical evaluation of such synergistic global environmental drivers is warranted²⁸.

Ocean sampling and methods

The Hydrostation S dataset represents more than 1381 cruises undertaken during the 1954–2020 period, while the BATS dataset

represents more than 450 cruises from October 1988 to the end of 2019. Sampling and analytical methods are detailed in the “Methods” section (Supplemental Materials). Measurements of seawater CO₂–carbonate chemistry include those of dissolved inorganic carbon (DIC), and total alkalinity (TA) since 1983 (1983–1988, Hydrostation S; 1988–present, BATS), with detailed sampling protocols, analytical approaches and the computation of other parameters such as pH and $p\text{CO}_2$ described in the “Methods” section. DIC and TA data are also normalised to a salinity of 36.6 (mean value in the Sargasso Sea) to remove the influence of evaporation and precipitation, and denoted as nDIC and nTA, respectively.

The suite of seawater CO₂–carbonate parameters (i.e., $f\text{CO}_2$, $p\text{CO}_2$, pH, Ω_{calcite} and $\Omega_{\text{aragonite}}$, Revelle factor) are determined for each sample using DIC and TA at in situ temperature and salinity (see “Methods”). pH, Ω_{calcite} , and $\Omega_{\text{aragonite}}$ in seawater are dimensionless and pH is expressed on the total scale.

Seasonally detrended anomalies of physical and chemical properties are shown to illustrate changes over time (Figs. 1 and 2) with trend analyses summarised in Table 1a for the past 40 years, and for the decades of the 1980s, 1990s, 2000s and 2010s. It is important to note that while surface data are shown and detailed here, seasonal variability and trends in the mixed layer are nearly identical to surface data. A tracer, TrOCA (Tracer combining Oxygen, inorganic Carbon and total Alkalinity)²⁹ is used to test

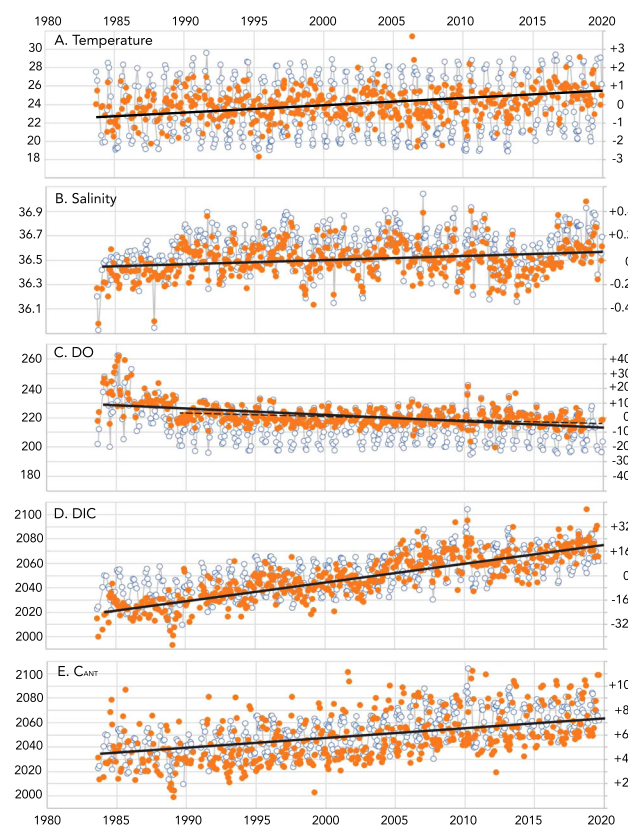


Fig. 1 | Hydrographic and seawater CO₂–carbonate properties at the BATS site (1988–present) with earlier data (1983–1988) from Hydrostation S. Observed data (left vertical axis; open blue symbols) and anomalies (right vertical axis; orange symbols) are shown. Regression lines are from anomaly data plotted in the right vertical axis and given in Table 1. **a** Surface temperature and anomalies (°C). **b** Surface salinity and anomalies. **c** Surface dissolved oxygen (DO) and DO anomalies (μmoles kg⁻¹); the dashed line shows the trend from 1990. **d** Surface DIC and salinity normalised DIC (nDIC; μmoles kg⁻¹). **e** nDIC and C_{ANT} values (μmoles kg⁻¹).

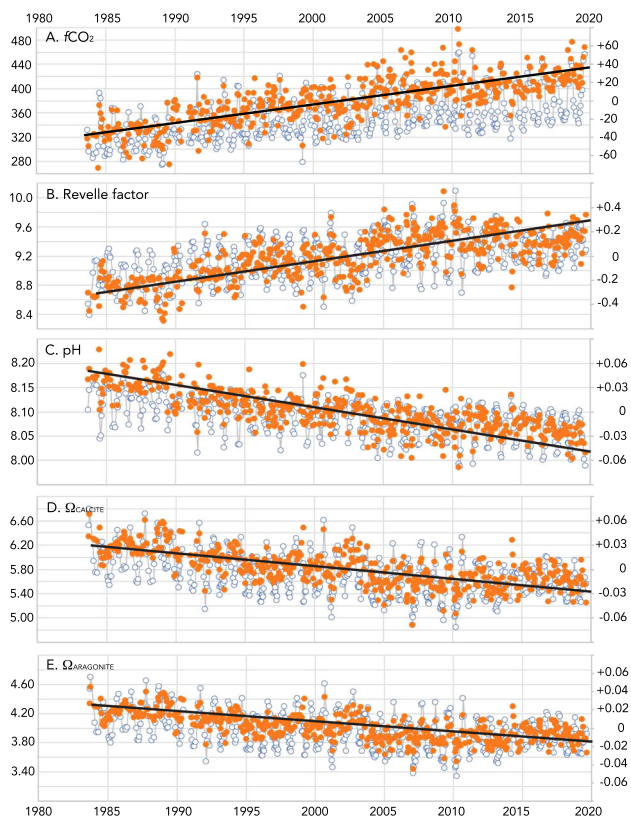


Fig. 2 Hydrographic and seawater CO₂–carbonate properties at the BATS site (1988–present) with earlier data (1983–1988) from Hydrostation S. Observed data (left vertical axis; open blue symbols) and anomalies (right vertical axis; orange symbols) are shown. Regression lines are from anomaly data plotted in the right vertical axis and given in Table 1. **a** Surface $f\text{CO}_2$ and anomalies (μatm). **b** Surface Revelle factor (β) and anomalies. **c** Surface pH and anomalies. **d** Surface Ω_{calcite} and anomalies. **e** Surface $\Omega_{\text{aragonite}}$ and anomalies.

the evidence for drivers of ocean seawater CO₂–carbonate changes, and the decadal variability in global ocean carbon uptake^{8,30}.

Results

Surface ocean warming and salinification. Menzel and Rytner¹ first documented seasonal cycles of the local convective mixed layer with subsequent papers improving our understanding of the year-to-year and multi-decadal variability of the physical seasonality of the subtropical gyre and relationships to climate drivers^{3,4,6}. The Sargasso Sea exhibits a seasonal sea surface temperature variation of ~ 9 – 11 °C (Fig. 3a) with higher surface salinities occurring in wintertime and freshening during the summertime, and low salinity events occasionally apparent (Fig. 3c).

Over the past nearly 40 years (1983–2019), surface temperatures have increased by 0.85 ± 0.12 °C (Table 1a; Figs. 1a and 3b). The summer surface temperatures have increased at a higher rate than winter (i.e., 0.26 ± 0.01 °C year⁻¹ compared to 0.10 ± 0.01 °C year⁻¹; Table 1a; note that winter rates are not statistically significant). During the same period, surface salinity also increased by $\sim 0.11 \pm 0.02$ (Table 1a; Fig. 1b). Importantly, these data show evidence of decadal variation in Table 1. Trend analysis of temperature, salinity, oxygen and ocean CO₂–carbonate chemistry from the BATS site (1988–2018; Hydrostation S data from 1983 to 1988) is included (see also Table 1b). Intermediate

and deep water variability and trends (including warming and cooling, and freshening) is the subject of a future comprehensive treatment, with, for example, warming below the mixed layer at 300 m at ~ 0.8 °C over the past three decades.

