

1 **The Bouraké semi-enclosed lagoon (New Caledonia). A natural**
2 **laboratory to study the life-long adaptation of a coral reef ecosystem**
3 **to extreme environmental conditions**
4

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14 **Key words:** natural laboratory, warming, climate change, ocean acidification, deoxygenation, benthic communities, coral
15 reefs, mangroves

16 **Abstract.** According to current experimental evidence, coral reefs could disappear within the century if CO₂ emissions remain
17 unabated. However, recent discoveries of diverse and high cover reefs that already live under extreme conditions suggest that
18 some corals might thrive well under hot, high pCO₂, and deoxygenated seawater. Volcanic CO₂ vents, semi-enclosed lagoons
19 and mangrove estuaries are unique study sites where one or more ecologically relevant parameters for life in the oceans are
20 close or even worse than currently projected for the year 2100. Although they do not perfectly mimic future conditions, these
21 natural laboratories offer unique opportunities to explore the mechanisms that reef species could use to keep pace with climate
22 change. To achieve this, it is essential to characterize their environment as a whole, and accurately consider all possible
23 environmental factors that may differ from what is expected in the future, possibly altering the ecosystem response.
24 This study focuses on the semi-enclosed lagoon of Bouraké (New Caledonia, SW Pacific Ocean) where a healthy reef
25 ecosystem thrives in warm, acidified and deoxygenated water. We used a multi-scale approach to characterize the main
26 physical-chemical parameters and mapped the benthic community composition (i.e., corals, sponges, and macroalgae). The
27 data revealed that most physical and chemical parameters are regulated by the tide, strongly fluctuate 3 to 4 times a day, and
28 are entirely predictable. The seawater pH and dissolved oxygen decrease during falling tide and reach extreme low values at
29 low tide (7.2 pH_T and 1.9 mg O₂ L⁻¹ at Bouraké, vs 7.9 pH_T and 5.5 mg O₂ L⁻¹ at reference reefs). Dissolved oxygen,
30 temperature, and pH fluctuates according to the tide of up to 4.91 mg O₂ L⁻¹, 6.50 °C, and 0.69 pH_T units on a single day.
31 Furthermore, the concentration of most of the chemical parameters was one to five times higher at the Bouraké lagoon,
32 particularly for organic and inorganic carbon and nitrogen, but also for some nutrients, notably silicates. Surprisingly, despite
33 extreme environmental conditions and altered seawater chemical composition measured at Bouraké, our results reveal a
34 diverse and high cover community of macroalgae, sponges and corals accounting for 28, 11 and 66 species, respectively. Both
35 environmental variability and nutrient imbalance might contribute to their survival under such extreme environmental
36 conditions. We describe the natural dynamics of the Bouraké ecosystem and its relevance as a natural laboratory to investigate
37 the benthic organism's adaptive responses to multiple extreme environmental conditions.

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39 **Copyright statement:** all figures used in this manuscript belong to authors and have never been published before. The satellite
40 image in Fig. 1 and 10 is from georep New Caledonia database (<https://georep.nc/>), and the QGIS software was used for
41 transect georeferencing.

42

43 **1 Introduction**

44 Atmospheric carbon dioxide (CO₂) has steadily increased over the industrial period (Gattuso et al., 2015), leading to ocean
45 warming, acidification and deoxygenation. Although the extent to which these stressors will affect marine life is still debated,
46 there is no doubt that their combination will negatively affect a range of marine organisms (e.g., Kroeker et al., 2011;
47 Wittmann and Pörtner, 2013; Hughes et al., 2018). Coral reefs are among the most productive and biodiverse marine
48 ecosystems on Earth. Their survival is expected to be compromised by climate change, whose impacts on reef structures and
49 associated communities span from biodiversity loss to ecosystem change shift (e.g., Fabricius et al., 2013; Sunday et al., 2017;
50 Agostini et al., 2018).

51 Marginal and extreme environments, where some species persist under sub-optimal environmental conditions, have become
52 a precious tool to investigate the potential resilience of marine organisms in the face of climate change (Camp et al., 2017,
53 2018). These sites may be used as natural laboratories where at least one or more environmental parameters naturally mimic
54 extreme environmental conditions over a large area of the ecosystem. They provide an opportunity to simultaneously
55 investigate changes in species responses and their ability to acclimatize and adapt to global environmental changes (Soares,
56 2020; Kurihara et al., 2020). Shallow-water volcanic CO₂ seeps, low pH springs, semi-enclosed bays, mangrove habitats,
57 shallow sheltered-bay reefs, macrotidal environments, and low-pH upwelling areas are all potential study systems where the
58 surrounding seawater is subject to a localized or widespread increase in either *p*CO₂, or temperature, and eventually a decrease
59 in dissolved oxygen (Camp et al., 2018). At these sites, general observation suggest ecosystem-level consequences of life-
60 long exposure to extreme conditions, such as reduced biological diversity, especially among calcifying organisms, decreased
61 rates of coral calcification, and high rates of bioerosion (e.g., Hall-Spencer et al., 2008; Manzello et al., 2008; Fabricius et al.,
62 2011; Crook et al., 2013; Kroeker et al., 2011; Iglesias-Prieto et al., 2014; Milazzo et al., 2014; Paytan et al., 2014). However,
63 some natural laboratories can host very rich reef communities. Examples of such sites have been documented in Palau (Golbuu
64 et al., 2016; Barkley et al., 2017; Shamberger et al., 2018; Kurihara et al., 2021), Papua New Guinea (Pichler et al., 2019),
65 the Kimberly region, Australia (Dandan et al., 2015; Schoepf et al., 2015), and in mangrove lagoons of New Caledonia (Camp
66 et al., 2017) and the US Virgin Islands (Yates et al., 2014).

67 These natural laboratories have become a common experimental asset in climate change research. However, the lack of
68 empirical characterization of the physical and biogeochemical conditions, including diurnal and seasonal fluctuations, have
69 been argued to bias the interpretation of the biological mechanisms that trigger the responses of organisms (e.g., Vizzini et
70 al., 2013; Camp et al., 2018; Aiuppa et al., 2021). Using limited environmental descriptors makes it difficult to unequivocally
71 identify the main driver(s) of the biological response among the primary factors (i.e., acidification, warming and/or
72 deoxygenation), the potential secondary factors (e.g., pollution, water flow, tide, seawater nutrients and organic content,
73 turbidity, etc.), and their combination. For instance, at CO₂ seeps, pH variability can unexpectedly go beyond projected future
74 values (e.g., Hall-Spencer et al., 2008; Kerrison et al., 2011) and the potential emission of toxic compounds, such as sulphur,
75 arsenic, and metal trace elements (Vizzini et al., 2013) compromises the attribution of specific responses to ocean
76 acidification. Water temperature, pH and dissolved oxygen can also co-vary negatively or positively and combine with other
77 secondary factors, acting synergistically or antagonistically with unknown effects on benthic community responses. Their
78 extreme values and the extent to which organisms are exposed are crucial in shaping biological responses (Boyd et al., 2016;
79 Rivest et al., 2017). For instance, early studies suggest that temperature fluctuations due to diel or tidal variations could expose
80 corals to stressful temperatures long enough to induce acclimatization or adaptation, but short enough to avoid coral mortality

81 (Craig et al., 2001; Oliver and Palumbi, 2011; Castillo et al., 2012; Palumbi et al., 2014; DeCarlo et al., 2019). Coral reef
82 organisms from such thermally variable environments are expected to respond positively to future heat events (Rivest et al.,
83 2017). Besides, corals naturally subjected to high $p\text{CO}_2$ variability have shown immune defences when experimentally
84 exposed to high temperatures, thus buffering the magnitude of thermal stress during heat waves (Wall et al., 2020). The
85 extreme tidal range in the Kimberly region (Northwest of Australia) exposes corals to short-term temperature maxima of up
86 to 37 °C, and fluctuations of up to 7 °C daily. Despite the high temperature, also combined with strong currents and turbid
87 waters, diverse and probably resilient coral reefs have been described there (Dandan et al., 2015; Schoepf et al., 2015).
88 Overall, only parts of these natural laboratories' physical and biogeochemical parameters have been quantified and only during
89 short periods of fieldwork due to logistic constraints. Long-term monitoring of seawater parameters and their fluctuation is
90 essential to better understand the mechanisms used by resilient reef organisms in these natural laboratories and interpret how
91 marine taxa will respond to future biogeochemical changes in the environment.

92 The semi-enclosed lagoon of Bouraké (New Caledonia, SW Pacific Ocean) has been considered one of the most suitable
93 natural laboratories for future extreme environmental conditions (Camp et al., 2019). In a preliminary study, Camp et al.,
94 (2017) reported a series of compelling short-term data of seawater carbonate chemistry and a general survey of the coral
95 populations. In this first assessment of the Bouraké lagoon, the authors measured diel fluctuations of temperature (ranging
96 from 25.9-33.1 °C), pH (ranging from 7.24-7.91 pH_T units) and dissolved oxygen (DO) (ranging from 1.80-6.97 mg L^{-1})
97 regulated by a one-meter tide. They revealed that about 20 species of corals were exposed 44 % of the time to a pH_T of 7.7-
98 7.8, and 71 % of the time to temperatures predicted for the end of the century under the IPCC scenario RCP4.5 (IPCC, 2014).
99 These striking preliminary findings qualified the Bouraké lagoon as a unique site where potentially adapted corals withstand
100 extreme environmental conditions.

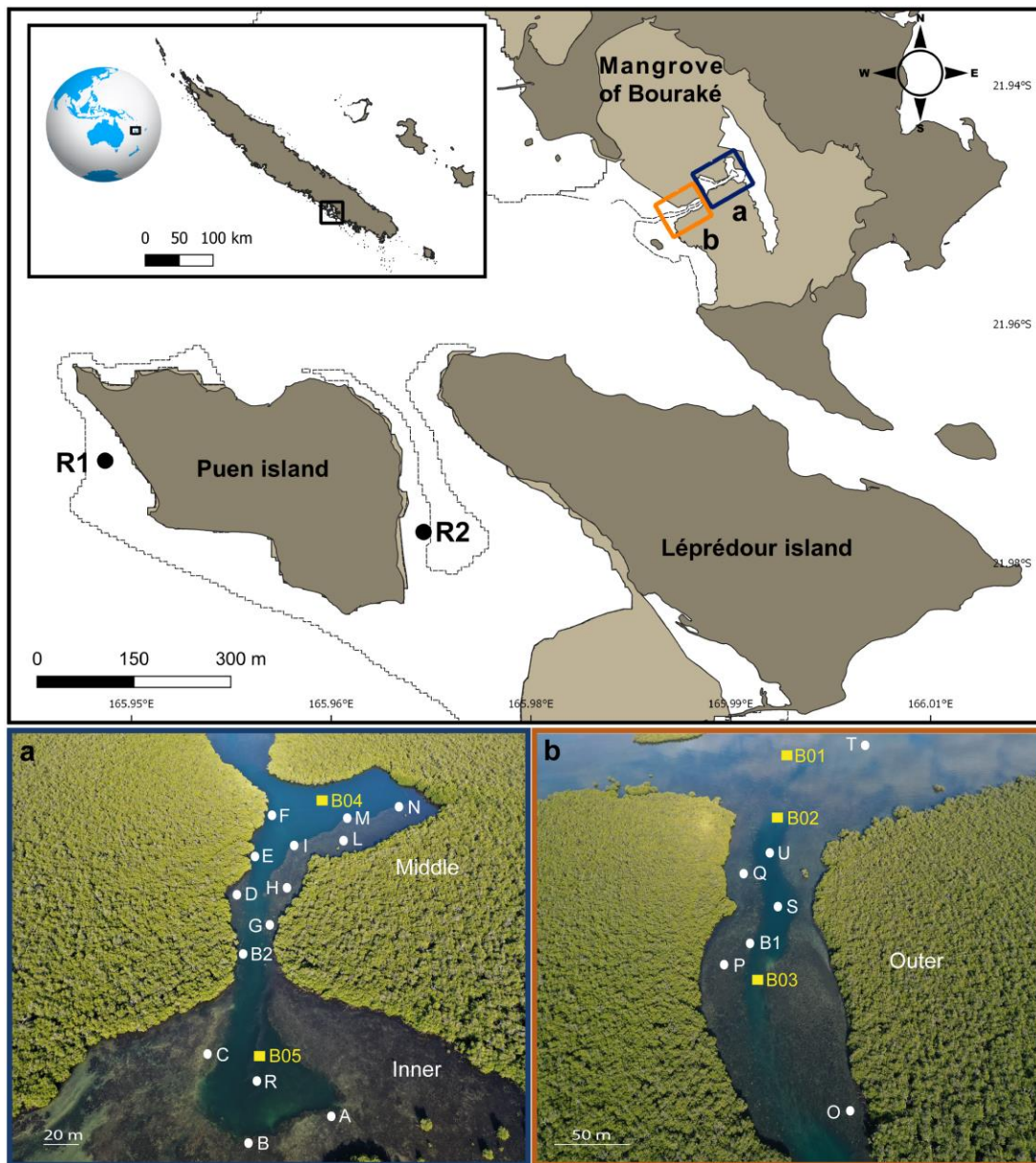
101 The extent to which the Bouraké species are exposed to sub optimal conditions remains unclear. We believe that the best way
102 to assess these very encouraging findings is to start with fully characterizing the main environmental parameters and the daily
103 and seasonal fluctuations to which reef species have been subjected during their entire life. Here we used a multi-scale
104 approach to map and describe the benthic community living in the Bouraké lagoon and report on new evidence based on three
105 years of data. Because of the size of the area, the close vicinity of a dense mangrove forest, and the demonstrated tide effect
106 on the local environmental conditions, we hypothesize that (1) environmental conditions fluctuate regularly but are spatially
107 heterogeneous throughout the Bouraké lagoon, and (2) only a limited number of species can resist to the extreme physical and
108 chemical conditions at the study site, when compared to adjacent bay-sheltered reefs.

109

110 **2 Methods**

111 **2.1 Study sites**

112 Starting in February 2016, we studied the semi-enclosed coral reef lagoon of Bouraké (South Province, Grande Terre, New
113 Caledonia) and adjacent reference reefs (Fig. 1). The mangrove of Bouraké is located in a semi-arid geographical area, and it
114 lacks rivers input typical of mangrove estuaries.



115
116 **Figure 1.** Map of the study site (top panel) showing the semi-enclosed mangrove lagoon of Bouraké and reference reefs (R1
117 and R2). Photos **a** and **b** (bottom panels) are aerial pictures (taken at 130 m above the Bouraké lagoon) of the inner (bottom)
118 and middle reefs (**a**) and the outer reef (**b**). White dots and yellow squares indicate the sampling stations used for physical
119 and chemical environmental monitoring, respectively. Georep New Caledonia database (<https://georep.nc/>) and QGIS software
120 were used to build the figure (top panel).

121 In Bouraké, a channel of more than 80 m wide and 0.5 to 6 m deep penetrates a dense mangrove forest made by *Avicennia*
122 *marina* and *Rhizophora stylosa*. It expands into side pools and a large reservoir in the inner part of the mangrove system. A
123 preliminary calculation of its surface and seawater volume of the main area (i.e., without the large seawater reservoir covered
124 by the mangrove forest) using the 3ds Max Model gives underestimated values of 192,100 m² and 246,060 m³, respectively
125 (Rodolfo-Metalpa et al., in prep.). Terraces extend from the mangrove, on both sides of the channel and form diverse and
126 compact reefs at their ends. Coral species are less abundant and diverse near the mangrove roots, where the bottom substrate
127 is composed of fine sediment and mud. There, reefs are exposed to air only at low tide during the spring tides (1.1 m maximum
128 tidal range). We subdivided the lagoon into three main areas: an external, an intermediate, and an inner reef (hereafter called
129 outer, middle and inner reefs; panels a & b in Fig. 1). A series of sampling stations were selected, and some of the main
130 seawater chemical and physical environmental parameters were measured at the study site and two adjacent reference reefs,
131 namely R1, a typical fringing reef, and R2, a semi-enclosed shallow and relatively large bay. Both reefs were located 4.8 and
132 3.2 km from the entrance of the Bouraké lagoon, respectively.

