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- 16
- 17 Abstract

Saltmarshes are increasingly at risk from sea level rise through climate change. Their vegetation is comprised of halophytes, salt-tolerant plants, that naturally regenerate from seed, yet the combination of functional traits associated with salinity tolerance in seeds is still poorly characterised. Combining approaches in microscopy, biochemistry and physiology, we focussed on elucidating two mechanisms that are commonly utilised by adult halophyte plants: ion accumulation and elemental localisation, and antioxidant protection against oxidative stress, in seeds of the saltmarsh halophyte *Suaeda maritima* during over-wintering and germination.

Fresh seeds were dormant, and required 19 weeks of submergence in artificial sea water (ASW) 26 at 5 °C, mimicking over-wintering in the field, to release dormancy. Seed ion contents were 27 quantified, along with the redox state of glutathione and low-molecular weight thiol 28 intermediates of its synthesis (Cys, Cys-Gly, γ-Glu-Cys), the production of the reactive oxygen 29 species H<sub>2</sub>O<sub>2</sub> and electrolyte leakage to indicate ion movement. Once dormancy was released, 30 seeds were germinated in solutions of ASW and iso-osmotic solutions of polyethylene (PEG) 31 for comparison. Temporal changes in the glutathione and low-molecular weight thiols redox 32 state, H<sub>2</sub>O<sub>2</sub> and electrolyte leakage were measured during germination. In addition, spatial 33

34 localisation of the seawater elements Na, Cl, Mg, Ca, K and S within the seed was resolved,

35 and seed moisture content measured to indicate the extent of osmotic adjustment.

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Fresh and pre-treated seeds had an abundance of Na<sup>+</sup> and Cl<sup>-</sup>, as well as H<sub>2</sub>O<sub>2</sub> and thiol 37 disulphides (i.e., oxidised thiols) that are indicative of oxidative stress, but this was 38 accompanied by a large pool of reduced glutathione and thiols suggesting a potentiated 39 antioxidant system in preparation for germination. During germination in ASW, the glutathione 40 and thiol redox state was highly reduced, with no consistent production of H<sub>2</sub>O<sub>2</sub>. Na and Cl 41 42 localised in the seed testa and tegmen, while Mg and K were only found in the embryo, and Ca and S had a broad distribution across the seed. Small differences in seed moisture content did 43 not account for a higher maximum threshold to germination of 2.22 times the concentration of 44 ASW compared to 1.76 times ASW-equivalents for PEG. Moreover,  $\theta_{\text{Halo}}$  was higher (i.e., 45 slower germination) in the presence of ASW than PEG, suggesting that slower germination 46 47 may give more time to initiate tolerance mechanisms in response to salinity that promote germination. Seeds of S. maritima appear to utilise seawater ions to their benefit during 48 49 germination in salinity, using elemental localisation to protect the embryo from potential toxicity of Na and Cl, while maintaining a highly reduced cellular redox state. 50

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#### 52 Keywords

53 Glutathione redox state, functional traits, halophyte, ion localisation, seed germination, 54 halotime model.

#### 55 1. Introduction

56 Coastal wetlands provide a natural buffer against coastal erosion and salt water intrusion, 57 providing a unique habitat for plants and animals, and make a huge contribution to global 58 carbon sequestration (Hulme, 2005; Duarte et al., 2013). However, with sea level rise projected 59 to increase by up to 2 m by 2100 (Bamber et al., 2019), coastal habitats and the plants which 60 inhabit them, including salt tolerant halophytes, are under threat from climate change. 61

- Whereas salt-sensitive plants, that include most of the world's economic crops, suffer osmotic 62 and ionic stress even under low salinity levels, halophytes complete their life cycle under high 63 saline conditions of around 200 mM NaCl or more (Flowers and Colmer, 2008; Munns and 64 Tester, 2008). Nearly all species of halophyte naturally regenerate from seed and therefore rely 65 on germination to establish the next generation of plants (Sealand Dantas, 2020). Tolerance to 66 salinity during germination is variable, and can range from species whose seeds only germinate 67 68 in fresh water to those that can tolerate over 1M NaCl (Gul et al., 2013). By exploiting the linear relationship of germination rate with salinity concentration, the maximum threshold of 69 70 salt for germination to occur has been estimated for a handful of halophytes, such as Atriplex prostrata (505 mM NaCl at 15 °C), Suaeda maritima (1280 mM NaCl at 5 °C) and Tecticornia 71 72 indica (2473 mM NaCl at 15 °C) (Allen et al. 2000; Seal et al., 2018; Seal and Dantas, 2020). Yet the combination of functional traits underlying salt tolerance during germination is still 73 74 largely unknown.
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One of the first challenges is for a seed to imbibe enough water for the germination process to 76 conclude, since a saline solution may have a lower osmotic potential than that of the seed (Seal 77 and Dantas, 2020). The seeds of some species are unable to do this, and may only complete 78 their germination once the salt is removed and replaced by fresh water (Gul et al., 2013). Of 79 80 the limited studies available, some species have overcome this gradient in osmotic potential by accumulating Na<sup>+</sup> within the whole seed (e.g., Suaeda maritima), which for some has been 81 resolved further as localisation into the seed coat (e.g., Suaeda physophora and Salicornia 82 pacifica var. utahensis), to aid osmotic adjustment under saline conditions (Khan et al., 1985; 83 Song et al., 2005; Seal et al., 2018). Localisation of Na<sup>+</sup> and Cl<sup>-</sup> away from the embryo is likely 84 to be advantageous since high concentrations can be toxic to metabolism (Flowers et al., 2015). 85 86

