Surface ocean pH variations since 1689 CE and recent ocean acidification in the tropical South Pacific

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Increasing atmospheric CO2 from man-made climate change is reducing surface ocean pH. Due to limited instrumental measurements and historical pH records in the world’s oceans, seawater pH variability at the decadal and centennial scale remains largely unknown and requires documentation. Here we present evidence of striking secular trends of decreasing pH since the late nineteenth century with pronounced interannual to decadal-interdecadal pH variability in the South Pacific Ocean from 1689 to 2011 CE. High-amplitude oceanic pH changes, likely related to atmospheric CO2 uptake and seawater dissolved inorganic carbon fluctuations, reveal a coupled relationship to sea surface temperature variations and highlight the marked influence of El Niño/Southern Oscillation and Interdecadal Pacific Oscillation. We suggest changing surface winds strength and zonal advection processes as the main drivers responsible for regional pH variability up to 1881 CE, followed by the prominent role of anthropogenic CO2 in accelerating the process of ocean acidification.
The concentration of atmospheric CO$_2$ is increasing at unrelenting rates in response to human activities in fossil fuel combustion and land-use practices\textsuperscript{1}. At present, the oceans take up more than 41% of anthropogenic CO$_2$ emissions\textsuperscript{2}, inducing the acidification of the ocean surface with major implications in marine carbonate chemistry and biological ecosystems\textsuperscript{3}. Many modelling and experimental studies have indeed shown clear trends of shallow water acidification in lockstep with increasing atmospheric CO$_2$\textsuperscript{4}. One of the major consequences is the decrease of carbonate saturation state that is crucial for calcifying organisms such as scleractinian corals to precipitate their aragonite skeletons\textsuperscript{5}. These critical threats of ocean acidification (OA) on such marine organisms and ecosystems have been documented on the Great Barrier Reef (GBR) in Australia that witnessed a 14% decrease in coral calcification since 1990\textsuperscript{6}. Future projections of coral calcification rates suggest a possible additional decrease by up to 30% in the twenty-first century\textsuperscript{7}. Moreover, end-of-century emission scenarios predicted by model simulations indicate unprecedented decrease by up to 0.4 pH units\textsuperscript{8} compared to the changes of only 0.2 pH units over glacial-interglacial cycles\textsuperscript{9}.

Presently, accurate measurements of pH in the oceans are near non-existent aside from selected ship track measurements\textsuperscript{10} and limited station-based instrumental time series available over the past few decades\textsuperscript{11}. The pH records that span only the most recent decades are far too short to precisely monitor the longer-term evolution of OA and to quantify its impacts on tropical ecosystems. To study the natural variability in oceanic CO$_2$ uptake and carbonate chemistry changes, proxy records in modern marine archives such as corals provide a unique opportunity to go further back in time at a high resolution on the centennial scale. Additionally, it is crucial to investigate pH changes with links to ocean CO$_2$ uptake as well as the relationship to sea surface temperature and forcing mechanisms at various timescales.

A method to document sub-seasonal to annual pH changes is based on the $\delta^{11}$B signature in scleractinian coral skeleton. Coral-based reconstructions of ocean pH suggest notable recent acidification\textsuperscript{12,13} and decadal variability\textsuperscript{12,14}. However, due to spatiotemporal constrains, the currently available coral-based pH reconstructions predominantly focus on the GBR and the marginal South China Sea\textsuperscript{15–19}, whereas longer timescale pH changes of the open surface ocean from the Pacific Ocean remain poorly documented\textsuperscript{17}.

In the south-western Pacific, large-scale atmospheric circulation features and the seasonally varying latitudinal position of the tropical convergence zones (Intertropical and South Pacific Convergence Zones (ITCZ and SPCZ) extending from the western Pacific warm pool (WPWP; Fig. 1) are primarily controlling climate variability. The large-scale atmospheric circulation features include the seasonal trade wind regime and strength in addition to the Hadley and Walker Circulations. The region is also intimately pulsed at the interannual timescale by the Pacific basin-wide phenomena of El Niño/Southern Oscillation (ENSO) and at decadal timescale by the Interdecadal Pacific Oscillation (IPO) both in relation to the modulation of Pacific sea surface temperature (SST). More specifically, New Caledonia and the surrounding region of the south-western Pacific are of great interest for investigation because more than 40% of the global anthropogenic CO$_2$ inventory is found in the region between 14°S and 50°S\textsuperscript{20} and about 60% of the total oceanic anthropogenic CO$_2$ is incorporated in the Southern Hemisphere oceans\textsuperscript{4} (Fig. 1). Moreover, prior geochemical studies with annually banded scleractinian corals have demonstrated its advantages as reliable recorders of climate change from this region\textsuperscript{19}.

Here we present 323 years of annually resolved $\delta^{18}$O, $\delta^{13}$C and $\delta^{11}$B results from a modern slow-growing massive scleractinian coral, Diploastrea heliopora, collected from the open-ocean island of New Caledonia in the South Pacific Ocean (22.21°S, 166.15°E; Fig. 1 and Supplementary Fig. 1). In this study, we provide evidence of the uninterrupted OA with secular decreasing pH trend in the South Pacific since the Industrial Revolution. We show that pronounced interannual to decadal-interdecadal pH variability has been regularly occurring in the South Pacific since 1689 CE. The high-amplitude oceanic pH changes over these timescales are likely related to atmospheric CO$_2$ uptake because of significant coherence to coral $\delta^{13}$C signature that tracked the changes in oceanic dissolved inorganic carbon (DIC) fluctuations. In addition, the timing of reconstructed pH variability in the South Pacific is coupled to prominent sea surface temperature

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**Fig. 1** Research location. Our research location at New Caledonia (22°21′47″ S, 166°15′29″ E; Star; Supplementary Fig. 1) is of high importance because it is located in the zone of transition from low to high amounts of anthropogenic CO$_2$ uptake\textsuperscript{23}. The western Pacific warm pool (WPWP) with a permanent sea surface temperature above 28 °C influences the climate of New Caledonia driving the steady convection in this region corresponding to the average precipitation axes of the Intertropical Convergence Zone (ITCZ) and South Pacific Convergence Zone (SPCZ). Map is produced using Ocean Data View ver. 4.7.4 (ref. \textsuperscript{74}) (http://odv.awi.de) and modified manually.
variations modulated by the phases of ENSO and IPO. We suggest changing surface wind strengths and zonal advection processes across the Pacific Ocean as the main drivers responsible for South Pacific pH variability that is superimposed by the prominent role of anthropogenic CO2 increase in accelerating the process of OA.

