1	Cold-induced changes affect	surviva	l after exposure to vitrification solution during	
2	cryopreservation in the south	1-west A	ustralian Mediterranean climate species Lomandra sonderi	
3	(Asparagaceae).			
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35 Abstract

There is limited knowledge of the effects of exposure to low temperatures in the unique Mediterranean 36 37 climate plant species of Western Australia. We have thus investigated the effect of low temperature on cryogenic tolerance in Lomandra sonderi, an endemic perennial species of southwest Western Australia. 38 Lomandra sonderi plants were preconditioned in tissue culture at constant 23 °C (12h light/dark cycle) or 39 alternating 20/-1 °C (16h light and 8h dark cycle). Shoot tips from both conditions were analysed for their 40 phospholipid, sterol and soluble sugar compositions. Shoot tips were also cryoexposed via a droplet-41 vitrification protocol. Survival in both preconditioning regimes for cryoexposed and non-cryoexposed 42 samples was the same, but plants from the 20/-1 °C regime displayed an improved tolerance to the overall 43 44 cryopreservation process in both cryoexposed and non-cryoexposed samples, thereby eliminating 45 exposure to liquid nitrogen as a primary cause of reduced post-cryogenic viability. Preconditioning of in *vitro* shoots of *L. sonderi* at 20/-1 °C induced significant increases in phosphatidylcholine (from $7.30 \pm$ 46 3.46 to 22.2 ± 7.80 ng mg⁻¹ FW) and increases in several soluble sugars (fructose, galactose, glucose, 47 sucrose) compared to shoots incubated at 23 °C - changes consistent with known cold acclimation 48 responses in plant species generally - but sterol content remained largely unchanged. Analysis of 49 electrolyte leakage in shoot tips from both preconditioning regimes generated a significantly lower LT₅₀ 50 value in the 20/-1 °C samples (-5.45 \pm 0.53 °C) over the 23 °C samples (-2.5 \pm 0.08 °C). Increased 51 tolerance to cryoexposure in L. sonderi appears to lie mainly with acclimation-induced changes in 52 membrane composition and promotion of membrane stability and hence increased resistance to freeze 53 54 damage.

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56 Keywords

57	Lomandra sonderi, phospholipid, sugar, cold acclimation, electrolyte leakage, cryopreservation
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69 Introduction

Continuous maintenance of valuable germplasm material of propagation-recalcitrant or rare and 70 71 endangered plants requires considerable resource allocation on the part of conservation and restoration 72 practitioners. Such material requires reliable long-term storage methods while awaiting utilisation in 73 conservation and restoration projects. Plants exhibiting low seed quality, or poor seed germination due to complex seed dormancy are often propagated via in vitro cultures of plantlets at considerable expense. 74 Consequently, the ongoing protection of each genetic line in culture is paramount. The high level of 75 76 manual input involved in a continuous micropropagation enterprise is prone to accidental contamination 77 through operator error and/or equipment failure, while other less common (but potentially catastrophic) 78 hazards such as fire may result in losses of irreplaceable material (Offord et al. 2009). Cryopreservation is 79 an efficient and ultimately safer alternative for the long-term storage and biosecurity of valuable culture 80 lines of plants (coupled with off-site storage). Agricultural, horticultural and endangered plants with low-81 seed production, complex dormancy or desiccation-intolerance properties that cannot otherwise be stored 82 in conventional seed banks can all benefit from cryogenic storage technology (Kaczmarczyk et al. 2011; Kaczmarczyk et al. 2012; Food and Agriculture Organisation of the United Nations [FAO] 2013). 83

Cryopreservation involves the storage of a range of plant tissues (meristems, excised zygotic 84 85 embryos, cell suspensions or callus tissues) at ultra-low temperatures in liquid nitrogen (LN) either at -135 °C in the vapour phase or -196 °C in the liquid phase (Kaczmarczyk et al. 2012). This process halts 86 all cellular metabolic processes and preserves viability over extended periods of time (Kaczmarczyk et al. 87 88 2012). However, cryopreservation is still limited by a lack of knowledge of how different plant species 89 respond to the cryoprotective agents (CPAs) or any other pre-cryopreservation treatment (Kaczmarczyk et al. 2012). Each step in the cryopreservation process can potentially impact on the survival of the 90 cryopreserved material for better or worse, depending on the cryo-capability of each species, particularly 91 92 as regards desiccation tolerance and ability to tolerate CPA toxicity (Arakawa et al. 1990; Sakai et al. 93 1990; Nishizawa et al. 1993).

94 The cell membrane is the primary site of cryo-injury when plant tissues are exposed to sub zero 95 temperatures (Steponkus 1984). In addition to intracellular ice formation and the mechanical damage 96 caused by this process, expansion-induced lysis, loss of osmotic responsiveness, membrane fusion and altered osmotic behaviour are other types of low temperature injury related to the cell membrane 97 (Steponkus 1984; Wolfe and Bryant 1999). The preconditioning of plants by exposure to low non-98 99 freezing temperatures has been shown to improve survival following cryostorage in various plant species, with studies focusing on agriculturally important species or those native to subtropical or temperate 100 climates, regardless of their ability to cold acclimate under natural conditions (Chang et al. 2000; 101 102 Leunufna and Keller 2005; Kaczmarczyk et al. 2008; Kushnarenko et al. 2009). Additionally, sub-lethal

exposure to low temperature has been shown to cause changes in the chemical composition of cell
membranes that coincides with improved resistance to cryo-injury (Steponkus 1984; Palta et al. 1993;
Uemura et al. 1995). However, the cold acclimation ability of endemic Australian plants, such as those
from the hot-summer Mediterranean climate regions of southwest Western Australia, has seldom been
investigated to the same extent (Funnekotter et al. 2013). Plant species in this region of Australia can
indeed be difficult to cryopreserve successfully (Kaczmarczyk et al. 2013).

