1 2 3	Effects of <i>in situ</i> CO ₂ enrichment on structural characteristics, photosynthesis, and growth of the Mediterranean seagrass <i>Posidonia oceanica</i>
4 5	T. E. Cox ^{1,2} , F. Gazeau ^{1,2} , S. Alliouane ^{1,2} , I. E. Hendriks ³ , P. Mahacek ^{1,2} , A. Le Fur ^{1,2} , and JP. Gattuso ^{1,2,4}
6 7 8	¹ Sorbonne Universités, UPMC Univ Paris 06, Observatoire Océanologique, F-06230 Villefranche-sur-mer, France, erincox@hawaii.edu
9 10 11	² CNRS, UMR 7093, Laboratoire d'Océanographie de Villefranche (LOV), F-06230 Villefranche-sur-mer, France
12 13 14	³ Global Change Department, IMEDEA (CSIC-UIB), Instituto Mediterraneo de Estudios Avanzados, C/Miquel Marques 21, 07190 Esporles, Mallorca, Spain
15 16 17 18	⁴ Institute for Sustainable Development and International Relations, Sciences Po, 27 rue Saint Guillaume, F- 75007 Paris, France
19	Running head: In situ ocean acidification effects on P. oceanica
20	Abstract
21	Seagrass are expected to benefit from increased carbon availability under future ocean
22	acidification. This hypothesis has been little tested by in situ manipulation. To test for ocean
23	acidification effects on seagrass meadows under controlled CO ₂ /pH conditions, we used a Free
24	Ocean Carbon Dioxide Enrichment (FOCE) system which allows for the manipulation of pH as
25	continuous offset from ambient. It was deployed in a Posidonia oceanica meadow at 11 m depth
26	in the Northwestern Mediterranean Sea. It consisted of two benthic enclosures, an experimental
27	and a control unit both 1.7 m^3 , and an additional reference plot in the ambient (2 m^2) to account
28	for structural artifacts. The meadow was monitored from April to November 2014. The pH of the
29	experimental enclosure was lowered by 0.26 pH units for the second half of the eight-month
30	study. The greatest magnitude of change in <i>P. oceanica</i> leaf biometrics, photosynthesis, and leaf
31	growth accompanied seasonal changes recorded in the environment and values were similar
32	between the two enclosures. Leaf thickness may change in response to lower pH but this requires

33	further testing. Results are congruent with other short-term and natural studies that have
34	investigated the response of <i>P. oceanica</i> over a wide range of pH. They suggest any benefit from
35	ocean acidification, over the next century (at a pH_T of ~7.7), on <i>Posidonia</i> physiology and growth
36	may be minimal and difficult to detect without increased replication or longer experimental
37	duration. The limited stimulation, which did not surpass any enclosure or seasonal effect, casts
38	doubts on speculations that elevated CO_2 would confer resistance to thermal stress and increase
39	buffering capacity of meadows.
40	

41 Keywords: buffering capacity, leaf biometrics, meadows, ocean acidification, oxygen fluxes,
42 PAM fluorescence, pH

44 **1 Introduction**

45 Ocean carbonate chemistry is being altered in ways that may affect future ocean ecology. 46 The ocean absorbs carbon dioxide (CO_2) from the atmosphere which increases the concentrations 47 of inorganic carbon and CO₂, and decreases pH in a process referred to as ocean acidification. 48 Surface ocean pH has decreased by 0.1 units since the beginning of the industrial era and a 49 further decline (0.06 to 0.32 units) is projected over the next century (Ciais et al., 2013). Through 50 this process, the relative proportions of dissolved inorganic carbon species are concurrently being 51 altered. By 2100, bicarbonate (HCO₃), already widely available, will increase along with CO_2 , 52 which will have the largest proportional increase from current day levels. An increase in carbon 53 availability may benefit some marine producers (Koch et al., 2013). In contrast, the concentration of carbonate ions (CO_3^{2-}) needed by calcifying organisms will decrease. Thus, ocean acidification 54 55 can alter competitive interactions which may cascade to alterations at the ecosystem level. 56 Seagrass meadows rank as one of the most productive ecosystems on Earth (Duarte et al., 57 2010; Duarte and Chiscano, 1999). They are highly valued for their ability to improve water 58 quality, stabilize sediment, and provide habitat for a diversity of organisms. Human-driven 59 changes to the seawater clarity and quality (e.g. eutrophication, ocean warming) are often related 60 to meadow decline (Jordà et al., 2012; Waycott et al., 2009). However, these habitat-forming 61 seagrasses are thought to benefit from ocean acidification because they are able to use both CO_2 and HCO_3^- for photosynthesis but, with a higher affinity for CO_2 and are often found to be 62 63 carbon-limited (Invers et al., 2001; Koch et al., 2013). 64 Experiments under elevated CO₂ have shown an increase in seagrass photosynthesis 65 (Apostolaki et al., 2010; Invers et al., 1997; Jiang et al., 2010; Ow et al., 2015; Zimmerman et al., 66 1997), below ground growth (Hall-Spencer et al., 2008; Zimmerman et al., 1997; Russell et al.,

67 2013) and flowering frequency (Palacios and Zimmerman, 2007). Yet the majority of these

68	studies were conducted in the laboratory over relatively short durations with single taxa or small
69	groups of taxa isolated from their surroundings. Although studies along carbon dioxide vents
70	allow for a whole ecosystem approach, the high spatial and temporal variability in CO_2 levels
71	hampers the determination of a reliable dose-response relationship (Hall-Spencer et al., 2008;
72	Kerrison et al., 2011). To the best of our knowledge, only Campbell and Fourqurean (2011,
73	2013a, 2014) have manipulated partial pressure of carbon dioxide (pCO_2) levels in a contained
74	(ie. as opposed to free flow CO ₂ bubbling) manner <i>in situ</i> within a <i>Thalassia</i> meadow to test the
75	response of seagrass to ocean acidification. After 6 months of exposure to lowered pH (-0.3 from
76	mean ambient), the seagrass had increased non-structural carbohydrate content by 29% in below
77	ground structures (Campbell and Fourqurean 2014). This finding generally supports the
78	hypothesis that plant production will be stimulated from the increased carbon availability.
79	Posidonia oceanica is the foundation species for mono-specific meadows in the
80	Mediterranean Sea where it covers up to 23% of shallow waters (0-50 m; Pasqualini et al., 1998)
81	and provide services valued at $172 \notin m^{-2}$ year ⁻¹ (Vassallo et al., 2013). These plants are largely
82	dependent upon abiotic factors as evident by its seasonal growth and physiology (Alcoverro et al.,
83	1995, 1998; Bay, 1984; Duarte, 1989). They have been studied under a range of pH in the
84	laboratory as well as along pH gradients near CO ₂ vents (Invers et al., 1997, 2001, 2002; Hall-
85	Spencer et al., 2008; Cox et al., 2015). Around natural CO ₂ vents in Ischia (Italy), P. oceanica
86	biomass was greatest at the station nearest the CO_2 source with a mean pH _T of 7.6 and minimum
87	of 6.98 (Hall-Spencer et al., 2008). Indeed, P. oceanica has a C3 photosynthetic pathway that is
88	hypothesized to benefit from increased carbon availability and its photosynthesis is not saturated
89	with respect to dissolved inorganic carbon at natural concentrations in seawater (Invers et al.,
90	1997, 2001). This is evident by their enhanced productivity in the laboratory under a pH range
91	from 9.0 to 7.9 and has been attributed to a less efficient use of widely available HCO_3^- and their

92	reliance on CO ₂ for about 50% of carbon for photosynthesis (Invers et al., 1997, 2001). External
93	carbonic anhydrase acts to dehydrate HCO_3^- to CO_2 which enters the cell by a diffusive process
94	(Invers et. al 2001). Thus CO ₂ limitation depends upon the thickness of the boundary layer and
95	can also occur at high pH with slow diffusion rates (Invers et al., 2001). However, the extent of
96	the stimulation at pCO_2 levels projected for the coming decades appears limited (Cox et al., 2015;
97	Invers et al., 2002). In addition, the environment and species dynamics in meadows are complex
98	and interactions can alter outcomes. For example, the leaves and roots are colonized by small
99	invertebrates and epiphytic algae (Borowitzka et al., 2006). These associated species, many
100	sensitive to dissolution, compete with the plants for resources (Cebrián et al., 1999; Martin et al.,
101	2008; Sand-Jensen et al., 1985). A laboratory investigation of this potential interaction under two
102	elevated pCO_2 levels (on the total scale, pH _T 7.7 and 7.3) was performed (Cox et al., 2015).
103	Despite loss of calcified photosynthetic epiphytes at pH _T 7.7, the effect on shoot productivity was
104	limited and seagrass photosynthesis (without epiphytes) was only stimulated at pH_T of 7.3, a
105	value unlikely to occur in the Mediterranean Sea in the next century (Cox et al., 2015). The long-
106	lived plants, however, were maintained for a relatively short duration of six weeks and only under
107	the irradiance, temperature, and nutrient conditions of February to March. From these studies it is
108	difficult to predict the impact of ocean acidification on P. oceanica.
109	Any alteration in <i>P. oceanica</i> productivity or abundance will likely have repercussions to
110	meadow function. Therefore the aim of the present study was to test the hypothesis that
111	Mediterranean seagrass, P. oceanica, will benefit from ocean acidification. We tested this
112	hypothesis in situ with a Free Ocean Carbon Dioxide Enrichment (FOCE) system (see Gattuso et
113	al., 2014) which consisted of two partially-open enclosures that were deployed in the Bay of
114	Villefranche (France) for eight months (April-November 2014). The pH was manipulated
115	continuously, in one enclosure, at a -0.26 pH unit offset from ambient between June and

116 November. Before and during pH manipulation, macrophyte abundance, *Posidonia* leaf
117 biometrics, photosynthesis, and growth were measured and environmental conditions were

118 monitored.

119 **2 Method**

120 **2.1 Experimental setup and system function**

This study used the European FOCE (eFOCE) system, an autonomous system which
allows for the *in situ* manipulation of pH in benthic enclosures as an offset from ambient pH
(Gattuso et al., 2014). The system was deployed in the Bay of Villefranche, approximately 300 m
from the Laboratoire d'Océanographie de Villefranche (NW Mediterranean Sea, France;
43°40.73'N, 07°19.39'E; Fig. 1). The eFOCE engineering design consisted of a surface buoy and
two underwater benthic enclosures (Fig. 1).