87 Another consequence of exposure to salinity is the increased production of reactive oxygen 88 species (ROS), such as hydrogen peroxide ( $H_2O_2$ ), superoxide and the hydroxyl radical,

originating mostly from respiration in germinating seeds (Kranner and Seal, 2013; Bose et al., 89 2014). At low levels, ROS can have a range of beneficial roles. During the imbibition of dry 90 seeds, an oxidative burst of ROS, such as H<sub>2</sub>O<sub>2</sub>, may occur as metabolic activity resumes 91 (Wojtyla et al., 2016). H<sub>2</sub>O<sub>2</sub> is an important messenger and regulator of key developmental 92 processes such as dormancy release, cell-wall loosening and reserve mobilisation (Wojtyla et 93 94 al., 2016) and is present in germinating seeds of the halophytes Arthrocnemum macrostachyum, Arthrocnemum indicum, Suaeda fruticosa and Limonium stocksii (Hameed et al., 2014; Nisar 95 et al., 2019). However, ROS levels require careful control by ROS-processing enzymes and 96 97 antioxidants, to avoid oxidative damage to macromolecules and cellular structures (Kranner and Seal, 2013; Bose et al., 2014). For example, peroxidation of membrane lipids leads to a 98 loss of membrane integrity and an increase in electrolyte leakage. Eventually, the accumulation 99 of damage may be too great to repair and can result in seed death (Kranner and Seal, 2013). 100 101

102 Glutathione ( $\gamma$ -glutamyl-cysteinyl-glycine) is a major water-soluble antioxidant. In unstressed cells, glutathione is present in a reduced form (GSH) and can donate electrons to detoxify free 103 104 radicals, forming glutathione disulphide (GSSG), as part of the ascorbate-glutathione cycle (Kranner and Seal, 2013; Bose et al., 2014). The glutathione redox state has been linked to seed 105 106 viability across a range of species, where seed viability loss is associated with a switch towards 107 more oxidising cellular conditions (Kranner et al., 2006). In halophytes, glutathione has 108 frequently been identified as an important component of the antioxidant response to salinity 109 exposure. For example, in plants of Suaeda maritima, a higher concentration of GSH, lower 110 GSSG concentration and more reducing cellular conditions were observed when grown in seawater with permanently flooded conditions than in drained conditions, suggesting a 111 112 heightened glutathione antioxidant system in response to waterlogging (Alhdad et al., 2013). During germination, Suaeda fruticosa and Limonium stocksii showed an increasing 113 114 GSH:GSSG ratio during germination in 200 and 400 mM NaCl (Hameed et al., 2014). However, in seedlings of Suaeda maritima germinated in 0 - 400 mM NaCl, no significant differences 115 were observed in the concentration of GSH, GSSG or the reduced and disulphide forms of 116 other low-molecular weight thiols such as cysteine (Boestfleisch et al., 2014). Recently, it has 117 118 been proposed that H<sub>2</sub>O<sub>2</sub> levels were closely linked to temporal changes in the GSH/GSSG redox state during seed germination (Gerna et al., 2017), further supporting the notion that 119 120 cellular redox changes are intricately involved in the germination process.

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122 In this study, we investigated the combination of traits that underlie tolerance to salinity in

halophyte seeds. Our focus is on seeds of the saltmarsh halophyte Suaeda maritima 123 (Amaranthaceae) that grows in the northern hemisphere, as well as in tropical habitats as an 124 125 introduced species (Seal et al., 2018). It frequently occupies the lower saltmarsh and is adapted to temporary flooding and hypoxic conditions in the soil (Alhdad et al., 2013; Behr et al., 2017). 126 As an annual plant, mature seeds are dispersed each autumn (typically October) in a dormant 127 128 state, and over-winter in cold-saline conditions in the field until they germinate in the spring 129 (March-April; Wetson et al., 2008). S. maritima is a colonising species and germinates well in bare patches resulting from disturbance (Chapman, 1947; Tessier et al., 2000), which is 130 131 important in establishing new saltmarsh habitats such as created by sea level rise (Hulme, 2005). 132

Taking a multidisciplinary approach combining physiology, biochemistry and microscopy, we 133 first investigated the role of ion accumulation and oxidative stress (through the glutathione 134 redox state, production of the ROS H<sub>2</sub>O<sub>2</sub> and electrolyte leakage) in protecting fresh (dormant) 135 136 and pre-treated (mimicking over-wintering in the field to release dormancy) seeds from salinity. Then, once dormancy was released, seeds were germinated in solutions of artificial seawater 137 138 (ASW) and iso-osmotic solutions of polyethylene (PEG), to simulate the osmotic constraint of salinity without ion toxicity, for comparison. The halotime model (Seal et al., 2018) was 139 140 adapted to estimate, for the first time, the maximum concentration of seawater for germination 141 to occur and the halo time constant (i.e., seawater accumulation over time at a specific 142 temperature for germination to complete). Temporal changes in the redox state of glutathione and low-molecular weight thiol intermediates of its synthesis (Cys, Cys-Gly,  $\gamma$ -Glu-Cys), as 143 well as H<sub>2</sub>O<sub>2</sub> production and electrolyte leakage were also measured during germination. 144 145 Finally, the spatial localisation of the seawater elements Na, Cl, Mg, Ca, K and S within the seed was resolved, and seed moisture content measured to indicate the extent of osmotic 146 147 adjustment.

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### 149 2. Materials and Methods

150 2.1. Seed collection and pre-treatment of seeds

Mature fruits of *Suaeda maritima* (Amaranthaceae) were collected from the saltmarsh at Cuckmere Haven, Seaford, East Sussex, United Kingdom (50° 45' 39.24" N, 0 08' 57.71" E) in October 2018. Seeds were removed from the fruits by hand. The relative humidity (RH) of fresh seeds was measured with a hand-held Rotronic (Hygrolog-D, Rotronic Instruments, Crawley, UK) and corresponded to 100 % RH at 20 °C. Seeds were stored at 75 % RH at 15 °C 156 until experiments commenced.

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Seeds of *S. maritima* correspond to a perispermic seed morphology (Fig. 1). The coiled embryo, which is chlorophyllous, is placed towards the centre of the seed. There is a layer of perisperm between the outer edge of the embryo and the testa inner surface located symmetrically on both sides, and a tegmen between the testa and the perisperm (Fig. 1c and d; Martin, 1946; Shepherd et al., 2005; Sukhorukov et al., 2018). The embryo almost completely occupies the internal area of the seed. The storage material is mostly nucellus-derived perisperm, consisting of "thinwalled, non-living cells densely packed with starch grains" (Shepherd et al., 2005).