109 A commonly used method of determining the relative stability or permeability state of the cell membrane is by the measurement of electrolyte leakage as a consequence of low temperature cellular 110 damage (Campos et al. 2003; Mancuso et al. 2004). Low, lethal temperatures cause a change in the semi-111 112 permeable nature of the cell membrane, which results in a large, irreversible loss of electrolytes from plant tissues (Campos et al. 2003; Mancuso et al. 2004). Plants able to acclimate via changes that stabilize 113 114 the cell membrane, can therefore tolerate lower temperatures by making their cell membranes "less 115 leaky", a response that can be easily measured and quantified using a conductivity probe and meter (Maier et al. 1994; Campos et al. 2003). The cell membrane typically consists of a lipid bilayer made up 116 primarily of phospholipids (PLs). The type of PL headgroup and fatty acid (diglyceride) chains, as well as 117 the level of unsaturation in these chains affect the stability, fluidity and permeability of the membrane 118 (van Meer et al. 2008). Major PL headgroup types found in plants include phosphatidylcholine (PC), 119 phosphatidylserine (PS), phosphatidylglycerol (PG), phosphatidylinositol (PI), phosphatidylethanolamine 120 (PE) and phosphatidic acid (PA) (Uemura and Steponkus 1994; Uemura et al. 1995). Cold acclimation 121 has shown to increase the concentration of PE and PC with unsaturated fatty acids as well as increased 122 123 total PL levels, which reduce the occurrence of deleterious membrane phase transitions and improve freeze tolerance (Uemura and Steponkus 1994; Uemura et al. 1995). 124

125 Sterols or steroid alcohols are also an integral component of cell membranes that restrict the 126 motion of the PL fatty acyl chains and thus control the fluidity and mobility of the membrane (Hartmann 127 1998; Dufourc 2008); this also reduces the likelihood of phase separation occurring in the PLs (Dufourc 128 2008). An increase in membrane sterol concentrations can also bring PL molecules closer together, thus 129 reducing the permeability of the membranes (Demel and De Kruyff 1976). The concentration of different 130 types of sterols also has an effect on cell membranes. β -sitosterol has a greater effect on membrane permeability and ordering than stigmasterol (Schuler et al. 1991; Hartmann 1998). Increased chill 131 tolerance in banana meristems and mung beans has been correlated to an increase in the β -sitosterol to 132 stigmasterol ratio (Guye 1989; Zhu et al. 2006). Intracellular sugars or sugar alcohols also have a 133 stabilising effect on cell membranes (Crowe et al. 1988; Wolfe and Bryant 1999; Turner et al. 2001). 134 Soluble sugars reduce the gel phase to fluid phase transition temperature (T_m) during dehydration of 135 cellular membranes (Wolfe and Bryant 1999; Koster et al. 2000; Lenné et al. 2010). These are known to 136

occur *via* osmotic and volumetric effects of the sugars on membranes (Lenné et al. 2010). In addition,
high intracellular sugar concentrations promote a vitrified state of water during cooling due to the highly
viscous nature of their aqueous solutions (Wolfe and Bryant 1999).

Using the south-west Australian endemic Lomandra sonderi as a model species (Menon et al. 140 2012) the overall aim of this study was to understand how low temperature preconditioning treatment 141 142 might induce cold acclimation and potentially improve post-cryogenic survival in a Mediterranean species 143 that might be expected to have a limited capacity to respond to low temperature acclimation. In this respect, changes to the temperature at which freeze damage occurs as well as the overall membrane PL 144 and sterol and intracellular sugar compositions were determined during this study. These attributes were 145 146 selected as indicators of cold acclimation in response to low temperature preconditioning due to their role 147 in membrane stability and freeze damage mitigation. We discuss how the changes in composition 148 observed relate to changes in post-cryogenic survival.

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150 Methods

151 Plant material

152 In vitro plants of Lomandra sonderi were made available from the tissue culture collection at Kings Park and Botanic Garden, Western Australia (originally sourced from BHP Billiton Worsley 153 Alumina Pty. Ltd., Boddington, Western Australia). Shoots of L. sonderii were maintained on a basal 154 tissue culture medium (BM), which consisted of half strength Murashige and Skoog (MS; Murashige and 155 Skoog 1962) macro- and micro-elements with 500 µM myo-inositol, 500 µM 4-morpholineethanesulfonic 156 157 acid, 100 µM NaFeEDTA, 4 µM niacin, 3 µM thiamine-HCl, 2.5 µM pyridoxine-HCl, 0.06 M sucrose, 0.8% (w/v) agar and pH adjusted to 6.5 prior to autoclaving (20 min at 121 °C). BM with 0.2 µM 6-158 benzylaminopurine (BAP) was used for shoot maintenance and multiplication, and as a pre-culture 159 160 treatment just prior to cryopreservation. Heat labile plant hormones including gibberellic acid (GA₃) and 161 zeatin (Z) [6-(4-hydroxy-3-methylbut-2-enylamino) purine] were filter-sterilised and added to BM 162 following autoclaving to prepare recovery medium (RM). Maintenance medium (BM \pm 0.2 μ M BAP) was 163 dispensed into 250 ml polycarbonate containers (~40-50ml per container) with polypropylene lids (each with 1 x 9 mm diam. vent covered with 0.2 micron micropore filter) then autoclaved; while RM was 164 autoclaved in 500 ml media bottles and dispensed into sterile 55 mm diam. Petri plates at \sim 10 ml per 165 plate. All media plate pouring and manipulations of in vitro plant material (for maintenance and 166 cryopreservation) were carried out under sterile conditions in a laminar flow cabinet. 167