133 **2.2 Environmental monitoring**

134 **2.2.1. Oceanographic sensors deployment and short- to long-term measurements**

135 From February 2016 to December 2020 up to eight YSI 600 OMS-M, three Seabird SeaFETTM pH loggers, and four Hobo
136 water temperature Pro V2 were set at 10-min logging intervals, and used individually or in combination to assess short-,
137 medium-, or long-term variations across depth (as a proxy of the tide), temperature (°C), dissolved oxygen concentration (DO,
138 mg L⁻¹), seawater pH_T (total scale), and salinity. A summary of the deployments is given in Supplementary Table S1. YSI
139 dissolved oxygen optical sensors were calibrated against zero and 100 % saturated seawater at 25 °C. Two SeaFETs were
140 calibrated by the manufacturer, while the third was corrected before deployment by measuring its deviation from the two
141 others in the same seawater.

142 Short-term (i.e., 24-h) spatial and depth-related variations in pH and DO were simultaneously measured at several
143 stations in the Bouraké lagoon in March and April 2018 (Fig. 1). Spatial variations were assessed i) within the inner (St A-C
144 for pH, and A-G for DO), the middle (St H-N for DO), and the outer (St O-Q for DO) reefs, ii) between the outer and the
145 inner reefs (St B1, B2 for pH), and iii) between the outer and the reef outside the semi-enclosed lagoon (St U vs T for pH). In
146 the Bouraké lagoon, we recorded differences between the surface (shallow) and the bottom water: i) at the outer reef (St S for
147 pH), ii) at the middle reef (St I and N for DO), and iii) at the inner reef (St R for pH, and DO). Salinity was measured in July
148 2019 (St R1, R2, B1 and B2) and December 2020 (St R1, R2, T and B2).

149 Medium-term measurements (i.e., 2-3 weeks) of the DO and pH were recorded at the reference (St R1, R2) and
150 Bouraké reefs (St B1, B2) between 2016 and 2019.

151 Long-term measurements (>1 year) of seawater temperature were recorded at R1 and R2 starting from January 2019
152 and September 2017, respectively, and at B1 and B2 from October 2018 and September 2017, respectively. Only temperature
153 data between October 2018 and April 2020 were compared between sites.

154 Short-term pH and DO data and long-term temperature data were compared between stations using General Linear
155 Modelling (GLM), and the Tukey' HSD *post hoc* test was used when significant factor effects were found. When data did not
156 conform to normality or homogeneity of variance, the Kruskal-Wallis test followed by the Dunn's multiple comparisons test

157 (Bonferroni-adjusted) or the Wilcoxon test were performed. Statistical analyses were carried out using either Statistica® or R
158 version 3.4.4 (R Core Team, 2018), the latter using "stats", "FSA", and "MASS" packages.

159 **2.2.2. Phase averaged and tidal harmonic analyses for diurnal and semidiurnal oscillations**

160 Medium-term pH and DO changes were investigated by averaging time and tidal phases for diurnal and semi-diurnal
161 oscillations. To do this, all data were overlaid on a daily period and a tidal phase. First, we calculated a predicted tide for the
162 study area using the Nouméa harbour tide (50 km south of our study site) modified with coefficients from the Naval
163 Hydrographic and Oceanographic Service (SHOM; <http://data.shom.fr>). The predicted tide was used to assign a semidiurnal
164 tidal phase (12 h) to each sampling time, and the data were averaged for each of these tidal phases. Similarly, the data were
165 averaged for each hour of the day (24 h). Because tides at sea are a sequence of sinusoidal harmonic components that are
166 different for each location, we performed a harmonic tidal analysis on the DO and pH data. We used the "UTide"-`ut_solv()`
167 tidal analysis package (Codiga, 2011) with the principal semidiurnal lunar constituent (M_2), principal semidiurnal solar
168 constituent (S_2), and solar diurnal constituent (S_1). For each parameter, the amplitudes of the tidal harmonics M_2 (12.4 h), S_2
169 (12 h), and S_1 (24 h) were calculated with a 95 % confidence interval based on the 200 Monte-Carlo simulations.

170 **2.2.3. Diel cycles of carbonate chemistry and chemical parameters**

171 Surface water samples were collected across a diel cycle in June 2017, and July 2019 for pH, total alkalinity (A_T), dissolved
172 inorganic carbon (DIC), nutrients (orthosilicic acid [$\text{Si}(\text{OH})_4$], nitrogen oxide [NO_x], ammonium [NH_4^+], phosphate [PO_4^{3-}],
173 dissolved organic carbon (DOC), particulate organic carbon (POC), and particulate organic nitrogen (PON). A summary of
174 the total sampling is given in Supplementary Table S2. Dissolved inorganic nitrogen (DIN) and total organic carbon (TOC)
175 were calculated as $[\text{NO}_x] + [\text{NH}_4^+]$, and $\text{DOC} + \text{POC}$, respectively. The most important ratios were calculated (e.g.,
176 $\text{DOC}:\text{TOC}$; $\text{Si}(\text{OH})_4:\text{DIN}$) and contrasted between stations to evaluate the availability of nutrients and organic matter in the
177 Bouraké lagoon (Jacquet et al., 2006; Leopold et al., 2017).

178 In 2017, during three consecutive days (from May 31st to June 02nd), seawater was sampled six times: twice during
179 both high and low tide and once at both rising and falling tide. In total, we sampled one reference station (R2), three stations
180 at the outer reef of the Bouraké lagoon (Outer: St B01-B03), one at the middle reef (Middle: St B04), and one at the inner reef
181 inside the lagoon (Inner: St B05) (Fig. 1). The whole collection lasted about 30 minutes.

182 In 2019, during three consecutive days (from July 16th to 18th), sampling was carried out every hour from 8 am to 3
183 pm. We sampled B1 and B2 on the first day, R1 on the second and R2 on the third day.

184 During diel cycles, at each station and sampling time, pH and temperature were measured at the surface (0.5 m deep) using a
185 portable pH-meter (913, Metrohm) calibrated with TRIS buffer (Dickson lab, batch #T28). A subsample (50 mL) was filtered
186 through 0.45 μm Whatman™ Puradisc CA filters using a syringe and poisoned™ with 20 μL saturated HgCl_2 to further measure
187 A_T . Two 20 mL subsamples were analysed using an auto titrator (EcoTitrator, Metrohm), and A_T was calculated from the Gran
188 function. Results were corrected against A_T standards (A. Dickson, batch #155, Scripps, USA). The seawater carbonate
189 parameters $p\text{CO}_2$, CO_3^{2-} , and aragonite saturation state (Ω_{ara}) were then calculated from the pH_T , A_T , temperature, and mean
190 salinity (35) using the free-access CO_2SYS package (Pierrot et al., 2006).

191 Ammonium concentration was determined on a 40 mL subsample of unfiltered seawater, collected using a 60 mL
192 Schott bottle and stored in the dark. Samples were processed using a fluorimeter (Turner Designs) between six and 18 h after
193 two mL of OPA reagent (o-phthaldialdehyde) was added (Holmes et al., 1999).

194 The sampling of nutrients was performed using two replicate 20 mL polypropylene vials, rinsed three times using
195 filtered seawater (Whatman™ Puradisc CA syringe filters 0.45 µm), filled with the sample and immediately poisoned with
196 20 µL saturated HgCl₂. Measurements of PO₄³⁻, NO_x, and Si(OH)₄ nutrients were performed by colourimetry (Seal
197 Analytical).

198 Seawater samples for DIC were collected in two replicate glass vials (20 mL), filled with unfiltered water and
199 poisoned with 10 µL saturated HgCl₂. The vials were immediately closed, the absence of bubbles was visually checked, and
200 the samples were stored in the dark at room temperature for later analysis on a Shimadzu TOC-L analyser (Non-Dispersive
201 Infrared, NDIR). Typical analytical precision was less than ± 2 µmol kg⁻¹. The accuracy was verified using regular
202 measurements of reference material (CRM) from A. Dickson's laboratory.

203 Seawater samples for DOC were collected in two pre-combusted (4h at 450 °C) glass ampoules filled with water
204 filtered using a glass syringe filtration system (SGE™) with two pre-combusted 25-mm GF/F Whatman™ filters. Samples
205 were then acidified with ultrapure orthophosphoric acid (H₃PO₄), sealed, and stored in the dark at room temperature for later
206 analysis by high-temperature catalytic oxidation (HTCO) (Sugimura and Suzuki, 1988; Cauwet, 1994) on a Shimadzu TOC-
207 L analyser. Typical analytical precision was ± 0.1-0.5 µM C (SD). Consensus reference materials
208 (<http://www.rsmas.miami.edu/groups/biogeochem/CRM.html>) were injected every 12 to 17 samples to ensure stable
209 operating conditions. DOC concentrations are only available for the 2017 sampling because of a sample's pollution in 2019.

210 Finally, one litre of unfiltered seawater was collected in a borosilicate glass bottle and stored on ice during sampling
211 for later measurement of POC and PON contents. In the lab, particulate matter was collected on pre-combusted (4h at 450
212 °C) Whatman™ GF/F filters using a Nalgene® vacuum system. The filters were dried at 60 °C in the oven for 24 h and stored
213 in airtight glass vials at 4 °C in the dark until analysis on a CHN Perkin Elmer 2400.

214 All glass bottles and vials used were pre-combusted, washed with HCl solutions (10 %) and rinsed using milliQ water.
215 Seawater chemistry data were pooled by sampling area (R1, R2, Outer, Middle, Inner), and differences were tested using the
216 Kruskal-Wallis test followed the Conover Multiple comparison test (Benjamini-Hochberg-adjusted). We focused on the effect
217 of the tidal phases (i.e., falling and rising tide) on the seawater chemical composition in the Bouraké lagoon only, by attributing
218 each sample a tidal phase between 0 (high tide) and 6 hours (low tide), and between 6 and 12 hours (high tide). Multiple linear
219 regression was used to assess the adjusted R² and significance ($p < 0.05$) of the data from 0 to 6 h (falling tide) and from 6 to
220 12 hours (rising tide) separately. Statistical analyses were performed using either Statistica® or R (version 3.2.4, R Core
221 Team, 2018), the latter using the "FSA", "stats", and "Conover.test" packages.

222 **2.3 Benthic community characterization and distribution**

223 The benthic community and bottom substrate of the Bouraké lagoon, referred hereafter as biotic and abiotic descriptors,
224 respectively, were assessed in April 2018. Twenty-four 30 m-long geo-referenced transects (T1-T24) were laid in the lagoon
225 along the terraces' edge at similar depths (i.e. ~ 1 m), targeting coral dominated benthic assemblages. On each transect, a 0.5
226 x 0.5 m PVC quadrat was placed every meter, and a picture was taken with a waterproof photo-camera (Nikon AW 130)
227 parallel to the substrate. We made a general description of the bottom (i.e. the various substrates) and a list of the most

228 common and identifiable sessile species for each transect. For each of the 835 pictures collected, we estimated the cover of
229 abiotic (i.e., mud, sand, rock, rubble, dead corals and unreadable) and biotic descriptors (i.e., branching, massive and soft
230 corals, sponges, macroalgae and "others") with photoQuad software both by automatic multi-scale image segmentation
231 regions and manual grid cell counts when necessary.

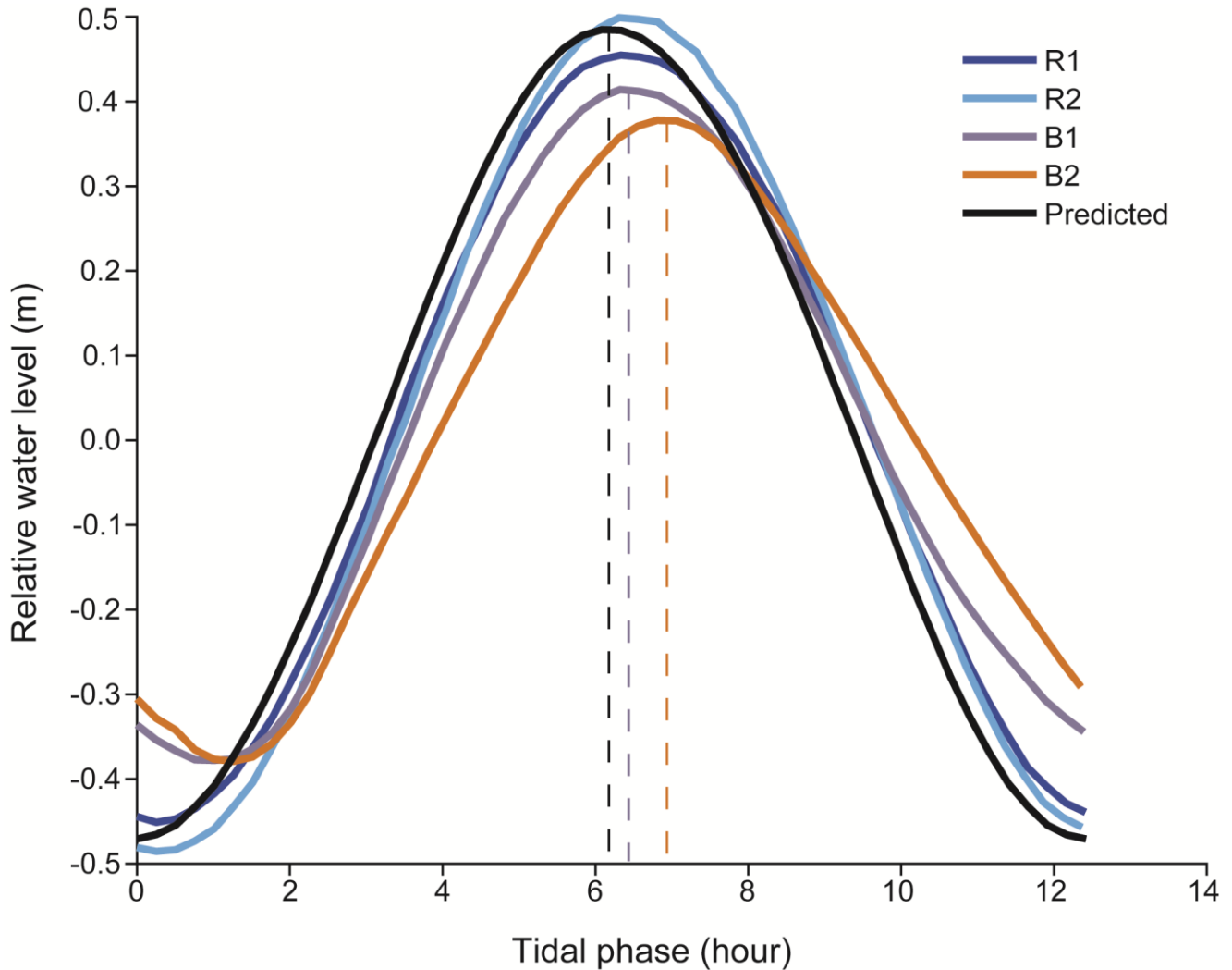
232 We used the photos of quadrats, the many other pictures collected during fieldwork, and laboratory morphological
233 observations on collected samples to produce a non-exhaustive species list of corals, macroalgae and sponges. Corals were
234 comprehensively sampled throughout the Bouraké lagoon and on the reference reef R2, while dominant macroalgal and
235 sponges species were collected in the Bouraké lagoon alone and only if they were encountered at least three times along a
236 transect, likely leading to an underestimation of their diversity. Coral diversity was assessed through photographic and
237 sampling during time-based open search swims and SCUBA dives (Hill and Wilkinson, 2004). Whenever possible,
238 scleractinian corals were identified to species level *in situ* by photographic sampling only. When identification was doubtful
239 *in vivo*, or when taxa were characterized by small corallite size (<1 mm in diameter) and required additional morphological
240 examination of the skeleton to confirm identification (e.g. genera *Acropora*, *Montipora* and *Porites*), a fragment of the colony
241 was collected, tagged, cleaned in sodium hypochlorite overnight, rinsed in freshwater and dried. The reference collection is
242 housed at IRD Nouméa. Microscopic examination of the skeletal features allowed species-level identification following the
243 reference literature (Veron, and Wallace, 1984; Wallace, 1999; Veron, 2000). Sponges were identified based on their spicules'
244 morphological characteristics (i.e., shape, length and width), or using a series of morphological descriptors (e.g., shape, size,
245 colour, texture, surface ornamentations, fibres) for species without spicules. In the lab, a subsample of the collected sponges
246 was immediately digested using HNO₃, and spicules were measured with an optical stereomicroscope. For species without
247 spicules, hand-cut sections of the choanosome and ectosome were observed under a stereomicroscope. Species were identified
248 using the taxonomical keys of Rützler (1978), Hooper and Van Soest (2002), and Pons et al. (2017). Macroalgae and corals
249 identification were based on morphological and anatomical observations following the dedicated literature and referring to
250 specimens housed at IRD Noumea.