Fresh seeds of S. maritima are dormant and require cold-wet stratification for a period of up to 166 20 weeks to break dormancy (Wetson et al., 2008). Following the protocol of Wetson et al., 167 (2008), seeds were submerged in full-strength ASW (1.0 ASW; containing 411 mM NaCl, 168 53.52 mM MgCl<sub>2</sub>, 10.16 mM CaCl<sub>2</sub>, 9 mM KCl and 28 mM Na<sub>2</sub>SO<sub>4</sub> (Harvey, 1966)) at 5 °C, 169 within 12 days of seed collection. Germination was monitored after 9, 14, 16 and 18 weeks of 170 171 submergence on subsets of 10 seeds sown onto one sheet of germination paper (90 mm diameter, Fisherbrand; Fisher Scientific, Loughborough, UK) moistened with 3.5 mL of dH<sub>2</sub>0 in Petri 172 173 dishes at 15/5 °C (12/12h light/dark, with the higher temperature corresponding to a light intensity of 11 W/m<sup>2</sup>), reflecting the temperature of spring germination in the natural habitat 174 175 (Wetson et al. 2008). Based on the data obtained (see Results), 19 weeks of submergence was 176 selected as sufficient time to release dormancy.

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#### 178 2.2. Ion content of fresh and pre-treated seeds

179 Cations and anions were extracted from seeds and analysed by ion chromatography (DIONEX ICS-1100, Waltham, MA, USA). Fresh and pre-treated (submerged in 1.0 ASW at 5 °C for 19 180 181 weeks) seeds were rinsed in dH<sub>2</sub>0 for 10 seconds and then dried at 103 °C for 17 h (ISTA, 2019). For quantification of anions, 0.5 g of dried material was extracted in 50 ml of anion 182 solution (3.5 mM Na<sub>2</sub>CO<sub>3</sub>/NaHCO<sub>3</sub>) under stirring for 20 min and filtered. For quantification 183 of cations, 1 g of dried material was ashed at 550 °C for 5-6 h in a porcelain capsule. The ash 184 185 was then mineralized for 30 min at 100 °C using 1M HCl solution and filtered. Analysis by ion chromatograph used the eluent 20 mM methane-sulfonic acid and ions were quantified using a 186 standard curve. All solvents and reagents were purchased from Panreac (Barcelona, Spain). 187

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189 2.3. Germination

All germination tests were conducted on pre-treated seeds (i.e., non-dormant) as described in 190 section 2.1. Four replicates of 25 were sown onto two layers of germination test paper 191 moistened with 7 mL of each treatment solution. The Petri dishes were placed in plastic bags 192 to prevent evaporation during the test and incubated at 15/5 °C. Seeds were germinated in the 193 solutions dH<sub>2</sub>O (0 ASW) and five different concentrations of ASW ranging from two times 194 195 full-strength ASW (2.0 ASW) to quarter-strength ASW (0.25 ASW). Iso-osmotic solutions of PEG 8000 were also prepared to correspond to each of the ASW solutions (0.25 ASW = -0.643) 196 MPa; 0.5 ASW = -1.277 MPa; 1.0 ASW = -2.614 MPa; 1.5 ASW = -3.954 MPa; 2.0 ASW = -197 198 5.508 MPa) using the equation (Michel, 1983; Eq. 4):

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200  $\Psi = 0.130[PEG]^2 T - 13.7[PEG]^2$ 

201

where  $\Psi$  = osmotic potential and *T* = temperature (10 °C, i.e. the average of 15/5 °C with a 12 h photoperiod). A freezing point osmometer (Roebling Micro-Osmometer Autocal Type 13 Osmometer, Camlab, Cambridge, UK) was used to confirm the osmotic potentials. 7 mL of PEG was also used to moisten two layers of germination test papers, since the ratio of solution volume to germination test paper weight was > 12 (Hardegree and Emmerich, 1990).

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Germination was scored daily as radicle emergence of at least 1 mm. Experiments were 208 209 conducted for up to 30 days where no further germination was observed. All non-germinated 210 seeds were rinsed five times in dH<sub>2</sub>O, blotted dry, and transferred to new germination test 211 papers moistened with 7 mL of dH<sub>2</sub>O and incubated at 15/5 °C. This recovery test was conducted for up to 25 days where no further germination was observed. For any remaining 212 213 non-germinated seeds, a small proportion of the seed coat was removed and seeds were submerged in 1 % 2,3,5-triphenyl tetrazolium chloride for 48 h in the dark (ISTA, 2019) at 214 215 25 °C. Seeds were rinsed for 10 seconds with dH<sub>2</sub>O and cut longitudinally to assess the embryo for staining (viable seeds stain red and non-viable seeds are unstained), as well as to identify if 216 217 seeds were mouldy or empty.

218

#### 219 2.4. Germination data analysis

220 For each treatment, cumulative germination percentages were plotted over time and a 221 Boltzmann curve was fitted. From this, the time to achieve 10 - 90 % germination was 222 calculated. A halotime model was created by regressing the reciprocal of time to achieve each 223 percentile  $(1/t_g)$  against the concentration of ASW or the osmotic potential of PEG (in units of

- ASW, termed ASW equivalent units or ASW<sub>equiv</sub>), according to the equation:
- 225

226  $1/t_g = ([ASW] - [ASW]_{max g}) / \theta_{Halo g}$ 

227

228 where  $[ASW]_{max g}$  is the threshold or base [ASW] of percentile g, above which there is no 229 germination.  $\theta_{Halo g}$  is the hydro time constant which can be calculated from the inverse of the

- 230 slope for percentile g (Seal et al., 2018).
- 231

233

232 A mean of  $\theta_{Halo g}$  was taken to estimate  $\theta_{Halo}$  for the seed population. The statistical difference

234 x-intercepts, representing [ASW]<sub>max g</sub> and [ASW<sub>equiv</sub>]<sub>max g</sub> were estimated for 50 % germination

in  $\theta_{\text{Halo}}$  between ASW and ASW<sub>equiv</sub> was assessed using a Student's test (*t*-test) (P < 0.05). The

235 ([ASW]<sub>max50</sub>; [ASW<sub>equiv</sub>]<sub>max50</sub>).

