- 1 Dynamics of environmental conditions during a decline of a *Cymodocea nodosa* meadow
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Abstract. The dynamics of the physicochemical and biological parameters were followed 16 during the decline of a *Cymodocea nodosa* meadow in the northern Adriatic Sea from July 17 2017 to October 2018. During the regular growth of C. nodosa from July 2017 to March 18 2018, C. nodosa successfully adapted to the changes of environmental conditions and 19 prevented  $H_2S$  accumulation by its re-oxidation, supplying the sediment with  $O_2$  from the 20 water column and/or leaf photosynthesis. The C. nodosa decline was most likely triggered in 21 April 2018 when light availability to the plant was drastically reduced due to increased 22 seawater turbidity that resulted from increased terrigenous input combined with resuspension 23 of sediment and elevated autotrophic biomass. Light reduction impaired photosynthesis of C. 24 nodosa and the oxidation capability of below-ground tissue. Simultaneously, a depletion of 25 oxygen due to intense oxidation of H<sub>2</sub>S occurred in the sediment, thus creating anoxic 26 conditions in most of the rooted areas. These linked negative effects on the plant performance 27 caused an accumulation of H<sub>2</sub>S in the sediments of the *C. nodosa* meadow. During the decay 28 of above- and below-ground tissues, culminating in August 2018, high concentrations of H<sub>2</sub>S 29 30 were reached and accumulated in the sediment as well as in bottom waters. The influx of oxygenated waters in September 2018 led to the re-establishment of H<sub>2</sub>S oxidation in the 31 32 sediment and remaining of the below-ground tissue. Our results indicate that if disturbance of environmental conditions, particularly those compromising the light availability, takes place 33 during the recruitment phase of plant growth when metabolic needs are at maximum and 34 stored reserves minimal, a sudden and drastic decline of the seagrass meadow occurs. 35

## 37 1 Introduction

Seagrasses are important ecosystem engineers constructing valuable coastal habitats which 38 play a key role in the preservation of marine biodiversity and carbon sequestration (Duarte et 39 al., 2013; Samper-Villareal et al., 2016). Seagrasses extend their active metabolic surfaces 40 (i.e., leaves, rhizomes and roots) into the water column and in the sediment, where root 41 activity might modify the chemical conditions (Marbà and Duarte, 2001). Their canopies and 42 dense meadows are responsible for trapping substantial amounts of sediment particles and 43 organic matter, enhancing water transparency and sediment stability with the dense network 44 formed by the rhizome (Gacia and Duarte, 2001; Hendriks et al., 2008; Widdows et al., 2008). 45 Seagrass rhizospheres store organic matter (Pedersen et al., 1997), promote sulfate reduction 46 (Holmer and Nielsen, 1997), release oxygen (Pedersen et al., 1998) and alter sediment redox 47 48 potential.

Seagrasses require some of the highest light levels of any plant worldwide to provide 49 oxygen to roots and rhizomes and support a large amount of non-photosynthetic tissue (Orth 50 et al., 2006). This make seagrasses sensitive to environmental changes, especially those that 51 52 deteriorate light availability, such as sediment loading, eutrophication or epiphyte cover on seagrass leaves (Terrados et al., 1998; Halun et al., 2002; Brodersen et al., 2015; Costa et al., 53 54 2015). Seagrasses have adapted to a highly variable light environment providing tolerance to short-term periods of low light conditions by balancing carbon supply and respiratory 55 56 requirements. In a healthy growing population this balance is achieved by increasing the 57 photosynthetic activity, re-allocation of carbohydrate reserves from rhizomes and slowing down growth rates (Collier et al., 2009). Beside metabolic and physiological changes, stress 58 responses under poor light conditions include shedding of leaves and shoots and production of 59 new, altered tissue. At sub-lethal light levels, these changes may be permanent. Below these 60 species-specific minimum light requirements seagrass populations are dying off (Collier et al., 61 2012). Membrane lipids, particularly polyunsaturated fatty acids (PUFA), as the most 62 responsive constituents have a major role in the adaptation processes of primary producers to 63 fluctuating environmental factors, such as temperature, irradiance or salinity (Viso et al., 64 1993; Lee et al., 2007; Schmid et al., 2014; Sousa et al., 2017; Beca-Carretero et al., 2018; 65 Beca-Carretero et al., 2019). The changes in the unsaturation degree (UND) of membrane 66 67 fatty acids affect the maintenance of membrane functions and its resistance to cold stress or

68 poor light conditions. UND depends mostly on the variation of α-linolenic (C18:3n-3, ALA)

and linoleic (C18:2n-6, LA), the major unsaturated fatty acids in leaves, implicated in the

volution of oxygen during photosynthesis. LA and ALA are derived from oleic acid by

71 desaturation in the chloroplast and this conversion considerably declines in the dark, being

completely inhibited by anaerobiosis (Harris and James, 1965).

Sediments inhabited by seagrasses are usually anoxic, highly reduced and rich in sulfide 73 (H<sub>2</sub>S), a strong phytotoxin (Koch and Erskine, 2001) which has been implicated in several 74 die-off events of seagrasses (Carlson et al., 1994; Borum et al., 2005; Krause-Jensen et al., 75 76 2011). H<sub>2</sub>S is produced by sulfate-reducing bacteria that use sulfate as a terminal electron acceptor for the mineralization of organic matter (Jørgensen, 1977; Capone and Kiene, 1988, 77 Canfield et al., 1993). High H<sub>2</sub>S concentrations may occur as a consequence of enhanced 78 mineralization due to increased temperature, organic loading or oxygen depletion (Moeslund 79 et al., 1994; Pérez et al., 2007; Mascaró et al., 2009). Under these conditions, sulfides may 80 intrude into plant. Re-oxidation of  $H_2S$  in the rhizosphere by incorporation of  $S^0$  in the below-81 82 ground tissue has been recognized as a major survival strategy of seagrasses in sulfidic sediments (Pedersen et al., 2004; Holmer et al., 2005; Hasler-Sheetal and Holmer, 2015). 83 Generally, the synergistic effect of oxygen depletion and other stresses, such as sulfide 84 toxicity may shorten the survival of benthic communities and possibly accelerate mortality 85 events (Vaquer-Sunyer and Duarte, 2010). 86

The seagrass Cymodocea nodosa (Ucria) Ascherson is widely distributed and common 87 species throughout the Mediterranean (Terrados and Ros 1992; Pedersen et al., 1997; 88 Cancemi et al., 2002; Agostini et al., 2003). For the northern Adriatic, however, only sparse 89 data are available on the standing crop, seasonal dynamics or natural/anthropogenic pressures 90 supporting the ecological or conservation status of C. nodosa meadows (Zavodnik et al., 91 1998; Orlando-Bonaca et al., 2015; 2016). Although C. nodosa show large phenotypic 92 93 plasticity adapting to diverse natural and anthropogenic stressors by physiological and morphological adaptations, a severe decline has been reported during the last decades in 94 95 coastal areas (Orth et al., 2006; Short et al., 2011; Tuya et al., 2002; 2014), including the northern Adriatic (Orlando-Bonaca et al., 2015; 2019). One of these declines was documented 96 in our study performed from July 2017 to October 2018 in Saline Bay (northern Adriatic Sea). 97 A series of monthly physicochemical and biological measurements were conducted in C. 98 nodosa tissues, sediment underlying the C. nodosa meadow, non-vegetated sediments and 99 4

- 100 surrounding water to i) determine the link between ambient seawater and sediment
- 101 environmental factors influencing the growth of *C. nodosa*, ii) document the response of *C*.
- 102 *nodosa* to the changes in environmental conditions that led to the meadow decline and iii)
- 103 evaluate the conditions leading to the decline of *C. nodosa*.

# 104 **2 Materials and methods**

105 2.1 Study site

Saline Bay is located 4 km northwest of Rovinj (Croatia) at the coast of the northern Adriatic 106 Sea (45°7′5″N; 13°37′20″E, Fig. S1). The bay represents the terminal shallow part of an 800 107 m long inlet, open towards the northwest. The southeastern coast of Saline Bay is 108 characterized by relatively pristine conditions, while the northwestern littoral part has been 109 110 completely modified by the excavation of coastal mud and the addition of large amounts of gravel to create an artificial beach. Large amounts of silty red soil (terra rossa) can be found 111 112 in the south eastern inner part of the bay in a large muddy flatland which is slowly being eroded by the sea and rain weathering. The main input of freshwater to the bay represents land 113 114 drainage canals since the year 2017. Even though Saline Bay is protected from the prevailing winds (from the NE and SE) circulations from the northwestern quadrant can occasionally 115 trigger bigger waves resuspending the surface sediments and giving the waters a muddy 116 appearance. At the beginning of this study, the seafloor was covered with large C. nodosa 117 meadows spreading from the southwestern coastal area (1.5 m depth) toward the central part 118 of the bay (4 m depth), while at the end of the study only a few small patches persisted in tiny 119 stripes along the shoreline. 120

121

122 2.2 Sampling

The sampling was performed for 15 months from July 2017 to October 2018. Seawater for 123 analyses of nutrients, chlorophyll a (Chl *a*), particulate matter concentration and prokaryotic 124 abundance was sampled using plastic containers (10 L). C. nodosa (3 – 4 m of depth) was 125 collected together with rhizomes, roots and epiphytic macroalgae by divers using the quadrat 126 127 sampling method. Three quadrats (20 x 20 cm) were randomly scattered in positions of maximum seagrass coverage (e.g. 100 %). Sediment samples were collected inside vegetated 128 and non-vegetated sediment by divers using plastic core samplers (15 cm, 15.9 cm<sup>2</sup>). For 129 granulometric composition, organic matter, prokaryotic abundance, total lipids and fatty acid 130 analyses, the cores were cut into 1 cm sections to a depth of 8 cm and lyophilized, except of 131

- sections for prokaryotic abundance analysis, that were weighted (approx. 2 g) and fixed with
- formaldehyde (final conc. 4% v/v) immediately after slicing the sediment core.
- 134
- 135 2.3 Temperature (T) and salinity (S) measurements
- 136 T was measured continuously (in 30 min. intervals) using HOBO pendant temp/light Data
- 137 Loggers (Onset, USA) which were replaced at each sampling. S was measured on sampling
- dates by a pIONneer 65 probe (Radiometer analytical, Copenhagen).
- 139