168

169 *Temperature preconditioning*

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L. sonderi plants were preconditioned in vitro on BM (+ 0.2 µM BAP) for 3-week incubation

171 periods using two temperature regimes: 20/-1 °C alternating (20°C during 16 h light period and -1°C 172 during 8 h dark period) and constant 23 °C with a 12 h photoperiod, prior to any cryoexposure 173 experiments. Preconditioning treatments were carried out in temperature controlled incubators with 174 lighting supplied by 36 W cool white fluorescent tubes (Philips Alto; photon flux density of ~30 μ mol m⁻² 175 s⁻¹).

176

177 Cryoexposure

Lomandra sonderi shoot tips were extracted from in vitro grown shoots and cryostored using the 178 method previously developed by Menon et al. (2012). Shoot tips from the two preconditioning regimes 179 180 (20/-1 °C and 23 °C) were transferred onto BM containing 0.2 µM BAP for 48 h as a pre-culture step, prior to loading treatment. Shoot tips were placed in ~ 10 ml of a loading solution (LS) comprised of 2 M 181 182 glycerol and 0.4 M sucrose in liquid BM (as described above, without agar) in a 55 mm sterile plate for 183 20 min, then treated in PVS2 [30% (w/v) glycerol, 15% (w/v) ethylene glycol, 15% (w/v) dimethyl sulfoxide (DMSO) in liquid BM with 0.4 M sucrose; (Sakai et al. 1990)] for 10 min at 23 °C. Each shoot 184 tip was individually transferred to a 1 µL droplet of PVS2 on a sterilised aluminium foil strip (~3 mm 185 wide x 20 mm long; 5 shoot tips per strip). Two foil strips were placed in an empty, pre-chilled cryovial, 186 187 plunged into LN and stored for a minimum of one hour.

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189 *Rewarming and recovery*

Cryovials containing L. sonderi shoot tips were rewarmed by immersion in a 40 °C water bath for 190 191 10 seconds. The foil strips with shoot tips were then quickly removed from vials and shoot tips released into washing solution (1M sucrose in liquid BM) and incubated for 20 min at $\sim 23^{\circ}$ C. Control shoot tips 192 were transferred directly after PVS2 treatment to this washing solution (no immersion in LN). All shoot 193 194 tips were then placed onto a recovery medium (solid BM containing 1.0 μ M Z and 0.5 μ M GA₃) for two weeks in darkness at ~25 °C, and then transferred to 23 °C/12 h photoperiod (as previously described). 195 196 The survival of shoot tips on RM was determined using visual indicators 6 weeks after the date of 197 rewarming. Green shoot tips showing leaf growth and increase in size were considered to be alive. Other 198 shoot tips that did not show any signs of greening or growth after six weeks were considered non-viable. 199 Percentage survival rates of the shoot tips were then determined.

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201 Electrolyte leakage and LT₅₀ calculation following staged cooling

Approximately 2 g of *L. sonderi* leaf material from the two temperature preconditioning regimes were excised per sample. The material per sample was placed in a glass test tube, and 200 μ L of deionised water was added to it. A total of 8 samples per preconditioning regime were placed into a cooling bath

205	(Model 9512 Refrigerating/Heating Circulator, PolySciences, Inc.), preset at -1 °C. The control samples			
206	were placed directly on ice. After one hour at -1 °C, the temperature of the cooling bath was gradually			
207	reduced at the rate of -1.5 °C h ⁻¹ to -10 °C, with samples taken out every hour and stored on ice (8 hour			
208	procedure). 500 μ L of deionised water was added to each sample before storage overnight at 4 °C. After			
209	24 h, 5 mL of deionised water was added to each sample before transferring to a shaker at 100 rpm			
210	overnight. 1 mL of each sample was transferred to 10 mL of deionised water and the conductivity of the			
211	diluted solution was measured (C1) using a 900-C Conductivity Meter (TPS Australia). The samples were			
212	boiled at 100 °C for 30 min. 1 mL of each boiled sample was then transferred to 10 mL of deionised			
213	water and the conductivity of the diluted solution was measured (C2). The electrolyte leakage during			
214	cooling was calculated as a percentage of total electrolyte concentration via the following formula:			
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216	% loss of electrolytes = $(C_1/C_2) * 100$			
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218	To account for the loss of electrolytes for reasons other than freeze damage, this value was then			
219	zero corrected via the following formula:			
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221	Zero correction = % leakage of sample - % leakage of control			
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223	These values were then normalized to the highest % electrolyte leakage value, to prevent			
224	percentage values above 100%, using the following formula:			
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226	Normalized value = ([zero correction]/[highest % leakage value]) * 100			
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228	The normalized values of percentage electrolyte leakage were used to construct a sigmoidal curve			
229	from which the LT_{50} was calculated (the temperature at which 50% of total ion leakage occurs) using			
230	GraphPad Prism version 5.03 for Windows, GraphPad Software, San Diego California USA.			
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232	Extraction of sterols and phospholipids			
233	Sterols and phospholipids were analysed as described by Funnekotter et al. (2013). L. sonderi			
234	shoot tips from the two temperature preconditioning regimes were isolated (9-11 mg, \sim 12 shoot tips).			
235	Shoot tips were homogenized in LN and 1 μ g cholestanol and 1 ng 14:0/14:0-PC and 14:0/14:0-PG were			
236	added as sterol and PL internal standards. In order to dissolve the samples, 400 μL of methanol was added			
237	followed by incubation at 75°C for 20 min. 400 μL of chloroform and 400 μL of water were added to			
238	each followed by centrifugation at 10,000 rpm for 3 min. The chloroform phase containing the lipids was			

then extracted. Solid phase extraction using SupelcleanTM LC-Si 1 ml columns (Supelco, Bellefonte,
USA) was performed on the chloroform phase. Neutral lipids were extracted using chloroform,
glycolipids were extracted using acetone and the PLs were extracted using methanol. The neutral lipid
and PL phases were dried under nitrogen gas, and the glycolipid phase was discarded.