236

237 2.5. Preparation of seeds for elemental microscopy and temporal analyses

Two contrasting ASW solutions, 1.0 and 1.5 ASW, were selected as imbibition treatments along 238 239 with their corresponding iso-osmotic PEG solutions and  $dH_2O$  (0 ASW) for physiological and biochemical analyses. These dilutions of ASW were chosen to reflect differences in 240 241 germination rate (based on  $t_{50}$ ) while > 80 % germination was still achieved (see Fig. 2a). Prior 242 to the start of all biochemical experiments, non-dormant seeds were imbibed for 20 % and 80 % 243 of the time to achieve at least 80 % germination, hereafter referred to as the early (EI) and late (LI) imbibition time points, respectively, in the five imbibition treatments at 15/5 °C (Table 1). 244 245 In addition, fresh (i.e., dormant) and pre-treated (i.e., non-dormant, to represent the start of 246 imbibition) seeds were also analysed for comparison.

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248 2.5.1. Seed moisture content

Seed moisture content (MC) of five replicates of 5 seeds was measured. Following imbibition,
seeds were rinsed with dH<sub>2</sub>O water for 10 seconds and transferred to an oven at 103 °C for 17
h (ISTA, 2019). MC was calculated gravimetrically and expressed on a fresh weight basis:

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253 MC (%) =  $[(FW - DW) / FW] \times 100$ 

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255 where FW is the fresh weight and DW is the dry weight.

256

257 2.5.2. Glutathione and low-molecular-weight thiols

Seeds were freeze-dried for five days. Four replicates of 30 mg of freeze-dried seeds were 258 ground using a pestle and mortar to a fine powder in liquid nitrogen. The extraction from seed 259 260 powder was carried out on ice in 1 mL of 0.1 M HCl with 30 mg polyvinylpolypyrrolidone 261 (PVPP) and 0.5 % (v/v) Tween. The mixture was centrifuged for 40 min at 4 °C and 15000 g, and the resulting supernatant was divided in two, one part for the determination of total GSH 262 263 content and the other part for the determination of GSSG content, following the protocol of Seal et al. (2010). Glutathione and the intermediates of glutathione metabolism, cysteine (Cys), 264 265 cysteinyl-glycine (Cys-Gly) and  $\gamma$ -glutamyl-cysteinyl ( $\gamma$ -Glu-Cys), were separated by reversed 266 HPLC (Agilent 1260; Cheshire, UK) on a Poroshell 120 EC-C18 column (150 × 3.0 mm i.d., 2.7 µm particle size; Agilent, Cheshire, UK), and detected fluorometrically (excitation: 380 nm; 267 emission: 480 nm) with a gradient elution of 0.25 % (v/v) acetic acid in distilled water at pH 268 3.9/methanol. Standards of these low-molecular-weight thiols (Sigma Aldrich, UK) at different 269 concentrations were prepared to construct calibration curves for quantification. 270

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### 272 2.5.3. Hydrogen peroxide

273 Hydrogen peroxide  $(H_2O_2)$  production was indirectly quantified via the formation of indamine, which is catalysed by the H<sub>2</sub>O<sub>2</sub>-dependent horseradish peroxidase (HRP) using a modified 274 275 method of Visscher et al., (2018). Briefly, three replicates of  $100 \pm 10$  mg FW of seed per treatment were ground in an ice-cold mortar and homogenized in 1 mL of 0.2 M perchloric 276 277 acid. After centrifugation at 13000 g for 15 min at 4 °C, 500 µL of the resulting supernatant was collected and neutralized to pH 7.5 with 1 M and 0.1 M KOH and then centrifuged again 278279 at 1000 g for 5 min at 4 °C. The supernatant obtained was immediately used for spectrophotometric assay using a WPA Lightwave s2000 UV/Vis spectrophotometer. Two 280 281 technical replicates for each biological replicate were measured. Absorbance values at 590 nm were quantified using a standard curve of known concentrations of H<sub>2</sub>O<sub>2</sub> within a 0 - 500 µM 282 283 range.

284

### 285 2.5.4. Electrolyte leakage

286 The electrical conductivity of three replicates of 20 seeds was measured. Seeds were rinsed 287 with dH<sub>2</sub>O for 10 seconds and transferred into 15 mL of dH<sub>2</sub>O. Conductivity was measured 288 using a meter (4520 Jenway; Fisher Scientific, Loughborough, UK) after two minutes (to 289 remove any electrolytes on the seed surface) and then after 5 h at room temperature (20 °C; 290 Marin et al. (2018)). The measurements at 2 min were subtracted from those at 5 h and 291 expressed per 20 seeds ( $\mu$ S).

#### 292

### 293 2.5.5. Spatial localisation of seawater elements within the seed

Seeds were imbibed in 0 ASW, 1.0 ASW and 1.5 ASW for the LI time point only (Table 1).
Seeds were then sectioned (longitudinally and transversely) with a scalpel and freeze-dried.
Sections were coated with 10 nm carbon (Cressington 208; Watford, UK). Sections were
analysed using scanning electron microscopy (FEI Quanta 650 FEG, Thermo Fisher Scientific;
Loughborough, UK) and mapped for elemental concentration (Na, Cl, Mg, Ca, K and S) using
an energy-dispersive X-ray detector (9 kV accelerating voltage, spot size of 2, 1.5h scan time;
Quantax annular flatQUAD, Bruker; Coventry, UK) equipped with ESPIRIT 2.1 software.