140 2.4 Inorganic nutrients, Chl *a* and particulate matter (PM) analysis

- 141 Seawater for all analysis was filtered through combusted Whatman GF/F filters. Nitrate
- 142 (NO<sub>3</sub>), nitrite (NO<sub>2</sub>), ammonia (NH<sub>4</sub>), phosphate (PO<sub>4</sub>) and silicate (SiO<sub>4</sub>) were analyzed
- spectrophotometrically according to Strickland and Parsons (1972). Chl *a* was determined on
- 144 filters by the fluorometric procedure after extraction in 90 % acetone (Holm-Hansen et al.,
- 145 1965). PM was determined gravimetrically after filtering up to 5L seawater on pre-weighed,
- 146 filters which were dried (at  $60^{\circ}$ C) and reweighed.
- 147
- 148 2.5 Determining prokaryotic abundance
- 149 For determining the prokaryotic abundance in seawater, 2 ml of formaldehyde (final conc. 4%
- 150 v/v) fixed samples were stained with 4,6-diamidino-2-phenylindol (DAPI, 1 µg mL<sup>-1</sup> final
- 151 conc.) for 10 min (Porter and Feig, 1980). In sediment samples, prokaryotes were detached
- 152 from the sediment particles by addition of Tween 80 (0.05 mL) and ultrasonicated for 15 min
- 153 (Epstein and Rossel, 1995). After sonication, 1 mL of the supernatant was stained with DAPI
- 154 (final conc. 5 µg/mL). DAPI stained samples were filtered onto black polycarbonate filters
- 155 (Whatman, Nuclepore, 0.22 μm) and counted under an epifluorescence microscope (Zeiss
- 156 Axio Imager Z1).
- 157
- 158 2.6 Biometry of *C. nodosa* and epiphytic macroalgae
- 159 The material from each quadrat was washed under running seawater to remove sediment.
- 160 From each quadrat algae, leaves and rhizomes with roots were separated. The length of the
- 161 longest leaf on each shoot was measured and the shoots were counted. Species of macroalgae
- 162 were determined, and their coverage was estimated according to the Braun-Blanquet scale.

163 Separated samples were washed with filtered and autoclaved seawater, weighed, dried at 60 164 °C for 48 h and re-weighed. The dry mass was calculated per area (g m<sup>-2</sup>).

165

166 2.7 Granulometric composition of the sediment and its organic matter content For granulometric analysis of the sediment, each sample was wet sieved through a set of 167 seven standard ASTM sieves (4-, 2-, 1-, 0.5-, 0.25-, 0.125-, 0.063-mm mesh size). The 168 169 fraction that passed through the 0.063-mm sieve was collected and analyzed following the 170 standard sedigraph procedure (Micromeritics, 2002). The material that was retained on the sieves was dried and weighted. The data obtained by both techniques were merged to obtain a 171 continuous grain size range and analyzed with the statistic package Gradistat v 6.0. Sediments 172 were classified according to Folk (1954). The sediment permeability was calculated based on 173 median grain size (dg) following the empirical relation by Gangi (1985). The organic matter 174 content was determined as ignition loss after heating dried sediment sections at 450°C for 4 h 175 176 in a muffle furnace.

177

178 2.8 Oxygen (O<sub>2</sub>), hydrogen sulfide (H<sub>2</sub>S) and redox potential (Eh) profiling

The microprofiles of O<sub>2</sub>, H<sub>2</sub>S and Eh were measured on intact cores immediately after
sampling using a motorized micromanipulator (MMS9083) equipped with microsensors OX-

181 100 and  $H_2S$ -200, redox microelectrode RD-200 coupled with reference electrode REF-RM

182 (Unisense A/S, Denmark). Prior to the measurements, the OX-100 microsensor was calibrated

using a two-point oxic – anoxic calibration;  $H_2S$ -200 was calibrated in fresh Na<sub>2</sub>S solutions

using eight-point calibration  $(1\mu M - 300 \mu M \text{ in a de-oxygenated calibration buffer})$ 

185 (NaAc/HAc, pH <4); RD-200 with REF-RM was calibrated using two point calibration by

simultaneous immersion of electrodes in quinhydrone redox buffers prepared in pH 4 and pH

187 7 buffers, all according to the manufacturer's recommendation. During measurements,

sediment cores were placed in a pool filled with seawater from the sampling site to maintain

in situ temperature. From July to October 2017  $H_2S$  was measured spectrophotometrically in

190 pore waters (Cline, 1969) squeezed out by centrifugation from each section (5 mm) of the

191 sediment cores.

192

193 2.9 Total lipids, fatty acid composition and elemental sulfur  $(S^0)$ 

Lyophilized samples of seagrass tissues, macroalgae, sediment or particulate matter were 194 weighed and extracted into a solvent mixture of dichloromethane/methanol (DCM: MeOH, 195 2:1) in an ultrasonic bath at 35°C with three solvent mixture changes. The extracts were 196 pooled and separated into layers by addition of 0.9% NaCl solution. Lower DCM layers 197 (containing lipids) were released over Na<sub>2</sub>SO<sub>4</sub> anhydride, collected in pre-weighed round 198 bottom flasks and evaporated to dryness using rotavapor. After evaporation, flasks were re-199 weighed, and total lipid concentrations (TL, mg  $g^{-1}$  DW) were calculated from the difference 200 in weight. For fatty acids determination, lipid extracts were saponified (1.2 M NaOH in 201 202 methanol), acidified (6 M HCl), methylated (14% BF<sub>3</sub> in methanol) and extracted into DCM. Fatty acid methyl esters (FAME) were analyzed by Agilent gas-liquid chromatography 203 (GLC) 6890 N GC System equipped with a 5973 Network Mass Selective Detector, capillary 204 column (30 m x 0.3 mm x 0.25 µm; cross-linked 5 % phenylmethylsiloxane) and ultra-high 205 206 purity helium as the carrier gas. The GLC settings were as follows: programmed column temperature rise from 145°C by 4°C/min to 215°C, then by 1°C/min to 225°C and finally by 207 208 4°C/min to 270°C at constant column pressure of 2.17 kPa. Retention times, peak areas and mass spectra were recorded on the ChemStation Software. FAME were identified by mass 209 210 spectral data and family plots of an equivalent chain length (ECL) for GC standards. Applied GC standards were: FAME mix C18-C20, PUFA1, PUFA3 standards (Supelco/Sigma-211 Aldrich, Bellefonte, PA, USA); C4–C24 FAME standard mix, cod liver oil and various 212 individual pure standards (Sigma, Neustadt, Germany). 213 The following indices of fatty acid profiles were calculated: saturated fatty acids (SAT), 214 monounsaturated fatty acids (MUFA), polyunsaturated fatty acids (PUFA) and the 215 unsaturation degree (UND). UND was employed to evaluate the degree of organic matter 216 degradation due to more susceptibility of unsaturated, particularly polyunsaturated, 217 components to degradation and calculated according to the formula 218 219 [1\*(% mono-)+2\*(% di-)+3\* (% tri-)+4\*(% tetra-)+5\*(% penta-)+6\*(% hexa-enoic)]/% SAT (Pirini et al., 2007). To evaluate the input of terrestrial organic matter relative to that of 220 marine origin in particulate matter, the terrestrial to aquatic acid ratio (TAR=C24+C26+C28 / 221 C12+C14+C16) was used (Cranwell et al., 1987; Bourbonniere and Meyers, 1996). 222 In FAME chromatograms elemental sulfur (S<sup>0</sup>), eluted as S<sub>8</sub> (m/z 256), was identified by 223 comparison of retention time and characteristic fragment ions in samples and standard 224 solutions. The concentration of  $S^0$  was estimated on the base of the calibration curve prepared 225

- for standard solution of  $S_8$  (Aldrich, Germany) in cyclohexane (2-20 mg L<sup>-1</sup>). The calibration
- 227 curve was determined under the same GLC settings as FAME. Limit of detection (LoD) and
- 228 limit of quantitation (LoQ) were calculated from the parameters of the calibration curve
- constructed on the basis of the 3 lowest concentrations in 3 replicates. LoD and LoQ (0.92 mg
- 230  $L^{-1}$  and 2.80 mg  $L^{-1}$ , respectively) were more than twice the values obtained by Rogowska et
- al. (2016) probably due to higher injector and column temperature used in this study than they
- 232 proposed as optimal for S determination.
- 233

234 2.10 Data analyses

A multivariate analysis, hierarchical clustering and K-means methods (Systat 12) was applied

to group *C. nodosa* above- and below-ground tissues according to the similarity of their fatty

- acid profiles and indices, i.e., physiological condition during the investigated period.
- 238 Sediment data were analyzed for two groups of sediment layers, the upper layer (0- 4 cm)
- where most of rhizomes and roots are located, and the lower layers (5-7 cm). Differences

between vegetated and non-vegetated sediment samples in each sediment layer were tested by

- one-way ANOVA. Correlations among parameters were tested using the Pearson's correlation
- coefficient (r). The level of statistical significance was p < 0.05. A multivariate principal
- component analysis (PCA, Primer 6) was applied to identify the most important variables

244 explaining differences between vegetated and non-vegetated sediments. Correlation matrices

245 were constructed using variables:  $H_2S$ , Eh,  $O_2$ ,  $S^0$ , PA, TL and UND. All variables were

- normalized due to their different scales. Only the principal components with eigenvalues >1
- 247 were considered.
- 248

# 249 **3 Results**

250 3.1 Water column

251 3.1.1 Environmental variables

252 During summer of 2017 daily means of sea-bottom temperature in *C. nodosa* meadow ranged

- between 26°C and 28°C. During autumn seawater temperatures decreased below 12°C until
- the end of December. The coldest period was recorded at the beginning of March lasting only
- for a few days (min. 8.62°C). From April to mid-July 2018, temperature increased with
- moderate fluctuations to the maximum of 29.26°C recorded in August 2018 (Fig. 1a).