**Results**

New Caledonia coral skeletal preservation and chronology. The studied *D. heliopora* coral was collected in March 2015 at the Fause Passe de Uitoé in New Caledonia (22°17′152 S, 166°10′992 E; Fig. 1 and Supplementary Fig. 1). The methodology applied for **230**Th/U-dating, δ^18^O, δ^13^C and δ^11^B isotopes measurements, chronology, climate reconstruction and statistical analyses, are described in detail in the Methods. Examination of coral skeletal material by X-radiograph imaging (Supplementary Fig. 2), powder X-ray diffraction (XRD) analysis (100% aragonite) and scanning electron microscopy (SEM; Supplementary Fig. 3), a prerequisite for coral-based palaeoclimatic work^1^, attested pristine preservation. The chronology of our coral is determined by

| Coral sample | **230**U (µg/g) | **232**Th (ng/g) | δ^234^U (measured)^a | **230**Th/**238**U (activity)^b | δ^234^U (initial)^c | Age corrected (before 2016)^d | Age (CE) |
|--------------|-----------------|-----------------|----------------------|-------------------------------|-------------------|--------------------------|---------|
| 1            | 3.007 ± 0.004   | 0.054 ± 0.0001  | 146.93 ± 0.68        | 0.0025 ± 0.00002              | 413.24 ± 3.71     | 0.231 ± 0.004            | 1785 ± 4.0 |
| 2            | 2.884 ± 0.001   | 0.080 ± 0.0001  | 146.77 ± 0.93        | 0.0034 ± 0.00004              | 366.86 ± 4.96     | 0.313 ± 0.008            | 1704 ± 7.5 |

Ages are presented as both age corrected (years before 2016) and age (CE) with the corresponding 2σ-uncertainty

δ^234^U = ([234U/238U]activity − 1) × 1000

^a Decay constants are 9.1705 × 10^−6 yr^−1^ for **230**Th, 2.8221 × 10^−6 yr^−1^ for **234**U and 1.55125 × 10^−10 yr^−1^ for **238**U

^b δ^234^U initial was calculated based on **230**Th age (t), i.e., δ^234^U_Initial = δ^234^U_measured × e^(λ^234^U × t), and t is corrected age

^d Age corrections were calculated using **230**Th/**232**Th activity ratio of 10 ± 3

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**Fig. 2** *Diploastrea heliopora* coral proxy records and δ^11^B-pH reconstruction. New Caledonia *D. heliopora* annually resolved records over the period 1689–2011 CE from precisely dated **230**Th/U-age (see Methods). a Coral δ^11^B signature (left y axis) with estimated seawater pH (pHSW) on the right y axis using the δ^11^BSW = 39.61‰ (ref. 67) and isotopic fractionation factor (α[B3-B4]) of 1.0272 (ref. 35) following the conversion equations of refs. 34 and 39 (see Methods for Eqs. (1) and (2)). b Coral δ^13^C (‰ VPDB) and c δ^18^O (‰ VPDB) ratios. To highlight the interannual variability, we applied a 7-year smoothing to the records (bold lines). The long-term secular trends of each time series (dashed lines) indicate the long-term changes and analysed by Bayesian Change Point Analysis algorithm^72, revealing significant times of emergence (see Methods)
δ¹⁸O calibration and SST reconstruction. We generated annually resolved records of coral δ¹⁸O, δ¹³C and δ¹⁸O values for a total of 323 continuous years over the Common Era (1689–2011 CE; Fig. 2 and Supplementary Fig. 5). On seasonal to sub-seasonal timescales, physiological processes such as photosynthesis, respiration and heterotrophy influenced by metabolic fractionation clearly affected coral δ¹³C levels32–34. However, over longer-term centennial timescales, we observed δ¹³C depletion towards more negative values that can be attributed to the uptake of ¹³C. Suez effect35 and may be used as an indicator of changes in ocean carbon content or the DIC in seawater24 and anthropogenic CO₂ uptake25. Coral δ¹⁸O signature is one of the most frequently used coral climate proxies because it is influenced by the combination of temperature-dependent isotope fractionation and the δ¹⁸O values of seawater (δ¹⁸Osw)26. Such value is directly related to sea surface salinity (SSS) responding to evaporation and freshwater input. Previous studies from the Pacific have demonstrated that coral δ¹⁸O variations over multiple timescales represent changes of SST, SSS, precipitation, or a mixture of those parameters depending on the environmental setting19,27,28.

The calibration of our D. heliopora coral δ¹⁸O ratio to SST from New Caledonia was completed using annual gridded SST from the National Oceanic and Atmospheric Administration (NOAA), National Centers for Environmental Information (NCEI) and Optimum Interpolation SST version 2 (OI-SST)29. A 4-grid averaged OI-SST (Supplementary Table 1) was chosen because coral δ¹⁸O-based SST is not only a local archive, but is in fact encapsulating a larger regional signal19,27. The least squares linear regression calibration of annually averaged δ¹⁸O to the 4-grid OI-SST is significant (p < 0.001). Even though coral δ¹⁸O record is a combination of both SST and the δ¹⁸Osw, the coral δ¹⁸O-SST sensitivity of −0.24‰ °C⁻¹ is in agreement to the typically described relationship across the Pacific even with the use of annual resolution sampling (Supplementary Fig. 6a). D. heliopora species-specific calibrations using in situ and gridded or satellite-based SST indicated a SST-sensitivity relationship ranging from −0.16 to −0.22‰ per °C for sub-seasonal calibrations30–32, which is similar to our finding.

