Ocean acidification enhances primary productivity and nocturnal carbonate dissolution in intertidal rock pools

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Abstract. Human CO₂ emissions are modifying ocean carbonate chemistry, causing ocean acidification and likely already impacting marine ecosystems. In particular, there is concern that coastal, benthic calcifying organisms will be negatively affected by ocean acidification, a hypothesis largely supported by laboratory studies. The interrelationships between carbonate chemistry and marine calcifying communities in situ are complex, and natural mesocosms such as tidal pools can provide useful community-level insights. In this study, we manipulated the carbonate chemistry of intertidal pools to investigate the influence of future ocean acidification on net community production (NCP) and calcification (NCC) at emersion. Adding CO₂ at the start of the tidal emersion to simulate future acidification (+1500 µatm pCO₂, target pH 7.5) modified net production and calcification rates in the pools. By day, pools were fertilized by the increased CO₂ (+20 % increase in NCP, from 10 to 12 mmol O₂ m⁻² h⁻¹), while there was no measurable impact on NCC. During the night, pools experienced net community dissolution (NCC < 0), even under present-day conditions, when waters were supersaturated with regard to aragonite. Adding CO₂ to the pools increased nocturnal dissolution rates by 40 % (from −0.7 to −1.0 mmol CaCO₃ m⁻² h⁻¹) with no consistent impact on nocturnal community respiration. Our results suggest that ocean acidification is likely to alter temperate intertidal community metabolism on sub-daily timescales, enhancing both diurnal community production and nocturnal calcium carbonate dissolution.

1 Introduction

The ongoing increase in anthropogenic carbon dioxide (CO₂) in the atmosphere and the ocean – resulting in ocean acidification – is likely to create adverse living conditions for marine coastal communities (IPCC, 2019). Ocean acidification is projected to further decrease average surface pH by up to 0.4 units by 2100 (SSP5-8.5 scenario; Kwiatkowski et al., 2020) and is identified as a major threat to marine ecosystems (IPCC, 2019). Lower seawater pH has significant effects on marine organisms’ physiology and fitness, from altered survival and reduced growth (see review by Kroeker et al., 2013) to changes in pH homeostasis (e.g., Kottmeier et al., 2022) and metabolic rates, energy trade-offs (e.g., Dorey et al., 2013; Pan et al., 2015), and reduced feeding efficiency (e.g., Stumpp et al., 2013). Marine calcifiers – the builders of calcified structures (CaCO₃) – have been a focus of ocean acidification research due to the sensitivity of calcification to the carbonate saturation state (Ω), defined as follows:

$$\Omega = [Ca^{2+}][CO_3^{2-}] / K'_{sp},$$

where $K'_{sp}$ is the stoichiometric solubility product for the considered carbonate polymorph (i.e., $\Omega_a$ for aragonite or $\Omega_c$ for calcite). The saturation state depends on temperature, pH and pressure (lower Ω when pH or temperature decreases, and pressure increases). When Ω < 1, inert carbonate minerals tend to dissolve. The polymorphs composing the calcified structure like calcite and to a greater extent aragonite and high-magnesium calcite are prone to dissolution when pH...
decreases. For instance, in Atlantic surface waters (at 20 °C), saturation state equilibrium (Ω = 1) is reached at pH 7.3 (pCO$_2$ = 2650 μatm) for calcite but at pH 7.6 (1250 μatm) for aragonite. For high-magnesium calcite, experiments from Yamamoto et al. (2012) demonstrate that inert (dead) high-magnesium calcite from coralline algae passively dissolves at Ω$_a$ values between 3.0 and 3.2 (see also Ries et al., 2016). Organisms with calcified structures are thus likely to experience reduced net calcification due to ocean acidification through both enhanced dissolution and reduced gross calcification rates.

Aside from acidifying the ocean (increased H$^+$), increased ocean CO$_2$ uptake could affect the productivity of algae and marine plants. As CO$_2$ dissolves in the ocean, the dissolved inorganic carbon (DIC; CO$_2$, HCO$_3^-$ and CO$_3^{2-}$) concentration increases. DIC is the substrate for marine photosynthesis (mainly CO$_2$ and HCO$_3^-$), and as such, it can limit photosynthetic rates when scarce. In algae and marine plants that are carbon-limited (permanently or periodically), elevated DIC could also directly increase photosynthetic rates, and Mackey et al. (2015) propose that these rates could be further increased by the higher concentration gradient between water and the photosynthetic cells. However, the authors point out that while positive effects are theoretically expected, they may be small, specific to species’ biology and the environment they live in, and difficult to predict (see also Hurd et al., 2019). In terrestrial ecosystems, the Intergovernmental Panel on Climate Change defines CO$_2$ fertilization as “the enhancement of plant growth as a result of increased atmospheric CO$_2$ concentration” (Jia et al., 2019) and reports that CO$_2$ fertilization has likely already happened, although the magnitude of this effect depends on the plants, assemblages or ecosystems considered (and on other factors constraining growth).

The response of single species to changes such as ocean acidification and increased DIC concentrations is often insufficient to predict community-level impacts. Ecological interactions such as competition or predation can affect the outcome of perturbation experiments (Kroeker et al., 2013). For instance, Paiva et al. (2021) showed that the laboratory growth of an isopod species was an order of magnitude slower than when raised in the presence of other species from its community. In another study, Legrand et al. (2019) showed that the presence of grazers increased coralline algal calcification (+50 % in winter and +100 % in summer), but when grazers were combined with ocean acidification, algal calcification decreased more than with acidification alone. Not taking into account such interactions can therefore result in poor characterization of the effects of ocean acidification. Furthermore, while critical for a mechanistic understanding of the processes affecting marine biota, laboratory studies are seldom realistic. Typically performed under controlled, simplified and stable conditions (e.g., with respect to temperature and food), laboratory studies can better assess the effect of pH alone (Widdicombe et al., 2010). However, exposure to a stable pH (e.g., 7.6 vs. 8.0) fails to reflect the daily and seasonal variability observed in natural ecosystems, in particular coastal ones (Torres et al., 2021). Natural mesocosm perturbation experiments are thus essential tools to investigate future changes in variable and complex ecosystems, which are difficult to capture in the lab (Barry et al., 2010; Andersson et al., 2015).

Most in situ mesocosm experiments investigating the effect of ocean acidification have been conducted on planktonic communities, kept in large “bags” equilibrated to the desired pH (Riebesell et al., 2013). These studies demonstrate that adding CO$_2$ can significantly change the organization of the plankton community (Spisla et al., 2021) and increase autotrophic biomass under high-nutrient conditions (Schulz et al., 2013). Due to the technical challenges, however, benthic calcifying communities are seldom manipulated this way in situ (Widdicombe et al., 2010). Two such manipulation experiments are the studies by Albright et al. (2016, 2018), where the authors used NaOH and CO$_2$ to reproduce pre-industrial and future pH conditions on a coral reef and found evidence that reef growth had been reduced by 7 % over the industrial era and was likely to decline further. Other studies have investigated such community-level effects by either simulating “artificial”, simpler assemblages in laboratory setups (e.g., Cox et al., 2015; Pansch et al., 2016) or using phenomena such as natural CO$_2$ vents. For instance, in the vents of Ischia, as pH decreases, the presence of calcifying species declines (see review by Foo et al., 2018). Alternatively, Kwiatkowski et al. (2016) used locally induced acidification due to respiration (no CO$_2$ addition) in tidal pools, a naturally closed system, and demonstrated that nighttime dissolution of these communities was positively correlated with Ω$_a$. Here, we used tidal pools of the English Channel as ephemeral mesocosms, where we modified carbonate chemistry conditions at the start of emersion through CO$_2$ addition.