#### 302 2.6. Additional data analysis

Differences of means between two groups was performed using a Student's t-test. The 303 normality of residues and the homogeneity of variances were verified using a Shapiro-Wilk 304 305 test (shapiro-test) and Fisher's test (var-test). To compare data for pre-treated seeds with ASW 306 and PEG treatments, and to compare the effect of the iso-osmotic solutions 1.0 ASW and 1.0 307 PEG, and 1.5 ASW and 1.5 PEG, a one-way ANOVA with post-hoc t-test was used. The assumption of normal distribution was verified by analysis of quantile-quantile plots and the 308 309 normality of residues and the homogeneity of variances were verified using Shapiro's test 310 (shapiro-test) and Levene's test (var-test). When the assumption of a normal distribution was 311 not met, data were arcsine-transformed and the analysis repeated. If the assumption of normal 312 distribution was still not met, a Kruskal-Wallis test with post-hoc Dunn's test was used. 313 Statistical analyses were performed in R (version 4.0.3) or Origin 8.6 (Origin Lab Corporation, 314 Northampton, MA, USA).

315

### 316 3. Results

#### 317 3.1. Seed pre-treatment and germination

318 Fresh seeds were dormant and did not germinate at 15/5 °C. Periodic sampling during 319 submergence in 1.0 ASW at 5 °C revealed the following germination: 40 % after 9 weeks, 50 % 320 after 14 weeks, 80 % after 16 weeks and 100 % after 18 weeks of submergence. Since each test 321 was run for at least 7 d, there was a minimum of a one-week lag to obtain the germination 322 results and therefore 19 weeks was selected for the dormancy breaking pre-treatment. 323

324 Concentrations of Na<sup>+</sup> and Cl<sup>-</sup> were the highest among the salt ions detected in fresh seeds and 325 significantly decreased after 19 weeks pre-treatment (Table 2). Concentrations of Mg, Ca and 326 K significantly decreased during pre-treatment except for SO<sub>4</sub><sup>2-</sup> that slightly increased.

327

328 The germination of pre-treated seeds varied with ASW concentration and their corresponding iso-osmotic PEG solutions (Fig. 2a). The percentage of germination was highest in 0.25 PEG 329 and 0.5 PEG at 97 % and 96 % respectively, and was over 88 % in 0 ASW, 0.25 ASW and 0.5 330 ASW. Germination was similarly high in 1.0 ASW (85 %), 1.0 PEG (88 %) and 1.5 ASW 331 332 (82%), three of the imbibition treatments selected for further analyses. In contrast, germination was lower in 1.5 PEG (52 %) and 2.0 ASW (33 %), with no germination occurring at 2.0 PEG. 333 334 Following recovery of non-germinated seeds on water, all treatments reached a total of 86 -100 % germination (Fig. 2b). The few seeds that remained non-germinated after recovery were 335 mostly mouldy, with very few empty or non-viable seeds (Supplementary Table 1). 336 337

Halo time parameters were estimated by regressing germination rate against ASW 338 concentration and iso-osmotic solutions of PEG (calculated as ASW equivalent units, ASW<sub>equiv</sub>) 339 340 for different percentiles (10 - 90 %) of the seed population (Fig. 3). Germination rate decreased linearly with increasing concentration of ASW and ASW<sub>equiv</sub>. Germination in ASW was the 341 slowest ( $\theta_{Halo} = 6.43$  ASW d) but had the highest 50 % maximum threshold of 2.22 x ASW 342 343 ([ASW]<sub>max50</sub>; Fig. 3a). This was in contrast to germination in iso-osmotic solutions of PEG that was associated with the fastest germination ( $\theta_{Halo} = 5.06 \text{ ASW}_{equiv}$  d) but a lower 344 345 [ASW<sub>equiv</sub>]<sub>max50</sub> of 1.76 (Fig. 3b).

346

#### 347 3.2. Seed moisture content and spatial localisation of seawater elements

The MC of fresh seeds was 17 % and pre-treated seeds was 40 % (Fig. 4). Compared to pretreated seeds, MC significantly increased to 45 % in seeds imbibed in 1.0 ASW at EI, and decreased to 35 % MC in seeds imbibed in 1.0 PEG at LI and at both imbibition time points in 1.5 ASW and 1.5 PEG. There were significant differences between ASW and iso-osmotic solutions of PEG, with a 5 % higher MC of seeds imbibed at LI in 1.0 ASW than in 1.0 PEG. At EI, seeds imbibed in 1.5 ASW had a significantly lower MC than in 1.5 PEG, but by LI, MC was the same between these treatments.

355

356 Microscopic analysis of seeds imbibed in 1.5 ASW revealed that Na accumulated in the testa

357 and Cl in the tegmen (Fig. 5b). Moreover, K and Mg were concentrated in the embryo (Fig.

358 5c,e) but Ca and S were more broadly distributed within the seed (Fig. 5d,f). Localisation of

359 Na, Cl and K was also observed in seeds imbibed in 1.0 ASW and to a lesser extent in seeds

360 imbibed in 0 ASW (Supplementary Fig. 1).

361

362 3.3. Temporal changes in glutathione, other LMW thiols and indicators of oxidative stress

Compared to pre-treated seeds, GSH was significantly higher in seeds imbibed in all treatments 363 except 0 ASW at EI. The concentration of GSH was significantly lower in seeds imbibed in 364 ASW than in iso-osmotic solutions of PEG at EI (for example, 2187 and 2492 nmol g<sup>-1</sup> DW in 365 1.5 ASW and 1.5 PEG respectively). However, by LI, concentrations of GSH were significantly 366 higher in seeds imbibed in ASW (for example, 2960 and 2533 in 1.5 ASW and 1.5 PEG 367 respectively). GSSG was present in fresh seeds at 424 nmol g<sup>-1</sup> DW but was not detected in 368 pre-treated seeds. Except for seeds imbibed in 1.0 PEG at EI, seeds imbibed in all other 369 treatments showed an increase in GSSG concentration. This was especially apparent in seeds 370 imbibed in 1.0 PEG at LI, and in 1.5 ASW and 1.5 PEG, where the highest concentration of 371 816 nmol g<sup>-1</sup> DW was detected in seeds imbibed in 1.5 PEG at LI. Comparing seeds imbibed 372 in iso-osmotic solutions of ASW and PEG revealed that the concentration of GSSG was 373 significantly lower in seeds imbibed in ASW than in PEG except in 1.0 ASW/1.0 PEG at EI. 374 375 Consistent with this, the percentage of GSSG of the total glutathione (GSH + GSSG) was lower in seeds imbibed in ASW than in iso-osmotic solutions of PEG (Table 3). The highest 376 377 percentage of GSSG was found in seeds imbibed in 1.5 PEG (24.4 % at LI), over four-times higher than present in seeds imbibed in iso-osmotic 1.5 ASW (5.7% at LI). 378