In the 1990s and 2000s, no significant trends in surface temperature and salinity were observed (Table 1a). In contrast, warming and salinification occurred during the 1980s and 2010s (Fig. 1a, b). In the 1980s, salinity increased by 0.27 (Table 1a) associated with the 1989–1990 transition from lower to higher salinity in the subtropical gyre (Fig. 3c, d). During the last decade (2010–2019), rapid warming of 1.18 °C and salinification of 0.14 has occurred. Temperature and salinity trends go back to 1954 (Table 1b). The warming of 1.18 °C in the last decade may at first appear inconsistent with overall warming of $\sim 0.85 \pm 0.12$ °C over the past 40 years. The longer trend estimate since 1983 reflects periods of interannual/decadal warming and cooling (e.g., the decades of the 1990s and 2000s had no statistical change) with overall warming since the early 1980s.

Surface ocean deoxygenation. The upper-ocean record of DO shows an overall decline of $\sim 17.8 \pm 2.4$ $\mu\text{moles kg}^{-1}$ over the past 40 years (Table 1a; Fig. 1c), with short-term year-to-year variability (Fig. 3e, f). The longer-term data represents a loss of DO of $\sim 8.1\%$ or $\sim 2\%$ per decade. The trend of DO decline in the Sargasso Sea since the 1980s is in the range shown for global ODO rates of ~ 0.3 to ~ 0.7 $\mu\text{moles kg}^{-1}$ year⁻¹ (~ 3 – 7 $\mu\text{moles kg}^{-1}$ decade⁻¹) in the thermocline^{31,32}.

Apparent oxygen utilisation (AOU) represents the difference between measured oxygen concentration and oxygen concentration at saturation in seawater with the same physical and chemical properties. AOU, as a calculated parameter²², offers a few clues as it reflects changes in oxygen due to the combined effect of biological processes such as primary production and respiration³², but also diffusion, circulation, and ventilation¹⁹. Here, the contribution of nitrification and denitrification to AOU is considered minor in surface waters. The trend in AOU in the upper 500 m is about -0.54 $\mu\text{moles kg}^{-1}$ year⁻¹, which is slightly higher than the contemporaneous decrease in DO (Table 1a).

The cause for deoxygenation of the Sargasso Sea over the past 40 years appears only partly due to solubility changes associated with warming. Given the ocean warming observed in the Sargasso Sea (Table 1a), we estimate that the warming impact on DO solubility¹⁶ would likely have contributed to about 13% of the total decline of DO over the past nearly 40 years. The remaining deoxygenation ($\sim 87\%$) must result from the combined effect of changes in ocean biology and physics. Recent warming in the 2010s of 1.18 °C contributed about 18% of the concurrent decline in DO in that decade. From 1988 to 2017, open-ocean chlorophyll biomass ($\sim +22\%$ per decade³³) and rates of in situ primary production observed offshore at BATS increased³³. Coccolithophore (i.e., calcifying phytoplankton) biomass has increased at BATS³⁴ and over the entire subtropical gyre of the North Atlantic Ocean, but it is typically a small ($<10\%$) contributor to primary production. Furthermore, low nitrate values in the mixed layer have doubled (i.e., an increase of ~ 0.25 $\mu\text{moles kg}^{-1}$) over the past two and a half decades while suspended particulate organic matter and dissolved organic carbon have increased by $\sim 30\%$ and 2% per decade, respectively³³. Other studies indicate that the nutrient stream supporting production in the subtropical North Atlantic has reduced³⁵. These puzzles are not addressed here, but likely includes the loss of oxygen from the surface ocean due to gas flux, and perhaps supplies of new nitrogen sources from nitrogen fixation and dissolved organic nitrogen.

Table 1 Trend analysis of physical properties, seawater CO₂-carbonate chemistry, and ocean acidification indicators at the BATS-Hydrostation S site from 1983 to 2020.

	n	Anomaly data and trends			Change over period					
		r/r ²	p Value	Slope/error	Entire	%	1980s	1990s	2000s	2010s
(a) BATS (1983–2019)										
<i>Physical properties</i>										
Temperature (°C)	466	0.26/0.07	<0.001	+0.021 ± 0.003	+0.85	+3.6%	+0.26	−0.10	−0.17	+1.18
Temp (winter)	125	0.02/0.01	0.15	+0.10 ± 0.006	+0.40					
Temp (summer)	104	0.09/0.008	<0.001	+0.26 ± 0.006	+1.06					
Salinity	466	0.22/0.05	<0.001	+0.0029 ± 0.0008	+0.11	+3.1%	+0.270	+0.026	+0.007	+0.143
DO (μmoles kg ^{−1})	466	0.31/0.11	<0.001	−0.37 ± 0.05	−17.76	−8.3%	−30.22	−5.33	−4.90	−17.56
<i>Alkalinity (units are μmoles kg^{−1})</i>										
TA	457	0.40/0.16	<0.001	+0.26 ± 0.03	+11.72	+0.5%	+6.82	−2.66	+3.60	+17.40
nTA	457	0.10/0.01	<0.03	+0.07 ± 0.05	+3.76	+0.1%	−7.37	−0.83	+5.05	+7.39
<i>DIC and Anthropogenic CO₂ (units are μmoles kg^{−1})</i>										
DIC	458	0.83/0.69	<0.001	+1.26 ± 0.04	+50.28	+2.3%	+0.24	+9.02	+21.65	+12.65
nDIC	458	0.85/0.72	<0.001	+1.07 ± 0.03	+42.76	+2.1%	−11.87	+9.43	+21.38	+4.54
C _{ANT}	458	0.64/0.44	<0.001	+1.03 ± 0.05	+38.96	+72%	−19.40	+8.70	+18.10	+15.00
<i>fCO₂ and Revelle buffer factor (units are μatm for fCO₂)</i>										
fCO ₂	458	0.77/0.50	<0.001	+1.92 ± 0.08	+76.88	+22.3%	+1.90	+20.60	+32.14	+18.40
Revelle (β)	458	0.70/0.49	<0.001	+0.014 ± 0.001	+0.54	+5.8%	0.00	0.16	0.27	−0.05
fCO ₂ (winter)	125	0.76/0.57	<0.001	+1.61 ± 0.13	+64.66					
fCO ₂ (summer)	104	0.78/0.60	<0.001	+2.40 ± 0.19	+95.88					
[Winter–summer fCO ₂]							83.5 ± 17.2	87.9 ± 22.7	89.9 ± 14.7	99.9 ± 30.5
<i>Ocean acidification indicators</i>										
pH	458	1.2881356	<0.001	−0.0019 ± 0.0001	−0.08		−0.004	−0.022	−0.031	−0.016
Ω _{calcite}	458	1.547619	<0.001	−0.014 ± 0.001	−0.58	−10.3%	−0.08	−0.17	−0.28	−0.08
Ω _{aragonite}	458	1.575	<0.001	−0.009 ± 0.001	−0.37	−10.0%	−0.05	−0.12	−0.18	−0.06
Note: p value > 0.05 denotes not significant trend for the period										
(b) Hydrostation S trends (1954–2019)										
Temperature (°C)	1441	0.23	<0.01	+0.01 ± 0.001						
Salinity	1227	0.19	<0.01	+0.002 ± 0.001	+0.10					
DO (μmoles kg ^{−1})	1203	−0.13	<0.01	−0.07 ± 0.02	−4.8					

Data comprises BATS site (1988–present) with earlier data (1983–1988) from Hydrostation S. Linear regression using a least-squares method is given with regression statistics, residuals and coefficients, and with p values < 0.05 represent significance at the 95% level. In Table 1a, trends for seasonally detrended data are given, but the slopes are identical at the decimal place given to the observed data. The difference in regression statistics is that the r² is higher, and p value and standard error of the slope is reduced for seasonally detrended data. The winter and summer fCO₂ trends are determined for January to March (JFM) and July to September (JAS), respectively.

Surface ocean TA and DIC changes. The observations near Bermuda provide the longest record of seawater CO₂-carbonate chemistry in the global open-ocean. Surface TA shows similar seasonal patterns to salinity, and a slight increase associated with salinity increases over time (Fig. 3c). However, salinity normalised alkalinity (nTA) shows marginally statistically significant changes over time (Table 1a; p-value of 0.03).

Surface DIC exhibits typical summer to wintertime seasonality of ~30–40 μmoles kg^{−1}, (Fig. 4a, b; salinity normalised DIC also shows a similar pattern but with reduced seasonality^{13,36}). Surface DIC and nDIC have significantly increased at rates of 1.26 ± 0.04 and 1.07 ± 0.03 μmoles kg^{−1} year^{−1}, respectively, over the past 40 years (Fig. 1d; Table 1a). The difference between DIC and nDIC trends indicate that salinity changes contributed about 20% of the changes in DIC.