The underwater portion of eFOCE consisted of two clear, 1.7 m^3 (2 m long x 1 m width x 127 128 0.85 m tall) perspex enclosures that were open on the bottom to partially enclose a portion of a P. 129 oceanica meadow. They were located at 11 m depth, were placed end to end approximately 1.5 m 130 apart and faced south. The pH in one enclosure, referred to as the experimental enclosure, was 131 lowered by ~0.25 units as an offset from ambient pH. The second enclosure served as a control. 132 A third treatment consisted of an open fiberglass frame of the same dimensions as the enclosure 133 footprint (2 m^2) . It was placed nearby (3 m North of the experimental enclosure) and in the same 134 meadow. It is referred to as a reference plot and accounts for any artifacts from the structure of 135 the enclosures.

The surface component of eFOCE consisted of a buoy that housed solar panels, a wind turbine and 12 V batteries that provided energy to the system. It also housed three CO_2 tanks and a peristaltic pumping system which drew surface seawater into a 20 L container inside the buoy where pure CO_2 was added and mixed until a desired pH was reached (usually between 5.5 and

5.7 pH_T). A Seabird potentiometric 18-S pH sensor was used to monitor pH_T in this surface
container.

142 The two underwater enclosures (experimental and control) were mostly enclosed to 143 maintain the desired pH offset, with the exception of two openings (12 cm) on the upper, side 144 panels. The top of the enclosure could be removed to allow scuba divers to enter when needed. 145 Each enclosure had 10 openings (8 cm diameter) along the bottom sides that allowed tubes to 146 pass through. These 'injection' tubes passed through each enclosure into the ambient environment 147 where they were connected to a set of three underwater brushless centrifugal pumps and a mixing 148 tube (one for each enclosure). For the experimental enclosure, a hose ran from the surface to 149 depth and connected the surface low pH container to the underwater mixing tube. A second peristaltic pump on the buoy controlled the flow rate (up to 0.12 L min⁻¹) of the low-pH water 150 through this hose while the underwater centrifugal pumps (6.7 L min⁻¹ each) continuously brought 151 152 ambient seawater into the mixing tube. Each mixing tube also housed a potentiometric Seabird 153 18-S pH sensor that monitored pH. By sensing the pH of seawater before it enters the enclosure, 154 the system, via a feedback loop, could adjust the CO₂-saturated seawater pumping rate to 155 maintain seawater entering the experimental enclosure at the desired pH offset from ambient. 156 Once seawater reached the subsurface mixing tubes, it then entered the enclosures via the 157 injection tubes described above, where it was circulated by another set of centrifugal pumps (4 per chamber; 6.7 L min⁻¹ each). Water could then exit enclosures through the two openings (12 158 159 cm diameter) on the upper, side panels. The complete renewal time of seawater in each enclosure 160 was ca. 1.5 h.

161 **2.2 Field sensors and system maintenance**

162 The environment was characterized using sensors placed inside the enclosures and placed 163 within 5 m from the reference plot. Sensors were connected by cables to the surface electronic

164 hub. The surface electronic hub communicated 2 min averaged data by radio to the laboratory. 165 Underwater sensors (with their sampling frequency) included 4 potentiometric Seabird 18-S pH 166 sensors (8 measures in 1 s) located inside each enclosure and in each mixing tube, three Seabird 167 37 SMP-ODO CTD with SBE 63 oxygen (O_2) optodes one in each enclosure and one nearby in 168 the ambient (one sample, each, for salinity and temperature every 2 min, two samples for O_2 169 every 2 min), and three LI-COR-192 PAR sensors (2000 irradiance measurements every 5 s) also 170 located in each enclosure and in the nearby ambient environment. 171 The system required routine maintenance. Scuba divers lightly brushed the enclosure 172 surfaces and sensor probes at least once per week to remove sediment and fouling. On four 173 occasions throughout the experiment duration, CTDs were flushed by a syringe filled with clean 174 seawater to remove any debris inside the sampling ports. Tubes and pumps on the buoy and 175 subsea were also cleaned once a week of debris and replaced when heavily fouled. 176 The underwater 18-S pH sensors were calibrated one to three times per month by placing 177 them together in the ambient environment for 45 min, followed by collection of three, 100 mL 178 syringes of seawater drawn directly next to the probes. Seawater was immediately returned to the 179 laboratory and pH determined spectrophotometrically as described in Dickson et al. (2007). 180 Absorbances at peak wavelengths for purified meta-Cresol Purple (Liu et al., 2011) were 181 measured using an Ocean Optics[©] spectrophotometer model USB2000+VIS+NIR. The pH of 182 seawater samples was determined in triplicate (SD < 0.008) at 22 °C and recomputed at *in situ* 183 temperature using the R package seacarb (function pHinsi, Gattuso et al., 2015, seacarb: seawater 184 carbonate chemistry with R. R package version 3.0.2). The offset between the probe-sensed value 185 at the time of water collection and laboratory determined measures was used for correction. In 186 addition, pH sensors were refreshed every four to six weeks in a NBS buffer at pH 4 for 45 min.

187 **2.3 Timeline**

188 The experiment was conducted from April to November 2014. The experimental duration 189 can be divided into three periods: (1) the pre-acidification period, before pH was manipulated, 190 lasted from 5 April to 11 June, (2) the transition period from 12 to 21 June, where pH in the 191 experimental enclosure was slowly lowered by no more than 0.05 units per day until an offset of 192 approximately -0.25 units was reached and (3) the acidified period from 22 June to 3 November 193 during which pH in the experimental enclosure was maintained at the targeted offset of -0.25194 units. It should be noted that the pre-acidification period began on 5 April; however, data from all 195 sensors were available from 15 May. 196 2.4 Environment characterization 197 All sensed data were initially screened for quality. Any obvious outliers or missing data 198 that resulted from system or sensor malfunction were eliminated from the dataset. The mean (± 199 SD) pH_T and median (± median absolute deviation, MAD) diel pH changes for the two enclosures 200 and the ambient environment were calculated by time period and month. 201 Seawater samples for the determination of total alkalinity (A_T) levels in each enclosure 202 were taken one to five times per month from May to October (n = 11 to 12). Samples (300 mL) 203 were filtered on GF/F membranes (47 mm) and immediately poisoned with 100 µL of mercuric 204 chloride (HgCl₂). A_T was determined on triplicate 50 mL subsamples by potentiometric titration 205 on a Metrohm Titrando 888 titrator coupled to a glass electrode (Metrohm, ecotrode plus) and a 206 thermometer (pt1000). The pH electrode was calibrated on the total scale using TRIS buffers of 207 salinity 38, corresponding to salinity in the Bay of Villefranche. Measurements were carried out 208 at 22 °C and A_T was calculated as described by Dickson et al. (2007). During the experiment, 209 standards provided by A. Dickson (batch 132) were used to check precision (standard deviation)

and accuracy (deviation from the certified value provided by Dickson); which was 0.889 and 1.04

 μ mol kg⁻¹ (n = 6), respectively. As A_T variations during the experiment were very small, average 211 $A_{\rm T}$ (mean ± SD, experimental enclosure, n = 12, $A_{\rm T}$ = 2545.5 ± 8.0 µmol kg⁻¹; control enclosure, n 212 = 11, $A_{\rm T}$ = 2541.7 ± 12.2 µmol kg⁻¹) was used to calculate all carbonate chemistry parameters at a 213 214 high frequency, together with sensed temperature, salinity and pH_T, using seacarb. To calculate 215 carbonate chemistry of the ambient environment at high frequency, we used an $A_{\rm T}$ value of 2556 μ mol kg⁻¹ and the sensed ambient values of temperature, salinity, pH_T, using seacarb. This A_T 216 217 value is the mean for 2014 determined from weekly measures of seawater collected at 1 m depth 218 station, Point B, within the Bay (Point B data provided by Service d'Observation Rade de 219 Villefranche and the Service d'Observation en Milieu Littoral). All these parameters, as well as the O_2 concentration (mean \pm SD), median (\pm median absolute deviation, MAD) diel O_2 change 220 and photosynthetically active radiation (PAR, mean \pm SD, mol photons m⁻² d⁻¹) were summarized 221 222 by month and by time period for the two enclosures and the reference plot (ambient).

223

2.5 Shoot density and macrophyte abundance

224 After the enclosures had been deployed on the meadow for four weeks and before the 225 acidification period, scuba divers counted the number of shoots within each treatment. Shoot 226 density was determined twice by different divers and values were averaged, except for the 227 experimental treatment where an observer error was made and one count was eliminated. 228 Permanent quadrats were then used to determine any change in shoot density through time. On 11 April, three 0.25 x 0. 25 m² permanent quadrats were haphazardly placed inside each enclosure 229 230 and in the reference plot. The number of shoots per quadrat was then determined every 2 to 4 231 weeks throughout the experiment.

Percentage cover of benthic macrophytes was estimated every two to four weeks in three to five haphazardly placed, but not overlapping, $0.5 \times 0.5 \text{ m}^2$ quadrats within each treatment. The quadrats were also divided into four smaller squares $0.25 \times 0.25 \text{ m}^2$ to assist with estimation.

Prior to estimation, researchers practiced estimates on the same quadrat location to inter-calibrate
and limit observer bias. On some occasions, the cover and shoot density could not be estimated in
all 9 to 15 quadrat locations in one day. In these instances, divers returned to the treatments
within 15 d (most within 8 d) to complete sampling.

239 **2.6 Leaf biometrics**

240 The number of leaves per shoot, and leaf length, area, thickness and toughness were 241 monitored several times per month from April to November, before and during the acidification 242 period. On these occasions, scuba divers used a tape measure to measure the leaf length and 243 counted the number of leaves per shoot for five to fifteen shoots per enclosure and plot. In 244 addition, approximately every four weeks from 1 August, divers collected eight mature, six 245 intermediate and two to four young leaves from each enclosure and from the reference plot. To 246 limit destructive sampling yet get a baseline measurement, on 27 June (near the start of the 247 acidification period) leaves of about the same age were collected nearby. All leaves were 248 collected from different shoots and taken at their base above the meristem. They were brought 249 back to the laboratory and their length, width, and thickness measured with a tape measure and 250 caliper. The width and thickness was measured at the middle of the length of each leaf. On three occasions (in July, September, and October), the toughness of each leaf was determined in the 251 252 middle of the leaf length with a penetrometer (see Cherrett, 1968).

For all leaf biometric parameters, data collected over several days were pooled into one dataset for a comparison by month and among treatments (experimental, control and reference plot). Lab and field determined leaf lengths were combined and averaged by month into a leaf length parameter that is included graphically. The leaf area is included because it is a frequent meadow descriptor (Pergent-Martini et al., 2005). The leaf length, thickness and toughness were investigated for relatedness with a scatter plot.