We cross-verified the accuracy of our calibrated δ¹⁸O-SST record and chronology across the overlapping period (1860–2011 CE) to the NOAA NCEI Extended Reconstructed SST version 4 (ERSST)33. A 4-grid regional annual average SST for ERSST (Supplementary Table 1) was compiled and compared as a moving average, which increased the signal-to-noise ratio to track the prominent interannual to decadal-interdecadal changes. The calculated running correlation between our coral δ¹⁸O-SST and ERSST yielded significant (p < 0.01) and high correlation coefficients between the most recent period of the record with the most accurate instrumental SST period (1960–2011 CE; Supplementary Fig. 6b). This considerable coherence in the late twentieth century (Supplementary Fig. 6b) corresponds to the period when the number of SST observations is higher with highest degree of confidence (Supplementary Fig. 7). The misfit between coral δ¹⁸O-SST record and the gridded SST is greater in the middle of the twentieth century, a period when the number of International Comprehensive Ocean-Atmosphere Data Set (ICOADS; http://icoads.noaa.gov/products.html) observations in the gridded SST timeseries was low. The observed lack of significant coherence over the mid-twentieth century in our analysis and verification may be reflecting the inadequacies of the gridded instrumental SST product. Nonetheless, the result over the verification period provides confidence in our chronology and the potential of the coral δ¹⁸O-SST proxy although it is likely that the contribution of δ¹⁸Osw is of significance. Thus, our δ¹⁸O record from New Caledonia suggests a strong secular warming and or freshening trend consistent with many coral records from the western Pacific with similar timing for the onset of warming in the mid-nineteenth century19 (Fig. 2).

Multicentennial coral δ¹¹B pH variability. To reconstruct seawater carbonate chemistry changes and potential modern ocean acidification from anthropogenic CO₂ uptake at the sea surface, we utilised the δ¹¹B composition in our coral. In seawater, the relative abundance of the two aQueous boron species (boric acid and borate) as well as their isotopic composition are pH dependent34 with a constant fractionation factor between the two aqueous boron species35. On the assumption that marine carbonates reflect the isotopic composition of seawater borate26, the δ¹¹B signature of coral aragonite skeleton has been shown to record seawater pH12,27. Possible intra-colony differences of δ¹¹B signature of coral aragonite skeleton were taken into account as studies of sub-seasonally resolved D. heliopora coral δ¹⁸O and Sr/Ca ratios have shown offsets between skeletal materials (columellar and septal) and different coral polyps30–32. However, at the annual resolution, our test across multiple coral polyps revealed that skeletal material differences are inconsequential for δ¹¹B ratios (Supplementary Fig. 8). The three replicated sections of 20-year periods (1935–1955, 1856–1876 and 1768–1788 CE) were reproducible with negligible intra-colony offset that were within the analytical errors between the different polyps and sampling paths (Supplementary Fig. 8). The large amount of coral skeletal material used for δ¹¹B analysis (~50 mg) compared to traditional δ¹⁸O and δ¹³C analyses makes it statistically likely that the sample comprises all different skeletal structures (columellar and septal) and is thus representative of the whole coral skeleton. Our method demonstrates the reliability and reproducibility of annual δ¹¹B ratio results from a new south Pacific coral species (D. heliopora) with larger skeletal structure than the commonly used Porites coral genus.

Careful considerations were taken to convert our coral δ¹¹B signature to coral internal pH as studies indicated differences between empirically and theoretically determined fractionation factors for isotope exchange between boric acid and borate in seawater35. Furthermore, results based on a variety of coral species across temperate and tropical regions indicate consistent difference between coral internal calcification pH by approximately one-half of ambient seawater pH38,39. However, recent results also indicated possible modifications of coral internal pH due to external factors and species-specific processes30,41. Thus, we estimated the changes in seawater pH at New Caledonia using a moderate general calibration conversion39 (Fig. 2). To be conservative with our conversion, a calibration based on a similar massive coral species (Porites spp.) to the one used in this study (D. heliopora) was chosen allowing for the establishment of biological and seawater pH linkage to climate-driven patterns. We focused our discussion on the relative pH changes instead of the absolute pH values because rigorous palaeo-pH reconstructions from coral δ¹¹B signature require species-specific calibrations with robust quantification of physiological processes (vital effects). Nevertheless, independently from a possible offset in
our reconstructed pH using this calibration conversion, our observed trend and interannual to decadal variability from the measurements along the core will remain.

Between the independent proxies of \( \delta^{13}B \), \( \delta^{13}C \) and \( \delta^{18}O \), significant relationships were found. The significant correlation between \( \delta^{13}B \) and \( \delta^{13}C \) (\( R = 0.61; p < 0.01, n = 319 \); Supplementary Fig. 9a) indicates a strong linear relationship indirectly measuring the amount of atmospheric CO\(_2\) that is forced by fossil fuel emissions and subsequently taken up by the ocean as reflected in the changing oceanic DIC content. Moreover, the correlation between \( \delta^{13}B \) and \( \delta^{18}O \) is also significant (\( R = 0.42, p < 0.01, n = 319 \); Supplementary Fig. 9b) but indicates a weaker linear dependence because coral \( \delta^{18}O \) record integrates both SST and the \( \delta^{18}O_{sw} \) (water balance at the sea surface). Obvious perturbations outside of the regression line were observed and distinguished the most recent samples related to anthropogenic CO\(_2\) emissions in the atmosphere from those related to the older preindustrial era period (Supplementary Fig. 9).