Temperate rocky tidal pools – or rock pools – are highly dynamic systems that have been long studied by naturalists since they are easy to reach, and their ecosystem structure generally resembles subtidal benthic communities (Ganning, 1971). Tidal pool organisms from the upper shore, well adapted to pool conditions, form typical benthic communities: often low in diversity, they consist of a few characteristic macroalgae (e.g., Ulva sp.) and animal species (e.g., limpets). In winter, red macroalgae – including calcifying algae – often dominate the pools, and while they do not disappear in summer, a bloom of soft green macroalgae is generally observed during the warm season. Temperature, salinity, oxygen and pH in the pools are extremely variable, often far outside the seasonal range of nearby free-flowing seawater (Legrand et al., 2018a; Morris and Taylor, 1983). Tidal pools generally emerge from the ocean twice a day in regions of semidiurnal tides, with the duration dependent on shore location and the tidal coefficient. On short timescales tidal pools act as closed systems, with carbonate chemistry easily ma-
nipulated and temporal changes reflecting in situ community metabolism (no water mass transport and negligible air–sea gas exchange).

In the present study, we used tidal pools as natural mesocosms to investigate the effect of ocean acidification on communities dominated by calcifying red algae. We measured diurnal and nocturnal net community calcification and production (or respiration) following CO₂ addition across three seasons (winter, spring and summer) to assess how tidal pool community metabolism may respond to end-of-the-21st-century high ocean acidification (pH 7.5).

2 Material and methods

2.1 Field site

The experiments were performed on a rocky intertidal shore characterized by granitic substrate on the north coast of Brittany, France, between 2019 and 2021. The beach of Bloscon (48°43′30.0″N, 3°58′10.5″W) is situated in Roscoff at the entrance of the Bay of Morlaix and has a hydrology principally affected by the waters of the English Channel and to a lesser extent the Penzé and Morlaix rivers (Fig. 1). This area is characterized by strong, oscillating, semidiurnal tides of up to 9 m. Temperatures are generally low in the deeper-flowing water (from 9–10°C in winter to 16–17°C in summer), and salinity is close to that of the adjacent Atlantic (~35; see Fig. S1 in the Supplement for detailed temperature and salinity data from the two monitoring stations Estacade and Astan; these stations are part of the Service d’Observation en Milieu Littoral, SOMLIT, a network described in Cocquem-pot et al., 2019).

2.2 Tidal pool characterization

For this study, we chose five tidal pools with high coverage in calcifying algae (>30% of the pool surface area). Both crustose (CCA) and articulated (branching) coralline algae (ACA) were present. The field site has an eastern exposure, resulting in full morning sun and relatively early shade in the evening. Foreshore locations of the pools resulted in daily emersion year-round including during neap tides (mid-tide, approx. 5–6 m above chart datum). Pools emerged for 6–7 h during low-tide periods. During that time, pools were completely separated from the adjacent open water, and their depths were effectively constant in winter (low-evaporation season), an indication that there was no seawater leakage.

The volume of each of the five pools (from 16 to 39 L; Fig. 2) was estimated in April 2021, at the end of the emersion period just before high-tide flooding, by measuring salinity changes when a known volume of freshwater was added and well mixed. To estimate the pools’ initial volumes, we also took into consideration the measured salinity changes throughout the emersion period to estimate evaporative losses and combined this with the volume directly lost through water sampling (see below). The pool projected area and the relative area covered by each type of algae were estimated from scaled aerial photographs and analyzed using ImageJ (US National Institutes of Health, Bethesda, Maryland, USA, https://imagej.nih.gov/ij, last access: 1 June 2022). Pool area ranged from 0.3 to 0.7 m² (Fig. 2). The pools had slightly different community composition, with dominant calcifying red algae represented by Lithophyllum in-crustans (CCA: 30 to 77% of the benthic cover) and El-lisolandia elongata (ACA: 0 to 6% of the benthic cover). The remaining pool area was either free of algal cover with only bare granitic rock visible or covered by soft macroalgae. In summer (September 2020 and 2021), the pools also hosted the green algae Ulva sp. and Enteromorpha sp. (2% to 44% of the benthic cover; see Supplement file SP1-2 for results detailed by season) and, in Pool E, small single branches of the brown algae Sargassum muticum, covering less than 0.5% of the pool. One limit of this method of aerial photography is that it only takes into account what is visible from above (2D). These estimates may thus be biased against algal that were hidden under the green algae canopy in summer or that were in crevices/under rocks. We also noted the presence of diverse heterotrophs such as anemones, sea sponges, small gobies and shrimps. Calcifying invertebrates were represented by four gastropod species: Phorкус lineatus, Patella alysipponensis, Patella vulgata and Gibbula pennis.
Figure 1. Field site location on a map of Europe (left). The study site (Bloscon) is located in Roscoff, Brittany, France (top right: river mapping data from HydroSHEDS, Penzé river and Morlaix river; bottom: satellite image from © Google Earth, https://earth.google.com/web/, acquired in June 2022). The SOMLIT stations Astan and Estacade are indicated with triangles (https://www.somlit.fr/en, last access: 1 June 2023).

Table 1. Sampling schedule: the dates of each field session are presented. Pools were monitored throughout multiple low-tide emersion periods (diurnal and nocturnal).

<table>
<thead>
<tr>
<th>Season</th>
<th>Dates</th>
<th>Low-tide emersion periods (N)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Diurnal</td>
</tr>
<tr>
<td>Winter</td>
<td>14–17 February 2020</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>9–19 February 2021</td>
<td>8</td>
</tr>
<tr>
<td>Spring</td>
<td>28–29 April 2021</td>
<td>2</td>
</tr>
<tr>
<td>Summer</td>
<td>2–11 September 2020</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td>6–9 September 2021</td>
<td>0</td>
</tr>
</tbody>
</table>

pared by supersaturating adjacent seawater in CO₂ using a high-pressure CO₂ cylinder.

2.4 Sampling and measurement of seawater parameters

Temperature, salinity, pH, oxygen and ammonium. From the start of the emersion period, we measured five parameters periodically using Hach Lange (Loveland, Colorado, USA) probes: temperature, pH (<i>T</i> (IntelliCAL PHC101, accuracy:
±0.02 pH units), salinity (conductivity probe IntelliCAL CDC401, ±0.1 units), oxygen concentration (optical sensor IntelliCAL LDO101, accuracy: ±0.1 for 0 to 8 mg L\(^{-1}\), ±0.2 mg L\(^{-1}\) for values greater than 8 mg L\(^{-1}\), maximum 22 mg L\(^{-1}\)) and NH\(_4^+\) concentration (ion-selective electrode IntelliCAL ISENH4181, range: 0.018–9000 mg L\(^{-1}\) NH\(_4^+\)–N). Pools were well mixed before any measurement to assure no influence of gradients forming in the pools. The measurement frequency during the emersion periods was every 15–20 minutes during the day (n = 1392) and reduced to once an hour at night (n = 159), when temperature, pH and light variations were limited or absent; pH was calibrated on the total scale (pH\(_T\)) using a TRIS (2-aminooxy-1,3-propanediol) and AMP (2-aminopyridine) buffer solution with a salinity of 35.0, following the recommendations from Dickson et al. (2007).