251 Abiotic and biotic cover percentages averaged per transect, and species richness, calculated as the number of species
252 in the transect, were plotted using non-metric multidimensional scaling (nMDS) based on Bray-Curtis dissimilarities ("vegan"
253 package in R) of square-root transformed data. Finally, the best number of clusters for the whole Bouraké lagoon was
254 determined using the gap statistic method ("cluster" and "factorextra" packages in R) and used for the hierarchical clustering
255 representation (Ward, 1963). The cluster separation was verified with a two-way analysis of similarity ANOSIM. Within each
256 cluster, the benthic community and bottom substrate were averaged between transects, and the dominant biotic and abiotic
257 descriptors were selected and used to define the cluster.

258 Principal component analysis (PCA) was used to visualize the correlation between environmental parameters and
259 the benthic descriptors of transects inside the Bouraké lagoon. The analysis was performed in R (package "FactorMineR",
260 version 3.2.4, R Core Team, 2018) using data of biotic descriptors (averaged per transect) and the seawater parameters
261 averaged for each sampling area (i.e., outer, middle and inner reefs).

262

264 3.1 Variability of physical environmental parameters

265 *Tidal phases.* The predicted tide in the study area (SHOM data) is close to what we measured at R1 and R2 (Fig. 2).

266

267 **Figure 2.** Temporal shift between the averaged tides recorded at the reference (St R1 and R2) and at the Bouraké reefs (St B1
 268 and B2) and the predicted tide calculated for the study area (see SHOM: <http://data.shom.fr> for the Nouméa harbour).

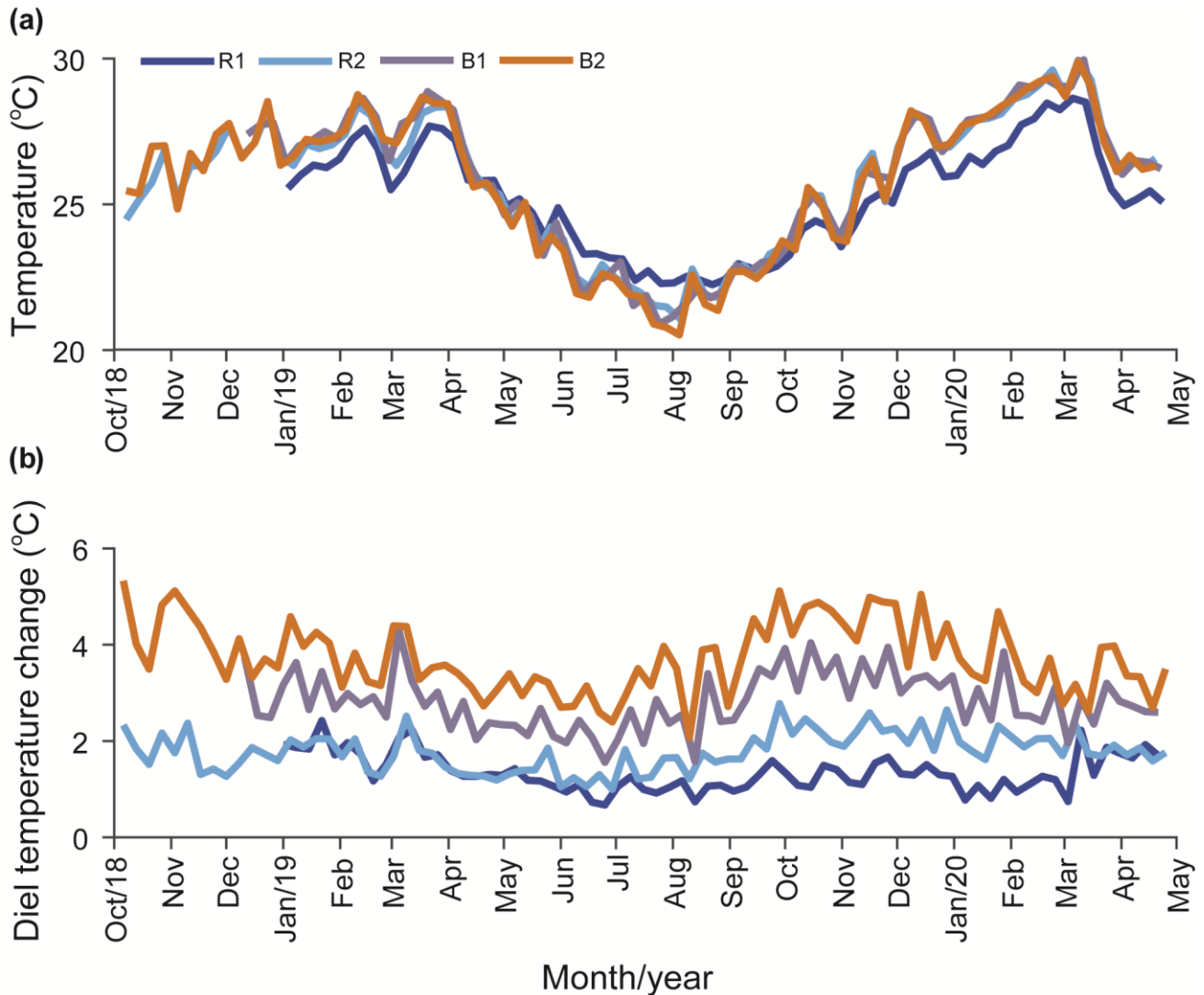
269

270 It is semidiurnal with a diurnal inequality ($M_2 = 0.44$ m, $S_2 = 0.16$ m) and has a small diurnal component ($S_1 = 0.15$ m). It
 271 varies between 0.4 and 1.7 m, depending on the phase of the moon, with a mean of 1.1 m. The tidal range inside the Bouraké
 272 lagoon is lower and varies between 0.4 and 1.2 m, with a mean of 0.9 m. At B1 and B2, the tidal signal lags the predicted and
 273 measured tide at the reference stations by an average of 1.5 h for low tide and up to 45 min for high tide.

274

275 *Temperature.* Our results indicate that the water temperature has an annual cycle with lower values during winter
 276 (May-October) and higher values in summer (November-April). Daily averaged temperatures at R2, B1 and B2 were not
 significantly different, while the temperature at R1 was cooler in summer and warmer in winter (Fig. 3a; Tables 1) compared

277 to the other stations. On a weekly basis, the averaged diel temperature variation was significantly different between stations:
 278 1.34 ± 0.39 °C at R1 and 3.73 ± 0.74 °C at B2 (Fig. 3b; Tables 1). In a single day, we recorded temperature fluctuations of up
 279 to 3.4 °C at R1 and 6.5 °C at B2.
 280



281
 282 **Figure 3.** Long-term temperature monitoring from October 2018 to April 2020 at the reference (St R1 and R2) and Bouraké
 283 reefs (St B1 and B2). Data are plotted using weekly averaged temperature (a) and weekly averaged diel changes (b).

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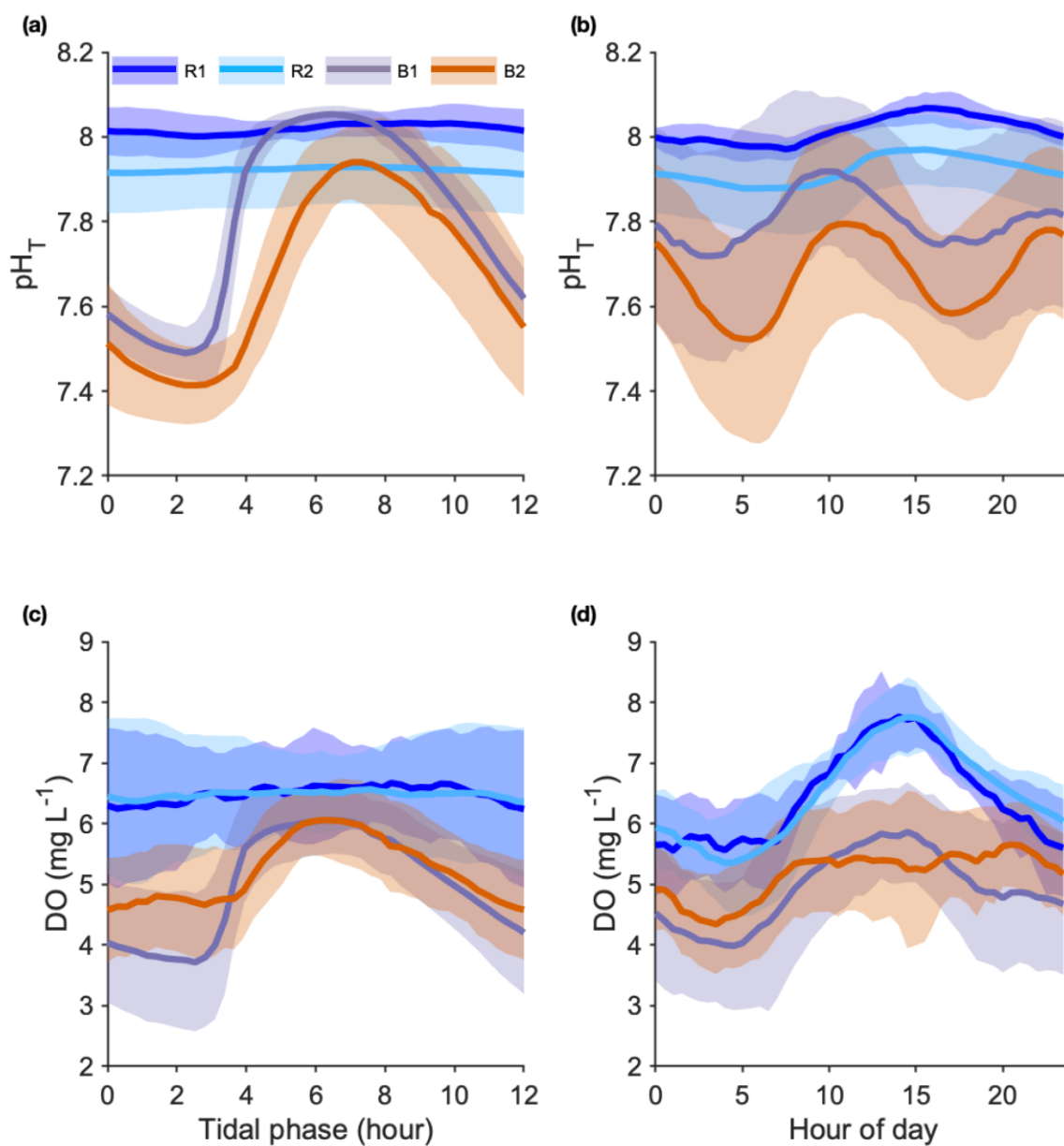
289 **Table 1.** Summary of the statistical analyses applied to the seawater physical and chemical parameters collected between
 290 2016 and 2020 at the Bouraké lagoon (St B1, B2, St A-R, outer, middle and inner) and at reference reefs (St R1 and R2).
 291 Differences in temperature, salinity and chemical parameters between the Bouraké lagoon and reference reefs and spatial and
 292 vertical differences in pH and DO inside the Bouraké lagoon. **K-W:** Kruskal-Wallis's test; **W:** Wilcoxon test; **GLM:** General
 293 Linear Model; **D:** Dunn's test; **T:** Tukey's test; **C:** Conover's test.

Parameters	(n)	K-W	W	GLM	p	post hoc (test)
<i>Seawater temperature (St R1, R2, B1, B2)</i>						
Daily means	2968	68.46			< 0.001	(D) $p < 0.001$ R1 < R2 = B1 = B2
Weekly oscillations	305	234.63			< 0.001	(D) $p < 0.01$ R1 < R2 < B1 < B2
<i>Seawater pH (Stations)</i>						
Spatial INNER (A-C)	711			5.32	0.005	(T) $p < 0.01$ A < C
Spatial (B1 vs B2)	1820			17.62	< 0.001	
Spatial (U vs T)	7280			189.8	< 0.001	
Surface vs deep (R)	664			26.88	< 0.001	
Surface vs deep (S)	692			41.48	< 0.001	
<i>Seawater DO (Stations)</i>						
Spatial INNER (A-G)	1031			1.145	0.330	
Spatial MIDDLE (H-N)	700			0.636	0.636	
Spatial OUTER (O-Q)	399			1.636	0.196	
Surface vs deep (I)	280			0.000	0.994	
Surface vs deep (N)	280			22.05	< 0.001	
Surface vs deep (R)	258			0.443	0.839	
<i>Seawater Salinity (Stations)</i>						
Winter (R1, R2, B1, B2)	1732			463.5	< 0.001	(T) $p < 0.01$ R1 < R2 < B1 < B2
Summer (R1, R2, T, B2)	5105			1583	< 0.001	(T) $p < 0.01$ R1 < R2 < T < B2
Winter vs Summer (R1)	1872		287264		< 0.001	
Winter vs Summer (R2)	1728		253060		< 0.001	
Winter vs Summer (B2)	1872		150175		< 0.001	
<i>Seawater carbonate chemistry (St R1, R2, outer, middle, and inner)</i>						
pH _T	66	36.48			< 0.001	(C) $p < 0.05$ R1 > R2 > Inn = Mid = Out
pCO ₂	66	36.48			< 0.001	(C) $p < 0.01$ R1 = R2 < Inn = Mid = Out
CO ₃ ²⁻	66	33.54			< 0.001	(C) $p < 0.05$ R1 > R2 > Inn = Mid = Out
Ω _{ara}	66	33.27			< 0.001	(C) $p < 0.05$ R1 > R2 > Inn = Mid = Out
DIC	66	25.95			< 0.001	(C) $p < 0.01$ R1 = R2 < Inn = Mid = Out
A _T	66	20.94			< 0.001	(C) $p < 0.05$ All ≠; Mid = R1; Mid = R2; Out = Inn; R1 = R2
<i>Seawater chemical parameters (St R1, R2, outer, middle, inner)</i>						
SiOH ₄	66	45.37			< 0.001	(C) $p < 0.05$ R1 < R2 < Out = Mid = Inn; Inn > Out
NO _x	66	26.46			< 0.001	(C) $p < 0.05$; R1 < R2 = Inn = Mid; Inn < Out; Out > R1; Out > R2
NH ₄ ⁺	66	26.46			< 0.001	(C) $p < 0.01$ R1 < Inn = Mid = Out = R2
PO ₄ ³⁻	66	29.43			< 0.001	(C) $p < 0.05$ All ≠; R2 = Mid
DOC	36	16.12			0.001	(C) $p < 0.05$ All ≠; Mid = Out
POC	66	52.46			< 0.001	(C) $p < 0.05$ R1 = R2 < Out < Mid < Inn
PON	66	43.79			< 0.001	(C) $p < 0.05$ R1 = R2 < Out < Mid < Inn
DIN	66	20.13			< 0.001	(C) $p < 0.01$ R2 < R1 < Inn = Mid = Out
TOC	36	19.48			< 0.001	(C) $p < 0.01$ All ≠; Mid = Out
DIC:DIN	66	17.40			< 0.001	(C) $p < 0.01$ R1 > R2 = Inn = Mid = Out
POC:PON	66	44.87			< 0.001	(C) $p < 0.01$ R1 = R2 < Inn = Mid = Out
POC:DOC	36	18.47			< 0.001	(C) $p < 0.05$ All ≠; Inn = Mid