**Discussion**

As the ocean is one of the major global sinks of anthropogenic CO\(_2\) emission, the incursion of isotopically light carbon (\( ^{12}\)C) from the burning of fossil fuel into the ocean has caused considerable decrease in the \( \delta^{13}C \) of seawater DIC, known as the \( ^{13}C \) Suess effect. The \( \delta^{13}C \) ratio depletion trend in atmospheric CO\(_2\) at five Pacific stations\(^{42} \), the Law Dome ice core record\(^{43} \), and archived in numerous Pacific coral records (Supplementary Tables 2 and 3) are remarkably indistinguishable (Fig. 3). Since 1978, both instrumental and proxy records indicate a consistent average of \(-0.027\%\text{e} \) per year decrease in \( \delta^{13}C \) (Supplementary Table 3). We can easily rule out wind-driven upwelling influence on our \( \delta^{13}C \) record because its dominant effects are experienced on daily to weekly timescales\(^{44} \) and not at annual to longer timescales. The geochemical variations observed in our coral cannot be attributed to the linear growth rate because the vertical extension rates remained relatively constant (2.68 ± 0.64 mm per year) throughout the interval of this study. For this particular coral genus, similar vertical extension rates have been reported throughout the Pacific\(^{36,31} \).

This long-term secular trend towards lower \( \delta^{13}C \) values is neither related to tropical Pacific solar radiative forcing\(^{45} \) nor tropical volcanic forcing\(^{46} \) (Fig. 3 and Supplementary Table 4). The emissions of CO\(_2\) from volcanic eruptions since 1750 (Supplementary Table 4) are mostly inconsequential for climate on centennial to millennial timescales as it is at least 100 times smaller in magnitude than the Suess effect and other atmospheric perturbations outside of the regression line were observed and distinguished the most recent samples related to anthropogenic CO\(_2\) emissions in the atmosphere from those related to the older preindustrial era period.

**Fig. 3** Tropical Pacific coral \( \delta^{13}C \) records and external forcings. a The major tropical Pacific volcanic eruptions (Supplementary Table 4) and total radiative forcing from volcanic events and b solar variability\(^{45,46} \) indicate a lack of direct influence on the atmospheric and coral \( \delta^{13}C \) values. c Atmospheric \( \delta^{13}C \) values recorded at observation stations in the Pacific\(^{42} \) (blue; Supplementary Table 3) and in the Law Dome Antarctica ice core (grey; mean \( \delta^{13}C \) record\(^{43} \)) indicate strong coherence and similar timing of significant secular depletion of \( \delta^{13}C \) since the industrial revolution. d Independent records of 22 published coral \( \delta^{13}C \) signature from the central and western Pacific (Supplementary Table 2) also recorded notable acceleration in long-term \( \delta^{13}C \) depletion due to increase in anthropogenic CO\(_2\) output and oceanic uptake. e The individual coral time series vary in their native temporal resolution (monthly to 5-year average) and are all downscaled to 5-year moving means to facilitate comparison. Due to known inter-colony and inter-species offsets in coral \( \delta^{13}C \) records, we removed the mean \( \delta^{13}C \) value from each individual record over the twentieth century to centre the records.
smaller than anthropogenic emissions. These consistent secular trends from instrumental measurements and the rapid rate of coral δ¹³C depletion found at New Caledonia (−0.024‰ per year; Supplementary Table 3) recording the δ¹³C of seawater DIC over the same period (1978–2011; Fig. 3 and Supplementary Table 3) demonstrate the significant absorption of anthropogenic CO₂ emission by the oceans. The pronounced long-term depletion since 1843 by at least 1.0‰ in our D. heliopora δ¹³C record is in agreement with the nearest New Caledonia Porites sp. δ¹³C record that found a depletion of 0.9‰ since the late nineteenth century, confirming the modern uptake of anthropogenic atmospheric CO₂ by the tropical oceans.

Fig. 4 Pacific Ocean modern coral δ¹³B and pH reconstructions. The published and publicly available coral δ¹³B and pH reconstructions from the Pacific Ocean are shown in the annual gridded mean estimated pH map and listed on Supplementary Table 5. Each individual symbol corresponds to each respective record’s research location. The records are presented relative to the location of the cores from the north to the south. The locations include two records from a, b the South China Sea, c one from Guam, and d, e, f three sites in the Great Barrier Reef (GBR). The records from f Havannah Island, GBR (magenta dot) are the results of six individual Porites coral colonies. All δ¹³B records are shown in their native temporal resolution as given in the original publication. The accompanying pH reconstructions are indicated on a secondary y axis if provided by the original publication and were not converted to seawater pH based on our methodology (see Methods). Map is produced using Ocean Data View ver. 4.7.4 (ref. http://odv.awi.de) and modified manually.
Over the period 1689–2011 CE, our unique δ¹¹B record from New Caledonia (Supplementary Fig. 1) provides the first pH variability reconstruction reflecting open ocean conditions as the currently available limited reconstructions are derived from enclosed reef sites and marginal seas in the Pacific. On both interannual and decadal-interdecadal timescales, the annually resolved δ¹¹B values as a proxy of pH show pronounced oscillations (Fig. 2). These high-amplitude δ¹¹B or estimated pH variations in the surface waters in the South Pacific indicate large interannual changes of up to 0.8‰ equating to a 0.16 pH unit change in seawater (Fig. 2). The extremes of interdecadal fluctuations are smaller than those observed at the higher interannual frequencies with an average change of 0.4‰ in δ¹¹B or 0.08 pH unit in seawater. The swings between 0.08 and 0.16 pH units in South Pacific pH over interannual to interdecadal timescales are not approaching the high benchmark of 0.4 pH units for end-of-century emission scenarios as predicted by model simulations.16 The full amplitude of our δ¹¹B and reconstructed seawater pH changes are well-within the magnitude of previously reported coral-based pH reconstructions,12,13,17 regardless of the absolute scale pH reconstruction, such as the 0.3 pH units of interdecadal variability found at Flandres Reef, GBR12 (Figs. 2, 4 and Supplementary Table 5). Moreover, studies from the South China Sea14 and the GBR15 recorded large interannual to interdecadal fluctuations (0.6–0.7 pH units) that are even greater than the 0.4 pH units predicted for the end-of-the-century (Fig. 4). The large pH discrepancy of the near-shore records from the South China Sea14 and GBR15 may indicate the large spatial variability of pH and possible coastal riverine influence16 or local reef effect as compared to our open-ocean record.