**Total alkalinity.** Discrete samples for total alkalinity (TA) analysis were collected hourly. The average time between two samples was 1.0 ± 0.2 h (n = 492, median = 1.0) during daytime and 1.4 ± 0.9 h (n = 135, median = 1.0) during nighttime. Seawater (150 mL) was filtered with 0.7 µm GF/F borosilicate filters directly after sampling. These samples were stored in a dark cool box until the end of the tide (max. 7 h). Upon return to the lab, they were stored at 4°C in the dark until they were either analyzed within the week or poisoned with 50 µL of saturated HgCl\(_2\). TA was assessed potentiometrically using 50.0 ± 0.5 g of seawater and a semi-automated titration system (0.1 M HCl, Titrino 848 plus by Metrohm, Switzerland; electrode calibrated on the National Bureau of Standards scale). TA was determined using Gran titration (Gran, 1952) according to the method of Haraldsson et al. (1997) and verified against reference standards provided by (Scrippps Institute of Oceanography, University of California, San Diego, United States). TA samples were analyzed with single (n = 312) or duplicate (n = 320) measurements (the median of the standard deviation between duplicates was 1.05 µmol kg\(^{-1}\)). TA was salinity-normalized before further calculations to take into account possible dilution from rain or concentration from evaporation.

To take into account the influence of the changes in nutrients (NO\(_3^-\), NO\(_2^-\), PO\(_4^{3-}\) and NH\(_4^+\)) on the changes in TA (Gazeau et al., 2015), we sampled seawater for nutrients in two samples was 1.0 µmol kg\(^{-1}\), i.e., < 2 % of the observed change in TA; see full details in “Nutrients” in the Supplement) and thus are ignored here.

**Light measurements.** Surface irradiance (photosynthetically active radiation, PAR) was continuously recorded (every minute) during experiments at the field station using a LI-COR flat quantum light sensor (LI-190R) and logger (LI-1500, LI-COR, Germany).

**Adjacent waters.** Temperature, salinity, pH and TA (n = 5) were similarly sampled and measured at the sampling site during ebb tide for the three seasons.

### 2.5 Carbonate chemistry calculations

The carbonate system parameters (e.g., pCO\(_2\); DIC concentration; CO\(_3^{2-}\) concentration; and Ω\(_{ar}\), the aragonite saturation state) were calculated from the measurements of pH\(_T\), TA, temperature and salinity using the R package seacarb (Gatutso et al., 2021) with the default dissociation constants recommended by Dickson et al. (2007), except for the low temperatures encountered in February 2021, where the refined constants of Sulpis et al. (2020) were used. When salinity decreased by more than 1.5 units h\(^{-1}\), data were excluded to avoid rain effects in the present study. When calculated DIC and Ω\(_{ar}\) were negative, likely due to inaccuracies in the measurement and computation of the carbonate system, values were approximated to be 0 (7/627 values). The air–sea gas fluxes due to net diffusive transport were considered to be negligible in the pools (see detailed explanation the interactive discussion).

### 2.6 Biological activity calculations

The rates of net community calcification (NCC; mmol CaCO\(_3\) m\(^{-2}\) h\(^{-1}\)) and net community production (NCP) or community respiration (CR; mmol O\(_2\) m\(^{-2}\) h\(^{-1}\) or mmol C m\(^{-2}\) h\(^{-1}\)) were calculated between two consecutive sampling times. These rates respectively represent the measured changes in net CaCO\(_3\) precipitation and net organic carbon production (or oxygen consumption) by the community. Positive NCC represents net CaCO\(_3\) precipitation (gross precipitation > dissolution), and negative rates represent net dissolution (dissolution > precipitation). NCP is positive when the community primary production exceeds respiration and negative when community primary production is less than respiration. We use CR for nights, when there is no primary production (oxygen consumption and carbon release only).

NCC was calculated using the alkalinity anomaly method (Smith and Key, 1975). Briefly, for each mole of CaCO\(_3\) precipitated, 2 mol of HCO\(_3^-\) combine with Ca\(^{2+}\), and TA decreases by 2 mol (Eq. 1). Two independent estimates of NCP (or CR) were calculated, one derived from changes in ΔDIC (NCP\(_{DIC}\) or CR\(_{DIC}\)) and one derived from ΔDIC and NCC (NCP\(_{DIC}\) or CR\(_{DIC}\)).
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NCC and NCP (or CR) were thus calculated as follows:

\[
\text{NCC} = -\frac{\Delta \text{TA}}{2\Delta t} \times \frac{V}{S} \quad (1)
\]

\[
\text{NCP} \text{ (or CR)}_{\text{O}_2} = \frac{\Delta \text{O}_2}{\Delta t} \times \frac{V}{S} \quad (2)
\]

\[
\text{NCP} \text{ (or CR)}_{\text{DIC}} = -\frac{\Delta \text{DIC}}{\Delta t} \times \frac{V}{S} - \text{NCC}, \quad (3)
\]

with \(\Delta \text{TA}\) (mmol L\(^{-1}\)), \(\Delta \text{DIC}\) (mmol L\(^{-1}\)) and \(\Delta \text{O}_2\) (mmol L\(^{-1}\)) the change in concentration of TA, DIC and O\(_2\) between consecutive samples and \(\Delta t\) the duration between consecutive samples (h), \(V\) the pool volume (L) and \(S\) the pool surface area (m\(^2\)).

Up to seven NCC and NCP (or CR) rates were calculated for each pool during each emersion period (one per hour). These rates were used to investigate the direct correlation between biological activity and environmental factors such as light intensity or \(\Omega_a\). Rates calculated this way are however not independent from each other (i.e., the rate measured at \(t + 2\) is dependent on the rate at \(t + 1\)), limiting further statistical analyses of the effect of the treatment. This is why, to investigate the effect of pH treatment (present vs. future) on community biological activity, we also calculated NCC and NCP or CR using linear regressions (NCC\(_{\text{lm}}\) and NCP\(_{\text{lm}}\) or CR\(_{\text{lm}}\)) between TA, [DIC] and [O\(_2\)] and time after the start of the emersion period (for detailed results of the regressions, e.g., goodness of fit, see LM1-3 in the Supplement). The few data from diurnal tides that were taken after sunset were excluded from these regressions. For oxygen, data were limited to the first 3 h of emersion as high O\(_2\) concentrations (\(> 22\) mg L\(^{-1}\)) and supersaturation (\(> 200\%) led to inaccurate measurements and/or possible oxygen degassing afterwards (see LM2 in the Supplement). This regression approach provides a single estimate of the rate of NCC, NCP\(_{\text{DIC}}\) (or CR\(_{\text{DIC}}\)) and NCP\(_{\text{O}_2}\) (or CR\(_{\text{O}_2}\)) for each pool during each emersion period (\(n = 17\) diurnal and 6 nocturnal low-tide emersion periods \(\times 5\) pools = 115). These rates were then used in generalized linear mixed models (GLMMs) to assess the effect of pH treatment on diurnal and nocturnal biological activity (see “Statistical analyses” below).

We calculated community calcification budget and production budget (respectively CCB and CPB) at emersion as an indication of the night–day balance in calcification and production: when CCB (CPB) is positive, the pool community calcifies (produces) more by day than it dissolves (respires) at night. Both were calculated for winter (February 2020 and 2021) and summer (September 2020 and 2021) for each pool as follows:

\[
\text{CCB} = \text{NCC}_{\text{D}} + \text{NCC}_{\text{N}} \quad (4)
\]

\[
\text{CPB} = \text{NCP}_{\text{D}} + \text{CR}_{\text{N}}, \quad (5)
\]

with NCC\(_{\text{D}}\) and NCP\(_{\text{D}}\) (> 0) the average diurnal NCC and NCP for a given pool for a treatment and a season and NCC\(_{\text{N}}\) and CR\(_{\text{N}}\) (< 0) the average nocturnal NCC and dark respiration for the same conditions. Three approaches were used for estimating CPB, given the uncertainties in each NCP estimate (see Discussion): (1) O\(_2\)-derived estimates of NCP (CPB\(_{\text{O}_2}\)), (2) DIC-derived estimates (CPB\(_{\text{DIC}}\)), and (3) a “mixed” approach that combined nocturnal CR\(_{\text{O}_2}\) and diurnal NCP\(_{\text{DIC}}\) (CPB\(_{\text{m}}\)) under the assumption that 1 mol of carbon is produced/consumed when 1 mol of O\(_2\) is produced/consumed. Although CPB resembles gross community production in the way the rates are calculated (difference between light and dark net production/respiration rates), if one wanted to reuse these rates for gross community production, they should do so with care due to differences in night and day temperature (see extended discussion on this subject in Bracken et al., 2022). The treatment effect was assessed on CCB and CPB by comparing the change due to the future treatment in each pool.