- 257 Concentrations of inorganic nutrients and Chl *a* were generally low. The highest
- 258 concentrations (DIN: 8.27  $\mu$ M; PO<sub>4</sub>: 0.18  $\mu$ M; SiO<sub>4</sub>: 9.82  $\mu$ M; Chl *a*: 0.89  $\mu$ g L<sup>-1</sup>) associated
- with the lowest salinity (34.2) were found in September 2017 (Table S1). The abundance of
- prokaryotes (2.6-11.3 x  $10^5$  cell mL<sup>-1</sup>) varied seasonally and significantly correlated to
- seawater temperatures (r = 0.618; p < 0.05). In contrast, salinity (S: 34.2 38.5) and
- 262 concentrations of particulate matter (PM:  $3.84 14.21 \text{ mg L}^{-1}$ ) showed irregular variations
- 263 (Fig. 1b) and a significant opposite trend (r = -0.630; p < 0.05).
- The particulate lipids exhibited the highest unsaturation degree (UND) during
  summer/early autumn 2017 and small increases of UND in April and September/October
- 205 Summer/early autumn 2017 and sman mereases of 01(D in riph and September/000000
- 266 2018 (Fig. 1c). UND was significantly correlated with Chl *a* (r = 0.603; p < 0.05). In contrast,
- terrestrial to aquatic ratio (TAR) considerably increased in April and was the highest in
- August 2018 (Fig. 1c). TAR was negatively correlated to UND (r = -0.644, p < 0.05) and
- 269 positively to particulate matter (r = 0.641, p < 0.05). Although PUFA with 18 C atoms made
- the largest contribution to the total PUFA pool, C20 PUFA, mainly of phytoplankton origin,
- showed a similar trend as observed for UND (Fig. S2, Table S2).
- 272
- 273 *3.2 Cymodocea nodosa* meadow
- 274 3.2.1 Biometry

275 *C. nodosa* leaves and shoots reached the highest biomass  $(285.3 \pm 57.4 \text{ g m}^{-2})$ , length  $(102.4 \pm 26.6 \text{ mm})$  and shoot density  $(3703\pm334 \text{ shoots m}^{-2})$  in October 2017 (Fig. 2a). After the 277 appearance of the regular vegetation minimum in November 2017, biometric indices further

- decreased reflecting the decay of the meadow in summer 2018. In August 2018, only yellow
- to brownish leaves on sparse shoots were collected ( $4.5 \pm 1.3 \text{ g m}^{-2}$ ,  $5.4 \pm 1.3 \text{ mm}$  and  $30 \pm 35$
- shoots  $m^{-2}$ ). In September and October 2018, no shoots or leaves were observed (Fig. 2a). The
- biomass of rhizomes and roots reached also its maximum in October 2017 (599.7  $\pm$  36.8 g m<sup>-</sup>
- <sup>2</sup>). In contrast to leaves and shoots, the belowground biomass was stable until March 2018
- when a decline was observed that continued until October 2018 ( $30.5 \pm 6.8 \text{ g m}^{-2}$ ) (Fig. 2a).
- 284
- 285 3.2.2 Total lipid (TL) concentrations and fatty acid composition
- TL in the *C. nodosa* above-ground tissue  $(6.7 25.3 \pm 2.4 \text{ mg g}^{-1} \text{ DW})$  increased until
- February 2018, when maximum TL concentrations were measured (Fig. 2b). Thereafter, TL
- 288 concentrations decreased until August 2018. During this period, the belowground TL

concentration  $(6.3 \pm 1.9 - 15.9 \pm 1.1 \text{ mg g}^{-1} \text{ DW})$  was generally lower than the above-ground TL concentrations and the trend was similar to that of leaves. The minimum concentrations of TL were observed in September 2018, while in October 2018, concentrations similar to that measured in October 2017 were observed (Fig. 2b).

The major fatty acid components in C. nodosa tissues were palmitic (C16:0) amongst the 293 saturated (SAT) and oleic (C18:1n-9) in monounsaturated fatty acids (MUFA). In the above-294 ground tissue, the main polyunsaturated fatty acids (PUFA) were  $\alpha$ -linolenic (C18:3 n-3, 295 ALA) and linoleic (C18:2 n-6, LA), while in the belowground tissue LA was dominant (Fig. 296 2b). The dynamics of UND in the above-ground tissue was principally influenced by changes 297 in ALA and LA. LA/ALA ratios were < 1 from July 2017 to March 2018, and > 1 from April 298 to July 2018 (Fig. 2b). In August 2018, the LA/ALA ratio was infinite due to the absence of 299 ALA (Fig. 2b). Elemental sulfur ( $S^0$ ) was detected only in decaying leaves in August 2018 300  $(0.21 \text{ mg g}^{-1} \text{ DW})$ . In the belowground tissue, S<sup>0</sup> was detected in all samples (Fig. 2b). Higher 301 concentrations were measured during summer 2017 (up to  $0.39 \pm 0.06$  mg g<sup>-1</sup> DW). S<sup>0</sup> 302 increased from minimum concentrations in April  $(0.02 \pm 0.01 \text{ mg g}^{-1} \text{ DW})$  until September 303 2018 reaching 1.42 mg  $g^{-1}$  DW (Fig. 2b). 304

305 According to the fatty acid profiles, C. nodosa leaves were classified in three groups, except for the leaves collected in August 2018 (Fig. 3). The most distinguishing features 306 307 specifying physiological differences between Group 1 (July - October 2017 and February -March 2018), Group 2 (November - December 2017 and April - May 2018) and Group 3 308 (June and July 2018) were decreasing mean values of PUFA, UND, ALA and LA and 309 increasing means of SAT and the proportion of long-chain saturated fatty acids (C  $\geq$  24). In 310 the ungrouped leaves from August 2018 ALA was not found, PUFA and UND were at a 311 minimum, while SAT and  $C \ge 24$  at a maximum (Table S3). Three groups of rhizomes and 312 roots (Group 1: July - October 2017 and February - March 2018; Group 2: November -313 December 2017 and April - May 2018 and Group 3: (June - October 2018) showed similar 314 characteristics to the groups 1, 2 and 3 of related leaves (Table S4). 315

316

317 3.2.3 Epiphytic macroalgae

- From July 2017 to February 2018 different taxa of macroalgae belonging to the three phyla
- 319 Chlorophyta (Halimeda tuna, Dasycladus vermicularis, Cladophora prolifera, Udotea
- 320 petiolata), Rhodophyta (Rytiphlaea tinctoria, Peyssonnelia spp, Gelidium sp.) and

Ochrophyta (*Dictyota dichotoma*) were covering the meadow in varying proportions and
abundances (Fig. 4). After March 2018, when only few individuals of *Peyssonnelia* sp. were
found, macroalgae were no longer present in the *C. nodosa* meadow.

Although the fatty acid profiles of macroalgal communities were highly variable, the 324 contribution of 18- and 20 PUFA to the total PUFA pool generally depended on the prevailing 325 phyla and their characteristic PUFA pattern. The algae belonging to Rhodophyta and 326 Ochrophyta are richer in 20 PUFA (C20:5n-3, C20:4n-6), while Chlorophyta are generally 327 showing prevalence of 18 PUFA (C18:3n-3, C18:2n-6) (Schmid et al., 2014, Gao et al., 328 2018). Furthermore, their contribution to biomass varied due to large differences in 329 morphology, which most likely also contributed to the variability of fatty acid profiles. 18 330 PUFA and 20 PUFA showed the highest contribution to the total PUFA pool during the 331 dominance of Chlorophyta and Rhodophyta in the macroalgal community, respectively. In 332 most samples, the lowest contribution to the total PUFA pool was observed for 16 PUFA and 333 22 PUFA (Fig. S3). 334

335

336 3.3 Sediment

337 3.3.1 Granulometric composition

According to the granulometric composition, median grain sizes  $(d_g)$  and permeability (k) the 338 vegetated and non-vegetated sediments were classified as slightly gravelly sandy mud (g)sM, 339 fine grained (d<sub>g</sub> < 165  $\mu$ m) and low permeable to impermeable sediment (k < 2 \cdot 10^{-11} m<sup>2</sup>). In 340 general, the C. nodosa sediment consisted of a significantly higher proportion of sand (Sa), 341 and lower proportion of silt (Si) and clay (C) (Sa,  $41.11 \pm 4.34$  %; Si,  $46.44 \pm 2.86$  %; C, 9.63 342  $\pm 2.76$  %) in comparison to non-vegetated sediment (Sa,  $20.53 \pm 10.49$  %; Si,  $53.24 \pm 6.76$  %; 343 C,  $23.29 \pm 4.86$  %). The median grain size and permeability in C. nodosa sediment (d<sub>g</sub>, 37.51 344  $\pm 17.97 \,\mu\text{m}$ , k,  $1.22 \cdot 10^{-12} \pm 1.13 \cdot 10^{-12} \,\text{m}^2$ ) were significantly higher than in non-vegetated 345 sediment (dg,  $10.86 \pm 5.34 \,\mu\text{m}$ ; k,  $1.04 \cdot 10^{-13} \pm 1.02 \cdot 10^{-13} \,\text{m}^2$ ). The upper layers of both cores 346 (0 - 4 cm) had larger particles, while the lower layers (5 - 8 cm) showed a uniform distribution 347 348 of smaller grain sizes (Fig. 5).

349

350 3.3.2  $O_2$ ,  $E_h$ ,  $H_2S$  and  $S^0$ 

351 Oxygen concentrations  $(O_2)$  in the bottom water of the *C. nodosa* meadow varied in a wide

range (0  $\mu$ M - 171.4 ± 17.6  $\mu$ M) and generally followed the O<sub>2</sub> saturation trend (Fig. 6a).