In contrast to our coral-based pH reconstruction, hindcast model simulations of historical pH from the Pacific are unable to reproduce any significant modes of longer-term lower frequency variability.48 Questions arise on the reliability of these model reconstructions and simulations based solely on Pacific SST and atmospheric CO₂ for the pre-industrial period. In addition, calculated pH reconstructions based on in situ pCO₂ measurements of the past decades also lack significant low frequency variability.49,50 It is likely that the underlying observation periods used for these reconstructions and model simulations are insufficient to capture the longer timescale variability that is taking place in the Pacific Ocean. Thus, our coral δ¹¹B record appears to document the longer timescale pH changes in the Pacific that cannot be archived from the marginal seas or expressed in reanalysis studies and model simulations.

Our annually resolved δ¹³C also reveals significant interannual and interdecadal variability of up to 0.4–0.5‰. The temporal variability in δ¹³C, which can be mainly attributed to changes in seawater DIC, is nearly synchronous to those of δ¹¹B. The lower frequency variability and the magnitude of change is consistent with that found in tropical shallow water corals from both the Atlantic and the Pacific oceans.25,27 (Fig. 3). Despite the independent nature and behaviour of both proxies (δ¹¹B and δ¹³C), 37% of the variability in δ¹¹B can be explained by δ¹³C from correlation analysis (Supplementary Fig. 9). This interrelationship over interannual to interdecadal timescales thus indicates a possible common driver that contributed to the changes of both proxies. We suggest that the relationship between coral δ¹¹B and δ¹³C records at these timescales illustrates consistent and considerable marine carbonate chemistry changes in the surface ocean of the South Pacific. The shallow water mixing and uptake/release of CO₂ is mostly controlled by the temperature-dependent solubility of CO₂ over interannual timescales and it is likely that surface wind strength and hydrodynamic changes of the Pacific Ocean is the dominant process over longer decadal-interdecadal timescales. Working in combination, this dominant process of the Pacific may dampen the interannual signature of the shallow water mixing and CO₂ uptake/release changes.

Another striking result from our coral reconstruction is the long-term decreasing secular trend of the δ¹¹B signature from 24.72 to 24.09‰ over the most recent century. This major depletion corresponds to a consistent decreasing pH of 0.12 pH units since the late nineteenth century. The trend of decreasing δ¹¹B, as well as the marked increasing uptake of CO₂ depicted in the lowering of coral δ¹³C values, substantially matches the expected trend for the greater Pacific (Figs. 2 and 3). Bayesian Change-Point Analysis (CPA; see Methods for details) determined the date of the initiation of this decreasing trend for our δ¹¹B and δ¹³C records. For δ¹¹B, the timing of significant modern OA emergence occurred at 1886, between the confidence interval of 1881–1891, and after the significant depletion emergence of δ¹³C in 1843 (confidence interval: 1829–1850; Figs. 2 and 3). The timing of this event emergence between δ¹¹B and δ¹³C is thus lagged by ~40 years, which indicates the link of the slow ocean pH and carbonate chemistry changes following the uptake of atmospheric CO₂.

Before this onset of modern anthropogenic-driven OA, a coral δ¹¹B pH maximum was reached in the late 1700s (Fig. 2). The increase in δ¹¹B at the centennial scale from the sixteenth to the seventeenth century appears to be decoupled from coral δ¹³C trend as the seawater DIC signature remained relatively level. We observed that the progressive increase in δ¹¹B pH during this time period (1701–1761 CE) coincided with changes in temperature most likely linked to the termination of the LIA, a period that was documented to be 1.4 °C cooler at New Caledonia. The end of this period was however characterised by a maximum δ¹¹B pH that is contrary to the recorded maximum δ¹³C enrichment (cooler and/or more saline conditions; Fig. 2). It is possible that as a consequence to the termination of the LIA, a redistribution of water masses occurred near New Caledonia, which experienced an intrusion of cooler and/or more saline water to the region from the enhanced subtropical countercurrent.52 (Supplementary Fig. 10). These connections point to the substantial linked behaviour of pH, temperature and salinity, at the longer-term centennial timescales.

The coral-based high-amplitude pH variations occurring on interannual to decadal timescales in New Caledonia are accompanied by strong coherent SST or δ¹⁸O records for most time intervals inferred from the coral δ¹³C proxy (Fig. 2). In general, coral δ¹¹B and δ¹³C records display anti-phase coherency with higher SST concomitant with more acidic conditions. Spectral analyses reveal significant variability for both δ¹¹B and δ¹³C coral records on interannual (3–7 years) and decadal-interdecadal (11–41 years) timescales (Supplementary Fig. 11). These frequencies are typical of the well-documented ENSO and IPO climate modes of variability as observed in both instrumental53 and coral proxy records27,54 in the South Pacific. Instrumental observations suggest that ENSO events are the major control mechanisms for seawater pCO₂ variation in the equatorial Pacific on interannual to decadal timescales. During most El Niño (EN) events, the δ¹¹B signature decreases from the preceding year translating into a surface seawater decrease in pH, and the opposite occurs during La Niña (LN) events with an increase in δ¹¹B signature from the preceding year (increase in surface pH; Fig. 5 and Table 2).