243

244 Derivatisation and gas chromatography mass spectrometry (GC/MS) of sterol phase

245 A solution of 10% potassium hydroxide (KOH) dissolved in methanol (500 µl) was used to dissolve the extracted neutral lipid phase. This was followed by incubation at 75°C for one hour and then 246 followed by a triple hexane extraction. Sterols were derivatised by incubation in 20 µL methoxypyridine 247 248 at 65°C with shaking at 750 rpm for one hour and then incubation in 30 µL bis-249 (trimethylsilyl)trifluoroacetamide (BSTFA) at 65°C with shaking for one hour. An Agilent 7890A GC 250 system coupled to an Agilent 5975C inertXL MSD (Agilent Technologies; Palo Alto, CA, USA) was 251 utilised for the identification of sterols. A Varian VF-5MS column (30 m x 250 µm x 0.25 µm; Agilent Technologies) was used for separation of the samples. A 1 µL splitless injection was used to load the 252 derivatised sterol extracts. The pressure of helium gas was 21.28 psi with a flow rate of 2.34 mol min⁻¹ 253 within the column. Temperature of the column was increased from 70 °C to 325 °C. Agilent Technologies 254 255 MSD ChemStation G1701EA E.02.00.493 was used for identification and quantitation of sterol species.

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Liquid chromatography electrospray ionisation tandem mass spectrometry (LC-ESI-MS-MS) of phospholipids

259 100 μ L of 50% v/v acetonitrile (in water) was used to dissolve the PL samples obtained from the solid phase extraction. An Agilent ZORBAX 300 Extended-C18 capillary column attached to an Agilent 260 6340 ion trap LC/MS was used for the separation of PLs. A model 1200 capillary pump system from 261 262 Agilent Technologies (Palo Alto, CA, USA) was used to inject the samples into the column. The method 263 from Funnekotter et al. (2013) modified from Kim et al. (2009) and Min et al. (2010) was used for the 264 runs. The solvents components of the mobile binary phase used were 50% v/v acetonitrile in water (A) 265 and 90:10 (v/v) isopropanol/acetonitrile (B). The solvents contained 0.1% (v/v) formic acid and 0.05% ammonia for positive ion and negative ion detection, respectively. A separation gradient was used to 266 separate the PL species. The gradient equilibrated to 55% solvent B as the column was loaded with the 267 sample. The gradient ramped to 70% solvent B over 40 min and was held at 70% solvent B for an 268 additional 5 min. Re-equilibration occurred via a 10 min post-run. A capillary voltage of 3500 V with a 269 nebulizer pressure of 15 psi and dry nitrogen gas flow of 7 L min⁻¹ at 325 °C was used for electrospray 270 ionisation. The ion scanning range for the tandem MS runs was 170-860 amu and 220-950 amu for the 271 272 positive and negative modes, respectively. The scanning range for the quantitative MS runs was 620-860

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278 Double-bond Index (DBI)

The DBI was calculated for the fatty acyl chains of the PLs detected, with a formula modified from(Quartacci et al. 2001):

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$= \frac{[(1 \times \% \text{monoenes}) + (2 \times \% \text{dienes}) + (3 \times \% \text{trienes}) + (4 \times \% \text{tetraenes} + (5 \times \% \text{pentaenes})]}{\Sigma\% \text{ saturated fatty acids}}$

amu for both the positive and negative modes. Agilent Technologies Data Analysis for 6300 Series Ion

Trap LC/MS v3.4 (Build 175) was used for identification and quantification of PL species. Calibration

standards of 18:0/18:0-PA, 18:0/18:0-PC, 18:0/18:0-PE, 18:0/18:0-PG, 18:1/18:1-PI and 16:0/16:0-PS

(Avanti Polar Lipids, Alabaster, AL, USA) were used to determine the quantities of species identified.

282 283

284 *Extraction of sugars*

DBI

L. sonderi shoot tips from the two temperature preconditioning regimes were isolated (9-11 mg). Shoot tips were homogenised in LN and sugars were extracted in 500 μ L 20:2:0.2 methanol/water/ribitol solution. Ribitol was used as an internal standard with a concentration of 0.2 mg mL⁻¹. Samples were then incubated at 75°C for 20 min. 100 μ L of the samples were then transferred to GC vials and dried in a Speed Vac®.

290

291 Derivatisation and GC/MS of sugars

Derivatisation of sugars was also conducted in methoxypyridine and BSTFA as per the sterols. Sugars were also identified using the Agilent 7890A GC system coupled to the Agilent 5975C inert XL MSD as per the sterols. Flow rate of helium gas in the column was modified to 1 mol min⁻¹ at 8.8085 psi. All other parameters remained unchanged from the sterol runs. Fructose, galactose, sucrose, glucose, sorbitol, trehalose and raffinose (Sigma-Aldrich, St. Louis, MO, USA) were used as calibration standards for the quantification of data. Agilent Technologies MSD ChemStation G1701EA E.02.00.493 was used for identification and quantitation of sugar species.