Earlier studies at BATS^{36–40}, elsewhere off Hawaii⁴¹ and the Canary Islands⁴², attribute the increase in DIC to the uptake of anthropogenic CO₂ from the atmosphere⁷. Here, we use the TrOCA method to determine if the remaining 80% of DIC

increase is due to anthropogenic CO₂ uptake. The TrOCA method²⁹ has limitations in use for anthropogenic CO₂ quantification⁴³ but nevertheless, provides a useful tracer of ocean chemical changes when used at a fixed location. In the Sargasso Sea, TrOCA derived anthropogenic carbon concentration (C_{ANT}) surface values exhibit seasonal ranges of ~30–40 μmoles kg^{−1} (Fig. 4c, d). Importantly, C_{ANT} has increased by ~38 μmoles kg^{−1} (+72%) over the past 30 years, with anthropogenic CO₂ concentration almost doubling in the upper ocean (Fig. 1d; Table 1a). The rate of increase of C_{ANT} and nDIC are very similar (1.03 ± 0.05 vs. 1.07 ± 0.03 μmoles kg^{−1} year^{−1}; Table 1a) and indicative that ocean uptake of anthropogenic CO₂ uptake has substantially changed ocean carbon chemistry over the past four decades at least.

Surface ocean fCO₂ and Revelle factor changes. Surface seawater fCO₂ also exhibits substantial seasonality with typical ranges of ~40–60 μatm (Figs. 2a and 4e, f). This variability reflects the underlying seasonality of temperature and DIC (similar trends

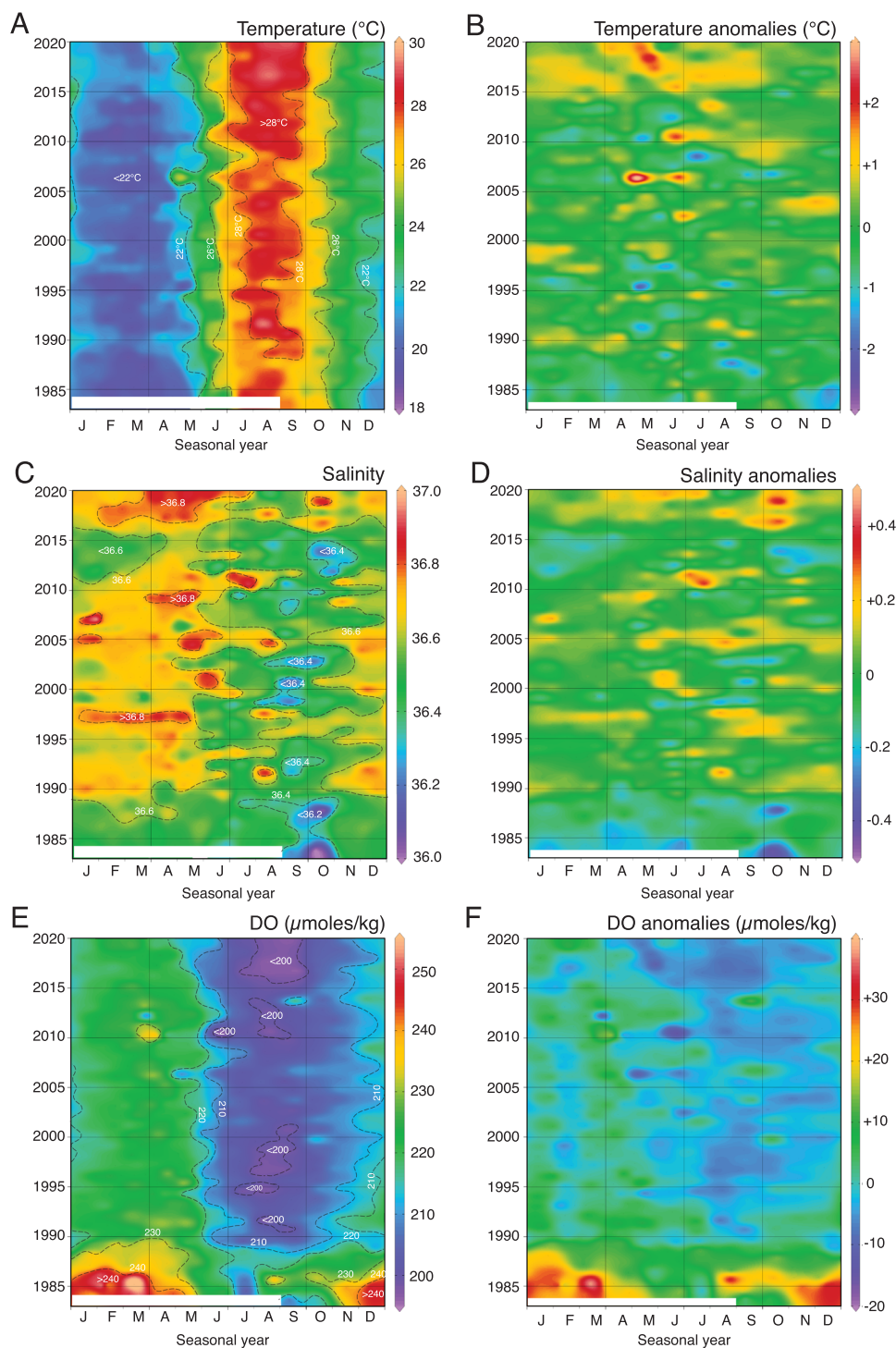


Fig. 3 Physical properties at BATS (1988–present) with earlier data (1983–1988) from Hydrostation S. **a** Surface temperature (°C). **b** Surface temperature anomalies (°C). **c** Surface salinity. **d** Surface salinity anomalies. **e** Surface dissolved oxygen DO ($\mu\text{moles kg}^{-1}$). **f** Surface dissolved oxygen DO anomalies ($\mu\text{moles kg}^{-1}$). Please note that the occasional high and low summertime DO events observed in the 2000s occur both at hydrostation S and BATS indicating that these features are “real” surface phenomena. Dashed lines for contours are included.

and patterns observed for $p\text{CO}_2$, or partial pressure of CO_2 , with typically less than $1 \mu\text{atm}$ difference between $f\text{CO}_2$ and $p\text{CO}_2$). In response to anthropogenic CO_2 uptake by surface waters and an increase in DIC, surface seawater $f\text{CO}_2$ exhibited significant gain over the 40 years with $f\text{CO}_2$ anomalies increasing by $\sim 76 \mu\text{atm}$ (Fig. 4f; Table 1a). The rate of change of $f\text{CO}_2$ is $1.92 \pm 0.08 \mu\text{atm year}^{-1}$ (Table 1a), representing an increase of nearly 25% from 1980 to 2020. The winter and summer $f\text{CO}_2$ values in recent years

are higher and outside the range of surface $f\text{CO}_2$ seasonality observed in the 1980s.

The increasing trend in seawater $f\text{CO}_2$ observed in the North Atlantic Ocean is similar to observed changes in atmospheric $f\text{CO}_2$ ^{16,41}, and, over shorter time-scales, similar trends in surface seawater $f\text{CO}_2$ have been reported elsewhere^{41,42}. The observation that surface and atmospheric $f\text{CO}_2$ have tracked each other over time implies that the driving force for air–sea CO_2 gas exchange