- From May to June 2018, O<sub>2</sub> decreased below 62.5 μM, considered as severe hypoxia (Vaquer-
- Sunyer and Duarte 2008) and was completely depleted in July 2018 (Fig. 6a). From August to
- 355 October 2018,  $O_2$  increased again. The variations of  $O_2$  in the bottom water of the non-
- vegetated sediment were similar to those in the *C. nodosa* meadow albeit generally higher
- 357  $(79.4 \pm 10.4 \,\mu\text{M} 212.2 \pm 33.4 \,\mu\text{M})$  than in the vegetated sediment except for September and
- 358 October 2018 (Fig. 6a).
- In general, O<sub>2</sub> penetration depth in the vegetated and non-vegetated sediment co-varied with the O<sub>2</sub> concentration in the bottom layer, penetrating deeper when its concentration in the bottom water was higher (Fig. 6b). In the vegetated sediment, O<sub>2</sub> was mainly depleted down to 1 cm of depth. In the non-vegetated sediment, the oxygen penetration depth was up to 4 times higher than in vegetated sediments, except for the period from August 2018 to October
- 364 2018 when the penetration depths were similar (Fig. 6b).
- The thickness of the oxic (Eh > 150 mV) and suboxic (150 mV > Eh > 0 mV) layers in the 365 vegetated sediment increased from July 2017 (~ 0.5 cm) to March 2018 (~ 4 cm), and 366 367 decreased progressively from April (~ 0.8 cm) towards the surface in July 2018, when the entire sediment core was anoxic (Eh < 0). From August (~ 1 cm) to October 2018 (~ 2.5 cm) 368 369 the oxic and suboxic layer thickness increased again (Fig. 7). Oxic conditions (Eh > 0)generally reflected O<sub>2</sub> concentrations in the bottom waters. The dynamics of Eh in non-370 vegetated sediment were similar to those in the vegetated sediment. However, the thickness of 371 the oxic layer was considerably larger than in the vegetated sediment. Reducing conditions 372 (Eh < 0) were only recorded in July and August 2017 (Fig. 7). 373
- 374Concentrations of free  $H_2S$  in the pore water of the vegetated sediment generally increased375with depth creating an accumulation zone mainly within the upper sediment layers (1 4 cm)
- 376 (Fig. 7). From July to November 2017,  $H_2S$  concentrations increased up to 120  $\mu$ M (at 4 5
- 377 cm). In December 2017,  $H_2S$  was low and uniformly distributed throughout the core (< 5
- $\mu$ M). H<sub>2</sub>S concentrations increased and the accumulation layer was ascending from March (up
- to  $34.2 \pm 12.8 \ \mu\text{M}$ ; 5 7 cm) to April 2018 (up to  $177.2 \pm 125.1 \ \mu\text{M}$ ; 3.5 4.5 cm). During
- 380 May 2018 (up to  $107.8 \pm 75.9 \,\mu\text{M}$ ; 2.5 4 cm), June (up to  $199.0 \pm 6.3 \,\mu\text{M}$ ; 1.5 6 cm) and
- July (up to  $210.1 \pm 138.9 \,\mu$ M; bottom water 6 cm) a propagation of the accumulation zone
- 382 was observed in addition to an increase in  $H_2S$  (Fig. 7). In August 2018 (up to 1164.1 ± 702.1
- $\mu$ M; bottom water 7 cm) extremely high concentrations over the entire sediment core were
- recorded. In September and October 2018,  $H_2S$  concentrations decreased (down to 140.0 ±

- 25.3 and  $72.7 \pm 52.7 \mu$ M; bottom water 7 cm and 1 7 cm, respectively). In the nonvegetated sediment, H<sub>2</sub>S depth profiles were similar to those in vegetated sediments, but the concentrations were generally lower, except for the summer of 2017 when the concentrations were comparable but the accumulation zones deeper (Fig. 7).
- S<sup>0</sup> mainly occurred in oxic (Eh > 150 mV) and suboxic (150 mV > Eh > 0 mV) layers of both, vegetated and non-vegetated sediments (Fig. 7). Generally, the ranges of approximated S<sup>0</sup> concentrations in vegetated sediment ( $8.5 \cdot 10^{-5} - 0.39 \text{ mg} \cdot \text{g}^{-1} \text{ DW} \sim 2.6 \cdot 10^{-3} - 12.1 \mu \text{mol} \cdot \text{g}^{-1}$ DW), except for the extreme value in April 2018 (0.99 mg \cdot \text{g}^{-1} DW ~ 30.8 \mu \text{mol} \cdot \text{g}^{-1} DW), were similar to those found at the non-vegetated sites ( $2.9 \cdot 10^{-4} - 0.28 \text{ mg} \cdot \text{g}^{-1} \text{ DW} \sim 9.2 \cdot 10^{-3} - 8.9 \mu \text{mol} \cdot \text{g}^{-1} \text{ DW}$ ).
- 395

396 3.3.3 Prokaryotic abundance

Prokaryotic abundance varied largely in vegetated  $(2.1 - 39.9 \cdot 10^7 \text{ cells g}^{-1} \text{ fresh weight, FW})$ 397 and non-vegetated sediments  $(3.7 - 24.1 \cdot 10^7 \text{ cells g}^{-1} \text{ FW})$ . Prokaryotic abundance was 398 significantly higher in the upper than the lower layers of vegetated (F = 40.553, p < 0.05) and 399 non-vegetated (F = 52.531, p < 0.05) sediments (Fig. 8). Prokaryotic abundance showed 400 401 significant monthly changes in the upper (F = 3.053, p < 0.05) and lower layer (F = 5.035, p < (0.05) of vegetated sediments, in contrast to both layers of non-vegetated sediments (p > 0.05). 402 Prokaryotic abundances were significantly higher in the upper layers (F = 44.577, p < 0.05) 403 and significantly lower in the lower layers (F = 5.986, p < 0.05) of vegetated than in the 404 respective layers of non-vegetated sediments (Fig. 8). In the upper sediment layer, prokaryotic 405 abundances were significantly higher in the vegetated than in the non-vegetated sediments 406 from July to October 2017 and from June to August 2018 (Fig. 8). In the lower layers of 407 vegetated sediments, prokaryotic abundance was significantly higher than in the non-408 vegetated sediments in October 2017 and in August and September 2018 (Fig. 8). 409 410

411 3.3.4 Organic matter, total lipids and fatty acid composition

412 The concentrations of organic matter (OM) and total lipids (TL) were highly correlated in

413 vegetated (OM: 37.6 - 231.1 mg/g DW, TL: 0.15 - 2.75 mg/g DW; F = 214.172, p < 0.05) as

- 414 well as in non-vegetated sediments (OM: 56.7 160.3 mg/g DW, TL: 0.33 2.39 mg/g DW; F
- 415 = 45.569, p < 0.05). OM and TL generally decreased with depth and exhibited similar

- 416 changes throughout the investigated period with significantly higher concentrations in upper 417 than in lower sediment layers (p < 0.05) (Fig. 9).
- In the vegetated sediment, TL showed significant monthly changes in the upper (F =418 11.418, p < 0.05) and lower sediment layers (F = 3.186, p < 0.05), in contrast to both layers of 419 non-vegetated sediment (p > 0.05). From July to October 2017, in the upper layer of vegetated 420 sediments, TL was significantly higher than in non-vegetated sediments (Fig. 9). From 421 November 2017 onwards, TL decreased slightly until April 2018, reaching similar 422 concentrations as TL in non-vegetated sediments (Fig. 9). TL concentrations decreased 423 markedly in May and continued until August 2018. During that period, TL in vegetated 424 sediments was significantly lower than in non-vegetated sediments. In September and October 425 2018, TL concentrations in vegetated sediments were similar to those in non-vegetated 426 sediment (Fig. 9). 427 428 The fatty acid composition of vegetated and non-vegetated sediments was similar and in both layers characterized by the prevalence of SAT (vegetated upper: 71.2 - 90.4%, lower: 429 430 75.9-89.1%; non-vegetated upper: 71.2-80.7%, lower: 78.2-82.5%) over MUFA (vegetated
- 431 upper: 7.6-22.9%, lower: 9.0-19.9%; non-vegetated upper: 17.8-24.1%, lower: 15.3-18.2%)
- and PUFA (vegetated upper: 1.9-6.9%, lower: 1.9-5.1%; non-vegetated upper: 1.7-4.8%,
- lower: 1.7-3.9%). The trends of the monthly changes in UND were similar in both layers of
- both sediment types. Those variations were less pronounced in the non-vegetated sediment
  - where UND varied in narrower ranges in both layers (upper: 0.26-0.51, lower: 0.23-0.33) than
- 436 in vegetated sediment (upper: 0.13-0.57, lower: 0.14-0.37). From July to October 2017 and in
- 437 April 2018, UND was higher in the upper layers of vegetated sediment than in non-vegetated
- 438 one, while from November 2017 to March 2018, UNDs of both sediments were lower than in
- 439 previous period (Fig. 9). From June to August 2018, UND decreased considerably in
- 440 vegetated sediment, being lower than in non-vegetated sediments. During September and
- 441 October 2018, an increase of UND was observed in both sediments. In the lower layers,
- 442 UNDs were similar, except for July and August 2018 when a considerable decrease of UND443 was observed in vegetated sediments (Fig. 9).
- The proportions of PUFAs with chain lengths of 16, 18, 20, and 22 C atoms within the PUFA pool were similar between the respective layers of both sediments. Throughout the study period, the highest contribution of 18PUFA originated from *C. nodosa* detritus and Chlorophyta was observed (Fig. S4, Table S2). From July to October 2017, April to May
- 15

- 448 2018 and September to October 2018, a contribution of 20PUFA attributed to phytoplankton
- and Rhodophyta was also detected. 16PUFA and 22PUFA accounted for the smallest
- 450 contribution to the PUFA pool and were found in seston and macroalgae (Fig. S4, Table S2).
- 451 The similarities between the sediments were also observed in the contribution of the main
- 452 SAT components to the SAT pool from July 2017 to March 2018 and from September to
- 453 October 2018 (Fig. S4, Table S2). From April to August 2018, an increase of the long-chain
- 454  $(C \ge 24)$  and common (C16:0 + C18:0) fatty acids followed by the decrease of bacterial fatty
- acids (BACT) contribution to the SAT pool was observed in both layers of the vegetated
- 456 sediment. In contrast, the contribution of these components to the SAT pool was fairly
- 457 invariable in non-vegetated sediments during the same period (Fig. S4, Table S2).
- 458
- 459 3.3.5 Relationship between different physicochemical parameters
- 460 The relationships between  $H_2S$ ,  $O_2$ , TL,  $S^0$ , PA, Eh and UND in vegetated and non-vegetated
- sediment are shown in the principal component analysis, where PC1 explained 42.5 % and
- 462 PC2 14.4 % of variability (Fig. 10). The loadings for positive relationships were obtained for
- 463  $H_2S$  (0.298) on PC1 and Eh (0.541) and O<sub>2</sub> (0.327) on PC2. For the negative relationships, the
- 464 loadings were for TL (-0.534), UND (-0.494),  $S^0$  (-0.388), Eh (-0.327), PA (-0.296) and  $O_2$  (-
- 465 0.191) on PC1, and H<sub>2</sub>S (-0.536), S<sup>0</sup> (-0.485), TL (-0.165) and UND (-0.221) on PC2.
- 466 PC1 separated most of the upper sediment layers (July 2017 May 2018, September -
- 467 October 2018) according to the higher concentrations of TL and  $S^0$ , higher UND and more
- 468 positive Eh from the most of the lower layers and upper layers of vegetated sediments (June -
- August 2018) with increased  $H_2S$  concentrations. On PC2, the vegetated was separated from
- 470 the non-vegetated sediment due to higher concentrations of  $H_2S$ ,  $S^0$  and more negative Eh,
- 471 which characterized vegetated sediments during almost the entire study period. The extreme
- 472 concentrations of  $S^0$  and  $H_2S$  found in the upper layer in April and the lower layer in August
- 2018, respectively, were responsible for the considerable separation of these layers from all
- 474 other vegetated layers (Fig. 10).
- 475