299

300 *Statistics*

The cryoexposure experiments were performed using six replicates (6 x 15 shoot tips) for each treatment. The cooling bath experiments were performed with three replicates per treatment. The sterol and sugar experiments were performed with three replicates each (three times ~12 shoot tips) per treatment. PL experiments were performed with four replicates (four times ~12 shoot tips) per treatment. The Shapiro-Wilk test was used to check normality of data. No transformations were applied if data was statistically normal. Statistical comparisons were made using independent t-tests for two-condition
comparisons using SigmaPlot for Windows Version 11.0 Build 11.0.0.77 (Systat Software, Inc. Chicago,
IL, USA).

309

310 **Results**

311 Cryoexposure

As previously determined (Menon et al. 2012), there was a significant difference in postcryoexposure survival of shoot tips derived from the two preconditioning temperatures investigated (23 and 20/-1 °C). Plants that had undergone the 20/-1 °C preconditioning had shoot tip survival of ~32.2% for samples not exposed to LN (-LN) and samples exposed to LN (+LN). Samples from the 23 °C preconditioning regime showed only ~14.4% survival for both -LN and +LN samples (Menon et al. 2012). However, within each regime, there was no significant difference between -LN and +LN samples (Menon et al. 2012).

- 319
- **320** *LT*₅₀

Alternating temperature preconditioning of plants resulted in altered electrolyte leakage readings from shoot samples, indicating that a change in cell membrane characteristics had occurred. The mean LT₅₀ value of shoot cultures incubated at 23°C was -2.50 \pm 0.08 °C compared to -5.45 \pm 0.53 °C for shoots incubated at 20/-1 °C.

325

326 Phospholipid analysis

PC and PE accounted for the highest content of PLs in both preconditioning regimes (over 97%). 327 The remaining PL classes (PA, PG, PI and PS) were less abundant, being less than 3% of the total PLs 328 329 detected. There was a significant increase (P < 0.05) in the amount of PC in shoot tips from the 20/-1 °C preconditioning regime compared to those from the 23 °C preconditioning regime. The amounts of all the 330 331 other PLs detected, particularly PE, did not significantly change between temperature treatments (Fig. 1). Due to the large increase in PC content, the total PL content in shoot tips from the 20/-1 °C 332 preconditioning (25.0 ng mg⁻¹ FW) was significantly higher (P < 0.05) than in those from the 23 °C 333 preconditioning (9.8 ng mg⁻¹ FW). See Table S1 in Supporting Information for content of individual PL 334 classes measured. 335

A total of 51 different PL species were identified and quantified (see Table S2 in Supporting Information). The widest range of PL species was detected in PC and PE, with only one type of PA species detected. Table 1 reports the amounts of each individual fatty acyl chain type detected in the PLs. Across both preconditioning regimes, palmitoleic acid (16:1) was the most common fatty acid detected, constituting 20.3% and 18.7% of all fatty acyl chains detected in the 23 and 20/-1 °C samples, respectively (Table 1). The other fatty acyl chains found in high amounts in the 23 °C samples were oleic acid (18:1), linoleic acid (18:2), and linolenic acid (18:3), all of which were present in significantly higher amounts in the 20/-1 °C samples (Table 1). Large, significant increases were also seen in fatty acyl chains 16:0, 18:0, 18:4, 20:2, 20:3 and 20:4 in the 20/-1 °C samples. The amount of C16 chains of the PLs had the largest increase after preconditioning, rising by 333% (Table 1).

Table 2 shows a comparison of the degrees of saturation and unsaturation in fatty acids between preconditioning regimes. There were large, significant increases in both the saturated and the unsaturated monoenes, dienes, trienes and tetraenes in the 20/-1 °C samples (Table 2). There was also an increase in pentaenes but this was not statistically significant. The associated DBIs were computed, revealing a large (50%), significant drop in DBI after preconditioning at 20/-1 °C (9.93 \pm 0.81) compared to the 23 °C regime (20.32 \pm 1.23).

352

353 Sterol Analysis

354 Cholesterol, stigmaterol and β -sitosterol were the three sterols detected. Any other sterol types 355 (such as campesterol) were either present in trace quantities or non-existent. β -sitosterol accounted for the 356 highest content of sterols in both preconditioning regimes. There was no significant difference in the 357 content of any of the three sterol types detected between the two temperature pre-treatments.

The total content of sterols measured for the 23 °C samples $(383.9 \pm 120.6 \text{ ng mg}^{-1} \text{ FW})$ was ~8.5% higher than the 20/-1 °C samples $(353.7 \pm 80.6 \text{ ng mg}^{-1} \text{ FW})$, but was not significantly different. See Table S3 in Supporting Information for content of individual sterol types measured.

361

362 Soluble Sugar Analysis

There were seven types of sugar identified in the shoot tips. Fructose, galactose, sucrose and glucose were the main sugars present (>30 ng mg⁻¹ FW) and their concentrations were significantly different between the two preconditioning regimes (Fig. 2). The other three sugars identified (sorbitol, trehalose and raffinose) were detected in significantly smaller amounts (<10 ng mg⁻¹ FW) (Fig. 2). See Table S4 in Supporting Information for content of individual sugar types measured.

The total sugar contents in the two preconditioning regimes investigated showed large variations (P < 0.001). The total sugar content of the 20/-1 °C samples (310.1 ± 65.3 ng mg⁻¹ FW) was ~15 times higher than that of the 23 °C samples (20.6 ± 8.4 ng mg⁻¹ FW).