259 **2.7 Fluorescence**, photosynthesis, and respiration

A diving pulse amplitude modulated fluorometer (diving-PAM, Walz, Germany) equipped with a red light emitting diode and an internal halogen lamp to provide actinic light, was used to measure the fluorescence in illuminated and dark-adapted leaves *in situ* throughout the experiment. These fluorescence values were used to produce rapid light curves (RLCs, *r*ETR, relative electron transport rate *vs* actinic light) and dark-adapted quantum yields (F_u/F_m).

All fluorescence and photosynthesis measures were performed on a randomly selected secondary leaf from enclosures and reference plot. Dark-adapted yields and RLCs were measured *in situ* between 10-12:00 hr (local time) over two to three consecutive days to produce a sample size of three to ten leaves per enclosure and reference for May (pre-acidification), July, September, and October (acidification period for experimental enclosure). For all fluorescence measures, the fiber optic cable was attached 8 cm above the leaf meristem and held at a standard distance of 3 mm and at a 90° angle from the blade.

272 RLCs were produced following the procedures outlined in Cox and Smith (2015). The actinic irradiance levels ranged up to 895 μ mol photons m⁻² s⁻¹ and were applied on the leaf 273 274 surface for 10 s followed by a 0.8 s saturating pulse. Actinic range was also adjusted by month to 275 account for the changing abilities of plants and corrected each time for battery decline. We 276 determined the absorption factor (AF), used in *r*ETR calculations, following the methods and 277 assumptions described in Beer and Björk (2000). Measurements were conducted one to three 278 times each sampled month and monthly averages were used in calculations. Curves were fitted 279 with the exponential model proposed by Platt et al. (1980). Parameters derived from the curves include (1) α , the initial slope before the onset of saturation (µmol electrons m⁻² s⁻¹ / µmol 280 photons $m^{-2} s^{-1}$, (2) the relative maximum electron transport rate, $rETR_{max}$ (µmol electrons $m^{-2} s^{-1}$) 281

¹) and (3) E_k , optimal irradiance for maximal electron transport (μ mol photons m⁻² s⁻¹) which is determined by the equation $E_k = r \text{ETR}_{\text{max}} / \alpha$.

For dark-adapted quantum yield, leaves were placed in the dark for five minutes using the dark-adapter then leaves were exposed to a 0.8 s white saturating light pulse (saturation intensity setting of 8). Then the maximum PSII quantum yield was calculated using the equation Genty et al. (1989) for dark adaption.

288 In addition, the photosynthesis versus irradiance (PE) curves of experimental and control 289 leaf segments were produced in the laboratory using O₂ evolution within a series of incubations. 290 These incubations were performed over two consecutive days in September and November to 291 produce four PE curves per enclosure each month. Leaf segments (5 cm) collected from ~10 cm 292 length leaf were collected from the enclosures in the morning and incubated in the afternoon 293 (13:00 - 19:00 h, local time). Immediately after collection, leaves were stored underwater in 294 plastic bags, and transported to the laboratory in a dark mesh bag. Leaves were held for up to 3 h 295 in dim light within a temperature-controlled laboratory (20 °C) in two open top cylindrical 296 aquaria (1.5 L). Ambient water from the nearby bay was pumped into two header tanks that fed 297 the aquaria and allowed excess water to overflow into a drainage basin. The pH in one header 298 tank was maintained at a pH_T of ~7.8, corresponding to pH levels in the experimental enclosure 299 by metered additions of pure CO₂ controlled by a pH-stat system (IKS, Aquastar Aquatic 300 Products).

After carefully removing all epiphytes, segments were individually placed inside 60 mL biological oxygen demand (BOD) bottles submerged into a 50 L aquarium maintained 1 to 2° C to the mean monthly seawater temperature at the time of collection (21.2 °C \pm 0.2 SD). BOD bottles were filled between each incubation with fresh seawater from the respective header tank (ambient, or lowered pH) with a stirrer below. Light was provided at a 90° angle to the leaf

306	surface by a 250 W metal-halide lamp and adjusted to nine increasing irradiance levels (5 to 200
307	μ mol photons m ⁻² s ⁻¹ measured directly at the leaf surface). This range of irradiance is within and
308	above irradiance observed at the depth of collection. Plants were maintained at each irradiance or
309	in darkness (to measure respiration, R) for 15-30 min while the concentration of O_2 was
310	continuously monitored with a PreSens OXY-4 O2 meter with PSt3 fiber-optic mini-sensors.
311	After the incubations, leaf segments were ground in a chilled room using a glass homogenizer
312	with 90% acetone that had been previously chilled for 12 h. The extract was left for 24 h in
313	darkness, centrifuged at 3000 rpm for 15 min, and the absorbance of the supernatant measured in
314	quartz-glass cuvette with a UV/VIS spectrophotometer (Lambda 2, Perkin 366 Elmer). The
315	concentrations of Chl a and b were determined by measuring the absorbance at 647 and 664 nm
316	and the concentrations calculated from the equations in Jeffrey and Humphrey (1975).
317	Rates of changes in O_2 normalised to total chlorophyll (Chl a + b) were plotted against
318	irradiance levels. Parameters of the PE curves were estimated using an hyperbolic tangent model
319	(Jassby and Platt, 1976), assuming that R is similar in the light and dark:
320	$P_{net} = P_{g \max} x \tanh (-E/E_k) + R$
321	with:
322	P_{net} : rate of net photosynthesis (μ mol O ₂ (mg Chl) ⁻¹ min ⁻¹)
323	$P_{g max}$, rate of maximal gross photosynthesis ($\mu mol O_2 (mg Chl)^{-1} min^{-1}$)
324	E, irradiance (μ mol photons m ⁻² s ⁻¹)
325	E_k , irradiance at which α intersects $P_{g max}$ (µmol photons m ⁻² s ⁻¹)
326	R, respiration rate
327	The initial slope, $\alpha \ (\mu mol O_2 \ (mg \ Chl)^{-1} \ min^{-1} / \mu mol \ photons \ m^{-2} \ s^{-1})$ was calculated as P_g
328	$_{max}$ / E_k and E_c , the irradiance at which gross photosynthesis equals respiration and above which
329	plants exhibit a positive net photosynthesis, was determined from R/α .

330

2.8 Growth and biomass

331 Leaf production and leaf plastochrone interval were determined using the Zieman method 332 modified by Short and Duarte (2001). Three to eight shoots in both enclosures and in the 333 reference plot were marked with a plastic tag with a unique number in July, August, and 334 September. A hypodermic needle was used to punch a hole in the meristem region. These tagged 335 shoots were again located 33 to 46 d later. The distance from the puncture to the meristem was 336 measured and any new leaves that lacked a puncture were enumerated. Using these methods, it 337 was possible to calculate the number of days to produce a new leaf (plastochrone interval) and 338 leaf production per day for each shoot. Leaf production incorporates the new length added to the 339 shoot from both, the newly produced leaf (or leaves) and from the growth of older leaves.

340 Above-ground and below-ground biomass was determined for each enclosure and for the reference plot at the conclusion of the study. A fourth 2 m^2 area was also sampled for biomass in 341 342 a nearby seagrass habitat located approximately 6 m from the enclosures. This area was added to 343 further account for natural spatial variation. Three to five, 10 cm diameter cores of 12 cm height 344 were hammered into haphazardly selected locations within the treatment area. They were brought 345 back to the laboratory, stored in 5% formalin, and later sorted into above-ground and below-346 ground plant parts, blotted dry, and weighed. An one-way ANOVA was used to test for 347 differences in above- and below- ground biomass when data met parametric requirements.

348 **2.9 Pseudoreplication**

349 Samples were collected or measured inside the plot or enclosure through time, often both 350 before and after the pH manipulation. Thus the replication is equal to one for each treatment. 351 True replication was sacrificed at the expense of controlling pH as an offset, at the spatial scale of 352 the plants. Traditional inferential statistics could, therefore, not be rigorously applied and we

353 compare results graphically, paying careful attention to any divergence in values between the354 enclosures and the reference plot.

355 **3 Results**

356 **3.1 Environment characterization**

The pH in the experimental enclosure was maintained at a -0.26 unit offset from the control enclosure during the acidification period (Table 1). Values summarized by month showed that the difference between the two enclosures was maintained close to the targeted offset (range: -0.22 to -0.29 pH units). Before the pH was manipulated the offset between enclosures was smaller, -0.1 to 0.06 pH units.

The pH_T in ambient ranged from a mean of 7.98 (\pm 0.06 SD) in September to 8.11 (\pm 0.04 SD) in June (Table 1). The ambient pH_T was similar to the pH_T in the control enclosure, which ranged from 7.97 (\pm 0.07 SD) in September to 8.12 (\pm 0.06 SD) in June. The greatest difference between control and ambient, in monthly mean pH_T values was 0.06 units. The differences in *p*CO₂ reflected the magnitude of difference in pH_T, as *A*_T levels were rather constant during the study (see method section).

368 The mean O₂ concentration was similar in enclosures and in the ambient (Table S1). For 369 example, the mean O_2 concentration (\pm SD) before acidification for ambient, control and experimental respectively was 258 ± 18 , 254 ± 34 , $258 \pm 32 \mu \text{mol kg}^{-1}$. In the ambient and in the 370 371 enclosures, the O₂ concentration fluctuated over the course of the day (data not shown). After 372 sunset, O₂ concentration declined to a night-time minimum. In the morning, the O₂ began to 373 increase to a daily afternoon maximum; then it declined with decreasing irradiance. Over the months of the experiment, this diel O_2 change ranged from 21 to 72 μ mol kg⁻¹ in the ambient, 34 374 to 95 μ mol kg⁻¹ in the control enclosure, and 34.5 to 100.5 μ mol kg⁻¹ in the experimental 375 376 enclosure (Table 1). The difference in diel change between the ambient and the enclosures was

most likely due to the amplification of a metabolic signal inside a partially enclosed space (similar to the example of a larger O_2 fluctuation when a similar sized plant is contained in a relatively smaller volume of water) as was evidenced by the more similar, and greater diel change in the two enclosures. The largest difference in median values between enclosures was 14 μ mol kg⁻¹ in May, prior to the perturbation.

382 The diel pH_T change in the meadow corresponded to the daily change in O₂ concentration. 383 The natural diel pH_T for this meadow was evident from the measures in the ambient which 384 median values show it fluctuated by 0.09 (\pm 0.02 MAD) and 0.08 (\pm 0.02 MAD) units per day in 385 the pre- and acidification period, respectively. The diel change in pH_T for the control enclosure 386 was slightly greater but consistent in the pre- and during acidification period $(0.14 \pm 0.06 \text{ MAD})$ 387 and 0.14 \pm 0.06 MAD). In contrast, the diel pH_T change for the experimental enclosure increased 388 from a median of 0.16 (\pm 0.06 MAD) before pH manipulation to 0.28 (\pm 0.14 MAD) during the 389 acidification period.