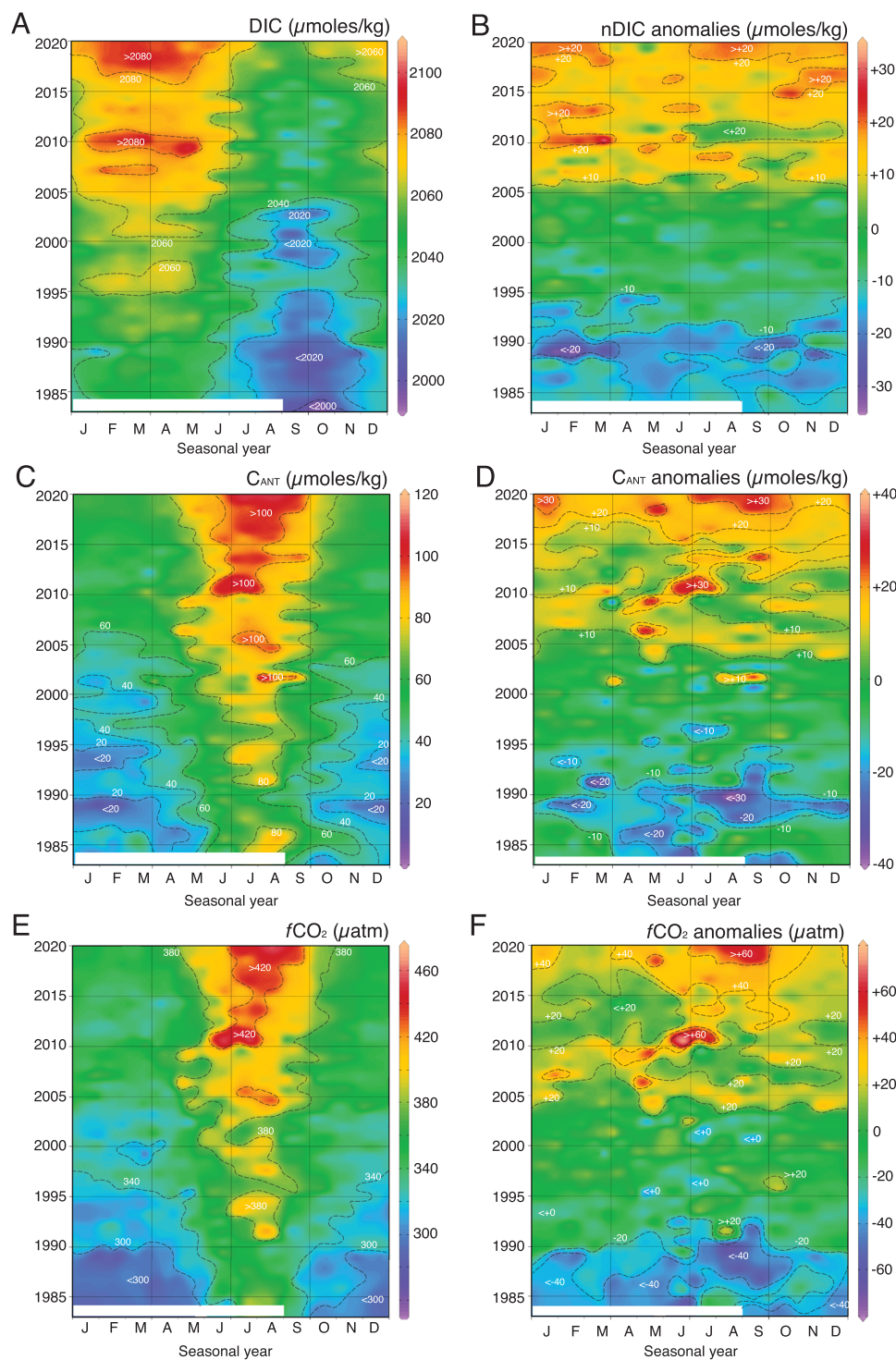


Fig. 4 Seawater CO₂-carbonate properties at BATS (1988–present) with earlier data (1983–1988) from Hydrostation S. **a** Scatter plot of DIC ($\mu\text{moles kg}^{-1}$). **b** Scatter plot of nDIC anomalies ($\mu\text{moles kg}^{-1}$). **c** Scatter plot of C_{ANT} ($\mu\text{moles kg}^{-1}$). **d** Scatter plot of C_{ANT} anomalies ($\mu\text{moles kg}^{-1}$). **e** Surface $f\text{CO}_2$ (μatm). **f** Surface $f\text{CO}_2$ anomalies (μatm). Dashed lines for contours are included.

(i.e., $\Delta f\text{CO}_2$; the $f\text{CO}_2$ difference between atmosphere and seawater) and the ocean CO₂ sink in the subtropical gyre has not changed significantly over the past 40 years. Earlier studies^{7,40} have concluded that the increases in $f\text{CO}_2$ largely reflect uptake of anthropogenic CO₂ and increasing DIC, with minor influences due to the small changes in temperature, salinity and alkalinity over time. As shown later, while surface and atmospheric $f\text{CO}_2$ have tracked each other over the past 40 years, there is substantial decade-to-decade variation of seawater $f\text{CO}_2$ increase (Table 1a;

low of +1.9 μatm per decade in the 1980s to +32 μatm per decade in the 2000s) and ocean CO₂ uptake.

The Revelle factor (β) reflects the underlying chemical state that can facilitate air–sea CO₂ exchange⁴⁴. In simple terms, the ocean’s ability to absorb CO₂ decreases with higher β values^{12,45}, and the chemical buffering capacity reduces. In the Sargasso Sea, the seasonality of β is ~ 0.4 , with wintertime values higher than summertime values. Lower β conditions reflect the underlying capacity for seawater to absorb CO₂, and yet the lowest $f\text{CO}_2$

occurs in winter coincide with seasonally higher β values. This finding demonstrates the influence of other factors on $f\text{CO}_2$, such as temperature. An important observation is that the Revelle factor, β , is increasing over time ($\sim 0.54 \pm 0.04$; Fig. 2b, Table 1a). These changes in ocean chemistry illustrate that the ocean capacity to absorb CO_2 from the atmosphere is gradually declining ($\sim 6\%$ since 1980), which provides additional future feedback for assessments of the quantitative fate of anthropogenic CO_2 in the atmosphere and its transfer to the ocean.

Recently, Landschutzer et al.⁴⁶ reported winter to summer $f\text{CO}_2$ difference has increased from the 1980s to the 2010s by an average of $2.2 \mu\text{atm}$ per decade with a rate of 1.5 and $3.8 \mu\text{atm}$ per decade report for BATS and the Hawaii Ocean Time Series (but without details in the paper about how these rates were estimated). Such changes are attributed primarily to the long-term increase in the mean concentration of CO_2 in the surface ocean, $[\text{CO}_2]_{\text{aq}}$, and increase in the Revelle factor due to the reaction of added CO_2 with carbonate ion to reduce the capacity of the surface ocean CO_2 -carbonate system to buffer against changes. At the BATS-Hydrostation S site, it is difficult to capture the peak maximum (summer) and minimum (winter) $f\text{CO}_2$ with monthly shipboard observations (i.e., the maximum and minimum $f\text{CO}_2$ may occur between cruises). Nonetheless, the mean winter to summer $f\text{CO}_2$ difference increases from the 1980s to 2010s (e.g., mean and standard error; 1980s, $83.6 \pm 17.2 \mu\text{atm}$; 1990s, $87.9 \pm 22.7 \mu\text{atm}$; 2000s, $89.9 \pm 14.7 \mu\text{atm}$; 2010s, $99.2 \pm 11.4 \mu\text{atm}$; Table 1a) contemporaneously with increasing $[\text{CO}_2]_{\text{aq}}$ and Revelle factor. The increase in mean winter to summer $f\text{CO}_2$ difference is $4.1 \mu\text{atm}$ per decade. Over the past nearly 40 years, the winter and summer $f\text{CO}_2$ have increased at different rates of $1.62 \pm 0.13 \mu\text{atm}$, and $2.40 \pm 0.19 \mu\text{atm}$, respectively (Table 1a; total $f\text{CO}_2$ change of $\sim 65 \mu\text{atm}$ and $\sim 96 \mu\text{atm}$ in winter and summer). The winter and summer $f\text{CO}_2$ difference has increased by $\sim 30 \mu\text{atm}$ which is $\sim 7.9 \mu\text{atm}$ per decade. The two approaches thus indicate increases of winter and summer $f\text{CO}_2$ difference in the range of 4.1 – $7.9 \mu\text{atm}$ per decade, and three to five times the rates reported by Landschutzer et al.⁴⁶ (their data covers the period 1988–2014). The divergent winter and summer $f\text{CO}_2$ conditions appear associated with reduction in winter length and lower rate of warming (the season with water less than 22°C has reduced from 157.7 ± 10.8 days in the 1980s to 124.7 ± 24.7 days in the 2010s; $0.10^\circ\text{C decade}^{-1}$; Table 1a), and increase in summer length and higher rate of warming (the season with water greater than 25°C has reduced from 121.0 ± 23.4 days in the 1980s to 140.3 ± 11.3 days in the 2010s; $0.26^\circ\text{C decade}^{-1}$; Table 1a). The BATS-Hydrostation S winter to summer $f\text{CO}_2$ difference also substantially increases by $\sim 10 \mu\text{atm}$ in the 2010s (due to summertime warming, Fig. 3a, and higher $f\text{CO}_2$, Fig. 4e).