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The low molecular weight thiols Cys, Cys-Gly and  $\gamma$ -Glu-Cys were also detected in seeds. Cys 380 381 was the second most abundant thiol present after GSH, with up to 469 nmol gDW<sup>-1</sup>. The concentration of Cys was significantly higher than in pre-treated seeds at the start of 382 383 germination in nearly all of the imbibition treatments (Fig. 6b). However, Cys, Cys-Gly and  $\gamma$ -Glu-Cys did not change in such a predictable pattern as GSH or GSSG during imbibition and 384 385 with treatment (Fig. 6 b,c,d). Where differences were observed in seeds imbibed in iso-osmotic solutions of ASW and PEG, reduced thiol concentrations were typically higher in seeds 386 imbibed in ASW and the concentration and percentage of thiol disulphides were higher in seeds 387 imbibed in PEG (Table 3). Thiol disulphides were not detected in seeds imbibed in every 388 389 treatment, particularly for Cys-bis-Gly (Table 3).

390

391 The highest concentration of  $H_2O_2$  was 2.4  $\mu$ M g DW<sup>-1</sup> in fresh seeds (Table 4). Compared to 392 pre-treated seeds, there was a significant production of  $H_2O_2$  only in seeds imbibed in 1.5 ASW, 393 however values never exceeded 2.15  $\mu$ M g DW<sup>-1</sup>. 394

395 The electrical conductivity was assessed to measure the leakage of solutes from seeds (Table 396 4). Fresh seeds had the highest conductivity of 40  $\mu$ S and was lower at less than 12.46  $\mu$ S in all 397 other imbibition treatments. The conductivity of seeds did not significantly change from pre-398 treated seeds in any of the imbibition treatments, but was higher in seeds imbibed in 1.5 ASW 399 than in 1.5 PEG.

400

#### 401 4. Discussion

402 Saltmarshes are a hostile environment for any plant to survive, as salinity imposes a plethora 403 of osmotic and toxic effects on plant growth and development (Flowers and Colmer, 2008, 404 Munns and Tester 2008). Whereas the different strategies for plants to tolerate these conditions 405 are reasonably well identified, such as the accumulation of salt ions in leaf vacuoles to aid osmotic adjustment and a potentiated antioxidant system to quench the excessive formation of 406 407 ROS (Flowers and Colmer, 2008; Kranner and Seal, 2013), the functional traits underlying salt 408 tolerance in seeds are less resolved. Here, evidence is presented for ion accumulation and a 409 reduced cellular redox state in response to early signs of oxidative stress, in freshly matured seeds of S. maritima that over-winter in cold-saline conditions, and elemental localisation, the 410 411 maintenance of a reduced cellular redox state and beneficial effects of seawater ions during 412 germination.

413

#### 414 4.1 Tolerance strategies of fresh seeds and seeds over-wintering in cold-saline conditions

415 The seeds of saltmarsh halophytes in the northern hemisphere are often shed in the autumn and 416 require an over-wintering period of cold-wet conditions for dormancy to be released, so that 417 germination can begin in the spring when temperatures start to rise but the concentration of salinity is low (Ungar, 1978, Wetson et al., 2008). The conditions experienced during such 418 419 periods not only impact on the timing of these events (Wetson et al. 2008), but exposure to 420 potentially toxic ions may have permanent cellular damage. In dormant seeds of S. maritima collected fresh from the field, Na<sup>+</sup> and Cl<sup>-</sup>, the most abundant ions in seawater, were prevalent 421 along with a mixture of other ions such as K<sup>+</sup>. When seeds were transferred to water to test for 422 423 electrolyte leakage, the data suggests that these ions readily moved out of the seeds down an osmotic gradient. Their presence was accompanied by biochemical changes. In mature, dry 424 425 seeds, a high proportion of total glutathione is present as GSSG (e.g., 24 % GSSG in pea seeds), which typically accumulates during seed maturation and is reduced to GSH within the first 426 hours of imbibition (Colville and Kranner, 2010; Gerna et al., 2017). Although fresh seeds of 427

S. maritima are desiccation tolerant (RBG Kew, 2021), and therefore undergo maturation 428 drying, the saltmarsh environment is not dry and seed MC reflected this at 17% MC. 429 430 Nonetheless, 13.6 % GSSG was present, and these seeds had one of the highest concentrations of GSH, Cys-Gly and  $\gamma$ -Glu-Cys of all the treatments. This was also associated with high 431 concentrations of the ROS H<sub>2</sub>O<sub>2</sub>. After 19 weeks of pre-treatment, ion concentrations were 432 significantly lower for all ions except SO<sub>4</sub><sup>2-</sup>, and with this, GSSG was not detected, the 433 434 proportion of other disulphides were < 1 % and H<sub>2</sub>O<sub>2</sub> was barely detectable. This implicates ion accumulation within the seed and oxidative stress before the event of germination, and 435 436 highlights the importance of maintaining a reduced cellular redox state that will likely contribute towards the persistence of seeds to over-wintering in cold-saline field conditions. 437