(continue Table 1)

343					
344					
345	DOC TOC	36	18.21	< 0.001	(C) $p < 0.01$ $R2 < Out < Inn < Mid$;
346					Inn=Mid; Inn=Out
347	Si(OH) ₄ :DIN	66	43.77	< 0.001	(C) $p < 0.01$ $R2 < Inn > Mid$;
348					Mid=Out; $R1 = R2$
349	DIN:DIP	65	24.59	< 0.001	(C) $p < 0.05$ $R1 < R2 > Inn$;
350					$R2 = Out = Mid$; Inn=Mid
351	Si(OH) ₄ :DIP	65	36.62	< 0.001	(C) $p < 0.01$
352					$R1 < R2 < Inn = Mid = Out$;
353					

354 *Seawater pH.* During the entire study period, pH was measured during 22, 72, 31, and 72 semidiurnal tidal cycles at
355 R1, R2, B1, and B2, respectively. We overlaid all data on a single tidal phase of 12 h (Fig. 4a) and a 24-hours cycle (Fig. 4b).
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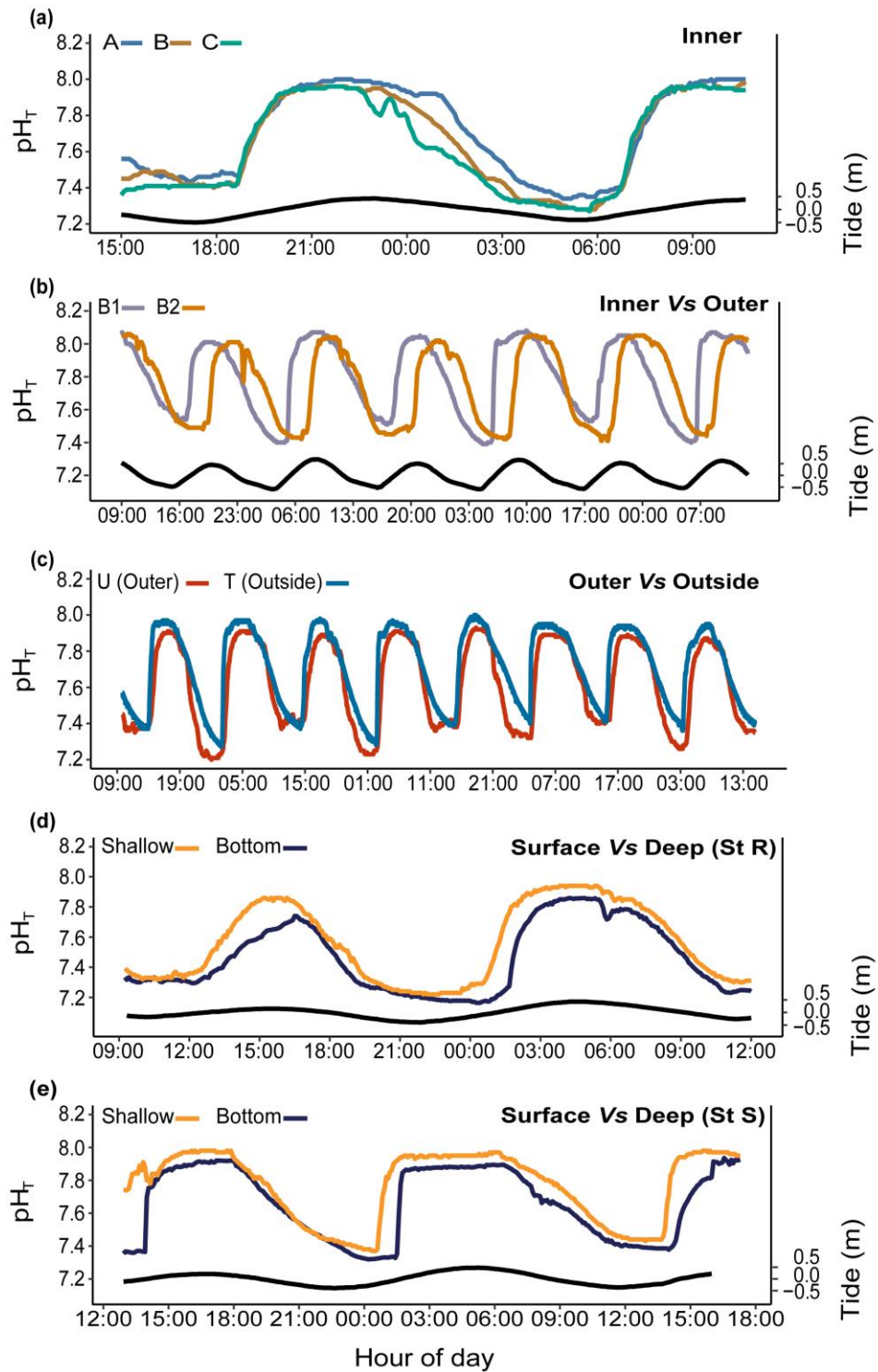
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358 **Figure 4.** Seawater pH_T (a), (b) and dissolved oxygen (DO) (c), (d) variations recorded at the reference (St R1 and R2) and
 359 Bouraké reefs (St B1 and B2). Data were overlaid in a single tidal phase (12 h) (a), (c); and in a 24-hour cycle (b), (d). Data
 360 are 22, 72, 31, and 72 semidiurnal tidal cycles for pH, and 36, 79, 34 and 42 semidiurnal tidal cycles for DO, for R1, R2, B1
 361 and B2, respectively.

362

363 The pH differed significantly between stations. During the studied period, we recorded means of 8.01 ± 0.04 and 7.89 ± 0.08
 364 pH_T units at R1 and R2, and 7.80 ± 0.22 and 7.67 ± 0.23 pH_T units at B1 and B2, respectively. At both B1 and B2, pH was
 365 strongly correlated with the tidal cycle (Fig. 4a, 82 and 73 % of the total variance were explained by the tidal harmonic
 366 analysis, respectively). In contrast, it was only marginally correlated with the 24-hour cycle (Fig. 4b). During each tidal phase,

367 the pH changed on average by about 0.6 units and reached a minimum of 7.23 and a maximum of 8.06 at B2 at low and high
368 tide, respectively (data not shown). The pH oscillations were mainly semidiurnal ($M_2 = 0.27 \pm 0.01$, $S_2 = 0.05 \pm 0.01$ pH_T
369 units, for B1; $M_2 = 0.24 \pm 0.01$, $S_2 = 0.09 \pm 0.01$ pH_T units for B2). At stations R1 and R2, pH changed on average by about
370 0.1 pH_T units and was mostly dependent on the 24-hour cycle (Fig. 4b, $S_1 = 0.058 \pm 0.004$ and $S_1 = 0.049 \pm 0.007$ pH_T units
371 for R1 and R2, respectively). Simultaneous short-term pH measurements showed significant spatial differences (Fig. 5; Tables
372 1), between i) stations A and C, ii) stations B1 and B2, and iii) the outer reef in the Bouraké lagoon (St U) and the station
373 outside the system (St T). There were also significant depth-related differences between shallow reefs and bottom water at
374 stations R and S. In both spatial and depth-related analyses, differences were approximately 0.05 to 0.1 pH_T units, and we
375 found the lowest values at stations C and B2 and the bottom of the channel (St R and S).

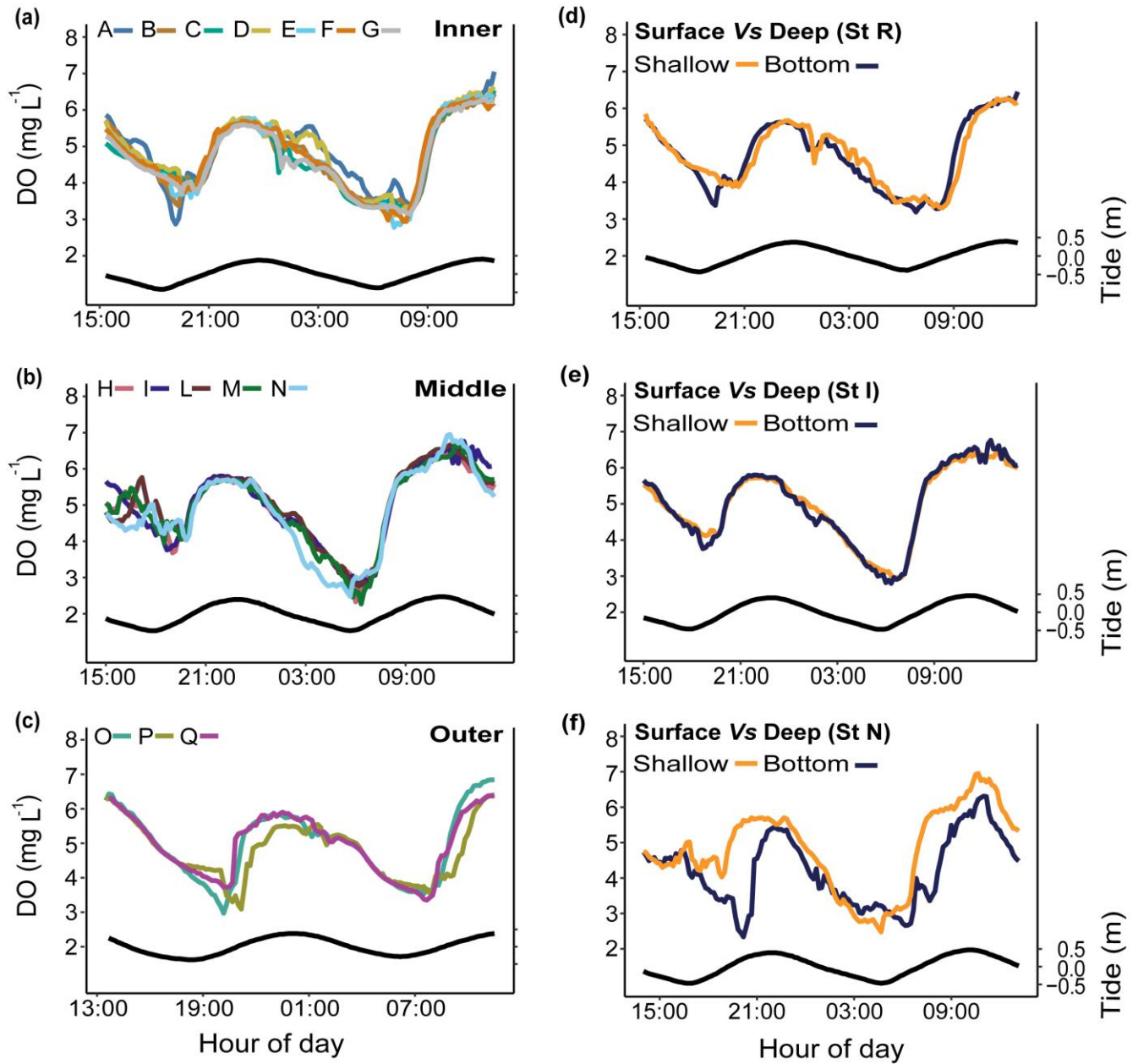


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377 **Figure 5.** Spatial and depth-related short-term changes (24 h) in pH_T at stations in the Bouraké lagoon (see Figure 1 for the
 378 stations' location). Spatial variations were assessed: **(a)** within the inner reef (St A-C); **(b)** between the inner and the outer
 379 reefs (St B1, B2); **(c)** between the outer reef (but still inside the Bouraké lagoon), and the reef outside the semi-enclosed
 380 lagoon (St U vs T). Depth-related variations were assessed between shallow reefs and the bottom of the channel: **(d)** at the
 381 inner reef (St R); and **(e)** at the outer reef (St S). Tide (black line) refers to St B2.

382

383 *Dissolved oxygen (DO)*. DO was measured during 36, 79, 34 and 42 semidiurnal tidal cycles at R1, R2, B1 and B2,
384 respectively. We overlaid all data on a single tidal phase of 12 h (Fig. 4c) and a 24-hour cycle (Fig. 4d). As with pH, the mean
385 diel DO was higher at the reference reefs than in the Bouraké lagoon. During the studied period, mean DO values were 4.89
386 ± 1.18 and 5.23 ± 0.89 mg L⁻¹ at B1 and B2, respectively, and 6.45 ± 0.95 and 6.48 ± 1.05 mg L⁻¹ at R1 and R2, respectively.
387 At stations B1 and B2, DO was strongly correlated with the tidal cycle (Fig. 4c, 82 and 72 % of the total DO variance were
388 explained by the tidal harmonic analysis, respectively), and only marginally with the 24-hours cycle (Fig. 4d). DO oscillations
389 were mainly semidiurnal ($M_2 = 1.17 \pm 0.08$ mg L⁻¹, and $M_2 = 0.83 \pm 0.09$ mg L⁻¹ for B1 and B2, respectively) with a substantial
390 diurnal component ($S_1 = 1.12 \pm 0.04$ mg L⁻¹, and $S_1 = 0.681 \pm 0.04$ mg L⁻¹ for B1 and B2, respectively). During a semidiurnal
391 tidal cycle, DO was lower at low tide (Fig. 4c, 3.7 mg L⁻¹ and 4.6 mg L⁻¹ at B1 and B2, respectively) and higher at high tide
392 (Fig. 4c, 6.0 mg L⁻¹ and 6.1 mg L⁻¹ at B1 and B2, respectively). The minimum (1.89 mg L⁻¹), and the maximum (7.24 mg L⁻¹)
393 DO value were both measured at B1 during low and high tide, respectively (data not shown). During a 24-hour cycle, DO
394 was lower in the early morning (Fig. 4d, 4.0 and 4.3 mg L⁻¹ at B1 and B2, respectively) and higher in the middle of the day
395 (Fig. 4d, 5.8 and 5.4 mg L⁻¹ at B1 and B2, respectively). In a single day, we recorded DO fluctuations of up to 6.37 mg L⁻¹ at
396 R1 and 4.91 mg L⁻¹ at B2. At stations R1 and R2, DO was mostly dependent on the 24-hour cycle ($S_1 = 1.34 \pm 0.06$ mg L⁻¹ and
397 $S_1 = 1.53 \pm 0.09$ mg L⁻¹ for R1 and R2, respectively) with lower values during the night (Fig. 4d, 5.5 mg L⁻¹ at both stations),
398 and higher values in the middle of the day (Fig. 4d, 7.8 mg L⁻¹, both stations).
399 Simultaneous short-term DO measurements (Fig. 6; Table 1) did not show significant differences over a 24-hour cycle at the
400 inner (St A-G; except for some specific deviations, e.g., St A), the middle (St H-N) or the outer reef (St O-Q). Dissolved
401 oxygen did not change on a vertical gradient at stations R and I, but the bottom DO was significantly lower at station N.