## 476 **4 Discussion**

- 477 Saline Bay is a shallow, highly dynamic coastal area characterized by frequent turbid waters
- 478 due to the combined effect of land run-off and wind-driven resuspension of fine sediment.
- 479 Nutrients and Chl a (as a proxy for autotrophic biomass) varied in the ranges characteristic for

the oligotrophic coastal waters off Rovinj (Ivančić et al., 2018). The dynamics of particulate 480 matter was associated with freshwater input. The higher contribution from autochthonous 481 sources was observed during the increases of autotrophic biomass. However, only in 482 September 2017, this increase was supported by nutrients from the water column, while all 483 other increases were most likely connected to bottom waters where phytoplankton could have 484 been supplied with nutrients through sediment resuspension. The considerable increase in the 485 particulate matter of terrigenous origin from April to August 2018 suggested the enhanced 486 487 land run-off in that period.

488 In temperate Mediterranean coastal waters C. nodosa meadows show a clear unimodal annual growth cycle, reaching maximum development in summer, and minima during winter 489 and a particularly active growth phase in spring (Terrados and Ross, 1992; Zavodnik et al., 490 1998; Agostini et al., 2003). In Saline Bay, the maximum biomass was measured in October 491 492 2017. This shift from summer to early autumn was most likely due to an intense grazing activities (Cebrian et al., 1996; Valentine and Duffy, 2006) suggested by a prevalence of 493 494 visibly grazed leaves during July and August 2017. A minimum growth occurred during late autumn/winter, as commonly observed. However, during the spring 2018, phenological 495 496 parameters continued to decrease in spite of established favorable environmental conditions 497 for growth, i.e., increase in water temperature, intensity and period of solar radiation. This decrease continued until the complete extinction of the above-ground tissue in August 2018. 498 The belowground tissue followed a similar trend, but with less expressed changes. Still, their 499 recognizable remnants were found after the loss of the above-ground tissues. 500

Organic matter and closely correlated total lipids in the sediment of C. nodosa rooted area 501 changed significantly throughout the investigated period, in contrast to organic matter in non-502 vegetated sediment. Nevertheless, considerable similarity in the quality and degradation of 503 lipid matter at both, the vegetated and the non-vegetated sites indicates an important 504 505 contribution of detritus imported from the meadow as a source of organic matter for prokaryotes in non-vegetated sediments. This close coupling could be expected due to site 506 proximity and lower organic content of the non-vegetated sediment, which should enhance the 507 dependence of prokaryotes on the imports of seagrass detritus from the adjacent meadows 508 (Holmer et al., 2004). Significant enrichment of C. nodosa sediment with unsaturated, more 509 labile components only during abundant growth of meadow could be explained by more 510 efficient entrapment of seston material within the meadow (Gacia and Duarte, 2001). Such 511

easily utilizable organic matter, including dissolved monomeric carbohydrates, leaching out
during decomposition of *C. nodosa* leaves stimulated prokaryotic growth as previously
observed (Peduzzi and Herndl, 1991).

From July 2017 to March 2018, an adaptation of *C. nodosa* leaves to the decreasing light 515 516 and temperature occurred. Until October 2017, the temperature of the water column was still optimal for elongation of the leaves and biomass increase, while the ambient light intensities 517 were continuously decreasing. An additional reduction of available light might occur from the 518 self-shading effect due to high canopy biomass, and/or shading due to epiphytic macroalgae 519 growth. Desaturation of low and fairly invariable lipids during the most active growth phase 520 suggested an increase in the membrane fluidity to optimize photosynthetic activity under low 521 light conditions. Such physiological adaptation was found in seagrasses living along a depth 522 gradient (Beca-Carretero et al., 2019) and macroalgae in contrasting seasons (Schmid et al., 523 524 2014). In late autumn 2017 g 2018, the decrease in desaturation indicated a reduced fluidity and activity of photosynthetically active membranes (Quigg et al., 2006; Wacker et al., 2016). 525 526 This was associated with a decreased abundance of shoots and above-ground biomass. By shedding leaves and shoots the plant further balances metabolic requirements and mobilize 527 528 energy from the carbohydrate reserves stored in the belowground tissue (Alcoverro et al., 529 2001; Lee et al., 2007). During the winter, due to a sharp and continuous decrease in water temperature, rapid desaturation of increasing lipids provided a cold resistance, as regularly 530 observed in algae and plants (Terrados and Lopezjimenez, 1996; Iveša et al., 2004; Upchurch, 531 2008). 532

In a healthy seagrass meadow, the oxygen generated by seagrass photosynthesis is 533 transported to belowground tissues to maintain an oxic microsphere around roots and 534 rhizomes, re-oxidize sulfide to non-toxic  $S^0$ , thus preventing an invasion of H<sub>2</sub>S into the plant 535 (Pedersen et al., 1998; Holmer et al. 2005). S<sup>0</sup> was found in the *C. nodosa* below-ground 536 537 tissue during the entire investigation period, as already observed in seagrasses living in sulfidic sediments (Holmer and Hasler-Sheetal, 2014; Hasler-Sheetal and Holmer, 2015). The 538 relatively low accumulation of  $H_2S$  (< 30  $\mu$ M) during the summer and early autumn 2017 539 540 indicated that H<sub>2</sub>S was apparently rapidly recycled within the rooted area via re-oxidation by  $O_2$  to  $S^0$  and/or removal by precipitation with iron compounds. Most of  $S^0$  was found in oxic 541 layers or suboxic/anoxic boundaries, being in ranges typical for sulfidic coastal sediments 542 (Troelsen and Jørgensen, 1982; Panutrakul et al., 2001; Pjevac et al., 2014). The oxidation of 543 18

H<sub>2</sub>S could occur spontaneously by chemical reaction with free oxygen or mediated by sulfide-544 oxidizing bacteria surrounding or being attached to seagrass roots (Jørgensen, 1977; Cucio et 545 al., 2016; Ugarelli et al., 2017; Fahimipour et al., 2017). In November, due to the degradation 546 of organic matter and reduced oxygen production and leakage in the rooted zone caused by C. 547 nodosa senescence, the re-oxidation capacity of the sediment was greatly decreased. This 548 resulted in considerable accumulation of  $H_2S$  (> 100  $\mu$ M) which extended up to the sediment 549 surface. During winter and early spring, H<sub>2</sub>S production generally decreased, likely due to the 550 reduced activity of sulfate reducing prokaryotes at lower temperatures, and the sediment 551 552 gradually shifted towards a more oxidized state. H<sub>2</sub>S detected even in within the oxic sediment and in the rooted area in February 2018 could be attributed to the sediment 553 554 heterogeneity and the presence of reducing micro-niches where anaerobic metabolism could occur regardless of surrounding redox conditions (Jørgensen, 1977; Frederiksen and Glud, 555 556 2006).

In April 2018, C. nodosa had been most probably exposed to increased siltation, due to an 557 558 increase in terrigenous input combined with resuspension of sediment provoking elevated autotrophic growth. The intensive siltation is associated with the increased light attenuation, 559 560 both through the direct shading effect of suspended sediments and through the promotion of phytoplankton and epiphyte growth by the associated increase in nutrients (Terrados et al., 561 1998; Halun et al., 2002; Brodersen et al., 2015). Therefore, the increase in seawater turbidity 562 and considerable sediment re-deposition on the leaves might have severely impaired the light 563 availability and slowed down the plant's photosynthetic activity as indicates LA/ALA > 1 in 564 the above-ground tissue resulting from decreased conversion of LA to ALA (Harris and 565 James, 1965). When the minimum light requirements (~14% of incidence light) are not met, 566 C. nodosa intensely sheds leaves and shoots (Collier et al., 2012). Such light condition 567 apparently persisted until May 2018 and most likely prevented the re-establishment of 568 photosynthesis and C. nodosa continued to shed shoots and leaves. The reduced 569 photosynthesis and therefore O<sub>2</sub> transport from the leaves to the rhizome-root system 570 571 probably minimized root respiration. The maintenance of the oxic rhizosphere and the internal  $O_2$  partial pressure in the lacunae further depended mainly on the diffusion of  $O_2$  from the 572 water column. From April to June 2018, O<sub>2</sub> in the bottom water drastically decreased. Due to 573 poor supply, O<sub>2</sub> content of the belowground tissue was too low to maintain the oxic 574

microenvironment and therefore, the plant tissues became potentially accessible to sulfideintrusion (Pedersen et al., 2004).

At the same time, the sediment was enriched with fresh organic matter derived from 577 578 increased autotrophic biomass in bottom waters. In addition to the induction of the bloom, strong sediment resuspension, most likely by aeration, stimulated the intense oxidation of H<sub>2</sub>S 579 that started to produce in the rooted zone (up to 180 µM), due to increased activity of sulfate 580 reducing prokaryotes possibly triggered by the increase in temperature. An increase in  $S^0$ 581 concentration that reached its maximum in the same layer suggests a simultaneous oxidation 582 583 of the produced H<sub>2</sub>S. The sulfide oxidation probably caused oxygen depletion in the rooted zone and anoxic zone extension up to the sediment subsurface. In May 2018, the excess of 584 organic matter accumulated in April 2018 was degraded. The concentrations of S<sup>0</sup>, detected 585 only in the suboxic layer, considerably decreased possibly by disproportionation or respiration 586 587 by members of the sulfate reducing bacteria (Pjevac et al., 2014).