2.7 Statistical analyses

All data are presented as mean ± standard deviation (SD). The analyses were made using the software R (R Core Team, 2017). The level of significance used was 5%. Because data were measured in the same five pools but on different days for different treatments, we used GLMM to test for the effect of treatment on NCC\(_{\text{lm}}\) and on O\(_2\) and DIC-derived NCP\(_{\text{lm}}\) (or CR\(_{\text{lm}}\)), assigning sampling days (i.e., low-tide emersion periods) as the random factor and pools (five levels), mean temperature of the pool during low-tide emersion period (a continuous proxy for season) and treatment (treat: future vs. present) as fixed factors. This was performed using the R package nlme (Pinheiro et al., 2018). Models with and without standardized residuals were compared using analyses of variance (ANOVAs), and, when different, the Akaike information criterion (AIC) was used to choose the best-fitted model of the two. For GLMMs, mean daily PAR was not used as it has strong collinearity with mean daily temperature/season. We used ANOVAs to test the effect of temperature, pool and treatment on initial (averaged over the first hour of emersion) and final (averaged for > 5 h after emersion) carbonate chemistry conditions. The normality of the data was tested using Shapiro–Wilk tests and qq plots, while variance homogeneity was tested with Bartlett tests.

3 Results

3.1 Environmental conditions

Adjacent waters. Temperatures (and salinity) measured in the seawater adjacent to the pools were 6–7 °C in winter (February; salinity \(S = 35.0\)), 11–12 °C in spring (April; \(S = 35.5\)) and 17–18 °C in summer (September; \(S = 36.0\)). This seawater was characterized by an average pH\(_T\) of 8.01 ± 0.06 units, total alkalinity of 2319 ± 6 mmol kg\(^{-1}\), \(pCO_2\) of
445 ± 69 µatm, \( \Omega_a \) of 2.2 ± 0.3 and \([O_2]\) of 100 ± 1 % of air saturation (or 10.1 ± 1.5 mg L\(^{-1}\); \( n = 5 \)).

**Light duration and intensity.** In Roscoff, day–night (i.e., no light) periods are typically 10 h of day, 14 h of night in February and 14 h of day, 10 h of night in April and September. Photosynthetically active radiation (PAR) was 2 to 3 times higher in spring/summer (April/September ~ 1500 µmol m\(^{-2}\) s\(^{-1}\); Fig. 3a) than in winter (February ~ 500 µmol m\(^{-2}\) s\(^{-1}\)).

**Carbonate chemistry conditions at the start of the emersion period (<1 h post-emersion).** For both diurnal and nocturnal tides, the initial pH was significantly lower in pools with added CO\(_2\) than in the present-day pools (pH\( _T \) = 8.2 ± 0.1 vs. 7.5 ± 0.2 units during the day and 8.0 ± 0.1 vs. 7.4 ± 0.1 units at night for present and future pools respectively; treat: \( p < 0.001 \); detailed results in Figs. S2–S3, Table S1–S2). This corresponds to pCO\(_2\) of 260 ± 100 vs. 1900 ± 835 µatm (day) and 510 ± 90 vs. 2310 ± 410 µatm (night) for pools under present and future conditions respectively. Adding CO\(_2\) to the pools increased the mean DIC concentration by 320 µmol kg\(^{-1}\) during the day and 240 µmol kg\(^{-1}\) during the night. Under present-day conditions, the pools started at supersaturated levels with regard to aragonite (day: \( \Omega_a = 3.3 ± 1.3; \) night: 2.2 ± 0.3). Adding CO\(_2\) significantly decreased \( \Omega_a \) (treat: \( p < 0.001 \); Table S1), leading to initial future conditions often undersaturated with regard to aragonite (\( \Omega_a = 0.8 ± 0.5 \) by day and always undersaturated conditions by night (\( \Omega_a = 0.6 ± 0.1 \)). Furthermore, under future diurnal conditions, pools were always undersaturated with respect to aragonite from the start of the emersion period in February (\( \Omega_a = 0.5 ± 0.2 \)) but not in April (\( \Omega_a = 1.1 ± 0.7 \)) and September (\( \Omega_a = 1.2 ± 0.5 \); Table S1). At the start of emersion, total alkalinity was 2303 ± 34 µmol kg\(^{-1}\) (similar to adjacent seawater) and uncorrelated with treatment (\( p > 0.05 \)) and temperature (\( p > 0.6 \)).

As data were averaged on the first hour post-emersion; the mean initial oxygen concentration calculated was already affected by NCP by day (14.0 ± 2 mg O\(_2\) L\(^{-1}\)) and CR by night (9.5 ± 1.5 mg O\(_2\) L\(^{-1}\) vs. 10.1 ± 1.5 mg O\(_2\) L\(^{-1}\) for adjacent seawater). This was also visible in CO\(_2\) partial pressure, with lower pCO\(_2\) than expected during the first hour post-emersion by day (262 ± 102 µatm vs. 445 ± 69 µatm for adjacent seawater) and higher pCO\(_2\) at night (508 ± 88 µatm) under the present-day conditions.

### 3.2 Diurnal tides

**Diurnal pool chemistry.** Starting from the aforementioned values at emersion, the pools followed a clear temporal evolution due to solar irradiance and community metabolism (Fig. 3). Firstly, we observed increases in salinity (+1.5 units on average; Fig. 3a) and temperature (+4 °C in September, +6 °C in April on average) in summer and spring. In winter, temperatures tended to decrease (−1.7 °C on average), with air temperatures colder than that of the seawater; salinity was stable (35.5 ± 0.8).

Secondly, we observed positive NCP, corroborated by a doubling in oxygen concentration (Fig. 3a) a few hours after the start of emersion. In parallel, the seawater DIC concentration decreased by 50 % from the initial concentration (from 2130 ± 195 to 1140 ± 560 µmol kg\(^{-1}\); Fig. 3b), the range of which largely depended on the season (Fig. S2). For instance, in February, DIC consumption in pool seawater averaged ~ 700 µmol kg\(^{-1}\) over a low-tide period, while it averaged ~ 1500 µmol kg\(^{-1}\) in September. Particularly extreme conditions, with DIC concentrations effectively reaching 0 µmol kg\(^{-1}\), were observed in two of the pools at three tides in September 2020 (see further details below in “The particular case of September 2020 tides”). At the end of diurnal emissions, average pCO\(_2\) was always below 100 µatm, reaching as low as 1 ± 2 µatm in September (Fig. 3b, Table S1). As a result, diurnal pH\( _T \) increased to 9.1 ± 0.6 by the end of emersion, with maximum values up to 10.3 in summer (Fig. 3b). At the end of a diurnal emersion period, the pools’ pH was stable, either reaching a plateau or decreasing after sunset (see PAR in Fig. 3a). Similarly, at the end of diurnal emersion periods, \( \Omega_a \) was high (5.6 ± 3.0 on average, max 10.4). Lastly, we observed a diurnal decrease in TA by 415 µmol kg\(^{-1}\) on average, indicative of net calcification.