The most recent ENSO events indicate possible changes of up to ±0.35‰ in coral δ¹¹B (Fig. 5 and Table 2), which can equate to ±0.07 pH units change in seawater pH for a single event under the most severe conditions. An additional important observation is that not all severe or very severe EN events result in the same magnitude of δ¹¹B depletion. Some moderate EN events (based on ERSST amplitude; Oceanic Niño Index) displayed more
ENSO event is broken down into categories of moderate (±1.0°C), severe (±1.5–2.0°C) and very severe (>±2.0°C) that is above or below the 0.5°C threshold for a period of at least 5 months in the Niño 3.4 region based on ERSST version 4 (ref.33). The threshold of ENSO event is broken down into categories of moderate (±1.0–1.4°C), severe (±1.5–2.0°C) and very severe (±2.0°C).

Fluctuations in the pH over ENSO and decadal-interdecadal timescales indicated by our coral suggests a dominant role of surface wind strength (trade winds and westerlies) changes and zonal advection of surface heat that modulated the uptake of CO2. More frequent occurrence of equatorial westerlies during rapid warming periods involve weakened trade winds, supporting the connection between South Pacific pH and tropical Pacific SST variability. Therefore, the pronounced and coherent high-amplitude variability in SST and pH over ENSO and decadal-interdecadal timescales indicated by our coral suggests a dominant role of surface wind strength (trade winds and westerlies) changes and zonal advection of surface heat in the Pacific Ocean as the primary cause and mechanism of surface ocean carbonate chemistry mixing and ultimately pH variability.

Cross-wavelets coherence analysis detecting frequency and relative phase relationship between our proxy records indicates significant non-stationary relationships between pH and SST/δ18Osw of the South Pacific (Fig. 7). Dominant low-frequency coherence at the decadal-interdecadal variability (~40 years) was observed between our coral δ13C and δ18O records from the beginning of the record up to the emergence of acidification (mid-1800s). This relationship then shifted to a more prominent higher-frequency variability of 7 years (ENSO-related) to 16 years (mid-1800s). This relationship then shifted to a more prominent higher-frequency variability of 7 years (ENSO-related) to 16 years (mid-1800s). This relationship then shifted to a more prominent higher-frequency variability of 7 years (ENSO-related) to 16 years (mid-1800s). This relationship then shifted to a more prominent higher-frequency variability of 7 years (ENSO-related) to 16 years (mid-1800s).
the ENSO frequency (4–8 years) in the last 100 years. Such observations are consistent with Central Pacific and Australian coral-based ENSO studies, where ENSO variance is significantly higher in the twentieth century than in any other time since the late Holocene. We conclude that the non-stationary behaviour of long-term pH and SST variability observed in New Caledonia, which shifted to more pronounced interannual fluctuations after ~1850 cannot be due to simple local ocean circulation changes. This prominent shift rather suggests a likely anthropogenic influence with strong CO2 uptake by the ocean because the significant trend changes and emergence timing coincides with these frequency evolution in the post-industrial era.

In summary, our reconstructed pH record from the South Pacific highlights that marine carbonate chemistry in the South Pacific is highly variable with pronounced high-amplitude pH oscillations since 1689 CE. We observed a significant emergence of secular decreasing pH trend since 1886 together with the dramatic depletion of oceanic δ13C that is related to the atmospheric 13C Suess Effect caused by the uptake of anthropogenic CO2 by the ocean. We argue that the onset of this modern acidification in the mid-1800s and the temporal coherence of δ11B pH and δ13C or marine carbonate chemistry changes are coupled to SST variability and have been consistently occurring for the last 300 years. The temporal coherence of the proxy records also has significantly changed in frequency since the emergence of the modern acidification. We implicate the changing strength of the westerlies and trade winds as important drivers forcing the pH and SST changes across the greater Pacific under possible decadal-enhanced ENSO variability (11–12 years). The changing surface ocean condition is a distinct process that modified the carbonate chemistry of the Pacific over the last three centuries with recent changes likely dominated by anthropogenic influence. Our results provide a new detailed perspective on the changes in modern OA that was likely influenced by the inseparable combination of the changing surface winds strength (trade winds and westerlies) and ENSO-related SST conditions driving the uptake of anthropogenic CO2 in the South Pacific.

**Methods**

**Coral sampling and skeletal verification.** In March 2015, a *D. heliopora* coral was cored in 7–8 m water depth at the Fausse Passe de Uitoé, New Caledonia in the southwestern Pacific (22°17′152 S, 166°10′992 E; Fig. 1 and Supplementary Fig. 1). The 7-cm diameter core was 1.3 m in total length and returned to Institut de Recherche pour le Développement (IRD) in Bondy, France in August 2015 for processing and analyses. The coral core was halved down the central growth axis and identical ca. 7 mm slabs were removed from each core half. Each coral slab was cleaned in an ultrasonic bath with 18.2 MΩ milli-Q water for 30 min and air-dried at room temperature. X-radiographs of the coral micro-sampling slab were completed at the L’Institut Multiusuliste Montsouris Paris (Supplementary Fig. 2).

Five subsamples from the coral core were systematically removed down the length of the coral core. Each subsample was further divided into individual portions for powder X-ray diffraction (XRD) and SEM analyses at the IRD, Laboratoire d’Océanographie et du Climat (LOCEAN) on the Analytical Platform AYZES (IRD-UPMC) at Bondy, France. Approximately 500 mg of crushed coral powder (<20 μm) for all subsamples were analysed on a PANalytical X’Pert Powder X-ray Diffractometer that revealed 100% aragonite. Verification of the coral’s suitability for palaeoclimate research by representative SEM analysis was conducted on a Zeiss EVO LS15 SEM. The samples were carbon coated and analytical results demonstrated good preservation of primary porosity, clear dissepiments and well-defined centres of calcification (Supplementary Fig. 3). Minor evidence and presence of secondary aragonite away from the regions of micro-sampling indicated alterations that are well within the limits of minor alterations in corals for palaeoclimate reconstruction studies.

Based on the X-radiograph positives of the coral slab with well-defined annual growth axis and identical ca. 7 mm slabs were removed from each core half. Each coral slab was cleaned in an ultrasonic bath with 18.2 MΩ milli-Q water for 30 min and air-dried at room temperature. X-radiographs of the coral micro-sampling slab were completed at the L’Institut Multiusuliste Montsouris Paris (Supplementary Fig. 2).