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374 Discussion

Overall, the preconditioning treatment resulted in substantial increases in sugar levels and PC content 375 376 of shoot tissues, accompanied by a two-fold improvement in the ability of shoot tips to survive desiccation. In a previous study following preconditioning at 20/-1 °C shoot tips of L. sonderi showed an 377 optimal mean survival of $32.2 \pm 11.5\%$ following cryopreservation (Menon et al. 2012). In comparison, 378 only $14.4 \pm 2.7\%$ of shoot tips derived from the 23 °C preconditioning treatment survived after the same 379 cryogenic treatment (Menon et al. 2012). The mean survival rates of the -LN controls were very similar to 380 the corresponding mean +LN survival rates in shoot tips with the same preconditioning regime (Menon et 381 al. 2012), suggesting that factors other than the cryopreservation protocol used (which was identical apart 382 383 from the LN step) are responsible for the differences in survival observed. As previously determined (Menon et al. 2012), the lack of difference in survival between +LN and -LN samples could be attributed 384 385 to the effective elimination of ice formation through the use of CPAs. Any reduction in survival would 386 have therefore not been directly attributable to the LN immersion step, but rather to other processes such 387 as toxic or desiccation stresses experienced by the plant material during the preparatory steps leading to cryopreservation. To temper these deleterious effects, cold acclimation stimulates plants to develop 388 tolerance mechanisms that enhance their capacity to survive adverse conditions such as desiccation, to 389 which plant tissues are subject during the cryogenic steps leading up to LN immersion (Thomashow 390 1999). These tolerance mechanisms arise primarily via improved stability of cell membranes and 391 accumulation of sugars (Thomashow 1999). Sugar accumulation is one of the mechanisms of desiccation 392 tolerance in 'resurrection plants', such as *Craterostigma plantagineum*, as well as changes to phospholipid 393 metabolism due to differential activation of phospholipase D genes (Bartels and Salamini 2001). Similar 394 families of genes responsible for related processes, such as sugar transporters, sugar synthases and LEA 395 396 proteins are upregulated as a response to both cold and desiccation stresses (Ingram and Bartels 1996; 397 Thomashow 1999; Chinnusamy et al. 2007). It is therefore reasonable to postulate that improving the 398 tolerance of a plant to one kind of stress, i.e. cold stresses, via preconditioning, would also improve 399 tolerance to other stresses such as desiccation.

The LT₅₀ values for the 23 °C and 20/-1 °C preconditioning regimes differed significantly. Similar 400 findings were reported by Mancuso et al. (2004), who investigated the freezing tolerance of the Australian 401 402 genera Grevillea and Callistemon using a similar electrolyte leakage method and determined that plants that were acclimated naturally to cold winter conditions exhibited significantly lower LT₅₀ values. 403 According to the plant hardiness map of Australia, divided into seven zones (Dawson 1991), the two 404 Grevillea species studied by Mancuso et al. (2004; Table 3) grow in Northwest Western Australia (near 405 Shark Bay), i.e. zone 5, with minimum temperatures during winter from 5 °C to 0 °C, and the four 406 Callistemon species grow in Eastern New South Wales, a zone 3 region, with minimum temperatures 407

408 during winter from 0 °C to -5 °C. Lomandra sonderi naturally grows in southwest Western Australia, a zone 3 region, with similar winter conditions to the Callistemon species. Nevertheless, a major difference 409 410 between both studies is that L. sonderi cultures were acclimated in vitro under controlled laboratory conditions intended to simulate natural average winter conditions, as opposed to the acclimation of pot-411 grown plants tested by Mancuso et al. (2004), which were maintained in greenhouses during winter. 412 Regardless of the differences in methodology, Table 3 shows that the LT₅₀ values of non-acclimated and 413 414 acclimated L. sonderi plant material are not greatly different from either Callistemon or Grevillea species, 415 as described by Mancuso et al. (2004).

Cold acclimation is, however, known to be more effective in species that naturally grow in low 416 417 average temperature climates than those from warmer origins (Flint 1972). For instance, the electrolyte leakage analysis of xylem tissue of evergreen Peach (Prunus persica [L.] Batsch) sampled in May (i.e. 418 419 non-acclimated or deacclimated) generated an LT₅₀ value of -3.0 compared to samples taken in January, 420 (i.e. cold acclimated) which had an LT₅₀ value of -17.0 °C (Arora et al. 1992). The plants were grown in 421 Kearneysville, West Virginia, USA, which has an average annual minimum temperature of -17.8 °C to -20.6 °C (USDA 2012). This indicates that although cold acclimation can occur in species from warmer 422 423 Mediterranean climates, the degree of acclimation possible appears to be far lower than in species from much colder climatic regions. Our findings with L. sonderi tend to reflect this difference. 424

The lower LT_{50} values for *L. sonderi* in the 20/-1 °C regime are strong indicators of reduced membrane permeability and improved membrane stability, which may partly explain the improved cryogenic tolerance of shoot tips following a suitable acclimation treatment, as found for other species (Flint et al. 1967; Thomashow 1999; Campos et al. 2003; Mancuso et al. 2004).