OA indicators. The BATS and Hydrostation S time-series data allow direct detection of the signal of OA in surface waters of the North Atlantic Ocean. CO_2 dissolved into seawater forms carbonic acid which rapidly dissociates into bicarbonate $[\text{HCO}_3^-]$ and in turn to carbonate $[\text{CO}_3^{2-}]$. Chemical equilibria reactions maintain mean contributions of $[\text{HCO}_3^-]$, $[\text{CO}_3^{2-}]$, and $[\text{CO}_2]_{\text{aq}}$ to DIC at $\sim 88\%$, 11% and 1% , respectively. pH and $[\text{CO}_3^{2-}]$ decrease as seawater absorbs CO_2 , a process termed OA¹⁵. OA is potentially impactful for those organisms that secrete CaCO_3 skeletons, tests or shells and for marine ecosystems where calcification and pH controls on biogeochemical processes are important factors⁴⁷. The typical pH range of surface waters in the 1980s ranged from wintertime highs of ~ 8.2 to summertime lows of ~ 8.08 – 8.10 (Figs. 2c and 5a, b), with the ocean remaining

Table 2 Mean annual values for DO and ocean carbonate chemistry in 1983, 2019, and predictions for mid-century (2050) and end-of-century scenario (2100) with extrapolation of current trends to 2050 and 2100.

	1983	2019	2050	2100
DO	228	212	199	177
DIC	2031	2077	2115	2177
C_{ANT}	25	61	90	138
$f\text{CO}_2$	310	381	439	535
Revelle (β)	8.89	9.39	9.8	10.48
pH	8.138	8.062	8.005	7.91
Ω_{calcite}	6.18	5.5	5.07	4.29
$\Omega_{\text{aragonite}}$	3.86	3.62	3.34	2.88

Note: DO, DIC and C are expressed units of $\mu\text{moles kg}^{-1}$, while $f\text{CO}_2$ is expressed in units of μatm . The Revelle (β) factor, pH, Ω_{calcite} and $\Omega_{\text{aragonite}}$ are dimensionless and do not have units.

mildly alkaline at present (~ 7.98 – 8.05). Concurrent with the changes in DIC, C_{ANT} , $f\text{CO}_2$ and β values, pH has also declined by approximately 0.07 from 1983 to 2020 (Fig. 2c; Table 1a).

The rate of pH change is $\sim 0.0019 \pm 0.0001 \text{ year}^{-1}$, which is a more negative rate than previously reported⁷, and represents a 20% increase in hydrogen ion concentration since 1983). Both Ω_{calcite} and $\Omega_{\text{aragonite}}$ exhibit seasonal changes of ~ 0.3 – 0.5 (Fig. 5c, e), Ω_{calcite} and $\Omega_{\text{aragonite}}$ have decreased by 0.52 and 0.35, the past 40 years (Figs. 2d, e and 5d, f; Table 1a) at rates of $0.014 \pm 0.001 \text{ year}^{-1}$ and $0.009 \pm 0.001 \text{ year}^{-1}$, respectively.

Present ocean carbon chemistry is now outside of 1980s values.

In forty years, seawater CO_2 -carbonate chemistry conditions are now altered beyond the seasonal chemical ranges observed in the 1980s (Table 2). For DIC and $f\text{CO}_2$, the low winter values in 2020 were the high summer values in the 1980s. Similarly, for pH, Ω_{calcite} and $\Omega_{\text{aragonite}}$, the high summer values are at present the low winter values. The statistically significant time-scale of emergence for carbonate parameters in the Sargasso Sea is typically less than 10 years⁴⁸ (Table 2). The modification of seawater CO_2 -carbonate chemistry will continue with future anthropogenic CO_2 emissions. Current chemical trends extended to 2050 indicate the following: Mean DO will be less than $200 \mu\text{moles kg}^{-1}$, mean DIC $> 2100 \mu\text{moles kg}^{-1}$, and $f\text{CO}_2$ and pH approaching $450 \mu\text{atm}$ and 8.000 , respectively (Table 2). If current rates of anthropogenic CO_2 emissions continue until the end of the century, $f\text{CO}_2$ will likely be at least $535 \mu\text{atm}$, C_{ANT} at $> 138 \mu\text{moles kg}^{-1}$ (nearly five times the 1983 concentration) and pH and $\Omega_{\text{aragonite}}$ close to 7.9 and below 3 , respectively (Table 2). This end-of-century scenario is hypothetical given that CO_2 emissions are likely to peak and decline with future climate policy instruments, with the result that ocean carbonate chemistry is not expected to reach such levels.

Discussion

Decadal variability of sargasso sea ocean carbon sinks. The Sargasso Sea observations off Bermuda reveal the substantial decadal variations. As with temperature and salinity, the 1990s and 2000s were periods of modest changes with low rates of oxygen decrease. In the decade of the 1980s and 2010s, the decline of oxygen was about 8% (Table 1a), but with potentially different causes. In the 1980s, warming of $\sim 0.26^\circ\text{C}$ contributed perhaps 5% to lower oxygen concentration. During the 2010s, oxygen concentration decreased by $\sim 8\%$. Approximately, 18% of that change was due to warming and the influence of temperature on oxygen solubility¹⁶. A combination of changes in gyre circulation, wind-driven ventilation and gas exchange may partly

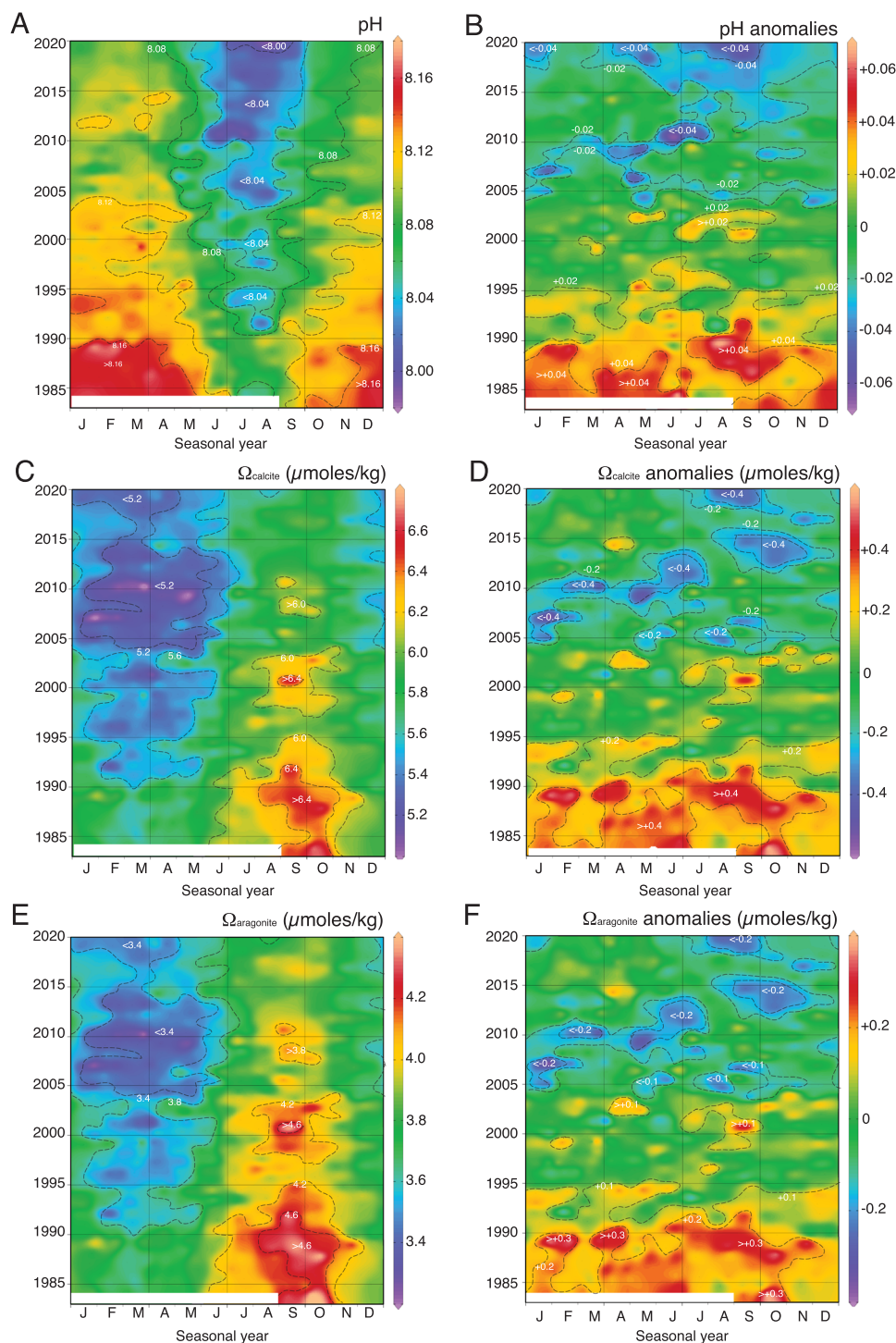


Fig. 5 Seawater CO_2 -carbonate properties at BATS (1988–present) with earlier data (1983–1988) from Hydrostation S. **a** Surface pH. **b** Surface pH anomalies. **c** Surface Ω_{calcite} . **d** Surface Ω_{calcite} anomalies. **e** Surface $\Omega_{\text{aragonite}}$. **f** Surface $\Omega_{\text{aragonite}}$ anomalies. Dashed lines for contours are included.

explain such decade to decade variation on DO loss. The slowdown of Atlantic Meridional Overturning Circulation remains unproven at present, and it is challenging to link loss of oxygen in the subtropical gyre to basin-scale changes. The underlying drivers for loss of oxygen in the subtropical gyre are the scope of future work.