It is noteworthy that the carbonate chemistry conditions experienced at the end of diurnal emersion converged whatever the initial treatment (Fig. S2, Table S1). For instance, while \( \Omega_a \) was significantly different between treatments at the start of the emersion period, both treatments reached similar \( \Omega_a \) at the end of emersion (> 5 h) of around 5.3 ± 2.2 (ANOVA: treat – \( p = 0.1 \), temp – \( p = 0.002 \), pool – \( p = 0.01 \)). There was less convergence for pH\( _T \); even 5 h after emersion, the treatments were still statistically significant, although the differences in pH were small (\( p < 0.001 \) for pH\( _T \) with 9.2 ± 0.6 for present and 9.0 ± 0.6 for future pools).

**Diurnal biological activity.** Net community production was positive during daytime, except at sunset (Fig. 3c). NCP was significantly correlated with light intensity (PAR), and further results for hourly NCP and their correlation with hourly averaged PAR, \( \Omega_a \), and temperature can be found in the Supplement (Figs. S6 and S7).

As expected, seasons/temperature affected net oxygen production (O\(_2\)-derived NCP\(_{\text{im}}\)), increasing from 7 ± 3 mmol O\(_2\) m\(^{-2}\) h\(^{-1}\) in February to 18 ± 11 mmol O\(_2\) m\(^{-2}\) h\(^{-1}\) in September (GLMM, \( p < 0.001 \); Fig. 4a and Table 2A). CO\(_2\) addition increased O\(_2\)-derived NCP\(_{\text{im}}\) by 20 % on average over all seasons, from 10 ± 7 mmol O\(_2\) m\(^{-2}\) h\(^{-1}\) under present conditions to 12 ± 9 mmol O\(_2\) m\(^{-2}\) h\(^{-1}\) (\( p = 0.0015 \)). Net oxygen production differed across pools (\( p = 0.003 \)), with significantly more productivity in Pools C (17.6 ± 12.7 mmol O\(_2\) m\(^{-2}\) h\(^{-1}\)) and D.
Figure 3. Composite 24h conditions and biological activity for all pools. (a) Temperature (°C), salinity and oxygen concentration (mg L⁻¹) and photosynthetically active radiation (PAR; µmol m⁻² s⁻¹); (b) pH, total alkalinity (TA; µmol kg⁻¹), pCO₂ (µatm), dissolved inorganic carbon (DIC; µmol kg⁻¹) and aragonite saturation state (Ωa); and (c) DIC- and O₂-derived NCP or CR (mmol C or O m⁻² h⁻¹) and NCC (mmol CaCO₃ m⁻² h⁻¹). Colors represent seasons (a: blue for February, orange for April, red for September) and treatment (b and c: purple for present and green for future). Horizontal dotted gray lines represent the mean values of the adjacent ocean. Curves were fitted by season for PAR and for diurnal NCP and NCC using a local polynomial regression (LOESS) with a 95 % confidence interval. Number of observations: n = 1551 for temperature, salinity and pH; n = 1169 for oxygen concentration (data recorded < 22 mg L⁻¹); and n = 632 (hourly data) for the carbonate chemistry parameters NCC and NCP or CR (b). All pools are shown.
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(10.6 ± 5.6 mmol O$_2$ m$^{-2}$ h$^{-1}$) compared to Pools A, B and E (8.1 ± 4.1 mmol O$_2$ m$^{-2}$ h$^{-1}$).

Results are similar for DIC-derived NCP$_{lm}$ (Fig. 4a and Table 2b), with primary production ranging from 6 ± 2 mmol C m$^{-2}$ h$^{-1}$ in February to up to 12 ± 5 mmol C m$^{-2}$ h$^{-1}$ in September ($p < 0.001$). As for O$_2$-derived NCP$_{lm}$, CO$_2$ addition increased DIC-derived NCP$_{lm}$ by 20% on average over all seasons ($p < 0.001$; Fig. 4a). This increase was particularly apparent in the summer, when NCP$_{lm}$ increased from 11 ± 4 mmol m$^{-2}$ h$^{-1}$ in the present treatment to 15 ± 5 mmol C m$^{-2}$ h$^{-1}$ in the future treatment (+35%). Productivity (NCP$_{lm}$) was significantly lower in Pools B and E than in Pool A and significantly higher in Pools C and D ($p < 0.003$).

By day, with the exception of sunset, net community calcification was positive (NCC and NCC$_{lm}$ > 0; Figs. 3c and 4b) and occurred in an environment that was supersaturated with regard to aragonite (Fig. 3b). This was with the exception of a few emersion periods in September 2020 where dissolution was observed despite high-saturation-state conditions (further details below). Similar to NCP$_{lm}$, diurnal net calcification rates (NCC$_{lm}$) were strongly influenced by temperature/season (GLMM, $p < 0.001$; Fig. 4b and Table 2c), ranging from 1.2 ± 0.5 mmol CaCO$_3$ m$^{-2}$ h$^{-1}$ in February to 3.3 ± 1.3 mmol CaCO$_3$ m$^{-2}$ h$^{-1}$ in September. NCC hourly rates positively correlated with averaged $\Omega_a$ ($p < 0.0001$, NCC = 0.15 $\times$ $\Omega_a$ + 0.85; linear regression presented in Fig. S7) significantly but not strongly ($R^2 = 10\%$). CO$_2$ addition did not influence NCC$_{lm}$ rates during the day ($p = 0.47$). However, NCC$_{lm}$ did differ across pools ($p < 0.003$): rates were relatively low in Pool E (lowest CCA cover: 30%; 1.4 ± 1.4 mmol CaCO$_3$ m$^{-2}$ h$^{-1}$) and high in Pool D (highest CCA cover: 70%; 2.2 ± 0.8 mmol CaCO$_3$ m$^{-2}$ h$^{-1}$) compared to the three other pools (2.0 ± 1.25 mmol CaCO$_3$ m$^{-2}$ h$^{-1}$).

3.3 Nocturnal tides

Nocturnal pool chemistry. Seawater temperatures during the nights were stable (Fig. 3a) throughout the emersion period in summer (from 17.3 ± 0.4°C < 1 h post-emersion to 17.2 ± 0.2°C > 5 h post-emersion) and winter (from 8.4 ± 1.4 to 7.8 ± 2.7°C in February; no April nights). We highlight the wide range of winter seawater temperatures with an exceptionally cold tidal cycle (5°C on 13 February 2021) due to air temperatures of 3–4°C (observations from the Île de Batz meteorological station). There was a decline in salinity at night in some winter emersion periods (Fig. 3a) due to high air humidity and/or rain. Data where salinity dropped by more than 1.5 units in less than an hour were removed from further analyses of net community calcification and respiration.

After 5 h of emersion, O$_2$ concentration had decreased by 50% (from 10.1 ± 1.5 mg O$_2$ L$^{-1}$ to 4.9 ± 3.3 mg O$_2$ L$^{-1}$) (Fig. 3a) due to community respiration. Simultaneously, pH$_I$ decreased to 7.6 ± 0.2 (present) or stayed at 7.4 ± 0.2 (future; Figs. 3b and S2, Table S1), with significant effects of pools, treatment and temperature ($p < 0.001$ for all three). DIC concentration increased by +256 µmol kg$^{-1}$ on average over an emersion period. The range of this increase depended on the temperature and the pool: in winter (5–10°C), present-day pool seawater gained +130 µmol kg$^{-1}$ (+60 for future pools) of DIC over an emersion period, when in summer it gained +370 µmol kg$^{-1}$ for present (future: +310 µmol kg$^{-1}$) pools (Fig. S3). Saturation state converged towards similar undersaturated levels at night (Figs. 3b and S2, Table S1): $\Omega_a$ stayed stable in the future treatment (0.7 ± 0.2 units on average) and decreased in the present-day treatment (−1.2 units from initial $\Omega_a$). At the end of nocturnal emersion $\Omega_a$ values were still statistically different due to the initial treatment ($p < 0.001$ for treat, temp and pools).