438

### 439 4.2 The beneficial effects of seawater ions on germination

Non-dormant seeds of S. maritima germinated readily in ASW, with a maximum threshold of 440 441 2.22 times full strength ASW. This is consistent with a previous study where the maximum 442 germination threshold was around 1M NaCl (where there is 411 mM NaCl in 1.0 ASW) at 443 temperatures between 5 and 15 °C (Seal et al., 2018). This ranks seeds of S. maritima among some of the most tolerant to salinity during germination, similar to other coastal halophytes 444 445 such as Suaeda japonica, Spartina alterniflora and Arthrocnemum indicum (Gul et al., 2013). 446 However, caution is needed when making inter-specific comparisons since temperature can 447 have a large effect on tolerance to salinity during germination (Allen et al., 2000; Zhang et al., 2012; Seal et al., 2018; Seal and Dantas, 2020). In temperate habitats, seeds of S. maritima 448 449 germinate during the spring when temperatures are mild (15/5 °C) and salinity levels are low 450 (Wetson et al., 2008). This is further reflected in the near full recovery of non-germinated seeds of S. maritima in water once salinity is removed, which is typical of many halophyte species 451 (Gul et al., 2013). Thus, although seeds of S. maritima show a broad physiological tolerance to 452 453 salinity ex situ, germination timing in nature will be governed by a combination of environmental factors. 454

455

The localisation of Na and Cl into testa and tegmen, respectively, suggests that these tissues formed a physical barrier to protect the embryo against their accumulation and toxicity. This has also been observed in the testa for Na<sup>+</sup> and Cl<sup>-</sup> in the halophytes *Salicornia pacifica* var. *utahensis* and *Atriplex canescens* (Khan et al., 1985) and for Na<sup>+</sup> *Suaeda physophora* and *Haloxylon ammodendron* (Song et al., 2005). During germination, the accumulation of Na<sup>+</sup> within the seed has been linked to the maintenance of seed fresh weight in several halophyte

species (Song et al., 2005) and the maintenance of MC in seeds of S. maritima (Seal et al., 462 2018), suggesting a role in osmotic adjustment. Although we observed slight, yet significant, 463 differences in MC (no more than 5%) in seeds imbibed in 1.0 ASW compared to those imbibed 464 in 1.0 PEG, this was not reflected in the final germination percentage (85 and 88 % 465 respectively). However, the MC of seeds imbibed in 1.5 ASW and 1.5 PEG was the same by 466 467 LI, at 5 % lower than in pre-treated seeds, yet final germination was higher in seeds imbibed in ASW (82 %) than in PEG (52 %). As germination will only complete (i.e., the emergence of 468 the radicle) above a critical water threshold (Seal et al., 2018), it can be assumed that enough 469 470 water was available within the seed to achieve this in the 1.5 ASW and 1.5 PEG imbibition 471 treatments.

472

Na and Cl were not the only elements to be detected within the seed. In addition, K and Mg 473 localised in the embryo. There are mixed reports as to whether  $K^+$  and  $Mg^{2+}$  promote 474 germination in salinity, although both can be more toxic than Na<sup>+</sup> (Manzoor et al., 2017; Tobe 475 476 et al., 2002), and their beneficial effects on germination may only be seen when in combination with Ca<sup>2+</sup> (Tobe et al., 2002; Gao et al., 2018), which appeared to have a broad distribution 477 across the seed. Nonetheless, both  $K^+$  and  $Mg^{2+}$  and are considered plant macronutrients.  $K^+$ , 478 479 fundamental to plant growth and metabolism, may experience efflux from the cytosol through ROS mediated channels in response to NaCl, making cytosolic K<sup>+</sup> retention an important trait 480 of salt tolerance in plants (Wu et al., 2018). The localisation of K in the embryo could also be 481 an important trait of salt tolerance in seeds. Mg<sup>2+</sup> has multiple roles in plants especially in 482 relation to photosynthetic carbon fixation (Hauer-Jákli et al., 2019), which could be important 483 for S. maritima since the embryo is chlorophyllous.  $Mg^{2+}$  deficiency has also been linked to a 484 lower GSH:GSSG ratio (i.e., more oxidised state) in leaves of Arabidopsis, as impairment of 485 CO2 fixation causing ROS production (Hermans et al., 2010), implicating an association 486 between  $Mg^{2+}$  and the cellular redox state. 487

488

489 During germination, evidence is presented for the synthesis of GSH and GSSG in seeds 490 imbibed in nearly all of the treatments. The accumulation of GSH was slower in seeds imbibed 491 in ASW than in PEG, so that although seeds imbibed in PEG had a higher concentration than 492 in seeds imbibed in ASW at EI, this was reversed by LI. Moreover, the concentration and 493 percentage of GSSG was higher in seeds imbibed in PEG than in ASW at LI. Thus, it appears 494 that seawater ions had a beneficial effect on germination that was not osmotic. This was further 495 apparent when halotime parameters were considered. Although  $\theta_{\text{Halo}}$  was higher (i.e., slower 496 germination) in the presence of ASW than PEG, the maximum threshold to germination was 497 also higher in ASW ([ASW]<sub>max50</sub> = 2.22) than PEG ([ASW<sub>equiv</sub>]<sub>max50</sub> = 1.76). Slower 498 germination may give more time to initiate tolerance mechanisms in response to ASW that 499 promote germination.

500

501 During GSH synthesis, the bond formation between GSH and Cys is the rate-limiting step, and may explain why the intermediates Cys-Gly and  $\gamma$ -Glu-Cys were not abundant during 502 503 germination (Zagorchev et al., 2012). In contrast, Cys was abundant in seeds. Cys has been 504 associated with stress tolerance, including as an osmoprotectant acting as a sulphur donor for the amino-acid methionine (Hesse et al., 2004) and as a ROS scavenger (Soares et al., 2019). 505 In addition, CySS can form complexes with alkali-metals such as Na<sup>+</sup>, suggesting a possible 506 role in Na<sup>+</sup> ion sequestration (Zagorchev et al., 2012), which could explain why CySS was 507 more abundant in seeds imbibed with PEG than ASW. 508