PLs are the major component of the plasma membrane (van Meer et al. 2008). The increase 429 measured in the total PL content in the 20/-1 °C preconditioning samples was largely due to the large 430 431 (two-fold) increase in the amount of PC (Fig. 1). Funnekotter et al. (2013) observed a similar trend in the 432 PC content of the Australian species they studied where it rose by 77% in Grevillea scapigera and by 433 65% in Loxocarya cinerea, after preconditioning at 20/10 °C for three weeks. This suggests that 434 preconditioning using low temperature regimes seems to increase PC content to stabilise cell membranes as a mechanism of chill tolerance. It is known that strong hydrating PL classes like PC are more likely to 435 436 retain a lamellar phase in the cell membrane as compared to PL classes like PE, which are less hydrating 437 and tend to form hexagonal II phases instead (Sen and Hui 1988). The total amount of PE, which is substantially lower than that of PC, remained unchanged in the two preconditioning regimes. These 438 findings agree with the model of strongly hydrating PL classes, such as PC, which are more likely to 439 maintain a functional liquid crystalline (lamellar) phase during cooling. Membranes containing higher 440 amounts of PC are therefore more likely to withstand the stresses experienced during the entire 441

cryopreservation process, which is reflected in the increase observed in both cryopreserved (+LN) andcontrol (-LN) shoot tip survival.

444 The gel-to-liquid crystalline transition temperature, i.e. the temperature at which the gel phase of a PL bilayer melts to form a lamellar phase, is affected by the fluidity of the membrane, which in turn 445 446 affects its stability. The strength of attractive van der Waals forces between lipid molecules largely controls the phase behaviour of lipid bilayers. These forces are affected by the length and saturation of 447 fatty acid tails of PLs. Longer acyl chains result in higher transition temperatures and reduced fluidity; 448 conversely, higher degrees of unsaturation result in lower transition temperatures and increased fluidity 449 (Silvius 1982). Interestingly, recent studies have revealed that increases in unsaturation in model cell 450 451 membranes increase resistance to the deleterious effects of CPAs (Hughes et al. 2012; Malajczuk et al. 452 2013) We would, therefore, expect to see a rearrangement in the composition of fatty acyl chains of PLs 453 after 20/-1 °C preconditioning, as the plant attempts to increase the fluidity (and, hence, lower the transition temperature) of its cell membranes as a stress response, either by accumulating more short-454 455 tailed fatty acids or by increasing the degree of unsaturation. Indeed, the largest increase was seen in 16C chains in the PLs detected (increased by 333%), followed by 18C chains (increased by 195%) and then 456 20C chains (increased by 150%; no significant increases were detected in 20:0, 20:1 and 20:5 fatty acyl 457 chains). It would appear that an accumulation of PLs containing shorter fatty acyl chains is favoured 458 during the cold acclimation of L. sonderi. 459

The DBI is a measure of the overall degree of unsaturation in fatty acyl chains. Increases in DBI 460 indicate a higher degree of unsaturation, which in turn mean increased disorder in the PL bilayer structure 461 due to the increase in the number of kinks in the chains, which consequently leads to increased fluidity 462 (and reduced transition temperature) (Silvius 1982; Nishida and Murata 1996). Higher DBI has been 463 related to stress tolerance in other plants (Nishida and Murata 1996). However, in the case of L. sonderi a 464 465 large 50% decrease in DBI was observed after preconditioning at 20/-1 °C (9.93 \pm 0.81) compared to preconditioning at 23 °C (20.32 \pm 1.23). This trend was similarly observed by Funnekotter et al. (2013) in 466 467 Loxocarya cinerea, which had a significantly lower DBI after preconditioning using an alternating 20/10 °C regime, compared to preconditioning using a constant 23 °C regime. Zhu et al. (2006), who utilised a 468 sucrose-pretreatment regime to improve cryoexposure tolerance in different banana varieties, found a 469 significant decrease of DBI in PL fatty acids in the pretreated samples and simultaneously observed 470 471 significantly improved cryogenic survival as a result of this pretreatment. It becomes clear that the loss of fluidity due to the decrease in DBI is counteracted by the accumulation of short, saturated fatty acyl chain 472 PLs, in particular the more strongly hydrating PC, as discussed above. The level of contribution of each 473 factor to membrane fluidity is, however, unknown and merits further research. 474

475 Membrane sterols are known to improve membrane stability via ordering of PLs (Demel and De Kruvff 1976), which can also protect membranes against damage by cryosolvents such as DMSO and 476 polyols (Hughes and Mancera 2013; Hughes et al. 2013). The sterols identified in shoot tips of L. sonderi 477 include stigmasterol, cholesterol and β -sitosterol. There was no statistically significant change measured 478 479 in total sterol content across the two preconditioning regimes. Similarly, O'Neill et al. (1981) and Funnekotter et al. (2013) detected no changes in free sterols in Fragraria virginiana and Arabidopsis 480 481 thaliana, in their respective studies, after cold acclimation. This may not hold true for all plant species, however, as Funnekotter et al. (2013) also found that G. scapigera and L. cinerea showed significantly 482 higher amounts of sterols after preconditioning at 20/10 °C (69% and 88% increases, respectively). This 483 484 suggests that, in the case of L. sonderi, membrane stability and/or permeability is not necessarily 485 regulated exclusively by or reliant on changes in sterol content, at least in response to the temperature 486 range treatments in this study.

487 Sugars are known to have a stabilising effect on cell membranes and promote the vitrification of water at low temperature (Wolfe and Bryant 1999). Preconditioning at 20/-1 °C largely increased the total 488 content of sugar species in shoot tips of L. sonderi compared to the total content of sugars after 23 °C 489 preconditioning. This may be due to the physiological responses to cold stress when exposed to low 490 temperature in the 20/-1 °C preconditioning regimes. Increased intracellular sugar content may be a stress 491 492 response that favours the vitrification of water (and hence the reduction of ice formation). In addition, the increased solute potential within the cells as a consequence of the increased sugar concentration would 493 494 greatly reduce the desiccation effects of the cryopreservation processes. This can be associated with the 495 greater survival rates that were recorded in both the cryopreserved (+LN) and control (-LN) shoot tips exposed to 20/-1 °C preconditioning. The large change in total sugar content was caused primarily by 496 significant increases in glucose, fructose, galactose and sucrose, with minor, insignificant changes in 497 498 trehalose, raffinose and sorbitol (which were detected in relatively small amounts). These changes 499 indicate a robust response to the low temperature preconditioning regimes. This was similarly observed 500 by Funnekotter et al. (2013) in G. scapigera and L. cinerea, which showed increases in total sugar content 501 after preconditioning at 20/10 °C. These observations relate to previous findings, which showed an 502 increase in sugar content as a survival response to abiotic stresses (Sasaki et al. 1996, Zhu et al. 2006).