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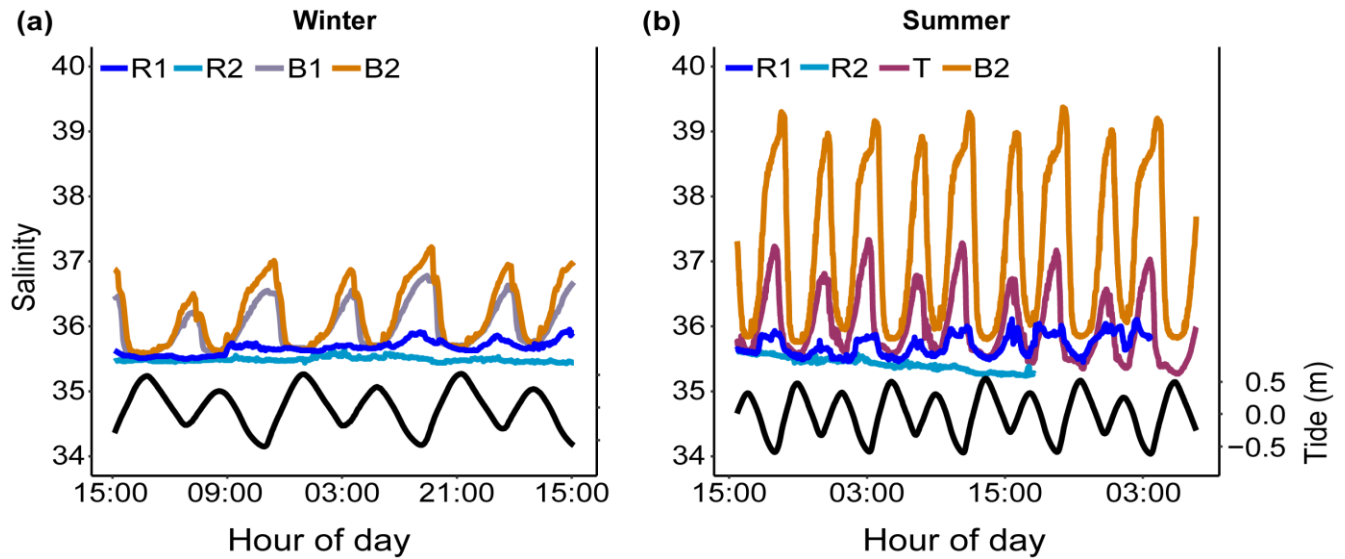
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Figure 6. Spatial and depth-related short-term measurements (24 h) of dissolved oxygen (DO) at stations in the Bouraké lagoon (see Figure 1 for the location of the stations). Spatial variations were assessed on (a) the inner reef (St A-G), (b) the middle reef (St H-N), and (c) the outer reef (St O-Q). Depth-related variations between shallow reefs and bottom water within the Bouraké lagoon were assessed (d) at the inner reef (St R) and (e, f) at the middle reef (St I and N). Tide (black line) refers to St B2.

412

Salinity (*S*). Salinity was measured only during two short-term periods in winter 2019 and summer 2020 (Fig. 7).



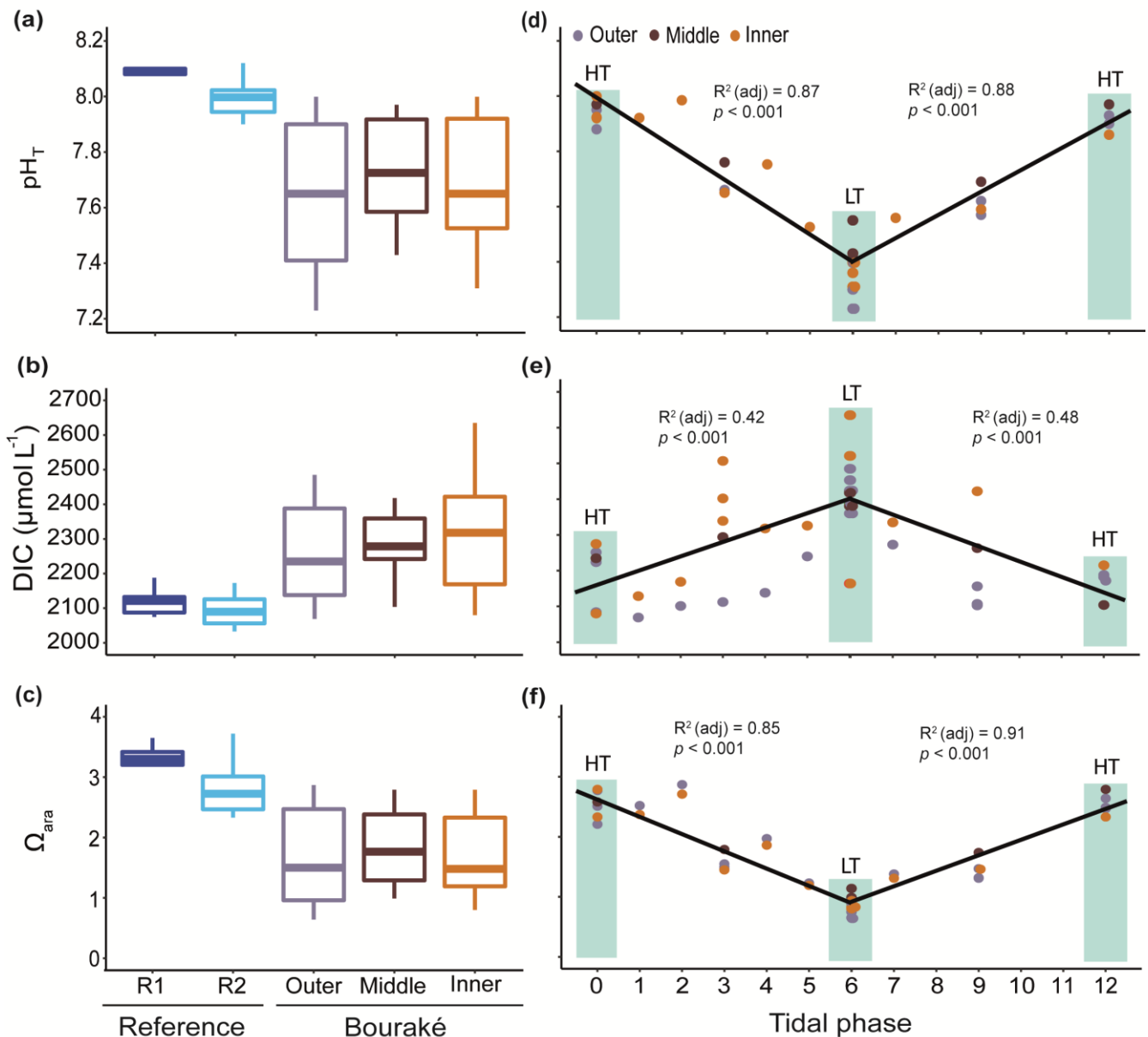
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Figure 7. Short-term salinity monitoring during (a) the winter of 2019 (from 15/07/2019 to 18/07/2019) at the reference (St R1 and R2) and Bouraké reefs (St B1 and B2); (b) the summer of 2020 (from 29/11/2020 to 04/12/2020) at the reference (St R1 and R2) and Bouraké reefs (St T and B2). Tide (black line) refers to St B2.

Salinity variations in the Bouraké lagoon were strongly correlated with the tidal cycle, and the highest mean values were measured during both winter and summer. Salinity was lower at reference reefs during both the winter and summer seasons (Fig. 7a-b). The mean salinity during the winter of 2019 was 35.49 ± 0.04 at R1, 35.67 ± 0.10 at R2, 36.00 ± 0.35 at B1, and 36.17 ± 0.47 at B2 with significant differences between stations (Tables 1). Salinity in the Bouraké lagoon peaked at 37.22 at B2 during low tide, while it was 35.65 at R1. During the summer of 2020, the mean salinity increased significantly in the Bouraké lagoon, with 37.22 ± 0.53 at B2 reaching 39.37, while it remained lower at R1 with 35.42 ± 0.10 and R2 with 35.71 ± 0.16 . During summer 2020, we also measured short-term variations in salinity at station T, which is outside the Bouraké system. Although its distance from the lagoon entrance, the water mass discharged during falling tide increased seawater salinity also at this station where we measured a maximum value of 37.33 during low tide.

3.2 Diel cycles of carbonate chemistry and chemical parameters

We monitored diel cycles in June 2017 and July 2019 at two reference reefs and three reefs within the semi-enclosed lagoon of Bouraké (i.e., the outer, the middle and the inner reefs). The seawater carbonate chemistry differed significantly between the reference and the Bouraké lagoon reefs (Fig. 8a-c; Supplementary Fig. S1; Tables 1, 2).



430
 431 **Figure 8.** Seawater carbonate chemistry measured (pH_T) and calculated (DIC and Ω_{ara}) during diel cycles in 2017 and 2019
 432 (pooled data, see also Supplementary Figure S1) at R1, R2 and the Bouraké reefs (outer, middle and inner) (a-c). Boxes (n =
 433 6-14) represent the interquartile range (25th and 75th percentile); the horizontal line is the median, and the whiskers represent
 434 the data range (i.e., minimum and maximum). Changes are illustrated across a 12 h tidal phase in the Bouraké lagoon (d-f).
 435 Linear regression lines are plotted for each falling (HT to LT) and rising tide (LT to HT). Shaded boxes have only a graphical
 436 significance and only suggest the tide changes during measurements.

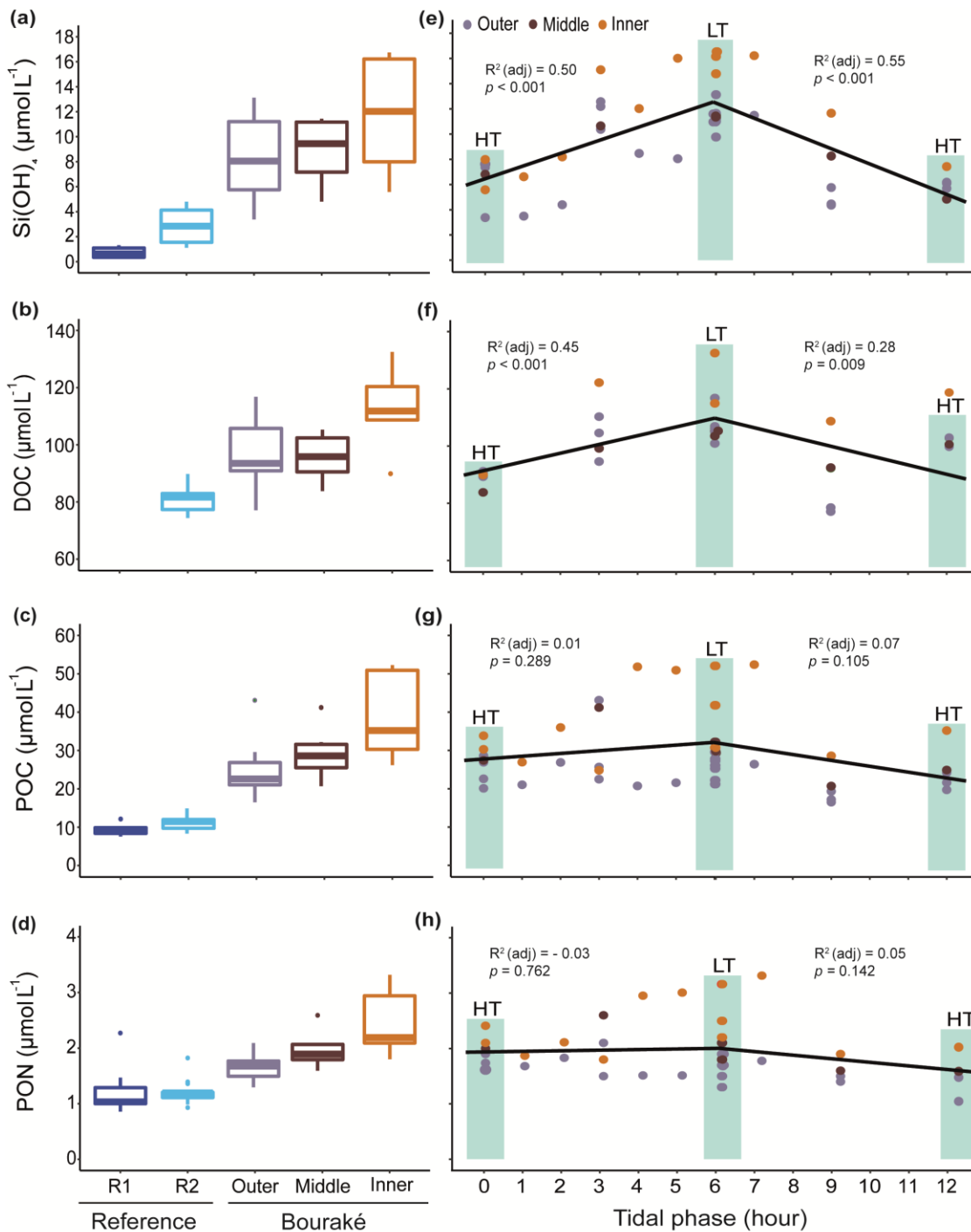
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440 **Table 2.** Summary of the seawater physical and chemical data (mean \pm SD) measured in July 2017 and June 2019 and calculated
 441 using the CO₂SYS package. Data were pooled and averaged per station. nd = not determined. Reference reefs: St R1 and R2;
 442 Bouraké lagoon stations: outer, middle and inner.
 443

444	R1	R2	Outer	Middle	Inner	
445	<i>Measured</i>					
446	Temp (°C)	22.54 \pm 0.31	22.69 \pm 1.50	23.57 \pm 1.31	24.05 \pm 1.20	23.00 \pm 1.46
447	pH _T	8.09 \pm 0.01	8.02 \pm 0.07	7.66 \pm 0.25	7.73 \pm 0.22	7.68 \pm 0.24
448	DO (mg L ⁻¹)	7.20 \pm 0.64	6.91 \pm 1.01	6.11 \pm 0.68	nd	5.98 \pm 0.67
449	A _T (μmol kg ⁻¹)	2294.1 \pm 0.3	2261.2 \pm 0.04	2357.8 \pm 0.1	2256.3 \pm 0.06	2393.7 \pm 0.08
450	Si(OH) ₄ (μmol L ⁻¹)	0.71 \pm 0.42	2.86 \pm 1.41	8.22 \pm 3.14	8.88 \pm 2.72	11.93 \pm 4.27
451	NO _x (μmol L ⁻¹)	0.21 \pm 0.05	0.51 \pm 0.16	0.65 \pm 0.20	0.54 \pm 0.09	0.54 \pm 0.12
452	PO ₄ ³⁻ (μmol L ⁻¹)	0.63 \pm 0.08	0.19 \pm 0.05	0.26 \pm 0.13	0.19 \pm 0.01	0.42 \pm 0.23
453	NH ₄ ⁺ (μmol L ⁻¹)	0.08 \pm 0.06	0.23 \pm 0.09	0.27 \pm 0.12	0.34 \pm 0.12	0.23 \pm 0.14
454	DOC (μmol L ⁻¹)	nd	81.15 \pm 5.49	96.89 \pm 10.78	95.70 \pm 8.38	112.82 \pm 14.41
455	DIC (μmol L ⁻¹)	2117.7 \pm 39.4	2092.4 \pm 42.9	2251.1 \pm 132.2	2282.5 \pm 112.0	2315.1 \pm 168.1
456	POC (μmol L ⁻¹)	9.28 \pm 1.47	11.15 \pm 1.97	24.07 \pm 5.31	29.37 \pm 7.03	38.11 \pm 10.43
457	PON (μmol L ⁻¹)	1.24 \pm 0.46	1.21 \pm 0.22	1.64 \pm 0.22	1.98 \pm 0.35	2.43 \pm 0.52
458	<i>Calculated</i>					
459	pCO ₂ (μatm)	353.42 \pm 7.24	467.76 \pm 112.03	1402.92 \pm 896.42	1047.06 \pm 599.07	1318.95 \pm 819.79
460	CO ₃ ²⁻ (μmol kg ⁻¹)	211.92 \pm 9.79	176.80 \pm 30.05	105.84 \pm 46.46	115.57 \pm 46.18	109.02 \pm 45.59
461	Ω _{aragonite}	3.35 \pm 0.16	2.80 \pm 0.49	1.68 \pm 0.73	1.84 \pm 0.73	1.72 \pm 0.71
462	TOC (μmol L ⁻¹)	nd	93.22 \pm 6.18	121.63 \pm 14.07	125.07 \pm 13.32	144.72 \pm 17.5
463	DIN (μmol L ⁻¹)	0.29 \pm 0.07	0.74 \pm 0.24	0.85 \pm 0.33	0.88 \pm 0.12	0.77 \pm 0.24
464	DIC:DIN	7371.9 \pm 1940.8	3132.9 \pm 999.1	4068.5 \pm 4621.8	2625.3 \pm 318.8	3276.9 \pm 1031.0
465	POC:PON	8.10 \pm 2.41	9.26 \pm 1.07	14.64 \pm 2.04	14.70 \pm 1.42	15.55 \pm 1.33
466	POC:DOC	nd	0.15 \pm 0.02	0.26 \pm 0.06	0.31 \pm 0.06	0.28 \pm 0.05
467	DOC:TOC	nd	0.87 \pm 0.01	0.80 \pm 0.03	0.77 \pm 0.02	0.78 \pm 0.03
468	Si(OH) ₄ :DIN	2.43 \pm 1.31	4.14 \pm 2.18	12.17 \pm 8.99	9.90 \pm 2.10	16.34 \pm 6.38
469	DIN:DIP	0.47 \pm 0.12	4.22 \pm 1.28	nd	4.65 \pm 0.52	2.78 \pm 1.21
470	Si(OH) ₄ :DIP	1.11 \pm 0.63	15.11 \pm 6.36	37.01 \pm 17.16	46.79 \pm 14.23	37.51 \pm 24.46