Five subsamples from the coral core were systematically removed down the length of the coral core. Each subsample was further divided into individual portions for powder X-ray diffraction (XRD) and SEM analyses at the IRD, Laboratoire d’Océanographie et du Climat (LOCEAN) on the Analytical Platform AYZES (IRD-UPMC) at Bondy, France. Approximately 500 mg of crushed coral powder (<20 μm) for all subsamples were analysed on a PANalytical X’Pert Powder X-ray Diffractometer that revealed 100% aragonite. Verification of the coral’s suitability for palaeoclimate research by representative SEM analysis was conducted on a Zeiss EVO LS15 SEM. The samples were carbon coated and analytical results demonstrated good preservation of primary porosity, clear dissepiments and well-defined centres of calcification (Supplementary Fig. 3). Minor evidence and presence of secondary aragonite away from the regions of micro-sampling indicated alterations that are well within the limits of minor alterations in corals for palaeoclimate reconstruction studies.

Based on the X-radiograph positives of the coral slab with well-defined annual skeletal density banding, the chronology and age model of the coral colony was developed based on band counting and then verified by 230Th/U-ages. The most recent 3 complete years of growth (2011–2014 CE) were omitted because of the presence of organic tissue layer remnants (~9–10 mm thick), which would have biased the geochemical results. Similarly, the coral section older than 1689 CE was avoided because the coral growth departed from the central vertical axis.
Fig. 7 Cross-wavelet results of proxy variability in the South Pacific. a The coral $\delta^{18}O$ (secular trend removed; trend shown in Fig. 2) compared with $\delta^{13}C$ with e cross-wavelet coherence analysis between the two time series with black contour lines enclosing time-periodicity regions with significant concentrations of coherence. Arrows to the right indicate positive coherence. d Coral $\delta^{18}O$ compared with $\delta^{10}B$ with f cross-wavelet coherence results indicating a significant shift in periodicity concentration from the earlier part of the record with longer-term decadal-interdecadal variability to the most recent years that are dominated by interannual changes.

230Th/U-age determination. For chronology verification, two samples were removed from the oldest sections of the coral as close to the sampling transect as possible. Due to sample size requirement for analysis, each sample contained ca. 4–5 years of coral growth (Supplementary Fig. 2). 230Th/U-age determinations of the corals were completed on a multi-collector inductively coupled plasma mass spectrometer (Thermo Neptune Plus MC-ICP-MS) on the Analytical Platform PANOPLY (LSCE-GEOPS), at the Laboratoire des Sciences du Climat et de l’Environnement (LSCE), Centre National de la Recherche Scientifique (CNRS), Gif-sur-Yvette, France. Details of the procedure (chemical preparation and analysis) can be found in ref. 62. The initial (234U/238U) activity ratios of samples are in agreement with the initial (234U/238U) activity ratio of modern seawater (i.e., 146 ± 0.8‰; ref. 61). The ages presented herein (Table 1) are considered reliable and criteria for the reliability of the coral 230Th/U-age are fulfilled within their 2σ-error or 95% confidence interval61.

Coral stable isotope ratios analyses ($\delta^{18}O$ and $\delta^{13}C$). Coral powder samples for $\delta^{18}O$ and $\delta^{13}C$ ratios were analysed at annual resolution and individually homogenised using mortar and pestle at the same time as the samples used for $\delta^{10}B$ analyses. About 100 μg of powder from each sample was analysed for $\delta^{18}O$ and $\delta^{13}C$ ratios using a VG Optima Stable Isotope-Ratio Mass Spectrometer on the Analytical Platform PANOPLY (LSCE-GEOPS), at the LSCE, CNRS, Gif-sur-Yvette, France. The isotope ratios are reported in ‰ deviation relative to the Vienna Pee Dee Belemnite (V-PDB). Long-term analytical precision based on repeated measurements of an in-house marble carbonate standard were ±0.04‰ (±1σ standard deviation, SD; n = 116) for $\delta^{18}O$ and ±0.02‰ (±1σ SD; n = 116) for $\delta^{13}C$ verified against NBS 19. Fifty samples or ~15% of the entire time series were analysed in duplicate and triplicate with robust average reproducibility better than ±0.05‰ (±1σ SD) for $\delta^{18}O$ and ±0.02‰ (±1σ SD) for $\delta^{13}C$, which verified the homogeneity of our measured samples (Supplementary Fig. 5).