390 Monthly differences were evident particularly for temperature, oxygen concentration, and 391 PAR (Table S1) but were similar in the ambient, control and experimental enclosures. For 392 example, the mean \pm SD during the acidification period for temperature in ambient, control and 393 experimental enclosures was 23.9 °C \pm 0.01 (for each) and for PAR, 4.6 \pm 1.9, 4.6 \pm 2.0, 4.1 \pm 1.7 mol photons m⁻² d⁻¹, respectively. Temperature increased approximately by 6 °C from May 394 395 through August and declined by 4 °C until November. Oxygen concentrations and PAR 396 fluctuated similarly with higher values in May to August (mean monthly range: 212 to 270 µmol kg⁻¹, 4.7 to 7.7 mol photons $m^{-2} d^{-1}$) and decreases in September to November (mean monthly 397 range: 193 to 211 μ mol kg⁻¹, 1.4 to 4.4 mol photons m⁻² d⁻¹). 398

399 **3.2 Shoot density and macrophyte abundance**

Initial shoot densities were similar in both enclosures and reference plot and ranged from 150 to 175 shoots m⁻² (Fig. 2). There was no obvious change in shoot number (as determined in permanent quadrates re-sampled through time) related to the lowered pH in the experimental enclosure. For both enclosures and the reference plot, the number of shoots (initially 6 to 27 in permanent quadrates) tended to decline with time.

405 The reference plot as well as the enclosures had very low diversity of benthic 406 macrophytes as measured by estimates conducted within haphazardly placed quadrats at each 407 sampling interval (Fig. 2). Posidonia oceanica was the dominant species, with a surface cover of 408 18 to 35%. Peyssonnelia, a red alga, covered between 1 and 11% of the substratum. Their 409 abundances were similar between months throughout the experiment. There was a slightly greater 410 percentage of *P. oceanica* in the experimental enclosure (experimental enclosure, $31.6 \pm 0.6\%$; 411 control enclosure, $27.9 \pm 1.7\%$; reference plot, $28.9 \pm 1.3\%$) throughout the experiment duration 412 that did not appear to be related to the timing of the pH manipulation.

413 **3.3 Leaf biometrics**

There was no large difference in shoot height among the enclosures and reference plot but there were large differences in shoot height between the sampled months (Fig. 3). A similar monthly pattern in leaf length was observed between the three treatments, for the minimum, average and maximum leaf length. From April through August, average leaf length and average shoot height both increased and then declined between August and September. For example, the overall average shoot height increased from 40.6 cm in April to 73.4 cm in August then declined to 24.8 cm in November.

421 Shoots had between 2 and 8 leaves (Fig. 3). The reference and control plants differed 422 slightly in leaf number per shoot (reference, 5.4 ± 0.1 vs control, 5.1 ± 0.1), but control and 423 experimental plants (5.2 ± 0.2) were highly similar indicating an absence of pH effect.

Furthermore, the number of leaves per shoot in the experimental enclosure did not consistently
increase or decrease after the pH was manipulated. Instead, leaf number per shoot in enclosures
and plot increased during months when leaf height was lower (April, May and then October,
November: 6 to 7) and tended to be lower in June and August (4 to 5) when leaf height was
elevated.

429 Leaf thickness and leaf toughness increased with leaf length. However, leaf thickness did 430 not appear to be correlated to leaf toughness. Both parameters varied among the enclosures and 431 reference plot, and between months (Fig. 3). Leaves in the experimental enclosure were slightly 432 thicker $(2.5 \pm 0.1 \text{ mm})$ than leaves in the control enclosure and the reference plot $(2.2 \pm 0.08 \text{ mm})$ 433 and 2.1 ± 0.1 mm, respectively). Leaves in control enclosure and the reference plot for the month 434 of November had decreased and more variable thickness. Leaves inside the enclosures appeared 435 to be weaker than the leaves in the reference plot. Furthermore, leaves appeared weaker in 436 October compared to July and September. For example, the weakest leaf in July could withstand 437 34 g of force as compared to the weakest leaf in October which could only withstand 12 g of 438 force. The ambient leaf samples collected in June also had a greater mean value of toughness than 439 the October values from enclosures and the reference plot.

440 **3.4 Fluorescence**, photosynthesis, and respiration

The dark-adapted quantum yield obviously differed by month but not according to pH
(Fig. 4). The overall dark-adapted quantum yield ranged from 0.72 to 0.88 (n = 69). The mean
values were similar in the enclosures and the reference plot. Mean yield was 0.8, 0.789, and 0.799
for leaves measured in the experimental, control, and reference treatments, respectively. Yield
values increased over the duration of the experiment.

The AF factor for the calculation of *r*ETR changed with month. The determined values (as a mean \pm SD) were as follows: May: 74.5; July: 65.0; September: 69.6 \pm 1.5 (n = 3); October, 54.2 \pm 0.0 (n = 2).

449 The photosynthetic RLCs in Fig. 4 (A-D), show that the shape of the curve changed with 450 month. Leaves from the control and experimental enclosures have similar *r*ETR values that were 451 slightly lower at elevated irradiance relative to the leaves in the reference plot.

The initial slope (α , μ mol electrons m⁻² s⁻¹ / μ mol photons m⁻² s⁻¹) ranged between 0.23 452 453 and 0.58 (n = 57). It decreased substantially as a function of time (Table S6) from elevated values 454 in May (0.43 ± 0.01) and July (0.48 ± 0.01) to lower values in September (0.31 ± 0.01) and October (0.27 \pm 0.01). Overall (n = 57), rETR_{max} values (in μ mol electrons m⁻² s⁻¹) ranged from 455 4.3 to 27.4 and E_k (µmol photons m⁻² s⁻¹) ranged from 12.0 to 63.6. The leaves from the reference 456 plot had $r \text{ETR}_{\text{max}}$ (12.3 ± 0.6) and E_k (33.7 ± 2.0) that were more different than the leaves from 457 458 the control (rETR_{max} = 10.8 \pm 0.7, E_k = 29.8 \pm 2.0) and experimental (rETR_{max} = 12.0 \pm 0.7, E_k = 459 30.9 ± 0.7). However, these parameters differed by a greater amount by month than among plants 460 from the enclosures and the plot. $rETR_{max}$ values were substantially higher in May (22.1 ± 1.4) 461 than in July (10.9 \pm 0.8), September (7.2 \pm 0.6), and October (7.5 \pm 0.8). Overall, E_k was 462 obviously greater in May (50.2 ± 2.2) than in July (23.1 ± 2.6) , September (24.5 ± 2.1) , and 463 October (28.1 ± 2.5) .

464 The parameters of the PE curves of leaves collected from the experimental and control 465 enclosures also did not greatly differ (Fig. 5). α , P_{g max}, and R were greater for leaves measured in 466 September than November.

467 The mean total concentration of chlorophyll in leaves did not greatly differ among 468 enclosures. It was 0.36 ± 0.04 , 0.38 ± 0.04 mg Chl cm⁻² in the experimental and control 469 enclosures, respectively. It was greater in November than in September (0.46 ± 0.03 vs $0.28 \pm$ 470 0.04 mg Chl cm⁻²). The Chl *a:b* ratio of leaves from the control and experimental enclosures did
471 not obviously differ, with an overall Chl *a:b* ratio of 0.64.

472 **3.5 Growth and biomass**

473 Leaf production and plastochrone interval of shoots in the reference plot and in the 474 enclosures appeared to differ (Fig. 6). Differences are most congruent with an effect caused by 475 the structure of enclosures and not from the lowered pH. The shoots in the reference plot were 476 able to produce more leaf material than in the experimental and control enclosures. From July to September, reference shoots grew new leaf material at a mean rate of 0.89 (\pm 0.06) cm d⁻¹ 477 478 compared to the reference plot and control enclosure, which both produced 0.66 (\pm 0.05 to 0.06) cm d⁻¹. Furthermore, reference shoots produced a new leaf in a fewer number of days than shoots 479 480 in the experimental and control enclosures. From August to September, it took 11 days to 481 produce a new leaf in the reference plot while it took between 23 to 29 days for shoots that grew 482 in the experimental and control enclosures, respectively. Overall, leaf production (the growth of 483 all leaves per shoot) was also seasonal. It was greater per day from September to October (1 cm d⁻ ¹) than during the periods July-August (0.5 cm d⁻¹) and August-September (0.6 cm d⁻¹). 484 485 At the end of the experiment, the above- and below-ground biomass was highly variable

(Fig. 6). The above- and below-ground biomass ranged from 318 to 1484 and from 348 to 1584 g FW m⁻², respectively. The control and experiment enclosures tended to have less above-ground biomass (630 and 530 g FW m⁻²) than the two external plots (reference: 850 and extra ambient plot: 870 g FW m⁻²).

490 4 Discussion

491 No overwhelming impact was observed on macrophyte abundance, *P. oceanica* leaf 492 biometrics, biomass, and photosynthesis after four months of elevated pCO_2 . Leaf thickness may 493 change in response to lowered pH but requires further testing. Many of the leaf biometrics and

494 physiology parameters varied seasonally with the varying temperature and irradiance. Posidonia 495 *oceanica* abundance did not substantially change over eight months as expected for a seagrass 496 with slow rates of colonization (Marbà and Duarte 1998). However, under elevated pCO_2 , no 497 other benthic macrophyte or epiphyte proliferated or decreased to alter the macro-community 498 structure. The similarity in leaf biometrics, photosynthesis, biomass and growth between 499 enclosures support the conclusion of limited stimulation for P. oceanica under future ocean 500 acidification. However, due to tradeoffs related to experimental design, there were limitations to 501 our conclusions.