472 The reference reefs R1 and R2 had higher pH, CO₃²⁻ and Ω_{ara}, and lower DIC and pCO₂ than the outer, middle or inner reefs
 473 in the Bouraké lagoon. The range of values in the lagoon was similar to our previous measurements (see above), with levels
 474 of pCO₂ > 2000 μatm and levels of Ω_{ara} < 1. By assigning a tidal phase (from 0 to 12 h) to each sample taken in the Bouraké
 475 lagoon (all reefs combined), our diel measurements showed significant regressions, either positive or negative, depending on
 476 the seawater carbonate parameter and the tide phase (Fig. 8d-f; Supplementary Fig. S1; Supplementary Table S3). For
 477 example, the DIC reached the highest value of 2635 μmol L⁻¹ during low tide on the inner reef with an average of 2315.1 \pm
 478 168.1 μmol L⁻¹, while CO₃²⁻ reached the lowest value of 50.10 μmol kg⁻¹ during low tide on the inner reef with an average of
 479 109.02 \pm 45.59 μmol kg⁻¹.

480 Most of the chemical parameters were, in general, more concentrated (up to 5-times) in the Bouraké lagoon than at the
 481 reference reefs R1 and R2, and increased from the outer to the inner reef (Fig. 9a-d; Supplementary Fig. S2; Tables 1, 2).



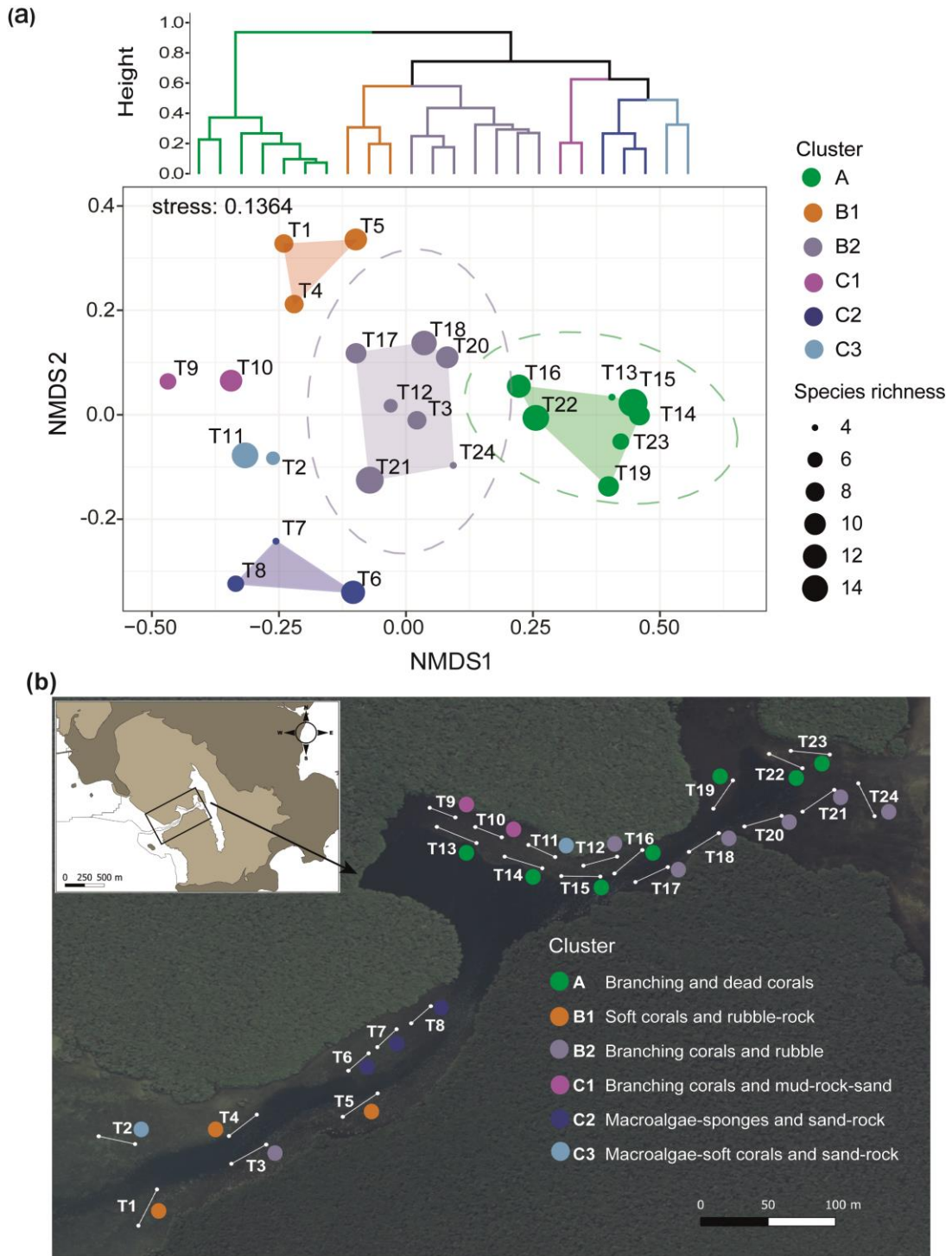
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483 **Figure 9.** Orthosilicic acid, organic carbon and nitrogen parameters measured during diel cycles in 2017 and 2019 (pooled
 484 data, see also Supplementary Figure S2) at R1, R2 and the Bouraké reefs (outer, middle and inner) (a-d). Boxes (n = 6-14)
 485 represent the interquartile range (25th and 75th percentile); the horizontal line is the median, and the whiskers represent the
 486 data range (i.e., minimum and maximum). Changes are illustrated across a 12 h tidal phase in Bouraké (e-h). Linear regression
 487 lines are plotted for each falling (HT to LT) and rising tide (LT to HT). Shaded boxes have only a graphical significance and
 488 only suggest the tide changes during measurements.

489 In addition, despite the Bouraké lagoon receives new seawater during the rising tide, only Si(OH)_4 had a decent regression
490 coefficient showing either positive and negative correlation with the falling and the rising tide (Fig. 9e-h; Supplementary Fig.
491 S2; Supplementary Table S3). It reached the highest value of $16.74 \mu\text{mol L}^{-1}$ during low tide on the inner reef with an average
492 of $11.93 \pm 4.27 \mu\text{mol L}^{-1}$, and the lowest value of $3.38 \mu\text{mol L}^{-1}$ during high tide on the outer reef with an average of $8.22 \pm$
493 $3.14 \mu\text{mol L}^{-1}$, 3- to 6-time higher than at the reference stations R2 and R1. The lack of a clear effect of the tide for the other
494 chemicals causes their accumulation inside the lagoon system. Ratios of organic and inorganic carbon, nitrogen, and nutrients
495 showed significant differences between the Bouraké lagoon and the reference reefs (Tables 1, 2) with higher average
496 POC:PON, POC:DOC, Si(OH)_4 :DIN, Si(OH)_4 :DIP and DIN:DIP(PO_4^{3-}), and lower average DIC:DIN and DOC:TOC in the
497 Bouraké lagoon compared to reference reefs.

498 **3.3 Benthic community distribution and species identification**

499 *Benthic community distribution.* The cluster analysis and nMDS allowed grouping the 24 transects in the Bouraké lagoon in
500 six clusters corresponding to habitat descriptors, namely clusters A, B1, B2, C1, C2, and C3 (Fig. 10a; see Supplementary
501 Fig. S3 for details of the community descriptors, and Table S4 for transect averaged data). The nMDS produced an exhaustive
502 representation of the Bouraké lagoon benthic communities and abiotic features (2-dimensional stress = 0.136) confirmed by
503 ANOSIM (Global R = 0.948, $p = 0.001$). For the benthic community distribution, a clear separation (95% confidence interval
504 represented by the ellipse in Fig. 10a) was found for cluster A and B2, including most transects located on the middle and
505 inner reefs (Fig. 10b).

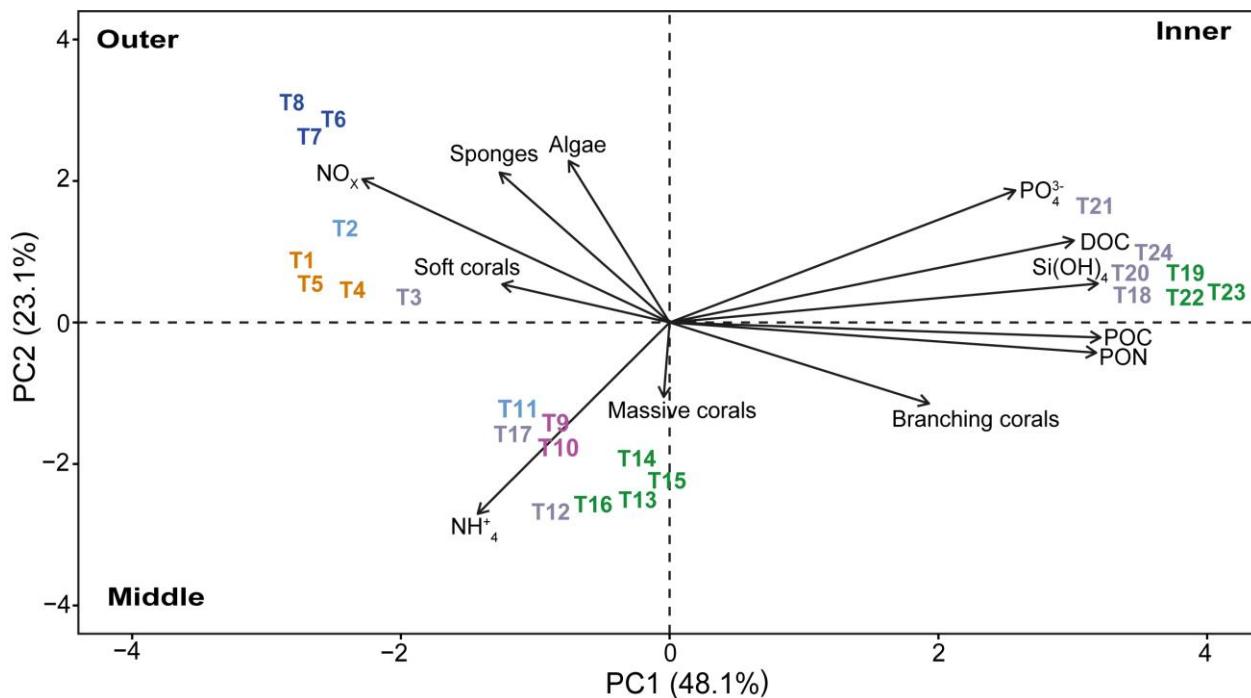


506

507 **Figure 10.** Benthic community and bottom substrate characterization in the Bouraké lagoon. Hierarchical Ward's method
 508 cluster dendrogram (top) and non-metric multidimensional scaling (nMDS) of abiotic and biotic percent cover and species
 509 richness per transect (T1-T24) (a). Colored circles represent clusters of transects identified in the dendrogram; circle size
 510 corresponds to species richness. Ellipses represent 95% confidence intervals. Dominant biotic and abiotic descriptors were
 511 used to describe the benthic community and bottom substrate for each cluster in (b). The satellite image is from georep New
 512 Caledonia database (<https://georep.nc/>), and the QGIS software was used for transect georeferencing.

513 For both, branching corals were the dominant biotic descriptor with an average of 81 % and 31 % coverage, respectively
 514 (Supplementary Fig. S3), with a maximum of 96 % in cluster A (T23, Supplementary Table S4). Among branching corals,
 515 the two most abundant genera were *Acropora* and *Montipora* (data not shown). For the abiotic features of the substrate, dead
 516 corals and rubbles characterized transects in clusters A and B2, respectively. Species richness (Fig. 10a) was high in all
 517 transects of the Bouraké lagoon, except T13 and T24, both found at the middle and inner reefs' limits. The abiotic substrate
 518 of cluster C1 was characterized by 66 % mud, 12 % rocks and 10 % sand (see Supplementary Table S4 for detailed cover data
 519 per transect). Only a few branching corals (<10 %) were found, but species richness was relatively high (6 at T9 and 10 at
 520 T10). Cluster C2 was distinct, which is not surprising due to its location in a relatively shallow convergence zone that divides
 521 the lagoon into two parts. There, the substrate is made of coarse sand (13 %) and rocks (12 %) and is mainly colonized by
 522 macroalgae and sponges (31 % and 32 %, respectively). Species richness in the area was heterogeneous and ranged from 4 to
 523 12. *Dictyota* spp. and *Halimeda discoidea* were the main macroalgal species, while *Rhabdastrella globostellata* was the
 524 dominant sponge species in the area. Cluster B1, located on the outer reef, is characterized by an abundance of soft corals (48
 525 %) and rubbles (21 %), and high biological richness (Fig. 10a). Cluster C3 is characterized by coarse sand (49 %), rocks (17
 526 %) and a few benthic organisms such as macroalgae (10 %) and soft corals (8 %).

527 Principal component analysis (PCA) reduced the multicollinearity problem and the first two Principal Components
 528 accounted for a cumulative 71.2 % of the dataset variance. In particular. The PCA plot (Fig. 11) allowed us to assign the
 529 transects to three distinct groups depending on their position in the study area.



530
 531 **Figure 11.** Principal component analysis (PCA) between benthic community (macroalgae, sponges, corals, branching corals
 532 and massive corals) data and environmental parameters (PON, POC, DOC, Si(OH)₄, PO₄³⁻, NO_x, and NH₄⁺) measured for
 533 each transect (n = 24). The length of the vectors indicates the contribution of each parameter. The first dimension explains
 534 48.1 % of the variance, and the second dimension explains 23.1 % of the variance. Colours refer to the cluster subdivision
 535 (Fig. 10).

536 The distinction among transects groups in the biplot is clear and they are grouped based on their location in the outer, middle
537 or inner lagoon reefs. The outer reef sites are characterized by sponges, macroalgae and soft corals, which appear to correlate
538 with NO_x. Conversely, branching corals are the shared and characteristic feature of the middle and inner reef sites. However,
539 these two reef groups differ in their major correlation to NH₄⁺ for middle reef sites and POC, DOC, Si(OH)₄, and PO₄³⁻ for
540 the inner sites.