Nocturnal biological activity. At night, oxygen was consumed; i.e., we observed dark respiration (CR; Fig. 3c). Community respiration (O$_2$-derived CR$_{lm}$) varied according to season ($p < 0.001$; Fig. 4a and Table 2a): temperature linearly increased nocturnal respiration rates from −1.0 ± 1.2 mmol O$_2$ m$^{-2}$ h$^{-1}$ in February to −4.7 ± 1.3 mmol O$_2$ m$^{-2}$ h$^{-1}$ in September. The CO$_2$ treatment did not influence night respiration ($p = 0.39$). Respiration rates were significantly influenced by pools ($p = 0.03$), probably linked to the relative biomass of heterotrophs and autotrophs; respiration was significantly higher in Pool C (−4.6 ± 2.8 mmol O$_2$ m$^{-2}$ h$^{-1}$) and significantly lower in Pool E (−2.4 ± 1.4 mmol O$_2$ m$^{-2}$ h$^{-1}$) than in Pools A, B and D (−3.4 ± 2.1 mmol O$_2$ m$^{-2}$ h$^{-1}$).

Night respiration estimated using DIC and NCC was near zero (CR$_{lm} = −0.2 ± 0.7$ mmol m$^{-2}$ h$^{-1}$). At these low rates, uncertainties associated with much higher rates of net dissolution (negative NCC) sometimes led to spuriously positive DIC-derived CR estimates, hindering interpretation. Nevertheless, DIC-derived community respiration was 10 times lower in February than in September (−0.2 ± 0.7 and −2.3 ± 1.1 mmol C m$^{-2}$ h$^{-1}$ respectively), although it was not linearly driven by temperature ($p = 0.053$; Fig. 4B and Table 2B). Adding CO$_2$ to the pools influenced DIC-derived community respiration in a way that was inverse to that seen with O$_2$, but as stated above, this was likely an artifact of subtracting NCC from small DIC changes. As for O$_2$, DIC-derived CR$_{lm}$ significantly changed depending on the pools.

At night, the pools experienced significant net community dissolution (NCC < 0; Fig. 3c) even when waters were supersaturated with regard to aragonite in the present treatment ($\Omega_a > 1$; Fig. 3b). Nocturnal net dissolution rates (NCC$_{lm}$) were not significantly affected by temperature in the range investigated (5–18°C, $p = 0.57$; Fig. 4c and Table 2c). However, adding CO$_2$ in the pools increased net dissolution rates ($p = 0.0017$) from −0.7 ± 0.3 mmol CaCO$_3$ m$^{-2}$ h$^{-1}$ to −1.0 ± 0.4 mmol CaCO$_3$ m$^{-2}$ h$^{-1}$ (+40%). Similarly, looking instead at hourly rates (NCC), dissolution correlated significantly ($p < 0.0001$ with $\Omega_a$ (NCC = 0.34 x $\Omega_a$).
Adding CO$_2$ in summer (FEB: CPB$_{W}$ rates were calculated for each pool, each tide and each treatment: $n = 50$ (FEB, day), $n = 10$ (FEB, night), $n = 10$ (APR, day), $n = 25$ (SEP, day), $n = 20$ (SEP, night). Seasons: FEB for winter (pooled February 2020 and 2021), APR for spring (April 2021) and SEP for summer (pooled September 2020 and 2021). Note that for NCC$_{lm}$, nights ($< 0$) and days ($> 0$) have different $y$-axis scales for better visualization of night differences. Statistical details of the linear regressions can be found in the corresponding Supplement. For O$_2$-derived NCP$_{lm}$, in September, three rates were out of the range plotted, and their values are indicated next to the small arrow.

- $1.22$, $R^2 = 11\%$; Fig. S7). The strength of this correlation depended on seasons and pools (Fig. S8). Net dissolution rates (NCC$_{lm}$) significantly differed by pool ($p < 0.0017$): the lowest rates were observed in Pool E ($-0.4 \pm 0.2$ mmol CaCO$_3$ m$^{-2}$ h$^{-1}$) – the pool with the lowest CCA cover – and the highest dissolution in Pool D – the pool with the highest CCA cover ($-1.0 \pm 0.4$ vs. $-0.9 \pm 0.3$ mmol CaCO$_3$ m$^{-2}$ h$^{-1}$ for A, B and C).

### 3.4 Influence of the treatment on CPB and CCB

Pools fixed more carbon during the day than they respired at night; i.e., the community production budget (CPB: balance between night and day) was positive in all the pools, in both winter and summer and whatever the treatment (Fig. 5). CPB$_{DIC}$ and CPB$_{lm}$ estimates were typically lower than CPB$_{O_2}$ (in 14/20 cases and 18/20 cases respectively). The production budget was significantly lower in winter than in summer (FEB: $\text{CPB}_{O_2} = 3 \pm 1$ mmol O$_2$ m$^{-2}$ h$^{-1}$; SEP: $7 \pm 5$ mmol O$_2$ m$^{-2}$ h$^{-1}$; $t = -2.4$, $df = 9.8$, $p = 0.03$). Adding CO$_2$ increased CPB in all the pools in summer by $+ 3.0 \pm 2.1$ mmol O$_2$ m$^{-2}$ h$^{-1}$, an increase in production by 50% to 80% ($\Delta$CPB; Fig. 5). In winter, there was no evidence of such a “fertilization effect” across the most accurate CPB estimates for this season (CPB$_{O_2}$, CPB$_{lm}$): we only observed a significant increase in production due to CO$_2$ addition in two of the pools ($+60\%$ to $+120\%$ for A and B). For the three other pools, CPB induced either minimal changes ($< 20\%$ for C and E) or a decrease in production (D: down to $-34\%$). DIC-derived $\Delta$CPB in winter (all positive) should be interpreted with caution since some nocturnal CR$_{lm}$ values were spuriously positive in the future treatment (see “Nocturnal biological activity” above).

The pools calcified more during the day than they dissolved at night (CCB $> 0$), in both summer and winter (Fig. 5). CCB was significantly lower in winter than in summer (FEB: CCB $= 0.2 \pm 0.2$ mmol CaCO$_3$ m$^{-2}$ h$^{-1}$; SEP: $1.2 \pm 0.6$ mmol CaCO$_3$ m$^{-2}$ h$^{-1}$; $t = -5.2$, $df = 11.7$, $p = 0.0002$). In winter, adding CO$_2$ decreased CCB by more than 80% in Pools C, D and E (Fig. 5). The CO$_2$ addition even resulted in a transition from a positive community calcification balance to dissolution in Pool C (133% change, from $+0.5$ to $-0.2$ mmol CaCO$_3$ m$^{-2}$ h$^{-1}$). For the two other pools (A and B), winter CO$_2$ addition increased their relatively small calcification balance (A: $+87\%$, from 0.1 to 0.2 mmol CaCO$_3$ m$^{-2}$ h$^{-1}$; B: $+71\%$, from 0.2 to 0.3 mmol CaCO$_3$ m$^{-2}$ h$^{-1}$). In summer, changes in CCB due to treatment appeared minimal in Pools A, B and E ($< 15\%$ change) and either increased (C: $+67\%$) or decreased (D: $-57\%$) in the two other pools.
3.5 The particular case of September 2020 tides