502 Thickness and toughness are two structural factors related to mechanical strain (Harder et 503 al., 2006; Littler and Littler, 1980; Padilla, 1985) and both traits were altered. Flexibility and 504 strength are needed in environments with strong wave forces (de los Santos et al., 2013). In 505 Cymodocea nodosa, another Mediterranean seagrass, leaf cross-sectional area varies with 506 hydrodynamical forces (de los Santos et al., 2013). Therefore, observed differences in leaf 507 toughness for plants maintained in the enclosures support the notion that mechanical abrasion was 508 less than in ambient. This finding is an artifact of the structure that could not be avoided. In P. 509 oceanica, thickness changes along the leaf axis and leaves are thinner with depth (Colombo et al., 510 1983). Given that the experiment was conducted at the same depth and leaves were measured at 511 their center, it is interesting to note that leaf thickness was greatest for the shoots collected from 512 the experimental enclosure and that this effect was driven by measures in November. An increase 513 in seagrass leaf thickness would be an opposing effect to those observed for the upright calcified 514 alga, Acetabularia acetabulum, which lost skeletal support under ocean acidification conditions 515 (Newcomb et al., 2015). There are several possible interpretations of these results. First, leaves at 516 the lower pH may have increased their carbon content as observed for below-ground plant 517 structure of the seagrass *Thalassia testudinum* under elevated pCO_2 (Campbell and Fourgurean,

518 2013a). Secondly, lowered pH could result in a delay of leaf shedding. Plants from the 519 experimental enclosure had a tendency towards relatively greater leaf length and maintenance of 520 number of leaves in November. A prolonged leaf life-span could allow plants to scavenge 521 nutrients from senescing leaves to maintain C/N ratio (Gobert et al., 2002). However, 522 photosynthesis measures were not elevated by the lowered pH and thus there would be no need 523 for increased nutrients. Additionally, increased pCO_2 and high light increased leaf shedding for the seagrass Amphibolis antarctica (Burnell et al., 2014). The response was linked to proliferation 524 525 of filamentous epiphytes, which did not occur in this study. Alternately, increased leaf thickness 526 could be the result of chance. The plausible relationship warrants further investigation in field 527 experiments with prolonged duration and increased replication.

528 If indeed leaf thickness increases with ocean acidification, it is unclear how this would 529 impact herbivore feeding. The main herbivores, the fish, Sarpa salpa, and the sea urchin 530 *Paracentrotus lividus*, feed preferentially on the adult and thicker leaves (Peirano et al., 2001). 531 These herbivores were prevented from grazing in enclosures. Arnold et al. (2012) have reported 532 increased rates of fish grazing on the plant at proximity of a CO_2 vent, presumably due to the 533 significant decreases in the production of phenolics. To date, very few studies have focused on 534 plant-herbivore interactions under elevated pCO₂ levels (Asnaghi et al., 2013; Campbell and 535 Fourgurean, 2014; Poore et al., 2013) and as plant-herbivore interactions were not the focus of 536 this study, it is not known how this would have impacted the results.

To our knowledge, this is the first *in situ* study to repeatedly and over several months (6) measure *P. oceanica* fluorescence to find that the second rank leaves showed a typical seasonal pattern of plant acclimation (Boardman, 1977). Leaves were more sun-adapted (relatively higher *r*ETR_{max} and E_k) in periods with elevated irradiance and more shade-adapted when irradiance and photoperiod were reduced. The relatively lowered Fo/F_m in May and July compared to October

542 indicates a down-regulation of PSII activity (Campbell et al., 2003; Henley, 1993) that

543 corresponds with elevated irradiance in warmer months. Findings are in agreement with Figuero 544 (2002) where ETR and E_k were higher in September than in February. Although there have been 545 some concerns on the ability of fluorescence techniques to indicate seagrass carbon stimulation 546 (see Cox et al., 2015; Jiang et al., 2010), *P. oceanica* productivity as a function of increasing 547 irradiance was in agreement with fluorescence results.

548 The results of the present study add to the growing evidence that the pH change predicted 549 over the next century may result in limited production stimulation for *P. oceanica*. The 550 relationship between pH and P. oceanica photosynthesis was established over wide range of pH_T 551 from 9.0 to 7.9 (scale unknown, Invers et al., 1997), or with more extreme low levels (6.98 pH_T, 552 Hall-Spencer et al., 2008; 7.5 scale unknown, Invers et al., 2002). Within the range 7.9 to 9.0, the 553 slope of the pH-photosynthesis relationship was significant but, the two variables were 554 moderately related (Inverset al., 1997). Along CO_2 vents, there was no indication of 555 photosynthetic stimulation at stations with a pH range of 6.98 to 8.17 but, shoot density was 30% 556 greater than nearby areas at the lowest mean pH station (Hall-Spencer et al., 2008). In a 557 laboratory incubation of *P. oceanica* shoots with their attached epiphytes, at a similar pH_T as this 558 study (~7.7-7.8), there was also limited stimulation of productivity (Cox et al., 2015). Similarly, 559 modeled outcomes from laboratory studies of leaf segments by Invers et al. (1997, 2001) 560 predicted that elevating pCO_2 by the amount used in this experiment would increase productivity 561 by only 10%. This first *in situ* experiment confirms previous results obtained on isolated plants or 562 leaf segments in the laboratory and is interpreted as in agreement with observations at CO₂ vents. 563 *Posidonia oceanica* has shoot lifespan estimated up to 50 years (Gobert et al., 2006). In 564 carbon budgets there is thought to be asynchrony between fixation (photosynthesis) and use 565 (respiration or growth), which is balanced by the storage of carbohydrate reserves (Alcoverro et

566 al., 2001). Because of this asynchronicity, the photosynthetic benefit of CO₂ may translate into 567 the following season or year as it did for the seagrass Zostera marina (Palacios and Zimmerman, 568 2007). In the present study, there was no indication of increased productivity as gauged by RLCs, 569 PE curves, and measures of leaf chlorophyll. Therefore there is no available evidence that carbon 570 availability translated into increased carbon storage as occurred for *T. testudinum* under elevated 571 pCO_2 (Campbell and Fourgurean, 2013a). Carbohydrates can be translocated to other ramets 572 (Marbà et al., 2002) which can lessen observed effects but, in this case, enclosure area captured 573 the 20 cm maximum translocation distance detected by Marbà and Duarte (1998) and edges 574 severed (designed to penetrate ~ 8 cm) several outside to inside shoot connections. The most 575 productive period for above-ground growth occurred from April to August; a pattern consistent 576 with increased growth induced from the greater availability of both light and nutrients in early 577 spring and increased storage in July to August (Alcoverro et al., 1995, 1998, 2001; Bay, 1984; 578 Duarte, 1989; Ott, 1980). Therefore it is possible that if the experiment were initiated earlier, in a 579 period more conducive for biomass production, or prolonged to capture any lagging effects the 580 outcome may have been different. Only two of six studies support a pulsed seasonal-pH 581 interaction that could result in long-term gains yet, these were found at pH < 7.7 (see Hall-582 Spencer et al., 2008; Invers et al., 2002).

We caution that conclusions should not be applied to other seagrasses and that outcomes may vary with differences in community composition and environment. Presumably due to differences in their evolutionary past, some species are comparatively more responsive to lowered pH (Campbell and Fourqurean, 2013b; Invers et al., 2001; Koch et al., 2013). *Posidonia oceanica* is less sensitive to pCO_2 and can rely heavily on bicarbonate compared to two other Pacific seagrass species (Invers et al., 2001). In addition, at CO₂ seeps in Papua New Guinea, two seagrass species (*Cymodocea serrulata* and *Halophila ovalis*) occur in mixed stands and while

590 both species had increased productivity along the lowered pH gradient, it was only C. serrulata 591 with dense below ground biomass that had increased abundance (Russell et al., 2013); 592 demonstrating that outcomes may be species specific, related to the plant physiology and 593 structure, and vary with competition. Biological communities and environmental conditions are 594 variable both within (e.g. depth) and among meadows (Hemming and Duarte, 2000). For 595 example, epiphyte coverage and thus level of competition were reported to be greater along 596 control stations at Ischia, Italy (Martin et al., 2008) than in our study site, however, differences in 597 methodology prevent direct coverage comparisons. Nutrient concentration can also alter the 598 response of seagrass to CO₂ additions (Burnell et al., 2014; Martínez-Crego et al., 2014). Clearly 599 our understanding of meadow dynamics under ocean acidification conditions could benefit from 600 repeated *in situ* studies that address issues such as species differences, more prolonged durations, 601 herbivore-plant interactions, and temporal and spatial effects.

602 Performing this experiment *in situ*, over several months, is an advancement for 603 understanding the response of *P. oceanica* to ocean acidification. The eFOCE design has 604 advantages to other mesocosm systems such as its large size which allows for measuring 605 processes at the scale of a meadow, its ability to monitor the environment in real-time, and its 606 ability to maintain pH as an offset. Though replicated enclosures would have been preferred and 607 are recommended for future use, their implementation was not feasible at this stage. However, 608 several steps were taken to eliminate possible erroneous conclusions including: (1) the 609 environment was continuously monitored to ensure comparisons were valid, (2) repeated 610 measurements were made at the same location through time both before and after acidification (3) 611 comparisons from the pH manipulated enclosure were made to at least two different spatial 612 locations and (4) results obtained in laboratory and natural experiments were compared and are in 613 general agreement. The duration of this study was longer than any previous pH perturbation

614 carried out on *P. oceanica* and it was performed in the most natural conditions possible. This
615 study addresses a need for manipulative experiments done *in situ* for longer durations to make
616 best predictions of future marine ecology (see Gattuso et al., 2014).

617 Our findings have implications for the function of future meadows. Seagrasses through 618 their metabolic activity alter the chemical properties of the meadow. In daylight, seagrasses draw 619 down the available dissolved inorganic carbon and at night their respiration has the opposite 620 effect (Hendriks et al., 2014a). The daily change in pH has been shown to be up to 0.24 pH units 621 and to be related to the density and length of leaves (Hendriks et al., 2014a). In the current study, 622 the decline in leaf length and 3°C difference in temperature likely contributed to the decline of 623 ambient pH_T from 8.10 to 8.01 from May to November. Hendriks et al. (2014b) has suggested 624 that (1) organisms within the meadow may not be as vulnerable to ocean acidification because 625 they are adapted to large diel pH changes (2) the productivity of *Posidonia* during the day may 626 buffer the impacts of ocean acidification, particularly for calcifiers by providing a daily window 627 of maximum calcium carbonate saturation where calcification can be more efficient and (3) ocean 628 acidification could stimulate seagrass productivity and thus increase buffering capacity; which 629 was not supported by the results of this present study. Considering the two previous proposed 630 hypotheses, the median diel pH variation for the meadow in this study was ~0.1 and also 631 appeared to be driven by plant metabolism. However, the median diel pH range in the 632 experimental enclosure was two to three times larger than the control (0.09 to 0.29 pH units) and 633 exhibited greater variability; a finding that would be missed in typical experiments which lower 634 pH and maintain it at a constant future level(s). The variation in diel pH cannot solely be 635 explained by O_2 fluxes. The increased diel pH fluctuation could largely be the result of the 636 reduced buffering capacity of seawater at lowered pH (Shaw et al., 2013). The lowered and larger

637	diel pH variation and lack of productivity stimulation casts doubt on the adaptability of organisms
638	to future pH change and the ability of a <i>P. oceanica</i> meadow to serve as a future refuge.
639	Ocean acidification is not occurring in isolation, warming has been predicted to result in a
640	complete extinction of <i>P. oceanica</i> meadows by the year 2049 (Jordà et al., 2012). The
641	speculation that increased CO_2 availability would enhance seagrass production and help to

642 alleviate thermal stress (Zimmerman et al., 2015) was not supported. Jordà et al. (2012) also

643 draws attention to the continuing decline of *P. oceanica* meadows from 1990 despite the increase

644 in CO₂ as a demonstration of the limited capacity of ocean acidification to buffer seagrass

645 vulnerability to disturbances. It confirms observations after an explosive episode at a CO₂ vent

646 which resulted in an extreme lowering of pH (4.7 to 5.4) and elevated temperatures (28-30 °C, 3

647 to 5 °C above ambient). Along this vent, *P. oceanica* experienced a decrease in growth that

648 persisted for three years (Vizzini et al., 2010). The extreme nature of the vent activity,

649 confounding biological differences found at vent sites (e.g. Vizzini et al., 2013), and the possible

650 change in physiology under combined stressors make it difficult to predict future meadow

651 ecology. It underscores the need to investigate stressors concurrently and *in situ*. The FOCE

652 systems are tools that can be used to investigate these types of impacts.