Coral boron chemistry preparation. For $\delta^{10}B$ extraction and analysis, ~50–55 mg of coral powder was milled from each annual banding as previously described. $\delta^{10}B$ extraction from 50 ± 1 mg of coral powder per sample was slightly modified from ref. 62. All powder samples were pre-treated with a cleaning process prior to dissolution and analytical measurements to ensure that seawater salt crystals did not contaminate the powder samples. Each individual annual coral powder sample was washed in 50 ml centrifuge tubes with 2 ml of 18.2 MQ milli-Q water on a benchtop shaker for 10 min and centrifuged at 6000 rpm for 5 min to concentrate the washed powder at the bottom so the water can be carefully removed by pipet. The pre-treated coral powder samples were then dissolved in 1.25 ml of 1N HNO₃ (Optima-grade Ultrapure), the corbic acid reference standard (NBS SRM 951) were purified with 60–80 mg of anion exchange resin Amberlite IRA 743 following the Batch Method of ref. 62. The amount of resin added to each solution was sufficient for 100% extraction of the boron. Briefly, each individual batch of tubes (coral samples and standards solutions) were slowly neutralised to a pH of 7.9–8.0 with 1N NH₄OH over 4 h on a benchtop shaker with constant pH checks every 30 min. Once the pH remained stable between 8 and 9 pH units, the resin-added solutions were allowed to dissolve and analytical measurements to ensure that seawater salt crystals did not contaminate the powder samples. Each individual annual coral powder sample was washed in 50 ml centrifuge tubes with 2 ml of 18.2 MQ milli-Q water on a benchtop shaker for 10 min and centrifuged at 6000 rpm for 5 min to concentrate the washed powder at the bottom so the water can be carefully removed by pipet. The pre-treated coral powder samples were then dissolved in 1.25 ml of 1N HNO₃ (Optima-grade Ultrapure). Each coral powder solution as well as the international coral standard JCp-163, North Atlantic Seawater Standard VI (NASS VI), and the corbic acid reference standard (NBS SRM 951) were purified with 60–80 mg of anion exchange resin Amberlite IRA 743 following the Batch Method of ref. 62. The amount of resin added to each solution was sufficient for 100% extraction of the boron. Briefly, each individual batch of tubes (coral samples and standards solutions) were slowly neutralised to a pH of 7.9–8.0 with 1N NH₄OH over 4 h on a benchtop shaker with constant pH checks every 30 min. Once the pH remained stable between 8 and 9 pH units, the resin-added solutions were allowed to absorb the boron overnight. After overnight absorption, the individual tubes were rinsed five times with 18.2 MQ milli-Q water on the benchtop shaker prior to boron extraction from the resin. The concentrated boron was eluted from the resin using four successive volumes of 1.25 ml 0.1N HNO₃ (Optima-grade Ultrapure) for a total extracted volume of 5 ml. Each individual extraction steps (elution of 1.25
ml 0.1N HNO₃ lasted at least 4 h to maximize boron extraction yield. Finally, the boron concentrations were adjusted to 150–200 ppb in 0.1N HNO₃ (Optima grade Ultrapure) for δ¹¹B ratio analysis on the MC-ICP-MS.

**Coral boron isotope analysis.** Boron isotope composition (δ¹¹B) was determined using a double-focus sector-field Multi-Collector Inductively Coupled Plasma Mass Spectrometer (Thermo Neptune Plus MC-ICP-MS) on the Analytical Platform PANOPLY (LSCE-GEOPS), at LSCE, Gif-sur-Yvette, France following established methods. The δ¹¹B ratio MC-ICP-MS analysis results are reported in % deviation relative to the NBS SRM 951 standard (boric acid isotopic standard). The standards from each individual batch (ICP-1, NBS SRM 951 and NASS VI) were first measured using the ‘classic’ method in the following sequence per standard: (a) rinse solution (0.1N HNO₃), (b) blank solution (0.1N HNO₃, same solution as the boron extraction and final dilution); typical blank contribution is <−0.5%, and (c) the standard (ICP-1 or NBS SRM 951 or NASS VI). This process sequence is repeated until all the standards from each individual batch or batches are analysed. Upon the completion of this ‘classic’ method, a ‘rapid’ method sequence is initiated following the common standard-sample bracketing. The standard-sample bracketing analytical method is used to systematically control the δ¹¹B ratio mass drift over time. All the individual coral samples were prepared and measured in random sequences to exclude possible effects of chemistry by batch or analytical bias linked to MC-ICP-MS sequences. Each individual solution was measured in triplicate that included triplicate blanks. This solution was analysed nine times achieving excellent reproducibility and stability. The analytical uncertainty of δ¹¹B, ratio measurements for ICP-1 was robust (24.28 ± 0.15‰, n = 22, 2σ SD; Supplementary Fig. 8) and within the previously published 2σ SD range for MC-ICP-MS from ref. 94.

Studies using *D. heliopora* coral at sub-seasonal resolutions have shown offsets between the different skeletal materials (columnar and septal) for sub-seasonally resolved isotope-based coral reconstructions. Consequently, three sections of 20-year periods (1935–1954, 1956–1975 and 1976–1987) were removed from the same coral core slabs for replication and test of reproducibility. The powdered samples were collected on the same coral slabs from a different coral polyp on the same coral core slabs for replication and test of reproducibility. The powdered samples were processed under the same conditions to examine intra-colony reproducibility of the δ¹¹B results.

**Coral-based seawater pH calculations.** Studies have shown that coral δ¹¹B values record the pH of the extracellular calcifying fluid space or surface between the BV and the calicoblastic ectoderm and indirectly the ambient seawater pH9,65. Thus, the estimation of coral calcifying pH values (pHₐ) were calculated from coral skeletal δ¹¹B ratio following the established equation, Eq. (1) of ref. 34:

\[
\text{pH}_a = \text{pK}_B - \log \left( \frac{[\text{B}^{3+}_{\text{BW}}] + [\text{B}^{3+}_{\text{BC}}]}{[\text{B}^{3+}_{\text{BW}}] + [\text{B}^{3+}_{\text{BC}}] + 10^{(\text{pH}_a - \text{pK}_B)}} \right)
\]

(1)

The parameter pK₃,6⁶ is the equilibrium (dissociation) constant for boron in seawater. For this study, we used the constant value, 8.58, because of the limited solubility of boron carbonate and interactions with seasonal change in temperature/alkalinity, and CaCO₃ saturation in the global surface ocean, and temporal scale changes in selected locations. Mar. Chem. 164, 95–125 (2014).

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Author contributions
D.D. and A.T. initiated the original project in collaboration with E.D., D.B., C.E.L. and F. L.C. D.D. and H.C.W. designed the research. H.C.W. completed the coral sampling, sample processing, data analysis and served as primary author on this manuscript. C.E.L. through a collaborative project with E.D., D.D. and F.L.C. recovered the coral core. C.E.L. assisted H.C.W. in SEM and XRD analyses. L.B. and H.C.W. performed the $\delta^{11}$B extraction chemistry. L.B., H.C.W., A.D. and E.D. executed the $\delta^{11}$B ratio analysis. D.B. and H.C.W. completed the $\delta^{13}$C and $\delta^{18}$O ratios analyses. E.P.B. completed the $^{230}$Th/U- age determination. All of the authors assisted in interpretation, editing, discussed the results and wrote the manuscript.

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