During June and July 2018, a sudden and significant deterioration of C. nodosa 588 589 physiological condition was indicated by the further increase in LA/ALA ratio in the leaves and overall saturation of decreasing lipids in above- and below-ground tissues. Additionally, 590 591 the loss of leaf tissue negatively impacted the photosynthetic carbon fixation and therefore oxygen production, including the transport of oxygen to below-ground tissue (Lee and 592 Dunton, 1997; Lee et al., 2007). The below-ground tissue that was not supported by 593 photosynthetically derived oxygen became anoxic. Thus induced anaerobiosis most likely 594 caused a complete inhibition of the fatty acid desaturation chain (Harris and James, 1965) and 595 a permanent breakdown of photosynthesis leading to the final decay of the above-ground 596 biomass and considerable loss of below-ground biomass. As the bottom waters were 597 completely depleted in O<sub>2</sub> the whole plant was probably exposed to sulfides. H<sub>2</sub>S inhibit 598 cytochrome c oxidase by binding to regulatory sites on the enzyme, reducing the rate of 599 600 cellular respiration and leading to the chemical asphyxiation (Nichols et al., 2013). From June to August 2018, the decomposition of organic matter, encompassing the entire 601 sediment core, was intensified and accompanied by a large increase in H<sub>2</sub>S concentrations (up 602 to 1200  $\mu$ M). The degradation process involved rhizomes and roots, as suggested by the 603

apparent loss of belowground biomass. Such loss typically occurs in the first stage of plant

- decay, the leaching phase (Trevathan-Tackett et al., 2017). Readily available, soluble
- 606 carbohydrates that largely contribute to the leachate mass (Vichkovitten and Holmer, 2004)

- most probably supported the increase in prokaryotic abundance observed in June and July
- 608 2018. However, the significant decrease in prokaryotic abundance that coincided with a
- 609 maximum degradation of organic matter and  $H_2S$  production in August 2018 might indicate
- 610 that remaining compounds were not degradable by the sulfate reduction pathway (Arndt et al.,
- 611 2013) and needed the presence of prokaryotes specialized in the anaerobic degradation of
- 612 refractory compounds, including cellulose and lignin.
- During September and October 2018, H<sub>2</sub>S concentrations drastically decreased, and the 613 sediment was gradually enriched in fresh organic matter. Due to the combined effect of 614 615 freshened oxygenated water inflow and resuspension which gradually deepened the oxic layer, re-oxidation of H<sub>2</sub>S increased. Biogeochemical studies suggest that most sulfides (80 – 616 90 %) are eventually re-oxidized; 10 - 20 % are ultimately buried as complexes with iron (i.e. 617 FeS, FeS<sub>2</sub>) or with organic matter after sulfurization (Jørgensen, 1977; 1982). H<sub>2</sub>S scavenging 618 619 with iron and formation of iron sulfides might be more important in Saline Bay, since terrestrial waters are washing out terra rossa, rich in Fe-oxides and oxyhydroxides (Durn, 620 621 2003). For this reason, sediment cores were most likely always black with sulfuric odor,
- 622 irrespective of  $H_2S$  concentrations or presence of vegetation.
- 623

### 624 5 Conclusions

Our results provide insights into the interaction of multiple stressors that have led to the meadow decay, triggered in the sensitive recruitment phase of meadow growth. Even after the improvement of the sediment conditions by the end of the summer 2018, *C. nodosa* was not able to recolonize its previously occupied areas. This finding combined with a visible alteration of the water column and sediment indicates a considerable loss of the *C. nodosa* habitat. Further research is needed to examine the fate of Saline Bay meadows and an eventual recolonization of the area.

Beyond seagrass itself, this loss had extensive consequences as it has endangered many species that depend on seagrass for food, shelter and nursery. Given the lack of data on the ecological and conservation status of the still numerous seagrass meadows along the northern Adriatic coast, the identification and monitoring of the main pressures acting on them are needed to protect such valuable habitats from degradation and extinction.

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- 639 II, LJI, IF and MN; Formal analysis and Writing original draft: MN; Writing review &
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#### References 646

- Agostini, S., Pergent, G., and Marchand, B.: Growth and primary production of *Cymodocea* 647 nodosa in a coastal lagoon. Aquat. Bot., 76, 185-193, 2003. 648
- Alcoverro, T., Manzanera, M., and Romero J.: Annual metabolic carbon balance of the 649
- 650 seagrass Posidonia oceanica: the importance of carbohydrate reserves. Mar. Ecol. Prog.
- Ser., 211, 105-116, 2001. 651
- Arndt, S., Jørgensen, B.B., LaRowe, D.E., Middelburg, J.J., Pancost, R.D., and Regnier, P.: 652 Quantifying the degradation of organic matter in marine sediments: A review and 653 synthesis. Earth-Science Rev., 123, 53-86, 2013.
- 654
- Beca-Carretero, P., Guihéneuf, F., Marín-Guirao, L., Bernardeau-Esteller, J., García-Muñoz, 655 R., Stengel, D.B., and Ruiz, J.M. Effects of an experimental heat wave on fatty acid 656
- composition in two Mediterranean seagrass species. Mar. Pollut. Bull., 134, 27-37, 2018. 657
- Beca-Carretero, P., Guihéneuf, F., Winters, G., and Stengel, D.B.: Depth-induced adjustment 658
- of fatty acid and pigment composition suggests high biochemical plasticity in the tropical 659 660 seagrass Halophila stipulacea. Mar. Ecol. Prog. Ser., 608, 105-117, 2019.
- Borum, J., Pedersen, O., Greve, T.M. Frankovich, T.A., Zieman, J.C., Fourgurean, J.W., and 661 662 Madden, C.J.: The potential role of plant oxygen and sulphide dynamics in die-off events of the tropical seagrass, Thalassia testudinum. J. Ecol., 93, 148-158, 2005. 663
- Bourbonniere, R.A., and Meyers, P.A.: Sedimentary geolipid records of historical changes in 664
- the watersheds and productivities of Lakes Ontario and Erie. Limnol. Oceanogr., 41, 352-665 359, 1996. 666
- Brodersen, K.E., Lichtenberg, M., Paz, L-C., and Kühl, M.: Epiphyte-cover on seagrass 667 (Zostera marina L.) leaves impedes plant performance and radial O<sub>2</sub> loss from the bellow-668
- ground tissue. Front. Mar. Sci., 2, 58 doi: 10.3389/fmars.201500058, 2015. 669
- Cancemi, G., Buia, M.C., and Mazzella, L.: Structure and growth dynamics of Cymodocea 670 nodosa meadows. Sci. Mar., 66: 365-373, 2002. 671
- Canfield, D.E., Jørgensen, B.B., Fossing, H., Glud, R., Gundersen, J., Ramsing, N.B., 672
- Thamdrup, B., Hansen, J.W., Nielsen, L.P., and Hall, P.O.J.: Pathways of organic carbon 673 oxidation in three continental margin sediments. Mar. Geol., 113, 27-40, 1993. 674
- Capone, D.G., and Kiene, R.P.: Comparison of microbial dynamics in marine and freshwater 675
- sediments: Contrasts in anaerobic carbon catabolism. Limnol. Oceanogr., 33, 725-749, 676

1988. 677

- Carlson, P.R., Yarbro, L.A., and Barber, T.R.: Relationship of sediment sulfide to mortality of
   *Thalassia testudinum*, Florida Bay. Bull. Mar. Sci., 54, 733-746, 1994.
- 680 Cebrian, J., Duarte, C.M., and Marbà, N.: Herbivory on the seagrass Cymodocea nodosa
- (Ucria) Ascherson in contrasting Spanish Mediterranean habitats. J. Exp. Mar. Biol. Ecol.,
  204, 103-111, 1996.
- 683 Cline, J.D.: Spectrophotometric determination of hydrogen sulfide in natural waters. Limnol.
  684 Oceanogr., 14, 454-458, 1969.
- Collier, C.J., Lavery, P.S., Masini, R.J., and Ralph, P.J.: Shade-induced response and recovery
  of the seagrass *Posidonia sinuosa*. J. Exp. Mar. Biol. Ecol., 370, 89-103, 2009.
- 687 Collier, J.C., Waycott, M., and Giraldo Ospina, A.: Responses of four Indo-West Pacific
  688 seagrass species to shading. Mar. Pollut. Bull., 65, 342-354, 2012.
- 689 Costa, M.M., Barrote, I., Silva, J., Olivé, I., Alexandre, A., Albano, S., and Santos, R.:
- 690 Epiphytes modulate *Posidonia oceanica* photosynthetic production, energetic balance,
- antioxidant mechanisms, and oxidative damage. Front. Mar. Sci. 2:111, 2015.
- 692 Cranwell, P.A., Eglinton, G., and Robinson, N.: Lipids of aquatic organisms as potential
  693 contributors to lacustrine sediments. Org. Geochem., 11, 513-527, 1987.
- Cúcio, C., Engelen, A.H., Costa, R., and Muyzer, G.: Rhizosphere Microbiomes of European
  Seagrasses Are Selected by the Plant, But Are Not Species Specific. Front. Microbiol., 7,
  440. doi: 10.3389/fmicb.2016.00440, 2012.
- 697 Duarte, C.M., Kennedy, H., Marbà, N., Gacia, E., Fourqurean, J.W., Beggins, J., Barrón, C.,
- 698 Apostolaki, E.T.: Seagrass community metabolism: Assessing the capacity of seagrass
- meadows for carbon burial: Current limitations and future strategies. Ocean Coast. Manag.,
  83, 32-38, 2013.
- Durn, G.: *Terra Rossa* in the Mediterranean Region: Parent Materials, Composition and
   Origin. Geologia Croatica, 56, 83-100, 2003.
- Epstein, S.S., and Rossel J.: Enumeration of sandy sediment bacteria: search for optimal
  protocol. Mar. Ecol. Prog. Ser., 117, 289-298, 1995.
- Fahimipour, A.K., Kardish, M.R., Lang, J.M., Green, J.L., Eisen, J.A., and Stachowicz, J.J.:
  Global-Scale Structure of the Eelgrass Microbiome. Appl. Environ. Microbiol., 83,
  e03391-16, 2017.
- Folk, R.L.: The distinction between grain size and mineral composition in sedimentary-rock
  nomenclature. J. Geol., 62, 344-359, 1954.