653 4.1 Summary, caveats, and perspectives

Any benefit from ocean acidification, over the next century, on *Posidonia* physiology and growth appears minimal. This conclusion is supported by the similarity of measures between enclosures and in context of results from other studies. We have cautioned that the eFOCE study, like all studies, has limitations. There may be small gains in plant productivity which are masked by an enclosure effect or difficult to identify without replication or more prolonged duration. We recommend that future *in situ* manipulative efforts use FOCE systems to control pH as an offset, as we did, and increase replication. The field of ocean acidification and future seagrass ecology

661 could benefit from further in situ experiments that focus on combined stressors, extended 662 experiment duration, and differences which occur over varying spatial and temporal scales (eg. 663

within a season promoting above-ground biomass).

664 Author contribution

665 All authors contributed to the research in this manuscript. J.-P. Gattuso and F. Gazeau 666 were co-principle investigators that had the idea, oversaw the project, and were involved in data 667 collection. P. Mahacek was responsible for eFOCE system design. A. Le Fur ensured the system 668 functioned with assistance from S. Alliouane, T.E. Cox, J.-P. Gattuso, and F. Gazeau. T.E. Cox 669 was responsible for the seagrass protocol and data collection with assistance from S. Alliouane 670 and advice given by I.E. Hendriks who contributed to fluorescence measures. T.E. Cox wrote the 671 manuscript with J.-P. Gattuso and F. Gazeau and all other authors contributed editorial

672 comments.

673 **Acknowledgements**

674 We would like to acknowledge the following people who assisted in the laboratory, in the 675 field, or with system engineering or maintenance: E. Beck Acain, J. Acain, J. Delille, L. van der 676 Heijden, M. Maillot, F. Moullec, S. Schenone, L. Urbini, K. Walzyńska. We also thank J.-J. 677 Pangrazi, R. Patrix, and E. Tanguy for aide in construction of the enclosures. Éric Béraud, G. de 678 Liege, D. Luquet, L. Mangialajo, S. Reynaud, and D. Robin kindly assisted in diving activities. 679 We are grateful to C. Ferrier-Pagès and her research team for use of their PAM fluorometer. We 680 also thank B. Kirkwood at Monterey Bay Aquarium Research Institute who advised in system 681 design. We thank the Service d'Observation Rade de Villefranche and the Service d'Observation 682 en Milieu Littoral for their kind permission to use Point B data. We also thank the Service 683 National d'Analyse des Paramètres Océaniques du CO₂ for performing the determination of A_T at 684 Point B. This work was funded by the 'European Free Ocean Carbon Enrichment' (eFOCE; BNP

- 685 Paribas Foundation), the European Commission through the project 'Mediterranean Sea
- 686 Acidification in a changing climate' (MedSeA; grant agreement 265103) and the MISTRALS-
- 687 MERMEX (INSU, CNRS) program.
- 688 References
- Alcoverro, T., Duarte, C. and Romero, J.: Annual growth dynamics of *Posidonia oceanica*:
 contribution of large-scale versus local factors to seasonality, Mar. Ecol. Prog. Ser., 120,
 203–210, doi:10.3354/meps120203, 1995.
- Alcoverro, T., Manzanera, M. and Romero, J.: Seasonal and age-dependent variability of
 Posidonia oceanica (L.) Delile photosynthetic parameters, J. Exp. Mar. Biol. Ecol.,
 230(1), 1–13, 1998.
- Alcoverro, T., Manzanera, M. and Romero, J.: Annual metabolic carbon balance of the seagrass
 Posidonia oceanica: the importance of carbohydrate reserves, Mar. Ecol. Prog. Ser., 211,
 105–116, 2001.
- Apostolaki, E. T., Holmer, M., Marba, N. and Karakassis, I.: Metabolic imbalance in coastal
 vegetated (*Posidonia oceanica*) and unvegetated benthic ecosystems, Ecosystems, 13,
 459–471, 2010.
- Arnold, T., Mealey, C., Leahey, H., Miller, A. W., Hall-Spencer, J. M., Milazzo, M. and Maers,
 K.: Ocean acidification and the loss of phenolic substances in marine plants, PLoS ONE,
 703 7(4), e35107, doi:10.1371/journal.pone.0035107, 2012.
- Asnaghi, V., Chiantore, M., Mangialajo, L., Gazeau, F., Francour, P., Alliouane, S. and Gattuso,
 J.-P.: Cascading effects of ocean acidification in a rocky subtidal community, PLoS ONE,
 8(4), e61978, doi:10.1371/journal.pone.0061978, 2013.
- Bay, D.: A field stuy of the growth dynamics and productivity of *Posidonia oceanica* (L.) Delile
 in Calvi Bay, Corsica, Aquat. Bot., 20(1-2), 43–64, doi:10.1016/0304-3770(84)90026-3,
 1984.
- Beer, S. and Björk, M.: Measuring rates of photosynthesis of two tropical seagrasses by pulse
 amplitude modulated (PAM) fluorometry, Aquat. Bot., 66(1), 69–76, 2000.
- Boardman, N. K.: Comparative photosynthesis of sun and shade plants, Annu. Rev. Plant
 Physiol., 28(1), 355–377, doi:10.1146/annurev.pp.28.060177.002035, 1977.
- Borowitzka, M. A., Lavery, P. S. and van Keulen, M.: Epiphytes of seagrasses, in Seagrasses:
 Biology, Ecology and Conservation, edited by A. W. D. Larkum, R. J. Orth, and C. M.
 Duarte, pp. 441–461, Springer, Dordrecht, The Netherlands., 2006.

- Burnell, O., Russell, B., Irving, A. and Connell, S.: Seagrass response to CO₂ contingent on
 epiphytic algae: indirect effects can overwhelm direct effects, Oecologia, 176, 871–882,
 2014.
- Campbell, J. E. and Fourqurean, J. W.: Novel methodology for *in situ* carbon dioxide enrichment
 of benthic ecosystems, Limnol. Oceanogr. Methods, 9, 97–109,
 doi:10.4319/lom.2011.9.97, 2011.
- Campbell, J. E. and Fourqurean, J. W.: Effects of *in situ* CO₂ enrichment on the structural and
 chemical characteristics of the seagrass *Thalassia testudinum*, Mar. Biol., 160, 1465–
 1475, 2013a.
- Campbell, J. E. and Fourqurean, J. W.: Mechanisms of bicarbonate use influence the
 photosynthetic carbon dioxide sensitivity of tropical seagrasses, Limnol. Oceanogr., 58,
 839–848, 2013b.
- Campbell, J. E. and Fourqurean, J. W.: Ocean acidification outweighs nutrient effects in
 structuring seagrass epiphyte communities, J. Ecol., 102, 730–737, doi:10.1111/13652745.12233, 2014.
- Campbell, S., Miller, C., Steven, A. and Stephens, A.: Photosynthetic responses of two temperate
 seagrasses across a water quality gradient using chlorophyll fluorescence, J. Exp. Mar.
 Biol. Ecol., 291(1), 57–78, doi:10.1016/S0022-0981(03)00090-X, 2003.
- Cebrián, J., Enríquez, S., Fortes, M. D., Agawin, N., Vermaat, J. E. and Duarte, C. M.: Epiphyte
 accrual on *Posidonia oceanica* (L.) Delile leaves: implications for light absorption, Bot.
 Mar., 42(2), 123–128, doi:10.1515/BOT.1999.015, 1999.
- Cherrett, J. M.: A simple penetrometer for measuring leaf toughness in insect feeding studies, J.
 Econ. Entomol., 66, 1736–1738, 1968.
- Ciais, P., Sabine, C., Bala, G., Bopp, L., Brovkin, V., Canadell, J., Chhabra, A., DeFries, R.,
 Galloway, J., Heimann, M., Jones, C., Le Quéré, C., Myneni, R. B., Piao, S. and
 Thornton, P.: Carbon and other biogeochemical cycles, Cambridge University Press,
 Cambridge, United Kingdom and New York, NY, USA., 2013.
- Colombo, P. M., Rascio, N. and Cinelli, F.: *Posidonia oceanica* (L.) Delile: a structural study of
 the photosynthetic apparatus, Mar. Ecol., 4(2), 133–145, doi:10.1111/j.14390485.1983.tb00292.x, 1983.
- Cox, T. E. and Smith, C. M.: Photosynthetic rapid light curves for *Padina sanctae-crucis* vary
 with irradiance, aerial exposure, and tides in Hawaii's micro-intertidal zones, Mar. Biol.,
 162(5), 1061–1076, doi:10.1007/s00227-015-2649-1, 2015.

Cox, T. E., Schenone, S., Delille, J., Díaz-Castañeda, V., Alliouane, S., Gattuso, J. P. and Gazeau, F.: Effects of ocean acidification on *Posidonia oceanica* epiphytic community and shoot productivity, J. Ecol., 103(6)1594-1609, doi:10.1111/1365-2745.12477, 2015.