541 *Species identification.* A total of 66 coral species were identified from the Bouraké lagoon (Supplementary Table
542 S5). Most of the species belong to the family Acroporidae (17 *Acropora* species, 2 *Anacropora* and 4 *Montipora*) and
543 Merulinidae (15 species). In total, 58 species were identified from the reference reef R2, 29 of which were also found in the
544 Bouraké lagoon. Overall, the hard coral fauna within the lagoon was characterized by species commonly found around the
545 Grande Terre, with some of them typically found in turbid coastal environments such as *Pseudosiderastrea tayamai* and
546 *Heliofungia actiniformis*. They were not encountered at the reference site. Remarkably, two New Caledonia endemic coral
547 species, *Cantharellus noumeae* and *Polycyathus fulvus*, both described from coastal habitats characterized by terrigenous
548 sediment inputs (Noumea lagoon and Prony Bay, respectively), were common in the innermost reef of the Bouraké lagoon.
549 The Bouraké lagoon also had 28 species of macroalgae (14 Phaeophyceae, 9 Chlorophyta and 5 Rhodophyta), including the
550 crustose coralline algae *Lithothamnium* sp., *Lithophyllum* sp., and *Hydrolithon reinboldii*, and 11 sponge species belonging
551 to eight families of the class Demospongiae (Supplementary Tables S5 and S6), which were found mainly in the coral matrix
552 and sediment.

553

554 **4 Discussion**

555 Marginal and extreme natural environments are increasingly used to predict the future of reefs in a changing world (Camp et al., 2018). In these environments, one or the combination of more environmental conditions differ from present-day values, 556 providing an opportunity to assess the resilience of organisms and to study their adaptive mechanisms in a natural 557 environment. Coral reefs, exposed to seawater pH and temperature values that are close to or even worse than those expected 558 for the future, have likely developed physiological trade-offs and expressed molecular changes that allow them to survive 559 sub-optimal and extreme conditions (Kurihara et al., 2021). When using these natural laboratories to predict species responses 560 to future environmental conditions, it is essential to take a multi-scale approach that incorporates the spatial and temporal 561 variability of the key physical and chemical parameters characterizing the study site (e.g., Vizzini et al., 2013; Camp et al., 562 2018; Aiuppa et al., 2021). Here, we mapped the spatial and temporal variability of the physical and chemical parameters in 563 the semi-enclosed lagoon of Bouraké, which is likely one of the most suitable natural laboratories to study the adaptation of 564 corals to the combination of acidification, warming and deoxygenation (Camp et al., 2017). There, we found an unprecedented 565 number of benthic species, including two New Caledonia endemics, thriving under chronic suboptimal conditions that 566 fluctuate with the tide. While the exact mechanism/s explaining their resilience remains to be discovered, our study provides 567 a compelling basis and fundamental baseline for using this site as a natural laboratory to investigate species' responses to a 568 combination of stresses in their natural environment. 569

570 **4.1 Physical and chemical characteristics of the Bouraké lagoon**

571 The Bouraké lagoon covers an estimated area of *ca.* 20 ha that penetrates a mangrove forest, large enough to assess the 572 combined effects of extreme environmental conditions on an established coral reef ecosystem. Our multi-scale approach 573 confirmed previous findings (Camp et al., 2017), showing that the Bouraké lagoon is hottest (ranging from 17.5-33.8 °C), 574 deoxygenated (ranging from 1.87-7.24 mg L⁻¹), and acid (ranging from 7.23-7.92 pH_T units) when compared to neighboring 575 reefs. Besides, we found that salinity was significantly higher than at the reference reefs during both winter and summer (with 576 maxima of 37.22 and 39.37, respectively).

577 We found several marked differences in the environmental conditions between the Bouraké lagoon and the reference reefs, 578 both in the absolute range, and in the variability of the measured environmental parameters.

579 First, the seawater temperature is higher in summer in the Bouraké lagoon (Fig. 3), but it is also colder during winter, resulting 580 in an annual temperature range of 17.5-33.8 °C. We compared temperatures recorded at Bouraké lagoon to those of the 581 reference St R2, which showed the most typical temperature range for shallow water temperatures in the south of New 582 Caledonia (i.e., 22-28 °C, Varillon et al., 2021). We notice that in Bouraké, temperatures were 40 % of the time above 28 °C 583 during the summer of 2020, while winter temperatures were on average 46.5 % of the time lower than 22 °C. While warming 584 is considered the main threat for coral reefs, low temperatures (< 20 °C) can cause coral bleaching by inducing responses 585 similar to high temperatures, including a reduction in the *Symbiodiniaceae* cell density and chlorophyll *a* content (e.g., Saxby 586 et al., 2003; Hoegh-Guldberg and Fine, 2004; Hoegh-Guldberg et al., 2005; Kemp et al., 2011; Bellworthy and Fine, 2021). 587 The negative effect of cold temperatures is even more substantial during neap tides when colonies on the reef crest are exposed 588 to air for hours at low temperatures during cold winters. For example, Porter et al. (1982) and Davis (1982) reported > 90 % 589 coral mortality in shallow (< 2 m) reefs of the Dry Tortugas following the winter of 1976 - 1977 when temperatures reached

590 14 °C. This is consistent with our observations (data not shown) during the cold winter of 2019 when we found that the upper
591 12-20 cm of several massive and branching corals had died.

592 Second, in the Bouraké lagoon, benthic assemblages are continuously exposed to large fluctuations in the main environmental
593 parameters, toward sub-optimal values. Some of these environmental fluctuations are entirely predictable. For instance,
594 marine organisms are exposed to a temperature fluctuation of about twice the reference reefs' amplitude (up to 6.5 °C vs 3.5
595 °C) in a single day. Dissolved oxygen fluctuations were similar between stations but in a significantly different range: 3.7 to
596 6.8 mg L⁻¹ at B2 and 5.4 to 7.8 mg L⁻¹ at the reference stations. According to the organisms tolerance to DO fluctuation, which
597 is quite unknown in corals, low DO concentrations can change fish tidal migration in the mangrove (Dubuc et al., 2019).
598 Besides, our tidal modelling revealed that, at the reference reefs, pH, DO (Fig. 4), and temperatures (data not shown) slightly
599 increased in the afternoon and decreased during the night. This finding agrees with what should be expected from reef
600 metabolic activities and daily cycles but, in the Bouraké lagoon, these parameters, including salinity (Fig. 7), are entirely
601 driven by tides. Here, seawater pH and DO varied between extremely low values at low tide and close-to-normal values during
602 high tide (see also Fig. 8a-c for pH). Finally, we found that the timing of the tide was out of phase between sites, with a delay
603 of about 45 minutes at high tide and 1.5 h at low tide in the Bouraké lagoon (Fig. 2).

604 The unique environmental conditions measured in the Bouraké lagoon are linked to its unique topographical and
605 geomorphological characteristics, the resulting water circulation and the direction of the tide. New water from the lagoon
606 enters through the channel at each rising tide and flows into the semi-enclosed lagoon towards the large mangrove area behind
607 it. This water initially had ambient values of pH, temperature and dissolved oxygen, but, during the trip, it mixes with the
608 acidic, warm and deoxygenated water in the system and the mangrove area, therefore gradually changing from its original
609 values. Inside the mangrove forest, we hypothesize that the water chemistry further changes due to the metabolic reactions in
610 the sediments, and mangrove roots (e.g., Alongi et al., 2004; Bouillon et al., 2007; Gleeson et al., 2013; Call et al., 2015).
611 Conversely, on a falling tide, the seawater becomes gradually more acidic, hot and oxygen-depleted because the water that
612 resided in the mangrove area gradually drains out of the system. This takes about 6 hours, during which the vast reservoir of
613 shallow mangrove water continues to be chemically altered, becoming increasingly acidic, oxygen-depleted and hot. As a
614 result, we measured significant spatial differences in pH between the outer reef (the entry of the lagoon) and the inner reef
615 (near the mangrove forest), as well as a considerable delay in the synchronization of the tidal shift (Fig. 5b). Interestingly,
616 because the volume of seawater discharged in 6 hours is so large, it affects also the area outside the system where we measured
617 similar seawater conditions as inside, even if it mixes with the main lagoon's water (see Fig. 5c, St U vs St T). It means that
618 the area (and the organisms) affected by the suboptimal parameters is larger than previously thought. The species living in
619 this area have likely developed specific mechanisms to withstand the very fluctuating environmental conditions, and as such,
620 they warrant further attention.

621 Since the fluctuations are linked to tidal phases, it could be argued that organisms living in the Bouraké lagoon may benefit
622 from periods of normal conditions at high tide, during which they can recover from the stress they have experienced at low
623 tide (e.g., Rivest et al., 2017). While this could be partially the case for species living on the outer reef, close to the main
624 lagoon, the environmental conditions inside the Bouraké lagoon rarely reach normal values (Fig. 8a-c), and persist longer as
625 the low tide is delayed by 1.5 h compared to the reference reef (Fig. 2).

626 Preliminary results from a hydrodynamic model of the study site suggest that tide-associated water mass movements are
627 spatially heterogeneous and likely to play an essential role in shaping coral resilience to extreme conditions (see discussion

628 below). Indeed, one can imagine a single water mass moving with the same physical characteristics from the mangrove area
629 towards the outer reefs or in the opposite direction depending on the tide. However, the complex geomorphology of the
630 Bouraké lagoon, its bottom topography and the complex web of coral reefs and mangrove trees on the edges deviate and
631 probably change the seawater physical and chemical properties. We measured significant spatial differences in pH within
632 each reef area (inner, middle and outer reefs; Fig. 5a, b), as well as throughout the water column (i.e., between the surface
633 and the bottom; Fig. 5d,e). In general, bottom seawater was 0.1-0.2 pH_T units lower than the surface, probably due to a
634 pumping mechanism by the water mass of more acidic pore water from the sediments. The pH also differed spatially within
635 the inner reef, by up to 0.3 pH_T units (for instance St. A vs St. C; Fig. 5a), perhaps due to the influence of stagnant water on
636 the mangrove edges or a more intense metabolic activity by the local benthic community. Dissolved oxygen concentrations
637 between the surface and the bottom were only significantly different at St. N on the middle reef (Fig. 6f). This station is
638 mainly characterized by mud as the current is lower than anywhere else in the Bouraké lagoon. Here, the sediment's biological
639 activity possibly traps the oxygen making it less available to the water column.

640 We can assume that, throughout the Bouraké lagoon, organisms are exposed to extreme and fluctuating suboptimal physical
641 parameters, such as pH, and DO which are more pronounced on the bottom and last longer, and with more extreme values,
642 on the inner reefs. It is also true for some of the seawater chemical parameters we measured, which show higher concentrations
643 in the Bouraké lagoon than on the reference reefs (see Table 2). For instance, we found that orthosilicic acid, phosphate,
644 dissolved and particulate organic carbon, and particulate organic nitrogen were 1.4 to 4.2-fold more concentrated on the inner
645 reef than at station R2. Within the Bouraké lagoon in general, chemicals increased from the outer to the inner reef, and except
646 for $[\text{NO}_x]$, $[\text{NH}_4]^+$, $[\text{PO}_4]^{3-}$, they never return to “normal” values despite the Bouraké system receives new seawater during the
647 rising tide. The high concentrations in orthosilicic acid and organic compounds, both dissolved and particulate are due to the
648 combined effect of more acidic and organic-rich water coming out of the mangrove forest during a falling tide (Fig. 9) and
649 the system's particular morphology, limiting the complete renewal of the seawater, especially at the inner reefs.

650 **4.2 Effects of physical and chemical conditions on species distribution**

651 Extreme environmental conditions, such as those measured in the Bouraké lagoon, are known to strongly affect the
652 metabolism, growth and even survival of several marine organisms, particularly those tolerating only a narrow range of
653 environmental changes such as scleractinian corals (e.g., Coles and Jokiel, 1977; Hoegh-Guldberg and Smith, 1989; Hoegh-
654 Guldberg, 1999; Fitt et al., 2001). For instance, *in situ* studies at volcanic CO_2 seeps have shown that chronic exposure to
655 ocean acidification (near-future $p\text{CO}_2$ levels) can cause a reduction in coral diversity and lower the recruitment and
656 abundances of structurally complex hermatypic corals. Moreover, shifts in competitive interactions between taxa and a
657 decrease in cover and richness of soft corals and sponges were also observed (e.g., Fabricius et al., 2011; Enochs et al., 2015;
658 Sunday et al., 2017; Agostini et al., 2018). However, the Bouraké lagoon features high coral, sponge, macroalgae and CCA
659 cover and species richness, adding at the check-list proposed by Camp et al. (2017) an additional 46 coral species (totaling 66
660 species and 33 genera), 28 species of macroalgae and 11 species of sponges (Supplementary Table S5). Such a high number
661 of species has never been reported for mangrove systems characterized by marginal conditions, at least concerning the three
662 major benthic groups taxa investigated here (e.g., Yates et al., 2014; Camp et al., 2019). In general, given the extensive
663 literature demonstrating the negative effects of suboptimal abiotic conditions on corals, our findings are unexpected and
664 suggest that corals in the Bouraké lagoon may have developed unique survival and adaptive mechanisms. Some studies have

665 reported similar findings, but none has ever reported such high coral species diversity (e.g., Yates et al., 2014; Schoepf et al.,
666 2015; Shamberger et al., 2018; Camp et al., 2019). For instance, in Palau (Micronesia), the highest coral cover (> 60%) and
667 species diversity (21 scleractinian genera) were found at the lowest pH study site of the Nikko Bay station (Barkley et al.,
668 2015). Studies in the Virgin Islands' mangrove system have reported over 30 coral species growing in marginal conditions
669 (Yates et al., 2014; Rogers, 2017). Similarly, in a recent study of two Australian mangrove lagoons, characterized by relatively
670 extreme environmental conditions (low pH, low oxygen and high temperature), Camp et al., (2019) identified 12 and 29
671 scleractinian coral species in the Woody Isles and at Howick Island, respectively. Among the 66 species we identified, 29 of
672 the most abundant coral species in New Caledonia were found at both the reference reef and in the Bouraké lagoon, indicating
673 that species living at our study site are not very different from a typical bay-sheltered fringing reef. There is not an apparent
674 selection in the coral form since massive (e.g., *Porites* spp.), corymbose (e.g., *Pocillopora digitata*), phaceloid (*Galaxea*
675 *fascicularis*) and branching (*Acropora* spp.) species are equally abundant, as well as in their thermal and pH tolerance. Further
676 experiments are underway at both the molecular and phenotypic level to better understand the mechanisms of resilience used
677 by the corals from Bouraké.

678 Remarkably, the two coral species, currently considered New Caledonia endemics, thrive in the innermost benthic
679 assemblages of the Bouraké lagoon, making it not only a unique natural laboratory as well as a potential conservation priority
680 site. In the Bouraké lagoon, benthos species richness was very high throughout the system and the distribution of branching
681 and massive corals was spatially heterogeneous (Fig. 10b; Supplementary Fig. S3). Coral cover decreased near the mangrove,
682 where the accumulation of fine sediments and exposure to air during low tide do not allow corals' survival (Supplementary
683 Fig. S3). Branching corals were particularly abundant on the inner and middle reefs, with the highest cover of 96 % in the
684 inner reef (T23, Supplementary Table S4). They became sparse on the outer reef, even absent at the system's entrance where
685 soft and massive corals dominated. Macroalgae and sponges, including CCA (data not shown), were quite cryptic throughout
686 the Bouraké lagoon but found almost everywhere in the coral matrix and buried in the sediment. They are particularly abundant
687 in a shallow area that divides the lagoon into two parts (transects T6-T8). There, a Venturi effect generates a strong current,
688 and the substrate consists of coarse sand and rocks. The high flow and the bottom characteristics are both likely to promote
689 macroalgae and sponge occurrences (averaged 30 % and 32 % cover, respectively). The sponge *Rhabdastrella globostellata*
690 is particularly abundant (up to 40 % cover) and forms massive banks embedding adjacent coral colonies. In general,
691 macroalgae and sponges diversity was not particularly rich, with only 28 and 11 species, respectively, likely due to the
692 sampling method. However, in the Indo-pacific mangrove systems, sponge diversity is, in general, not extremely high,
693 especially if compared to the Caribbean (Nagelkerken et al., 2008). Few studies are available from Indo-Pacific mangrove
694 sponges. The highest diversity was reported in a study in the mangrove of Banka Island (North Sulawesi, Indonesia), where
695 19 species were found (Calcinai et al., 2017). Our findings on mangrove sponges in such a unique mangrove area in New
696 Caledonia add to the limited knowledge of sponge species diversity in Pacific mangrove systems.