- 710 Frederiksen, M.S., Holmer, M., Pérez, M., Invers, O., Ruiz, J.M., and Knudsen, B.: Effect of
- 711 increased sediment sulfide concentrations on the composition of stable sulfur isotopes
- 712  $(\delta^{34}S)$  and sulfur accumulation in the seagrasses *Zostera marina* and *Posidonia oceanica*.
- 713 J. Exp. Mar. Biol. Ecol., 358, 98-109, 2008.
- Frederiksen, M.S., and Glud, R.N.: Oxygen dynamics in the rhizosphere of *Zostera marina*: A
  two-dimensional planar optode study. Limnol. Oceanogr., 51, 1072-1083, 2006.
- 716 Gacia, E., and Duarte, C.M.: Sediment Retention by a Mediterranean *Posidonia oceanica*
- Meadow: The Balance between Deposition and Resuspension. Estuar. Coast. Shelf. Sci.,
  52, 505–514, 2001.
- Gangi, A.F.: Permeability of unconsolidated sands and porous rocks. J. Geophys. Res. –Solid,
  90, 3099-3104, 1985.
- Gao, G., Clare, A.S., Chatzidimitriou, E., Rose, C., and Caldwell, G.: Effects of ocean
- warming and acidification, combined with nutrient enrichment, on chemical composition
  and functional properties of *Ulva rigida*. Food Chem., 258, 71-78, 2018.
- Greve, T.M., Borum, J., and Pedersen, O.: Meristematic oxygen variability in eelgrass
  (*Zostera marina*). Limnol. Oceanogr., 48, 210-216, 2003.
- Halun, Z., Terrados, J., Borum, J., Kamp-Nielsen, J., Duarte, C.M., and Fortes, M.D.:
- Experimental evaluation of the effects of siltation-derived changes in sediment conditions
- on the Philippine seagrass *Cymodocea rotundata*. J. Exp. Mar. Biol. Ecol., 279, 73-87,
- **729** 2002.
- Harris, R.V., and James, A.T.: Linoleic and α-linolenic acid biosynthesis in plant leaves and a
  green alga. Biochim. Biophys. Acta, 106, 456-464, 1965.
- Hasler-Sheetal, H., and Holmer, M.: Sulfide intrusion and detoxification in the seagrass *Zostera marina*. Plos One 10(6): e0129136, 2015.
- Hendriks, I.E., Sintes, T., Bouma, T.J., and Duarte, C.M.: Experimental assessment and
- modeling evaluation of the effects of the seagrass *Posidonia oceanica* on flow and particle
- 736 trapping. Mar. Ecol. Prog. Ser., 356, 163-173, 2008.
- Holm-Hansen, O., Lorenzen, C. J., Holmes, R. W., and Strickland, J. D. H.: Fluorometric
  determination of chlorophyll. J. Conseil., 301, 3-15, 1965.
- Holmer, M., Duarte, C.M., Boschker, H.T.S., and Barrón, C.: Carbon cycling and bacterial
- carbon sources in pristine and impacted Mediterranean seagrass sediments. Aquat. Microb.
- 741 Ecol., 36, 227-237, 2004.

- 742 Holmer, M., and Hasler-Sheetal, H.: Sulfide intrusion in seagrasses assessed by stable
- isotopes- a synthesis of current results. Front. Mar. Sci., doi: 10.3389/fmars.2014.00064,
  2014.
- Holmer, M., and Nielsen, S.L.: Sediment sulfur dynamics related to biomass-density pattern
  in *Zostera marina* (eelgrass) beds. Mar. Ecol. Prog. Ser., 146, 163-171, 1997.
- Holmer, M., Frederiksen, M.S., and Møllegaard, H.: Sulfur accumulation in eelgrass (*Zostera marina*) and effect of sulfur on eelgrass growth. Aquat. Bot., 81, 367-379, 2005.
- 749 Holmer, M., Pedersen, O., and Ikejima, K.: Sulfur cycling and sulfide intrusion in mixed
- Southeast Asian tropical seagrass meadows. Bot. Mar., 49, 91-102, 2006.
- 751 Ivančić, I., Paliaga, P., Pfannkuchen, M., Đakovac, T., Najdek, M., Steiner, P., Korlević, M.,
- 752 Markovski, M., Baričević, A., Smodlaka Tanković, M., and Herndl, G.J.: Seasonal
- variations in extracellular enzymatic activity in marine snow-associated microbial
- communities and their impact on the surrounding water. FEMS Microbiol. Ecol., 94,

755 fyi198, 2018.

- Iveša, Lj., Blažina, M., and Najdek, M.: Seasonal variations in fatty acid composition of *Caulerpa taxifolia* (M. Vahl.) C. Ag. in the northern Adriatic Sea (Malinska, Croatia). Bot.
  Mar., 47, 209-214, 2004.
- Jørgensen, B.B.: The sulfur cycle of a coastal marine sediment (Limfjorden, Denmark).
  Limnol. Oceanogr., 22, 814-832, 1977.
- Jørgensen, B.B.: Mineralization of organic matter in the sea bed the role of sulphate
  reduction. Nature, 296, 643-645, 1982.
- Koch, M.S., and Erskine, J.M.: Sulfide as a Phytotoxin to the Tropical Seagrass *Thalassia testudinum*: Interactions with Light, Salinity and Temperature. J. Exp. Mar. Biol. Ecol.,
   266, 81-95, 2001.
- 766 Krause-Jensen, D., Carstensen, J., Nielsen, S.L., Dalsgaard, T., Christensen, P.B., Fossing, H.,
- and Rasmussen, M.B.: Sea bottom characteristics affect depth limits of eelgrass *Zostera marina*. Mar. Ecol. Prog. Ser., 425, 91-102, 2011.
- Lee, K-S., and Dunton, K.H.: Diurnal changes in pore water sulfide concentrations in the
  seagrass *Thalassia testudinum* beds: the effects of seagrasses on sulfide dynamics. J. Exp.
  Mar. Biol. Ecol., 255, 201-214, 2000.
- 772 Lee, K-S., Park, S.R., and Kim, Y.K.: Effects of irradiance, temperature, and nutrients on
- growth dynamics of seagrasses: A review. J. Exp. Mar. Biol. Ecol., 350, 144-175, 2007.

- Marbà, N., and Duarte, C.M.: Growth and sediment space occupation by seagrass *Cymodocea nodosa* roots. Mar. Ecol. Prog. Ser., 224, 291-298, 2001.
- 776 Mascaró, O., Valdemarsen, T., Holmer, M., Pérez, M., and Romero, J.: Experimental
- manipulation of sediment organic content and water column aeration reduces *Zostera marina* (eelgrass) growth and survival. J. Exp. Mar. Biol. Ecol., 373, 26-34, 2009.
- 110 marina (cergrass) growin and survival. J. Exp. Mar. Diol. Ecol., 575, 20-54, 2007.
- Micromeritics: SediGraph 5100 particle size analysis system operator' manual. Micromeritics
  Instrument Corporation, Norcross, 2002.
- 781 Moeslund, L., Thamdrup, B., and Jørgensen, B.B.: Sulfur and iron cycling in a coastal
- sediment—radiotracer studies and seasonal dynamics. Biogeochemistry, 27, 129-152,
  1994.
- Nicholls, P., Marshall, D.C., Cooper, C.E., and Wilson, M.T.: Sulfide inhibition of and
- metabolism by cytochrome c oxidase. Biochem. Soc. Transact., 41, 1312-1316, 2013.
- 786 Orlando-Bonaca, M., Francé, J., Mavrič, B., Grego, M., Lipej, L., Flander Putrle, V., Šiško,
- M., and Falace, A.: A new index (MediSkew) for the assessment of the *Cymodocea nodosa*(Ucria) Ascherson meadow's status. Mar. Environ. Res., 110, 132-141, 2015.
- Orlando-Bonaca, M., Francé, J., Mavrič, B., and Lipej, L.: Impact of the Port of Koper on
  Cymodocea nodosa meadow. Annales, 29, 187-194, 2019.
- 791 Orth, R.J., Carruthers, T.J.B., Dennison, W.C., Duarte, C.M., Fourqurean, J.W., Heck Jr.,
- K.L., Hughes, A.R., Kendrick, G.A., Kenworthy, E.J., Olyarnik, S., Short, F.T., Waycott,
- M., and Williams, S.L.: A Global Crisis for Seagrass Ecosystems. BioScience, 56, 987996, 2006.
- Panutrakul, S., Monteny, F., and Baeyens, W.: Seasonal Variations in Sediment Sulfur
  Cycling in the Ballastplaat Mudflat, Belgium. Estuaries 24, 257-265, 2001.
- 797 Pedersen, M.F., Duarte, C.M., and Cebrián, J.: Rates of changes in organic matter and nutrient
- stocks during seagrass *Cymodocea nodosa* colonization and stand development. Mar. Ecol.
- 799 Prog. Ser., 159, 29-36, 1997.
- Pedersen, O., Borum, J., Duarte, C.M., and Fortes, M.D.: Oxygen dynamics in the rhizosphere
  of *Cymodocea rotundata*. Mar. Ecol. Prog. Ser., 169, 283-288, 1988.
- 802 Pedersen, O., Binzer, T., and Borum, J.: Sulphide intrusion in eelgrass (*Zostera marina* L.).
- 803 Plant Cell Environ., 27, 595-602, 2004.