753 Dickson, A. G., Sabine, C. L. and Christian, J. R.: Guide to best practices for ocean CO₂ 754 measurements., PICES Special Publication 3, British Columbia, Canada., 2007. 755 Duarte, C. M.: Temporal biomass variability and production/biomass relationships of seagrass 756 communities, Mar. Ecol. Prog. Ser., 51, 269-276, doi:10.3354/meps051269, 1989. 757 Duarte, C. M. and Chiscano, C. L.: Seagrass biomass and production: a reassessment, Aquat. 758 Bot., 65, 159–174, 1999. 759 Duarte, C. M., Marba, N., Gacia, E., Fourqurean, J. W., Beggins, J., Barron, C. and Apostolaki, 760 E. T.: Seagrass community metabolism: Assessing the carbon sink capacity of seagrass 761 meadows., Glob. Biogeochem. Cycles, 24(4), 1-9, doi:10.1029/2010GB003793, 2010. 762 Figueroa, F. L., Jiménez, C., Viñegla, B., Pérez-Rodríguez, E., Aguilera, J., Flores-Moya, A., 763 Altamirano, M., Lebert, M. and Häder, D. P.: Effects of solar UV radiation on 764 photosynthesis of the marine angiosperm Posidonia oceanica from southern Spain, Mar. 765 Ecol. Prog. Ser., 230, 59-70, 2002. 766 Gattuso, J.-P., Kirkwood, W., Barry, J. P., Cox, T. E., Gazeau, F., Hansson, L., Hendriks, I., 767 Kline, D. I., Mahacek, P., Martin, S., McElhany, P., Peltzer, E. T., Reeve, J., Roberts, D., 768 Saderne, V., Tait, K., Widdicombe, S. and Brewer, P. G.: Free-ocean CO₂ enrichment 769 (FOCE) systems: present status and future developments, Biogeosciences, 11, 4057–4075, 770 2014. 771 Gattuso, J. P., Epitalon, J. M. and Lavigne, H.: Seacarb: seawater carbonate chemistry with R., 772 [online] Available from: ran.r-project.org/package=seacarb, 2015. 773 Genty, B., Briantais, J.-M. and Baker, N. R.: The relationship between the quantum yield of 774 photosynthetic electron transport and photochemical quenching of chlorophyll 775 fluorescence, Biochem. Biophys. Acta, 990, 87-92, 1989. 776 Gobert, S., Laumont, N. and Bouquegneau, J.-M.: Posidonia oceanica meadow: a low nutrient 777 high chlorophyll (LNHC) system?, BMC Ecol., 2(1), 9, 2002. 778 Gobert, S., Cambridge, M. L., Velimirov, B., Pergent, G., Lepoint, G., Bouquegneau, J.-M., 779 Duaby, P., Pergent-Martini, C. and Walker, D. I.: Biology of Posidonia, in Seagrasses: 780 biology, ecology and conservation, edited by A. W. D. Larkum, R. J. Orth, and C. M. 781 Duarte, pp. 387–408, Springer, Dordrecht, The Netherlands., 2006. 782 Hall-Spencer, J. M., Rodolfo-Metalpa, R., Martin, S., Ransome, E., Fine, M., Turner, S. M., 783 Rowley, S. J., Tedesco, D. and Buia, M. C.: Volcanic carbon dioxide vents show 784 ecosystem effects of ocean acidification., Nature, 454, 96–99, 2008. 785 Harder, D. L., Hurd, C. L. and Speck, T.: Comparison of mechanical properties of four large, 786 wave-exposed seaweeds, Am. J. Bot., 93(10), 1426-1432, doi:10.3732/ajb.93.10.1426, 787 2006.

- Hemminga, M. A. and Duarte, C. M.: Seagrass ecology, University of Cambridge, Cambridge,
 United Kingdom., 2000.
- Hendriks, I. E., Olsen, Y. S., Ramajo, L., Basso, L., Steckbauer, A., Moore, T. S., Howard, J. and
 Duarte, C. M.: Photosynthetic activity buffers ocean acidification in seagrass meadows,
 Biogeosciences, 11, 333-346, doi:10.5194/bg-11-333-2014, 2014a.
- Hendriks, I. E., Duarte, C. M., Olsen, Y. S., Steckbauer, A., Ramajo, L., Moore, T. S., Trotter, J.
 A. and McCulloch, M.: Biological mechanisms supporting adaptation to ocean
 acidification in coastal ecosystems, Estuar. Coast. Shelf Sci., 152, 1–8,
 doi:10.1016/j.ecss.2014.07.019, 2014b.
- Henley, W. J.: Measurement and interpretation of photosynthetic light-response curves in algae in
 the context of photoinhibition and diel changes, J. Phycol., 29(6), 729–739,
 doi:10.1111/j.0022-3646.1993.00729.x, 1993.
- Invers, O., Romero, J., Perez, M. and Pérez, M.: Effects of pH on seagrass photosynthesis: a
 laboratory and field assessment, Aquat. Bot., 59(3-4), 185–194, doi:10.1016/S03043770(97)00072-7, 1997.
- Invers, O., Zimmerman, R., Alberte, R. S., Perez, M. and Romero, J.: Inorganic carbon sources
 for seagrass photosynthesis: an experimental evaluation of bicarbonate use in species
 inhabiting temperate waters, J. Exp. Mar. Biol. Ecol., 265, 203–217, 2001.
- Invers, O., Tomas, F., Perez, M., Romero, J., Tomàs, F., Pérez, M. and Romero, J.: Potential
 effect of increased global CO₂ availability on the depth distribution of the seagrass
 Posidonia oceanica (L.) Delile: a tentative assessment using a carbon balance model,
 Bull. Mar. Sci., 71(3), 1191–1198, 2002.
- Invers, O., Kraemer, G. P., Pérez, M. and Romero, J.: Effects of nitrogen addition on nitrogen
 metabolism and carbon reserves in the temperate seagrass *Posidonia oceanica*, J. Exp.
 Mar. Biol. Ecol., 303(1), 97–114, doi:10.1016/j.jembe.2003.11.005, 2004.
- Jassby, A. D. and Platt, T.: Mathematical formulation of the relationship between photosynthesis
 and light for phytoplankton, Limnol. Oceanogr., 21, 540–547, 1976.
- Jeffrey, S. and Humphrey, G.: New spectrophotometric equations for the determination of
 chlorophylls a, b, c1 and c2 in higher plants, algae and natural phytoplankton. 167, 191–
 194., Biochem. Physiol. Pflanz., 167, 191–194, 1975.
- Jiang, Z. J., Huang, X.-P. and Zhang, J.-P.: Effects of CO₂ enrichment on photosynthesis, growth,
 and biochemical composition of seagrass *Thalassia hemprichii* (Ehrenb.) Aschers, J.
 Integr. Plant Biol., 52, 904–913, 2010.
- Jordà, G., Marbà, N. and Duarte, C. M.: Mediterranean seagrass vulnerable to regional climate
 warming, Nat. Clim. Change, 2(11), 821–824, doi:10.1038/nclimate1533, 2012.

- Kerrison, P., Hall-Spencer, J. M., Suggett, D. J., Hepburn, L. J. and Steinke, M.: Assessment of pH variability at a coastal CO₂ vent for ocean acidification studies, Estuar. Coast. Shelf
 Sci., 94, 129–137, 2011.
- Koch, M., Bowes, G., Ross, C. and Zhang, X. H.: Climate change and ocean acidification effects
 on seagrasses and marine macroalgae, Glob. Change Biol., 19(1), 103–132,
 doi:10.1111/j.1365-2486.2012.02791.x, 2013.
- Libes, M.: Productivity-irradiance relationship of *Posidonia oceanica* and its epiphytes, Aquat.
 Bot., 26, 285–306, 1986.
- Littler, M. M. and Littler, D. S.: The evolution of thallus form and survival strategies in benthic
 marine macroalgae: field and laboratory tests of a functional form model, Am. Nat., 116,
 25–44, 1980.
- Liu, X., Patasavas, M. C. and Byrne, R. H.: Purification and characterization of meta-Cresol
 Purple for spectrophotometric seawater pH measurements, Environ. Sci. Technol., 45,
 4862–4868, 2011.
- Marbà, N. and Duarte, C. M.: Rhizome elongation and seagrass clonal growth, Mar. Ecol. Prog.
 Ser., 174, 269–280, 1998.
- Marbà, N., Hemminga, M. A., Mateo, M. A., Duarte, C. M., Mass, Y. E. M., Terrados, J. and
 Gacia, E.: Carbon and nitrogen translocation between seagrass ramets, Mar. Ecol. Prog.
 Ser., 226, 287–300, 2002.
- Martin, S., Rodolfo-Metalpa, R., Ransome, E., Rowley, S., Buia, M.-C. C., Gattuso, J.-P. and
 Hall-Spencer, J.: Effects of naturally acidified seawater on seagrass calcareous epibionts,
 Biol. Lett., 4(6), 689–692, doi:10.1098/rsbl.2008.0412, 2008.
- Martínez-Crego, B., Olivé, I. and Santos, R.: CO₂ and nutrient-driven changes across multiple
 levels of organization in *Zostera noltii* ecosystems, Biogeosciences, 11, 7237–7249, 2014.
- Newcomb, L. A., Milazzo, M., Hall-Spencer, J. M. and Carrington, E.: Ocean acidification bends
 the mermaid's wineglass, Biol. Lett., 11(9), 20141075, doi:10.1098/rsbl.2014.1075, 2015.
- 849 Ott, J. A.: Growth and production in *Posidonia oceanica* (L.) Delile, Mar. Ecol., 1(1), 47–64,
 850 doi:10.1111/j.1439-0485.1980.tb00221.x, 1980.
- Ow, Y. X., Collier, C. J. and Uthicke, S.: Response of three tropical seagrass species to CO₂
 enrichment, Mar. Biol., 162(5), 1005–1017, doi:10.1007/s00227-015-2644-6, 2015.
- Padilla, D. K.: Structural resistance of algae to herbivores: a biomechanical approach, Mar. Biol.,
 90(1), 103–109, doi:10.1007/BF00428220, 1985.
- Palacios, S. L. and Zimmerman, R.: Response of eelgrass *Zostera marina* to CO₂ enrichment:
 possible impacts of climate change and potential for remediation of coastal habitats, Mar.
 Ecol. Prog. Ser., 344, 1–13, 2007.