During diurnal tides of September 2020 (high-PAR and high-temperature summer conditions), we observed an unexpected phenomenon: dissolution occurred at extremely high pH values (9–10) in Pools C and E (Fig. 6). Under these conditions effectively all the seawater DIC in these pools was consumed by photosynthesis and calcification (DIC ≈ 0 mmol kg⁻¹) 4 h after emersion. As such, the CO₂⁻³ concentration was also effectively zero, and the pools reached very low saturation states (Ω ≈ 0) despite high pH (Fig. 6). These conditions were quickly followed by indicators of CaCO₃ dissolution (increasing TA and DIC) instead of the expected diurnal precipitation. It is therefore notewor-
4 Discussion

Temperate tidal pools are environments of extreme variability. In our pools, we observed seawater temperatures that could increase by up to 10°C in a few hours compared to the adjacent ocean. During diurnal emersion periods, oxygen concentrations doubled, and pH could increase to pH 10 under present-day summer conditions. At night, pH routinely reached levels usually used as the “treatment” for ocean acidification perturbation experiments (∼7.6). Organisms present in the tidal pools may therefore already be adapted or acclimatized to extreme variability in pH and saturation state, which could affect their responses to ocean acidification (Andersson et al., 2015). For example, CCA from a site with naturally high pCO₂ variability calcified ∼50% more than individuals from a nearby site of low variability when subjected to oscillating high pCO₂ treatments (Johnson et al., 2014). Here we show that, even in intertidal communities likely already acclimated or adapted to variable conditions, with potentially large phenotypic plasticity, acidification can still modify net community production and calcification rates.

4.1 Diurnal fertilization under CO₂ addition

Adding CO₂ to simulate future seawater acidification in the pools led to a diurnal fertilization effect. The community’s net primary production increased by 20% on average across all seasons, which was particularly visible in summer (∼+35%), when temperatures/metabolic rates were high. Adding CO₂, we also added substrate for photosynthesis in the form of DIC (Fig. 3b) that the algae of the pools can assimilate, potentially supporting higher DIC use and algal primary production. This effect was apparent from the start of the emersion, suggesting a direct effect of increasing DIC concentration in the pools. It seems that photosynthesis in the pools was carbon-limited and that carbon addition therefore enhanced primary production in winter and to an even greater extent in summer. During photosynthesis, the uptake of inorganic carbon leads to a significant decrease in DIC – even under present-day conditions. Intertidal algae are typically adapted to this, with coralline algae in particular containing CCMs (CO₂ concentrating mechanisms) that allow them to achieve primary production in low DIC concentrations (Raven, 2011). Increasing seawater DIC may however promote an increase in active and/or passive CO₂ and HCO₃⁻ fluxes towards photosynthetic compartments. Borowitzka (1981) found that the photosynthetic rate of an intertidal CCA was highest at pH 6.5 to 7.5 (increased from pH 8.1), a change in pH that was achieved using HCl, suggesting that increased photosynthetic activity could also be linked to proton gradients/pumps and/or decreased energy expenditure.

Figure 5. Community production budget (CPB) (a) and calcification budget (CCB; mmol CaCO₃ m⁻² h⁻¹) (b) by treatment (purple for present and green for future) for each pool and season (same legend as Fig. 4). CPB > 0 if diurnal NCP > nocturnal respiration, and CCB > 0 if diurnal NCC > nocturnal dissolution. CPB was estimated three different ways: from O₂-derived NCP (bars), from DIC-derived NCP (round symbols) and from nocturnal O₂-derived CR combined with diurnal DIC-derived NCP (mixed, vertical segments). Panels (c) and (d) present the change (%) in diel production (ΔCPB, a, c) and diel calcification (ΔCCB, b, d) due to CO₂ addition. Positive ΔCPB indicates a fertilization effect due to the CO₂ addition; negative ΔCCB is expected if the CO₂ addition decreases net calcification/increases net dissolution. All three methods to estimate CPB indicate a fertilization effect in summer.

thy that dissolution may happen at high pH and that pH and Ω can decorrelate (Fig. 7) in situations with high photosynthesis and limited mixing of water masses.
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Figure 6. Time series for September 2020 diurnal data only: total alkalinity (TA; \(\text{µmol kg}^{-1}\)), dissolved inorganic carbon, CO\(_2\)\(^{-}\) concentration (\(\text{µmol kg}^{-1}\)), aragonite saturation state (\(\Omega_a\)) and NCC (mmol m\(^{-2}\) h\(^{-1}\)) with time after emersion by treatment (purple for present and green for future). The lines in bold represent individual Pools C and E, which switched from calcification to dissolution when pH\(_T\) was still above 9. A similar figure in the Supplement (Fig. S4) shows that sunset and irradiance are not correlated with the sudden change towards dissolution.

Figure 7. At very high pH there was both fast net calcification (red) and rapid net dissolution (blue). In some extreme cases, pH\(_T\) was not a good indicator of seawater saturation state (\(\Omega_a\)). A selected dataset of diurnal low-tide emersion periods from September 2020. Colors represent NCC (mmol CaCO\(_3\) m\(^{-2}\) h\(^{-1}\), as presented in Fig. 3c). Dashed horizontal lines represent saturation state for aragonite (\(\Omega_a = 1\)) and for high-Mg calcite (\(\Omega_a = 2\)).

needed to operate CCMs rather than directly related to CO\(_2\) gradients or higher substrate availability.

In winter and summer, pools under present-day and future conditions were autotrophic at emersion (NCP\(_D\) > CR\(_N\); Fig. 5). If we consider the CPB to be integrated diurnal NCP and nocturnal CR over 24 h (assuming equal day–night duration), this means that the pools always fixed more carbon during the day than they respired at night at emersion (NCP \(\gg\) CR), regardless of treatment. One methodological uncertainty we highlight regarding net production is that diurnal DIC-derived NCP estimations were 50 % higher than O\(_2\)-derived NCP estimates (NCP\(_{DIC}\) = 1.6 ± 0.05 NCP\(_O2\) by day, \(R^2 = 75\%\); Figs. 3c and 4). This discrepancy was far less apparent during nights, when methods agreed on respiration rates (CR\(_{DIC}\) = 1.0 ± 0.09 CR\(_O2\), \(R^2 = 56\%\)). While O\(_2\)-derived NCP appears accurate during the night, O\(_2\) production during the day is likely to have been underestimated due to degassing (e.g., visible formation of oxygen bubbles at the surface of algae, > 150 % air saturation by day vs. < 100 % at night). Thus, estimating diurnal net production using oxygen measurements may not be appropriate in algaedominated environments such as these tidal pools. Nevertheless, despite the difference in absolute NCP estimates, both approaches indicate a diurnal fertilization effect.

4.2 Nocturnal dissolution under CO\(_2\) addition

In the present study, natural mesocosms – temperate coralline-dominated tidal pools – were used to investigate the effect of ocean acidification on net calcification at the community level. As we observed a fertilization effect of CO\(_2\) addition by day, we could have expected that it would also enhance diurnal calcification, as photosynthesis and calcification are tightly linked (Martin et al., 2013a, b; Williamson et al., 2017), but this was not observed. Treatment had no significant effect on the daytime net calcification rates, and diurnal variability in calcification appears to be predominately driven by PAR, temperature and metabolic activity (NCP). Increasing metabolic rates – in turn increasing calcification rates – however may have counterbalanced any calcification suppression or increased dissolution due to acidification, making its effect invisible. Noisette et al. (2013) similarly found no effect of \(pCO_2\) treatment on light calcification for E. elongata. However, the authors reported a significant de-
increase in light calcification in *L. incrustans*; net calcification even switched to net dissolution in 750 and 1000 µatm* p*CO₂ treatments. While our future treatments started at *p*CO₂ levels higher than 1000 µatm, the fact that *CO₂* addition did not influence diurnal calcification could also be due to favorable saturation state conditions in the micro-environment in which calcification occurs. The diffusive boundary layer (DBL) can enhance CaCO₃ precipitation micro-environment conditions due to the uptake of *CO₂* OR DIC for photosynthesis. For instance, under light conditions, CCA surface pH has been shown to reach as high as 8.6 (Houlihan et al., 2020) in surrounding seawater at pH 7.7 (+1.1 pH units), which would be highly favorable to calcification. Although there are conflicting results indicating that saturation state of the ambient seawater is a key driver of coralline algal calcification, the biomineralization process in coralline algae has also been shown to present a certain degree of biological control (de Carvalho et al., 2017; Nash et al., 2019). Recent work using boron isotopes (δ11B) as a proxy for pH showed that coralline algae have the ability to elevate pH at their site of calcification (Cornwall et al., 2017). But more complex interactions may also be at work; e.g., CCA may use increases in HCO₃⁻ (due to *CO₂* dissolution) to calcify, making them more resistant to ocean acidification, as suggested by Comeau et al. (2013).