697 Water flow may play a critical role in the response of organisms to acidification (Cornwall et al., 2014; Hurd, 2015; Comeau
698 et al., 2019), warming (Schoepf et al., 2018) and deoxygenation (Hughes et al., 2020). It has been reported to affect the
699 diffusion boundary layer (i.e., a thin layer of stagnant water located around aquatic organisms) of corals, CCA and other
700 calcareous macroalgae, altering their ability to calcify at low pH and to excrete metabolic wastes (reviewed in Nelson and
701 Altieri, 2019). Knowing that flow speed could play a critical role and that deserves further investigations and measurements
702 at the study site, we found two hypotheses to explain the high diversity of the Bouraké species and their resilience to

703 suboptimal parameters: i) the species may benefit from the constant fluctuations of the physical and chemical parameters, ii)
704 the species may benefit from heterotrophic inputs of the mangrove (in term of nutrients and organic matter).
705 Regarding environmental fluctuations, we measured averaged daily variations in temperature, DO, and pH of about 3.79 °C,
706 3.80 mg L⁻¹, and 0.63 pH_T units, respectively. The temperature fluctuated by up to 6.55 °C in a single day. Frequent exposure
707 to stressful temperatures can induce acclimatization or adaptation in corals if the period of exposure is short enough to avoid
708 mortality (Oliver and Palumbi, 2011; Palumbi et al., 2014; Schoepf et al., 2015; Rivest et al., 2017; Safaie et al., 2018). For
709 example, in the Kimberly region (Australia), corals regularly exposed to temperatures up to 37 °C, with daily variations of
710 up to 7 °C, appear less prone to bleaching and grow and calcify at rates comparable to corals in more thermally stable
711 environments (Dandan et al., 2015; Schoepf et al., 2015; 2020). However, they are not immune to extreme heat stress events
712 (Le Nohaïc et al., 2017). Another example is the corals in the back reef and shallow pools around the island of Ofu (American
713 Samoa), which despite occasional daily fluctuations of up to 6 °C, and an average daily temperature of 29 °C in summer
714 (Piniak and Brown, 2009; Oliver and Palumbi, 2011) sustain reasonable levels of coral cover (25-26 %) and high diversity
715 (Craig et al., 2001). Some studies tend to indicate that corals depended on the *a priori* 'frontloading' of genes involved in heat
716 resistance in the coral host and/or the host's ability to adjust its physiology during short-term (weeks) environmental changes
717 (Barshis et al., 2013; Palumbi et al., 2014). However, physiological adjustments have an energetic cost that corals cannot
718 sustain in the long term, affecting other metabolic functions such as calcification. It could explain in part the lower
719 calcification rates observed in some coral species, as seems to be the case for corals in the Bouraké lagoon (e.g., Camp et al.,
720 2017). There is also evidence that exposure to short-term oscillations in low pH, alone or in combination with high
721 temperature, can mitigate the impact of extreme environmental stressors on corals (Warner et al., 1996; Oliver and Palumbi,
722 2011; Dufault et al., 2012; Schoepf et al., 2015; Safaie et al., 2018). Species in the Bouraké lagoon appear to be thriving
723 despite the large seawater pH variability (up to 0.6 pH_T units). In general, species exposed to pCO₂ fluctuations above 500
724 µatm in their natural environment display enhanced plasticity to future ocean acidification scenarios (Vargas et al., 2017;
725 Carstensen and Duarte, 2019). For example, fluctuations in pH could increase CCA tolerance to ocean acidification by
726 providing respite periods at ambient pH (Rivest et al., 2017), during which organisms can calcify at a standard rate,
727 compensating for decreased calcification during periods of lower pH (Comeau et al., 2013; Cornwall et al., 2018). Diurnal
728 variability in pH, typical of more variable pH habitats, may confer tolerance to resident calcifying species via the selection of
729 individuals better adapted to survive in these environments (Rivest et al., 2017; Kapsenberg and Cyronak, 2019), although
730 transgenerational acclimatization is required (Cornwall et al., 2020).
731 In addition to changes in temperature and pH, which are the two most commonly tested environmental parameters, oxygen
732 must also be considered. Despite the impact of reduced oxygen levels and its natural fluctuations on coral reefs has yet to be
733 established, low DO (2-4 mg L⁻¹) seems to increase the sensitivity of branching corals, resulting in a decline in coral health,
734 bleaching, and tissue loss (Haas et al., 2014; Hughes et al., 2020; Alderdice et al., 2021). Our study shows that although the
735 Bouraké system can reach conditions close to hypoxia for several coral species (< 3 mg L⁻¹; Fig. 4), these latter seem to have
736 promoted compensation mechanisms that allow them to survive in these conditions. The natural laboratory of Bouraké, where
737 DO fluctuates with the tide, in combination with other environmental stressors, offers a perfect setting to test the practically
738 unknown effects of deoxygenation in reef-building corals exposed to acid and hot conditions (Nelson and Altieri, 2019;
739 Hughes et al., 2020).

740 Although corals appear to possess cellular mechanisms to counteract short-term osmotic changes (Mayfield and Gates, 2007),
741 high and fluctuating salinity is possible additional stress that corals living in the Bouraké lagoon have to face daily, adding
742 up to the already long list of suboptimal environmental parameters. *In situ* studies have demonstrated that acute and prolonged
743 decrease in salinity can affect the coral photosynthetic efficiency, resulting in a reduction of the amount of energy transferred
744 to corals (Muthiga and Szmant, 1987; Manzello and Lirman, 2003) and induce coral death (e.g., Jokiel et al., 1993). The
745 response of corals to a change in salinity is related to the strength and duration of the hypo-hypersaline exposure and the
746 species tolerance. For example, *Stylophora pistillata* seems able to acclimate more effectively to hypo- rather than to hyper-
747 saline conditions (Ferrier-Pagès et al., 1999). Further experiments are needed to assess the effect of high and fluctuating
748 salinity on the physiology of corals in Bouraké.

749 Besides the hypothesis that environmental variability improves the metabolism of organisms, particularly their resilience to
750 extreme conditions, a series of other physical (e.g., current flow) and chemical parameters (e.g., organic matter) in the Bouraké
751 lagoon may work in combination to offset or enhance these effects. Mangrove habitats are highly productive ecosystems and
752 are sites of intense carbon processing, with a high potential impact on the global carbon budget (e.g., Borges et al., 2003;
753 Dittmar et al., 2006; Bouillon et al., 2014). In the Bouraké lagoon, benthic communities might have access to a range of
754 heterotrophic inputs, nutrients, carbon, and nitrogen sources. These sources can be metabolized by the species to increase
755 their energy budget and cope with the suboptimal parameters, but they can also become toxic if too concentrated or depleted,
756 leading to functional limitations. We measured exceptionally high concentrations of organic and inorganic carbon and
757 nitrogen, but also of some nutrients, notably silicates and phosphorus, and we confirmed the potential contribution of the
758 mangrove in those inputs, especially during the falling tide (Fig. 8 and 9; Supplementary Fig. S1 and S2; Table 2). We found
759 that dissolved organic carbon contributes significantly to the TOC pool (POC+DOC), with a concentration increase between
760 the reference sites and the Bouraké lagoon, reaching maximum values on the inner reef. High organic matter content can
761 increase DOC availability to corals, providing the sustainable energy to withstand extreme environmental conditions (Levas
762 et al., 2015). Some studies showed that the high organic matter of turbid reefs can support elevated coral heterotrophy that
763 can facilitate energy maintenance during periods of stress (Anthony and Fabricius, 2000).

764 We also found that nutrients could partially explain the distribution of organisms throughout the Bouraké lagoon (Fig. 11).
765 Indeed, NO_x concentrations were higher on the outer reef, dominated by sponges, macroalgae and soft corals, while NH_4^+ was
766 high on the middle reef and POC, DOC, $\text{Si}(\text{OH})_4$, and PO_4^{3-} were higher on the inner reef. Both the middle and inner reefs are
767 characterized by the highest branching coral cover. Nitrates can accumulate inside the host cells, possibly favoring sponges
768 and macroalgae, while ammonium is the preferred source used by coral symbionts (Raven et al., 1992). However, the nutrient
769 negative or positive effect on corals' physiology is difficult to demonstrate experimentally (Atkinson et al., 1995; Szmant,
770 2002; Bongiorni et al., 2003). Van De Waal et al., (2009) have shown that the performance of organisms depends on
771 concentrations of, and ratios between, different nutrients, and between organic and inorganic matter/components and the
772 possible imbalance due to environmental changes. We found that the Bouraké lagoon is mostly N-limited ($\text{DIN:DIP} < 10$ and
773 $\text{Si}(\text{OH})_4:\text{DIN} > 1$), which confirms the findings of Justić et al. (1995) and is similar to the conditions observed in other New
774 Caledonian mangroves (Jacquet et al., 2006). Nutrients limitation has been demonstrated to lower the temperature effect at
775 which coral bleaching occurs (Wiedenmann et al., 2013; Ezzat et al., 2016a; 2019), which contrasts with the resilience of
776 Bouraké corals to the warming in the summer of 2016 (10-20 % bleaching only) compared to other reefs in New Caledonia
777 (up to 90 % bleaching) (Benzoni et al., 2017). Corals symbionts recycle their host's metabolic wastes and take up dissolved

778 inorganic nitrogen (DIN) and phosphorus (DIP) from seawater (Grover et al., 2003; Pernice et al., 2012; Rosset et al., 2015),
779 both of which are used to produce vital organic molecules. Nutrient starvation can occur when the availability of one type of
780 essential nutrient decreases, resulting in an imbalanced N:P ratio of inorganic nutrients in seawater (Wiedenmann et al., 2013;
781 D'Angelo and Wiedenmann, 2014). Based on our measurements, the N:P ratio in the Bouraké lagoon was 2.8:1 and 4.6:1 at
782 the middle and the reference reefs, respectively, which is lower than the range calculated for average reef waters of 4.3:1 to
783 7.2:1 (Crossland et al., 1984, Furnas et al., 1995). Although these data should be considered cautiously, they suggest that the
784 Bouraké lagoon seawater is not limited in phosphorus, an essential nutrient in coral resilience to bleaching (Ezzat et al., 2016b;
785 Rosset et al., 2017).

786 Seawater in the Bouraké lagoon was not limited in Si (Si(OH)_4 :DIP > 10 and Si(OH)_4 :DIN > 1; Justić et al., 1995), and Si was
787 six times more abundant on the inner reef than elsewhere in the system (Fig. 9; Table 2). Silicates are an important source of
788 nutrition and skeletal construction for primary producers such as sponges. Orthosilicic acid is a biologically available form of
789 silicon that is poorly soluble at a very low pH. In the Bouraké lagoon, pH fluctuates with the tide, and one can expect that
790 orthosilicic acid would rise during the falling tide. This elevated concentration of orthosilicic acid could be involved in the
791 growth and persistence of the Bouraké large banks of *Rhabdastrella globostellata*, as previously reported for other sponges
792 in the shallow hydrothermal vents in the Pacific Ocean (Maldonado et al., 1999; Cárdenas and Rapp, 2013; Bertolino et al.,
793 2017).

794 We are still missing information about light, turbidity, current, bacteria biomass, phytoplankton sources as well as other
795 biological communities to fully understand this complex and dynamic system and the functioning of this complex and dynamic
796 coral ecosystem. However, our data already demonstrate the Bouraké lagoon's uniqueness as a natural laboratory for studying
797 the adaptive responses of corals and other reef species to the combination of multiple suboptimal environmental parameters,
798 which are, to some extent, worse than those projected for the future. Our investigations indicate that the geomorphology of
799 this unique site has not changed for the last 80 to 100 years, certainly longer. With this in mind, we assume that, at least in
800 the previous century, the environmental conditions remained unchanged, and the corals of the Bouraké lagoon have
801 experienced the current extreme conditions for several generations. Most importantly, the species found in the Bouraké lagoon
802 are not unique to the mangrove habitat. They are common throughout New Caledonia and the SW Pacific region, suggesting
803 that they have used specific strategies to cope with the suboptimal environmental conditions.

804 We are also unsure how the different parameters will fluctuate at bay-sheltered reefs over time and under future climate
805 change conditions and if the variability measured in the Bouraké lagoon is representative of the natural fluctuations expected
806 for coastal habitats in the future. Indeed, earth system model simulations, which are mainly based on open-ocean system
807 models, project that the seasonal amplitude of pH and $p\text{CO}_2$ will increase by 81 % in the future climate (Kwiatkowski and
808 Orr, 2018). Projections also indicate that ocean acidity extremes will be more frequent (Burger et al., 2020), which could
809 mean that future diel pH variability will increase even more at sites with the most significant variability today. The situation
810 could be similar for seawater temperature, a parameter that is already high in Bouraké, and likely close to the coral's thermal
811 tolerance threshold. If this were to happen in the following decades, the coral reef of Bouraké would face incredibly harsh
812 environmental conditions since physical and biogeochemical parameters measured during this study already exceed future
813 climatic simulations.

814

815 **5 Conclusion**

816 We used a multi-scale approach to characterize the physical and chemical environmental parameters of one of the most
817 realistic natural laboratory for extreme environmental conditions, the semi-enclosed lagoon of Bouraké (New Caledonia), and
818 accurately map its benthic community for the first time. We studied several physical and chemical parameters such as pH,
819 dissolved oxygen, temperature, and salinity, but also nutrients and organic matter, and we found that: i) they fluctuate between
820 low and high tides, ranging from suboptimal-extreme to near normal values; ii) although predictable according to the tide,
821 they differed spatially, and iii) suboptimal values persisted longer and were more acute on the inner reef. Our data clearly
822 show that fluctuations are: i) predictable, at least for some of the physical parameters for which we have enough data (i.e.,
823 pH, DO, temperature and salinity); ii) mainly driven by the tide; and iii) that seawater nutrient imbalance and organic inputs
824 increase during the falling tide and originate from the mangrove forest and associated sediments. Although several studies
825 suggest that ocean acidification, warming and, to some extent, deoxygenation will lead to a reduction in biodiversity, increase
826 in bleaching and reef dissolution, in the Bouraké lagoon, we found healthy reef with high coral cover and species richness,
827 but also sponges and macroalgae (including CCA). It was beyond the scope of this study to assess the contribution of
828 environmental variability and nutrient imbalance to the organism' stress tolerance under extreme conditions. However, both
829 coexist in the Bouraké lagoon, and we believe there is evidence of their contribution to the survival of organisms in extreme
830 environmental conditions. Our study provides evidence that this is possible in existing natural habitats, giving a glimmer of
831 hope for the future of coral reefs. Further experiments are needed to reveal the mechanisms involved in the organisms'
832 resilience to such conditions. Finally, we provide the compelling basis for using this site as a natural laboratory and study
833 better the multitude of complex stressors acting together on life-long adapted coral reefs.

834

835 *Author contribution.* R.R.-M. conceived and designed the project with input from F.M. for the benthic community distribution
836 study. F.M. and R.R.M. collected the data. F.B., C.P., C.C. and B.C. identified the corals, macroalgae and sponges,
837 respectively. M.P.-P. performed all the chemical analyses. F.M. conducted the data analysis with the help of J.A. and R.R.-M.
838 F.M. drafted the manuscript in collaboration with R.R.-M. All co-authors read and edited the final version of the manuscript.

839

840 *Competing interests.* The authors declare that they have no conflict of interest.

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848

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