An important additional finding from the BATS–Hydrostation S time-series is that although long-term trends of DIC result from anthropogenic CO_2 uptake, there is a substantial decade to decade variation in changes. In the 1980s; associated with salinification

and an 8% DO decrease, both nDIC and nTA also declined by 11.8 and 7.3 $\mu\text{moles kg}^{-1}$, respectively (Table 1a). Prior studies of nDIC and nTA show that both decrease meridionally by 15–30 $\mu\text{moles kg}^{-1}$, from subpolar waters into the subtropical gyres of the Atlantic, Pacific and Indian Oceans⁴⁹. The salinification event at the end of the 1980s may mark a transition in North Atlantic subtropical gyre circulation and wind-driven ventilation, and perhaps a change to water masses with a more “southern” subtropical gyre physical and biogeochemical character (i.e., warmer, higher salinity and lower oxygen contents). If we assume

Table 3 Comparison of atmospheric CO₂ increase, global ocean carbon sinks and Sargasso Sea carbon sink for recent decades.

		Change over period			
		1980s	1990s	2000s	2010s
Atmospheric CO ₂ increase	[De Vries et al. ⁸ [McKinley et al. ³⁰]	n/a	Higher ⁸	Lower ⁸	n/a
Global ocean carbon sinks	[De Vries et al. ⁸ [McKinley et al. ³⁰]	n/a	Weak (−8 to −16%)	Strong (+35 to +43%)	n/a
Sargasso Sea carbon sink	[This study; C _{ANT}]	n/a	Weak (−17 to −28%)	Strong (+24 to +69%)	n/a
	[C _{ANT} increase]	−1.94 ± 1.23	+0.87 ± 0.50	+1.81 ± 0.53	+1.50 ± 0.58
	[fCO ₂ increase]	+0.19 ± 0.08	+2.06 ± 0.08	+3.21 ± 0.08	+1.84 ± 0.08
	[Winter–summer fCO ₂]	83.5 ± 17.2	87.9 ± 22.7	89.9 ± 14.7	99.9 ± 30.5

Note: the change is given in units of μmoles kg^{−1} year^{−1}, while fCO₂ increase is μatm year^{−1}. The winter–summer fCO₂ difference is given units of μatm. The accumulation of CO₂ in the atmosphere relative to emissions is higher in the 1990s compared to the 2000s⁸. In Table 3, the designation of “weak” and “strong” global ocean uptake is taken from DeVries et al.⁸. The % change in global ocean uptake is taken from DeVries et al.⁸ and McKinley et al.³⁰, and represents the change in ocean sink between the beginning and end of the decade, and the range determined from the various approaches used in each of their studies. The change in C_{ANT} in the anthropogenic CO₂ relative to the long-term increase of −1.03 ± 0.05 μmoles kg^{−1}. Positive increases denoted in % indicate that the C_{ANT} has increased at a higher rate relative to the long-term trend and thus indicate increase ocean uptake of anthropogenic CO₂.

an increase of DIC due to uptake of anthropogenic CO₂ (~9 μmoles kg^{−1}) for the decade of the 1980s in the Sargasso Sea, the nDIC content of a more “southern” water mass would likely have been ~20 μmoles kg^{−1} lower.

Since the 1990s, the three decades have exhibited different DIC, nDIC, TA and nTA changes (Table 3). The largest increase in nDIC occurred in the 2000s. The finding of such significant decade to decade variation in seawater CO₂–carbonate chemistry has implications for determination of trends from shorter duration observations which likely have interannual to decadal change. Trend analysis of seawater CO₂–carbonate chemistry, as with other ocean physical and biogeochemical properties, thus requires long-term data.

Recently, DeVries et al.⁸ and McKinley et al.³⁰ reported substantial decadal variation in the global ocean carbon sink with climate variability a highly influential driver of ocean carbon uptake. In the 1990s, enhanced accumulation of CO₂ in the atmosphere relative to CO₂ emissions⁸ was accompanied by a period of reduced global ocean carbon uptake (~−8 to −28%^{8,30} from beginning to end of the decade; Table 3). In contrast, in the 2000s, the atmospheric CO₂ growth rate slowed relative to higher anthropogenic CO₂ emissions⁸ while the global ocean carbon sink increased (~+24 to +69% from beginning to end of the decade; Table 3), with indications of a strengthening terrestrial carbon sink⁸. The Sargasso Sea C_{ANT} data adds regional weight to these findings if we use C_{ANT} changes as a proxy for ocean carbon uptake. In the 1990s, the rate of C_{ANT} increase was lower by 16% (i.e., +0.87 ± 0.50 μmoles kg^{−1} year^{−1}; Table 3), compared to the 1983–2020 change of +1.03 μmoles kg^{−1} year^{−1}, indicative of lower anthropogenic CO₂ accumulation (and ocean uptake) during the decade. In contrast, in the 2000s the rate of C_{ANT} increase was higher by 74% (i.e., +1.81 ± 0.53 μmoles kg^{−1} year^{−1}; Table 3) were indicative of doubling of CO₂ accumulation (and ocean uptake) during the decade. Higher C_{ANT} accumulation in the Sargasso Sea (i.e., +1.50 ± 0.58 μmoles kg^{−1} year^{−1}; Table 3) may reflect the continuation of higher ocean carbon uptake in the 2010s from the 2000s.

The decade to decade variability in C_{ANT} also covaries with rates of fCO₂ change and winter–summer fCO₂ difference. For example, the highest C_{ANT} increase occurs in the 2000s along with higher fCO₂ increases and winter–summer fCO₂ difference (Table 3). This finding adds regional evidence to the study of Landschützer et al.⁴⁶ that winter–summer fCO₂ difference is increases. Following the methods of Landschützer et al.⁴⁶ that

derive the thermal and non-thermal components that drive the differential winter and summer changes, the winter and summer DIC changes at BATS have a near identical influence on fCO₂. The thermal changes have a one-third and two thirds influence on fCO₂ in winter and summer, respectively. The non-thermal changes of fCO₂ are driven by an increase in nTA (0.29 μmoles kg^{−1} year^{−1}) in winter and decrease in nTA in summer (~−0.13 μmoles kg^{−1} year^{−1}). At BATS, the differential changes in winter and summer temperature, and increase/decrease in alkalinity drive the 4–8 μatm per decade increase in winter–summer fCO₂ difference over the past forty years. In contrast, Landschützer et al.⁴⁶ reported a change of 1.5 μatm per decade, but critically, ignored the effect of alkalinity changes in their analysis. Here, we demonstrate that observed changes in alkalinity do make a substantial difference for evaluating changes in winter–summer fCO₂ difference over time and the need to incorporate TA for global syntheses⁴⁶.

Nearly, 40 years of data at BATS–hydrostation S also indicate that the ocean is not uniformly changing over time, and there is a substantial decade to decade variation in CO₂–carbonate chemistry. And that the ocean carbon sink is not stable over recent time and varies from decade to decade. If the coupling of atmospheric CO₂ accumulation, ocean carbon sink and climate variability^{8,30} is proven, the Sargasso Sea appears to act in concert with global decade to decade changes in ocean carbon sinks.

In summary, these data reveal recent warming in the last decade, associated with salinification and substantial deoxygenation. These physical changes were accompanied with significant increases of DIC and fCO₂, and 20–40% decreases in pH, Ω_{calcite} and Ω_{aragonite}, respectively, since the 1980s. Apparent reduced ocean carbon uptake in the decades of the 1990s and higher CO₂ uptake 2000s/2010s in the Sargasso Sea agree with the findings of decadal variability in “weaker”/“stronger” global ocean carbon uptake^{8,30} and points to reduced ocean uptake in the Sargasso Sea in the 2010s. Such data on ocean physics and chemistry provide critically needed and unparalleled observational tests of coupled ocean–atmosphere models and future refinement of drivers for a model-data decade to decade assessment of the global carbon cycle.