509

Even though there was a partial conversion of reduced thiols to their disulphides, especially in 510 511 seeds imbibed in 1.0 PEG, 1.5 PEG and 1.5 ASW (of up to 25% for GSSG, 13% for CySS, 25 % for Cys-bis-Gly and 27 % for bis-Glu-Cys), the cellular redox state remained highly 512 513 reduced. This is in contrast to seedlings of S. maritima germinated in NaCl that contained about 514 50 % GSSG at ca. 35 % germination (Boestfleisch et al., 2014) and NaCl-stressed cells of Dactylis glomerata with 40 % GSSG during cell differentiation (Zagorchev et al., 2012). It is 515 likely that our treatments selected for biochemical analyses were not "stressful" to reflect a 516 517 period where the antioxidant mechanisms were insufficient to cope with any stress, characterised by a major shift towards an oxidised cellular redox state (Kranner et al., 2006), 518 519 as seeds subjected to all treatments achieved at least 80 % germination. There was also no obvious indication of damage to cellular membranes during imbibition, with higher electrolyte 520 521 leakage from seeds imbibed in 1.5 ASW than in 1.5 PEG most likely reflecting the movement 522 of salt ions out of the seed down an osmotic gradient, since electrolyte leakage was tested in 523 water. Nonetheless, differences in glutathione in seeds imbibed in PEG and in ASW were 524 evident during germination. In contrast, there was no consistent evidence of involvement of H<sub>2</sub>O<sub>2</sub> during germination, unlike H<sub>2</sub>O<sub>2</sub> that was observed to increase during the germination of 525 the subtropical halophytes Suaeda fruticosa and Limonium stocksii in NaCl (Hameed et al., 526 527 2014).

528

529 In conclusion, seeds of S. maritima were highly adapted to saline conditions, through a

combination of strategies involving ion accumulation and localisation, a potentiated 530 glutathione and low-molecular weight antioxidant system to prevent oxidative stress and the 531 532 maintenance of seed moisture content to levels sufficient for germination to complete. This culminated in a higher  $\theta_{\text{Halo}}$  in seeds imbibed in ASW than in iso-osmotic solutions of PEG, 533 that was tolerant to over twice the concentration of ASW. The localisation of potentially toxic 534 535 Na and Cl away from the embryo may be paramount to achieving germination under salinity, yet it remains to be explored whether it is a common feature of all halophyte species or specific 536 537 to certain habitats or lineages.

538

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- 545 None.
- 546

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550

# 551 Author Contributions

552 CES and CMS conceived the study and designed the experiments. CMS conducted the 553 experiments and statistical analyses. MY carried out the seed cleaning/pre-treatment and 554 provided training of some biochemical analyses. AM conducted the ion analysis. IC and TS 555 conducted the scanning electron microscopy with elemental analysis. CMS, CES and LC 556 drafted the manuscript, and all authors contributed towards critically revising the manuscript 557 and give their final approval.

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and the glycophyte, *Digitaria sanguinalis*. S. African J. Bot. 78, 203–210.

683 Figure legends

684

Fig. 1. *Suaeda maritima* seed morphology and anatomy. (a) and (b) External views of the seed;(c) longitudinal and (d) cross-section of the seed. Scale bars represent 0.50 mm.

687

Fig. 2. (a) Cumulative germination of pre-treated seeds in solutions of ASW and iso-osmotic
PEG at 15/5 °C. Data points are mean values of four replicates of 25 seeds, and the lines are
the fitted Boltzmann curves. (b) Recovery of non-germinated seeds from Fig. 2a at the end of

691 the germination test. Seeds were recovered on  $dH_2O$  and incubated at 15/5 °C.

692

693 **Fig. 3.** Germination rate  $(1/t_g)$  in solutions of (a) ASW and (b) iso-osmotic PEG (calculated as 694 ASW equivalent units, ASW<sub>equiv</sub>). Values are the mean halo time  $(\theta_{\text{Halo}}) \pm$  standard deviation 695 for the population, and the maximum concentration of ASW and ASW<sub>equiv</sub> for 50 % 696 germination ([ASW]<sub>max50</sub>; [ASW<sub>equiv</sub>]<sub>max50</sub>). Lower case letters show significantly different 697 values using a Student's test (P < 0.05).

698

**Fig. 4.** Seed moisture content of fresh (F) and pre-treated (PT) seeds exposed to different treatments during imbibition in ASW and PEG (PT = start of imbibition, EI = early imbibition, II = late imbibition; see Table 1 for details). Values are the means of five replicates ± standard deviation. \* indicates significant differences from pre-treated seeds; # indicates significant differences between corresponding EI or LI of 1.0 ASW and 1.0 PEG, and † of 1.5 ASW and 1.5 PEG (one-way ANOVA, F(11,44) = 60.88, P < 0.0001; post-hoc t-test at P < 0.05).

705

Fig. 5. (a) Scanning electron micrograph of a seed, showing localisation of (b) Na and Cl, (c)
Mg and Cl, (d) Ca and Cl, (e) K and Cl, and (f) S, O, Na and Cl. Seeds were imbibed in 1.5
ASW for LI time point. Scale bars represent 50 µm.

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**Fig. 6.** Concentration of low molecular weight thiols of fresh (F) and pre-treated (PT) seeds imbibed in different concentrations of ASW and PEG (PT = start of imbibition, EI = early imbibition, LI = late imbibition; see Table 1 for details). (a) Concentration of glutathione (GSH) and glutathione disulphide (GSSG); (b) cysteine (Cys) and cystine (CySS); (c) cysteinylglycine (Cys-Gly) and cystinyl-bis-glycine (Cys-bis-Gly); and (d) γ-glutamyl-cysteinyl (γ-Glu-715 Cys) and bis-γ-glutamyl-cystine (bis-γ-Glu-Cys). Data are mean values of four replicates ± 716 standard deviation. \* indicates significant differences from pre-treated seeds; # indicates

- 717 significant differences between corresponding EI or LI of 1.0 ASW and 1.0 PEG, and † of 1.5
- 718 ASW and 1.5 PEG (one-way ANOVA with post-hoc t-test or Kruskal-Wallis test with post-hoc
- 719 Dunn's test; P < 0.05; see Supplementary Table 2).