- Pasqualini, V., Pergent-Martini, C., Clabaut, P. and Pergent, G.: Mapping of *Posidonia oceanica*using aerial photographs and side scan sonar: application off the island of Corsica
 (France), Estuar. Coast. Shelf Sci., 47, 359–367, 1998.
- Peirano, A., Niccolai, I., Mauro, R. and Bianchi, C. N.: Seasonal grazing and food preference of
 herbivores in a *Posidonia oceanica* meadow, Sci. Mar., 65(4), 367–374, 2001.
- 863 Pergent-Martini, C., Leoni, V., Pasqualini, V., Ardizzone, G. D., Balestri, E., Bedini, R., 864 Belluscio, A., Belsher, T., Borg, J., Boudouresque, C. F., Boumaza, S., Bouquegneau, J. 865 M., Buia, M. C., Calvo, S., Cebrian, J., Charbonnel, E., Cinelli, F., Cossu, A., Maida, D. 866 I., Dural, B., Francour, P., Gobert, S., Lepoint, G., Meinesz, A., Molenaar, H., Mansour, 867 H., Panayotidis, M. P., Peirano, A., Pergent, G., Piazzi, L., Pirrotta, M., Relini, G., 868 Romero, J., Sanchez-Lizaso, J. L., Semroud, R., Shembri, P., Shili, A., Tomasello, A. and 869 Velimirov, B.: Descriptors of *Posidonia oceanica meadows*: Use and application, Ecol. 870 Indic., 5, 213–230, 2005.
- Platt, T., Gallegos, C. and Harrison, W.: Photoinhibition of photosynthesis in natural assemblages
 of marine phytoplankton., J. Mar. Res., 38, 687–701, 1980.
- Poore, A. G. B., Graba-Landry, A., Favret, M., Sheppard Brennand, H., Byrne, M. and
 Dworjanyn, S. A.: Direct and indirect effects of ocean acidification and warming on a
 marine plant-herbivore interaction, Oecologia, 173(3), 1113–1124, doi:10.1007/s00442013-2683-y, 2013.
- Russell, B. D., Connell, S. D., Uthicke, S., Muehllehner, N., Fabricius, K. E. and Hall-Spencer, J.
 M.: Future seagrass beds: Can increased productivity lead to increased carbon storage?,
 Mar. Pollut. Bull., 73(2), 463–469, doi:10.1016/j.marpolbul.2013.01.031, 2013.
- Sand-Jensen, K., Revsbech, N. P. and Jorgensen, B. B.: Microprofiles of oxygen in epiphyte
 communities on submerged macrophytes, Mar. Biol., 89, 55–62, 1985.
- de los Santos, C. B., Brun, F. G., Vergara, J. J. and Perez-Llorens, J. L.: New aspect in seagrass
 acclimation: leaf mechanical properties vary spatially and seasonally in the temperate
 species *Cymodocea nodosa* Ucria (Ascherson), Mar. Ecol. Prog. Ser., 1–13, 2013.
- Shaw, E. C., McNeil, B. I., Tilbrook, B., Matear, R. and Bates, M. L.: Anthropogenic changes to
 seawater buffer capacity combined with natural reef metabolism induce extreme future
 coral reef CO₂ conditions, Glob. Change Biol., 19(5), 1632–1641, doi:10.1111/gcb.12154,
 2013.
- Short, F. T. and Duarte, C. M.: Methods for the measurement of seagrass growth and production,
 in Global seagrass research method, edited by F. T. Short and R. G. Coles, pp. 155–180,
 Elsevier, Amsterdam, The Netherands., 2001.

Vassallo, P., Paoli, C., Rovere, A., Montefalcone, M., Morri, C. and Bianchi, C. N.: The value of the seagrass *Posidonia oceanica*: A natural capital assessment, Mar. Pollut. Bull., 75, 157–167, 2013.

- Vizzini, S., Tomasello, A., Di Maida, G., Pirrotta, M., Mazzola, A. and Calvo, S.: Effect of
 explosive shallow hydrothermal vents on d13C and growth performance in the seagrass *Posidonia oceanica*, J. Ecol., 98(6), 1284–1291, doi:10.1111/j.1365-2745.2010.01730.x,
 2010.
- Vizzini, S., Di Leonardo, R., Costa, V., Tramati, C. D., Luzzu, F. and Mazzola, A.: Trace element
 bias in the use of CO₂ vents as analogues for low pH environments: implications for
 contamination levels in acidified oceans, Estuar. Coast. Shelf Sci., 134, 19–30,
 doi:10.1016/j.ecss.2013.09.015, 2013.
- Waycott, M., Duarte, C. M., Carruthers, T. J. B., Orth, R. J., Dennison, W. C., Olyarnik, S.,
 Calladine, A., Fourqurean, J. W., Heck, K. L., Hughes, A. R., Kendrick, G. A.,
 Kenworthy, W. J., Short, F. T. and Williams, S. L.: Accelerating loss of seagrasses across
 the globe threatens coastal ecosystems., Proc. Natl. Acad. Sci. U. S. A., 106, 12377–
 12381, 2009.
- Zimmerman, R. C. A., Kohrs, D. G. A., Steller, D. L. B. and Alberte, R. S. A.: Impacts of CO₂
 enrichment on productivity and light requirements of eelgrass, Plant Physiol., 115, 599–
 607, 1997.
- 911 Zimmerman, R. C., Hill, V. J. and Gallegos, C. L.: Predicting effects of ocean warming,
- 912 acidification, and water quality on Chesapeake region eelgrass: Predicting eelgrass
- 913 response to climate change, Limnol. Oceanogr., 60(5), 1781–1804,
- 914 doi:10.1002/lno.10139, 2015.

915 Figure captions

Figure 1. Schematic of the system and study design (A) see text for details (B): the pH (total
scale) inside the enclosures and in ambient during the week-long transition to the targeted offset
(-0.25 units).

919

Figure 2. Macrophyte abundance throughout the experiment; A: enclosures and reference plot had initially similar *P. oceanica* shoot density m^{-2} (mean ± SE). B: mean shoot number with time within three permanently located quadrats (0.25 m²) per reference plot (top), control (middle) and experimental (bottom) enclosures. C, D, E: coverage (%) of benthic macrophytes and unoccupied sediment or rocks (bare space) before and during the acidification period (x-axis after the dashed vertical line).

926

Figure 3. Leaf biometrics (mean ± SE) before and during the acidification period for the
reference and enclosure plants. Measures through time: average shoot height (A), leaf length (B),
minimum (C) and maximum leaf length (D), number of leaves per shoot (E), leaf area (F), leaf
thickness (G) and leaf toughness (H) are shown. The dashed vertical line indicates when the pH
was lowered in the experimental enclosure. Additional leaves were collected in June from the
meadow and are referred to as ambient leaves.

933

Figure 4. Photosynthetic rapid light curves (RLCs, A-D), dark-adapted quantum yield (E), and the derived RLC parameters (F-H) measured on 2^{nd} rank leaves in enclosures and reference plot before (May) and during (July, September, and October) the acidification period. Symbols represent the mean (±SE) relative electron transport rate (*r*ETR) at each mean photosynthetic

938	active radiation (PAR) value. Curved lines represent the Jassby and Platt (1976) regression based
939	on mean values. The dashed outline encloses the acidification period.

940

941	Figure 5.	Photosynthesis	versus irradiance	(PE) curves	produced from	laboratory incubations	of
-----	-----------	----------------	-------------------	-------------	---------------	------------------------	----

942 *P. oceanica* leaf segments collected from the enclosures after two (September, A) and four

943 (November, B) months of acidification. The derived parameters from the curves are shown in944 panels C-G.

945

946 Fi	gure 6.	Growth as <i>P</i> .	<i>oceanica</i> leaf	production ((A) and	leaf	plastochrone ir	nterval (B) during	the
--------	---------	----------------------	----------------------	--------------	---------	------	-----------------	------------	----------	-----

947 acidification period. After 4 months of acidification, biomass (above-ground, C; below-ground,

948 D) was determined from replicate cores collected from enclosures and the reference plot. A fourth

949 nearby ambient area was additionally sampled to better account for spatial variation.

Table 1. A comparison of the carbonate chemistry and diel changes within the ambient and enclosures: the mean (\pm standard deviation, SD) pH (on the total scale), the maintained pH offset between experimental and control enclosures as a difference (Diff), the partial pressure of carbon dioxide (pCO_2), and the median (Med \pm median absolute deviation, MAD) diel pH and oxygen (O_2) change for each month and the period before and during the acidification.

		pH _T										pCO ₂	(µatm)			Δ Diel pH _T							\triangle Diel O ₂					
		Ν	Am	bient	Con	trol	Experin	nental	Di	ff	Amb	ient	Control		Experimental		N Amł		bient	Control		Experimental		Ambient		Control		Experimental	
Months	in Period	Samples	Mean	SD	Mean	SD	Mean	SD	Mean	SD	Mean	SD	Mean	SD	Mean	SD	Days	Med	MAD	Med	MAD	Med	MAD	Med	MAD	Med	MAD	Med	MAD
Before																													
	May	11840	8.10	0.03	8.12	0.06	8.01	0.05	-0.10	0.03	374	30	358	55	477	74	17	0.08	0.02	0.10	0.06	0.12	0.05	52.0	11.0	68.0	7.0	82.0	14.0
Acidificatio	June on	8119	8.11	0.04	8.04	0.05	8.10	0.06	0.06	0.05	369	38	443	63	378	65	11	0.10	0.03	0.15	0.04	0.16	0.02	72.0	9.0	91.0	8.0	101.0	9.0
	June	6226	8.05	0.03	8.02	0.04	7.79	0.13	-0.23	0.13	430	42	470	57	868	318	9	0.12	0.03	0.12	0.04	0.27	0.08	72.0	7.0	85.0	10.0	92.0	11.0
	July	21007	8.03	0.03	8.03	0.06	7.79	0.12	-0.24	0.11	454	46	453	81	870	254	30	0.09	0.02	0.17	0.05	0.27	0.06	60.5	14.0	95.5	18.0	100.5	18.0
	August	22682	8.00	0.03	8.04	0.07	7.81	0.12	-0.23	0.09	489	42	445	85	834	253	31	0.09	0.02	0.18	0.05	0.29	0.06	55.0	8.0	77.0	12.0	86.0	12.0
	September	21854	7.98	0.07	7.97	0.06	7.70	0.11	-0.27	0.10	521	96	536	87	1098	288	30	0.07	0.01	0.11	0.06	0.28	0.10	37.5	5.5	62.5	15.5	54.0	11.5
	October	22420	8.01	0.04	8.00	0.04	7.70	0.13	-0.29	0.14	480	52	497	64	1086	390	31	0.06	0.02	0.09	0.04	0.29	0.08	27.0	3.0	34.0	5.0	44.0	5.0
	November	5377	8.02	0.03	8.02	0.02	7.80	0.15	-0.22	0.15	469	48	467	22	836	305	10	0.04	0.01	0.06	0.03	0.09	0.05	21.0	5.5	34.0	22.5	45.5	29.5
Before		24334	8.10	0.04	8.05	0.07	8.06	0.07	0.01	0.09	380	39	434	85	426	87	34	0.09	0.02	0.14	0.06	0.16	0.07	63.5	13.0	80.5	13.5	88.0	13.0
Acidificati	on	95711	8.01	0.05	8.01	0.06	7.75	0.13	-0.26	0.11	483	67	482	86	971	323	132	0.08	0.02	0.14	0.06	0.28	0.14	44.0	14.5	68.5	23.5	74.0	23.0







959960 Figure 3.



961962 Figure 4.





Treatment and Month

963964 Figure 5.