Methods

Sampling methods. Ocean sampling at hydrostation S and BATS started in 1954 and 1988, respectively. Hydrostation S began with the pioneering efforts of Hank Stommel (Woods Hole Oceanographic Institution) and colleagues⁵⁰ at a site approximately 26 km southeast of Bermuda (32° 10'N, 64° 30'W, Fig. 1). The first

water-column sampling occurred on the 7th June 1954 from the 61' R.V. *Panularis* with more than 1381 cruises conducted up to the present time from the R.V. *Panularis II* (1967–1983), R.V. *Weatherbird* (1983–1989); R.V. *Weatherbird II* (1989–2006) and R.V. *Atlantic Explorer* (2006–present).

Since the arrival of the R.V. *Weatherbird*, Hydrostation S has been occupied at near biweekly intervals, with multiple CTD–hydrocasts through the water-column to ~2600 m, and to 4500 m at BATS (more than 450 cruises to the site). Nansen bottles were used for water sampling at first, then 5 L Niskin samplers until October 1988. Thereafter sampling has been conducted with a Seabird 9/11 CTD equipped with 12 L Niskin and Ocean Test Equipment (OTE) samplers.

Water sampling, temperature and CTD measurements. The sampling format has remained substantially consistent for the past 65 years, but with the introduction of CTD–hydrocast sampling in October 1988. From 1954 to 1988, reversing mercury thermometers were used for measurements of temperature until replaced by CTD measurements using a Sea-Bird 9/11 system. Bottle samples before 2012 were taken down to 2600 m at Hydrostation S. Since the addition of reliable altimeters to the CTD package, sampling was extended to full ocean depth at both the hydrostation S (~3400 m) and BATS (~4500 m) sites. Numerous sensor configurations have been used on the CTD package (e.g., dual temperature, dual conductivity, dual DO sensors, transmissometer, fluorometer, PAR and altimeter). In contrast, the CTD sampling system has predominately been a Seabird 24-place rosette using 12 L Ocean Test bottles. Before profiling, the CTD is allowed to stabilise at 10 m and once stable, the CTD returns to the surface to start the profile with typical descent rates of 0.5–1.0 m s⁻¹, depending on weather conditions. Water samples are collected on the upcast, whereby the OTE bottles are closed at the target depth after a waiting period of 45 s. The CTD is held at the target depth for another 10 s to allow the SBE35-RT sensor to take an 8 s average. The CTD continues with the upcast at an ascent rate of 0.7–1.0 m s⁻¹. Temperature, conductivity and DO sensors are routinely returned to SeaBird every 6–9 months for routine calibration. The differences between primary and secondary temperature sensors in the deep ocean at BATS (>3000 m) were 0.002–0.006 °C regardless of time since most recent factory calibration.

Determination of salinity. Salinity samples are typically taken from the OTE bottles at all depths. These samples are collected immediately following DO and CO₂ sampling. Samples are taken in 125–250 ml borosilicate glass bottles (Ocean Scientific, UK) that use plastic thimbles to form a better seal. The sample remaining from the previous use is left in the bottles between cruises to prevent salt crystal buildup due to evaporation. When drawing a new sample, the old sample is first discarded over the sampling spigot, and the bottle is rinsed three times with water from the new sample. The bottle is then filled to the shoulder with the sample, and the thimble inserted into the container. The neck of the bottle and the inside of the cap are dried, then the thimble is inserted, and the cap is replaced and firmly tightened. These samples are stored in a temperature-controlled laboratory for later analysis (typically within 1–2 weeks of their collection).

Salinity measurements have been made with a Guildline salinometer at BIOS from 1981 to present (calibrated with IAPSO standard water⁵¹) for both BATS and Hydrostation S samples. At present, samples for salinity are analysed on a Guildline Autosal 8400B laboratory salinometer using the manufacturer's recommended techniques. All readings (10 s average) are taken using the Ocean Scientific interface box and PC software. The salinometer drift during and between successive runs tends to be zero (room temperature carefully controlled and monitored). Bottle salinities are used to calibrate the profiling CTD SBE-04 sensors, and additionally, they are also compared with the downcast CTD profiles to search for possible outliers. Deep-water samples (>2000 m) are replicated for precision estimates (typically < 0.002 salinity units).

Determination of DO. Early samples for DO were analysed by manual endpoint detection (1954–1988) though more recently, automated titration systems have been used (1988–present). On each hydrostation S and BATS cruise, the first samples from the rosette are taken for DO, immediately following the opening of the valve on the OTE bottle and confirmation that the OTE bottle has not leaked. The sample flasks used for this measurement are Pyrex iodine determination flasks of 140 ml nominal capacity with ground glass barrel stoppers. The precise volume of each stopper/bottle pair is determined gravimetrically.

Seawater is carefully sampled to avoid the introduction of air bubbles with a minimum of five bottle volumes overflow. The samples are then immediately fixed with 1 ml of manganous chloride and 1 ml of sodium iodide/sodium hydroxide solution. With the stopper in place, and after vigorously shaking, the necks of the flasks are sealed with surface seawater. The samples are stored upright in the dark at 21–24 °C. The temperature of the water from the OTE bottle is measured to allow the conversion to units of mass.

The samples are analysed after 6–8 h, based on the method proposed by Winkler (1888; modified by Strickland and Parsons 1968⁵²). For this analysis, BIOS/BATS currently uses an automated temperature-controlled titration system (developed by SIO) using an ultra-violet light endpoint detection system with a

Metrohm 665 Dosimat burette for precise delivery of the sodium thiosulfate (0.18 M; reagent grade).

Before running the samples, the precise concentration of the thiosulfate and the chemical blanks are determined. Typically, six to eight standards are run for determination of a mean value. Blanks are prepared in the same manner as the standards. However, they are based on the 1 ml standard addition of KIO₃. The samples are run following the sodium thiosulfate normality determination. The water in the neck of the sample is carefully removed, taking care to avoid disturbing the precipitate. The stopper is then removed, and 1 ml of 50% sulphuric acid added slowly, then a stir bar, and the sample titrated immediately. A 30% triplication is performed for estimates of precision (typically < 0.4 μmoles kg⁻¹; ~0.1%).

Determination of DIC and TA. The description of seawater CO₂–carbonate chemistry used in this paper follows long-established knowledge of the marine carbon cycle^{53–55}. In this paper, DIC and TA were directly determined, and the chemical description of DIC is as follows^{53,54}:

$$\text{DIC} = [\text{CO}_2^*] + [\text{HCO}_3^-] + [\text{CO}_3^{2-}] \quad [\text{equation S1}]$$

The term [CO₂*] denotes the summed concentration of dissolved H₂CO₃ and CO₂. The chemical description of alkalinity of seawater (TA) is defined^{53,54} as:

$$\text{TA} = [\text{HCO}_3^-] + 2[\text{CO}_3^{2-}] + [\text{B}(\text{OH})_4^-] + [\text{OH}^-] + [\text{HPO}_4^{2-}] + 2[\text{PO}_4^{3-}] + [\text{SiO}(\text{OH})_3^-] + [\text{HS}^-] + [\text{NH}_3] + \text{minor species} - [\text{H}^+] - [\text{HSO}_4^-] - [\text{HF}] - [\text{H}_3\text{PO}_4] - \text{minor species} \quad [\text{equation S2}]$$

In this second chemical definition, the summed concentrations of [HCO₃⁻] + 2 [CO₃²⁻] + [B(OH)₄⁻] represent the major chemical components of TA and the proton/charge balance of seawater. Other chemical contributors to alkalinity are typically considered minor constituents in seawater^{54–56}. Given DIC and TA are both expressed as μmoles kg⁻¹ following long-established guidelines (Dickson et al., 2007⁵⁴ and references therein).

Initially, at hydrostation S, samples for DIC and TA were collected into 11 Pyrex bottles with a change to 500 ml bottles in the early 1990s, poisoned with Hg₂Cl, sealed with ground glass stoppers and then shipped to Scripps Institution of Oceanography (SIO) for analysis^{56,57} (1983–1988). Storage time for samples before analysis ranged from a few months to several years. Similar sampling protocols were established at BIOS for sampling at the BATS^{36,58,59}, but in the early 2000s, smaller Pyrex bottles (350 ml) were used. Samples for DIC and TA have been typically analysed within a few months of collection at BIOS since 1988.