- Peduzzi, P., and Herndl, G.J.: Decomposition and significance of seagrass litter (*Cymodocea nodosa*) for the microbial food web in coastal waters (Gulf of Trieste, Northern Adriatic
  Sea). Mar. Ecol. Prog. Ser., 71, 163-174, 1991.
- Pérez, M., Invers, O., Ruiz Fernandez, J.M., Frederiksen, M., and Holmer, M.: Physiological
  responses of the seagrass *Posidonia oceanica* to elevated organic matter content in
  sediments: An experimental assessment. J. Exp. Mar. Biol. Ecol., 344,149-160, 2007.
- 810 Pirini, M., Manuzzi, M.P., Pagliarani, A., Trombetti, F., Borgatti, A.R., and Ventrella, V.:
- 811 Changes in fatty acid composition of *Mytilus galloprovincialis* (Lmk) fed on microalgal
- and wheat germ diets. Comp. Biochem. Physiol. B, 147, 616-626, 2007.
- Pjevac, P., Kamyshny Jr., A., Dyksma, S., and Mussmann, M.: Microbial consumption of
  zero-valence sulfur in marine benthic habitats. Environ. Microbiol., 16, 3416-3430, 2014.
- 815 Porter, K.G., and Feig, Y.S.: The use of DAPI for identification and counting aquatic
- 816 microflora. Limnol. Oceanogr., 25, 943-984, 1980.
- 817 Quigg, A., Kevekordes, K., Raven, J., and Beardall, J.: Limitations on microalgal growth at
- very low photon fluency rates: the role of energy slippage. Photosynth. Res., 88, 299-310,
  2006.
- Rogowska, J., Sychowska, J., Cieszynska-Semenowicz, M., and Wolska, L.: Elemental sulfur
  in sediments: analytical problems. Environ. Sci. Pollut. Res., 23, 24871-24879, 2016.
- 822 Samper-Villarreal, J., Lovelock, C.E., Saunders, M.I., Roelfsema, C., and Mumby, P.J.:
- 823 Organic carbon in seagrass sediment is influenced by seagrass canopy complexity,
- turbidity, wave height, and water depth. Limnol. Oceanogr., 61, 938-952, 2016.
- 825 Sand-Jensen, K., Pedersen, O., Binzer, T., and Borum, J.: Contrasting Oxygen Dynamics in
- the Freshwater Isoetid *Lobelia dortmanna* and the Marine Seagrass *Zostera marina*. Ann.
  Bot., 96, 613-623, 2005.
- Schmid, M., Guihéneuf, F., and Stengel, D.B.: Fatty acid contents and profiles of 16
  macroalgae collected from the Irish Coast at two seasons. J. Appl. Phycol., 26, 451-463,
- 830 2014.
- 831 Short, F.T., Polidoro, B., Livingstone, S.R., Carpenter, K.E., Bandeira, S., Bujang, J.S.,
- Calumpong, H.P., Carruthers, T.J.B., Coles, R.G., Dennison, W.C., Erftemeijer, P.L.A.,
- Fortes, M.D., Freeman, A.S., Jagtap, T.G., Kamal, A.M., Kendrick, G.A., Kenworthy,
- 834 W.J., La Nafie, Y.A., Nasution, I.M., Orth, R.J., Prathep, A., Sanciangco, J.C., van

- Tussenbroek, B., and Vergara, S.G.: Extinction risk assessment of the world's seagrass
  species. Biol. Conserv., 144, 1961-1971, 2011.
- 837 Sousa, A.I., Calado, R., Cleary, D.F.R., Nunes, C., Coimbra, M.A., Serôdio, J., and Lillebø,
- A.I.: Effect of spatio-temporal shifts in salinity combined with other environmental
- variables on the ecological processes provided by *Zostera noltii* meadows. Sci. Rep., 7,
  1336, 2017.
- Strickland, J.D.H., and Parsons, T.R.: A practical handbook of seawater analysis. Bull. Fish.
  Res. Board. Can., 167,1-310, 1972.
- 843 Terrados, J., and Lopez-Jimenez, J.A.: Fatty acid composition and chilling resistance in the
  844 green alga *Caulerpa prolifera* (Forrskal) Lamouroux (Chlorophyta, Caulerpales).Biochem.

845 Molecul. Biol. Internatl., 39, 863-869, 1996.

- 846 Terrados, J., Duarte, C.M., Fortes, M.D., Borum, J., Agawin, N.S.R., Bach, S., Thampanya,
- 847 U., Kamp-Nielsen, L., Kenworthy, W.J., Geertz-Hansen, O., and Vermaat, J.: Changes in
- 848 Community Structure and Biomass of Seagrass Communities along Gradients of Siltation
- 849 in SE Asia. Estuar. Coast. Shelf Sci., 46, 757-768, 1998.
- Terrados, J., and Ros, J.D.: Growth and primary production of *Cymodocea nodosa* (Ucria)
  Ascherson in a Mediterranean coastal lagoon: the Mar Menor (SE Spain). Aquat. Bot., 43,
  63-74, 1992.
- 853 Trevathan-Tackett, S.M., Seymour, J.R., Nielsen, D.A., Macreadie, P.I., Jeffries, T.C.,
- Sanderman, J., Baldock, J., Howes, J.M., Steven, A.D.L., and Ralph, P.J.: Sediment anoxia
- 855 limits microbial-driven seagrass carbon remineralization under warming conditions. FEMS
  856 Microbiol. Ecol., 93,fix033, 2017.
- Troelsen, H., and Jørgensen, B.B.: Seasonal dynamics of elemental sulfur in two coastal
  sediments. Estuar. Coast. Shelf Sci., 15, 255-266, 1982.
- Tuya, F., Martín, J.A., and Luque, A.: Impact of a marina construction on seagrass bed at
  Lanzarote (Canary Islands). J. Coast. Conserv., 8, 157-162, 2002.
- Tuya, F., Ribeiro-Leite, L., Arto-Cuesta, N., Coca, J., Haroun, R., and Espino, F.: Decadal
  changes in the structure of *Cymodocea nodosa* seagrass meadows: Natural vs. human
- 863 influences. Estuar. Coast. Shelf Sci., 137, 41-49 (2014).
- Ugarelli, K., Chakrabarti, S., Laas, P., and Stingl, U.: The seagrass holobiont and its
  microbiome. Microorganisms 5, 81, 2017.

- Upchurch, R.G.: Fatty acid unsaturation, mobilization, and regulation in the response of plants
  to stress. Biotechnol. Lett., 30, 967-977, 2008.
- Valentine, J.F., and Duffy, J.E.: The central role of grazing in seagrass ecology. In:
- 869 Seagrasses: Biology, Ecology and Conservation, Springer, Netherlands, pp 431-501, 2006.
- Vaquer-Sunyer, R., and Duarte, C.M.: Thresholds of hypoxia for marine biodiversity. PNAS,
  105, 15452-15457, 2008.
- Vaquer-Sunyer, R., and Duarte, C.M.: Sulfide exposure accelerates hypoxia-driven mortality.
  Limnol. Oceanogr., 55, 1075-1082, 2010.
- Viso, A.C., Pesando, D., Bernard, P., and Marty, J.C.: Lipid components of the Mediterranean
  seagrass *Posidonia oceanica*. Mar. Pollut. Bull., 34, 381-387, 1993.
- Vichkovitten, T., and Holmer, M.: Contribution of plant carbohydrates to sedimentary carbon
  mineralization. Org. Geochem., 35,1053-1066, 2004.
- 878 Wacker, A., Piepho, M., Harwood, J.L., Guschina, I.A., and Arts, M.T.: Light-induced
- changes in fatty acid profiles of specific lipid classes in several freshwater phytoplankton
  species. Front. Plant Sci., 7, 264, 2016.
- Widdows, J., Pope, N.D., Brinsley, M.D., Asmus, H., and Asmus, R.M.: Effects of seagrass
  beds (*Zostera noltii* and *Z. marina*) on near-bed hydrodynamics and sediment
- resuspension. Mar. Ecol. Prog. Ser., 358, 125-136, 2008.
- 884 Zavodnik, N., Travizi, A., and De Rosa, S.: Seasonal variations in the rate of photosynthetic
- activity and chemical composition of the seagrass *Cymodocea nodosa* (Ucr.) Asch. Sci.
- 886 Mar., 62, 301-309, 1998.



Figure 1. Temperature (a); salinity (b), particulate matter concentration (b); unsaturation degree (UND) and terrestrial to aquatic ratio (TAR) of the particulate lipid matter (c) in seawater.



Figure 2. Above- and below-ground tissue biomasses and shoot density (a), total lipid concentrations (TL) and linoleic to  $\alpha$ -linolenic fatty acids ratios (LA/ALA, an arrow indicates an infinite value) in above-ground tissue and TL and approximated concentrations of elemental sulfur (S<sup>0</sup>) in below-ground tissue (b).





Figure 3. Cluster analysis dendrogram of fatty acid composition of *C. nodosa* leaves.

899 Summary statistics is given in Table S3.





Figure 4. The contribution of macroalgal phyla in a meadow cover and total macroalgal

902 biomass. After February 2018 macroalgae were no longer present in the *C. nodosa* meadow.

903



Figure 5. Granulometric composition and median grain size  $(d_g)$  of vegetated (a) and nonvegetated sediment (b).





Figure 6. Oxygen concentrations (O<sub>2</sub>) in bottom waters (a) and O<sub>2</sub> penetration depths (b)

above and in vegetated and non-vegetated sediment, respectively.  $O_2$  at the saturation level

910 was calculated according to the temperature and salinity measured in seawater at the sampling

911 dates;  $O_2$  at the hypoxic frontier (~ 62.5  $\mu$ M) was taken from Vaquer-Sanyer and Duarte

912 (2008).



913

Figure 7. Depth profiles of  $H_2S$  and  $S^0$  concentrations in vegetated and non-vegetated sediment (adjacent narrow graphs). The redox potential (Eh) in both sediments is shown as areas corresponding to oxic (Eh > 150 mV), suboxic (150 > Eh > 0 mV) and anoxic (Eh < 0 mV) conditions.



Figure 8. Prokaryotic abundance (PA) in the upper (0 - 4 cm) and lower (5 - 8 cm) layers of

918 vegetated and non-vegetated sediments; significant differences in PA between the sediments



920



922 Figure 9. Total lipid concentrations (TL) and unsaturation degree (UND) in the upper (0 - 4

923 cm) and lower (5 - 8 cm) layers of vegetated and non-vegetated sediments. Significant

924 differences in TL between the sediments are indicated by asterisks.



#### 926

927 Figure 10. PCA plot of redox potential (Eh), oxygen (O<sub>2</sub>), hydrogen sulfide (H<sub>2</sub>S), sulfur (S), total lipids (TL) and prokaryotes (PA)

concentrations and unsaturation degree (UND) in the upper  $(0 - 4 \text{ cm}; \Delta, \diamond)$  and lower  $(5 - 7 \text{ cm}; \blacktriangle, \blacklozenge)$  layers of vegetated and non-vegetated sediments, respectively. Projections of variables are given in circle