The preconditioning regimes used in this study caused several significant changes to the plant shoot tip cells that are likely to be related to the improvement in post-cryoexposure survival. Changes to membrane composition, and by extension, its stability, as demonstrated by electrolyte leakage and PL/sterol analysis, appear to be involved in the reduction of freeze, osmotic and toxic stresses (Menon et al. 2012). 508 Studies describing acclimation effects on cryostorage success have been published with temperate and sub-tropical species (Chang et al. 2000; Leunufna and Keller 2005; Kaczmarczyk et al. 2008; 509 510 Kushnarenko et al. 2009), however the number of studies on Mediterranean spp. especially Australian taxa is very limited by comparison. Indeed, given that five of the 34 internationally recognised 511 biodiversity hotspots are found in small but floristically diverse Mediterranean climate regions including 512 513 southwest Western Australia, South Africa's Cape floristic region, the European Mediterranean basin, the 514 Californian Floristic Province and the Chilean Matorral it is imperative the underlying physiology of plants from these environments be better understood as a means to aid and enhance current conservation 515 measures (Myers et al. 2000; Mittermeier et al. 2005). It might be argued that temperate or subtropical 516 517 species are 'related enough' to Mediterranean species in terms of climatic extremes (particularly temperature) to void claims of any discernable physiological differences, however in the context of south-518 519 west Australian taxa the type of hot-summer Mediterranean climate is not directly comparable to classical 520 temperate or sub-tropical climes. In particular, summer temperatures can be extreme and drought frequent, unlike most 'temperate' climates; with short cool wet winters this is again quite different from 521 sub-tropical climates with warm dry 'winters' and hot wet summers (McKnight and Hess 2000). 522 Therefore it is logical to expect that the adaptive physiology of south-west Australian Mediterranean 523 plants would also express a different range (and possibly intensity) of responses to acclimation treatments 524 compared to temperate or sub-tropical taxa. From this perspective while the current study reflects some 525 subtle differences in adaptive physiology for acclimated L. sonderi plants, it also illustrates several key 526 527 similarities to other studies on cold acclimation in species from more temperate northerly regions, 528 suggesting that basic acclimation responses are conserved but adaptations are likely to have evolved in response to unique local environmental stresses. 529

The information gained on the effects of preconditioning with respect to changes in the membrane PL and cell sugar contents provide an excellent framework for further research on membrane stability and its relation to improved stress tolerance. Overall this study provides valuable insight into the underlying processes governing effects of low temperature preconditioning on post-cryogenic survival of native Australian plant species and enhances our current cryogenic approaches for the long term storage of valuable and endangered plant germplasm, as well as other plant material currently maintained under tissue culture conditions used for post-mining restoration.

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538 Acknowledgements.

The authors declare that there are no conflicts of interest. This work was funded by ARC Linkage
Grant LP0884027 with further financial support from Alcoa of Australia Ltd and BHP Billiton Worsley
Alumina Pty Ltd. Dr. David Willyams of Alcoa of Australia's Marrinup Nursery is acknowledged for

providing the plant cultures used to conduct all experiments. The support of Matthew Timmins(Metabolomics Australia) for the analytical work is appreciated.

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- 706 Supporting Information
- 707
- **Table S1.** Phospholipids (ng mg⁻¹ FW) detected in *Lomandra sonderi* shoot tips after temperature
 preconditioning at 23 °C and 20/-1 °C for 3 weeks.

- **Table S2.** Identified phospholipid species (ng mg⁻¹ fresh weight \pm SD) found in *Lomandra sonderi* shoot
- tips after preconditioning at 23 °C or 20/-1 °C for 3 weeks.
- 712 **Table S3.** Sterols (ng mg⁻¹ FW) detected in *Lomandra sonderi* shoot tips after temperature
- 713 preconditioning at 23 °C and 20/-1 °C for 3 weeks.
- **Table S4.** Sugars (ng mg⁻¹ FW) detected in *Lomandra sonderi* shoot tips after temperature
- 715 preconditioning at 23 °C and 20/-1 °C for 3 weeks.
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717 Figure Legends

- **Fig. 1** Content of phospholipid (PL) classes (ng mg⁻¹ fresh weight) analysed in *Lomandra sonderi* shoot
- tips exposed to three-week preconditioning at 23 °C or 20/-1 °C. Bars represent average \pm SE of four
- replicates per treatment assessed. Bars labelled with different letters differ significantly when compared
- 721 across treatments (P < 0.05). PC phosphatidylcholine, PE phosphatidylethanolamine, PA -
- phosphatidic acid PG phosphatidylglycerol, PI phosphatidylinositol, PS phosphatidylserine
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- Fig. 2 Content of each sugar (ng mg⁻¹ fresh weight) analysed in *Lomandra sonderi* shoot tips exposed to
 three-week preconditioning at 23 °C and 20/-1 °C. Bars represent average ± SE of three replicates of
 treatments assessed. Bars labelled with different letters differ significantly when compared across
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