The sampling frequency of the combined dataset from hydrostation S and BATS was not uniform in time. In the 1980s, samples were collected 9–12 times a year, while since 1992, sampling increased to 14–15 times a year^{7,40}. The increase in sampling since the middle 1990s was due to supplemental BATS bloom cruises (1–4 in number) conducted during the January to April period in addition to BATS core cruises. The increase in sampling frequency over time weights the time-series to springtime when determining non-seasonally aliased trend^{7,40}.

Potentiometric titration methods were also used for determination of TA at BIOS⁵⁸. At the beginning of the 1990s, a manual alkalinity titrator was used for determination of TA at BIOS. This was replaced by an automated VINDTA 2S (Versatile Instrument for the Determination of Titration Alkalinity) in the early 2000s⁴⁰. For both manual and automated TA systems, 15–20 titration points past the carbonic acid endpoint were determined for each sample, with TA computed from these titration data using nonlinear least-squares methods⁵⁴. Surface samples of Sargasso Sea water were also analysed each day before sample analyses, and certified reference materials (CRMs) were used routinely to calibrate the TA measurements.

At BIOS, DIC was determined using coulometric methods with a SOMMA system^{7,40,54}. During the first 2 years of sampling, DIC samples were analysed at WHOI (e.g., BATS cruise 1–21), and subsequently at BIOS. DIC measurements were calibrated with known volumes of pure CO₂ gas while CRM⁵⁵ were routinely analysed each day of analysis from 1991. Hydrostation S samples were analysed for DIC at SIO from 1983 to 1988 using manometric methods⁵⁶. Potentiometric titration methods were used for determination of TA⁵⁶ at SIO, and after that at BIOS. No significant bias has been noted between these earlier methods (also analysed by A.G. Dickson to present)⁴⁰. Analytical precision for DIC and TA at BIOS was typically <0.03% and <0.05%, respectively for within the bottle and between bottle replicate analyses of more than 5,000 samples⁴⁰. Analytical accuracy for both DIC and TA using CRM analyses is better than 0.1%.

Seawater CO₂–carbonate chemistry computations and errors. The suite of seawater CO₂–carbonate parameters (i.e., [CO₂], [H₂CO₃], [HCO₃⁻], [CO₃²⁻], [H⁺]) for each sample can be determined from the measurement of any two parameters (i.e., DIC, TA, *f*CO₂, *p*CO₂ and pH), along with temperature and salinity^{33,54,55}. Seawater *f*CO₂ (fugacity of CO₂) is used here rather than the partial pressure of CO₂ (i.e., *p*CO₂; note that this parameter was also computed in data files), and with units of μatm. pH, Ω_{calcite}, Ω_{aragonite} in seawater are dimensionless and pH expressed on the total scale⁵⁴. All parameters are computed at in situ temperature and salinity.

Here, seawater *f*CO₂, pH, [CO₃²⁻], mineral saturation states for Ω_{calcite} and Ω_{aragonite}, the Revelle factor (*β*) were computed from DIC, TA, temperature and salinity data using the programme CO₂calc⁶⁰. Carbonic acid dissociation constants (i.e., *p*K₁ and *p*K₂)⁶¹, as refit by Dickson and Millero (1987)⁶² were used for the computation, as well as dissociation constants for KHSO₄⁻⁶³, borate⁶⁴.

The computation error estimates for calculation of $p\text{CO}_2$, pH, Ω_{calcite} and $\Omega_{\text{aragonite}}$ were undertaken using standard procedures for the propagation of uncertainty⁵⁴. Given the analytical uncertainty of the two seawater carbonate chemistry parameters (i.e., DIC and TA; $\pm 1 \mu\text{moles kg}^{-1}$) from replicate measurements of CRM's, the error of the computation is estimated at $3 \mu\text{atm}$, 0.003, 0.018, and 0.012, for $p\text{CO}_2$, pH, and Ω_{calcite} and $\Omega_{\text{aragonite}}$ for surface waters, respectively⁶⁵.

Use of the tracer TrOCA. TrOCA and TA is a method first proposed by Touratier and Goyet (2004a, b)^{66,67}, and updated by Touratier et al. (2007)²⁹ to quantify anthropogenic carbon (i.e., C_{ANT}) in seawater. The quasi-conservative tracer, TrOCA in water masses accounts for changes in carbonate chemistry due to biological influences (e.g., remineralization of organic matter) and abiotic processes (e.g., dissolution of CaCO_3), and as such, is also useful for following ocean chemistry changes in the surface ocean. Here, the formulation proposed by Touratier et al. (2007²⁹, their equation 11) is used as a water tracer:

$$C_{\text{ANT}} = ([\text{O}_2] + 1.279 \times (\text{DIC} - \text{TA}/2) - \text{EXP}(7.511 - 1.087 \times 10^{-2} \times \Phi - 7.81 \times 10^5 / \text{TA}^2)) / 1.279 \quad [\text{equation S3}]$$

Φ is the potential temperature. The uncertainty of the estimate is $\sim 6 \mu\text{moles kg}^{-1}$ (29). The C_{ANT} estimates determined using the TrOCA method for the mixed layer are modified by euphotic zone biological activity and air-sea gas exchange²³, but, over time, provide additional evidence for changes in ocean chemistry.

Trend analyses. Trend analyses were conducted of the time-series of surface temperature and salinity, seawater carbonate chemistry (DIC, TA, $p\text{CO}_2$ and Revelle factor) and OA indicators (pH, $[\text{CO}_3^{2-}]$, Ω_{calcite} and $\Omega_{\text{aragonite}}$). Here, trend analysis of salinity normalised DIC (nDIC), and TA (nTA) data were also made to account for local evaporation and precipitation changes. These data were normalised to salinity of 36.6, as this represents the mean salinity observed at the BATS site^{37,40}. Trend analysis was performed with observed data, and seasonally detrended data is given in Table 1) Statistics generated from least-squares regression approaches were slope, coefficients, error, multiple r , r^2 , p value and n . Trends with p values greater than 0.01 were deemed statistically not significant at the 99% confidence level.

Trend analysis for the period from 1983 to 2020, and for each decade (i.e., the 1980s, 1990s, 2000s, 2010s) was determined using linear regression methods with physical and biogeochemical data are the dependent variable and time as independent, and p values < 0.05 used to determine statistical significance of trend. Trend analyses with observed data exhibit seasonal aliasing due to sampling weighting to spring conditions^{12,13}. To account for seasonal weighting, the data were also seasonally detrended⁶⁸. Seasonal detrending of the BATS/Hydrostation S data was accomplished by binning data into the appropriate month, with mean values calculated from two or more cruises conducted within a representative month each year. This provides a uniform time step of approximately 1 month (i.e., 365 or 366 days/12) throughout the time-series, thereby removing any potential seasonal weighting especially to springtime conditions. Secondly, a mean and standard deviation is then determined each month for the 1983–2020, and anomalies computed from monthly data minus mean values. Trends and regression statistics anomaly data determined from seasonally detrended data are given in Table 1. Removing any potentially seasonal weighting allowed trend analysis of data that had any non-temporal uniformity reduced as much as possible.

BATS and Hydrostation S data. The BATS and Hydrostation S sites have been sampled on a monthly and twice-monthly basis, respectively, since October 1988. The data is publicly and permanently available at <http://bats.bios.edu/data/> with a transition to BCO-DMO (Biological and Chemical Oceanography Data Management Office; Woods Hole, USA). The data used are primarily fully processed CTD profiles and bottle data with both dataset available to the end of 2019. The surface data is also available as Supplemental Data accompanying this paper. Information regarding CTD processing and QC routines are detailed at http://bats.bios.edu/wp-content/uploads/2017/07/report_methods.pdf.

Data availability

The BATS data used in this paper are available as a supplemental data file accompanying the paper. All BATS and Hydrostation S bottle and CTD data are freely available up to the end of 2019 at <http://www.bios.edu/research/projects/bats/>. Information regarding CTD processing and QC routines are detailed at http://bats.bios.edu/wp-content/uploads/2017/07/report_methods.pdf. The CTD and bottle data used here have been separately uploaded to BCO-DMO (Biological & Chemical Oceanography Data Management Office) data repository (<https://www.bco-dmo.org/>) and will be freely available with a doi to CTD and bottle data. Please note that observed data are given (e.g., temperature, salinity, dissolved oxygen, DIC and TA) but not calculated parameters in accordance with BCO-DMO policies.

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Author contributions

N.R.B. and R.J.J. are principal investigators for the BATS/Hydrostation S projects and coordinate the sampling and analysis of data. Both authors contributed to designing the study and composing the analyses.

Competing interests

The authors declare no competing interests.

Additional information

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