There was net CaCO₃ dissolution in the pools at night (−0.7 mmol CaCO₃ m⁻² h⁻¹), even when waters were still supersaturated with regard to aragonite under present-day conditions. Night dissolution may be a sign that the DBL of the calcifiers inhabiting the pools is undersaturated, possibly as a result of respiration. Indeed, Houlihan et al. (2020) observed that nocturnal algal respiration by CCA increased *CO₂* in the DBL, decreasing pH of the DBL by 0.1 units. Such a small pH decrease alone is unlikely to explain an undersaturation of the calcifying environment as aragonite saturation state was still above 1.2 under most of the present-day conditions. However, given that the solubility of high-Mg calcite – the mineral composing *L. incrustans* and *E. elongata* in particular (Ries, 2011) – can be twice that of aragonite (Sulpis et al., 2021; Yamamoto et al., 2012), it is possible that undersaturation already occurs at night for this mineral even for *Ω₃* > 1. Another reason for night dissolution might be linked to the patellid limpet, an opportunist (Schaal and Grall, 2015) and dominant grazer in rock pools that can be particularly active at night (Lorenzen, 2007). Encrusting coralline algae can be an important food source for these herbivores, and the large percentage of grazed coralline algal CaCO₃ in their gut (Maneveldt et al., 2006) could dissolve easily at night.

Adding *CO₂* (from 445 to 1500–2000 µatm) at the start of emersion significantly increased net dissolution (NCC₀), by 40% in summer and 70% in winter. In a previous single-species experiment, Noisette et al. (2013) demonstrated that, for *L. incrustans* from an area close to our site, dark dissolution doubled with increasing *p*CO₂ (1000 µatm vs. 380 µatm), unlike *E. elongata*, for which there was no effect of *p*CO₂: the ACA even calcified in the dark up to 750 µatm (see also similar results from Egilsdottir et al., 2013). Since *L. incrustans* is the major calcifying species of the tidal pools we studied, it is likely that this species drives the results we observed at the pool community scale. Regardless of the treatment, nocturnal net dissolution rates (NCC) were also significantly correlated with *Ω₄*: the results are similar to those found by Kwiatkowski et al. (2016) in temperate tidal pools of California, without *CO₂* addition.

In summer, pools under present-day and future conditions were precipitative (CCB > 0), meaning that diurnal net calcification exceeded nocturnal net dissolution, regardless of treatment. Adding *CO₂* in summer did not consistently change CCB, with most pools showing little change in CCB due to treatment. By contrast, in the colder winter, the calcification budget was at least 50% lower than in summer (present), with some pools having comparable net calcification during the day to net dissolution at night. During this season, adding *CO₂* had variable impacts on CCB, decreasing it in three of the pools by more than 80% and increasing relatively small CCB in two. These variable effects may be due to differences in community composition and highlight the difficulty in generalizing the results of natural mesocosm manipulations in which the initial community composition is not controlled. Nevertheless, we expected *CO₂* addition to have a greater negative effect in winter (more dissolution) than in summer, with saturation states being lower due to colder temperatures, making it more of a “crucial”/“bottleneck” season. This emphasizes the need to study the effect of ocean acidification across seasons and temperature ranges, especially given the associated changes in algal community composition and metabolic activity.

A limit of the CCB in the current study is that we only considered the tidal pools to be closed (emersed) systems. However, in an acidifying ocean, tidal pool communities are submersed for nearly 12 h d⁻¹, resulting in long exposure to low pH. More accurate and realistic budgets would need to integrate these immersion periods, which might have additive negative effects on calcification (see, e.g., Legrand et al., 2018b, for tidal assemblage experiments on net production/respiration).

### 4.3 Instances of aragonite undersaturation at high pH

An unexpected phenomenon happened in Pools C and E in summer: although we measured very high pH values, we observed that total alkalinity suddenly increased, a sign of fast net dissolution. When we then computed the carbonate chemistry, the saturation states were surprisingly low (*Ω₃* = 0 for pH₇ = 10), which was due to near-zero DIC concentrations – and thus near-zero CO₃²⁻ concentrations. Under these particular conditions, which occurred towards the end of the tidal emersion period, any CaCO₃ precipitation was less than dissolution; precipitation may even have
been impossible due to a lack of DIC substrate. In intertidal pools with a high density of *Zostera marina*, Miller and Kelley (2021) observed a similar decoupling between pH and $\Omega_a$, with increases in pH not leading to an increase in saturation state at high pH values due to a lack of DIC OR CO$_3^{2-}$. In our study, we observed even more drastic decoupling between expected changes in pH, $\Omega_a$, and NCC, with some of the fastest net dissolution rates observed at very high pH and very low $\Omega_a$ values that were a consequence of near-complete consumption of DIC by community production (Fig. 7). Macroalgae cultivation has been proposed as a method of bioremediation for local acidification, in particular to improve aquaculture environments (e.g., Bergstrom et al., 2019; Gao and Beardall, 2022): an increase in algal or marine plant cover would reverse or buffer the negative effects of acidification on heterotroph calcifiers. Our results and those of Miller and Kelley (2021) suggest that phytoremediation should not consider pH to be the sole indicator for “acidification remediation” and that periodical decreases in saturation state in macroalgae- or seaweed-dominated environments in summer (and during marine heat waves) may need to be considered for these proposed types of remediations.

5 Conclusion

Relative to its area, human societies are disproportionately reliant on the coastal ocean for the provision of natural resources and climate regulation. Yet our understanding of how anthropogenic carbon emissions and associated ocean acidification will influence natural coastal ecosystems and community metabolism remains limited. In the present study, we manipulated the carbonate chemistry of natural temperate intertidal pools to explore the potential impact of future ocean acidification on community-level calcification and production. We find evidence of large seasonal, diel and community-specific differences in the sensitivity of intertidal community metabolism to acidification. Diurnally, acidification was found to enhance net community production, with this fertilization effect indicating that algal photosynthesis is naturally carbon-limited in such environments at emergence. Diurnal net community calcification was unaffected by acidification. In contrast, nocturnal acidification resulted in greater net community dissolution in the intertidal pools yet had no consistent effect on community respiration. Integrated over day/night emersion periods, the intertidal mesocosms maintained positive net community calcification and production under both present-day and future conditions. Despite considerable differences between individual pools and strong seasonal dependencies, our results indicate that the net calcification and production of temperate intertidal communities – likely acclimated/adapted to variable conditions – could be affected by future acidification.

Data availability. Raw data and linear regression model results are provided in the Supplement.

Supplement. The supplement related to this article is available online at: https://doi.org/10.5194/bg-20-4289-2023-supplement.

Author contributions. ND, SM and LK designed the experiments, and ND carried them out with help from all co-authors. ND analyzed the data and prepared the manuscript, with contributions from all co-authors.

Competing interests. The contact author has declared that none of the authors has any competing interests.

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